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A biomimetic magnetosome: formation of iron oxide within carboxylic acid terminated polymersomes†

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Bioinspired macromolecules can aid nucleation and crystallisation of minerals by mirroring processes observed in nature. Specifically, the iron oxide magnetite (Fe_3O_4) is produced in a dedicated liposome (called a magnetosome) within magnetic bacteria. This process is controlled by a suite of proteins embedded within the liposome membrane. In this study we look to synthetically mimic both the liposome and nucleation proteins embedded within it using preferential orientation polymer design. Amphiphilic block co-polymers self-assemble into vesicles (polymersomes) and have been used to successfully mimic liposomes. Carboxylic acid residue-rich motifs are common place in biomineralisation nucleating proteins and several magnetosome membrane specific (Mms) proteins (namely Mms6) have a specific carboxylic acid motifs that are found to bind both ferrous and ferric iron ions and nucleate the formation of magnetite. Here we use a combination of 2 diblock co-polymers: Both have the hydrophobic 2-hydroxypropyl methacrylate (PHPMA) block with either a poly(ethylene glycol) (PEG) block or a carboxylic acid terminated poly(2-methacryloyloxyethyl phosphorylcholine) (PMPC) block. These copolymers ((PEG₁₁₃-PHPMA₄₀₀) and (PMPC₂₈-PHPMA₄₀₀) respectively) self-assemble *in situ* to form polymersomes, with PEG₁₁₃-PHPMA₄₀₀ displaying favourably on the outer surface and PMPC₂₈-PHPMA₄₀₀ on the inner lumen, exposing numerous acidic iron binding carboxylates on the inner membrane. This is a polymersome mimic of a magnetosome (PMM₂₈) containing interior nucleation sites. The resulting PMM₂₈ were found to be 246 ± 137 nm in size. When the PMM₂₈ were subjected to electroporation (5 pulses at 750 V) in an iron solution, iron ions were transported into the PMM₂₈ polymersome core where magnetic iron-oxide was crystallised to fill the core; mimicking a magnetosome. Furthermore it has been shown that PMM₂₈ magnetopolymersomes (PMM₂₈Fe) exhibit a 6 °C temperature increase during *in vitro* magnetic hyperthermia yielding an intrinsic loss power (ILP) of $3.7 \text{ nHm}^2 \text{ kg}^{-1}$. Such values are comparable to commercially available nanoparticles, but, offer the added potential for further tuning and functionalisation with respect to drug delivery.

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Introduction

Biological iron oxide precipitation

The presence of iron oxide particles has been found in various species of birds and dolphins.^{1,2} While the true function and mechanism of action of these particles has never been properly explained, it is widely believed that in most cases they play an essential role in navigation by acting as “internal compasses”. The formation of such particles is the result of biomineralization. Biomineralization involves precipitation of an

inorganic material within or around living organisms.³ These minerals typically fulfil a specific function, such as forming a protective shell or a structural skeleton.² Biominerals often possess highly controlled size, shape and crystal structure compared to the equivalent synthetic materials.⁴ This level of structural control can be difficult to achieve in the laboratory because most biomineralization processes are governed by a complex suite of proteins, usually embedded within templating membranes.⁴

Precipitation of magnetite in magnetotactic bacteria

Magnetotactic bacteria are the smallest organisms that are known to perform biomineralization. Most strains precipitate pure magnetite nanoparticles (MNPs) within internally formed vesicular organelles, known as “magnetosomes”. Precise mag-

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Mimicking membrane/protein surface of the magnetosome. The inner lumen of the magnetosome membrane contains a range of biomineralisation proteins. Importantly, carboxylic acid rich nucleation proteins such as Mms6, which display a negatively charged surface for iron ion binding to nucleate magnetite formation. In this work we have advanced our pre-

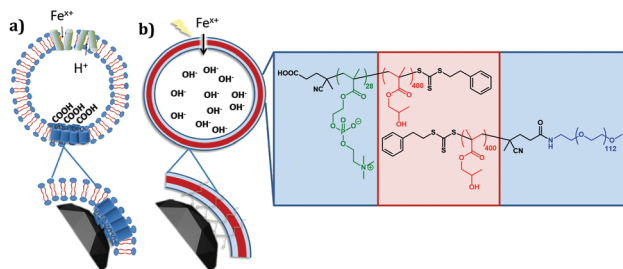


Fig. 1 Schematic of natural and synthetic vesicle membranes (blue = hydrophilic membrane leaflet, red = hydrophobic membrane leaflet) representing, (a) the bacterial magnetosome in which iron is recruited into the core of the magnetosome using iron transporters, which also acts as an antiporter pumping out internal protons. The efflux of protons causes a pH increase resulting in the precipitation of the magnetite crystal. The magnetite nucleation protein Mms6 (inset bottom left) which accumulates iron ions on the acidic amino acids on the inner lumen of the magnetosome. (b) Chemical composition of PEG-PHPMA and PMPC-PHPMA polymer which represent the hydrophilic (blue) and hydrophobic (red) leaflets of the vesicles, which features a carboxylic acid group on the PMPC monomer to mimic the iron ion binding/mineral nucleating sites of the Mms6 protein. The lighting represents electroporation to mimic the protein iron transporter, while the insert shows how the acidic end groups on PMPC mimic Mms6.

vious polymersomes by designing and creating a new di-block co-polymersome that will better mimic the magnetosome. Specifically, we have designed one copolymer to include a short hydrophilic block that displays carboxylate groups, while the other will instead include a longer neutral block. Due to the curvature of the polymersome we predict that steric factors will orientate the shorter acidic copolymer on the interior, displaying its acidic groups on the inner surface of the polymersome, while the bulkier copolymer will preferentially orientate on the exterior. The polymer system designed and presented in this work should now enable more preferential and controlled particle formation inside the polymersome (as opposed to within the membrane) by mimicking the action of nucleating proteins on the inner membrane (Fig. 1).

PEG-PHPMA/PMPC-PHPMA polymersomes

Poly(2-(methacryloyloxy)ethyl phosphorylcholine)-poly(2-hydroxypropyl methacrylate) (PMPC-PHPMA) is a vesicle-forming diblock copolymer that has a terminal carboxylic acid group situated on the hydrophilic PMPC block. In principle, addition of such groups should mimic the iron ion binding sites found in the Mms6 protein (mimicked by the COOH end group) within the magnetosomes liposome (lipid mimicked by the phosphorylcholine). It should be noted that carboxybetaine zwitterionic groups similar to phosphorylcholine have been shown to not interact with calcite mineralisation, to the extent they are not incorporated into calcite mineral whereas COOH groups alone do, so interaction with the negative charge on zwitterionic phosphorylcholine is improbable.³⁷ Moreover, preferential location of these carboxylic acid liposome-mimicking groups on the polymersome inner leaflet should create a syn-

thetic mimic of a nucleation protein within the magnetosome membrane (Fig. 1). In practice, this is achieved using a binary mixture of the PMPC-PHPMA chains with a second diblock copolymer, poly(ethylene glycol)-poly(2-hydroxypropyl methacrylate) (PEG-PHPMA). The two weakly hydrophobic PHPMA blocks have the same mean degree of polymerisation and co-assemble to form the vesicle membrane. However, the PMPC and PEG chains are thermodynamically incompatible.³⁸ Hence microphase separation occurs across the vesicle membrane, with the shorter PMPC block being preferentially expressed at the inner leaflet, while the longer PEG block is located mainly at the outer leaflet.

It is well known that amphiphilic diblock copolymers undergo spontaneous self-assembly in water.^{39–41} Traditionally diblock copolymers are prepared in a solvent with ideal solubility for both blocks before being transferred into water.⁴² Such copolymers can assemble into spheres, worms or vesicles depending on the relative volume fractions of each block.^{41,43} More recently several groups have utilised polymerisation-induced self-assembly (PISA) to prepare nano-objects *in situ*.^{44–46} More specifically, a soluble homopolymer is chain-extended with a monomer which when polymerised forms the insoluble core-forming block. For example, a poly(ethylene glycol) macromolecular chain transfer agent (PEG macro-CTA) has been prepared *via* end-group derivatization and subsequently chain-extended with HPMA *via* reversible addition-fragmentation chain transfer (RAFT) aqueous dispersion polymerization.⁴⁵ By varying the degree of polymerization (DP) of the core-forming PHPMA block and also the copolymer concentration, pure spheres, worms or polymersomes could be prepared in concentrated aqueous solution. Similarly, chain-extending a PMPC macro-CTA with HPMA enabled the synthesis of a similar series of PMPC-PHPMA nano-objects.⁴⁷ In both cases, PISA syntheses had to be conducted at relatively high copolymer concentration (20–25% w/w) in order to produce a pure polymersome phase. According to Gonzato and co-workers,⁴⁸ using a binary mixture comprising a short macro-CTA and a long macro-CTA should produce low-polydispersity polymersomes, whereby the short macro-CTA chains are mainly located within the inner lumen, while the long macro-CTA chains are preferentially expressed at the outer surface of the polymersomes.⁴⁵ In the present work, we have explored the use of a relatively long PEG macro-CTA in conjunction with a relatively short PMPC macro-CTA to polymerize HPMA *via* RAFT aqueous dispersion polymerization (Fig. 2). Microphase separation of these two steric stabilizer blocks across the membrane was anticipated, not least because Blanazs and co-workers have shown that PEG and PMPC homopolymers are enthalpically incompatible.³⁸ Moreover, judicious selection of a carboxylic acid-functionalized RAFT agent for the synthesis of the PMPC macro-CTA enables the construction of polymersomes in which the acid end-groups on this stabilizer block are mainly located within the lumen.⁴⁹ Thus such polymersomes mimic the structure of naturally-occurring bacterial magnetosomes containing the Mms6 protein. Following the work of Chambon *et al.*,⁵⁰ such poly-



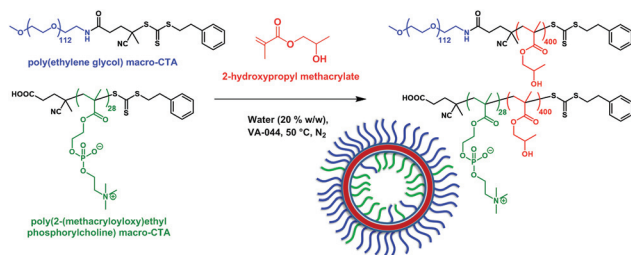


Fig. 2 Poly(ethylene glycol)-poly(2-hydroxypropyl methacrylate)/poly(2-(methacryloyloxy)ethyl phosphorylcholine)-poly(2-hydroxypropyl methacrylate) (PEG + PMPC)-PHPMA polymersomes are prepared by chain extension of a binary mixture of PEG₁₁₃ and PMPC₂₈ macro-CTAs via reversible addition–fragmentation chain transfer (RAFT) aqueous dispersion polymerisation of 2-hydroxypropyl methacrylate (HPMA).⁴⁵

mersomes can be covalently stabilized by addition of a bi-functional cross-linker at high HPMA conversions (>99%). PEG-PHPMA/PMPC-PHPMA vesicles assemble as part of the polymer synthesis process. Therefore, it is not possible to encapsulate base during rehydration. Instead, we report the novel methodology of soaking pre-formed polymersomes in NaOH.

Using this di-block system we have designed and are able to produce synthetic polymersomes that mimic magnetosomes (PMM₂₈).

Materials and methods

Materials

All solutions used were degassed and sparged with N₂ prior to use and all experiments were carried out under an inert atmosphere to minimise iron oxidation. 2-(Methacryloyloxy)ethyl phosphorylcholine (MPC; >99%) was donated by Biocompatibles Ltd (Farnham, UK). 2-Hydroxypropyl methacrylate (HPMA) was purchased from Alfa Aesar, 2,2'-azobis[2-(2-imidazolin-2-yl)propane] dihydro-chloride (VA-044) was purchased from Wako Pure Chemical Industries (Japan), ethylene glycol dimethacrylate (EGDMA) was purchased from Sigma Aldrich. All the above were used as received. PEG₁₁₃ macro-CTA was prepared as reported elsewhere.⁴⁵ All other chemicals were purchased from Sigma Aldrich, unless otherwise stated.

Polymersome synthesis

PMPC macro-CTA was prepared as reported elsewhere.⁴⁷ A typical protocol for the synthesis of PMM₂₈ polymersomes (self-assemblies of 70% PEG₁₁₃ PHPMA₄₀₀ and 30% PMPC₂₈PHPMA₄₀₀) is as follows: PEG₁₁₃ macro-CTA (0.0735 g, 1.4×10^{-5} mol), PMPC₂₈ macro-CTA (0.0516 g, 6.0×10^{-6} mol); vA-044 initiator (0.00187 g, 6.7×10^{-6} mol), ([combined CTA]/[VA-044] molar ratio = 3.0) and HPMA monomer (1.12 g, 8.0×10^{-3} mol, target DP = 400) were dissolved in deoxygenated water to generate a 20% w/w solution. The pH of the solution was adjusted to pH 6.8 using 0.1 M NaOH. This solution was then purged with nitrogen for 30 minutes. The flask

was sealed using a rubber septum and immersed in an oil bath at 50 °C. When the polymerisation had proceeded for 4 hours, the reaction was quenched by exposure to air and cooling to 20 °C. Cross-linked polymersomes, were synthesised as described above, but, after 4 hours degassed EGDMA (0.158 g, 8×10^{-4} mol) was added under N₂, and left to stir for 12 hours at 50 °C.

¹H NMR

¹H NMR was used to calculate conversion in the synthesis of 70% PEG₁₁₃-PHPMA₄₀₀, 30% PMPC₂₈-PHPMA₄₀₀ to form PMM₂₈. Spectra were acquired using a Bruker Avance III HD 400 spectrometer. Samples were prepared by dissolving 100 µL polymersomes solution in CD₃OD (3 mL).

Electroporation of polymersomes

PMM₂₈ polymersomes were incubated in 10 mM NaOH, then cleaned up using a size exclusion chromatography (SEC) column. A 1 : 2 Fe(II) : Fe(III) stock iron solution was prepared by mixing 1 ml of 10 mM FeCl₂·4H₂O solution (0.019 g, 1.00×10^{-4} mol, 10 mL, dissolved in degassed water) with 2 ml of 10 mM FeCl₃·6H₂O solution (0.027 g, 1.00×10^{-4} mol, 10 mL, dissolved in degassed water). The PMM₂₈ polymersomes with now basic cores were electroporated using a BioRad Micropulser™. 100 µL of basic PMM₂₈ were incubated with 100 µL of mixed valence iron solution (in a 1 : 1 v/v ratio), before being electroporated at a maximum volume of 800 µL. 5 pulses were applied at a voltage of 750 V, to facilitate transport of the iron ion solution as previously published.³⁰ SEC Cleaned electroporated PMM₂₈ were visualised unstained using TEM (Fig. 4).

Dynamic light scattering (DLS)

5 µL of the vesicle samples were diluted in 1 ml of MilliQ water before analysis in a disposable DLS cuvette using a scattering angle of 173° on a Zetasizer Nano (Malvern Instruments). Samples were scanned three times at a standard temperature of 25 °C, with each scan having 10–14 runs; data were analysed using Malvern Zetasizer software.

Transmission electron microscopy (TEM)

Transmission electron microscopy (TEM) was used to visually assess the polymersomes size, shape and integrity pre and post electroporation. The size and location of the iron oxide precipitation associated with the vesicle was assessed post-electroporation and selected area electron diffraction was used to assess crystallinity of the iron oxide. 5 µL of the polymer-some sample (diluted 1 : 1000 v/v) was applied to a carbon film grid and left to stand for approximately 1 minute, before blotting off excess liquid. Grids were then dried at ambient temperature. All post-electroporation samples were imaged unstained. Control samples assumed not to contain dense iron minerals were subsequently stained (0.75% uranyl formate) prior to imaging, before blotting off after a 12 seconds staining period. Samples were imaged using a Tecnai F20 and images processed using ImageJ.





Fig. 4 Unstained TEM images of (a) PMM_{28}Fe . The PMM_{28} sample was soaked in NaOH (10 mM) before electroporation at 750 V (5 pulses) in the presence of excess $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$. The polymersomes show dark electron dense cores, attributed to the precipitation of magnetic iron oxide.

is not seen in the PMM_{28} prior to electroporation (compare Fig. 4 with ESI 3†).

The electroporated PMM_{28}Fe are visibly more spherical uniformly across the whole population and are also smaller with a narrower size distribution (212 ± 79 nm). The filled cores appear to be made up of aggregates of smaller MNPs of iron oxide, with some discreet particles visible nearer the membrane as the density of iron oxide reduces radially from the centre (Fig. 5a). The size of these isolated particles was determined to 5.54 ± 0.55 nm (ESI 4†). Furthermore, the iron oxide-loaded polymersomes were found to be magnetic (attracted to a permanent magnet) and selected area electron diffraction confirms that the precipitation visible in the polymersome core was crystalline, indicating the iron oxide core is crystalline magnetite or maghemite (Fig. 5b). ICP elemental analysis gave

$27.5 \mu\text{g ml}^{-1}$ of Fe to 0.5 mg ml^{-1} of polymersomes giving approximately 5.5% iron content by weight in PMM_{28}Fe .

Comparison to other magnetopolymersomes

This is the first example of the precipitation of magnetic iron-oxide densely concentrated into the core of a polymersome (Fig. 5a). Our previous work utilising electroporation to produce magnetopolymersomes was conducted on poly(butadiene)-poly(ethylene oxide) (PBD-PEO) diblock co-polymersomes. PBD-PEO polymersomes possess no functional nucleation sites and as a result the MNPs simply form in the membrane where the two reagents (iron ions and NaOH) meet in the pores opened by the electroporation process (Fig. 5c).²² Similarly we have demonstrated electroporation can be used to produce magnetopolymersomes from Tri-block co-polymers by precipitate MNPs within an ABA Tri-block co-polymersome composed of PMOXA-PDMS-PMOXA (where PMOXA = poly(2-methyloxazoline) & PDMS = poly(dimethylsiloxane)) (Fig. 5d).³⁰ Again, there are no functional nucleation sites on this polymer and while we believe there may be more internal MNPs in this example (possibly due to the different structural configuration of pore formation for a tri-block under electroporation conditions), the size of particles formed in both these two previously reported examples is ~ 2.6 nm. This contrasts with this current study which shows a two-fold increase in the particle size (5.5 nm) and the overall quantity and quality of precipitation within the PMM_{28}Fe core. ICP elemental analysis shows an approximate 5-fold minimal increase in quantity of elemental iron compared with both the tri-block and di-block.^{29,30} This strongly indicates that the COOH groups promote more internal nucleation of larger MNPs.

Effect of crosslinking PMM_{28}

There is a clear electron dense “halo” on the outer leaflet of the polymersomes. The chromatography clean-up removes all un-associated iron oxide precipitate, showing the iron oxide halo to be strongly associated with the vesicles surface (Fig. 4 & 5a). This could be evidence for the presence of deprotonated carboxylic acid groups on the outer surface of the vesicle, nucleating and forming iron-oxides on the exterior. Ideally the acidic shorter polymer chains in the PMPC block would exclusively orientate to the inner lumen of the polymersome, however, a small amount of carboxylic acid groups may mis-orientate at the formation stage or flip to the exterior once formed. Crosslinking could reduce polymer flipping post-formation.

Ethylene glycol dimethylacrylate (EGDMA) cross-linker was added to the PMM_{28} during synthesis in an attempt to retain all carboxylic acid groups on the inner membrane leaflet,⁵⁰ to “lock” the chains in place. EGDMA-crosslinked PMM_{28} were electroporated at 750 V in the presence of mixed valence iron solution as described above. TEM images without negative staining showed magnetopolymersomes similar to those shown in Fig. 4 and 5a but of a smaller average diameter (80 nm) (ESI 5†), but importantly, the outer halo was still present showing little difference between polymersomes pro-

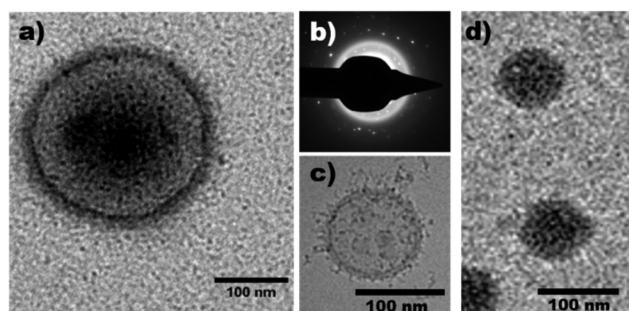


Fig. 5 (a) Unstained TEM image of electroporated PMM_{28}Fe polymersomes showing precipitation on the outer leaflet of the polymersome as well as in the vesicle core. (b) A selected area electron diffraction image of the same electroporated polymersome confirming the presence of a crystalline substance within the polymersome. (c) Cryo-EM micrograph of PEO-PBD polymersomes from the first reported use of the electroporation method²⁹ and (d) show comparison of use of the electroporation method on PMOXA-DMS-MOX tri-block polymersomes.³⁰





Fig. 6 Temperature increase vs. time curve obtained during magnetic hyperthermic heating of a 0.2% w/v aqueous dispersion of PMM₂₈Fe polymersomes (0.11 mg ml⁻¹ of iron). A temperature increase of 6 °C is observed over a period of 10 minutes (600 s).

duced with and without the EGDMA crosslinker. If anything there is less MNP precipitated in the core of the EGDMA cross-linked PMM₂₈. We can thus conclude flipping of the PMPC₂₈-PHPMA₄₀₀ block onto the outer surface post formation is not the dominant reason for surface iron oxide precipitation. Thus it is clear that either some of the carboxylic acid groups mis-orientation to the exterior during polymersome formation, or the precipitation on the surface is simply due to some NaOH leaking out during electroporation and is bound at the external membrane pore site.

Hyperthermic heating effect of PMM₂₈Fe magnetopolymersomes

The biomedical hyperthermia treatment potential of PMM₂₈Fe magnetopolymersomes was investigated. PMM₂₈Fe were exposed to an alternating magnetic field to determine their potential for magnetic hyperthermia by assessing their heating power (over a 10 minutes period). A control of the blank PMM₂₈ polymersome in PBS (ESI 6†) was subtracted from the final heating power of the magnetopolymersomes (Fig. 6). A temperature increase of 6 °C was observed over a 10 minutes period. A SAR value of 122 W g⁻¹ and an ILP of 3.7 nHm² kg⁻¹ was thus calculated. In principle, such localized heating can sensitize surrounding tissue towards a therapeutic payload that could be delivered within the same polymersomes.

Conclusions

In this work we have successfully designed and synthesized an asymmetric polymersomes with a selectively orientated smaller acidic block copolymer (PMPC₂₈-PHPMA₄₀₀) in the interior and a larger PEG₁₁₃-PHPMA₄₀₀ block copolymer forming the exterior of the vesicle to create a polymersome mimic of a magnetosome (PMM₂₈). Similar to our previous work producing magnetopolymersomes, we used electroporation to allow iron ion transport across the membrane and into the core of the vesicle to precipitate iron oxide. Unlike our pre-

vious work, here we have been successful in precipitating dense, magnetic, crystalline iron oxide within the core of the vesicle (closer to resembling a magnetosome) due to iron ion binding and iron-oxide nucleation capacity of the COO⁻ functional groups exposed on the inner leaflet in the core.

The iron oxide core of PMM₂₈Fe is not a single crystal but an aggregation of 5.5 nm sized MNPs of high density in the centre, with discreet MNP being visible near the membrane.

The addition of the acidic nucleation sites have not only targeted the MNP precipitation into the core of PMM₂₈ but has also lead to larger MNP within the magnetopolymersome. A small amount of exterior surface iron oxide precipitation is seen which was not reduced by cross-linking the PMM₂₈ during formation.

Finally, PMM₂₈Fe magnetopolymersomes were found to have advantageous properties with respect to potential future magnetic hyperthermia treatments, achieving a 6 °C heat increase in 10 minutes. In principle, such magnetite-loaded polymersomes can sensitize tissue for the *in vivo* delivery of a therapeutic payload or serve as a contrast agent for MRI diagnostic applications.²⁷ Furthermore, the use of polymersomes enable greater ability to design in multiple responsive functionalities for novel bioinspired smart nanomedical materials.

Conflicts of interest

There are no conflicts of interest to declare.

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