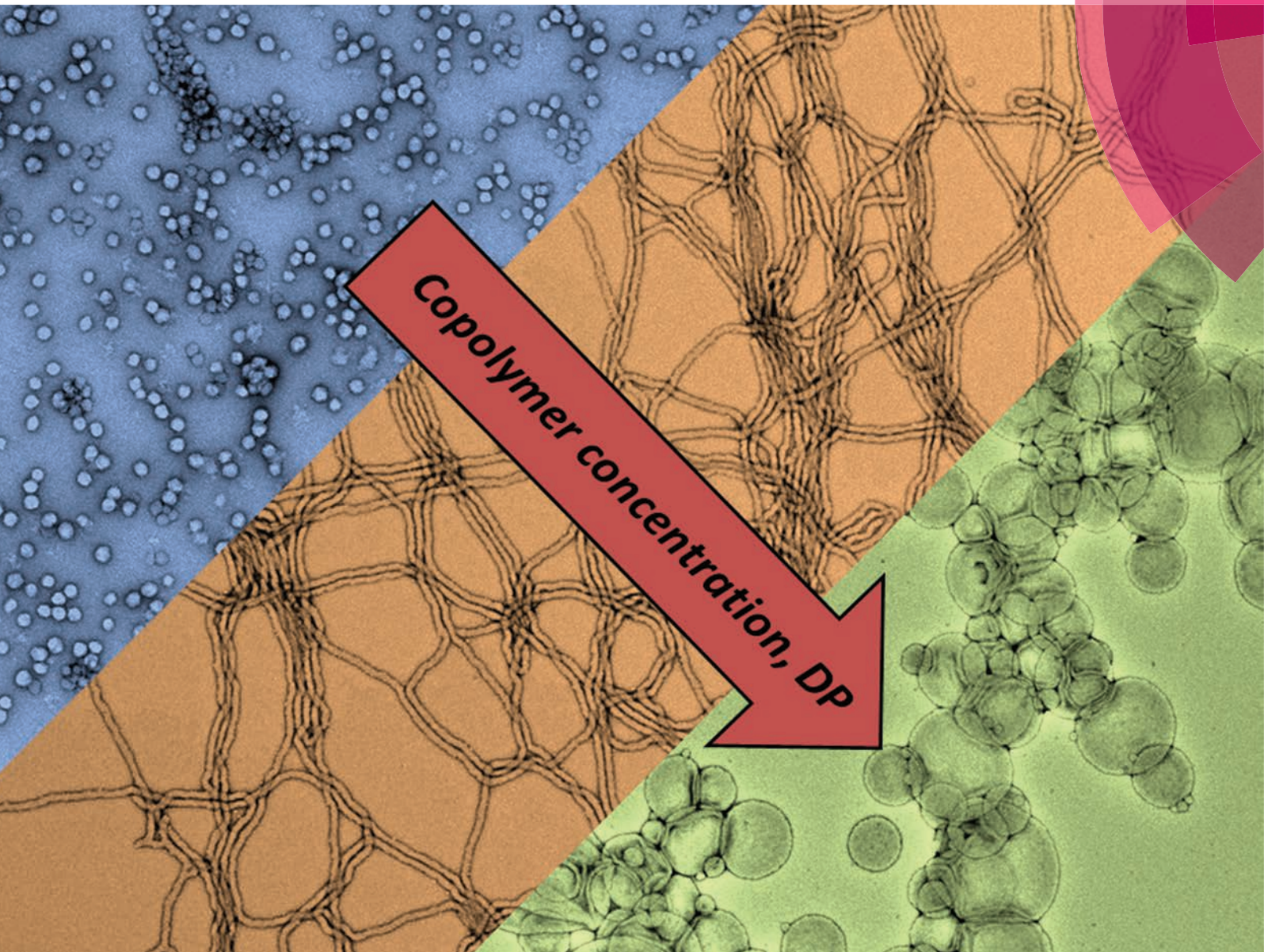


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Polysulfobetaine-based diblock copolymer nano-objects *via* polymerization-induced self-assembly†

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A zwitterionic polysulfobetaine-based macromolecular chain transfer agent (PSBMA₃₈) was prepared by reversible addition–fragmentation chain transfer (RAFT) solution polymerization of [2-(methacryloyloxy)ethyl] dimethyl(3-sulfopropyl) ammonium hydroxide (SBMA) in an aqueous solution containing 0.5 M NaCl at 70 °C. This PSBMA₃₈ macro-CTA was then utilized for the RAFT aqueous dispersion polymerization of a water-miscible monomer, 2-hydroxypropyl methacrylate (HPMA). The growing PHPMA block became hydrophobic *in situ*, leading to polymerization-induced self-assembly. Systematic variation of the mean degree of polymerization of the PHPMA block and the copolymer concentration enabled access to pure phases of spheres, worms or vesicles, as judged by transmission electron microscopy and dynamic light scattering studies. A detailed phase diagram was constructed and the thermo-responsive behavior of selected PSBMA₃₈-PHPMA_x nanoparticles was investigated. Finally, the salt tolerance of PSBMA₃₈-PHPMA₄₀₀ vesicles was compared to that of PGMA₇₁-PHPMA₄₀₀ vesicles; the former vesicles exhibit much better colloidal stability in the presence of 1 M MgSO₄.

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Introduction

It is well known that amphiphilic AB diblock copolymers spontaneously undergo self-assembly in aqueous solution in order to minimize the unfavorable interactions between the hydrophobic blocks and the solvent.¹ The resulting copolymer morphology depends on the so-called packing parameter, *p*, which is related to the relative volume fractions of the hydrophilic and hydrophobic blocks.² A wide range of copolymer morphologies have been reported, including spherical micelles,³ worm-like particles⁴ and vesicles.⁵

Traditionally, diblock copolymer self-assembly in solution has been achieved *via* post-polymerization processing techniques such as thin film rehydration^{6,7} or a solvent switch,⁸ where the copolymer chains are initially dissolved in a good solvent for both blocks and then a selective solvent for one of the blocks is added in order to induce self-assembly. This approach usually involves additional purification steps to remove the non-selective solvent, *e.g.* by dialysis or evaporation. This self-assembly route also suffers from a major dis-

advantage: it is almost invariably conducted in dilute solution (typically <1% w/w copolymer).

Recently, we^{9–13} and others^{14–18} have reported that polymerization-induced self-assembly (PISA) of various amphiphilic diblock copolymers can be readily achieved using either aqueous dispersion polymerization^{10,19} or aqueous emulsion polymerization^{20–22} based on reversible addition–fragmentation chain transfer (RAFT) chemistry.^{23–26} This approach allows various copolymer morphologies to be readily prepared *in situ* at relatively high copolymer concentrations and requires no post-polymerization processing. One early formulation based on RAFT aqueous dispersion polymerization utilized a zwitterionic phosphobetaine homopolymer, poly(2-(methacryloyloxy)ethyl phosphorylcholine) (PMPC), as the stabilizer block and poly(2-hydroxypropyl methacrylate) (PHPMA) as the core-forming block to access a range of copolymer morphologies.¹⁰ However, the relatively high mass of the MPC repeat units meant that highly anisotropic copolymer compositions had to be targeted at relatively high copolymer solids (*ca.* 16–25%) in order to access non-spherical copolymer morphologies (*e.g.* worms or vesicles).

Polysulfobetaines are closely related to polyphosphobetaines and both classes of polyzwitterions have been shown to be salt-responsive.^{27–35} The presence of salt generally leads to higher water solubility, which is sometimes known as the ‘anti-polyelectrolyte’ effect.³⁶ Certain polysulfobetaines also display thermo-responsive behavior, with their upper critical solution temperature (UCST) depending on both the copolymer molecular weight and copolymer concentration.^{29,37–39}

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† Electronic supplementary information (ESI) available: Additional kinetic data;

¹H NMR spectra for CTA, macro-CTA and diblock copolymer; TEM images of S₃₈-H₄₀₀ vesicles dispersed in pure water and in an aqueous solution of 1 M MgSO₄; digital photographs of S₃₈-H₄₀₀ and G₇₁-H₄₀₀ vesicles in aqueous solutions containing MgSO₄. See DOI: 10.1039/c5py00396b



Like polyphosphobetaines,^{40–42} polysulfobetaines have been shown to be highly biocompatible and exhibit anti-fouling properties.^{41,43–46} Importantly, polysulfobetaines are significantly cheaper than polyphosphobetaines.

In 2014 Pei and Lowe reported the synthesis of a range of diblock copolymer nano-objects comprising a polysulfobetaine stabilizer block and 2-phenylethyl methacrylate (PEMA) as the hydrophobic block. However, these copolymers were prepared *via* post-polymerization modification. First, a poly(2-(dimethylamino)ethyl methacrylate) (PDMA) macro-CTA was employed for the ethanolic RAFT dispersion polymerization of PEMA.¹⁶ These precursor diblock copolymers underwent self-assembly during their PISA synthesis to yield a range of copolymer morphologies, including spheres, worms and vesicles. The PDMA stabilizer block was subsequently quaternized using 1,3-propanesultone to yield polysulfobetaine-based nanoparticles. However, 1,3-propanesultone is known to be carcinogenic and the PISA synthesis was conducted in ethanol rather than water (with the purified copolymer nanoparticles being redispersed in water).

Herein we employ poly(2-(methacryloyloxy)ethyl dimethyl(3-sulfopropyl)ammonium hydroxide) (PSBMA) as a polysulfobetaine macro-CTA; this is used as a steric stabilizer to conduct the RAFT aqueous dispersion polymerization of 2-hydroxypropyl methacrylate (HPMA) in order to generate a thermo-responsive core-forming block. This formulation yields a range of PSBMA-HPMA diblock copolymer nano-objects directly *via* PISA (see Fig. 1). Our approach is both versatile and efficient: it avoids post-polymerization modification with carcinogens such as 1,3-propanesultone and affords the desired nano-objects directly in water (hence subsequent transfer from ethanol to water is not required). Moreover, thermo-responsive behaviour is conferred on the resulting diblock copolymer nano-objects.

Experimental

Materials

4-Cyanopentanoic acid dithiobenzoate (CADB), [2-(methacryloyloxy)ethyl] dimethyl(3-sulfopropyl) ammonium hydroxide (SBMA) and deuterium oxide (D₂O) were obtained from Sigma-Aldrich U.K. and were used as received. 4,4'-Azobis(4-cyanovaleic acid) (ACVA, 99%) and 2-hydroxypropyl methacrylate (HPMA, 98%) were obtained from Alfa Aesar (UK) and were used as received. Deuterated methanol (CD₃OD) was obtained from Cambridge Isotope laboratories. Dialysis tubing was received from SpectraPor. Deionized water was used in all experiments.

Methods

Synthesis of PSBMA₃₈ macro-CTA. CADB RAFT agent (0.75 g, 2.7 mmol), SBMA monomer (30.0 g, 108 mmol) and ACVA (151 mg, 0.54 mmol, CTA/ACVA molar ratio = 5.0) were weighed into a 500 ml round-bottomed flask containing a stir bar. 0.5 M NaCl solution (150 g) was added and the pH slowly adjusted to *ca.* pH 7 using dilute aqueous NaOH. The solution was purged with nitrogen for 45 minutes and sealed with a

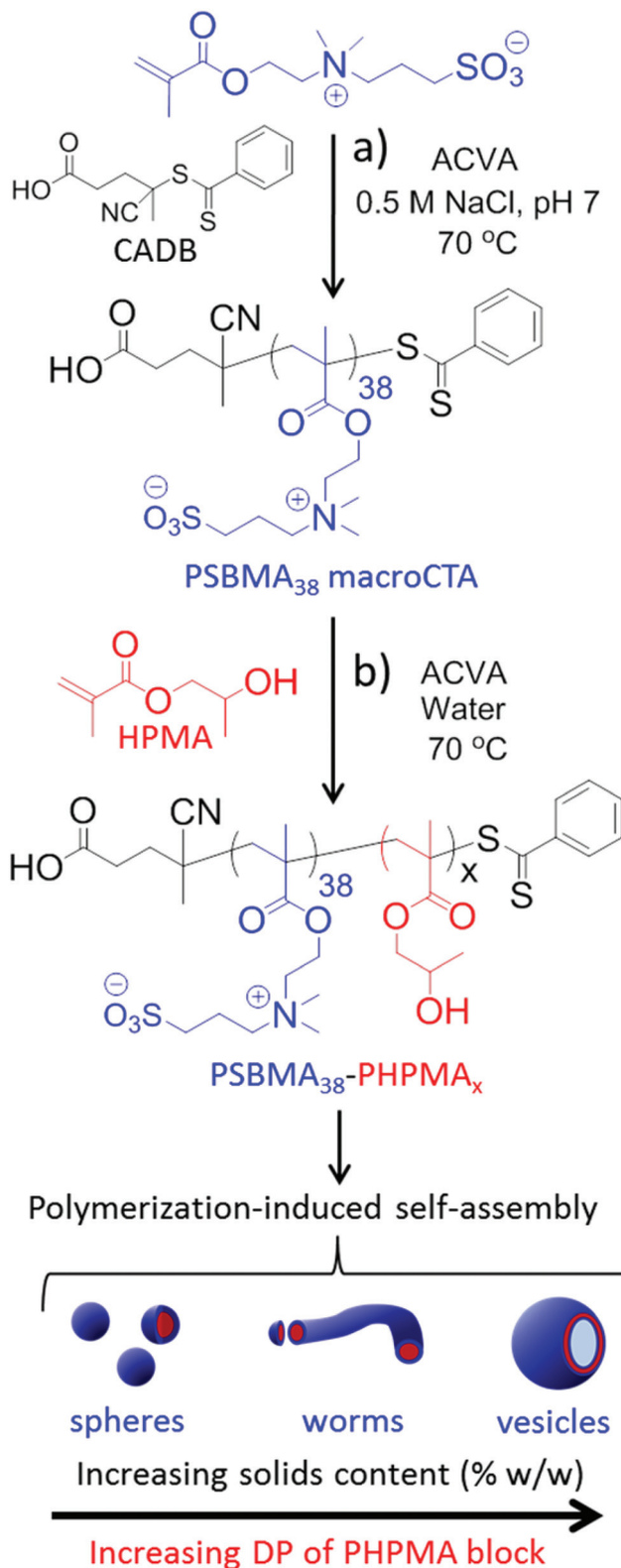


Fig. 1 (a) Synthesis of the PSBMA₃₈ macro-CTA *via* RAFT aqueous solution polymerization of SBMA. (b) RAFT aqueous dispersion polymerization of HPMA using this PSBMA₃₈ macro-CTA at 70 °C to afford various PSBMA-HPMA diblock copolymer nano-objects *via* polymerization-induced self-assembly (PISA).



rubber septum under a positive nitrogen pressure. The flask was then immersed in a pre-heated oil bath set at 70 °C. The polymerization was quenched after 2 h by rapid cooling in liquid nitrogen (final monomer conversion = 88%). The crude PSBMA was purified by exhaustive dialysis (SpectraPor membrane, MWCO = 3.5 kDa) against water, followed by lyophilization to yield **1** as a pink powder in 86% yield; M_n NMR = 11 kDa, M_n SEC = 6.9 kDa, M_w/M_n = 1.09.

Synthesis of PSBMA₃₈-PHPMA_x via RAFT aqueous dispersion polymerization. A typical protocol for the synthesis of PSBMA₄₇-PHPMA₂₀₀ is as follows: PSBMA₄₇ macro-CTA (0.44 g, 0.038 mmol), HPMA (0.82 g, 7.4 mmol), ACVA (2.0 mg, 0.008 mmol, CTA/ACVA molar ratio = 5.0) and water (4.5 g) were weighed into a round-bottomed flask containing a stir bar. The solution was purged with nitrogen for 30 minutes and sealed with a rubber septum under a positive nitrogen pressure, prior to immersion in a pre-heated oil bath at 70 °C for 6 h. The reaction was quenched by exposure to air and rapid cooling at 20 °C. ¹H NMR spectroscopy analysis in CD₃OD indicated less than 1% of residual HPMA monomer. For kinetic studies, aliquots were periodically removed for analysis by ¹H NMR spectroscopy. For brevity, the notation S₃₈-H_x-Y is used where S stands for PSBMA, H for PHPMA, x is the DP of the PHPMA block and Y is the solids content (% w/w) used for the copolymer synthesis.

Polymer characterization

Molecular weight distributions were assessed by aqueous gel permeation chromatography (GPC) at 40 °C using a guard column, a 5 μm PL Aquagel-OH 30 column and a 5 μm PL Aquagel-OH 40 column connected in series to an Agilent Technologies 1260 Infinity refractive index detector, using a phosphate buffer eluent (0.08 M Na₂HPO₄; adjusted to pH 8.9 using NaOH) at a flow rate of 1.0 ml min⁻¹. The number-average molecular weight (M_n) and polydispersity (M_w/M_n) were calculated using a series of near-monodisperse poly(ethylene oxide) calibration standards.

¹H NMR spectra were acquired using a Bruker AV1-400 MHz spectrometer in either D₂O or CD₃OD. At least 64 scans were recorded for each sample. All chemical shifts (δ) are reported in ppm.

DLS measurements were performed using a Zetasizer Nano-ZS instrument (Malvern Instruments, UK) equipped with a 4 mW He-Ne laser operating at 633 nm. Light scattering was detected at 173°. Hydrodynamic diameters were determined using the Stokes-Einstein equation and averaged over three consecutive runs. Copolymer dispersions were diluted to 0.2% w/v prior to analysis at 25 °C. Viscosities (and refractive indices) for aqueous solutions of 1 M NaCl, 2 M NaCl and 1 M MgSO₄ at 20 °C were calculated to be 1.096 mPa s (1.343), 1.219 mPa s (1.372) and 1.884 mPa s (1.355) respectively.⁴⁷

Transmission electron microscopy studies were conducted using a Phillips CM 100 TEM instrument operating at 100 kV, equipped with a Gatan 1k CCD camera. Copper/palladium TEM grids (Agar Scientific) were surfaced-coated in-house to produce a thin film of amorphous carbon. The grids were then

plasma glow-discharged for 30 seconds to yield a hydrophilic surface. 10 μL of the aqueous dispersion (diluted to 0.1% w/w) was placed on the grid, then blotted after 1 minute to remove excess sample. The grids were then negatively stained using uranyl formate solution (10 μL, 0.75% w/w) for 20 seconds, followed by blotting to remove excess staining solution and dried using a vacuum hose.

Rheology measurements were performed using an AR-G2 rheometer equipped with a variable temperature Peltier plate, a 40 mL aluminium cone and a solvent trap. Storage (G') and loss (G'') moduli were measured as a function of percentage strain and temperature in order to identify the linear viscoelastic region and also to determine the critical degelation temperature. Percentage strain sweeps were conducted at constant temperature (37 °C) using an angular frequency of 1.0 rad s⁻¹. Angular frequency sweeps were carried out at a constant applied strain of 1.0%. Temperature sweeps were conducted at an angular frequency of 1.0 rad s⁻¹ and an applied strain of 1.0%. The temperature was increased by 2 °C, with 5 minutes being allowed for thermal equilibrium between each measurement.

Results and discussion

Synthesis of the PSBMA₃₈ macro-CTA

Synthesis of a PSBMA₃₈ homopolymer using 4,4'-azocyanovaleic acid (ACVA) initiator and 4-cyanopentanoic acid dithiobenzoate (CADB) as the RAFT agent was conducted in aqueous solution containing 0.50 M NaCl at 70 °C. The reaction solution pH was adjusted to 7.2 to facilitate dissolution of the RAFT agent and electrolyte was added because it is well known that this enhances the water solubility of PSBMA.^{27,30,31} The polymerization kinetics for SBMA exhibited a linear semi-logarithmic plot and reached high conversion (90%) within 3 h (see Fig. S1 in ESI†). The M_n increased monotonically with conversion and the polydispersities remained low throughout the polymerization ($M_w/M_n < 1.30$). The crude PSBMA was purified by dialysis against water and recovered by lyophilization to yield macro-CTA **1** as a pink powder (M_n = 5 kDa, M_w/M_n = 1.09 vs. poly(ethylene oxide) standards).

Analysis of the ¹H NMR spectrum of **1** recorded in D₂O indicated a mean DP of 38, as calculated by comparing the integrals of the aromatic RAFT end-group signals (**k**, **n** and **m**) between 7.4 ppm and 8.0 ppm with the polymer side-chain signals at 2.98, 3.18, 3.54, 3.78 and 4.46 ppm (assigned as **j**, **f**, **g**, **e** and **d**; see Fig. 2).

Polymerization-induced self-assembly

A series of PSBMA₃₈-PHPMA_x (hereafter referred to as S₃₈-H_x) diblock copolymers was prepared using this PSBMA₃₈ macro-CTA for the RAFT aqueous dispersion polymerization of PHPMA in deionized water at 70 °C. It was decided to avoid using an aqueous salt solution to enable direct comparison of the results obtained with this PISA formulation to those previously reported for the PMPC-PPHMA system.¹⁰ It is known



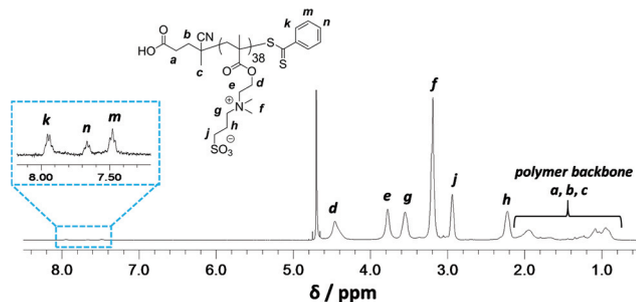


Fig. 2 Assigned ^1H NMR spectrum recorded for PSBMA macro-CTA **1** in D_2O .

that the UCST of PSBMA is reduced in the lower molecular weight limit.^{29,37,38} More specifically, Willcock *et al.* investigated the effect of chain length at a solution concentration of 1.0 g dm^{-3} : No UCST was observed for PSBMA homopolymers with molecular weights of 20 kDa ($\text{DP} = 70$) or below.³⁹ Therefore, given its relatively low molecular weight and anionic carboxylate end-group (which also suppresses UCST behavior),⁴⁸ the PSBMA₃₈ macro-CTA used in this work was not expected to exhibit a UCST in the absence of salt. Indeed, control experiments confirmed that a 15% w/w aqueous solution of PSBMA₃₈ remained water-soluble even at 4 °C and all aqueous dispersion polymerization syntheses were performed using an initial PSBMA₃₈ concentration of 12% w/w (or lower). The total solids content of the PISA formulation was varied from 10% w/w to 25% w/w. High HPMA conversions were obtained within 3 h at 70 °C. Three distinct regions can be observed in the semi-logarithmic kinetic plot (see Fig. 3).

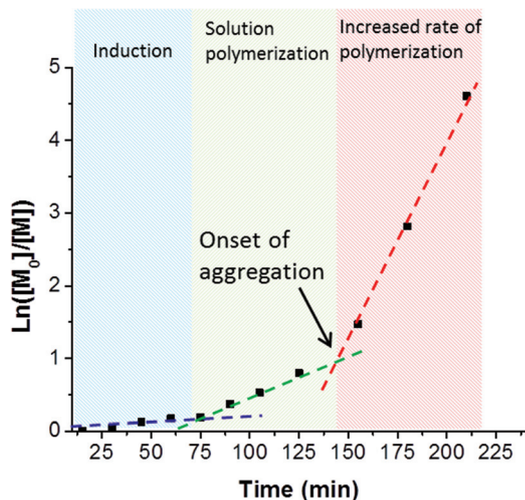


Fig. 3 First-order kinetic plot for the RAFT aqueous dispersion polymerization of HPMA conducted using a PSBMA₃₈ macro-CTA in deionized water at 70 °C and 20% w/w solids. Reaction conditions: $[\text{PSBMA}_{38}] : [\text{HPMA}] : [\text{ACVA}] = 200 : 1 : 0.20$. Three regimes can be identified: an induction period of approximately 75 min, a relatively slow rate of solution polymerization up to 145 min, and a relatively fast rate of dispersion polymerization after 145 min. The onset of micellization aggregation occurs at 150 min, which corresponds to a HPMA conversion of 67% and equates to a mean PHPMA DP of 133.

There is an induction period for the first 75 min, which is commonly observed for RAFT polymerizations of methacrylates conducted with dithiobenzoates,^{49,50} followed by a significant increase in rate for the next 70 min as solution polymerization occurs and water-soluble diblock copolymer chains are formed. The rate of polymerization further increases after 145 min, since this coincides with micellar nucleation.⁵¹ At this point, the HPMA monomer becomes partitioned within the micellar core, thereby increasing the local monomer concentration and hence accounting for the enhanced rate of polymerization.⁵¹ The point at which micellar nucleation occurs corresponds to a HPMA conversion of 67% for this particular PISA formulation, which equates to a mean DP of 133 for the growing PHPMA block. This value is significantly higher than the critical degree of polymerization of 92 reported by Blanazs *et al.* when polymerising HPMA using a poly(glycerol methacrylate) (PGMA₄₇) macro-CTA. This difference can be rationalized by the higher molecular weight of the PSBMA₃₈ block (10.9 kDa) compared to the PGMA₄₇ block (7.5 kDa). Thus the former block occupies a larger volume fraction than the latter and, although the PSBMA₃₈ stabilizer block is shorter, a longer core-forming PHPMA chain is needed to induce micellization.

Systematic variation of the mean DP of the core-forming PHPMA block and the total solids content (Y) enabled a detailed phase diagram to be constructed for the synthesis of $\text{S}_{38}\text{-H}_x\text{-Y}$ nanoparticles (see Fig. 4; full characterization data can be found in Table S1 in the ESI†). The final copolymer compositions were determined by ^1H NMR analysis (see Fig. S3 in ESI†). This phase diagram is rather similar to that reported by Sugihara *et al.* for a zwitterionic PMPC₂₅ stabilizer block¹⁰ and by Blanazs *et al.* for a PGMA₇₈ stabilizer block.¹¹ In each case, the final copolymer morphology is strongly concentration-dependent, with kinetically-trapped spheres being obtained at lower solids (since the probability of sphere-sphere fusion is reduced under these conditions) while equilibrium vesicular morphologies are formed at higher solids. As is usually the case,^{10–13} the pure worm phase occupies a relatively narrow region and is bounded by mixed phases. There is also a small region corresponding to around 14–16% w/w solids and a PHPMA DP of 250 that contains all three copolymer phases (*i.e.* spheres, worms and vesicles).

TEM studies indicate a strong correlation between the mean sphere diameter (D_{TEM}) and the target PHPMA block DP. Thus $D_{\text{TEM}} = 34 \pm 4 \text{ nm}$ for PHPMA₂₀₀, $D_{\text{TEM}} = 42 \pm 3 \text{ nm}$ for PHPMA₃₀₀ and $D_{\text{TEM}} = 54 \pm 6 \text{ nm}$ for PHPMA₄₀₀. TEM consistently undersizes these diblock copolymer nanoparticles relative to DLS. This is because the former technique examines dehydrated nanoparticles with collapsed stabilizer blocks, while the latter interrogates hydrated nanoparticles in aqueous solution. In addition, TEM reports a number-average diameter whereas DLS reports a z -average diameter, which is inherently biased towards larger nanoparticles. If $\text{S}_{38}\text{-H}_{400}$ nanoparticles are prepared at higher solids contents (12.5% w/w), then the DLS diameter increases up to 111 nm (polydispersity = 0.05). Along with the greater turbidity of the dispersion, this suggests



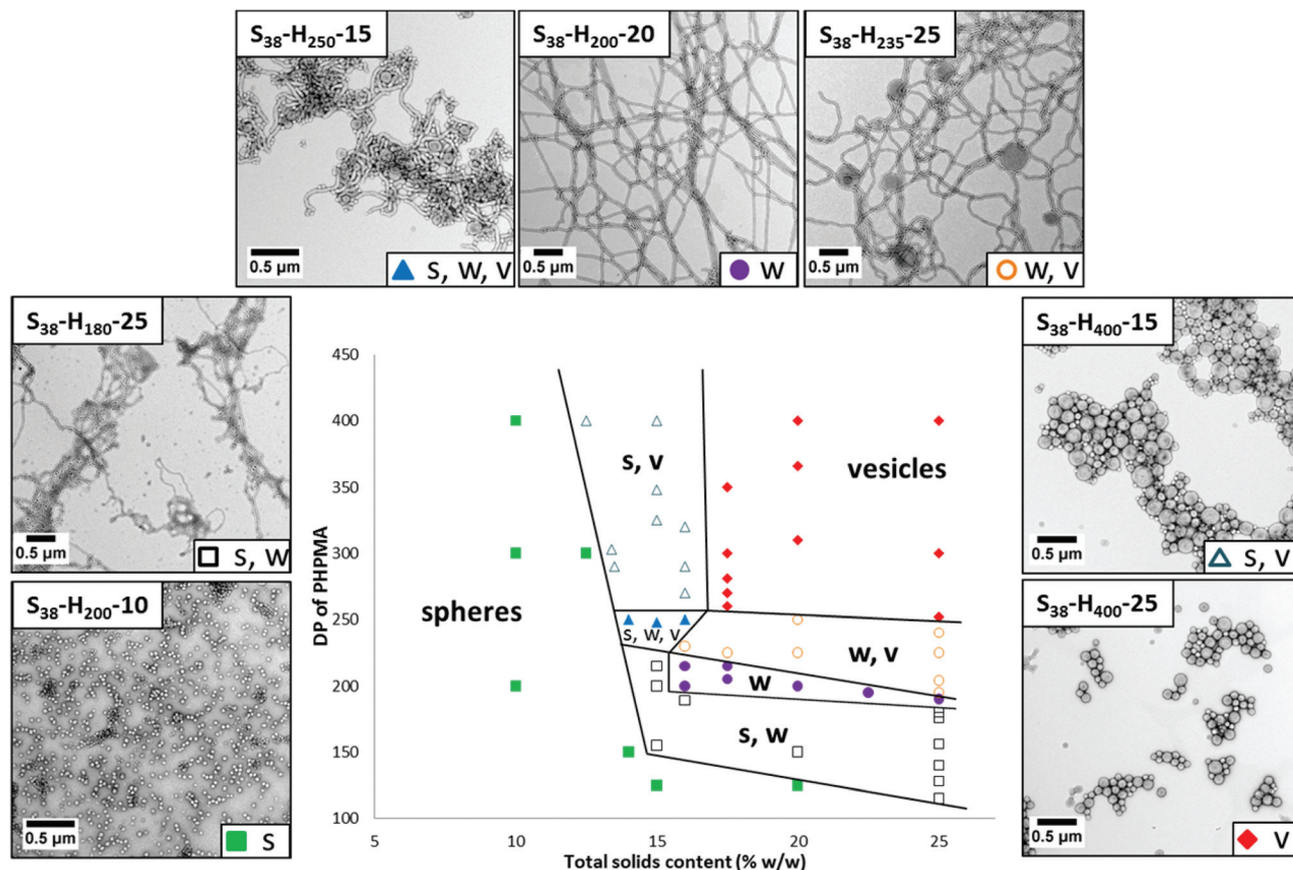


Fig. 4 Detailed phase diagram constructed for the preparation of $S_{38}\text{-H}_x\text{-Y}$ diblock copolymer nano-objects by RAFT aqueous dispersion polymerization of HPMA at 70 °C (for brevity, S denotes PSBMA and H denotes PHPMA). The target DP of the PHPMA block (x) and the total solids content (Y) were systematically varied and the diblock copolymer morphologies were assigned by *post mortem* TEM analysis of the final copolymer dispersions after appropriate dilution. Also shown are representative TEM images obtained for the various phases; S denotes spheres, W denotes worms and V denotes vesicles.

the presence of larger particles. TEM analysis indicates two distinct populations, corresponding to spheres ($D_{\text{TEM}} = 57 \pm 6$ nm) and relatively well-defined vesicle ($D_{\text{TEM}} = 97 \pm 7$ nm). The vesicles dimensions observed by TEM correlate quite well with those indicated by DLS. However, the latter technique is insensitive to spheres as the light scattering is dominated by the larger vesicles. Increasing the total solids concentration to 20% w/w for the synthesis of $S_{38}\text{-H}_{400}$ nanoparticles produces a pure vesicle phase ($D_{\text{TEM}} = 98 \pm 13$ nm), as indicated by TEM analysis, with DLS studies reporting a z -average diameter of 114 nm (polydispersity = 0.04). Thus the mean size of the pure vesicles is comparable to that of the vesicle population within the spheres plus vesicles mixed phase (Fig. 5 and 6).

Varying the target DP of the PHPMA block at a fixed total solids concentration of 20% w/w allows access to pure spheres, worms and vesicles, as well as various mixed phases comprising these two or more morphologies (see Fig. 7). For example, targeting $S_{38}\text{-H}_{125}$ leads to ill-defined spheres (TEM indicates $D_{\text{TEM}} = 27 \pm 5$ nm; DLS reports a z -average diameter of 34 nm (polydispersity = 0.14)). Increasing the target DP of the PHPMA block to 150 produces a mixed phase of spheres and

worms, as indicated by TEM analysis. Visual inspection of this dispersion confirms an increase in turbidity and formation of a free-standing gel. Targeting a PHPMA block DP of 200 results in the formation of a pure worm phase.

Both $S_{38}\text{-H}_{225}$ and $S_{38}\text{-H}_{250}$ formulations produced a mixed phase comprising worms and vesicles, with a higher proportion of vesicles being observed in the latter case as estimated by TEM analysis. The visual appearance and macroscopic behavior of these two dispersions were consistent with these TEM observations. $S_{38}\text{-H}_{225}$ formed an opaque free-standing gel, while $S_{38}\text{-H}_{250}$ comprised an opaque viscous solution. A pure vesicle phase was obtained when targeting a PHPMA block DP of 310 or higher. TEM studies indicate the formation of relatively polydisperse vesicles ranging from approximately 90 nm to 520 nm ($D_{\text{TEM}} = 214 \pm 122$ nm), which correspond quite well to the z -average diameter of 202 nm reported by DLS analysis.

Thermo-responsive behavior of $S_{38}\text{-H}_x\text{-Y}$ dispersions

There is considerable literature precedent for thermo-responsive PHPMA-based nano-objects in aqueous solution.^{12,13,52,53}



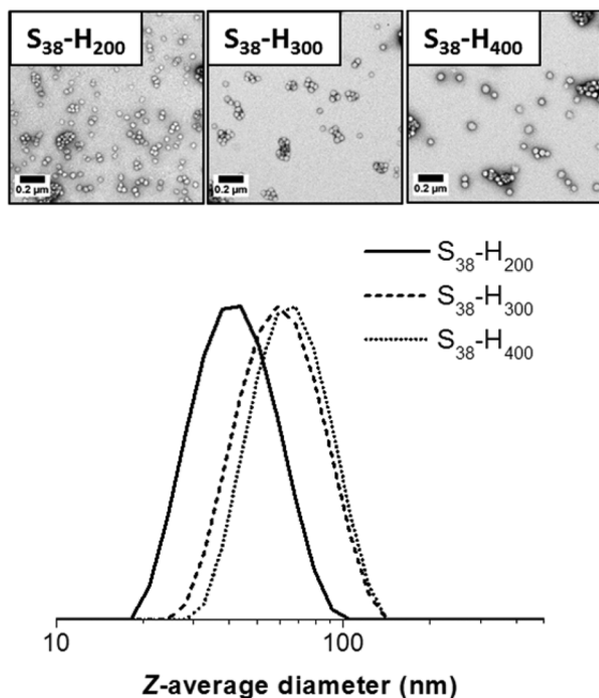


Fig. 5 TEM images and corresponding DLS particle size distributions obtained for spherical $S_{38}\text{-H}_x\text{-10}$ nanoparticles when $x = 200, 300$ or 400 .

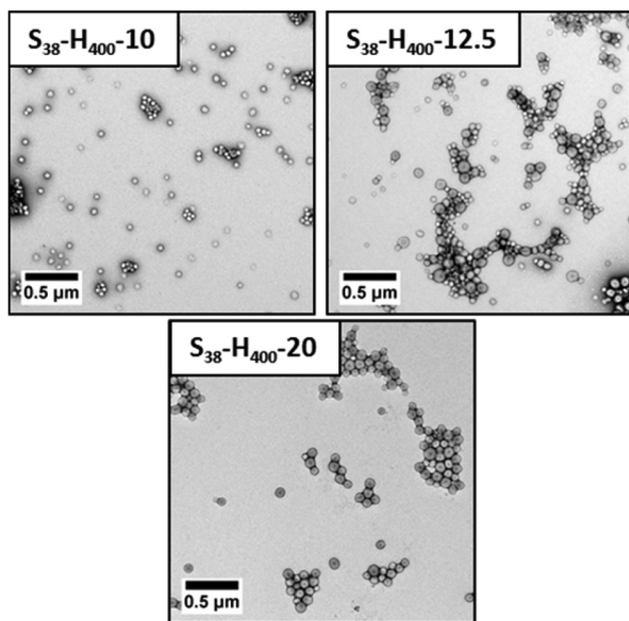


Fig. 6 Representative TEM images obtained for PSBMA₃₈-PPHMA₄₀₀ nanoparticles prepared at various solids contents. The copolymer morphology evolves from kinetically-trapped spheres (10% w/w) to equilibrium vesicles (20% w/w) via a spheres plus vesicles mixed phase (12.5% w/w).

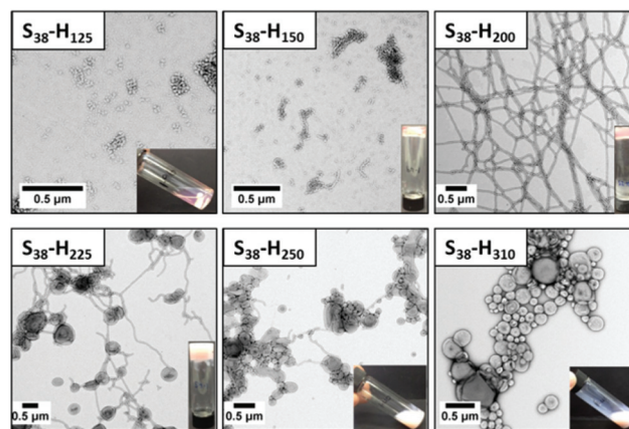


Fig. 7 Representative TEM images obtained for PSBMA₃₈-PPHMA_x (denoted as $S_{38}\text{-H}_x\text{-20}$ for brevity) nano-objects prepared by RAFT aqueous dispersion polymerization of HPMA at 70 °C at a total solids concentration of 20% w/w. A digital photograph of the physical appearance of each copolymer dispersion is shown as an inset in each TEM image. The scale bar in each case is 0.5 μm. The tube inversion test confirms the formation of free-standing gels in three cases, whereas the other three dispersions remained free-flowing.

Hence the thermo-responsive behavior of selected $S_{38}\text{-H}_x\text{-Y}$ dispersions were investigated using variable temperature oscillatory rheology. For example, a $S_{38}\text{-H}_{200}\text{-10}$ dispersion formed a rather soft worm gel ($G' \sim 15$ Pa) at 20 °C. A temperature sweep from 37 °C to 2 °C led to an increase in gel strength up to approximately 40 Pa, which is consistent with previous observations made by Warren *et al.*¹³ It is suggested that the initially short worms become longer upon cooling, hence a larger number of inter-worm contacts leads to a higher gel strength. G'' exceeded G' at around 2 °C, indicating degelation.

This is a result of greater solvation of the PPHMA core, which lowers the packing parameter¹ and results in a morphology transition from a worm gel to free-flowing spheres.^{52,53} However, this thermal transition proved to be irreversible, since G'' remained greater than G' throughout the subsequent heating cycle (see Fig. 8A). This thermo-responsive behavior is in contrast to that reported for PGMA₅₄-PPHMA₁₄₀ worm gels,⁵² but is similar to that observed by Warren *et al.* for a PEG₁₁₃-PPHMA₂₂₀ worm gel.¹³ During such thermal cycling, the temperature was lowered by 2 °C for each measurement, with 5 min being allowed between each data point to ensure thermal equilibrium. The time required to induce degelation at a given copolymer concentration was also examined. The same $S_{38}\text{-H}_{200}\text{-10}$ worm gel was held at 2 °C for 200 min (see Fig. 8B). For the first 60 min, the worm gel remained intact since G' exceeded G'' (and $G' \sim 65$ Pa). However, G'' became greater than G' after 60 min, producing a free-flowing liquid. Degelation was confirmed by visual inspection. TEM studies were undertaken, both for the original worm gel at 20 °C and also 2 h after this dispersion was held at 2 °C. There was a dramatic change in copolymer morphology from worms to spheres, which accounts for the observation of degelation.



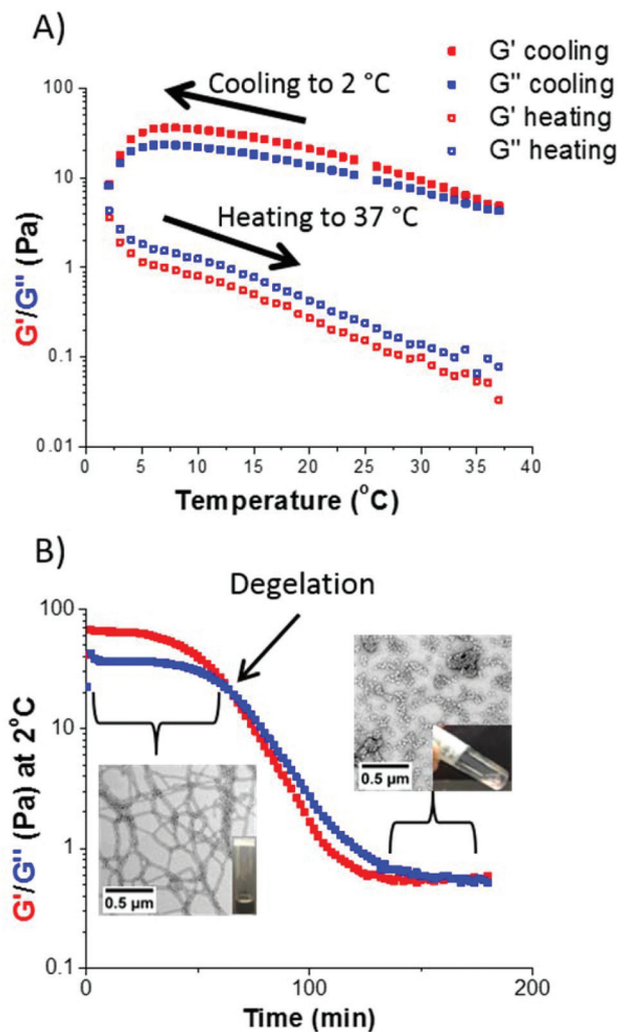


Fig. 8 (A) Storage (G') and loss (G'') moduli recorded for a 10% w/w $S_{38}\text{-H}_{200}$ worm gel upon cooling from 37 $^{\circ}\text{C}$ to 2 $^{\circ}\text{C}$ followed by heating up to 37 $^{\circ}\text{C}$. Irreversible degelation occurred at 2 $^{\circ}\text{C}$. (B) Storage (G') and loss (G'') moduli recorded for a 10% w/w $S_{38}\text{-H}_{200}$ worm gel held at 2 $^{\circ}\text{C}$ for 200 min. Degelation occurred after 60 min at this temperature to produce a free-flowing liquid. TEM images recorded before cooling and 2 h after being held at 2 $^{\circ}\text{C}$ confirmed a worm-to-sphere transition. Both sets of measurements were conducted at 1.0% applied strain using an angular frequency of 1.0 rad s^{-1} .

DLS studies of the diluted copolymer dispersions were also consistent with these TEM observations: the z -average diameter of the original worms at 20 $^{\circ}\text{C}$ was 420 nm (polydispersity = 0.29), whereas the z -average diameter was reduced to 30 nm (polydispersity = 0.25) after cooling to 2 $^{\circ}\text{C}$ for 2 h. The latter dimensions suggest the formation of polydisperse spheres.

A dispersion of $S_{38}\text{-H}_{310}\text{-25}$ vesicles was cooled to 4 $^{\circ}\text{C}$ for 48 h. The original dispersion was an opaque viscous liquid at 20 $^{\circ}\text{C}$. Visual inspection of the cold dispersion at 4 $^{\circ}\text{C}$ confirmed the formation of a turbid gel, as demonstrated by the tube inversion test. TEM studies indicated that the initial vesicles had transformed into a mixed phase comprising worms and vesicles (see Fig. 9).

Formation of the worm population accounts for the observation of macroscopic gelation. This dispersion did not undergo degelation on warming to room temperature, even on a time scale of several weeks. However, heating to 70 $^{\circ}\text{C}$ for 5 hours resulted in degelation. TEM studies indicated the presence of mainly vesicles, but some short worms are also observed. These short worms persist even after heating at 70 $^{\circ}\text{C}$ for 16 h.

Therefore this transition cannot be considered to be fully reversible. Again, this behavior is similar to that of the PEG₁₁₃-PHPMA₂₂₀ worms reported by Warren *et al.*¹³

The above examples suggest that $S_{38}\text{-H}_x$ nano-objects can be transformed into lower-order morphologies upon cooling. This indicates a reduction in the packing parameter under these conditions, which is consistent with a higher degree of plasticization of the PHPMA core-forming block. However, it appears that this thermo-responsive behavior is not fully reversible. This aspect is not yet understood and warrants further studies.

Behavior of $S_{38}\text{-H}_x$ nano-objects in the presence of added electrolyte

As previously discussed, polysulfobetaines such as PSBMA are known to exhibit enhanced aqueous solubility in the presence of salt compared to pure water.^{27,30,31} Therefore the tolerance of PSBMA-stabilized vesicles towards the addition of electrolyte was investigated. $S_{38}\text{-H}_{400}\text{-25}$ vesicles were diluted to 0.02% w/v using either water or various salt solutions and their colloidal stabilities were compared to that of PGMA₇₁-PHPMA₄₀₀-20 (denoted $G_{71}\text{-H}_{400}\text{-20}$) vesicles. Each aqueous dispersion was analyzed by DLS immediately after dilution, again after 24 h and finally after ageing for one week at 20 $^{\circ}\text{C}$. In such experiments, any apparent increase in size is taken to be evidence of particle aggregation caused by colloidal destabilization. The results are summarized in Table 1 (see overleaf).

$S_{38}\text{-H}_{400}\text{-25}$ vesicles remained stable in either 1 M or 2 M NaCl solution. In 1 M MgSO_4 solution, the vesicles actually appear *smaller* compared to that observed in water (119 nm vs. 140 nm). TEM studies confirmed that no discernible change in copolymer morphology occurred under these conditions (see Fig. S4 in ESI[†]). A possible explanation for these DLS observations is that the PSBMA chains expressed at the outer leaflet of the vesicles become less hydrated in the presence of 1 M MgSO_4 , which in turn reduces the mean hydrodynamic diameter of the vesicles. In contrast, exposure to 2 M MgSO_4 solution results in macroscopic precipitation of the $S_{38}\text{-H}_{400}$ vesicles (see Fig. S5 in ESI[†]). This is commonly known as the "salting out" effect. Dissolution of this 2:2 electrolyte produces a significantly higher ionic strength than 1:1 electrolytes such as NaCl, which explains why the $S_{38}\text{-H}_{400}$ vesicles remain colloiddally stable in 2 M NaCl, but become aggregated in the presence of 2 M MgSO_4 .

As a comparison, $G_{71}\text{-H}_{400}\text{-20}$ vesicles were also exposed to various salt solutions. These vesicles also exhibited reasonable colloidal stability in either 1 M or 2 M NaCl for at least a week (see Table 1), although a small degree of aggregation can be detected in the particle size distributions obtained by DLS (see



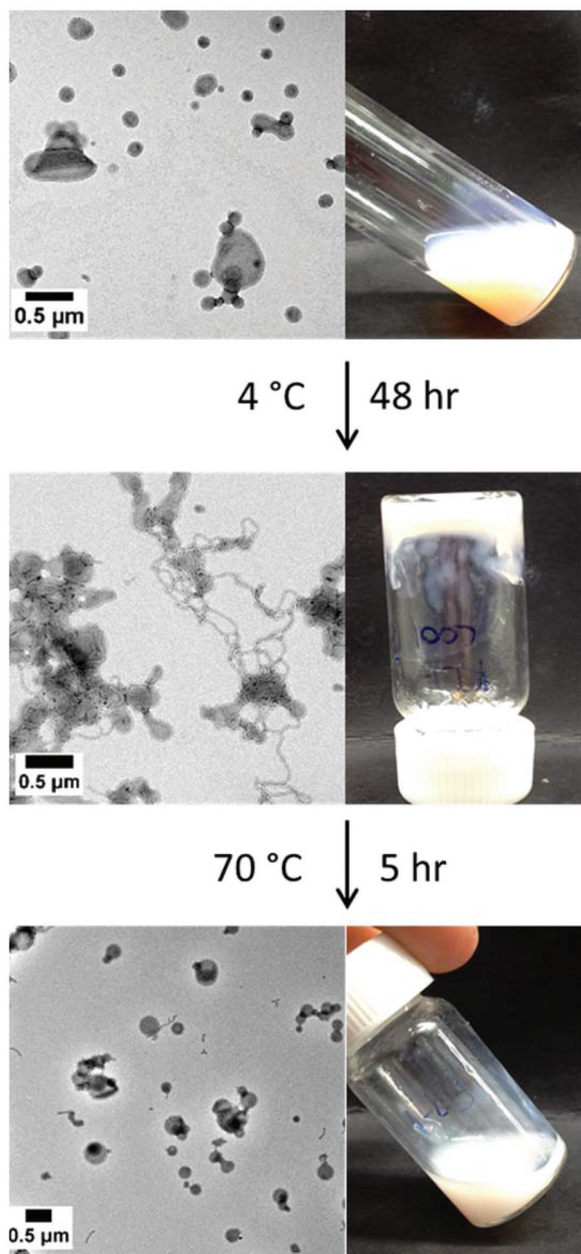


Fig. 9 TEM images and physical state of the dispersions obtained for (top) S_{38} - H_{310} -25 vesicles at 20 °C and (middle) a mixed phase of worms and vesicles formed after being held at 4 °C for 48 h and (bottom) a mixed phase of vesicles and short worms obtained after heating to 70 °C for 5 hours.

Fig. 10). However, unlike the S_{38} - H_{400} spheres and vesicles, the G_{71} - H_{400} vesicles proved to be unstable with respect to particle aggregation in the presence of 1 M $MgSO_4$, with immediate macroscopic precipitation being observed under these conditions (see Fig. S5 in the ESI†). In summary, these ‘added salt’ studies indicate that the zwitterionic PSBMA₃₈ block is a somewhat more salt-tolerant steric stabilizer than the non-ionic PGMA₇₁ block under the same conditions, which is consistent with earlier literature reports.^{27,37,54–56}

Table 1 Summary of z-average DLS diameters obtained for PSBMA₃₈-stabilized vesicles and PGMA₇₁-stabilized vesicles at 20 °C in the presence of either water or added salt

Nanoparticle type	Aqueous solution	Initial DLS diameter/nm (PDI)	DLS diameter after one week/nm (PDI)
S_{38} - H_{400} -25 vesicles	Water	140 (0.03)	140 (0.03)
	1 M NaCl	140 (0.04)	141 (0.03)
	2 M NaCl	140 (0.04)	145 (0.02)
	1 M $MgSO_4$	119 (0.05)	118 (0.09)
G_{71} - H_{400} -20 vesicles	Water	314 (0.14)	315 (0.12)
	1 M NaCl	307 (0.19)	310 (0.22)
	2 M NaCl	314 (0.19)	312 (0.15)
	1 M $MgSO_4$	Precipitate	—

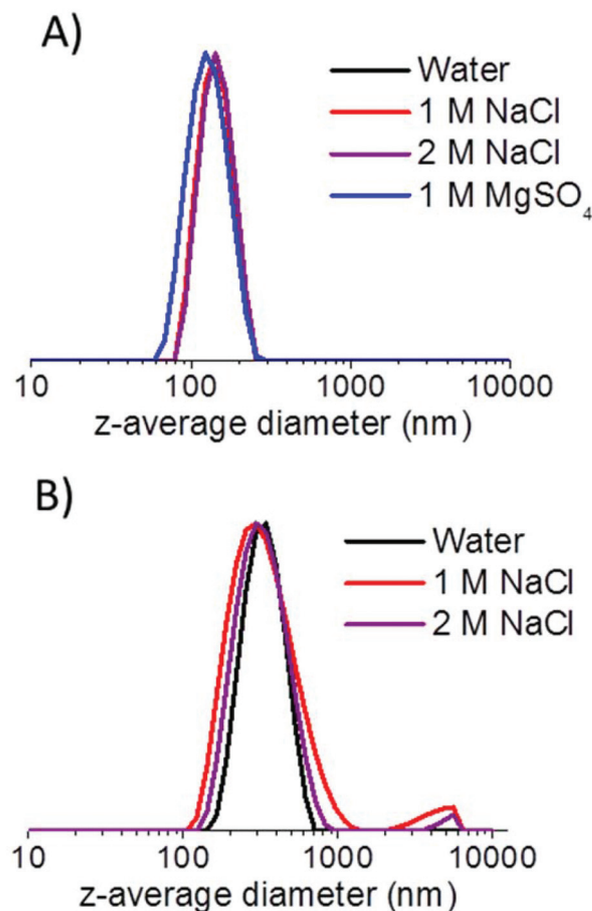


Fig. 10 Intensity-average particle size distributions obtained by DLS for (a) S_{38} - H_{400} vesicles and (b) G_{71} - H_{400} vesicles after being aged in various aqueous salt solutions for one week.

Conclusions

A near-monodisperse polysulfobetaine macro-CTA with a mean DP of 38 has been utilized as the water-soluble stabilizer block for the RAFT aqueous dispersion polymerization of HPMA to produce a range of copolymer morphologies *via* polymerization-induced self-assembly (PISA). Systematic variation of the



mean DP of the structure-directing hydrophobic PHPMA block and the copolymer concentration enables pure spheres, worms or vesicles to be obtained. However, construction of a detailed phase diagram is required for reproducible targeting of pure worms, since this anisotropic morphology occupies rather narrow phase space. Degelation of a $S_{38}\text{-H}_{200}$ worm gel occurs on cooling: this thermal transition is *irreversible* as judged by both the tube inversion test and temperature-dependent oscillatory rheology studies. TEM images recorded for dried diluted aqueous dispersions prepared before and after cooling suggests that degelation is the result of a worm-to-sphere transition. In addition, a pure vesicle phase undergoes a sol-gel transition upon cooling as a result of formation of a mixed phase comprising worms and vesicles. Finally, the colloidal stability of $S_{38}\text{-H}_{400}$ vesicles in the presence of added salt was compared to that of a control sample of $G_{71}\text{-H}_{400}$ vesicles. The $S_{38}\text{-H}_{400}$ vesicles remained stable with respect to aggregation in the presence of 1 M MgSO_4 , whereas $G_{71}\text{-H}_{400}$ vesicles underwent macroscopic precipitation under the same conditions. Thus, the zwitterionic PSBMA stabilizer confers significantly higher salt tolerance than a non-ionic poly(glycerol monomethacrylate) stabilizer.

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Notes and references

- 1 A. Blanazs, S. P. Armes and A. J. Ryan, *Macromol. Rapid Commun.*, 2009, **30**, 267–277.
- 2 J. Israelachvili, *Intermolecular and Surface Forces*, Academic Press, London, 2nd edn, 1991.
- 3 G. Riess, *Prog. Polym. Sci.*, 2003, **28**, 1107–1170.
- 4 L. Zhang and A. Eisenberg, *Science*, 1995, **268**, 1728–1731.
- 5 D. E. Discher and A. Eisenberg, *Science*, 2002, **297**, 967–973.
- 6 K. Kita-Tokarczyk, J. Grumelard, T. Haeefele and W. Meier, *Polymer*, 2005, **46**, 3540–3563.
- 7 J. Du and R. K. O'Reilly, *Soft Matter*, 2009, **5**, 3544–3561.
- 8 L. Zhang and A. Eisenberg, *J. Am. Chem. Soc.*, 1996, **118**, 3168–3181.
- 9 Y. Li and S. P. Armes, *Angew. Chem., Int. Ed.*, 2010, **49**, 4042–4046.
- 10 S. Sugihara, A. Blanazs, S. P. Armes, A. J. Ryan and A. L. Lewis, *J. Am. Chem. Soc.*, 2011, **133**, 15707–15713.
- 11 A. Blanazs, A. J. Ryan and S. P. Armes, *Macromolecules*, 2012, **45**, 5099–5107.
- 12 V. Ladmiral, M. Semsarilar, I. Canton and S. P. Armes, *J. Am. Chem. Soc.*, 2013, **135**, 13574–13581.
- 13 N. J. Warren, O. O. Mykhaylyk, D. Mahmood, A. J. Ryan and S. P. Armes, *J. Am. Chem. Soc.*, 2013, **136**, 1023–1033.
- 14 Z. An, Q. Shi, W. Tang, C.-K. Tsung, C. J. Hawker and G. D. Stucky, *J. Am. Chem. Soc.*, 2007, **129**, 14493–14499.
- 15 G. Delaittre, M. Save and B. Charleux, *Macromol. Rapid Commun.*, 2007, **28**, 1528–1533.
- 16 Y. Pei and A. B. Lowe, *Polym. Chem.*, 2014, **5**, 2342–2351.
- 17 C. A. Figg, A. Simula, K. A. Gebre, B. S. Tucker, D. M. Haddleton and B. S. Sumerlin, *Chem. Sci.*, 2015, **6**, 1230–1236.
- 18 J. Rieger, C. Gazon, B. Charleux, D. Alaimo and C. Jérôme, *J. Polym. Sci., Part A: Polym. Chem.*, 2009, **47**, 2373–2390.
- 19 N. J. Warren and S. P. Armes, *J. Am. Chem. Soc.*, 2014, **136**, 10174–10185.
- 20 J. Rieger, F. Stoffelbach, C. Bui, D. Alaimo, C. Jérôme and B. Charleux, *Macromolecules*, 2008, **41**, 4065–4068.
- 21 V. J. Cunningham, A. M. Alswieleh, K. L. Thompson, M. Williams, G. J. Leggett, S. P. Armes and O. M. Musa, *Macromolecules*, 2014, **47**, 5613–5623.
- 22 J. Rieger, G. Osterwinter, C. Bui, F. Stoffelbach and B. Charleux, *Macromolecules*, 2009, **42**, 5518–5525.
- 23 J. Chiefari, Y. Chong, F. Ercole, J. Krstina, J. Jeffery, T. Le, R. Mayadunne, G. Meijs, C. Moad and G. Moad, *Macromolecules*, 1998, **31**, 5559–5562.
- 24 G. Moad, E. Rizzardo and S. H. Thang, *Aust. J. Chem.*, 2005, **58**, 379–410.
- 25 G. Moad, E. Rizzardo and S. H. Thang, *Aust. J. Chem.*, 2012, **65**, 985–1076.
- 26 D. J. Keddie, *Chem. Soc. Rev.*, 2014, **43**, 496–505.
- 27 A. B. Lowe, N. C. Billingham and S. P. Armes, *Macromolecules*, 1999, **32**, 2141–2148.
- 28 A. B. Lowe and C. L. McCormick, *Chem. Rev.*, 2002, **102**, 4177–4190.
- 29 Y. J. Che, Y. Tan, J. Cao and G. Y. Xu, *J. Macromol. Sci., Phys.*, 2010, **49**, 695–710.
- 30 P. Mary, D. D. Bendejacq, M. P. Labeau and P. Dupuis, *J. Phys. Chem. B*, 2007, **111**, 7767–7777.
- 31 J. C. Salamone, W. Volksen, A. P. Olson and S. C. Israel, *Polymer*, 1978, **19**, 1157–1162.
- 32 J. Virtanen, M. Arotçaréna, B. Heise, S. Ishaya, A. Laschewsky and H. Tenhu, *Langmuir*, 2002, **18**, 5360–5365.
- 33 D. Wang, T. Wu, X. Wan, X. Wang and S. Liu, *Langmuir*, 2007, **23**, 11866–11874.
- 34 M. Kobayashi, Y. Terayama, M. Kikuchi and A. Takahara, *Soft Matter*, 2013, **9**, 5138–5148.
- 35 J. Rodríguez-Hernández and S. Lecommandoux, *J. Am. Chem. Soc.*, 2005, **127**, 2026–2027.
- 36 W. Schärtl, *Light Scattering from Polymer Solutions and Nanoparticle Dispersions*, Springer-Verlag, Berlin Heidelberg, Berlin, 2007.
- 37 D. N. Schulz, D. G. Peiffer, P. K. Agarwal, J. Larabee, J. J. Kaladas, L. Soni, B. Handwerker and R. T. Garner, *Polymer*, 1986, **27**, 1734–1742.
- 38 Z. Dong, J. Mao, D. Wang, M. Yang, W. Wang, S. Bo and X. Ji, *Macromol. Chem. Phys.*, 2014, **215**, 111–120.
- 39 H. Willcock, A. Lu, C. F. Hansell, E. Chapman, I. R. Collins and R. K. O'Reilly, *Polym. Chem.*, 2014, **5**, 1023–1030.



- 40 J. Du, Y. Tang, A. L. Lewis and S. P. Armes, *J. Am. Chem. Soc.*, 2005, **127**, 17982–17983.
- 41 S. L. West, J. P. Salvage, E. J. Lobb, S. P. Armes, N. C. Billingham, A. L. Lewis, G. W. Hanlon and A. W. Lloyd, *Biomaterials*, 2004, **25**, 1195–1204.
- 42 J. P. Salvage, S. F. Rose, G. J. Phillips, G. W. Hanlon, A. W. Lloyd, I. Y. Ma, S. P. Armes, N. C. Billingham and A. L. Lewis, *J. Controlled Release*, 2005, **104**, 259–270.
- 43 Y. J. Shih, Y. Chang, A. Deratani and D. Quemener, *Biomacromolecules*, 2012, **13**, 2849–2858.
- 44 Y. J. Shih and Y. Chang, *Langmuir*, 2010, **26**, 17286–17294.
- 45 Y. Chang, W. Y. Chen, W. Yandi, Y. J. Shih, W. L. Chu, Y. L. Liu, C. W. Chu, R. C. Ruaan and A. Higuchi, *Biomacromolecules*, 2009, **10**, 2092–2100.
- 46 Y. Chang, S. C. Liao, A. Higuchi, R. C. Ruaan, C. W. Chu and W. Y. Chen, *Langmuir*, 2008, **24**, 5453–5458.
- 47 F. Lee Rodkey, *Clin. Chem.*, 1966, **12**, 517–518.
- 48 F. Liu, J. Seuring and S. Agarwal, *J. Polym. Sci., Part A: Polym. Chem.*, 2012, **50**, 4920–4928.
- 49 C. Barner-Kowollik, M. Buback, B. Charleux, M. L. Coote, M. Drache, T. Fukuda, A. Goto, B. Klumperman, A. B. Lowe, J. B. McLeary, G. Moad, M. J. Monteiro, R. D. Sanderson, M. P. Tonge and P. Vana, *J. Polym. Sci., Part A: Polym. Chem.*, 2006, **44**, 5809–5831.
- 50 G. Moad, *Macromol. Chem. Phys.*, 2014, **215**, 9–26.
- 51 A. Blanazs, J. Madsen, G. Battaglia, A. J. Ryan and S. P. Armes, *J. Am. Chem. Soc.*, 2011, **133**, 16581–16587.
- 52 A. Blanazs, R. Verber, O. O. Mykhaylyk, A. J. Ryan, J. Z. Heath, C. W. I. Douglas and S. P. Armes, *J. Am. Chem. Soc.*, 2012, **134**, 9741–9748.
- 53 J. Madsen, S. P. Armes, K. Bertal, S. MacNeil and A. L. Lewis, *Biomacromolecules*, 2009, **10**, 1875–1887.
- 54 S. F. Lascelles, F. Malet, R. Mayada, N. C. Billingham and S. P. Armes, *Macromolecules*, 1999, **32**, 2462–2471.
- 55 A. B. Lowe, N. C. Billingham and S. P. Armes, *Chem. Commun.*, 1996, 1555–1556.
- 56 Z. Zhu, J. Xu, Y. Zhou, X. Jiang, S. P. Armes and S. Liu, *Macromolecules*, 2007, **40**, 6393–6400.

