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RAFT aqueous emulsion polymerization of methyl methacrylate: observation of unexpected constraints when employing a non-ionic steric stabilizer block†

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The RAFT aqueous emulsion polymerization of methyl methacrylate (MMA) is conducted at 70 °C using poly(glycerol monomethacrylate) (PGMA) as a steric stabilizer block. This non-ionic precursor has previously proved to be highly effective for the RAFT aqueous emulsion polymerization of various vinyl monomers such as benzyl methacrylate (BzMA), 2,2,2-trifluoroethyl methacrylate (TFEMA), isopropylidene glycerol monomethacrylate (IPGMA) or glycidyl methacrylate. However, an unexpected constraint was encountered in the case of MMA. Targeting a degree of polymerization (DP) of 20 to 100 for the PMMA block led to colloidal dispersions of kinetically-trapped spherical nanoparticles ranging in size from 17 nm to 31 nm. On the other hand, targeting DPs above 100 invariably led to the formation of highly flocculated spherical nanoparticles. This rather limited DP range is in striking contrast to the much higher DPs that can be targeted without loss of colloidal stability when using more hydrophobic monomers such as BzMA, TFEMA or IPGMA. The same flocculation problem was also evident when employing a PGMA precursor containing an anionic carboxylate end-group, but a series of colloidal stable dispersions could be obtained when using an anionic poly(methacrylic acid) stabilizer. Finally, the efficient removal of RAFT end-groups from PGMA₅₀-PMMA₈₀ nanoparticles was achieved by visible light irradiation using a blue LED source ($\lambda = 405$ nm). UV GPC studies confirmed that up to 87% dithiobenzoate end-groups can be removed from such nanoparticles within 12 h at 80 °C. On the other hand, using excess H₂O₂ under the same conditions only led to 24% end-group removal. This is because this water-soluble reagent has restricted access to the hydrophobic PMMA cores.

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Introduction

Reversible addition–fragmentation chain transfer (RAFT) polymerization is a well-established pseudo-living technique based on the principle of rapid reversible chain transfer that

enables the convenient synthesis of a wide range of controlled-structure vinyl polymers using various organosulfur-based reagents.^{1–5} RAFT polymerization is exceptionally tolerant of monomer functionality and is usually conducted in either organic solvents or aqueous solution to afford soluble polymer chains.⁶ However, it is equally well-suited to heterogeneous conditions such as suspension,⁷ dispersion^{8–12} or emulsion^{13–15} polymerization. In particular, RAFT aqueous emulsion polymerization – which is an important example of polymerization-induced self-assembly (PISA)^{16,17} – enables the controlled polymerization of a wide range of water-immiscible vinyl monomers in aqueous media.^{17–19} Typically, spherical nanoparticles of tunable particle size can be readily prepared at high solids,¹⁶ although so-called ‘higher order’ copolymer morphologies (e.g. worms or vesicles) are also accessible for at least some formulations.^{18–20} Potential applications for such nano-objects include surface modifiers for cellulose fibres,²¹ viscosity modifiers,²² organic opacifiers for paints²³ and additives for the reinforcement of latex films.²⁴

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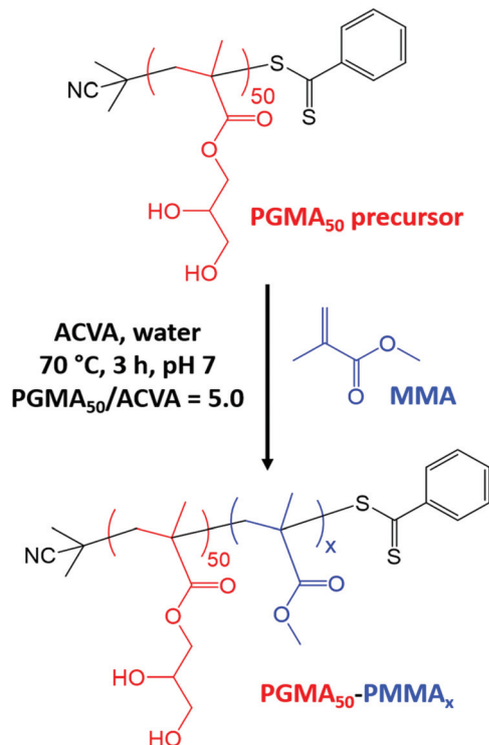
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† Electronic supplementary information (ESI) available: Summary of GPC, DLS and TEM data obtained for HOOC-PGMA₅₄-PMMA₈₀ nanoparticles and HOOC-PGMA₅₄-PMMA₁₅₀ nanoparticles synthesised with PETTC RAFT agent; synthesis of the PMETAC₄₆ precursor and the corresponding PMETAC₄₆-PMMA_x nanoparticles; DLS and TEM data obtained for PMETAC₄₆-PMMA_x nanoparticles; UV GPC curves ($\lambda = 309$ nm) recorded during kinetic studies of the removal of dithiobenzoate end-groups from an aqueous dispersion of PGMA₅₀-PMMA₈₀ nanoparticles using either excess H₂O₂ or no light irradiation at 80 °C. UV GPC curves ($\lambda = 309$ nm) recorded during the removal of trithiocarbonate end-groups from an aqueous dispersion of HOOC-PGMA₅₄-PMMA₈₀ nanoparticles using visible light irradiation, excess H₂O₂ or neither condition at 80 °C. See DOI: 10.1039/d1py01008e



There are many reports of the RAFT aqueous emulsion polymerization of commodity monomers such as styrene,^{12,23,25–27} *n*-butyl acrylate^{13,28–30} or vinyl acetate.³¹ Moreover, there are also various reports of the use of speciality vinyl monomers such as benzyl methacrylate,³² 2,2,2-trifluoroethyl methacrylate,³³ isopropylidenglycerol monomethacrylate,³⁴ 2-methoxyethyl methacrylate,³⁵ glycidyl methacrylate,^{36–39} or hydroxybutyl methacrylate.⁴⁰ However, given its undoubted importance as a commodity monomer, there are surprisingly few literature reports of the RAFT aqueous emulsion polymerization of methyl methacrylate (MMA).^{21,29,41–43} Moreover, given the moderately high aqueous solubility of this monomer (15.9 g dm⁻³ at 20 °C), such PISA formulations might be expected to provide convenient access to a wide range of block copolymer nano-objects (*e.g.* spheres, worms and vesicles).^{35,39,40,44}

In the present study, we examine the RAFT emulsion polymerization of MMA using poly(glycerol monomethacrylate) (PGMA) as a water-soluble precursor (Scheme 1), which is known to act as an effective non-ionic steric stabilizer for various other water-immiscible vinyl monomers under such conditions.^{32–36} In view of the unexpected limited utility of this PISA formulation, a PGMA stabilizer containing an anionic carboxylate end-group and an anionic poly(methacrylic acid) stabilizer were also evaluated for the RAFT emulsion polymerization of MMA.



Scheme 1 Synthesis of PGMA₅₀-PMMA_x diblock copolymer nanoparticles by RAFT aqueous emulsion polymerization of methyl methacrylate MMA at 70 °C using a dithiobenzoate-capped PGMA₅₀ precursor to target *x* = 20 to 130 at pH 7.

Experimental section

Materials

Glycerol monomethacrylate (GMA) was donated by GEO Specialty Chemicals (Hythe, UK). Methyl methacrylate (MMA; 99%), 4,4'-azobis(4-cyanopentanoic acid) (ACVA; 98%), methacrylic acid (MAA; 99%), [2-(methacryloyloxy)ethyl]triethylammonium chloride (METAC; 80% solution), 2-cyano-2-propyl benzodithioate (CPDB; 97%) and trimethylsilyl diazomethane solution (2.0 M in hexanes) were each purchased from Sigma-Aldrich (UK) and used as received. 4-Cyano-4-(2-phenylethanesulfanylthiocarbonyl) sulfanylpentanoic acid (PETTC) was prepared according to a literature protocol.⁴⁵ Deionized water from an Elga Medica DV25 water purification unit was used in all the experiments.

Synthesis of the PGMA₅₀ precursor by RAFT aqueous solution polymerization

GMA monomer (30.0 g, 187 mmol), CPDB RAFT agent (0.589 g, 2.66 mmol; target PGMA DP = 70) and ACVA initiator (0.149 g, 0.53 mmol; CPDB/ACVA molar ratio = 5.0) were weighed into a 250 mL round-bottom flask. Ethanol (46.5 g, 60% w/w) was added and the flask was cooled by immersing in an ice bath while degassed with N₂ gas for 30 min. The flask was then immersed in an oil bath at 70 °C for 165 min. The polymerization was quenched by exposing the reaction mixture to air and cooling the flask to 20 °C. ¹H NMR spectroscopy indicated a final GMA conversion of 71%. The reaction solution was diluted with methanol (30 mL) and then precipitated into a ten-fold excess of dichloromethane (three times). End-group analysis *via* ¹H NMR spectroscopy indicated a mean degree of polymerization (DP) of 50 (the integrated aromatic proton signals at 7.4–7.8 ppm were compared to the integrated methacrylic backbone proton signals at 0.7–2.5 ppm).

Synthesis of the HOOC-PGMA₅₄ precursor by RAFT aqueous solution polymerization

GMA monomer (10.0 g, 62.4 mmol), PETTC RAFT agent (0.302 g, 0.892 mmol; target PGMA DP = 70) and ACVA initiator (0.050 g, 0.18 mmol; PETTC/ACVA molar ratio = 5.0) were weighed into a 100 mL round-bottom flask. Ethanol (15.5 g, 60% w/w) was added and the flask was cooled by immersing in an ice bath while degassed with N₂ gas for 30 min. The flask was then immersed in an oil bath at 70 °C for 120 min. The polymerization was quenched by exposing the reaction mixture to air and cooling the flask to 20 °C. ¹H NMR spectroscopy indicated a final GMA conversion of 77%. The solution was diluted with methanol (10 mL) and then precipitated into a ten-fold excess of dichloromethane (three times). End-group analysis *via* ¹H NMR spectroscopy indicated a mean DP of 54.

Synthesis of PGMA₅₀-PMMA_x diblock copolymer nanoparticles by RAFT aqueous emulsion polymerization

A typical protocol for the synthesis of PGMA₅₀-PMMA_x diblock copolymer nanoparticles was as follows. PGMA₅₀ precursor



(0.150 g, 18.2 μmol), MMA monomer (0.146 g, 1.46 mmol, target DP = 80), ACVA initiator (1.00 mg, 3.65 μmol , PGMA₅₀/ACVA molar ratio = 5.0) and deionized water (2.675 g, 10% w/w solution) were added to a 20 mL round-bottom flask. The mixture was adjusted to pH 7 using 1 M NaOH. The flask was then placed in an ice bath and degassed with N₂ gas for 30 min, before immersing the flask in an oil bath set at 70 °C. After 3 h, the polymerization was quenched by exposing the reaction mixture to air while cooling the flask to 20 °C. Essentially the same protocol was employed for the synthesis of HOOC-PGMA₅₄-PMMA_x diblock copolymer nanoparticles but in this case the HOOC-PGMA₅₄ precursor was used instead of the PGMA₅₀ precursor.

Synthesis of the PMAA₅₆ precursor by RAFT aqueous solution polymerization

MAA monomer (45.0 g, 0.523 mol), PETTC RAFT agent (3.165 g, 9.34 mmol; target DP = 56) and ACVA initiator (0.523 g, 1.87 mmol; PETTC/ACVA molar ratio = 5.0) and ethanol (73.0 g, 40% w/w) were weighed into a 500 mL round-bottom flask. The solution was placed in an ice bath and degassed with N₂ gas for 30 min, before the flask was immersed in an oil bath at 70 °C. The polymerization was quenched after 3 h by exposing the reaction mixture to air while cooling the flask to 20 °C. The crude polymer was then precipitated into a ten-fold excess of diethyl ether. The insoluble polymer was redissolved in ethanol prior to a second precipitation step and then freeze-dried overnight. End-group analysis by ¹H NMR spectroscopy indicated a DP of 56 for this PMAA precursor.

Synthesis of PMAA₅₆-PMMA_x diblock copolymer nanoparticles by RAFT aqueous emulsion polymerization

A typical protocol for the synthesis of PMAA₅₆-PMMA₁₀₀ diblock copolymer nanoparticles was conducted as follows. PMAA₅₆ precursor (0.100 g, 19.4 μmol), MMA monomer (0.194 g, 1.94 mmol, target DP = 100), ACVA initiator (1.10 mg, 3.88 μmol , PMAA₅₆/ACVA molar ratio = 5.0) and deionized water (1.18 g, 20% w/w solution) were added to a 20 mL round-bottom flask. The mixture was adjusted to pH 5 using 1 M HCl. The reaction mixture was then placed in an ice bath and degassed with N₂ gas for 30 min, before immersing in an oil bath set at 70 °C. The polymerization was allowed to proceed at this temperature for 6 h and then quenched by exposing the reaction mixture to air while cooling the flask to 20 °C.

Methylation of PMAA₅₆-PMMA_x diblock copolymer nanoparticles

Methylation of the carboxylic acid groups on the PMAA block was performed before THF GPC analysis. PMAA₅₆-PMMA_x diblock copolymer powder (20 mg) was diluted in THF (2.0 mL). Excess trimethylsilyldiazomethane was then gradually added to this solution until it turned yellow. The reaction mixture was stirred overnight, dried and analysed by THF GPC.

Protocols for cleavage of RAFT end-groups

The dithiobenzoate end-groups within PGMA₅₀-PMMA₈₀ nanoparticles were cleaved using the following protocol. A 10% w/w

copolymer dispersion (1.00 g) was placed in a water-jacketed Schlenk tube wrapped in blue LED light strips ($\lambda = 405 \text{ nm}$, 0.37 mW cm^{-2}) with the temperature of the water within the recirculating jacket set to 80 °C. Aliquots of this reaction mixture were extracted periodically and analyzed using UV GPC (with the UV detector set at $\lambda = 308 \text{ nm}$). As a control experiment, the same protocol was used but without the blue LED light strips (*i.e.* no visible light irradiation). The same 10% w/w dispersion of PGMA₅₀-PMMA₈₀ nanoparticles was also treated with H₂O₂ (using a five-fold excess relative to the concentration of dithiobenzoate end-groups, as reported by Jesson and co-workers⁴⁶) at 80 °C with no visible light irradiation. The same protocol was used for cleaving trithiocarbonate RAFT end-groups from a 10% w/w aqueous dispersion of PGMA₅₄-PMMA₈₀ nanoparticles. This included cleavage at 80 °C using blue LED strips, H₂O₂ or no visible light irradiation for 24 h.

Dynamic light scattering and aqueous electrophoresis

DLS and aqueous electrophoresis studies were conducted on 0.50% w/w aqueous dispersions using a Malvern Zetasizer NanoZS instrument. The hydrodynamic z-average diameter was determined at 20 °C at a scattering angle of 173° and averaged over three measurements. Aqueous electrophoresis studies were conducted in the presence of 1 mM KCl as background electrolyte. The solution pH was adjusted using either NaOH or HCl. Zeta potentials (also averaged over three measurements) were calculated *via* the Henry equation using the Smoluchowski approximation.

Gel permeation chromatography (GPC)

Molecular weight distributions of the PGMA₅₀ and HOOC-PGMA₅₄ precursors and a series of PGMA₅₀-PMMA₈₀ diblock copolymers were assessed using DMF eluent (containing 10 mM LiBr) at 60 °C. Two Polymer Laboratories PL gel 5 μm Mixed-C columns were connected in series to a Varian 290-LC pump injection module and a Varian 390-LC multidetector suite (refractive index detector) at a flow rate of 1.0 mL min⁻¹. Near-monodisperse poly(methyl methacrylate) standards with M_p values ranging from 645 g mol⁻¹ to 618 000 g mol⁻¹ were used for calibration. Molecular weight distributions of methylated PMAA₅₆-PMMA_x diblock copolymers were assessed using THF eluent (containing 0.05% w/v butylhydroxytoluene and 2.0% v/v trimethylamine). Two Polymer Laboratories PL gel 5 μm Mixed-C columns were connected in series to a WellChrom K-2301 refractive index detector at a flow rate of 1.0 mL min⁻¹. Near-monodisperse poly(methyl methacrylate) standards were used for calibration.

Transmission electron microscopy (TEM)

Copper/palladium TEM grids (Agar Scientific, UK) were coated with a thin film of amorphous carbon. Grids were treated with a plasma glow discharge for 30 s to produce a hydrophilic surface. A 10 μL droplet of a 0.10% w/w aqueous dispersion was placed on a grid and left for 1 min before blotting. The adsorbed nanoparticles were then stained using uranyl formate (9.0 μL of a 0.75% w/w aqueous solution) for 20 s fol-



lowed by blotting to remove excess stain. Grids were carefully dried under vacuum and images were recorded at 100 kV using a Philips CM100 instrument equipped with a Gatan 1k CCD camera.

Small-angle X-ray scattering (SAXS)

SAXS patterns were collected using a Xeuss 2.0 (Xenocs) SAXS instrument equipped with a Dectris Pilatus 1 M detector and an Excillum liquid gallium MetalJet X-ray source ($\lambda = 1.34 \text{ \AA}$). SAXS patterns were recorded for 1.0% w/w aqueous dispersions of PGMA₅₀-PMMA_x nanoparticles over a scattering vector q range of 0.04–0.4 \AA^{-1} using 2.0 mm diameter glass capillary cells. The scattering of deionized water was used for absolute intensity calibration. Irena SAS macros for Igor Pro software were used to perform background subtraction, normalization and data analysis.⁴⁷

Results and discussion

RAFT aqueous emulsion polymerization of MMA

A PGMA precursor was prepared by RAFT solution polymerization of GMA in ethanol using a dithiobenzoate-based RAFT agent (CPDB) at 70 °C. End-group analysis by ¹H NMR spectroscopy indicated had a mean DP of 50. This PGMA₅₀ precursor was subsequently chain-extended *via* RAFT aqueous emulsion polymerization of MMA. A series of PGMA₅₀-PMMA_x nanoparticles were obtained by systematically varying the target DP of the PMMA core-forming block (Scheme 1). High MMA conversions were achieved within 3 h at 70 °C for all syntheses (Table 1). DMF GPC analysis indicated relatively high blocking efficiencies and unimodal molecular weight distributions in each case (Fig. 1). Increasing the target PMMA DP led to progressively higher M_n values, albeit with a gradual increase in dispersity (from $M_w/M_n = 1.17$ for PGMA₅₀-PMMA₂₀ up to $M_w/M_n = 1.37$ for PGMA₅₀-PMMA₁₃₀). Similar observations have been reported for various other PISA formulations in the literature.^{21,29,41}

TEM studies confirmed that kinetically-trapped spherical nanoparticles of increasing size were obtained when systematically varying the DP of the core-forming PMMA block (x) from 20 to 130 (Fig. 2). Furthermore, dynamic light scattering (DLS) studies revealed a systematic increase in the hydrodynamic z -average diameter while DLS polydispersities remained reasonably low up to $x = 100$, signifying relatively narrow par-

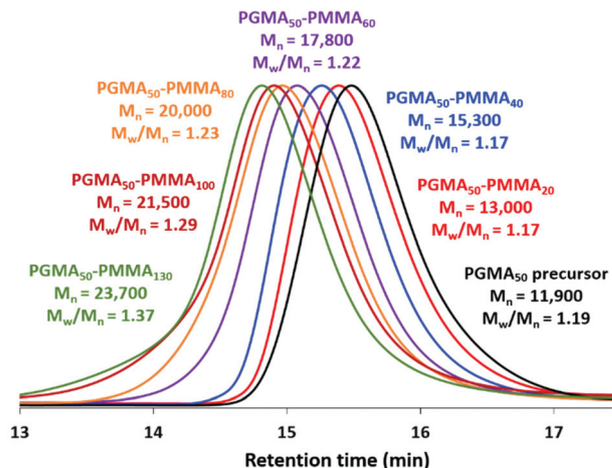


Fig. 1 DMF GPC curves recorded for the PGMA₅₀ precursor and a series of PGMA₅₀-PMMA_x nanoparticles for which the target PMMA DP (x) is 20, 40, 60, 80, 100 or 130 (see Table 1 for further details).

ticle size distributions (see Table 1 and Fig. 2). However, a substantial increase in both the apparent z -average diameter and DLS polydispersity was observed when targeting $x = 130$. Moreover, the TEM image recorded for this PISA synthesis indicates aggregates or clusters of nanoparticles on the grid, suggesting colloidal instability. Various PISA syntheses were conducted targeting $x > 100$ and similar results were invariably obtained (data not shown). This was an unexpected limitation, not least because we had previously reported that non-ionic dithiobenzoate-based PGMA precursors with similar (or lower) DPs were effective when performing RAFT aqueous emulsion polymerization syntheses with more hydrophobic methacrylic monomers. For example, Akpınar and co-workers³³ were able to prepare PGMA₂₈-PTFEMA₅₀₀ and PGMA₄₃-PTFEMA₁₀₀₀ spherical nanoparticles with no loss in colloidal stability. Similar results were also obtained by Jesson and co-workers when preparing PGMA₃₉-PIPGMA₁₀₀₀ spheres³⁴ and by Cunningham *et al.* when preparing PGMA₅₁-PBzMA₁₀₀₀ spheres.³²

Thus, at first sight, this unforeseen limitation appeared more likely to be associated with the choice of PMMA as the core-forming block, rather than the use of PGMA as a non-ionic stabilizer block. A few additional experiments were conducted using a longer PGMA₁₀₁ block as a steric stabilizer. However, nanoparticle aggregation was still observed when tar-

Table 1 Summary of monomer conversions and mean particle diameters obtained for the synthesis of a series of PGMA₅₀-PMMA_{20–130} nanoparticles *via* RAFT aqueous emulsion polymerization of MMA at 70 °C targeting 10% w/w solids

Target PMMA DP	Conversion (%)	DLS z -average diameter (nm)	DLS polydispersity	TEM morphology	SAXS core diameter (nm)
20	>99	18	0.04	Spheres	14
40	>99	20	0.04	Spheres	15
60	>99	29	0.08	Spheres	18
80	>99	38	0.10	Spheres	21
100	>99	66	0.12	Spheres	24
130	99	422	0.49	Spheres	30



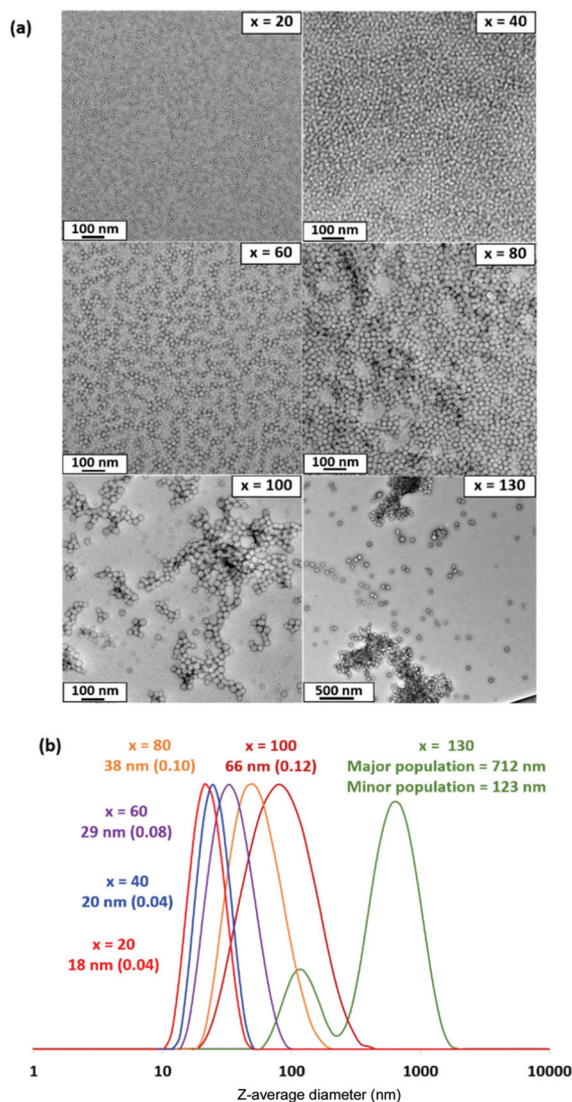


Fig. 2 (a) Representative TEM images and (b) DLS particle size distributions obtained for a series of PGMA₅₀-PMMA_x nanoparticles prepared by RAFT aqueous emulsion polymerization of MMA where the target PMMA DP (x) is 20, 40, 60, 80, 100 or 130. DLS indicates a bimodal particle size distribution in the latter case owing to nanoparticle flocculation.

getting PMMA DPs above 100. For example, DLS studies indicated incipient flocculation for PGMA₁₀₁-PMMA₂₀₀ nanoparticles prepared at 70 °C when targeting 10% w/w solids using the CPDB RAFT agent (data not shown).

SAXS patterns were recorded for 1.0% w/w aqueous dispersions of the PGMA₅₀-PMMA_{20–130} nanoparticles. Representative $I(q)$ vs. q plots for six dispersions are shown in Fig. 3. The local minimum observed for each pattern is shifted to lower q as higher PMMA DPs are targeted. This indicates a systematic increase in the nanoparticle core diameter, d , according to the well-known relation $q = 2\pi/d$.⁴⁸ These findings are consistent with the TEM and DLS data discussed above. Moreover, it is well-known that the low q gradient can be used

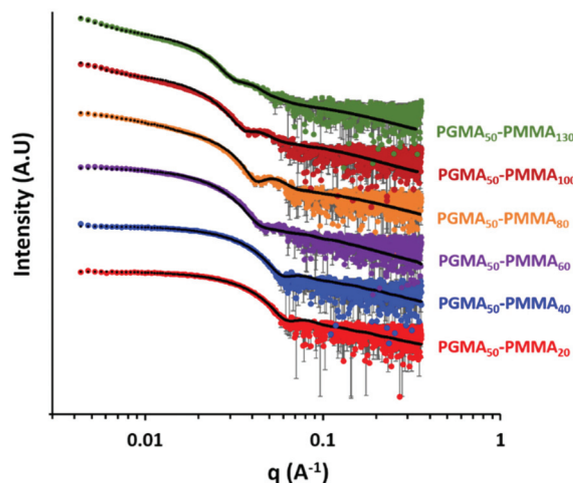


Fig. 3 SAXS patterns recorded for a series of 1.0% w/w aqueous dispersions of PGMA₅₀-PMMA_x nanoparticles for which the target PMMA DP has been systematically varied from 20 to 130.

to infer the predominant copolymer morphology.⁴⁹ More specifically, a low q gradient of zero indicates a spherical morphology, which is indeed observed when targeting PMMA DPs of between 20 and 80. However, non-zero low q gradients are observed for PGMA₅₀-PMMA₁₀₀ and PGMA₅₀-PMMA₁₃₀, which suggests incipient nanoparticle aggregation and the formation of mass fractals for these two dispersions. Again, this is consistent with the corresponding TEM and DLS data shown in Fig. 2. Fitting the SAXS patterns using a well-established spherical micelle model⁵⁰ and also a unified fit^{51–53} (to account for nanoparticle aggregation) enables the volume-average diameter of the PMMA cores to be determined in each case (Table 1).

Aqueous electrophoresis studies were performed on PGMA₅₀-PMMA₈₀ nanoparticles prepared using the dithiobenzoate-capped PGMA₅₀ precursor (Fig. 4a). These nanoparticles exhibited zeta potentials close to zero (approximately -3 mV) across the whole pH range, which is consistent with the non-ionic nature of this steric stabilizer block. Given the unexpected colloidal stability problems associated with this PISA formulation, an alternative PGMA stabilizer bearing a carboxylic acid end-group⁵⁴ and an anionic poly(methacrylic acid) stabilizer²⁹ were also evaluated for the RAFT aqueous emulsion polymerization of MMA (see Scheme 2). A trithiocarbonate-based RAFT agent, PETTC, was used to prepare a carboxylic acid-capped PGMA₅₄ precursor, which was then chain-extended *via* RAFT aqueous emulsion polymerization of MMA at pH 7 to produce either HOOC-PGMA₅₄-PMMA₈₀ or HOOC-PGMA₅₄-PMMA₁₅₀ spherical nanoparticles (Scheme 2a). The associated DLS, TEM and GPC data for these two dispersions are summarized in Fig. S1.†

The anionic charge conferred by the terminal ionized carboxylic group led to significant anionic character under the PISA synthesis conditions (pH 7) as judged by aqueous electrophoresis studies (see Fig. 4). More specifically, the nano-



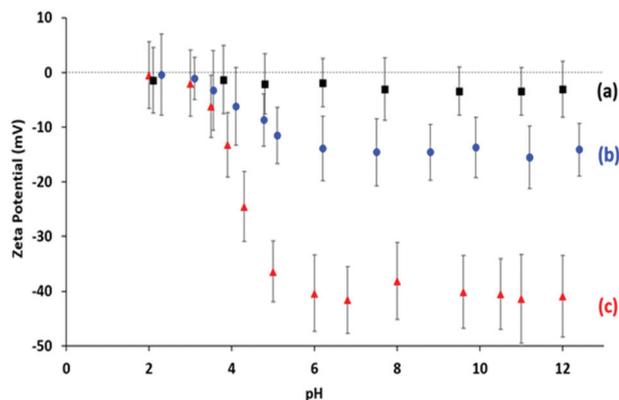
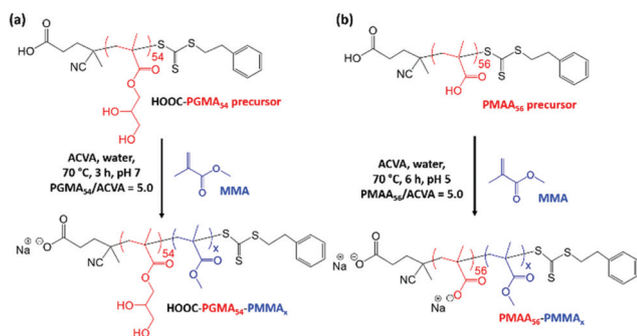


Fig. 4 Zeta potential vs. pH curves obtained for dilute aqueous dispersions of: (a) PGMA₅₀-PMMA₈₀ spheres prepared using the non-ionic dithiobenzoate-based PGMA₅₀ precursor shown in Scheme 1; (b) HOOC-PGMA₅₄-PMMA₈₀ spheres prepared using the carboxylic acid-functionalized trithiocarbonate-based PGMA₅₄ precursor shown in Scheme 2a; (c) PMAA₅₆-PMMA₁₀₀ spheres prepared using the anionic PMAA precursor shown in Scheme 2b.



Scheme 2 (a) Synthesis of HOOC-PGMA₅₄-PMMA_x diblock copolymer nanoparticles by RAFT aqueous emulsion polymerization of methyl methacrylate using a trithiocarbonate-based HOOC-PGMA₅₄ precursor at 70 °C and targeting $x = 80$ or 150 at pH 7 . (b) Synthesis of PMAA₅₆-PMMA_x diblock copolymer nanoparticles by RAFT aqueous emulsion polymerization of methyl methacrylate at 70 °C targeting $x = 50$ to 2000 at pH 5 .

particles become progressively more anionic between pH 2 and 6, before a maximum zeta potential of approximately -14 mV is observed at or above pH 6. DLS studies confirmed the formation of relatively small nanoparticles with a z-average diameter of 26 nm when targeting a PMMA DP of 80, which indicated good colloidal stability in this case (Fig. S1†). However, such anionic character was not sufficient to allow the PISA synthesis of colloidally stable spherical nanoparticles when targeting a PMMA DP of 150. In this case, DLS and TEM studies indicated extensive nanoparticle aggregation similar to that observed for the non-ionic PGMA₅₀-PMMA₁₃₀ nanoparticles.

Accordingly, a poly(methacrylic acid) (PMAA) precursor was evaluated for the RAFT aqueous emulsion polymerization of MMA.²⁹ A series of PMAA₅₆-PMMA_x nanoparticles was prepared using this polyelectrolytic steric stabilizer by systematic

variation of the target PMMA DP from 50 to 2000 (see Scheme 2b). All syntheses were performed at pH 5 targeting 20% w/w solids, as reported by Cockram and co-workers for aqueous PISA syntheses using a PMAA steric stabilizer.⁴⁰ ¹H NMR spectroscopy studies confirmed that relatively high conversions (94–99%) were obtained in each case, with a modest reduction in the final conversion being observed when targeting DPs above 500 (Table 2). Mean hydrodynamic diameters determined by DLS are summarized in Table 2 and the relationship between DLS diameter and the PMMA DP is shown in Fig. 5. As expected, the particle size increases monotonically as the target DP is increased from 50 to 2000. Indeed, an approximate linear relationship is observed between DP 50 (29 nm) and DP 500 (73 nm). Above DP 500, the particle size continues to increase up to DP 2000, albeit more slowly. TEM images for selected PMAA₅₆-PMMA_x nanoparticles are shown in Fig. 6 when targeting $x = 50$ to 2000. As expected, a kinetically-trapped spherical morphology was obtained in each case.

Selected PMAA₅₆-PMMA_x diblock copolymers were methylated to convert such copolymers into the corresponding PMMA_{56+x} homopolymers for THF GPC analysis using PMMA calibration standards (see Fig. 7). In each case, unimodal MWDs and high blocking efficiencies were observed and targeting higher PMMA DPs led to the expected monotonic increase in the GPC M_n . Notably, no systematic GPC error was incurred for this particular data set and the M_n values were reasonably close to the expected theoretical values. However, dispersities were relatively high ($M_w/M_n = 1.51$ – 1.76). Similar GPC data were reported by Chaduc and co-workers for the RAFT aqueous emulsion polymerization of MMA using a PMAA precursor at pH 3.5.²⁹ Clearly, this aqueous PISA formulation does not suffer from colloidal instability problems when targeting relatively high PMMA DPs. This is perhaps not surprising, because aqueous electrophoresis studies of PMAA₅₆-PMMA₁₀₀ nanoparticles (Fig. 4c) indicated highly negative zeta potentials (approximately -40 mV at or above pH 5) owing to the strongly anionic nature of the ionized PMAA₅₆ chains. Similar results were obtained when employing a cationic steric

Table 2 Summary of monomer conversions and z-average diameters obtained for a series of PMAA₅₆-PMMA_{50–2000} nanoparticles prepared via RAFT aqueous emulsion polymerization of MMA at 70 °C when targeting 20% w/w solids at pH 5

Target PMMA DP	Conversion (%)	Actual PMMA DP	DLS Diameter (nm)	DLS PDI	TEM morphology
50	99	50	29	0.14	Spheres
100	99	99	35	0.13	Spheres
200	99	198	42	0.16	Spheres
300	99	297	54	0.14	Spheres
400	99	396	59	0.10	Spheres
500	99	495	73	0.11	Spheres
800	98	784	84	0.19	Spheres
1000	97	970	92	0.17	Spheres
1500	94	1410	94	0.20	Spheres
2000	95	1900	99	0.24	Spheres



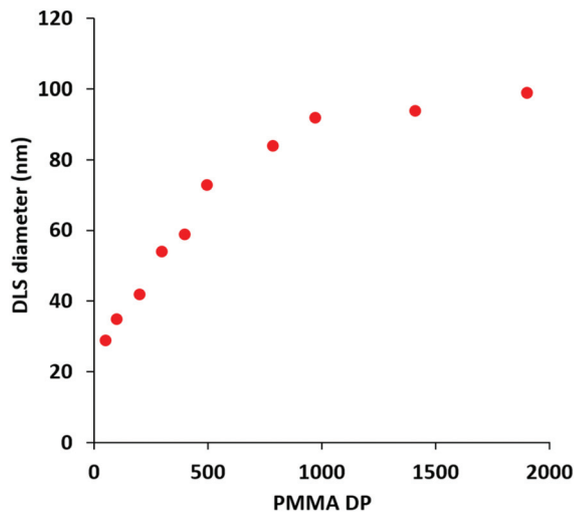


Fig. 5 Variation in z-average diameter with PMMA DP (x ; corrected for the final monomer conversion) for a series of PMAA₅₆-PMMA _{x} diblock copolymer nanoparticles prepared via RAFT aqueous emulsion polymerization of MMA at 70 °C targeting 20% w/w solids at pH 5.

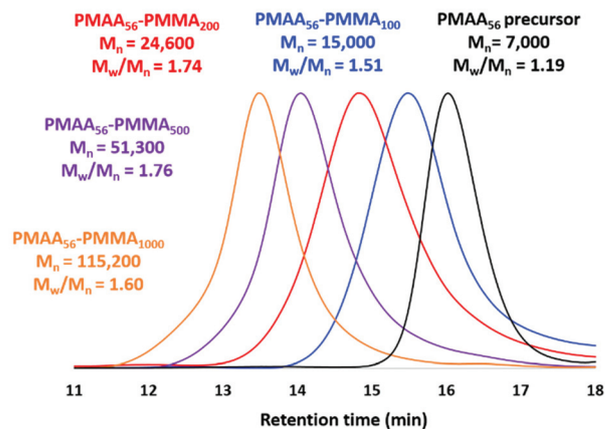


Fig. 7 THF GPC curves recorded for the methylated PMAA₅₆ precursor and a series of methylated PMAA₅₆-PMMA _{x} diblock copolymer nanoparticles for which the target PMMA DP (x) was 100, 200, 500 or 1000.

stabilizer comprising poly(2-(methacryloyloxy)ethyl trimethylammonium chloride) [PMETAC], see the ESI† for full synthesis details and also Fig. S2.† Thus, if relatively large PMMA-core nanoparticles are required for a given application, it is clear that polyelectrolytic stabilizers offer a decisive advantage over non-ionic stabilizers such as PGMA.

End-group removal from PGMA-PMMA nanoparticles using visible light irradiation

For many potential applications, it is desirable to remove the RAFT end-groups after the polymerization because these organosulfur groups confer color and malodour.⁵ For soluble copolymers, this is readily achieved using various chemistries.^{55–57} However, there are rather few studies of the removal of RAFT end-groups from diblock copolymer nanoparticles. In 2015, Destarac and co-workers demonstrated that xanthate groups could be efficiently cleaved from aqueous poly(*n*-butyl acrylate) latexes using ozone at ambient temperature.⁵⁸ Subsequently, Jesson *et al.*⁴⁶ reported that dithiobenzoate end-groups can be efficiently removed from PGMA₅₂-PHPMA₁₃₅ (where PHPMA denotes poly(2-hydroxypropyl methacrylate) diblock copolymer worms after treatment with H₂O₂ at 70 °C for 3 h using a H₂O₂/dithiobenzoate molar ratio of 5.0. However, removal of a trithiocarbonate end-group from essentially the same diblock copolymer proved to be much slower, with around 24% organosulfur groups remaining after 8 h under the same reaction conditions. Furthermore, the removal of dithiobenzoate end-groups from PGMA₆₁-PBzMA₁₀₀ (where PBzMA denotes poly(benzyl methacrylate) spheres using this H₂O₂ protocol was relatively ineffective, with UV GPC analysis indicating that more than 60% of the original end-groups remained intact after 8 h. This was attributed to the relatively hydrophobic nature of the core-forming PBzMA block, which is likely to impede ingress of the H₂O₂ into the nanoparticle cores. More recently, Gibson and co-workers⁵⁹ reported that visible light irradiation (blue LED; $\lambda = 405$ nm) at 50 °C removed dithiobenzoate end-groups from aqueous dispersions of PNMEP₂₈-

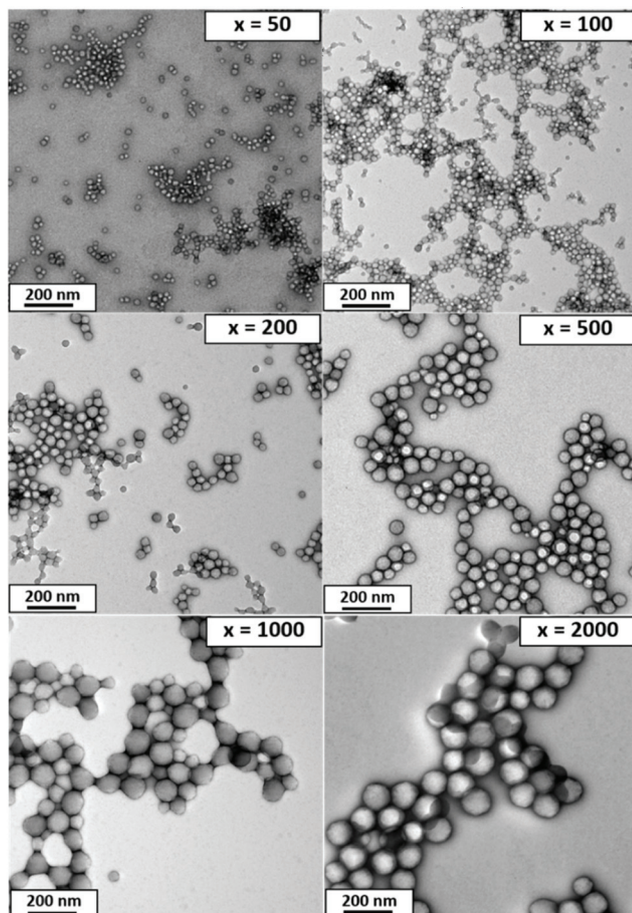


Fig. 6 TEM images recorded for a series of PMAA₅₆-PMMA _{x} diblock copolymer nanoparticles prepared by RAFT aqueous emulsion polymerization of MMA when targeting a PMMA DP (x) of 50, 100, 200, 500, 1000 or 2000 (see Table 2 for further details).



PLMA₈₇ [where PNMEP denotes poly(*N*-(2-methacryloyloxy) ethyl pyrrolidone) and PLMA denotes poly(lauryl methacrylate)] diblock copolymer vesicles. In principle, this approach is attractive because it should not suffer from the retarded diffusion of reagents observed by Jesson and co-workers.⁴⁶ Thus, we decided to study the removal of dithiobenzoate end-groups from 10% w/w aqueous dispersions of PGMA₅₀-PMMA₈₀ nanoparticles using similar visible light irradiation conditions as those reported by Gibson and co-workers.⁵⁹ Bearing in mind the aqueous solubilities of the respective monomers at room temperature, the hydrophobic character of the core-forming PMMA block is significantly greater than that of PHPMA but rather less than that of either PBzMA or PLMA. Moreover, it is perhaps worth emphasizing that PMMA has a significantly higher glass transition temperature (T_g) than these three alternative core-forming blocks and it is not obvious whether the glassy nature of the PMMA nanoparticle cores might impede removal of the RAFT end-groups.

In our initial RAFT end-group removal studies, visible light irradiation experiments were conducted at 70 °C. However, only rather slow and incomplete end-group cleavage (86% within 24 h; data not shown) was achieved under such conditions for PGMA₅₀-PMMA₈₀ nanoparticles prepared using CPDB. Fortunately, we found that significantly higher rates of end-group cleavage could be achieved at 80 °C. Accordingly, the kinetics of dithiobenzoate end-group removal by visible light irradiation (blue LED source; $\lambda = 405$ nm) of a 10% w/w aqueous dispersion of PGMA₅₀-PMMA₈₀ nanoparticles at 80 °C was monitored by periodic sampling of the reaction mixture followed by DMF GPC analysis using a UV detector set at 309 nm. Unlike visible absorption spectrophotometry, this technique enables the RAFT end-groups that remain attached to the copolymer chains to be distinguished from those that have been cleaved to produce small molecule by-products.⁴⁶ Representative UV GPC curves are shown in Fig. 8a and the fraction of remaining dithiobenzoate end-groups is plotted against time in Fig. 8b. Initially, relatively rapid cleavage occurs, with 87% of the original end-groups being removed within 12 h at 80 °C. After continuous irradiation for 24 h, 94% end-group removal can be achieved and the initial pink copolymer dispersion is converted into a colorless dispersion (see inset digital photographs). As a comparison, H₂O₂ was employed for oxidative end-group removal under the same conditions, as previously reported.⁴⁶ However, this only led to 24% end-group removal within 12 h and 58% after 24 h (Fig. 8b and S3†). This is presumably because this water-soluble reagent cannot readily diffuse into the glassy hydrophobic PMMA cores. The extent of end-group removal was also monitored for the PGMA₅₀-PMMA₈₀ nanoparticles at 80 °C in the absence of either visible light irradiation or H₂O₂. For this control experiment, UV GPC studies (Fig. S3†) indicated that 92% end-groups remained intact after 24 h, suggesting minimal thermally-induced hydrolysis.

Precisely the same protocol was adopted when studying trithiocarbonate end-group removal from a 10% w/w (aqueous dispersion of HOOC-PGMA₅₄-PMMA₈₀ nanoparticles. The GPC

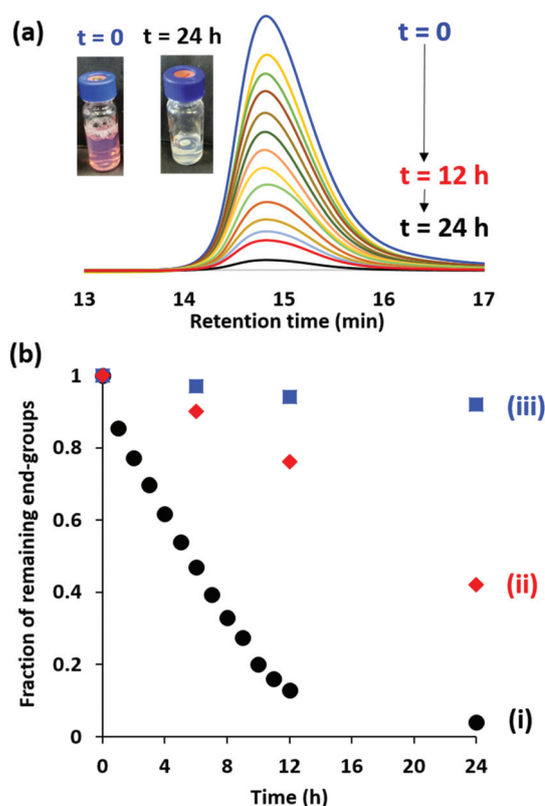


Fig. 8 (a) Representative UV GPC curves ($\lambda = 309$ nm) recorded during kinetic studies of the cleavage of dithiobenzoate end-groups for a 10% w/w aqueous dispersion of PGMA₅₀-PMMA₈₀ spherical nanoparticles at 80 °C. (b) Fraction of remaining dithiobenzoate RAFT end-groups over time determined by UV GPC analysis when a 10% w/w aqueous dispersion of PGMA₅₀-PMMA₈₀ spherical nanoparticles at 80 °C is exposed to: (i) continuous visible light irradiation ($\lambda = 405$ nm), (ii) H₂O₂ (using a H₂O₂/dithiobenzoate molar ratio of 5.0) and (iii) neither visible light nor H₂O₂ (control).

curves and kinetics are summarized in Fig. S4.† Cleavage of trithiocarbonate end-groups was achieved, albeit at a somewhat slower rate than that found for the dithiobenzoate end-groups. More specifically, 72% of the original trithiocarbonate groups were cleaved within 12 h at 80 °C with 87% being removed after 24 h. As a control experiment, the trithiocarbonate-capped HOOC-PGMA₅₄-PMMA₈₀ nanoparticles were exposed to H₂O₂ at 80 °C using a H₂O₂/trithiocarbonate molar ratio of 5.0 (see Fig. S4 and S5†). However, UV GPC studies indicated that more than 80% of the original RAFT end-groups remained intact within 24 h. Similarly, a control experiment conducted in the absence of either H₂O₂ or visible light irradiation indicated that 96% trithiocarbonate end-groups survived intact after 24 h at 80 °C.

Clearly, visible light irradiation can be a highly effective means of removing trithiocarbonate end-groups as well as dithiobenzoate end-groups. This approach works well for both high T_g core-forming blocks such as PMMA (as demonstrated herein) and low T_g core-forming blocks such as PLMA.⁵⁹ The only disadvantage appears to be the relatively long reaction



time required at 80 °C but presumably this could be reduced by increasing the intensity of the visible light source.

Conclusions

Sterically-stabilized diblock copolymer nanoparticles are prepared by RAFT aqueous emulsion polymerization of MMA using PGMA as a non-ionic steric stabilizer block. For a target PMMA DP of 20 to 100, kinetically-trapped spherical nanoparticles ranging in size from 17 nm to 31 nm are obtained. However, highly flocculated spherical nanoparticles are produced when targeting DPs above 100. Similar flocculation problems are encountered when employing a PGMA stabilizer block that possesses a terminal anionic carboxylate group. Moreover, this unexpected limitation does not appear to apply to various other RAFT aqueous emulsion polymerization formulations: core-forming block DPs of up to 1000 can be targeted without any loss of colloidal stability when employing alternative (and more hydrophobic) water-immiscible monomers such as benzyl methacrylate, 2,2,2-trifluoroethyl methacrylate or isopropylidenediglycerol monomethacrylate when using a non-ionic PGMA stabilizer block. This perplexing constraint appears to be related to the high T_g of the PMMA block, which exceeds the reaction temperature of 70 °C used for such PISA syntheses. However, we demonstrate that colloidal stable dispersions can be obtained when targeting PMMA DPs of up to 2000 using a highly anionic PMAA stabilizer block. Finally, visible light irradiation is used to cleave dithiobenzoate end-groups from a 10% w/w aqueous dispersion of PGMA₅₀-PMMA₈₀ nanoparticles. UV GPC studies indicated 87% end-group removal from such nanoparticles within 12 h at 80 °C. In contrast, using excess H₂O₂ only led to 24% end-group removal under the same conditions. This striking difference is attributed to the water-soluble reagent having restricted access to the hydrophobic PMMA nanoparticle cores. Furthermore, the same visible light irradiation protocol can be used to remove trithiocarbonate end-groups from HOOC-PGMA₅₀-PMMA₈₀ nanoparticles.

Conflicts of interest

There are no conflicts to declare.

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References

- J. Chiefari, Y. K. B. Chong, F. Ercole, J. Krstina, J. Jeffery, T. P. T. Le, R. T. A. Mayadunne, G. F. Meijs, C. L. Moad, G. Moad, E. Rizzardo and S. H. Thang, *Macromolecules*, 1998, **31**, 5559–5562.
- G. Moad, E. Rizzardo and S. H. Thang, *Aust. J. Chem.*, 2006, **59**, 669.
- G. Moad, E. Rizzardo and S. H. Thang, *Aust. J. Chem.*, 2009, **62**, 1402–1472.
- G. Moad, E. Rizzardo and S. H. Thang, *Aust. J. Chem.*, 2012, **65**, 985–1076.
- S. Perrier, *Macromolecules*, 2017, **50**, 7433–7447.
- D. J. Keddie, G. Moad, E. Rizzardo and S. H. Thang, *Macromolecules*, 2012, **45**, 5321–5342.
- J. D. Biasutti, T. P. Davis, F. P. Lucien and J. P. A. Heuts, *J. Polym. Sci., Part A: Polym. Chem.*, 2005, **43**, 2001–2012.
- W. Wan and C. Pan, *Polym. Chem.*, 2010, **1**, 1475–1484.
- Y. Li and S. P. Armes, *Angew. Chem., Int. Ed.*, 2010, **49**, 4042–4046.
- L. A. Fielding, M. J. Derry, V. Ladmiral, J. Rosselgong, A. M. Rodrigues, L. P. D. Ratcliffe and S. P. Armes, *Chem. Sci.*, 2013, **4**, 2081–2087.
- X. Zhang, J. Rieger and B. Charleux, *Polym. Chem.*, 2012, **3**, 1502–1509.
- O. J. Deane, O. M. Musa, A. Fernyhough and S. P. Armes, *Macromolecules*, 2020, **53**, 1422–1434.
- C. J. Ferguson, R. J. Hughes, B. T. T. Pham, B. S. Hawkett, R. G. Gilbert, A. K. Serelis and C. H. Such, *Macromolecules*, 2002, **35**, 9243–9245.
- C. J. Ferguson, R. J. Hughes, D. Nguyen, B. T. T. Pham, R. G. Gilbert, A. K. Serelis, C. H. Such and B. S. Hawkett, *Macromolecules*, 2005, **38**, 2191–2204.
- J. Rieger, F. Stoffelbach, C. Bui, D. Alaimo, C. Jérôme and B. Charleux, *Macromolecules*, 2008, **41**, 4065–4068.
- B. Charleux, G. Delaittre, J. Rieger and F. D'Agosto, *Macromolecules*, 2012, **45**, 6753–6765.
- S. L. Canning, G. N. Smith and S. P. Armes, *Macromolecules*, 2016, **49**, 1985–2001.
- X. Zhang, S. Boissé, W. Zhang, P. Beaunier, F. D'Agosto, J. Rieger and B. Charleux, *Macromolecules*, 2011, **44**, 4149–4158.
- S. Boissé, J. Rieger, G. Pembouong, P. Beaunier and B. Charleux, *J. Polym. Sci., Part A: Polym. Chem.*, 2011, **49**, 3346–3354.
- W. Zhang, F. D'Agosto, O. Boyron, J. Rieger and B. Charleux, *Macromolecules*, 2012, **45**, 4075–4084.
- L. Carlsson, A. Fall, I. Chaduc, L. Wågberg, B. Charleux, E. Malmström, F. D'Agosto, M. Lansalot and A. Carlmark, *Polym. Chem.*, 2014, **5**, 6076–6086.
- W. Zhang, B. Charleux and P. Cassagnau, *Macromolecules*, 2012, **45**, 5273–5280.
- B. T. T. Pham, D. Nguyen, V. T. Huynh, E. H. Pan, B. Shirodkar-Robinson, M. Carey, A. K. Serelis, G. G. Warr, T. Davey, C. H. Such and B. S. Hawkett, *Langmuir*, 2018, **34**, 4255–4263.
- R. Albigès, P. Klein, S. Roi, F. Stoffelbach, C. Creton, L. Bouteiller and J. Rieger, *Polym. Chem.*, 2017, **8**, 4992–4995.
- N. P. Truong, M. V. Dussert, M. R. Whittaker, J. F. Quinn and T. P. Davis, *Polym. Chem.*, 2015, **6**, 3865–3874.



- 26 W. Zhao, G. Gody, S. Dong, P. B. Zetterlund and S. Perrier, *Polym. Chem.*, 2014, **5**, 6990–7003.
- 27 I. Chaduc, O. Boyron, B. Charleux, F. D. Agosto and M. Lansalot, *Macromolecules*, 2013, **46**, 6013–6023.
- 28 J. Rieger, G. Osterwinter, C. Bui and B. Charleux, *Macromolecules*, 2009, **42**, 5518–5525.
- 29 I. Chaduc, M. Girod, R. Antoine, B. Charleux, F. D'Agosto and M. Lansalot, *Macromolecules*, 2012, **45**, 5881–5893.
- 30 M. Chenal, C. Véchambre, J. M. Chenal, L. Chazeau, V. Humblot, L. Bouteiller, C. Creton and J. Rieger, *Polymer*, 2017, **109**, 187–196.
- 31 S. Binauld, L. Delafresnaye, B. Charleux, F. Dagosto and M. Lansalot, *Macromolecules*, 2014, **47**, 3461–3472.
- 32 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.
- 33 B. Akpınar, L. A. Fielding, V. J. Cunningham, Y. Ning, O. O. Mykhaylyk, P. W. Fowler and S. P. Armes, *Macromolecules*, 2016, **49**, 5160–5171.
- 34 C. P. Jesson, V. J. Cunningham, M. J. Smallridge and S. P. Armes, *Macromolecules*, 2018, **51**, 3221–3232.
- 35 E. E. Brotherton, F. L. Hatton, A. A. Cockram, M. J. Derry, A. Czajka, E. J. Cornel, P. D. Topham, O. O. Mykhaylyk and S. P. Armes, *J. Am. Chem. Soc.*, 2019, **141**, 13664–13675.
- 36 F. L. Hatton, J. R. Lovett and S. P. Armes, *Polym. Chem.*, 2017, **8**, 4856–4868.
- 37 J. Tan, D. Liu, C. Huang, X. Li, J. He, Q. Xu and L. Zhang, *Macromol. Rapid Commun.*, 2017, **38**, 1–7.
- 38 F. L. Hatton, A. M. Park, Y. Zhang, G. D. Fuchs, C. K. Ober and S. P. Armes, *Polym. Chem.*, 2019, **10**, 194–200.
- 39 F. L. Hatton, M. J. Derry and S. P. Armes, *Polym. Chem.*, 2020, **11**, 6343–6355.
- 40 A. A. Cockram, T. J. Neal, M. J. Derry, O. O. Mykhaylyk, N. S. J. Williams, M. W. Murray, S. N. Emmett and S. P. Armes, *Macromolecules*, 2017, **50**, 796–802.
- 41 W. Zhang, F. D'Agosto, P. Y. Dugas, J. Rieger and B. Charleux, *Polymer*, 2013, **54**, 2011–2019.
- 42 F. L. Hatton, M. Ruda, M. Lansalot, F. D. Agosto, E. Malmstro and A. Carlmark, *Biomacromolecules*, 2016, **17**, 1414–1424.
- 43 J. Engström, H. Asem, H. Brismar, Y. Zhang, M. Malkoch and E. Malmström, *Macromol. Chem. Phys.*, 2020, **221**, 1900443.
- 44 S. J. Hunter, J. R. Lovett, O. O. Mykhaylyk, E. R. Jones and S. P. Armes, *Polym. Chem.*, 2021, **12**, 3629–3639.
- 45 M. Semsarilar, V. Ladmira, A. Blanazs and S. P. Armes, *Langmuir*, 2013, **29**, 7416–7424.
- 46 C. P. Jesson, C. M. Pearce, H. Simon, A. Werner, V. J. Cunningham, J. R. Lovett, M. J. Smallridge, N. J. Warren and S. P. Armes, *Macromolecules*, 2017, **50**, 182–191.
- 47 J. Ilavsky and P. R. Jemian, *J. Appl. Crystallogr.*, 2009, **42**, 347–353.
- 48 A. Guinier and G. Fournet, *Small-Angle Scattering of X-rays*, John Wiley & Sons, New York, 1955.
- 49 O. Glatter and O. Kratky, *Small-angle X-ray Scattering*, Academic Press, London, 1982.
- 50 J. S. Pedersen, *J. Appl. Crystallogr.*, 2000, **33**, 637–640.
- 51 G. Beaucage and D. W. Schaefer, *J. Non.-Cryst. Solids*, 1994, **172–174**, 797–805.
- 52 G. Beaucage, *J. Appl. Crystallogr.*, 1995, **28**, 717–728.
- 53 G. Beaucage, *J. Appl. Crystallogr.*, 1996, **29**, 134–146.
- 54 J. R. Lovett, N. J. Warren, L. P. D. Ratcliffe, M. K. Kocik and S. P. Armes, *Angew. Chem., Int. Ed.*, 2015, **54**, 1279–1283.
- 55 S. Perrier and P. Takolpuckdee, *J. Polym. Sci., Part A: Polym. Chem.*, 2005, **43**, 5347–5393.
- 56 H. Willcock and R. K. O'Reilly, *Polym. Chem.*, 2010, **1**, 149–157.
- 57 B. Quiclet-Sire and S. Z. Zard, *Bull. Korean Chem. Soc.*, 2010, **31**, 543–544.
- 58 D. Matioszek, P. E. Dufils, J. Vinas and M. Destarac, *Macromol. Rapid Commun.*, 2015, **36**, 1354–1361.
- 59 R. R. Gibson, E. J. Cornel, O. M. Musa, A. Fernyhough and S. P. Armes, *Polym. Chem.*, 2020, **11**, 1785–1796.

