Tricomponent thermoresponsive polymers based on an amine-containing monomer with tuneable hydrophobicity:
Effect of composition

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Abstract

In the present study, six dual-responsive ABC triblock copolymers were synthesised via group transfer polymerisation (GTP) and investigated through visual inspections in terms of their thermoresponsive behaviour. The copolymers consist of i) penta(ethylene glycol) methyl ether methacrylate (PEGMA), which is hydrophilic and thermoresponsive at high temperatures, ii) n-butyl methacrylate (BuMA) as the hydrophobic counterpart to promote self-assembly, and iii) 2-(diethylamino)ethyl methacrylate (DEAEMA), which is pH-responsive by adjusting its hydrophilicity depending on the pH. The effect of the degree of ionisation of DEAEMA units as well as the ionic strength effect on the self-assembly behaviour of the copolymers was tested via dynamic light scattering (DLS). The dissociation constants (pK_a) of the amine units of DEAEMA were determined via potentiometric titrations. The thermoresponse has been primarily been investigated in means of cloud points (CPs) at various pH values in deionised water. Detailed phase diagrams were constructed for all the polymer solutions in phosphate buffered saline (PBS), with the interest being focused on the gelation area. It has been clearly proven that gelation is promoted as the content in BuMA and DEAEMA is increased. The polymer that presented the widest gelation area has been further investigated via rheology in terms of its gelation temperature, gelation time and shear-thinning properties.

Keywords: thermoresponsive polymers, 2-(diethylamino)ethyl methacrylate, penta(ethylene glycol) methyl ether methacrylate, block terpolymers, thermoresponsive gels, group transfer polymerisation (GTP)

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Graphical Abstract:
Introduction

Thermoresponsive polymers are widely studied by scientists around the world, as they do present interesting changes in their properties depending on the temperature.[1-5] Thermoresponsive polymers can either be chemically crosslinked, thus forming thermoresponsive polymeric networks (also known as chemically crosslinked gels), or can be studied without further chemical crosslinking.[3] In both cases, they are categorized in two groups, depending on how their compatibility with the solvent varies with the temperature.[1] Thermoresponsive polymers or polymeric networks which are more soluble or compatible with a solvent as the temperature increases, present upper critical solution temperature (UCST).[1, 4, 6] A well-known example of a UCST thermoresponsive polymer is gelatin.[7] On the contrary, when the solubility/compatibility with the solvent decreases as the temperature increases, they present lower critical solution temperature (LCST).[1] LCST polymers with appropriate structural characteristics can form gels via physical interactions as the temperature increases.[2]

These thermoresponsive physical gels have attracted much scientific interest because of their potential use in tissue engineering as injectable gels.[2, 8-10] In this concept, the polymer solution at room temperature is loaded with cells, and it is easily transferred to a syringe. A physical gel is formed post-injection, which is triggered by the change of the environmental temperature to body temperature. This gel matrix that contains the cells acts as a scaffold, in which the cells will form the new tissue. Ideally, the rate of tissue growth should match the rate of polymer dissolution/ degradation. The use of injectable gels in the field of tissue engineering is advantageous as surgery to implant a scaffold can be avoided.[2, 8-10]

Thermoresponsive gels have also been studied as drug delivery systems.[11-15] It has been proven in several studies that incorporating the drug in the gel matrix is beneficial, as it controls the drug release rate for prolonged periods. Also, topical drug delivery, as opposed to systemic drug delivery, is advantageous as it minimises the side effects.[11-15]

More recently, thermoresponsive gels have been investigated as 3-D printable materials.[2, 16-26] The use of thermoresponsive gels in 3-D printing facilitates the easy loading into the syringe of the printer, which is then heated to the desired temperature to promote gelation. In 3-D printing, thermoresponsive gels are used as sacrificial materials, either to facilitate the printing of another material, e.g. graphene,[19] or to print more complicated structures. In the first case, the thermoresponsive gel is mixed with the material in interest, while in the second case, the thermoresponsive gel is printed within the permanent structure. In both cases, the temperature of the printed construct is lowered after printing, and thus the physical gel returns to the solution phase thus a structure consisting of the permanent material remains.[2, 16-26] In addition to its thermoresponsive properties, and thus its reversible gelation properties, the ink should possess shear-thinning properties, i.e. the viscosity should decrease with increased shear rate, and thus the easier the printing process is.[27-29] Therefore, as thermoresponsive gels are widely studied not
only by polymer chemists but also by material scientists and (bio-)engineers, there is an increase of interest in fabrication of new thermoresponsive polymers.

Several thermoresponsive polymers have been reported in the literature. Among the most popular thermoresponsive units, which have been investigated either as homopolymers or co-polymers are N-isopropylacrylamide (NIPAAm),[30-33] ethylene glycol (EG),[34, 35] 2-(dimethylamin) ethyl methacrylate (DMAEMA),[36-42] various EG-based (meth)acrylates, [33, 43-49] and poly(oxazolines)[50-55]. The most well-documented thermoresponsive polymer is Poloxamer 407 (also known with its trademark name Pluronic® F127), which is based on EG and propylene glycol (PG).[22, 56-63]

The synthesis of new thermoresponsive polymers by polymer chemists can be facilitated by various polymerisation techniques, either anionic or radical polymerisations. In either case, well-defined polymers (narrow dispersity) should be synthesised, as it is well-established that polymers with different molar mass (MM), and composition present different thermoresponse.[2] Therefore, “living” or controlled polymerisation methods should be implemented. In the current study, group transfer polymerisation (GTP) has been chosen as the fabrication method, as it fulfils the aforementioned criterion, and also it is time-effective, cost-effective and industrially applicable.[64-66] More specifically, GTP has been invented in DuPont in 1980’s, and it is completed within 15 mins (per block) and at room temperature. The monomer conversion is 100%, thus no intermediate purification steps are required.[64-66]

Our group has previously published several research papers on thermoresponsive polymers.[37-42, 67] More specifically, our studies have been focused on amphiphilic block copolymers (mostly diblock and triblock copolymers, or tetrablock copolymers), and clear trends have been established in each study. These studies have shown that each structural parameter, i.e. architecture,[37, 39, 41, 42] composition[38-40, 42, 67] and MM,[38] is decisive for the final thermoresponsive properties. In these studies, various monomers have been used, such as poly(ethylene glycol) (PEG)-based[67] and alkyl-based methacrylate monomers of various lengths,[37] as the hydrophilic and hydrophobic units, respectively. In all the studies, a hydrophilic and pH- and thermoresponsive amine-containing monomer was incorporated into the structure, namely DMAEMA.[37-42, 67] DMAEMA has been thoroughly investigated in the past for its thermoresponsive and pH-responsive properties, and its cloud point (CP) and dissociation constant (pKₐ) are well documented in the literature.[36, 68-70] More specifically, the pKₐ of the conjugate acid form of the DMAEMA amine is around 7.0, while its CP decreases from 46°C to 32°C, as the MM increases from 1450 g mol⁻¹ to 53000 g mol⁻¹.[36] In most of the studies, penta(ethylene glycol) methyl ether methacrylate (PEGMA) was used as the hydrophilic unit, which is also thermoresponsive but at higher temperatures than DMAEMA; it has been reported that its CP is 64°C, MM dependent.[71] Concerning the length of the alkyl group, it has been concluded that polymers containing n-butyl methacrylate (BuMA) form stronger gels.[37] For the systems that have been investigated by our group, it has been observed that the triblock copolymers with i) the hydrophobic block in the centre of the polymer chain,[39, 41] ii) target MM around 7600 to 8300 g mol⁻¹,[38] and iii) hydrophobic BuMA content between 30 w/w% and 40 w/w%.[39, 67] show clear sol-
gel transition, i.e. they are soluble at low temperatures and form a gel at higher temperatures, without presenting any solubility issues.

In our latest study, block copolymers based on PEGMA and the pH-responsive 2-(diethylamino)ethyl methacrylate (DEAEMA) have been investigated for their use as precursors for fabrication of gold nanoparticles (AuNPs).[72] DEAEMA is known to be pH-responsive, as its hydrophilicity changes depending on the pH.[36, 72-76] More specifically, at alkaline environment (pH higher than its $pK_a \sim 7.3$), DEAEMA homopolymer is hydrophobic, and thus water-insoluble.[36, 74, 76] On the other hand, at acidic conditions (pH lower than its $pK_a$), the amino groups are protonated and hydrophilic and thus the DEAEMA homopolymer is soluble in water.[36, 74, 76] When DEAEMA is compared to DMAEMA, DEAEMA is a stronger base, indicated by its higher $pK_a$ value (weaker conjugate acid).[36]

In the present study, we wanted to investigate the effect of replacing the DMAEMA units with DEAEMA units will affect the solution properties of these methacrylate based triblock copolymers. Thus, to study how the incorporation of the pH responsive DEAEMA units, which are hydrophobic at alkaline pH and hydrophilic at acidic pH, affect the thermoresponsive properties, as it is well established that the hydrophobicity affects the thermoresponsive/gelation properties. Therefore, we investigate in more detail the thermoresponsive behaviour of dual-responsive ABC triblock copolymers based on PEGMA (A unit), BuMA (B unit), and DEAEMA (C unit). Six triblock copolymers of various compositions were synthesised via GTP. The hydrophobic BuMA content was kept constant either at 10 w/w% (Family A), or at 20 w/w% (Family B). Since DEAEMA units are more hydrophobic than DMAEMA units, the hydrophobic content was kept at lower values than the optimum hydrophobic content determined for DMAEMA-containing polymers to ensure that the polymers are water-soluble. In the first family, three ABC compositions were targeted: i) 50-10-40 w/w%, ii) 45-10-45 w/w%, and iii) 40-10-50 w/w%, whereas in the second family, the compositions were: i) 55-20-25 w/w%, 50-20-30 w/w%, and 45-20-35 w/w%. In the present study, all the copolymers were extensively investigated through visual tests in terms of their thermoresponsive behaviour.

Experimental
Materials and Methods

The monomers, PEGMA (MM = 300 g mol⁻¹), BuMA (99%), and DEAEMA (99%), the initiator, methyl trimethylsilyl dimethylketene acetal (MTS, 95%), the purifying agents, aluminium oxide activated basic (Al₂O₃·KOH), and calcium hydride (CaH₂, ≥90%), the free-radical inhibitor, 2,2-diphenyl-1-picrylhydrazyl hydrate (DPPH), the polymerisation solvent, tetrahydrofuran (THF, HPLC grade, ≥99.9%), and sodium and potassium metals were purchased from Sigma Aldrich. The precipitation solvent, n-hexane, was purchased from VWR Chemicals, while phosphate buffered saline (PBS, 10x solution) was purchased from Fisher Scientific. Tetrabutylammonium hydroxide (40% in water), which was used for the catalyst synthesis, as it is discussed in the following paragraphs, was acquired from Acros Organics.

The monomers were purified by i) passing twice through basic aluminium oxide, and ii) stirring under CaH₂ for at least 3h. These steps were followed to ensure an acid-free environment, e.g. removal of the polymerisation inhibitor and methacrylic acid, and ii) a humidity-free environment. DPPH has been added in the low MM monomers, BuMA and DEAEMA, to prevent undesired polymerisation and these monomers were vacuum-distilled prior to use, whereas the high MM monomer PEGMA was diluted in THF (50 v/v%) prior to any purification step and was syringed directly in the polymerisation flask using 0.45μm PTFE filters. The initiator was vacuum-distilled prior to use, while the polymerisation solvent was dried by refluxing for 3 days in sodium-potassium alloy. All the chemicals were kept under inert argon conditions. The catalyst was synthesised by tetrabutylammonium hydroxide and benzoic acid, as reported previously,[77] it was recrystallised in diethyl ether and it was vacuum-dried prior to use. All the glassware used during the polymerisation and distillation were dried overnight at 140°C prior to use. The chemical structures of the monomers used in this study, the initiator and the catalyst are presented in Figure 1.
**Figure 1:** Chemical structures, names and abbreviations of the monomers, the initiator and the catalyst used in this study.

**Triblock Copolymer Synthesis and Recovery**

All the triblock copolymers were synthesised via sequential GTP, in a similar procedure as reported in the literature.\cite{64} The synthesis of Polymer 1, PEGMA\textsubscript{12.5}-b-BuMA\textsubscript{5}-b-DEAEMA\textsubscript{16}, is reported as an example. Specifically, 10mg of TBABB was added in a 250 mL round-bottom flask, which was sealed immediately with rubber septum and purged with argon to ensure inert atmosphere. Freshly distilled THF (37.9 mL) was injected in the polymerisation flask, followed by the injection of the initiator (0.4 mL, 0.3 g, 2.0 mmol). Then, the monomer additions followed as described: i) PEGMA solution (14.1 mL, 7.34 g, 25.0 mmol), ii) BuMA (1.7 mL, 1.5 g, 10 mmol), and iii) DEAEMA (6.4 mL, 5.9 g, 32 mmol). It should be mentioned that each monomer injection was performed every 15 mins, which is the time interval that ensures that the polymerisation is complete. The temperature variation was monitored after each addition. Two 0.1 mL aliquots were withdrawn after the completion of each block polymerisation for GPC and \textsuperscript{1}H NMR analysis. The other triblock copolymers were synthesised in a similar manner. In all the cases, the polymerisation yield was quantitative (i.e. 100\% yield), which is a characteristic of GTP. All the polymers were recovered by precipitation in cold \textit{n}-hexane and the traces of \textit{n}-hexane and THF were removed by drying in a vacuum oven at room temperature.
Characterisation in Organic Solvents

Gel Permeation Chromatography (GPC)

GPC was utilised to determine the MM and MMDs of the polymers and their linear precursors. The SECurity GPC system was purchased from Agilent, and it is equipped with a Polymer Standard Service (PSS) SDV analytical linear M column (SDA083005LIM), an Agilent 1260 refractive index (RI) detector, and a “1260 Iso” isocratic pump. The GPC solvent was THF/Et$_3$N solution (95/5 v/v%). Prior to the measurements, the GPC system was calibrated using six poly(methyl methacrylate) standard samples of well-defined MMs: 2000, 4000, 8000, 20000, 50000, and 100000 g mol$^{-1}$.

Proton Nuclear Magnetic Resonance (¹H NMR) Spectroscopy

An Avance Bruker 400 MHz NMR system was used to determine the compositions of the copolymers and their precursors (homopolymers and diblock copolymers). CDCl$_3$ was used as the NMR solvent.

Characterisation in Aqueous Solvents

Potentiometric titrations

Hydrogen ion titrations were performed on 1 w/w% polymer solutions in DI water to determine the pK$_a$ values of DEAEMA units. For this experiment, a portable HI98103 pH checker was used, which was calibrated at pH 4 and pH 7 prior to each experiment. The pK$_a$ is determined as the pH at which 50 mol% of the DEAEMA are protonated.

Dynamic Light Scattering (DLS)

Dynamic Light Scattering (DLS) was performed in all polymer solutions at a concentration of 1 w/w% in either DI water or PBS. A Zetasizer Nano ZSP instrument was utilised for this experiment. The DLS analysis in DI water was performed at three different pH values: i) initial pH, which varied between 7.1-7.6, ii) acidic pH (pH 5), and alkaline pH (pH 8). Prior to the measurement, the samples were filtered using nylon 0.45 μm for removing any dust and big aggregates. The experiments were conducted at room temperature and the results were obtained by collected the light at a backscatter angle of 173°. The results are reported as the average of the size corresponding to the maximum by intensity. In case more than one peak were detected, the results reported are the ones corresponding to the peak of micelles.

The experimental diameters were compared to the theoretical ones. Two different models were used for calculating the diameters of the triblock copolymers. According to the first model, the cores of the micelles were formed only by the hydrophobic BuMA blocks, while the corona of the micelle formed by both PEGMA blocks and DEAEMA blocks. This model assumes that DEAEMA is fully hydrophilic, which is the case at low pH values (below the pK$_a$ value). The equation used to calculate the hydrodynamic diameter is $d = [\text{DP}_{\text{BuMA}} + 2*\text{DP}_{\text{longest hydrophilic}}]*0.254$ nm; where DP stands for degree
of polymerisation. The second model assumes that both BuMA and DEAEMA blocks are hydrophobic, thus they contribute to the formation of the cores of the micelles, while the corona of the micelle was formed by the hydrophilic PEGMA blocks only; DEAEMA is considered hydrophobic at high pH values, at which the amino groups are not protonated (above the pKₐ value). The equation used to calculate the hydrodynamic diameter is \( d = \left( [D_{BuMA} + D_{DEAEMA}] + 2*D_{PEGMA} \right)*0.254 \) nm. The DPs used for these calculations are the experimental ones calculated by using the GPC results after precipitation and the \(^1\)H NMR results before precipitation.

**Visual Tests**

The polymer solutions were visually tested from 20°C to 80°C at various concentrations and aqueous solvents. The experiment was conducted using an IKA RCT stirrer hotplate and an IKA ETS-D5 temperature controller. The vials were immersed in a continuously stirred water-bath and observations were obtained every degree. 1 w/w% polymer solutions in deionised (DI) water, both at initial pH (pH varying between 7.1 – 7.5) and at pH 8, were investigated for a CP (temperature at which the transparent solutions turn to cloudy). The polymer solutions in phosphate buffered saline (PBS) were investigated at various concentration for gelation. More specifically, 1, 2, 5, 10, 15, 20, 25 and 30 w/w% of the polymer solutions were tested and detailed phase diagrams were constructed. Nine transitions are stated in the phase diagrams: i) solution phase (clear, slightly cloudy, and cloudy), ii) viscous solution phase (transparent and cloudy), iii) gel phase (transparent and cloudy), and iv) two-phases system (gel syneresis and precipitation). Detailed pictures that define each state are reported in our previous publication in Constantinou et al. 2017.[41]

**Rheology**

The best-performing polymer of this study was tested rheologically at 30 w/w% in PBS, as at which concentration presented the widest gelation area. The analysis was performed using a TA discovery HR-1 hybrid rheometer, which was equipped with a 40 mm parallel Peltier steel plate (996921). Firstly, a temperature ramp measurement was performed from 40°C to 60°C, as this was the area of interest according to the visual tests. Secondly, after determining the gelation temperature rheologically, the gelation time was tested at 49°C. During both measurements, the strain was kept constant at 1%, while the angular frequency applied was at 1 rads⁻¹. The gel was investigated further for its shear-thinning properties from 0.01 s⁻¹ to 100 s⁻¹. As all the measurements were performed at temperatures higher than room temperature, a solvent trap was used to prevent solvent evaporation.
Results and Discussion

This study investigates the synthesis of six novel ABC triblock copolymers via GTP. The A, B and C blocks consisted of PEGMA, BuMA and DEAEMA units, respectively. The MM of the copolymers was kept constant at 7600 g mol$^{-1}$. This value is within the optimum MM range proven determined by our previous study on DMAEMA-based polymer for producing copolymers with clear-sol gel transition.[38] The composition of the triblock copolymers was varied to investigate its effect on the thermoresponsive properties of the copolymers. More specifically, two families, each consisting of three copolymers, with different BuMA content were synthesised. The six different PEGMA-BuMA-DEAEMA compositions targeted are: i) first family with 10 w/w% BuMA content (50-10-40, 45-10-45, and 40-10-50 w/w%), and ii) second family with 20 w/w% BuMA content (55-20-25, 50-20-30, and 45-20-35 w/w%). The BuMA content in this study was targeted at lower values than the optimum range proven in the previous studies, in which the polymers consisted of DMAEMA instead of DEAEMA,[37-42, 67] to ensure solubility (DEAEMA is more hydrophobic than DMAEMA). The polymer structures studied in the current study are schematically illustrated in Figure 2, in which the PEGMA, BuMA, and DEAEMA units are represented by light blue, red, and yellow spheres, respectively.

![Figure 2: Schematic of structures of polymers investigated in this study. PEGMA, BuMA, and DEAEMA units are represented by light blue, red, and yellow spheres, respectively.](image)

Structural Properties

The synthesis was monitored via GPC and $^1$H NMR spectroscopy. The molar mass characteristics (resulted from GPC) of the polymers and their linear precursors, and the compositions (resulted from $^1$H NMR) are summarised in Table 1. Table 1 also shows the theoretical MM and composition values.
**Table 1**: Theoretical polymer structures, theoretical molar masses, experimental number-average molar masses and dispersity indices, and theoretical and experimental compositions of the terpolymers and their linear precursors.

<table>
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<th>MM theor. (g mol⁻¹)</th>
<th>Mₙ (g mol⁻¹)</th>
<th>D c</th>
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a PEGMA, BuMA and DEAEMA stand for penta(ethylene glycol) methyl ether methacrylate, n-butyl methacrylate and 2-(diethylamino)ethyl methacrylate, respectively.

b The theoretical molar mass (MM) is calculated by summing the MM of PEGMA, BuMA, and DEAEMA parts, and 100 g mol⁻¹ is added to the final result, as a part of the initiator stays on the polymer chain. The MM of each part is calculated by multiplying the degree of polymerisation (DP) by the MM of the corresponding repeated unit.

c The Mₙ and D values resulted after GPC analysis, which was based on linear calibration curve with 6 poly(methyl methacrylate) (PMMA) standard samples of MM equal to 2, 4, 8, 20, 50, and 100 kDa.

**Molar Mass characteristics**

As can be seen in Table 2, the experimental MM values (number-average MM, Mₙ) vary between 8900 and 11800 g mol⁻¹. These values are higher than the theoretical ones (7600 g mol⁻¹), but still within the desirable range which produces polymers with clear sol-gel transition. The discrepancy of the theoretical and experimental MM values is attributed to the deactivation of the initiator (MTS) molecules because of the presence of humidity and impurities. This effect has been previously observed in GTP-studies on PEG macromonomers, i.e. monomers with average molar mass, that cannot be distilled because of their high MM.[42, 78]

The D values of the final terpolymers are between 1.13 and 1.21. These values prove that the GTP was successful and well-defined polymers consisting of PEG macromonomers were produced, and they are in agreement with previous results published on GTP synthesis.[42, 78] It is noteworthy that the triblock copolymers with
higher PEGMA content present higher $D$ values, which shows that the dispersity results from the use of macromonomer, i.e. a monomer with average molar mass, and not well-defined monomers.

The GPC traces of PEGMA$_{12.5}$-$b$-BuMA$_5$-$b$-DEAEMA$_{16}$ and its precursors (i.e. PEGMA$_{12.5}$ and PEGMA$_{12.5}$-$b$-BuMA$_5$) are presented in Figure 3. More specifically, the GPC traces of the triblock terpolymer before and after precipitation are shown in yellow solid and dashed lines, whereas light blue and red show the MM profiles of the homopolymer and diblock copolymer, respectively. As can be observed, the peak moves to higher MM values as the polymerisation progresses. Also, the GPC profiles are of narrow distribution, which proves the successful “living” GTP. The traces of shoulder present on the GPC traces of the final terpolymer are associated with the PEGMA homopolymer, which slightly deactivated the initiator, as observed in previous studies.[37, 41, 42, 67, 72] The GPC traces of the syntheses of the other triblock terpolymers are presented in Figure S1 in the Supporting Information and they show similar profiles.

![Figure 3: GPC traces showing synthetic steps of Polymer 1, PEGMA$_{12.5}$-$b$-BuMA$_5$-$b$-DEAEMA$_{16}$. Its MM profile before and after precipitation are shown in yellow solid and dashed lines. The GPC traces of its precursors are shown in blue (PEGMA$_{12.5}$) and red (PEGMA$_{12.5}$-$b$-BuMA$_5$).](image)

**Polymer Composition**

The compositions of the final terpolymers, homopolymer precursors and diblock copolymer precursors were analysed via $^1$H NMR and an example of the NMR analysis of the synthesis of Polymer 1 is presented in Figure S2 in the Supporting Information. The calculations of the experimental values are based on three distinct peaks, each corresponding to a different repeated unit. In particular, the peak of the three methoxy protons in PEGMA units ($\text{CH}_3\text{O}$-) appears at 3.35 ppm, while the distinct peak of the two methyl protons on the second carbon from the ester bond ($\text{-OCH}_2\text{CH}_2\text{CH}_3$) appear in 1.5-1.6 ppm. Concerning the DEAEMA unit, two distinct peaks appear in the
area between 2-4 and 2-7 ppm which correspond to the protons closer to the nitrogen
atom. For the calculations, the second peak (higher ppm) was used, and it belongs to
the four protons of the final ethyl groups [-OCH$_2$CH$_2$N(CH$_3$)$_2$]. The experimental
composition, listed in Table 1, are in good agreement with the theoretical/targeted
values, within the experimental error of the NMR technique.

Properties in Aqueous Solutions

The self-assembly behaviour in various aqueous solvents and pH values, the strength
of the amine groups, as well as the thermoresponsive behaviour of the terpolymers in
aqueous solutions have been tested and are discussed in the following sections. The
results are summarised in Table 2.

Table 2: Theoretical polymer structures, theoretical diameters based on two different models
and experimental hydrodynamic diameters in DI water at various pH and in PBS, dissociation
constants, and cloud points in DI water.

<table>
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<th>No</th>
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<th>$^{1}$ pH initial $^f$</th>
<th>$^{2}$ pH 8</th>
<th>$^{3}$ pH 5</th>
<th>PBS</th>
<th>$^{pK_a} \pm 0.1$</th>
<th>$^{CP} \pm 2 (°C) ^g$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>PEGMA$_{12.5}$-b-BuMA$<em>5$-b-DEAEMA$</em>{16}$</td>
<td>14.3 17.4 21.0 28.2 18.2 24.4</td>
<td>6.8 61</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>PEGMA$_{11.1}$-b-BuMA$<em>5$-b-DEAEMA$</em>{18}$</td>
<td>16.9 18.8 24.4 32.7 21.0 28.2</td>
<td>7.1 66</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>PEGMA$_{10.4}$-b-BuMA$<em>5$-b-DEAEMA$</em>{20}$</td>
<td>16.4 15.5 18.2 28.2 18.2 24.4</td>
<td>7.1 68</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>4</td>
<td>PEGMA$<em>{14.4}$-b-BuMA$</em>{10.5}$-b-DEAEMA$_{10}$</td>
<td>15.0 18.9 43.8 32.7 15.7 32.7</td>
<td>6.2 62</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>5</td>
<td>PEGMA$<em>{12.1}$-b-BuMA$</em>{10.5}$-b-DEAEMA$_{12}$</td>
<td>12.7 16.9 24.4 32.7 13.5 28.2</td>
<td>6.4 61</td>
<td></td>
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<tr>
<td>6</td>
<td>PEGMA$<em>{12.1}$-b-BuMA$</em>{10.5}$-b-DEAEMA$_{14}$</td>
<td>12.0 15.5 24.4 15.7 24.4 21.0</td>
<td>6.7 53</td>
<td></td>
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</table>

$^a$ PEGMA, BuMA and DEAEMA are the abbreviations for penta(ethylene glycol) methyl ether
methacrylate, n-butyl methacrylate, and 2-(diethylamino)ethyl methacrylate, respectively.

$^b$ The theoretical calculations were based on the experimental degrees of polymerisations
(DPs), as resulted from GPC analysis after precipitation and $^1$H NMR analysis before
precipitation.

$^c$ In model 1, the cores of the micelles were formed only by the hydrophobic BuMA blocks,
while the corona of the micelle formed by both PEGMA blocks and DEAEMA blocks. The
equation used to calculate the hydrodynamic diameter is $d_H = [DP_{BuMA} + 2*DP_{longest hydrophilic}]^{0.254}$ nm.

$^d$ In model 2, both BuMA blocks and DEAEMA blocks contributed to the formation of the cores
of the micelles, while the corona of the micelle was formed by PEGMA blocks only. The
equation used to calculate the hydrodynamic diameter is $d_H = [(DP_{BuMA} + DP_{DEAEMA}) +
2*DP_{PEGMA}]^{0.254}$ nm.

$^e$ The results reported are the average of the size corresponding to the maximum by intensity.
In case more than one peak was detected, the results reported are the ones corresponding to
the peak of micelles.

$^f$ The initial pH of the polymer solutions varied between 7.1-7.6.

$^g$ The CPs were tested at the initial pH of the polymer solutions in DI water.

Hydrodynamic Diameters

As DEAEMA is a monomer with tunable hydrophobicity, the self-assembly behaviour
(micelle size and size distribution) of the copolymer solutions at 1 w/w% has been
thoroughly investigated in different aqueous media by DLS: i) DI water at initial pH (pH between 7.1 to 7.6, depending on the strength of the amine groups), ii) DI water at alkaline pH (pH 8) to ensure complete deprotonation of the DEAEMA units and thus hydrophobic character, iii) DI water at acidic pH (pH 5) at which complete protonation of the DEAEMA units takes place and thus they are hydrophilic, and iv) PBS. The hydrodynamic diameters have been experimentally determined and compared to the calculated ones.

For the theoretical calculations, two models are proposed and are illustrated in Figure 4 below. The proposed self-assembly structures of Polymer 1 are shown as an example. In the first model (structure shown on the left), the hydrophobic BuMA (shown in red) forms the core of the micelle, while the corona consists of both PEGMA (represented by light blue spheres) and DEAEMA (shown in yellow); in this model, it is assumed that DEAEMA is hydrophilic, which is the case at pH values below its pKᵦ. In the second model (structure shown on the right), the core is formed by both BuMA and DEAEMA, while PEGMA contributes to the corona of the micelle; this is the case when the pH is higher than the pKᵦ value of the DEAEMA units. In both models and derivation of the corresponding equations, it is assumed that i) the hydrophobic block(s) form(s) the core of the micelle, while the hydrophilic block(s) form the corona of the micelles, ii) the hydrophobic part fully overlaps, iii) only the methacrylate backbone contributes to the micelle size (i.e. length side chains, e.g. in PEGMA, are not taken into consideration), iv) the methacrylate backbone is fully extended, v) the projected length of a methacrylate backbone equal to 0.254 nm is considered.

![Figure 4: Schematic representation of the suggested micelle configuration adopted by Polymer 1. Model 1 (left) assumes that the core is formed only by the hydrophobic BuMA units (red spheres), while Model 2 (right) assumes that both BuMA and DEAEMA (yellow spheres) units contribute to the core of the micelle.](image-url)
As previously mentioned, Table 2 includes the experimental peak by intensity that corresponds to the micelle size (around $10 \leq d_H \leq 50$ nm). Depending on the polymer composition, the pH of the aqueous solution and the solvent (DI water versus PBS), more than one peaks might be detected, which indicates the presence of species of various sizes. Specifically, unimers ($d_H \leq 10$ nm) and aggregates ($d_H \geq 100$ nm) might be present. The PDI values, which indicate the homogeneity of the samples, are presented in the SI (Table S2), as well as the full histograms (Figures S3-S6). It is noteworthy, that at acidic pH, the micelle structures seem to be disrupted, as most of the polymer solutions consist of unimers and aggregates, in addition to the micelles. Remarkably, the polymer solutions in PBS, which is the solvent that the gelation is investigated in, are monodisperse, as indicated by the low PDI values (0.035-0.156), and the presence of only one peak that corresponds to the micelles.

When either of the calculated diameters are compared to the experimental hydrodynamic diameters by intensity, it is observed that the experimental values are in most of the cases higher than the theoretical ones, regardless the pH of the solution or the solvent (DI water versus PBS). This opposes to the expected trend,[38, 42, 67, 78, 79] and it can be ascribed to i) the poor overlap of the hydrophobic part due to either electrostatic repulsions and/or steric hindrance, and ii) the theoretical calculations not taking into consideration any contributions of the length of the side chain. Interestingly, in most of the cases, the hydrodynamic diameters at alkaline pH (at which the DEAEMA units are hydrophobic) are higher than those at acidic pH (at which the amine groups are hydrophilic). This can be explained upon consideration of the amine groups contributing to the core formation and thus leading to formation of bigger micelles (see Figure 4).

**Dissociation Constants (pKₐs)**

As the polymer contains an amine-based repeated unit, their aqueous solutions were characterised by hydrogen ion titrations and their pKₐs were calculated and listed in Table 2. As can be seen, the pKₐ values vary from 6.2 to 7.1, which are composition dependent. As can be seen, all the copolymers with 10 w/w% BuMA content show pKₐ between 6.8 and 7.1. This range is in high agreement with the results of our recently-published study, in which copolymers based only on DEAEMA and PEGMA were studied.[72] Nevertheless, an important drop on the pKₐ is observed when the content in BuMA is increased to 20 w/w%, with the values being between 6.2 and 6.7. The effect of the hydrophobicity on the pKₐ has been reported previously,[42] and it is explained by the reduced dielectric constant caused by the increased hydrophobicity.

**Cloud Points**

As this study focuses on the thermoresponsive properties of novel terpolymers by varying the composition, as well as the effect of the pH response of DEAEMA units on the thermoresponsive behaviour, the CPs were investigated at 1 w/w% polymer concentration in three different environments: i) DI water at initial pH, ii) DI water at
alkaline pH (pH 8), and iii) PBS; the results are presented in Figure 5 below in squares, triangles, and circles, respectively.

**Figure 5**: Effect of the pH (initial pH and pH 8) and effect of the solvent (DI water versus PBS) on the CP of the 1 w/w% copolymer solutions. The error of the technique is ± 2°C. Note: Polymers 1-3 have 10 w/w% hydrophobic BuMA content, while Polymers 4-5 have 20 w/w% BuMA content. In both families, the DEAEMA content increases from Polymers 1-3 and 4-6, respectively.

As can be seen in Figure 5, two effects can be drawn concerning the CPs at initial pH (squares). Regarding the copolymers with only 10 w/w% BuMA content (Polymer 1 to 3), the CP increases from 61°C to 68°C by decreasing the DEAEMA content increases (or the PEGMA content decreases). On the other hand, the CPs of the copolymers with 20 w/w% BuMA (Polymer 4 to 6) drops from 62°C to 53°C as the DEAEMA content increases (or the PEGMA content decreases); this trend has been observed before in PEGMA-DEAEMA based copolymers.[72] The CPs are comparable with the ones resulted in our latest study, in which they varied from 61°C to 57°C as the content in DEAEMA increased.[72] However, it should be reminded that the initial pH of the solutions varied from 7.1 to 7.6, polymer dependent, which indicates different degrees of ionisation of the DEAEMA units, which could affect the CP values.

To confirm our hypothesis and eliminate any effects resulting from varying degrees of ionisation on the CPs, the pH was adjusted to 8, to ensure complete deprotonation of the amine groups and to eliminate any effect of the electrostatic interactions on the CP. The CPs at alkaline pH are presented in Figure 5 (in triangles). As can be seen, the CPs at alkaline pH vary between 54°C to 51°C and they are consistently lower than the CPs at initial pH. This is expected as the complete deprotonation of the DEAEMA units ensures that they are at the hydrophobic state, which enhances the “hydrophobic” effect, thus lowering the CPs. Therefore, it is expected that decreasing the pH below the pK<sub>a</sub> will result to increase in the CP. This has been previously observed on DMAEMA-containing polymers.[41, 80, 81] The lowest CP is presented
by Polymer 6 (CP at 51°C), which might be attributed to its smaller micelle size formed at pH 8, i.e. 15.7 nm versus 28.2-32.7 nm. This agrees with the previous findings by our group on DMAEMA-containing polymers.[42] Interestingly, when comparing the CPs of the copolymers at pH 8 with PEGMA homopolymers of comparable molar mass as the PEGMA block (unpublished data), it is observed that the CPs of the triblock copolymers are much lower than those of the PEGMA homopolymers (around 77°C). Thus, it can be concluded that the incorporation of both BuMA and DEAEMA units in the polymer structure favours the thermoresponse by lowering the CPs.

The CPs in PBS are also shown in Figure 5 (in circles) and they vary between 49°C and 52°C. It is observed that the increase in ionic strength reduces the CP values. This has been observed before on ionisable polymers and can be attributed to the ionic strength effect.[80, 82]

**Phase Diagrams**

The concentrated polymer solutions have also been investigated for gelation and the results are presented in means of phase diagrams in Figure 6. The effect of BuMA composition is presented from the top to the bottom, while the effect of the PEGMA/DEAEMA composition is presented from the left to the right. The polymer solutions were inspected for presenting 9 different transitions: (i) runny solution phase in white (squares, triangles and circles for transparency, slight cloudiness and cloudiness, respectively), ii) viscous solution phase in red (triangles and circles for transparency and cloudiness, respectively), iii) stable gel phase in blue (triangles and circles for transparency and cloudiness, respectively), and iv) two phases in yellow (rhombi and squares for gel syneresis and precipitation, respectively).
Figure 6: Detailed phase diagrams of all the soluble polymers (P1-P6) in phosphate buffered saline (PBS). The effect of the BuMA content is shown from the top to the bottom, whereas the effect of the DEAEMA content is shown from the left to the right. The runny solution phase is presented in white symbols (square, triangle and circle show clear, slightly cloudy, and cloudy solution), while the viscous solution phase is symbolised in red triangles (transparent) and circles (cloudy). The stable gel phase is coloured in blue triangles (transparent) and circles (cloudy), whereas when two-phases are present, i.e. gel syneresis and precipitation, are represented by yellow rhombus and squares, respectively. The gelation area, i.e. temperature and concentration values that a stable gel is formed, is drawn in black dashed line. The corresponding polymer structures are also schematically illustrated, with PEGMA, BuMA, and DEAEMA represented by blue, red, and yellow spheres, respectively.

In Figure 6, it can be observed that all the copolymer solutions show thermoresponse upon heating. More specifically, all the copolymer solutions, regardless the polymer composition and concentration, present a CP as the temperature increases. Upon further heating, phase separation occurs at all cases.

Interestingly, clear effects of the BuMA and DEAEMA contents on the thermoresponse are observed. It can be clearly concluded that by increasing the BuMA content from 10 w/w% to 20 w/w%, the gelation is favoured. Concerning the family of 10 w/w% BuMA (Polymers 1, 2 and 3), only Polymer 3, with the highest content in DEAEMA,
forms gels at the highest concentration tested. In contrast, Polymers 1 and 2, which have the same BuMA content as Polymer 3 but lower DEAEMA content, only precipitate upon heating, with the exemption of their concentrated polymer solutions that pass through a viscous solution state prior to clear phase separation. On the other hand, concerning the copolymers with 20 w/w% BuMA, both Polymer 5 and 6 form gels upon heating, while Polymer 4, with the lowest DEAEMA content, precipitates regardless the concentration. Also, as can be seen in Figure 8, Polymer 6 presents the widest gelation area of all the copolymers tested. Specifically, Polymer 6 gels at a concentration of at least 15 w/w%, while its most concentrated polymer solution forms a gel from 48°C to 53°C. The better thermogelling properties of Polymer 6 are attributed to its high BuMA and DEAEMA content. Therefore, it can be concluded that both BuMA and DEAEMA favour the gelation, as they enhance the “hydrophobic effect” in thermoresponsive polymers, which supports the previous findings.\[38, 40, 42, 67\]

**Rheological Properties**

Polymer 6, which showed the best gelation properties visually, has been tested at 30 w/w% in PBS via rheology and the results are presented in Figure 7 below. Three different measurement types have been carried out: (a) temperature ramp measurements, (b) time sweep measurements and (c) flow sweep measurements.

![Figure 7: Rheological measurements performed at 30 w/w% solution of Polymer 6 (PEGMA$_{11}$-b-BuMA$_{10.5}$-b-DEAEMA$_{14}$) in phosphate buffered saline (PBS): (a) temperature ramp measurement (storage modulus, G’, in blue, loss modulus, G”, in red, and complex viscosity, η*, in light blue), (b) time sweep measurements at 49°C (storage modulus, G’, in blue, loss modulus, G”, in red, and complex viscosity, η*, in light blue), and (c) flow sweep measurements (viscosity versus shear rate) at 49°C.](image)
Concerning the temperature ramp measurement (Figure 7(a)), gelation is observed upon temperature increase, as the storage modulus (G', shown in blue) exceeds the loss modulus (G", shown in red). Further increase in temperature leads to gel destabilisation, as expected from the visual tests. More specifically, a gel is formed rheologically at 49°C, while it is disrupted at 55°C. These temperature values are in high agreement with the results by visual tests, within the error of the techniques, as a gel was formed visually at 48°C, gel syneresis was presented at 53°C, and precipitation occurred at 55°C. Therefore, it is concluded that rheology confirms the visual tests.

To expand the rheological evaluation, the same polymer solution has been also tested in terms of its gelation time and shear-thinning properties at 49°C, and the results are presented in Figure 7(b) and 7(c), respectively. As can be seen in Figure 7(b), both moduli (storage modulus, G', in blue and loss modulus, G", in red) increase in magnitude as the time passes, and at 130s there is a crossover of the two moduli (G'> G"), which indicates that gelation takes place. Interestingly, the gel possesses shear-thinning properties, as the viscosity decreases with increased shear rate (see Figure 7(c), which is important especially in 3D printing.[27-29]
Conclusions

This study presents the synthesis and characterisation of novel thermoresponsive polymers based on PEGMA, BuMA, and DEAEMA. Six triblock copolymers (PEGMA$_x$-b-BuMA$_y$-b-DEAEMA$_z$), of the same total molar mass and architecture but various compositions were systematically investigated in terms of their thermoresponsive properties; $x$, $y$, and $z$ are the degrees of polymerisation of PEGMA, BuMA, and DEAEMA units, respectively. All the copolymers were successfully synthesised via GTP and their solubility in aqueous media was tested. Their pK$_a$ values were investigated and it has been found to be affected by the BuMA content. The effects of the degree of ionisation and the ionic strength on the self-assembly behaviour have been thoroughly investigated by DLS. The aqueous polymer solutions have been extensively studied for visual changes at various temperatures. The detailed phase diagrams prove that the gelation is strongly governed by the content in DEAEMA and BuMA. The gelation of the best-performing polymer has been investigated rheologically, and its gelation temperature, gelation time and shear-thinning properties have been determined.

Declaration of Competing Interest

The authors declare no competing interest.

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References


