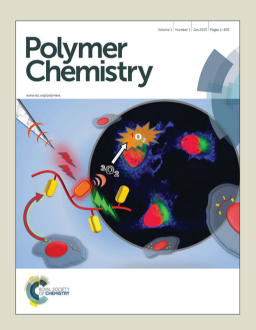
# Polymer Chemistry

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# ARTICLE TYPE

# One pot synthesis of higher order quasi-block copolymer libraries via sequential RAFT polymerization in an automated synthesizer†

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Recently developed sequential reversible addition-fragmentation chain transfer (RAFT) polymerization protocols allow the rapid, fully unattended preparation of quasi-block copolymer libraries that cover a wide range of copolymer composition in an automated synthesizer. This contribution explores the scope 10 and limitations of this sequential approach for the synthesis of higher order quasi-multiblock copolymers (including copolymer sequences of BAB, CBABC, ABC and ABCD). These syntheses illustrate the utility of this high-throughput approach for the one pot synthesis of functional polymers of increased complexity. Additionally, the use of this experimental technique for method development is highlighted.

#### Introduction

15 One of the major benefits to stem from reversible additionfragmentation chain transfer (RAFT) polymerization<sup>1</sup> is ready access to block copolymers.<sup>2</sup> However, the protocol for polymer synthesis, requiring multiple isolation and purification steps, can be demanding. Thus the development of one-pot methods for the 20 synthesis of block copolymers using RAFT polymerization<sup>3-5</sup> and other reversible-deactivation radical polymerization (RDRP)<sup>6</sup> methods, such as nitroxide mediated polymerization (NMP) and atom transfer radical polymerization (ATRP)<sup>8</sup>, is currently a research topic of substantial interest to industry and academia. 25 The adoption of such protocol inevitably leads to lower production costs through avoidance of expensive and timeconsuming intermediate purification steps. 3-5,7,8

During RAFT polymerization, every mole of initiator decomposed will produce between one and two moles of dead 30 chains. Those formed during synthesis of a first block of an A-B block will constitute a homopolymer A impurity in the block copolymer. It will also produce between one and two moles of initiator-derived chains. Those formed during synthesis of the second block of an A-B block will constitute a homopolymer B 35 impurity in the block copolymer. Thus to minimize impurity one should minimize the amount of initiator consumed and take the polymerization to form block A to <100 % conversion.<sup>9</sup>

In the synthesis of multi-block copolymers from high  $k_p$ monomers such as acrylates and acrylamides near quantitative 40 monomer conversions can be achieved rapidly with very low initiator concentrations and thus minimal formation of dead chains and/or homopolymer impurities. 5,10 The fraction of living chains at any stage of RAFT polymerization can be easily estimated with knowledge of the concentrations of RAFT agent 45 and initiator and readily available kinetic parameters. 26

For lower  $k_p$  monomers, which include methacrylates and

styrenes, where high conversions require longer polymerization times and/or higher initiator concentrations, a different strategy is required.

Two main strategies have been utilized to achieve the synthesis of block-like copolymers using one pot techniques via RDRP: 1) Exploitation of differing monomer reactivity (i.e., reactivity ratios) in limited comonomer systems<sup>3a,7</sup> and 2) utilizing sequential monomer addition. 3b,3c,4,5,8 These latter approach yields 55 quasi-block copolymers when <100% of the first monomer has been consumed prior to a second monomer being incorporated. This approach is necessary with the low  $k_p$  monomers where relatively high initiator concentrations are required to obtain acceptable rates of polymerization.3,4

The term quasi-block was introduced to refer to block copolymers formed by sequential addition of monomers A, B, ... where the second (and subsequent) blocks are, in the general case, some form of gradient copolymer poly(A)-block-poly(Agrad-B)... due to the incorporation of residual first block 65 monomer(s). For the case of an all methacrylate quasi-block where reactivity ratios are close to unity the product will be a poly(A)-block-poly(A-ran-B); i.e., the ratio of A:B will remain essentially constant. The determination of composition is not trivial from an experimental point of view. Nevertheless, it can be 70 estimated by simulation using the appropriate monomer reactivity ratios.11

In this work, we use high-throughput polymer synthesis<sup>12</sup> and build upon our previously developed one pot high-throughput synthetic strategy to develop protocols for the rapid synthesis of 75 quasi-multiblock copolymer libraries based on methacrylates. 4,13 Furthermore, we utilize this experimental technique for the optimization of the reaction conditions of the investigated systems. This method allows the rapid and systematic preparation of higher order (multi) quasi-block copolymer libraries with 80 "new" block combinations that expand over a comprehensive

copolymer composition range. Rapid access to these new materials is particularly pertinent for rapid screening of structureproperty relationships and development of novel applications. These are areas where quasi-block copolymers are ideally suited 5 and are currently being applied. 14,15

# **Experimental**

#### Materials

*n*-Butyl methacrylate (BMA) and methyl methacrylate (MMA), 10 di(ethylene glycol) methyl ether methacrylate (DEGMA) and benzyl methacrylate (BzMA) monomers were purchased from Sigma-Aldrich and purified by stirring in the presence of inhibitor-remover for hydroquinone or hydroquinone monomethyl ether (Aldrich) for 30 minutes prior to use. RAFT 4-Cyano-4-[(dodecylsulfanylthiocarbonyl)sulfanyl]pentanoic acid was purchased from Strem Chemicals and utilized as received. Bis-RAFT agent: 4-cyano-4(dodecylthiocarbonothioylthio)pentanoyloxy) butyl 4-cyano-4-20 (dodecylthiocarbonothioylthio)pentanoate prepared reported literature to procedures. 1.1′-Azobis(cyclohexanecarbonitrile) (ACHN) initiator (DuPont VAZO-88) was used as received. N,N-Dimethylformamide (DMF) (AR grade) was purchased from Merck.

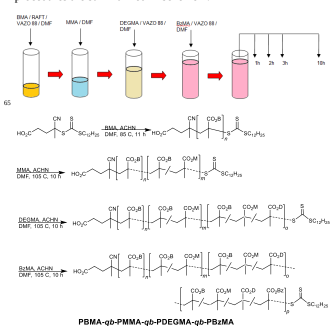
#### 25 Automated parallel synthesizer

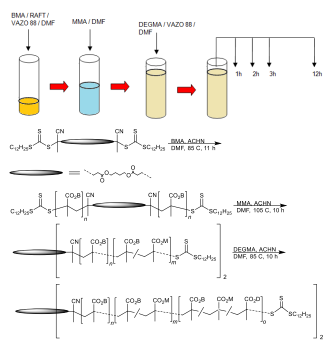
The commercially available synthesizer utilized in this work was a Chemspeed Swing-SLT automated parallel synthesizer. 13 The synthesizer was equipped with a glass reactor block consisting of 16 reaction vessels (13 mL) with thermal jackets connected in 30 series through the reaction block to a heating/cooling system (Hüber, -90 °C to 140 °C). In addition, all reaction vessels were equipped with cold-finger reflux condensers (~7 °C). Mixing was achieved by vortex agitation (up to 1400 rpm). Liquid transfers were handled by a 4-needle head (4-NH) capable of four 35 simultaneous sample transfers. The 4-NH was connected to a reservoir bottle (degassed DMF solvent) for needle rinsing after each liquid transfer step. This DMF solvent reservoir was degassed by continuous sparging with nitrogen and was also utilized to prime the tubing lines of the 4-NH. When experiments 40 were carried out, the synthesizer was maintained under an inert atmosphere by supplying a constant flow of nitrogen to the hood of the synthesizer. A nitrogen atmosphere was also applied to reactors and stock solutions at all times. Prior to the experiments, the reaction vessels were heated to 135 °C and subjected to 10 45 cycles of vacuum (2 min each) and filling with nitrogen (2 min each) to ensure the elimination of oxygen. After this pretreatment, the RAFT polymerization experiment was carried out following similar procedures to those reported elsewhere. 4,13 It is worth mentioning that the characteristics of the RAFT-50 synthesized polymers prepared in the automated parallel synthesizer are very similar to those obtained in conventional batch polymerization performed in sealed ampoules as demonstrated in a previous contribution. 13a

#### Automated synthesis of high order (multi) quasi-block 55 copolymer libraries

Monomers and solvent (BMA, MMA, DEGMA, BzMA and DMF), and stock solutions of ACHN (4.12 mg mL<sup>-1</sup> in DMF),

RAFT (136.24 mg mL<sup>-1</sup> in DMF) and bis-RAFT (140 mg mL<sup>-1</sup> in DMF) agents were prepared, degassed by sparging with nitrogen 60 for 15 min, and placed inside the automated synthesizer. The sequential RAFT polymerization method utilized in this work was adapted from that previously reported<sup>4</sup> and is described below. Schematic representations of the followed synthetic procedures are summarized in scheme 1.





 ${\tt PDEGMA-}qb{\tt -PMMA-}qb{\tt -PBMA-}qb{\tt -PMMA-}qb{\tt -PDEGMA}$ 

Scheme 1. Representation of the automated parallel synthesis of high 70 order (multi) quasi-block copolymer libraries in one pot via sequential RAFT polymerization utilizing a RAFT agent (top) and a bis-RAFT agent (bottom). Illustration of the two most complex cases: ABCD (top) and CBABC (bottom), where A, B, C and D represent monomers BMA, MMA, DEGMA and BzMA, respectively. The letters also indicate the 75 order of monomer incorporation.

The utilized characterization methods, *i.e.*, proton nuclear magnetic resonance (<sup>1</sup>H-NMR) and size exclusion chromatography (SEC), are described in the Electronic 5 Supporting Information (ESI). Figure S1 in the ESI displays representative <sup>1</sup>H-NMR spectra and their analysis to calculate conversions for the four different monomers investigated in this work

Synthesis of poly(butyl methacrylate) (PBMA) macro RAFT and 10 bis-macro-RAFT agents.

For each investigated case, aliquots of RAFT agent (for the ABC and ABCD cases) or bis-RAFT agent (for the BAB and CBABC cases) and ACHN stock solutions, BMA and solvent from the reservoir were transferred into different reactors (13 mL) of the 15 synthesizer with the automated liquid handling system to provide the desired concentrations of reagents. The total reaction volume was 1.75 mL corresponding to a BMA concentration of 2.143 M  $(3.75 \times 10^{-3} \text{ mol of BMA})$ . Different degrees of polymerization (DP) –assuming 100% monomer conversion– were targeted (50, 20 75, 100 and 150) by utilizing different RAFT agent concentrations  $(4.286 \times 10^{-2} \text{ M} (7.50 \times 10^{-5} \text{ mol of RAFT}), 2.857$  $\times 10^{-2} \text{ M} (5.00 \times 10^{-5} \text{ mol of RAFT}), 2.143 \times 10^{-2} \text{ M} (3.75 \times 10^{-5} \text{ mol of RAFT})$ mol of RAFT) and  $1.429 \times 10^{-2}$  M ( $2.50 \times 10^{-5}$  mol of RAFT), respectively). A [RAFT]/[Initiator] ratio of 10, 6.667, 5 and 3.333 25 were used for the DP of 50, 75, 100 and 150, respectively (i.e.,  $7.50 \times 10^{-6}$  mol of initiator ACHN in all cases). After all the liquid transfers were completed, the reaction solutions were degassed through three automated freeze-evacuate-thaw cycles carried out as follows: The reaction mixture was cooled to -90 °C 30 while applying vortex to the reaction block (200 rpm). Vacuum (~5 millibar) was then applied to the reactor block while warming the reactors to −10 °C with vortex (600 rpm, 2 min). Thereafter, the reactors were sealed under nitrogen and heated to the reaction temperature (85 °C) while applying vortex to the reaction block 35 (typically 300 rpm); the temperature of the reflux condensers was set at 7 °C. The onset of the polymerizations was considered the point in which the reaction temperature in the apparatus reached the set point. At the pre-established reaction time of 11 h for all cases except for the BAB and CBABC cases (12 h), an aliquot of 40 200 μL was withdrawn with the automated liquid handling system from each reaction mixture and transferred to nuclear magnetic resonance (NMR) tubes and size exclusion chromatography (SEC) vials (75 µL each) to determine monomer conversion, molar mass and dispersities of the formed polymer, 45 respectively.

SEC and NMR samples for analysis were prepared with the automated liquid handling system of the synthesizer at the end of each sampling sequence by adding the corresponding SEC and NMR solvents. Once the pre-established reaction time elapsed, 50 the polymerization mixture was cooled to 20 °C.

Synthesis of the BAB quasi-triblock copolymer libraries of poly(methyl methacrylate-quasi-block-butyl methacrylate-quasi-block-methyl methacrylate) (PMMA-qb-PBMA-qb-PMMA).

After the synthesis of the PBMA *bis*-macro-RAFT agent 55 described above, 1 mL of MMA (9.39 × 10<sup>-3</sup> mol of MMA) was added to the reactor, followed by 1.5 mL DMF to improve the homogeneity of the new reaction mixture. The total reaction volume of this second polymerization was 4.05 mL. Thereafter this new reaction mixture was degassed through three automated 60 freeze-evacuate-thaw cycles. The reactor was heated at 105 °C

for 11 h to carry out the chain extension reaction and to obtain different BAB quasi-triblock copolymers. During this step, aliquots (200  $\mu$ L) were automatically withdrawn from the reactors at different time intervals for NMR and SEC analysis (75  $\mu$ L each).

Synthesis of the CBABC quasi-pentablock copolymer libraries of poly(di(ethylene glycol) methyl ether methacrylate-quasi-block-methyl methacrylate-quasi-block-butyl methacrylate-quasi-block-methyl methacrylate-quasi-block-di(ethylene glycol) methyl ether methacrylate) (PDEGMA-qb-PMMA-qb-PBMA-qb-PMMA-qb-PDEGMA).

Following a similar procedure to the synthetic case of the BAB quasi-triblock copolymer materials, BAB bis-macro-RAFT agents were synthesized at 105 °C with a reaction time of 10 h. 75 After this period, an aliquot of 200 µL was automatically withdrawn from each reaction mixture for NMR and SEC analysis (75 µL each), and the remaining polymerization mixtures were cooled to 20 °C. Thereafter, 1 mL of DEGMA (5.419  $\times$  10<sup>-3</sup> mol of DEGMA) and 0.25 mL of ACHN stock solution (4.215  $\times$ <sub>80</sub> 10<sup>-6</sup> mol of initiator) were added to the reactor. 1.5 mL of DMF (in total) was also added to the reactor to improve the homogeneity of the new reaction mixture. The total reaction volume of this second polymerization was 6.35 mL. This new reaction mixture was degassed through three automated freeze-85 evacuate-thaw cycles. The reactor was heated at 85 °C for 8 h to carry out the second chain extension reaction and to obtain different CBABC quasi-pentablock copolymers. During this step, aliquots (200 µL) were automatically withdrawn from the reactors at different time intervals for NMR and SEC analysis (75 90 μL each).

Synthesis of the ABC quasi-triblock copolymer libraries of poly(butyl methacrylate-quasi-block-methyl methacrylate-quasi-block-di(ethylene glycol) methyl ether methacrylate) (PBMA-qb-PMMA-qb-PDEGMA).

95 Following similar procedures to the synthetic cases of the BAB quasi-triblock and CBABC quasi-pentablock copolymer materials explained above, AB macro RAFT agents were synthesized at 105 °C with a reaction time of 10 h. After this period, an aliquot of 200 µL was automatically withdrawn from each reaction 100 mixture for NMR and SEC analysis (75 μL each), and the remaining polymerization mixtures were cooled to 20 °C. Thereafter, 1 mL of DEGMA (5.419  $\times$  10<sup>-3</sup> mol of DEGMA) and 0.25 mL of ACHN stock solution 88 (4.215  $\times$  10<sup>-6</sup> mol of initiator) were added to the reactor. 1.5 mL of DMF (in total) was 105 also added to the reactor to improve the homogeneity of the new reaction mixture. The total reaction volume of this second polymerization was 6.35 mL. Next, this new reaction mixture was additionally degassed through three automated freeze-evacuatethaw cycles as explained above. The reactor was heated at 85 °C 110 for 8 h to carry out the second chain extension reaction and to obtain different ABC quasi-triblock copolymers. During this step, aliquots (200 µL) were automatically withdrawn from the reactors at different time intervals for NMR and SEC analysis (75 μL each).

115

Synthesis of the ABCD quasi-tetrablock copolymer libraries of poly(butyl methacrylate-quasi-block-methyl methacrylate-quasi-block-di(ethylene glycol) methyl ether methacrylate-quasi-block-benzyl methacrylate) (PBMA-qb-PMMA-qb-PDEGMA-qb-5 PBzMA).

Following a similar procedure to the synthetic case of the ABC quasi-triblock copolymer materials explained above, ABC macro-RAFT agents were synthesized at 85 °C with a reaction time of 12 h. At the onset of the respective second chain extension 10 reaction and after 12 h, aliquots of 200 μL were automatically withdrawn from each reaction mixture for NMR and SEC analysis (75 µL each). Thereafter, the remaining polymerization mixtures were cooled to 20 °C. Subsequently, 1 mL of BzMA  $(5.902 \times 10^{-3} \text{ mol of BzMA})$  and 0.25 mL of ACHN stock  $_{15}$  solution 88 (4.215  $\times$  10<sup>-6</sup> mol of initiator) were added to the reactor. 1.5 mL of DMF (in total) was also added to the reactor to improve the homogeneity of the new reaction mixture. The total reaction volume of this second polymerization was 8.45 mL. This new reaction mixture was additionally degassed through three 20 automated freeze-evacuate-thaw cycles. The reactor was heated at 85 °C for 10 h to carry out the third chain extension reaction and to obtain different ABCD quasi-triblock copolymers. During this step, aliquots (200 µL) were automatically withdrawn from the reactors at different time intervals for NMR and SEC analysis (75 25 µL each).

#### **Results and discussion**

Within this investigation the simplest library synthesis corresponds to the BAB block copolymer case, which utilizes a similar procedure to that described in our previous contribution 30 for quasi-diblock copolymer libraries. 4b For this a series of one pot (two step) sequential polymerizations were performed in the automated synthesizer using a bis-macro-RAFT agent to yield a library of 24 PMMA-qb-PBMA-qb-PMMA quasi-triblock copolymers derived from three different bis-macro-RAFT agents. 35 Table 1 summarizes the synthetic results and the reaction conditions utilized (see Table 1 footnote A) of this library and shows that all materials have D values below 1.27. Figure 1 displays representative SEC traces demonstrating an efficient chain extension process for each specific case. The same data is 40 presented with the SEC traces scaled against monomer conversion (MMA) during the chain extension reaction of bis-RAFT agent precursor 1 in Figure 1D.

Figure 2 displays kinetic plots of the chain extension reaction of one of the investigated cases (*bis*-macro-RAFT agent 2 in Table 1) where a linear relationship can be observed between the number average molar mass (*M*<sub>n</sub>) *vs*. conversion (x). and the  $-\ln(1-x)$  *vs*. reaction time indicating good control over the consecutive polymerizations. Additional kinetic plots for the cases of the *bis*-macro-RAFT agents 1 and 3 (Table 1) can be found in Figure S2 in the ESI. As full conversion was not achieved during the first polymerization the residual BMA is incorporated within the PMMA blocks of the PMMA-qb-PBMA-qb-PMMA quasi-triblock copolymers. The monomer conversions for the three different *bis*-macro-RAFT agents of Table 1 were 55 86, 91 and 89% for 1, 2 and 3, respectively. Based on these measurements and using a similar <sup>1</sup>H-NMR analysis as reported elsewhere, <sup>4</sup> the amount of BMA incorporated into the PMMA

blocks during the second polymerization step can be estimated. In all cases of Table 1 was found that the PMMA-*qb*-PBMA-*qb*<sup>60</sup> PMMA quasi-triblock copolymers have BMA units within the PMMA blocks below or at a value of 4 mol %.

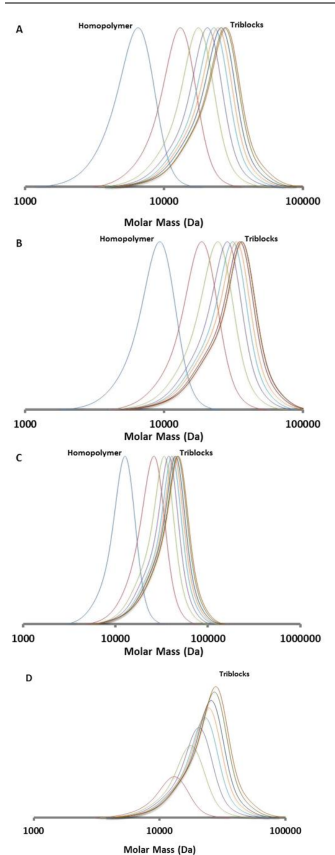
**Table 1.** Characteristics of PMMA-qb-PBMA-qb-PMMA materials obtained from the chain extension of three different *bis*-macro-RAFT agents in the automated parallel synthesizer. A

	<b>1.</b> A <i>bis</i> -macro-RAFT agent ( $M_n = 5400 \text{ g mol}^{-1}$ , $D = 1.14$ )						
	ID	Reaction	$M_{\rm n}$	Đ	$M_{\rm n(theory)}$	MMA	
_	ID	time (h)	(g mol <sup>-1</sup> )	D	(g mol <sup>-1</sup> )	conversion (%)	
	1A	1	11300	1.12	10632	29	
	1B	2	14900	1.14	13708	51	
	1C	3	17100	1.15	15428	64	
	1D	4	18700	1.16	16370	71	
	1E	5	19700	1.17	17358	78	
	1F	6	20900	1.14	18040	83	
	1G	8	21700	1.18	18860	89	
	1H	11	22400	1.19	19463	93	

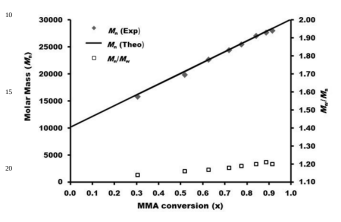
<b>2.</b> A <i>bis</i> -macro-RAFT agent ( $M_n = 7800 \text{ g mol}^{-1}$ , $D = 1.13$ )						
ID	Reaction	$M_{ m n}$	Đ	$M_{\rm n(theory)}$	MMA	
110	time (h)	(g mol <sup>-1</sup> )	Đ	(g mol <sup>-1</sup> )	conversion (%)	
2A	1	15800	1.14	16279	31	
2B	2	19900	1.16	20478	52	
2C	3	22600	1.17	22669	63	
2D	4	24400	1.18	24512	72	
2E	5	25500	1.19	25641	78	
2F	7	27000	1.20	26971	84	
2G	9	27700	1.21	27918	89	
2H	11	28000	1.20	28493	92	

TD	Reaction	$M_{ m n}$	D.	$M_{\rm n(theory)}$	MMA
ID	time (h)	(g mol <sup>-1</sup> )	Đ	(g mol <sup>-1</sup> )	conversion (%)
3A	1	20900	1.17	21408	31
3B	2	26200	1.19	26574	50
3C	3	28900	1.20	28926	59
3D	4	31100	1.21	30588	65
3E	5	32100	1.23	32256	72
3F	6	33500	1.25	34264	79
3G	8	34300	1.26	35558	84
3H	11	34900	1.26	36560	87

<sup>A</sup> Number average molar mass ( $M_n$ ) and dispersity ( $D = M_w / M_n$ ) were estimated by SEC and are reported as PMMA equivalents. The monomer to polymer conversion was determined by  $^1$ H-NMR.  $M_{n(theory)}$  was estimated using the formula:  $M_{n(theory)} = ([M_{BMA}]_o \times M_{BMA} \times \%$  conversion<sub>BMA</sub> +  $[M_{MMA}]_o \times M_{MMA} \times \%$  conversion<sub>MMA</sub>) / [bis-RAFT]<sub>o</sub> +  $M_{bis-RAFT}$  M<sub>BMA</sub>, M<sub>MMA</sub> and M<sub>bis-RAFT</sub> are the molar masses of BMA, MMA and bis-RAFT agent, respectively. [M<sub>BMA</sub>]<sub>o</sub>, [M<sub>BMA</sub>]<sub>o</sub> and [bis-RAFT]<sub>o</sub> are the initial concentrations of BMA, MMA and bis-RAFT agent, respectively. For the synthesis of the A bis-macro-RAFT agents, [M<sub>BMA</sub>]<sub>o</sub> = 2.143 M, reaction temperature = 85 °C and reaction time = 12 h; [RAFT]:[Initiator]:[M<sub>BMA</sub>] ratios of 1:0.1:50, 1:0.15:75 and 1:0.2:100 were utilized for the cases of bis-macro-RAFT agents 1, 2 and 3, ss respectively. For the chain extension reaction of the respective bis-macro-RAFT agents, [M<sub>MMA</sub>]<sub>o</sub> = 2.318 M, reaction temperature = 105 °C and reaction time = 11 h.



5 Figure 1. SEC traces of the chain extension bis-macro-RAFT polymerization (Table 1) of the synthesized PMMA-qb-PBMA-qb-PMMA quasi-triblock copolymer materials derived from bis-macro-RAFT A) precursor agent 1, B) precursor agent 2, C) precursor agent 3 and D) precursor agent 1 scaled against conversion of MMA.



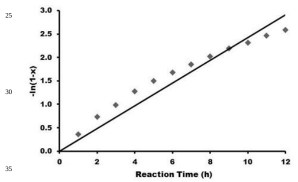


Figure 2. Kinetic data of the synthesized PMMA-qb-PBMA-qb-PMMA quasi-triblock copolymer materials derived from bis-macro-RAFT agent 2 (Table 1).  $M_n$  and D as a function of the MMA conversion (top) and 40 MMA conversion as a function of reaction time (bottom).

Next, we consider the synthesis of a PDEGMA-qb-PMMA-qb-PBMA-qb-PMMA-qb-PDEGMA quasi-pentablock copolymer library (CBABC case). For this CBABC case, three similar bis-45 macro-RAFT agents to those reported in Table 1 were sequentially chain extended with MMA to obtain the respective BAB bis-macro-RAFT agents of the type PMMA-qb-PBMA-qb-PMMA quasi-triblock copolymers. Thereafter, a second chain extension polymerization step with DEGMA to the three BAB 50 bis-macro-RAFT agents was sequentially undertaken to yield a library of 24 PDEGMA-qb-PMMA-qb-PBMA-qb-PMMA-qb-PDEGMA quasi-pentablock copolymers. Table 2 summarizes the synthetic results and the reaction conditions utilized (see Table 2 footnote B) of this library and shows that all the materials had  $\mathcal{D}$ 55 values below or at 1.37, whereas Figure 3 displays representative SEC traces demonstrating an efficient second chain extension process for each specific case. Figure 4 displays kinetic plots of the chain extension reaction of one of the investigated cases (BAB bis-macro-RAFT agent 4 in Table 2) where a linear 60 relationship can be observed between  $M_n$  vs. conversion (x) and the  $-\ln(1-x)$  vs. reaction time indicating good control over the consecutive polymerization. Additional kinetic plots for the cases of the BAB bis-macro-RAFT agents 5 and 6 (Table 2) can be found in Figure S3 in the ESI.

Similar to the previous analysis, full conversion was also not reached during the second polymerization step. Thus, the residual MMA was incorporated within the PDEGMA blocks of the PDEGMA-qb-PMMA-qb-PBMA-qb-PDEGMA

35

quasi-pentablock copolymers. The MMA monomer conversions for the three different BAB *bis*-macro-RAFT agents of Table 2 were 82, 86 and 82% for **4**, **5** and **6**, respectively. Based on these measurements and using the <sup>1</sup>H-NMR analysis described in the

**Table 2.** Characteristics of PDEGMA-*qb*-PMMA-*qb*-PBMA-*qb*-PMMA-*qb*-PDEGMA materials obtained from the chain extension of three different *bis*-macroRAFT agents in the automated parallel synthesizer. <sup>B</sup>

<b>4.</b> BAB bis-macro-RAFT agent ( $M_n = 18900 \text{ g mol}^{-1}$ ,	D = 1.15	)
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ID	Reaction	$M_{ m n}$	Đ	$M_{\rm n(theory)}$	DEGMA
110	time (h)	(g mol <sup>-1</sup> )	D	(g mol <sup>-1</sup> )	conversion (%)
4A	1	23500	1.18	22594	25
4B	2	26100	1.19	24471	37
4C	3	27800	1.20	26560	48
4D	4	27900	1.22	27554	55
4E	6	28900	1.25	29150	64
4F	8	30000	1.25	29869	68
4G	10	31600	1.26	30855	74
4H	12	32300	1.27	31538	78

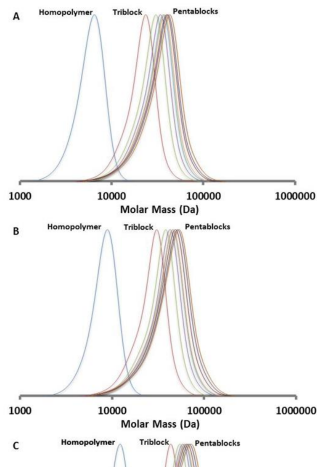
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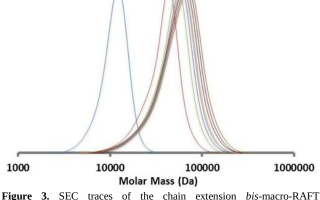
ID	Reaction	$M_{\rm n}$	Đ	$M_{\rm n(theory)}$	DEGMA
110	time (h)	(g mol <sup>-1</sup> )	D	(g mol <sup>-1</sup> )	conversion (%)
5A	1	29200	1.20	34428	25
5B	2	31400	1.24	37357	37
5C	3	33800	1.24	39536	47
5D	4	33400	1.26	41033	53
5E	6	35400	1.28	43676	63
5F	8	36900	1.29	45126	69
5G	10	38400	1.30	46421	75
5H	12	39600	1.30	47569	80

<b>6.</b> BAB <i>bis</i> -macro-RAFT agent ( $M_n = 32600 \text{ g mol}^{-1}$	, D = 1.22)
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ID	Reaction	$M_{\mathrm{n}}$	Đ	$M_{\rm n(theory)}$	DEGMA
	time (h)	(g mol <sup>-1</sup> )		(g mol <sup>-1</sup> )	conversion (%)
6A	1	38900	1.27	44392	25
6B	2	41000	1.30	47515	34
6C	3	42900	1.31	50151	42
6D	4	42200	1.34	52138	48
6E	6	44000	1.34	55626	58
6F	8	47000	1.33	57084	63
6G	10	48800	1.34	59149	69
6H	12	49700	1.37	60427	73

<sup>&</sup>lt;sup>B</sup> Number average molar mass  $(M_n)$  and dispersity  $(D = M_w / M_n)$  were estimated by SEC and are reported as PMMA equivalents. The monomer to polymer conversion was determined by  ${}^{1}H$ -NMR.  $M_{n(theory)}$  was estimated using the formula:  $M_{\text{n(theory)}} = ([M_{\text{BMA}}]_{\text{o}} \times M_{\text{BMA}} \times \%$ 20 conversion<sub>BMA</sub> +  $[M_{MMA}]_o \times M_{MMA} \times \%$  conversion<sub>MMA</sub> +  $[M_{DEGMA}]_o \times$  $M_{DEGMA} \times \%$  conversion<sub>DEGMA</sub>) / [bis-RAFT]<sub>o</sub> +  $M_{bis-RAFT}$ .  $M_{BMA}$ ,  $M_{MMA}$ , M<sub>DEGMA</sub> and M<sub>bis-RAFT</sub> are the molar masses of BMA, MMA, DEGMA and bis-RAFT agent, respectively. [MBMA]o, [MBMA]o, [MDEGMA]o and [bis-RAFT]<sub>o</sub> are the initial concentrations of BMA, MMA, DEGMA and bis-25 RAFT agent, respectively. For the synthesis of the A bis-macro-RAFT agents, [M<sub>BMA</sub>]<sub>o</sub> = 2.143 M, reaction temperature = 85 °C and reaction time = 12 h; [RAFT]:[Initiator]:[ $M_{BMA}$ ] ratios of 1:0.1:50, 1:0.15:75 and 1:0.2:100 were utilized for the cases of bis-macro-RAFT agents 4, 5 and 6, respectively. For the synthesis of the BAB bis-macro-RAFT agents,  $_{30}$  [M<sub>MMA</sub>]<sub>0</sub> = 2.318 M, reaction temperature = 105 °C and reaction time = 10 h. For the chain extension reaction of the respective BAB bis-macro-RAFT agents with DEGMA, [M<sub>DEGMA</sub>]<sub>o</sub> = 0.853 M, additional [Initiator]<sub>o</sub> =  $6.638 \times 10^{-4}$  M, reaction temperature = 85 °C and reaction time = 12h.

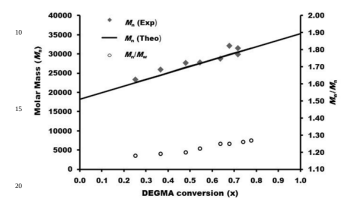


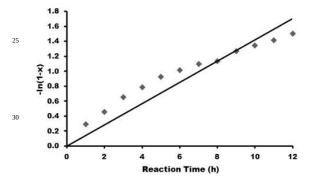


**Figure 3.** SEC traces of the chain extension *bis*-macro-RAFT polymerization (Table 2) of the synthesized PDEGMA-qb-PMMA-qb-40 PBMA-*qb*-PMMA-*qb*-PDEGMA quasi-pentablock copolymer materials derived from *bis*-macro-RAFT A) precursor agent **4** B) precursor agent **5** and C) precursor agent **6**.

experimental section, the amount of MMA incorporated into the PDEGMA blocks during the third polymerization step can be estimated. It was found that the PDEGMA-qb-PMMA-qb-PBMA-qb-PDEGMA quasi-pentablock copolymers have MMA units within the PDEGMA blocks in the range of 18 to 25 mol %. The relative high impurity found in the PDEGMA block can be ascribed to the lower MMA monomer conversion obtained during the second polymerization as well as to the low concentration of DEGMA monomer utilized for the third polymerization step. It was found that, within the investigated reaction conditions, higher conversions during the second

polymerization, higher concentration of third monomer values in the materials or dead polymer in the third sequential polymerization led to higher dispersity chains. This aspect is analysed in more detail below for the ABC quasi-triblock 5 copolymer library case. Nevertheless, further optimization to reduce these defects or impurities in the sequentially formed blocks could be easily achieved.





35 Figure 4. Kinetic data of the synthesized PDEGMA-qb-PMMA-qb-PBMA-qb-PMMA-qb-PDEGMA quasi-pentablock copolymer materials derived from BAB bis-macro-RAFT agent 4 (Table 2).  $M_n$  and D of as a function of the DEGMA conversion (top) and DEGMA conversion as a function of reaction time (bottom).

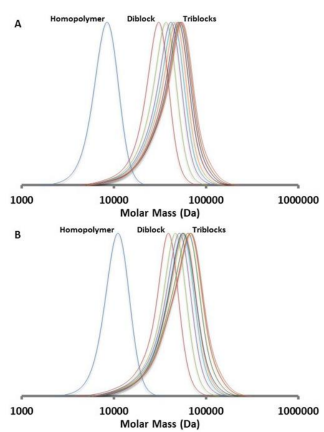
For the PBMA-qb-PMMA-qb-PDEGMA ABC case, two macro-RAFT agents were sequentially chain extended with MMA monomer utilizing the reaction conditions described in the experimental section to obtain the respective AB macro-RAFT 45 agents of the type PBMA-qb-PMMA quasi-diblock copolymers. Thereafter, a second chain extension with DEGMA monomer to the two AB macro-RAFT agents was sequentially undertaken to yield a library of 16 PBMA-qb-PMMA-qb-PDEGMA quasitriblock copolymers. Table 3 summarizes the synthetic results 50 and the utilized reaction conditions (see Table 3 footnote C) of this library and shows that all the materials have D values below or at 1.34, whereas Figure 5 displays representative SEC traces demonstrating an efficient second chain extension process for each specific case. Figure 6 displays kinetic plots of the chain 55 extension reaction of one of the investigated cases (AB macro RAFT agent 7 in Table 3) where a linear relationship can be observed between the  $M_n$  vs. conversion (x) and the  $-\ln(1-x)$  vs. reaction time indicating all in all a good control over the consecutive polymerization. Kinetic plots for the additional case 60 of the AB macro-RAFT agent 8 (Table 3) can be found in Figure S4 in the ESI. Similar to the previous analysis, full conversion was also not reached during the second polymerization step (synthesis of the AB macro-RAFT agents). Thus, the residual MMA was incorporated within the PDEGMA blocks of the 65 PBMA-qb-PMMA-qb-PDEGMA quasi-triblock copolymers. The MMA monomer conversions for the two different AB macro-RAFT agents of Table 3 were 80% for both cases (7 and 8). Attempts for bringing to higher conversions (> 80%) these polymerization reactions led to the appearance of a considerable 70 amount of dead polymer chains as revealed by SEC traces (appearance of shoulders and tails in the low molar mass range during the second chain extension reaction; see Figure S5 in the ESI for an example where a reaction time of 12 h was utilized instead of 11 h). Similar to our optimization of the reaction time, 75 to provide a reasonable balance between high monomer

**Table 3.** Characteristics PBMA-qb-PMMA-qb-PDEGMA materials obtained from the chain extension of two different macroRAFT agents in the automated parallel synthesizer.<sup>C</sup>

<b>7.</b> AB macro-RAFT agent ( $M_n = 24400 \text{ g mol}^{-1}$ , $D = 1.16$ )						
ID	Reaction	$M_{ m n}$	Đ	$M_{\rm n(theory)}$	DEGMA	
110	time (h)	(g mol <sup>-1</sup> )	D	(g mol <sup>-1</sup> )	conversion (%)	
7A	1	28200	1.19	30338	17	
7B	2	30800	1.22	33601	31	
7C	3	32900	1.23	35932	41	
7D	4	34300	1.24	37709	48	
7E	6	35900	1.26	40071	58	
7F	8	37400	1.27	41694	65	
7G	10	38100	1.29	43161	71	
7H	12	39100	1.30	44341	76	

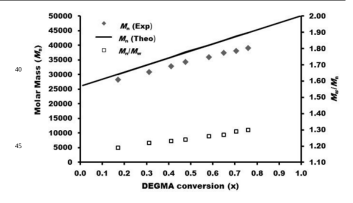
**8.** AB macro-RAFT agent  $(M_n = 29800 \text{ g mol}^{-1}, D = 1.21)$ Reaction **DEGMA**  $M_{\rm n}$  $M_{\rm n(theory)}$ ID Đ (g mol<sup>-1</sup>) (g mol<sup>-1</sup>) time (h) conversion (%) 17 8A 34800 1.22 39921 2 32 8B 37600 1.25 45265 3 39 8C 39400 1.27 47688 4 46 49938 8D41400 1.27 5 1.25 51583 52 8E 42600 60 8F 44100 1.31 54310 10 70 1.33 57458 8G 46100 59052 75 8H 47100 1.34

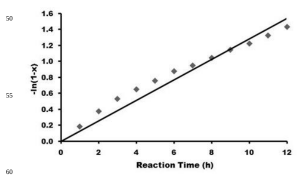
<sup>&</sup>lt;sub>85</sub> <sup>C</sup> Number average molar mass  $(M_{\rm n})$  and dispersity  $(D = M_{\rm w} / M_{\rm n})$  were estimated by SEC and are reported as PMMA equivalents. The monomer to polymer conversion was determined by  ${}^{1}H$ -NMR.  $M_{n(theory)}$  was estimated using the formula:  $M_{\rm n(theory)} = ([M_{\rm BMA}]_{\rm o} \times M_{\rm BMA} \times \%$  $conversion_{BMA} + [M_{MMA}]_o \times M_{MMA} \times \% conversion_{MMA} + [M_{DEGMA}]_o \times (M_{DEGMA})_o \times (M_{DEGMA})_$ 90  $M_{DEGMA} \times \%$  conversion<sub>DEGMA</sub>) / [RAFT]<sub>o</sub> +  $M_{RAFT}$   $M_{BMA}$ ,  $M_{MMA}$ ,  $M_{DEGMA}$ and M<sub>RAFT</sub> are the molar masses of BMA, MMA, DEGMA and RAFT agent, respectively. [M<sub>BMA</sub>]<sub>o</sub>, [M<sub>BMA</sub>]<sub>o</sub>, [M<sub>DEGMA</sub>]<sub>o</sub> and [RAFT]<sub>o</sub> are the initial concentrations of BMA, MMA, DEGMA and RAFT agent, respectively. For the synthesis of the A macro RAFT agents,  $[M_{BMA}]_o =$ 95 2.143 M, reaction temperature = 85 °C and reaction time = 11 h; [RAFT]:[Initiator]:[ $M_{BMA}$ ] ratios of 1:0.15:75 and 1:0.2:100 were utilized for the cases of macro RAFT agents 7 and 8, respectively. For the synthesis of the AB macro RAFT agents,  $[M_{MMA}]_o = 2.318$  M, reaction temperature = 105 °C and reaction time = 10 h. For the chain extension 100 reaction of the respective AB macro RAFT agents with DEGMA,  $[M_{DEGMA}]_o = 0.853 \text{ M}$ , additional [Initiator] $_o = 6.638 \times 10^{-4} \text{ M}$ , reaction temperature = 85 °C and reaction time = 12 h.



**Figure 5.** SEC traces of the chain extension RAFT polymerization (Table 3) of the synthesized PBMA-*qb*-PMMA-*qb*-PDEGMA quasi-triblock 5 copolymer materials derived from macro-RAFT A) precursor agent **7** and B) precursor agent **8**.

conversion and minimal dead polymer chains, optimization experiments could be performed using our high-throughput 10 methodology to establish the minimum amount of initiator required to reach a desired conversion and level of end group fidelity.<sup>5,10</sup> This optimal initiator level might also be estimated by simulations<sup>11,16</sup> for systems where kinetic parameters are available. 2b As a direct consequence of the relatively low 15 conversion in the first chain extension reaction (polymerization of MMA), higher amounts of MMA monomer will be incorporated into the PDEGMA block as "impurity" during the second chain extension reaction. Based on this and using the <sup>1</sup>H-NMR analysis, the amount of MMA incorporated into the PDEGMA blocks 20 during the third polymerization step was estimated to be in the range of 18 to 27 mol %. Similar to the previous discussed case, the relatively high impurity found in the PDEGMA block can be ascribed to the low MMA monomer conversion during the second polymerization and the relatively low concentration of DEGMA 25 monomer utilized for the third polymerization step.





**Figure 6.** Kinetic data of the synthesized PBMA-qb-PMMA-qb-PDEGMA quasi-triblock copolymer materials derived from AB macro RAFT agent **7** (Table 3).  $M_n$  and D of as a function of the DEGMA conversion (top) and DEGMA conversion as a function of reaction time (55 (bottom)).

For the PBMA-qb-PMMA-qb-PDEGMA-qb-PBzMA ABCD case, the ABC macro-RAFT agent was sequentially chain extended with BzMA monomer in a third polymerization step. 70 This resulted in 8 PBMA-qb-PMMA-qb-PDEGMA-qb-PBzMA quasi-tetrablock copolymers. Table 4 summarizes the synthetic results and the reaction conditions utilized (see Table 4 footnote D) of this library and shows that all the materials had D values below or at 1.46, whereas Figure 7 displays representative SEC 75 traces demonstrating an efficient third chain extension process for this case. Figure 8 displays kinetic plots of the chain extension reaction of these cases (ABC macro-RAFT agent 9 in Table 4) where a linear relationship can be observed between the  $M_n$  vs. conversion (x) and the  $-\ln(1-x)$  vs. reaction time indicating good 80 control over the consecutive polymerizations. However, after three consecutive polymerization reactions the D values of the materials become higher with tailing to low molar mass evident in the SEC traces due to the unavoidable contribution of initiatorderived dead chains from the necessary use of relatively high 85 initiator concentrations.

Furthermore, similar to the previous analysis, full conversion was also not reached during the third polymerization step (synthesis of the ABC macro-RAFT agent). Thus, the residual DEGMA was incorporated within the PBzMA blocks of the PBMA-qb-PMMA-qb-PDEGMA-qb-PBzMA quasi-tetrablock copolymers. The DEGMA monomer conversion for ABC macro-RAFT agent of Table 4 was 70%. Attempts to obtain higher conversion (> 70%) in these polymerization reactions led to the appearance of a considerable amount of dead chains evident strough SEC traces (see Figure S6 in the ESI for an example where a lower molar mass ABC macro-RAFT agent was obtained

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at the level of 74% conversion). As a direct consequence of the relatively low conversion in the second chain extension reaction (polymerization of DEGMA), higher amounts of DEGMA monomer will be incorporated to the PBzMA block as "impurity" 5 during the third chain extension reaction. Based on these findings and using the <sup>1</sup>H-NMR analysis described in the experimental section, the amount of DEGMA incorporated into the PBzMA blocks during the fourth polymerization step was estimated. It was found that the PBMA-qb-PMMA-qb-PDEGMA-qb-PBzMA 10 quasi-tetrablock copolymers have DEGMA units within the PBzMA blocks in the range of 22 to 35 mol %. Similar to the previous discussed case, the relatively high impurity found in the PBzMA block can be ascribed to the relative lower DEGMA monomer conversion obtained during the third polymerization as 15 well as to the relative low concentration of BzMA monomer utilized for the fourth polymerization step.

**Table 4.** Characteristics PBMA-qb-PMMA-qb-PDEGMA-qb-BzMA materials obtained from the chain extension of a macro-RAFT agent in 20 the automated parallel synthesizer. D

**9.** ABC macro-RAFT agent  $(M_n = 39600 \text{ g mol}^{-1}, D = 1.28)$ Reaction BzMA  $M_{\rm n}$  $M_{\text{n(theory)}}$ ID Đ time (h) (g mol<sup>-1</sup>) (g mol<sup>-1</sup>) conversion (%) 9A 1.34 50411 15 42900 29 2 9B 44700 1.36 53767 3 36 9C 46100 1.37 55615 4 43 9D 47300 1.39 57413 9E 6 48200 1.43 60183 54 9F 1.46 67 49600 63876 10 1.45 64955 70 9G 52000

 $^{\rm D}$  Number average molar mass (Mn) and dispersity (D = Mw / Mn) were 25 estimated by SEC and are reported as PMMA equivalents. The monomer to polymer conversion was determined by  ${}^{1}H$ -NMR.  $M_{n(theory)}$  was estimated using the formula:  $M_{\text{n(theory)}} = ([M_{\text{BMA}}]_{\text{o}} \times M_{\text{BMA}} \times \%$  $conversion_{BMA} \, + \, [M_{MMA}]_o \, \times \, M_{MMA} \, \times \, \% \ conversion_{MMA} \, + \, [M_{DEGMA}]_o \, \times \,$  $M_{\text{DEGMA}} \times \% \ conversion_{\text{DEGMA}} + [M_{\text{BzMA}}]_o \times M_{\text{BzMA}} \times \% \ conversion_{\text{BzMA}})) \ /$  $_{30}$  [RAFT]  $_{o}$  +  $M_{RAFT.}$   $M_{BMA},\,M_{MMA,}\,M_{DEGMA,}\,M_{BzMA}$  and  $M_{RAFT}$  are the molar masses of BMA, MMA, DEGMA, BzMA and RAFT agent, respectively. [M<sub>BMA</sub>]<sub>o</sub>, [M<sub>BMA</sub>]<sub>o</sub>, [M<sub>DEGMA</sub>]<sub>o</sub>, [M<sub>BzMA</sub>]<sub>o</sub> and [RAFT]<sub>o</sub> are the initial concentrations of BMA, MMA, DEGMA, BzMA and RAFT agent, respectively. The synthesis of the ABC macro RAFT agent was an 35 extension of 7 in Table 3,  $[M_{DEGMA}]_o = 0.853$  M, additional [Initiator] $_o =$  $6.638 \times 10^{-4}$  M, reaction temperature = 85 °C and reaction time = 12 h. For the chain extension reaction of this ABC macro RAFT agent with BzMA,  $[M_{BzMA}]_0 = 0.682$  M, additional [Initiator]<sub>0</sub> =  $5.848 \times 10^{-4}$  M, reaction temperature = 85 °C and reaction time = 10 h.

Hadjiantoniou et al. synthesized a series of methacrylate based "pure" block copolymers in consecutive polymerization steps utilizing the RAFT technique. 10e In specific, they synthesized di-, tri, tetra- and pentablock copolymers and obtained D values of 45 1.32, 1.48, 1.58 and 1.83, respectively. The results reported in this contribution clearly demonstrate that properly optimized one pot RAFT synthetic approaches can yield multi-block copolymers with lower D values as compared to other more demanding methods where intermediate purification steps are applied for the 50 synthesis of each block.

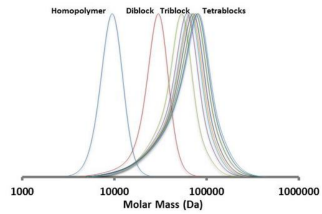
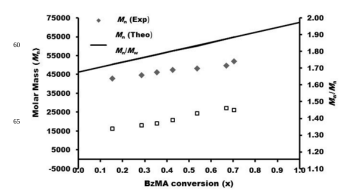


Figure 7. SEC trace of the chain extension RAFT polymerization (Table 4) of the synthesized PBMA-qb-PMMA-qb-PDEGMA-qb-PBzMA quasi-55 tetrablock copolymer materials derived from macro-RAFT precursor agent 9.



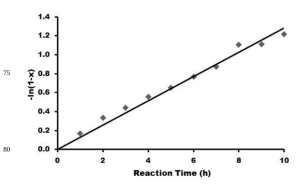


Figure 8. Kinetic data of the synthesized PBMA-qb-PMMA-qb-PDEGMA-qb-BzMA quasi-tetrablock copolymer materials derived from 85 ABC macro-RAFT agent 9 (Table 4).  $M_n$  and D of as a function of the BzMA conversion (top) and BzMA conversion as a function of reaction time (bottom).

### Conclusion

90 In this work, we have developed a convenient high-throughput approach for the one pot synthesis of quasi-block copolymer libraries of methacrylic monomers by RAFT polymerization. Utilizing this powerful technique, we could optimize reaction conditions and synthesize up to 71 different quasi-block 95 copolymers, including diblocks, triblocks, tetrablocks and pentablocks, demonstrating that highly comprehensive and

systematic polymer libraries can be obtained in a short period of time. Future efforts in this direction will include the synthesis of materials that utilize monomers of varying reactivity to further expand the range of materials that are easily accessible through 5 process automation.

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# **Notes and references**

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  - † Electronic supplementary information (ESI) available: Additional <sup>1</sup>H-NMR spectra, integration methods, SEC traces and kinetic plots of the discussed experiments See DOI:
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