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Effect of radical copolymerization of the (oxa) norbornene end-group of RAFT-prepared macromonomers on bottlebrush copolymer synthesis via ROMP†

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Bottlebrush polymers are attractive for use in a variety of different applications. Here we report the synthesis of two novel trithiocarbonate RAFT agents bearing either an oxanorbornenyl or norbornenyl moiety for bottlebrush synthesis *via* ROMP grafting-through polymerization. RAFT polymerization kinetics was evaluated as a function of the monomer type, number-average degree of polymerization (X_n) and RAFT agent structure. The correlation between the oxa/norbornenyl moiety and the type of RAFT monomer (methyl acrylate, *n*-butyl acrylate, and styrene) has been investigated. The reactivity of the oxa/norbornenyl group of the RAFT agent towards the radical propagating species during RAFT polymerization influences the molar mass, molar mass distribution and the residual olefinic end-group functionality of the resulting polymeric macromonomers. The RAFT synthesized macromonomers (MMs) are subjected to "grafting-through" ROMP using a Grubbs 3rd generation catalyst, resulting in bottlebrush polymers. The 'defects' in the MM structures have been found to be responsible for the higher amount of MM residue after the ROMP process and hence affect the microstructures of the synthesized bottlebrush polymers.

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Introduction

Due to their outstanding and tunable properties, molecular bottlebrush polymers have attracted great interest in the last few years.^{1–3} Molecular bottlebrushes consist of long linear polymeric backbones densely grafted with short side-chains. Based on their well-defined structures and fascinating properties they can be functionalized and utilized in many applications such as coatings, lubrication, nanomedicine, drug delivery, and photonics.^{4–6}

There are three main approaches for the preparation of bottlebrush polymers: (1) 'grafting-to' – attachment of side chains to the polymeric backbone; (2) 'grafting-from' – polymerization of monomers from the backbone; and (3) 'grafting-through' – polymerization of macromonomers.⁷ Each method has its own advantages and limitations. Generally, these well-defined bot-

tlebrushes are synthesised *via* a combination of two or more polymerization techniques.^{8,9}

The grafting-through strategy is the polymerization of macromonomers (MMs) that contain polymerizable end-groups. The grafting density and dispersity play a crucial role in the performance of these bottlebrush polymers. Unlike the other two strategies, the grafting-through technique ensures high grafting density as well as low brush dispersity and hence extraordinary properties and high efficacy toward applications. Often, ring-opening metathesis polymerization (ROMP), using Grubbs-type Ru catalysts, has been utilized for the grafting-through technique due to its rapid polymerization rates. In addition, the molar mass of MMs and the reactivity of polymerizable moieties are key factors in the grafting-through strategy.^{1,10}

For the preparation of functional MMs with tailored molecular properties (*i.e.* targeted molar mass, low dispersity) reversible deactivation radical polymerization (RDRP) techniques are highly attractive. Reversible addition–fragmentation chain transfer (RAFT) polymerization is arguably the most versatile RDRP method as it has a superior tolerance for a wide range of functional groups;¹¹ the RAFT technique is compatible with non-ionic, cationic, and anionic monomers. Through careful selection of the RAFT agent and reaction conditions, MMs amenable for polymerization by "grafting-through" ROMP can be prepared.^{1,10,12}

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In the broader scientific literature *exo*-norbornenes are the cyclic olefins (monomers) of choice for ROMP; their rapid ring opening metathesis kinetics and low incidence of chain transfer render their polymerization 'living'.¹³ An added advantage of norbornenes is their thermal stability. While being less commonly used, *exo*-oxanorbornenes (*i.e.* oxygen-bridged analogues of norbornenes) are also widely reported in ROMP.⁷

In the context of functional RAFT agent synthesis, *exo*-oxanorbornene derivatives are quite attractive starting materials as they are relatively inexpensive. In contrast, *exo*-norbornenes tend to be quite expensive and are often prepared 'in-house' by laborious isomerization methods from the cheaper *endo*-isomers. Whilst this suggests oxanorbornenes as an attractive alternative to norbornenes, for the synthesis of 'ROMP-able' RAFT agents, their main drawback is their thermal lability; they can readily undergo retro-Diels–Alder reactions extruding furan.¹⁴

Herein, we directly compare the utility of oxanorbornene- and norbornene-based trithiocarbonate RAFT agents for the preparation of bottlebrush polymers *via* sequential RAFT/ROMP. The influence of the identities of the RAFT polymerizable monomer (*i.e.* methyl acrylate (MA), *n*-butyl acrylate (BA), and styrene (St)) and the strained olefin end-group (*i.e.* oxanorbornenyl, norbornenyl) on macromonomer (MM) synthesis is investigated in detail. While both norbornene^{1,12,15–17} and oxanorbornene^{5,18} end-groups have been used for the synthesis of polymers *via* a sequential RAFT/ROMP strategy (or other RDRP/ROMP methods) in the past, the incidence and effect of radical propagation to the end-group has largely been ignored and a direct comparison is lacking; here we seek to remedy these points. Additionally, the effect of the resultant MM structure on the subsequent ROMP grafting-through polymerization is also investigated.

Experimental

Materials

2-Bromopropionyl bromide, carbon disulfide (CS₂), chlorobenzene, dodecanethiol, ethanolamine, *n*-heptane, neutral alumina Brockmann activity I (70–230 mesh), *cis*-5-norbornene-*endo*-2,3-dicarboxylic anhydride, potassium *tert*-butoxide, pyridine, triethylamine (TEA) and all solvents were purchased from Fisher Scientific. Solvents were of analytical reagent grade unless otherwise stated. *exo*-3,6-Epoxy-1,2,3,6-tetrahydrophthalic anhydride **1**, a Grubbs 3rd generation catalyst (**G3**) [dichloro[1,3-bis(2,4,6-trimethylphenyl)-2-imidazolidinylidene] (benzylidene)bis(3-bromopyridine) ruthenium(II)], and anhydrous sodium sulfate (Na₂SO₄) were obtained from Sigma-Aldrich. Methyl acrylate (MA), *n*-butyl acrylate (BA) and styrene (St) were purchased from Sigma-Aldrich and freed from inhibitors by passage through neutral alumina activity I (70–230 mesh). 2,2'-Azobis[2-methylpropionitrile] (AIBN) was purchased from Acros and purified by recrystallization twice from methanol prior to use. All NMR solvents were produced by Cambridge Isotope Laboratories and obtained through

Goss Scientific. Silica gel (ZEO prep 60 HYD 40–63 µm) was obtained from Apollo Scientific. Hydrochloric acid was purchased from Better Equipped.

Characterization

Nuclear magnetic resonance (NMR) spectra were recorded on a Jeol 400 MHz spectrometer at room temperature. ¹H and ¹³C NMR spectra were internally referenced to the residual solvent.¹⁹ Size exclusion chromatography (SEC) was conducted on an EcoSEC-HLC 8320GPC system with a dual flow RI detector and a TSKgel Super HZM-N 3 µm (4.6 × 150 mm) column. THF was used as the eluent at a flow rate of 0.35 mL min^{−1} at 40 °C and low dispersity polystyrene standards were used for the calibration.

Synthesis

Preparation of alcohol-functional oxanorbornene imide (3). To a mixture of *exo*-3,6-epoxy-1,2,3,6-tetrahydrophthalic anhydride **1** (15 g, 90 mmol) and methanol (375 mL), ethanolamine (6.6 mL, 6.53 g, 107 mmol, 1.2 equiv.) and trimethylamine (12.6 mL, 9.15 g, 90 mmol, 1 equiv.) were added. The reaction mixture was heated under reflux at 70 °C for 24 h. The solution was concentrated to half the volume under reduced pressure at ambient temperature, which upon crystallisation in the freezer (−18 °C) gave oxanorbornene imide **3** as white crystals (13.0 g, 62.1 mmol, 69%); ¹H NMR (400 MHz, CDCl₃) δ 2.92 (s, 2H, 2 × *endo*-CH), 3.89 (m, 4H, NCH₂ and OCH₂), 4.78 (br t, 1H, OH), 5.12 (s, 2H, 2 × bridgehead-CH), 6.55 (s, 2H, 2 × =CH). These data are in agreement with those of Kötteritzsch *et al.*²⁰

Preparation of the oxanorbornene alkylating agent (5). To a solution of *N*-(2-hydroxyethyl)-*cis*-5-oxanorbornene-*exo*-2,3-dicarboximide **3** (13.0 g, 62.1 mmol) in anhydrous tetrahydrofuran (THF) (125 mL), dry pyridine (5.65 g, 5.8 mL, 71.4 mmol, 1.15 equiv.) was added. Subsequently, a solution of 2-bromopropionyl bromide (14.75 g, 7.16 mL, 68.3 mmol, 1.1 equiv.) in dry THF (25 mL) was added dropwise to the reaction mixture at 0 °C. The reaction mixture was allowed to stir for 24 h at RT under nitrogen. The resulting reaction mixture was poured into a dilute aq. HCl solution (350 mL, 0.1 M) and was extracted with DCM (3 × 100 mL), and the combined organics was washed with dilute aq. HCl solution (100 mL, 0.1 M) and water (100 mL), and dried over anhydrous sodium sulfate (Na₂SO₄). The solvent was removed under reduced pressure giving the oxanorbornene-functional alkylating agent **5** as a greyish-white solid (20.3 g, 59.0 mmol, 95%); ¹H NMR (400 MHz, CDCl₃) δ 1.77 (d, *J* 7.0 Hz, 3H, CHCH₃), 2.85 (s, 2H, 2 × *endo*-CH), 3.77 (m, 2H, NCH₂), 4.31 (m, 3H, OCH₂ and BrCH), 5.24 (d, *J* 0.6 Hz, 2H, 2 × bridgehead-CH), 6.49 (d, *J* 0.6 Hz, 2H, 2 × =CH); ¹³C NMR (50 MHz, CDCl₃) δ 21.5, 37.6, 39.9, 47.5, 62.0, 80.9, 136.6, 170.0, 176.0. These data are in agreement with those of Runge and Bowden.²¹

Preparation of the oxanorbornene RAFT agent (7-ONb). Dodecane-1-thiol (6.36 g, 31.5 mmol, 1 equiv.) was added to dry THF (100 mL). After 10 min potassium *tert*-butoxide (KO'Bu) (3.54 g, 31.5 mmol, 1 equiv.) was added to the solution and kept under stirring for 15 min at ambient temperature.

Then, carbon disulfide (CS_2) (4.8 g, 63 mmol, 2 equiv.) was added and stirred for 30 min. Subsequently, the oxanorbornene-functional alkyl bromide **5** (10.89 g, 31.5 mmol, 1 equiv.) was added to the reaction mixture and allowed to stir for 20 hours at room temperature. Afterwards, the reaction mixture was diluted with water (200 mL), extracted with DCM (3×75 mL), washed with brine and dried over anhydrous sodium sulfate. The solvent was reduced *in vacuo* at ambient temperature and precipitated into 50 mL of *n*-heptane. The resulting yellow precipitate was isolated by filtration and dried under reduced pressure at ambient temperature to give the RAFT agent **7-ONb** (10.0 g, 18.5 mmol, 59%); ^1H NMR (400 MHz, CDCl_3) δ 0.87 (t, J 6.7 Hz, 3H, CH_3), 1.20–1.30 (m, 16H, alkyl CH_2), 1.37 (m, 2H, $\text{SCH}_2\text{CH}_2\text{CH}_2$), 1.56 (d, J 7.3, 3H, CHCH_3), 1.67 (p, J 7.4 Hz, 2H, SCH_2CH_2), 2.85 (s, 2H, 2 \times *endo*-CH), 3.33 (m, 2H, SCH_2), 3.76 (m, 2H, NCH_2), 4.28 (m, 2H, OCH_2), 4.79 (q, J 7.3, 1H, CHCH_3), 5.26 (d, J 0.6 Hz, 2H, 2 \times bridgehead-CH), 6.50 (d, J 0.6 Hz, 2H, 2 \times =CH); ^{13}C NMR (50 MHz, CDCl_3) δ 14.2, 16.8, 22.8, 27.9, 29.0, 29.2, 29.4, 29.5, 29.6, 29.7, 29.7, 32.0, 37.4, 37.7, 47.6, 47.8, 62.0, 77.3, 136.6, 171.0, 176.0, 222.1.

Synthesis of *exo*-5-norbornene-2,3-dicarboxylic anhydride (2) via thermal isomerisation. The isomerization of *endo*-5-norbornene-2,3-dicarboxylic anhydride to *exo*-5-norbornene-2,3-dicarboxylic anhydride **2** was adapted from a procedure reported by Yu *et al.*²² Briefly, *endo*-5-norbornene-2,3-dicarboxylic anhydride (20.0 g, 121.8 mmol) was heated under a nitrogen atmosphere at 200 °C for 12 h. The resulting material was recrystallized three times from toluene to give the *exo*-5-norbornene-2,3-dicarboxylic anhydride **2**; the residue can be recycled and treated in the same manner several times to increase the yield (after three cycles; 10 g, 60.9 mmol, 50%; isomeric ratio 95% *exo*/5% *endo*). ^1H NMR (400 MHz, CDCl_3) δ 1.44 (d, J 10.3 Hz, 1H, CHH), 1.66 (dm, J 10.3 Hz, 1H, CHH), 2.99 (m, 2H, 2 \times *endo*-CH), 3.45 (br s, 2H, 2 \times bridgehead-CH), 6.32 (br s, 2H, 2 \times =CH). These data are in agreement with those of Matson and Grubbs.²³

Preparation of *N*-(2-hydroxyethyl)-*cis*-5-norbornene-*exo*-2,3-dicarboximide (4). To a mixture of *exo*-5-norbornene-2,3-dicarboxylic anhydride **2** (10.0 g, 61 mmol) and toluene (100 mL), ethanolamine (4.06 mL, 67 mmol, 1.1 equiv.) and triethylamine (0.92 mL, 6.8 mmol, 0.11 equiv.) were added. A Dean–Stark trap was attached and the reaction mixture was refluxed at 125 °C for 20 h. The resulting mixture was concentrated and DCM (100 mL) was added. The combined organics was washed with aq. HCl (1 M, 30 mL) and brine (30 mL). The organic layer was dried over anhydrous sodium sulfate and the solvent was removed under reduced pressure to give *N*-2-(hydroxyethyl)-*cis*-5-norbornene-*exo*-2,3-dicarboximide **4** as a white solid (11.4 g, 54.5 mmol, 90%); ^1H NMR (400 MHz, CDCl_3) δ 1.32 (br d, J 9.9 Hz, 2H, CHH), 1.50 (dm, J 9.9 Hz, 1H, CHH), 2.19 (s, 1H, OH), 2.69 (d, J 1.2 Hz, 2H, 2 \times *endo*-CH), 3.26 (m, 2H, 2 \times bridgehead-CH), 3.67 (m, 2H, NCH_2), 3.76 (m, 2H, OCH_2), 6.27 (t, J 1.7 Hz, 2H, 2 \times =CH); ^{13}C NMR (50 MHz, CDCl_3) δ 41.4, 42.9, 45.3, 48.0, 60.3, 137.9, 178.9. These data are in agreement with those of Matson and Grubbs.²³

Preparation of the norbornene alkylating agent (6). *N*-(Hydroxyethyl)-*cis*-5-norbornene-*exo*-2,3-dicarboximide **4** (9 g, 43 mmol) was added to anhydrous THF (100 mL), followed by pyridine (4.2 mL, 4.1 g, 52 mmol, 1.2 equiv.). Then, 2-bromo-propionyl bromide (5.4 mL, 11.1 g, 52 mmol, 1.2 equiv.) in THF (20 mL) was added dropwise to the reaction mixture at 0 °C. The reaction mixture was allowed to stir for 48 h at RT under nitrogen. Then, the solvent was reduced and DCM (100 mL) was added. The combined organics was washed with aq. HCl (0.1 M, 30 mL) and brine (30 mL). The organic layer was dried over anhydrous sodium sulfate and concentrated to give the norbornene-functional alkylating agent **6** as a viscous oil (12.3 g, 35.9 mmol, 83%); ^1H NMR (400 MHz, CDCl_3) δ 1.28 (d, J 10.0 Hz, 1H, CHH), 1.50 (d, J 10.0 Hz, 1H, CHH), 1.77 (d, J 7.0 Hz, 3H, CHCH_3), 2.68 (s, 2H, 2 \times *endo*-CH), 3.25 (s, 2H, 2 \times bridgehead-CH), 3.76 (m, 2H, NCH_2), 4.31 (m, 3H, OCH_2 and BrCH), 6.26 (s, 2H, 2 \times =CH); ^{13}C NMR (50 MHz, CDCl_3) δ 20.4, 36.2, 38.6, 41.8, 44.2, 46.8, 61.2, 136.7, 168.8, 176.7.

Synthesis of the norbornene RAFT agent (8-Nb). Dodecane-1-thiol (4.93 mL, 20.5 mmol, 1 equiv.) was added to dry THF (80 mL). After 10 min potassium *tert*-butoxide (KO^tBu) (2.3 g, 20.5 mmol, 1 equiv.) was added to the solution and kept under stirring for 15 min at ambient temperature. Then, CS_2 (2.5 g, 32.3 mmol, 1.6 equiv.) was added and stirred for 30 min. Subsequently, the alkylating agent **6** (7.0 g, 20.5 mmol, 1 equiv.) in dry THF (10 mL) was added to the reaction mixture and allowed to stir for 20 hours at room temperature. Afterwards, the reaction mixture was diluted with water (200 mL), extracted with DCM (3×50 mL), washed with brine and dried over anhydrous sodium sulfate. The solvent was reduced *in vacuo* at ambient temperature. The crude product was purified by column chromatography (silica gel; ethyl acetate/*n*-hexane (1 : 2)). Removal of the solvent under reduced pressure gave the norbornene-functional RAFT agent **8-Nb** as a yellow solid (6.1 g, 11.3 mmol, 55%); ^1H NMR (400 MHz, CDCl_3) δ 0.86 (t, J 6.8 Hz, 3H, CH_3), 1.20–1.30 (m, 16H, alkyl CH_2), 1.29 (d, J 9.7 Hz, 1H, CHH), 1.37 (m, 2H, $\text{SCH}_2\text{CH}_2\text{CH}_2$), 1.51 (m, 1H, CHH), 1.56 (d, J 7.3 Hz, 3H, CHCH_3), 1.66 (p, J 7.3 Hz, 2H, SCH_2CH_2), 2.69 (d, J 1.2 Hz, 2H, 2 \times *endo*-CH), 3.28 (m, 2H, 2 \times bridgehead-CH), 3.33 (t, J 7.3 Hz, 2H, SCH_2), 3.76 (m, 2H, NCH_2), 4.38 (m, 2H, OCH_2), 4.80 (q, J 7.3, 1H, CHCH_3), 6.28 (t, J 1.8 Hz, 2H, 2 \times =CH); ^{13}C NMR (50 MHz, CDCl_3) δ 14.2, 16.8, 22.8, 28.0, 29.0, 29.2, 29.4, 29.6, 29.7, 32.0, 37.4, 37.5, 43.0, 45.4, 47.6, 48.0, 62.1, 137.9, 170.9, 177.8, 221.9.

RAFT polymerization

RAFT polymerization of methyl acrylate and butyl acrylate. Methyl acrylate (MA) or *n*-butyl acrylate (BA) (60 wt% in toluene), the oxanorbornene RAFT agent **7-ONb** or the norbornene RAFT agent **8-Nb**, and AIBN in a ratio of 200 : 1 : 0.1 or 50 : 1 : 0.1 were mixed in a 25 mL round bottomed flask (RBF), and the resulting solution was degassed by sparging with nitrogen for 30 min. The solution polymerization was initiated by increasing the temperature to 60 °C. For kinetic studies, an

aliquot of the reaction mixture (0.3 mL) was taken at predetermined times and quenched by rapid cooling in liquid nitrogen. The polymer was recovered by precipitation three times in a methanol/water solution.

RAFT polymerization of styrene. RAFT polymerization of styrene was performed in bulk. Styrene (St), the oxanorbornene RAFT agent **7-ONb** or the norbornene RAFT agent **8-Nb**, and AIBN in a ratio of 250:1:0.1 or 50:1:0.1 were mixed in a 25 mL RBF, and the resulting solution was degassed by sparging with nitrogen for 30 min. The polymerization was initiated by increasing the temperature to 65 °C. For kinetic studies, an aliquot of the reaction mixture (0.3 mL) was taken at predetermined times and quenched by rapid cooling in liquid nitrogen. The polymer was recovered by precipitation three times in methanol.

Grafting through polymerization *via* ring opening metathesis polymerization

Preparation of macromonomers. Defined macromonomers with low molar masses ($\sim 2000\text{--}4000\text{ g mol}^{-1}$) derived from both RAFT agents (**7-ONb** and **8-Nb**) were prepared as described above ($M:\text{RAFT}:I = 50:1:0.1$) and the polymerization was quenched after a certain time to obtain the desirable molar mass.

ROMP of macromonomers via ‘grafting through’ ROMP. The defined macromonomer was added to a dry 5 mL RBF charged with a stir bar. The flask was then degassed by applying vacuum for 30 min, and the desired amount of degassed, anhydrous THF was added (the total macromonomer concentration was ~ 0.03 M). The required amount of the degassed Grubbs catalyst G3 solution was transferred to the reaction flask containing the macromonomer to initiate the polymerization and stirred at room temperature for at least 3 h. The reaction was quenched by the addition of a few drops of ethyl vinyl ether. The product was collected by precipitation in methanol and dried under vacuum.

Results and discussion

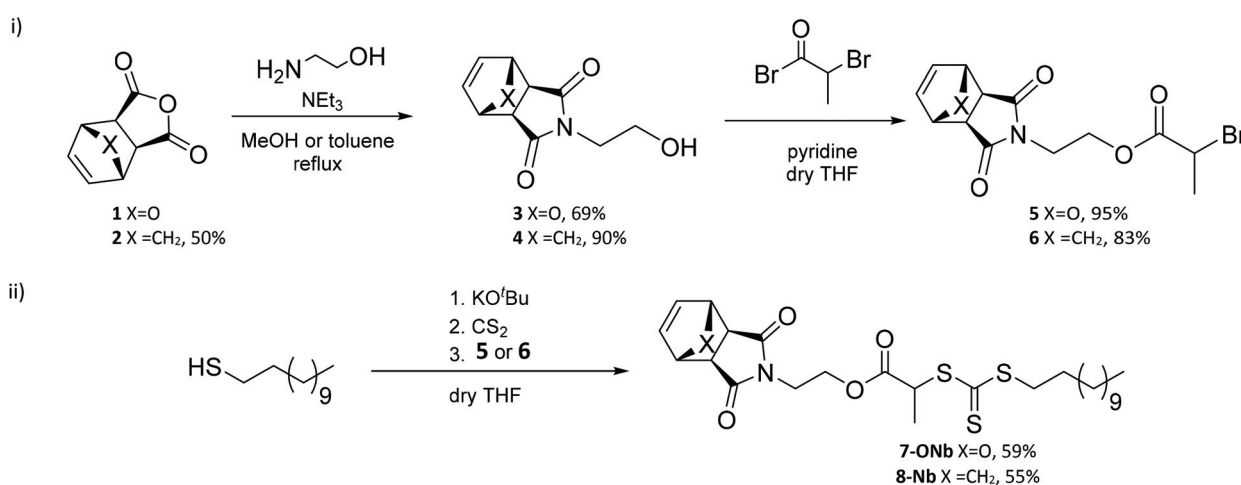
RAFT agent design and synthesis

To probe the differences in the performance between oxanorbornene and norbornene strained olefinic groups in both (a) RAFT-based macromonomer (MM) synthesis and (b) ROMP-based bottlebrush polymer synthesis *via* grafting-through polymerization, two RAFT agents were prepared; these were the oxanorbornenyl RAFT agent **7-ONb** and the norbornenyl RAFT agent **8-Nb** (see Scheme 1). Briefly, the alcohol functional imides **3** and **4** were obtained in moderate to high yields from the relevant *exo*-anhydrides **1** and **2** by treatment with ethanolamine as per published literature procedures.^{20,23} The subsequent reaction of alcohols **3** and **4** with 2-bromopropionyl bromide adapted from the procedure of Keddie *et al.*²⁴ delivered the alkylating agents **5** and **6**. The RAFT agents **7-ONb** and **8-Nb** were isolated in moderate yields by following standard RAFT agent syntheses,²⁵ *i.e.* by alkylation of the carbodithioate salt derived from dodecanethiol and CS₂.²⁶

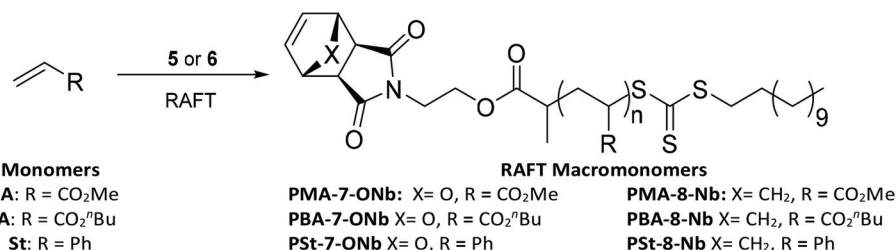
Kinetic analysis of RAFT polymerization

Three monosubstituted monomers of interest, namely, methyl acrylate (MA), *n*-butyl acrylate (BA), and styrene (St), were selected for the preparation of macromonomers *via* RAFT polymerization (see Scheme 2). Importantly, these monomers allow us to probe the effect of electronics and/or sterics in the preparation of (oxa)norbornenyl-functional macromonomers.

MA was the first monomer investigated. Initially, we targeted a number-average degree of polymerization (X_n) of 50 (i.e. $[\text{MA}]:[\text{RAFT}] = 50:1$), using the RAFT agents **7-ONb** (see Table 1, entry 1, and Fig. 1(a and b)) or **8-Nb** (see Table 1, entry 2, and Fig. 1(c and d)). From the SEC data, it can be clearly observed that the oxanorbornene RAFT agent **7-ONb** delivered polymers of higher molar masses and higher dispersity (see Table 1, entry 1, and Fig. 1(b)) than those of the analogous norbornene RAFT agent **8-Nb** (see Table 1, entry 2, and



Scheme 1 Synthesis of (i) alkylating agents 5 and 6 and (ii) RAFT agents 7-ONb and 8-Nb.



Scheme 2 RAFT polymerization of MA, BA and St using RAFT agents 7-ONb and 8-Nb

Table 1 Details of polymers prepared via RAFT Polymerization

Entry ^a	Monomer	[M] (mol L ⁻¹)	RAFT agent	T (°C)	Time (h)	M conv. ^b (%)	M _n ^c (calc)	M _n ^d	D ^d	L% (ω-end) ^e	DB% (NMR) ^{b,f}	DB % (SEC) ^{f,g}
1	MA	6.79	7-ONb	60	0.5	32	1920	2200	1.19	99.9	7	7.3
					1.5	69	3510	4500	1.37	99.6	24	24.3
					2.5	80	3980	5900	1.49	99.4	37	26.5
					3.5	85	4200	7000	1.64	99.2	42	32.7
					5	92	4500	7900	1.80	98.9	47	35.4
2	MA	6.79	8-Nb	60	0.5	19	1360	1100	1.12	99.9	4	0
					1.5	57	2990	3200	1.13	99.6	10	4.1
					2.5	76	3810	4400	1.17	99.4	13	9.7
					3.5	85	4200	5000	1.22	99.2	15	14.2
					5	90	4410	5500	1.25	98.9	15	19.2
3	BA	4.68	7-ONb	60	0.5	34	2720	2900	1.19	99.9	17	3.4
					1.5	68	4900	6300	1.33	99.6	33	35.3
					2.5	78	5540	7800	1.43	99.4	40	41.6
					3.5	84	5920	8900	1.51	99.2	45	41.6
					4.5	87	6120	9400	1.60	99.0	47	39.2
					6	90	6310	10 300	1.69	98.7	49	44.5
4	BA	4.68	8-Nb	60	0.5	23	2010	2100	1.14	99.9	5	0
					1.5	63	4580	5000	1.16	99.6	7	5.5
					2.5	77	5470	6200	1.19	99.4	10	9.9
					3.5	83	5860	6900	1.22	99.2	16	13.4
					4.5	87	6120	7400	1.24	99.0	18	15.5
					6	92	6440	7900	1.28	98.7	21	20.0
5	St	8.73	7-ONb	65	2	9	1010	—	—	99.1	7	—
					4	12	1160	1500	1.13	98.3	9	5.0
					7	31	2150	2200	1.18	97.4	15	9.4
					10	42	2730	2900	1.20	96.6	20	14.2
					22.5	68	4080	4800	1.33	94.7	34	27.9
6	St	8.73	8-Nb	65	2	11	1110	—	—	99.1	4	—
					4	21	1630	1400	1.09	98.3	5	0
					7	36	2410	2000	1.10	97.4	8	0
					10	47	2990	2500	1.09	96.6	11	0
					22.5	82	4910	4200	1.09	94.7	17	1.2

^a $[\text{M}]_0 : [\text{RAFT}]_0 : [\text{AIBN}]_0 = 50 : 1 : 0.1$. ^b Calculated from ^1H NMR. ^c $M_n(\text{calc}) = ([\text{M}]_0 - [\text{M}]_t)/([\text{RAFT}]_0) \times \text{MW}_{\text{monomer}} + \text{MW}_{\text{RAFT}}$. ^d SEC THF eluent, $T = 40^\circ\text{C}$ (data reported in polystyrene equivalents). ^e $L\% = ([\text{CTA}]_0/([\text{CTA}]_0 + df \times [\text{I}]_0 \times 1 - e^{k_d t}) \times 100\%$,^{27,28} where f is the initiator efficiency ($=0.7$),²⁹ d is the number of chains formed by radical-radical termination ($=1$),³⁰ and $k_d = 9.67 \times 10^{-6} \text{ s}^{-1}$ at 60°C (ref. 31) or $k_d = 1.95 \times 10^{-5} \text{ s}^{-1}$ at 65°C (calculated from Arrhenius parameters).³¹ ^f DB% = percentage degree of branching. ^g calculated following the deconvolution of SEC chromatograms.

Fig. 1(d)). Of note, the high percentage “livingness” ($L\%$)^{27,28} calculated from kinetic factors indicates that the high molar mass shoulder(s) observed in the SEC traces are due to ‘branching’, formed *via* the reaction of the olefinic RAFT end-groups, rather than chain-coupling *via* termination by combination (see Table 1). The degree of branching (DB%), quantification of the presence of branched polymers (*i.e.* polymer dimers, trimers, *etc.*) calculated by either NMR analysis of ole-

finic RAFT end-group consumption[‡] or SEC deconvolution, was significantly higher for **7-ONb** than for **8-Nb** (see Table 1 entries 1 and 2). This is also clearly evidenced by kinetic ana-

† The conversion of the (oxa)norbornene end-group is calculated in the same standard manner in which conversion of the vinyl monomer is achieved; the resonances from vinylic end-groups are well resolved in all cases. A representative example can be found in the ESI (Fig. S1†).

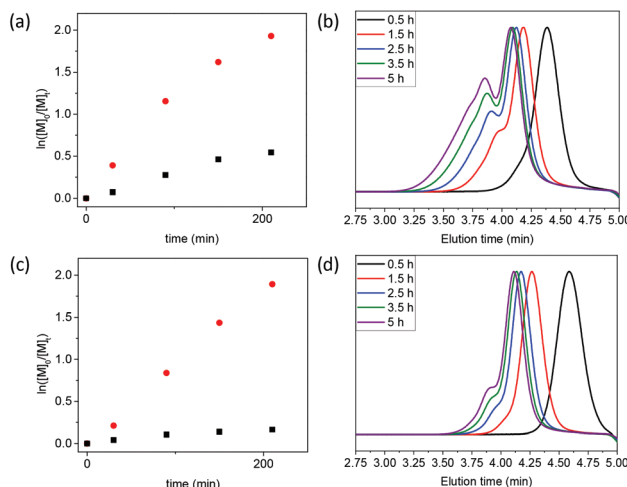


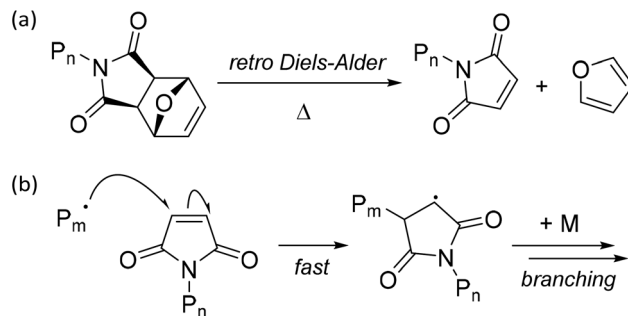
Fig. 1 Pseudo-first order kinetics plots (a and c) for the conversion of MA (red circles) and the (oxa)norbornene end-group (black squares), and SEC chromatograms illustrating the evolution of the molar mass distribution with the reaction time (b and d) for polymerizations of MA with the RAFT agents **7-ONb** (a and b) or **8-Nb** (c and d). All polymerizations were performed with $[MA] : [RAFT] = 50 : 1$.

lysis of the rate of olefinic end-group consumption during polymerization; the oxanorbornene end-group is consumed to a greater extent than that of norbornene (see Table 1, entries 1 and 2, and Fig. 1(a) and (c)). Clearly the propensity for the cross-propagation of the poly(methyl acrylate) propagating radical (PMA^\cdot) to the olefinic polymer end-group is higher for **7-ONb** based systems than those that use **8-Nb**. We postulate that this is likely due to a retro Diels–Alder extrusion of furan from **7-ONb** derived chain-ends during the reaction to produce a more reactive maleimide end-group,[§] which can then undergo rapid copolymerization with PMA^\cdot (see Scheme 3(a) and (b)). Indeed estimations of indicative copolymerization reactivity ratios, using the Alfrey-Price $Q-e$ system,^{32,33} indicate a significantly larger preference for PMA^\cdot to cross-propagate to a maleimide than to a norbornene.^{¶||} When targeting a higher chain length of $X_n = 200$ (*i.e.* $[MA] : [RAFT] = 200 : 1$) almost the same trends were observed as those for the $X_n = 50$ examples,

[§] No categorical evidence of *in situ* maleimide end-group formation could be observed in NMR analysis of the kinetic samples or of the final product polymers. We believe this is due to its rapid consumption upon its formation, in addition to the already relatively low concentration of RAFT groups in the polymerization mixture, rendering analysis of the molecular microstructure difficult. Additionally, the furan by-product was also not observed, presumably due to its volatility; all kinetic samples were taken directly from the polymerization reaction mixtures that were all at higher temperatures than the boiling point of furan (31 °C).

^{¶||} Indicative copolymerization reactivity ratios, calculated from $Q-e$ values, for norbornene systems: methyl acrylate (MA)/norbornene (Nor): $r_{MA} = 3.90$, $r_{Nor} = 0.181$; *n*-butyl acrylate (BA)/norbornene (Nor): $r_{BA} = 5.83$, $r_{Nor} = 0.146$; styrene (St)/norbornene (Nor): $r_{St} = 7.25$, $r_{Nor} = 0.045$.

^{||} Indicative copolymerization reactivity ratios, calculated from $Q-e$ values, for maleimide systems: methyl acrylate (MA)/maleimide (MI): $r_{MA} = 0.36$, $r_{MI} = 2.62$; *n*-butyl acrylate (BA)/maleimide (MI): $r_{BA} = 0.47$, $r_{MI} = 1.81$; styrene (St)/maleimide (MI): $r_{St} = 0.178$, $r_{MI} = 0.170$.



Scheme 3 (a) Proposed formation of a maleimide chain-end *via* a retro Diels–Alder reaction, and (b) cross-propagation of the propagating polymer species to the maleimide chain-end leading to branching, during RAFT polymerization reactions using the oxanorbornene RAFT agent **7-ONb**. P_n, P_m = polymer chains; M = monomer.

with the oxanorbornene-based materials displaying a higher number-average molar mass (M_n), molar mass dispersity (D) and degree of branching (DB%) (see Table S1, entries 1 and 2, and Fig. S1(a) and (b)†).

To probe the effect of the acrylate ester chain length on the RAFT system, BA was the next monomer investigated, targeting $X_n = 50$. Unsurprisingly, the additional sterics from the *n*-butyl ester of the monomer made a little difference to the reactivity of the poly(*n*-butyl acrylate) propagating species (PBA[·]) towards the different end-groups of **7-ONb** and **8-Nb** when compared to the PMA[·] systems. PBA prepared in the presence of oxanorbornene **7-ONb** displayed significantly higher M_n , D and DB% values than the PBA samples prepared in the presence of norbornene **8-Nb** (see Table 1, entries 3 and 4, and Fig. 2). Again, analogous outcomes were observed when targeting PBA of $X_n = 200$ (see Table S1, entries 3 and 4, and Fig. S1(c) and (d)†).

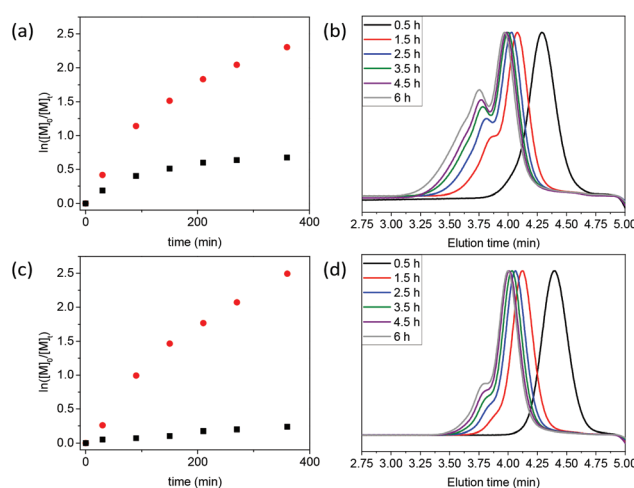


Fig. 2 Pseudo-first order kinetics plots (a and c) for the conversion of BA (red circles) and the (oxa)norbornene end-group (black squares), and SEC chromatograms illustrating the evolution of the molar mass distribution with the reaction time (b and d) for polymerizations of BA with the RAFT agents **7-ONb** (a and b) or **8-Nb** (c and d). All polymerizations were performed with $[BA] : [RAFT] = 50 : 1$.



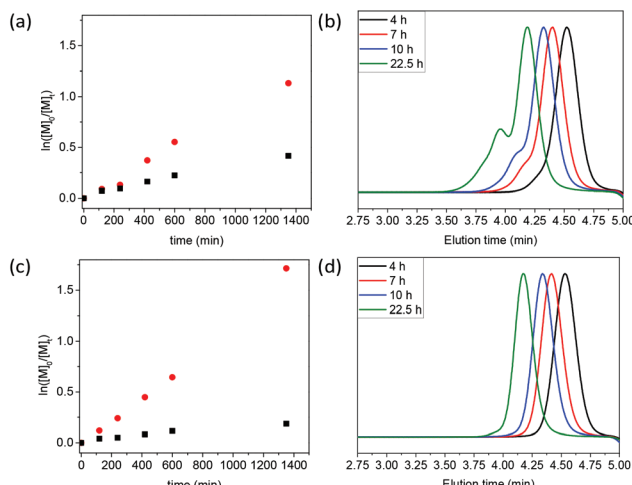


Fig. 3 Pseudo-first order kinetics plots (a and c) for the conversion of St (red circles) and the (oxa)norbornene end-group (black squares), and SEC chromatograms illustrating the evolution of the molar mass distribution with the reaction time (b and d) for polymerizations of St with the RAFT agents 7-ONb (a and b) or 8-Nb (c and d). All polymerizations were performed with $[St] : [RAFT] = 50 : 1$.

The final monomer investigated was St, again initially targeting $X_n = 50$ (*i.e.* $[St] : [RAFT] = 50 : 1$). As with the previous examples discussed above, for St polymerization the oxanorbornene RAFT agent **7-ONb** led to significantly higher M_n , D and DB% values than the norbornene RAFT agent **8-Nb** (see Table 1, entries 5 and 6, and Fig. 3). Interestingly, for the polymerization of St controlled with the norbornene RAFT agent **8-Nb** minimal branching (DB% = 1.2%) was observed, particularly when compared to the acrylate systems (*cf.* DB% $\sim 20\%$ for MA and BA at $\sim 90\%$ monomer conversion). We attributed this to the electronic differences between the electron-rich polystyryl radical (PSt^\cdot) and the electron-poor acrylate-based radicals (*i.e.* PMA^\cdot and PBA^\cdot). PSt^\cdot cross-propagates to the electron-rich norbornene end-group more slowly than do either of the acrylate-based radicals; this observation is in agreement with the indicative copolymerization reactivity ratios.^{32,33} ¶ It appears that electronics plays a less significant role in the DB% in the oxanorbornene case, which provides further indirect evidence for the contribution of retro Diels–Alder to branching. Indicative copolymerization reactivity ratios, between St and maleimide, suggest a tendency towards alternation which would lead to the consumption of a maleimide at the chain end. On the other hand an unreacted, electron-rich oxanorbornene would be expected to behave in almost the same way as norbornene in St polymerization (*i.e.* display conversion *via* cross-propagation to the chain end). Similar polymerization outcomes were obtained when targeting PSt of $X_n = 250$ (*i.e.* $[St] : [RAFT] = 250 : 1$) with **7-ONb** delivering materials with higher M_n and D , and significantly higher DB% than those with **8-Nb** (see Table S1, entries 5 and 6, and Fig. S1(e) and (f)†).

In summary, cross propagation of the propagating species (PMA^\cdot , PBA^\cdot and PSt^\cdot) to the olefinic RAFT chain-end was found to occur in all cases discussed, albeit in varying degrees.

This results in a proportion of branched structures, a generally undesired topological ‘impurity’ in the final polymer sample.

This is also expected to adversely impact the preparation of the targeted bottlebrush polymers due to the (partial) consumption of the ROMP polymerizable end-group. It is clear from the results discussed above that **8-Nb** is the preferred RAFT agent for preparing macromonomers from monosubstituted monomers *via* RAFT. Significantly more ‘defects’ in the macromonomer structure are due to the trade-off of using the more easily prepared **7-ONb** instead of **8-Nb**. It should be noted that even when using the norbornene-based RAFT agent **8-Nb** in the polymerization of the acrylates MA and BA significant end-group consumption was observed (up to 20%, see Table 1, entries 2 and 4). To decrease the incidence of these ‘defects’ in the synthesis of MMs we recommend targeting higher molar masses and quenching the reaction at a lower conversion; from a copolymerization standpoint this effectively decreases the monomer feed ratio of the norbornene chain-end reducing the rate of cross-propagation.

Macromonomer synthesis *via* RAFT polymerization

Following on from the kinetic investigations described above, we successfully prepared three macromonomers based on MA, BA, and St with low molar masses (~ 2000 – 4000 g mol $^{-1}$) by RAFT polymerization using the RAFT agents (**7-ONb** or **8-Nb**). The reactions used to prepare the MMs were quenched after the desired time and purified by precipitation three times in methanol to completely remove any unreacted monomers present. $^1\text{H-NMR}$ spectroscopy and size exclusion chromatography (SEC) were used to characterize the resulting MMs. The properties of the prepared MMs are summarized in Table 2 and the SEC chromatograms are shown in Fig. 4 (black traces).

Akin to the data described above, the norbornenyl RAFT agent **8-Nb** delivered MMs with the lowest D and DB% values in the case of each monomer (see Table 2, entries 2, 4 and 6.). The oxanorbornenyl RAFT agent **7-ONb** resulted in higher dispersities (and bimodality in the molar mass distribution) (see entries 1, 3 and 5, Table 2). Compared to styrene MMs, the acrylate MMs exhibit slightly higher dispersities and DB% (see Table 2 and Fig. 4-black traces).

ROMP of macromonomers

Grafting-through polymerization *via* ROMP of the (oxa)norbornene macromonomers (see Table 2) was carried out in THF using a MM to catalyst ratio of 25 : 1 with a Grubbs third generation catalyst, **G3** (see Scheme 4 and Table 3). Exhibiting rapid initiation kinetics and high functional-group tolerance, **G3** is well known to successfully polymerize sterically hindered substrates, allowing for the synthesis of polymers with narrow molar mass distributions.

Due to the difference in thermal stability between the RAFT agents **7-ONb** and **8-Nb** and their behaviour in RAFT polymerization (*i.e.* higher DB%) as described above, it was found that the bottlebrush polymers based on **7-ONb** have higher levels of residual MMs than those prepared using **8-Nb**; ROMP of MMs

Table 2 Details of macromonomers prepared via RAFT polymerization

Entry	Macromonomer	Time (h.)	M_n (NMR) ^a (g mol ⁻¹)	M_n (SEC) ^b (g mol ⁻¹)	D^b	L (ω -end) ^c	DB % (SEC) ^{d,e}
1	PMA-7-ONb	1	2700	3300	1.24	99.8	15
2	PMA-8-Nb	1.5	3100	3600	1.15	99.6	6
3	PBA-7-ONb	1	4300	5600	1.28	99.8	20
4	PBA-8-Nb	1	3400	3900	1.14	99.8	2
5	PSt-7-ONb	10	3100	2900	1.19	96.6	1
6	PSt-8-Nb	10	2800	2500	1.08	96.6	0

^a Calculated from ¹H NMR. ^b SEC THF eluent, $T = 40$ °C (data reported in polystyrene equivalents). ^c $L\% = ([\text{CTA}]_0/[\text{CTA}]_0 + df \times [\text{I}]_0 \times 1 - e^{kd\tau}) \times 100\%$,^{27,28} where f is the initiator efficiency ($=0.7$),²⁹ d is the number of chains formed by radical–radical termination ($=1$),³⁰ and $k_d = 9.67 \times 10^{-6}$ s⁻¹ at 60 °C³¹ or $k_d = 1.95 \times 10^{-5}$ s⁻¹ at 65 °C (calculated from Arrhenius parameters).³¹ ^d DB % = percentage degree of branching. ^e Calculated following the deconvolution of SEC chromatograms.

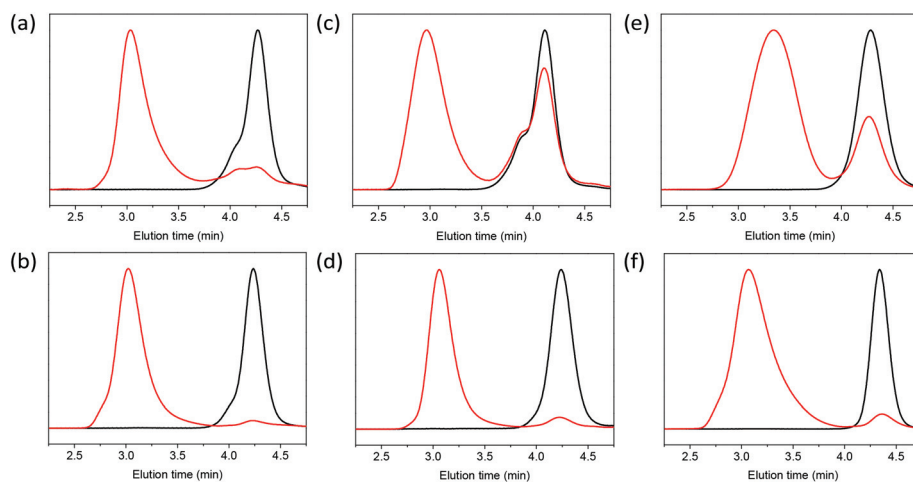
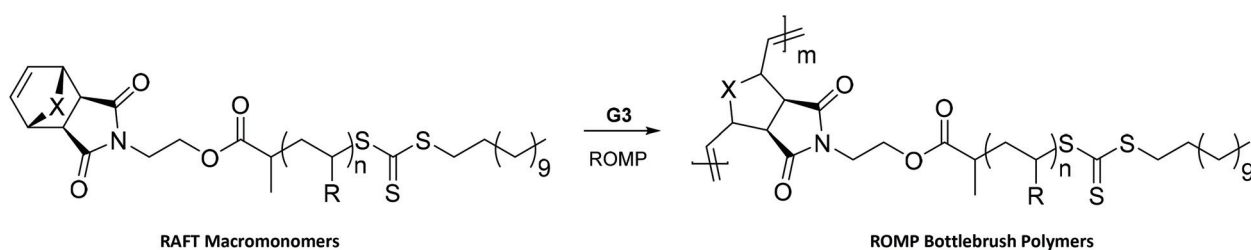


Fig. 4 SEC chromatograms of macromonomers (black) and bottlebrush polymers (red) for the polymerizations of (a) PMA-7-ONb, (b) PMA-8-Nb, (c) PBA-7-ONb, (d) PBA-8-Nb, (e) PSt-7-ONb and (f) PSt-8-Nb. Polymerizations were performed with [MM] : [G3] = 25 : 1.



Scheme 4 Grafting-through ROMP of RAFT Macromonomers (PMA-7-ONb, PBA-7-ONb, PSt-7-ONb, PMA-8-Nb, PBA-8-Nb, and PSt-8-Nb).

Table 3 Details of bottlebrush polymers prepared via ROMP

Entry ^a	Macromonomer (MM)	MM M_n ^b (g mol ⁻¹)	Residual MM %	M_n ^c (SEC) (g mol ⁻¹)	D^c
1	PMA-7-ONb	2700	11	117 000	1.29
2	PMA-8-Nb	3100	3	125 000	1.35
3	PBA-7-ONb	4300	37	162 000	1.27
4	PBA-8-Nb	3200	5	117 000	1.23
5	PSt-7-ONb	2900	20	53 000	1.40
6	PSt-8-Nb	2800	4	90 000	1.58

^a [MM] = 0.33 M in THF, [MM]/[G3] = 25, $t = 3$ h. ^b Calculated from ¹H NMR. ^c From SEC THF eluent, $T = 40$ °C (data reported in polystyrene equivalents).



based on **8-Nb** gives a bottlebrush polymer with a low amount of MM residue ($\leq 5\%$).

Bottlebrush polymers with a $[MM]/[I]$ ratio of 50:1 were prepared and full characterization was attempted, however, the polymers contained fractions that were larger than the exclusion limit of our SEC columns, limiting the ability to assess their molar masses and dispersity accurately (see Table S2 and Fig. S2 in the ES[†]). Keeping this drawback aside these materials displayed similar MM incorporation for the $[MM]/[I]$ 25:1 samples.

From these experiments it is clear that using less effective **7-ONb** in the RAFT synthesis of MMs also leads to less desirable outcomes (e.g. higher residual MM%) in the sequential RAFT/ROMP process for the preparation of bottlebrush polymers than the use of **8-Nb**. We postulate that the altered architecture of the MMs derived from **7-ONb** and the related decrease in the amount of “ROMP-able” end-groups per unit mass of MMs (both brought about by the higher incidence of cross-propagation to the olefin chain end during the RAFT syntheses) contribute to the poorer performance of oxanorbornene-based MMs in the preparation of bottlebrush polymers.

Conclusions

Two new RAFT agents with either an oxanorbornenyl or norbornenyl moiety were successfully prepared. The design of the RAFT agent has been found to affect the RAFT polymerization and hence the sequential ROMP process. Three different monomers (MA, BA, and St) were selected for RAFT polymerizations. The thermal stability of RAFT agents and the monomer electronics and sterics were found to affect the resultant MM structure and molar mass distribution.

In the case of the oxanorbornenyl RAFT agent **7-ONb**, the polymerization analysis revealed the increase of branching as a function of conversion, due to copolymerization to a maleimide end-group following thermal extrusion of furan. On the other hand, the RAFT agent **8-Nb** with the norbornenyl moiety is more thermally stable and yields MMs with low branching, due to the limited copolymerization reactivity of norbornenyl alkenes towards acrylate or styrene propagating species. The more significant “imperfections” in the **7-ONb** derived MMs (in comparison to those prepared using **8-Nb**) were found to affect the molecular properties of the bottlebrush polymers prepared *via* ROMP; NMR and SEC analyses revealed significant differences in the residual MM concentrations following polymerization.

Author contributions

MN: investigation, data analysis, drafted sections of the original manuscript, and review and editing of the manuscript; KN: data analysis, peak fitting and deconvolution of SEC data, review and editing of the manuscript; DJK: conceptualization, investigation, data analysis, drafted sections of the original manuscript, supervision, and review and editing of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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