NOTE

ATRP-RCM Synthesis of 8-Shaped Poly(methyl acrylate) Using a 4-Armed Star Telechelics

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KEY WORDS: 8-Shaped Polymer / Atom Transfer Radical Polymerization / Keck Allylation / Ring-closing Metathesis /

Cyclic and multicyclic polymers have drawn increasing attention due to their unique architectures, i.e., free of chain ends. A number of protocols including, in particular, the ring-expansion polymerization and the end-to-end polymer cyclization, have now been developed for the selective and effective synthesis of single cyclic polymers to realize novel properties and functions unattainable by conventional linear or branched counterparts.

We have also proposed a metathesis polymer cyclization (MPC) process, i.e., a ring-closing metathesis (RCM) reaction with a linear telechelic precursor having olefinic end groups under dilution. And the effective and selective synthesis of cyclic polyacrylate has been achieved through atom transfer radical polymerization (ATRP)-Keck allylation and the subsequent RCM reaction. In this process, the key telechelic prepolymer was produced by the ATRP using a difunctional initiator and the terminal bromide group was quantitatively converted to allyl group by the reaction with allyltributyltin. The subsequent intramolecular RCM reaction proceeded even under dilution in the presence of Grubbs catalyst to give a cyclic polymer in high efficiency.

On the other hand, such a dicyclic as 8-shaped polymer has yet been an ongoing challenge in the synthetic polymer chemistry. We have shown that 8-shaped polymers are obtainable through an electrostatic self-assembly formed by two units of telechelic precursor having moderately strained cyclic ammonium salt end groups carrying one unit of tetracarboxylate counter anion. Alternatively, inter- and intramolecular metathesis condensation process have been applied between two units of a cyclic prepolymer having an allyloxy group (Scheme 1-a), and with a twin-tailed tadpole polymer precursor (Scheme 1-b), as well as with a cyclic polymer precursor having two allyloxy groups at the opposite positions (Scheme 1-c), respectively. The intramolecular metathesis condensation has also been applied with an H-shaped precursor having allyloxy end groups at each chain ends, to produce other types of dicyclic polymers having theta and manacle topologies (Scheme 1-d).

As an extension of the preceding studies, we report in this paper on the synthesis of an 8-shaped polymer through the double ring-closing metathesis reaction using a 4-armed star polymer precursor, synthesized by means of the ATRP of methyl acrylate using a tetrafunctional initiator (Scheme 2). Since the ATRP has been recognized as a versatile means to give living star polymers up to hexafunctionalities by using multi-functional initiators, complex multicyclic polymers will be obtainable through the MPC process with end-functional star polymers.

EXPERIMENTAL

Materials

A tetrafunctional ATRP initiator, pentaerythritol tetrakis-(2-bromoisobutyrate), I, was synthesized by the procedure reported previously. Methyl acrylate (99.0%, Aldrich) was distilled under reduced pressure. Cuprous bromide (CuBr, Nacalai tesque), allyltri-n-butyltin (99.0%, Aldrich), Grubbs catalyst first generation (Aldrich) was used as received. Methylene chloride (99.0%, Kanto Chemical Co) was distilled over CaH2. Triethylamine (99.5%, Kanto Chemical Co) was distilled over Na wire. Alumina (Wako Chemical Co., Aluminium Oxide, activated, about 45 μm, ca. 300 mesh) and other chemicals are used as received.

Synthesis of 4-Armed Star PMA having Allyl End Groups, 2.

Methyl acrylate (560 mg, 5.6 mmol), 2,2'-bipyridine (bpy) (61 mg, 0.39 mmol), CuBr (21 mg, 0.15 mmol), tetrafunctional initiator, 1 (66 mg, 0.090 mmol) were placed in a 10 mL reaction tube, and the mixture was subjected to three freeze-pump thaw cycles. The mixture was then heated to 90 °C for 20 min, and benzene (0.5 mL) and allyltri-n-butyltin (320 mg, 0.96 mmol) were added. The reaction was allowed to continue at 90 °C under stirring for 8 h. The polymer product was recovered after the treatment with alumina, and finally purified by the precipitation twice into hexane. Yield = 362 mg.

1H NMR of 2 (CDCl3): δ 1.22–2.15 (br, CH2CH(OOC)H), 2.15–2.55 (br, CH2CH(OOC)H), 3.47–3.84 (br, CH2CH(OOC)H), 4.10 (s, CH2O), 5.04 (m, CH2CH=CH2), 5.67 (m, CH2CH=CH2).

Synthesis of 8-Shaped PMA, 3. The 4-armed star PMA, 2,
(80 mg) and methylene chloride (200 mL) were placed in a 300 mL flask. Thereupon Grubbs catalyst 1st generation (40 mg, 48/10 mmol) was added. The reaction was conducted under reflux for 48 h, and the solvent was evaporated. The polymer product was recovered after the treatment with the alumina column and finally isolated by the precipitation into hexane. Yield = 38 mg (48%). $^1$H NMR of 3 (CDCl$_3$): δ 1.22–2.15 (br, CH$_2$CHCOOCH$_3$), 2.15–2.55 (br, CH$_2$CHCOOCH$_3$), 3.47–3.84 (br, CH$_2$CHCOOCH$_3$), 4.10 (s, CCH$_2$O(CO)), 5.34 (m, CH$_2$CH=CH, cis and trans).

**Measurements**

$^1$H NMR spectra were recorded with a JEOL JNM AL300 apparatus in CDCl$_3$ at ambient temperature. The proton chemical shifts (ppm) were referenced from the signal of tetramethylsilane. MALDI-TOF MS spectra were taken on a SHIMADZU AXIMA-CFR mass spectrometer. The spectrometer was equipped with a nitrogen laser ($\lambda = 337$ nm) and with pulsed ion extraction. The operation was performed at an accelerating potential of 20 kV by a linear-positive ion mode. The sample polymer solution (10 g/L) was prepared in THF. The matrix, 1,8-dihydroxy-9(10H)anthracenone (dithranol, Aldrich) was dissolved in THF (30 g/L) and sodium trifluoroacetate was dissolved in THF (10 g/L). The polymer solution (90 μL) was then mixed with 30 μL of matrix solution. A 1 μL portion of the final solution was deposited onto a sample target plate and allowed to dry in air at room temperature. Mass values were calibrated by the three-point method with insulin $\beta$ (Sigma-Aldrich) at 3497.0 (plus H$^+$), bovine insulin (Aldrich) at 5733.5 (plus H$^+$), and myoglobin (Aldrich) at 16950.9 (plus H$^+$). SEC measurements were performed using a Shodex GPC101 equipped with a refractive index detector model Shodex RI-71S. A column of shodex LF-804 was employed with THF as an eluent at a flow rate of 1.0 mL/min. In a typical procedure, 30 μL of sample solution (sample concentration of 1 wt %) was injected.
RESULTS AND DISCUSSION

A 4-armed star poly(methyl acrylate), PMA, having allyl end groups has been prepared by the ATRP of methyl acrylate using pentaerythritol tetrakis(2-bromoisobutyrate), 1, as a tetrafunctional initiator, ([MA]:[1]:[Cu(I)Br]:[bpy] = 36:1.0:0.81:2.2, 90 µL, 20 min), and the subsequent in situ allylation of the end groups with allyltributyltin. 8d The 1H NMR spectrum of the product (Figure 1, top), showed signals due to the methylene protons in the core unit (4.10 ppm) and those due to allyl groups located at the chain ends (5.04 and 5.67 ppm). Methyne proton signals adjacent to the terminal bromide atom (4.23 ppm) was eliminated after the allylation reaction. 8d The MALDI-TOF mass spectra (Figure 2, top) showed a uniform series of peaks corresponding to PMA (peak interval of 86 mass units). Each peak corresponds exactly to the molar mass summing up the 4-armed star structure having allyl groups at each chain ends. As an example, the peak (assumed to be the adduct with Na+) at 3986.6 corresponds to the product with the number of monomer MA units, NMA of 40, (C4H6O2)n + 40 + C29H44O8, plus Na+ as 3987.28. Since the 8-shaped PMA product, 3, is produced from the 4-armed PMA precursor, 2, by the elimination of two ethylene molecule, their molecular weights differ by 56, (28/2), mass units. And this was indeed confirmed by comparing the two spectra in Figure 2. It is also remarkable that neither peak were detectable corresponding to twin-tail tadpole PMA formed by single metathesis condensation of 4-armed star PMA precursor, nor corresponding to unreacted 4-armed star PMA. Also due to the detection limit of the molar mass (around \(10^4\)) by the current MALDI condition, no peaks were visible for dimeric byproducts obtainable by intermolecular metathesis condensation. Thus it is confirmed that the 8-shaped PMA was obtained selectively.

SEC traces of the 4-armed star PMA precursor and the 8-shaped PMA were compared in Figure 3. Both traces retained unimodal distribution to indicate negligible chain degradation during the metathesis process. It has been shown theoretically as well as from SEC measurements that 4-armed star polymers have smaller hydrodynamic volume than
its linear counterpart, and 8-shaped polymers are smaller in their 3D sizes than either linear or simple ring counterparts. Moreover, the 8-shaped polymer could be smaller in its hydrodynamic volume than the 4-armed star precursor, since the former is produced by the double cyclization of the latter. However, the apparent peak molecular weight, $M_p$, as the measure of the 3D size, of the isolated 8-shaped PMA ($M_p = 8900$, PDI = 1.36, polystyrene standard) was almost unchanged from that of 4-armed star PMA ($M_p = 8300$, PDI = 1.35, polystyrene standard). This could be due to the loss of a part of the 8-shaped polymer product during the purification procedure (48% recovery yield), and the observed number-average molecular weights (estimated by NMR analysis) were not constant but reduced for the 8-shaped polymer (4600) from the 4-armed star precursor (5600). Thus in the present stage, we have reserved to undertake the rigorous examination of the 3D size of the 8-shaped polymer in comparison with the 4-armed star polymer.

**CONCLUSION**

An 8-shaped polymer was synthesized through double RCM reaction of a 4-armed star PMA precursor having four allyl end groups, obtained through the ATRP of methyl acrylate with tetrafunctional initiator and subsequent end-capping radical reaction (Keck allylation) with allyltributyltin. This study shows that the combination of ATRP and RCM technique provides a versatile means to produce new polymer material of unconventional topologies.
Acknowledgment. We are grateful to Professor M. Kakimoto for our access to the NMR and GPC apparatus. This work was supported partly by a grant from the Ministry of Education, Culture, Sports, Science, and Technology, Japan (17350054).

Received: February 5, 2008
Accepted: March 11, 2008
Published: April 23, 2008

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