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Thermoresponsive property of poly(*N,N*-bis(2-ethoxyethyl)acrylamide) and its multiblock copolymers with poly(*N,N*-dimethylacrylamide) prepared by hydrosilylation-promoted group transfer polymerization†

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Poly(*N,N*-bis(2-ethoxyethyl)acrylamide) (PEOEAm), a new thermoresponsive polyacrylamide, has been studied from the perspectives of synthesis and thermal phase transition. PEOEAm without any initiator residue at the chain end was synthesized via hydrosilylation-promoted group transfer polymerization (GTP) to exclude the effect of polymer end groups on the thermoresponsive properties of the polymer. To extend thermoresponsive PEOEAm to its copolymer systems, the GTP tendency of EOEAm was evaluated and compared with that of *N,N*-dimethylacrylamide (DMAm). There exists a significant difference between the polymerization reactivities of EOAm and DMAm, and this difference was applied to realize the random group transfer copolymerization (GTcoP) of EOAm ("A" unit) and DMAm ("B" unit) for the one-pot synthesis of a BA-block copolymer. A series of block copolymer architectures, such as AB-, ABA-, BAB-, and BABA-block copolymers, were similarly prepared. The thermoresponsive properties of the block copolymers were evaluated by measuring the cloud-point temperature and hydrodynamic radius of their aqueous solutions.

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Introduction

Smart polymers can respond to various stimuli such as heat, light, pH, and magnetic and electric fields. Changes have been observed in individual properties such as hydrophilicity and hydrophobicity, sol-gel properties, and crystallinity, as well as in combinations of these properties, of these materials before and after they respond to such stimuli.^{1–8} For example, aqueous solutions of certain polymers exhibit reversible phase

transition behavior, turning turbid when heated to a temperature above the lower critical solution temperature (LCST) and turning back to a clear solution upon dissolution when cooled below the LCST.^{9–14} One of the molecular design approaches adopted in the synthesis of thermoresponsive polymers is to balance the water solubility and hydrophobicity of the polymers.^{15–17} Among representative thermoresponsive polymers such as poly(*N*-substituted (meth)acrylamide), polyethers, and cellulose derivatives, poly(*N*-isopropylacrylamide) (PNIPAM) has shown potential for chemical and biological applications in controlled drug delivery, bioconjugation, tissue engineering, and bioseparation, as well as in biosensors.^{18–21} Therefore, the phase transition behavior of PNIPAM has been studied using analytical methods such as turbidimetric, calorimetric, nuclear magnetic resonance, fluorescence, light scattering, and neutron scattering analyses.^{22–24} With regard to the synthetic chemistry of thermoresponsive poly(*N*-substituted (meth)acrylamide), there are two important challenges: the expansion of applicable monomers and the synthesis of well-defined polymers through precise polymerization. Many thermoresponsive poly(*N*-substituted acrylamides), including PNIPAM, are known, and various polymer architectures have

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been designed, synthesized, and used in a variety of applications. In contrast, poly(*N,N*-disubstituted acrylamide) (PDSAm) has two different combinations of *N,N*-disubstituents: (1) one type comprises PDSAm with two different *N*-substituents to balance water solubility and hydrophobicity, and these *N,N*-disubstituents can cause thermal responses. Examples of this type are *N*-ethyl, *N*-methyl, *N*-methyl, *N*-*n*-propyl, *N*-methyl, *N*-isopropyl, *N*-ethyl, *N*-(2-methoxylethyl), *N*-(2-methoxylethyl), *N*-*n*-propyl, and *N*-(2-methoxylethyl), *N*-isopropyl groups.^{25–30} (2) The other type comprises thermo-responsive PDSAm with two identical *N*-substituents and is limited to poly(*N,N*-diethylacrylamide) (PDEAm), poly(*N,N*-bis(2-methoxylethyl)acrylamide) (PMOEAm), poly(*N*-acryloylpiperidine), and poly(*N*-acryloylpiperidine) because of the limited number of suitable *N*-substituents compared to the first type.^{31–34} Although PDEAm and PMOEAm have been used predominantly, it will be very interesting to explore the PDSAm family to determine how the balance between water solubility and hydrophobicity affects thermoresponsive polymers.

Precise synthesis of PDSAm and its derivatives has been realized using controlled/living radical and anionic polymerizations of *N,N*-disubstituted acrylamide (DSAm).^{35–38} We developed the hydrosilylation-promoted group transfer polymerization (GTP) of DSAm with hydrosilane (HSiR₃) using tris(pentafluorophenyl)borane (B(C₆F₅)₃) as a catalyst. This method is a certain and reliable method for producing PDSAm with both ends capped with hydrogen due to initiation and termination reactions. This method can provide important insights into the intrinsic thermoresponsive properties of PDSAm, unaffected by its chain-end groups.^{39–42} In this paper, we reported the synthesis of poly(*N,N*-bis(2-ethoxyethyl)acrylamide) (PEOEAm) by the hydrosilylation-promoted GTP of *N,N*-bis(2-ethoxyethyl)acrylamide (EOEAm) using Me₂EtSiH and B(C₆F₅)₃, as shown in Scheme 1. The polymerization reactivity of EOEAm was revealed by comparing the polymerization kinetics and monomer reactivity ratios of EOEAm and *N,N*-dimethylacrylamide (DMAm). The hydrosilylation-promoted group transfer copolymerization (GTcoP) of EOEAm and DMAm was applied to prepare di-, tri-, and tetra-block copolymers. The thermoresponsive behaviors of PEOEAm and its block copolymers were discussed by measuring the cloud-point temperature (*T*_{cp}) and the aggregation properties below and above the *T*_{cp}.

Experimental

Measurements

¹H nuclear magnetic resonance (NMR) spectra were recorded by using a Bruker Avance III HD 500. Polymerization solutions

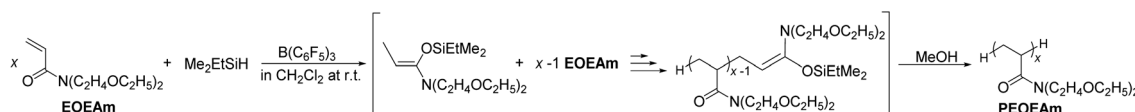
were prepared in a Mikrouna glove box equipped with a gas purification system (molecular sieves and a copper catalyst) under a dry Ar atmosphere (H₂O and O₂ contents <1 ppm). The moisture and oxygen contents in the glove box were monitored by using MK-XTR-100 and MK-OX-SEN-1 sensors, respectively. The number-average molecular mass (*M*_{n,SEC}) and molecular mass distribution (*D*) of the polymers were measured by size-exclusion chromatography (SEC) at 40 °C using an Agilent high-performance liquid chromatography system (1260 Infinity II) in *N,N*-dimethylformamide (DMF) containing lithium chloride (0.01 mol L^{−1}) at a flow rate of 1.0 mL min^{−1} using Agilent Polar Gel-M (exclusion limit, 2 × 10⁴ g mol^{−1}) and Polar Gel-M (exclusion limit, 4 × 10⁶ g mol^{−1}) columns (7.5 × 300 mm; average bead size, 5 μm). Cloud-point measurements were performed using an ultraviolet-visible (UV-vis) spectrophotometer (Jasco V-770, Tokyo, Japan) equipped with a Jasco CTU-100 temperature controller. The path length was 10 mm, and temperature increased at a rate of 1 °C min^{−1}. Changes in transmittance with increasing temperature were recorded at a wavelength of 500 nm. The hydrodynamic radii (*R*_h) of the obtained polymers were analyzed using a Dyna Pro Nanostar® (Wyatt Technology, Santa Barbara, CA, USA) with a heating rate of 10 °C min^{−1}.

Materials

Dichloromethane (CH₂Cl₂, >99.5%; water content, <0.001%), methanol (MeOH), and calcium hydride (CaH₂) were purchased from Kanto Chemicals Co., Inc. Bis(2-ethoxyethyl) amine, *N,N*-dimethylacrylamide (DMAm), and dimethyl-ethylsilane (Me₂EtSiH) were purchased from Tokyo Kasei Kogyo Co., Ltd. Tris(pentafluorophenyl)borane (B(C₆F₅)₃) was purchased from Wako Pure Chemical Industries, Ltd, and used after recrystallization from *n*-hexane at −30 °C. DMAm was distilled from CaH₂, degassed in three freeze–pump–thaw cycles, and stored under an Ar atmosphere prior to use. All other chemicals were purchased from available suppliers and used without purification. All polymerizations were performed in a glove box under an Ar atmosphere at 25 °C.

Synthesis of EOEAm

A solution of acryloyl chloride (22.6 mL, 25.2 g, 0.278 mol) in CH₂Cl₂ (80 mL) was added with stirring to a solution of bis(2-ethoxyethyl)amine (46.1 mL, 41.0 g, 0.254 mol) and triethylamine (53.0 mL, 38.6 g, 0.382 mol) in CH₂Cl₂ (700 mL) at 0 °C. The reaction mixture was allowed to warm up to ambient



Scheme 1 Synthetic pathway of poly(*N,N*-bis(2-ethoxyethyl)acrylamide) (PEOEAm) by the hydrosilylation-promoted group transfer polymerization (GTP) of *N,N*-bis(2-ethoxyethyl)acrylamide (EOEAm).

temperature and was stirred overnight. The precipitated triethylammonium hydrochloride was filtered and washed with CH_2Cl_2 (2×100 mL). Subsequently, the combined organic solutions were extracted with 1 M HCl (3×100 mL), water (100 mL), and brine (200 mL). The organic layer was dried over anhydrous MgSO_4 , and the solvent was removed under reduced pressure. The raw product was purified by vacuum distillation ($154^\circ\text{C}/945$ mbar) to obtain a viscous liquid. Yield, 32.9 g (60%). Anal. Calcd for $\text{C}_{12}\text{H}_{22}\text{O}_2\text{Si}$ (226.39): C, 61.37; H, 9.83; N, 6.51. Found: C, 61.41; H, 9.81; N, 6.49. ^1H NMR (500 MHz in CDCl_3 , δ in ppm): 1.18 (t, 6H, $-\text{CH}_3$), 3.47 (q, 4H, $-\text{O}-\text{CH}_2\text{CH}_3$), 3.55 (t, 2H, $-\text{NCH}_2-$), 3.62 (dd, 4H, $-\text{O}-\text{CH}_2\text{CH}_2\text{N}-$), 3.64 (t, 2H, $-\text{NCH}_2-$), 5.66 (dd, 1H, $\text{CH}^E=\text{CH}-\text{CO}$), 6.32 (dd, 1H, $\text{CH}^Z=\text{CH}-\text{CO}$), 6.68 (dd, 1H, $=\text{CH}-\text{C}=\text{O}$). ^{13}C NMR (125 MHz in CDCl_3 , δ in ppm): 15.10 ($-\text{CH}_3$), 47.32 and 49.09 ($-\text{NCH}_2-$), 66.40 and 66.78 ($-\text{OCH}_2\text{CH}_3$), 68.75 and 68.86 ($-\text{NCH}_2\text{CH}_2-$), 127.24 ($\text{CH}_2=\text{CH}-$), 128.14 ($=\text{CH}-\text{CO}$), 166.88 ($-\text{CO}$).

Synthesis of PEOEAm

A typical polymerization procedure was conducted as follows (Procedure A): a solution of EOEAm (861.2 mg, 4.0 mmol), Me_2EtSiH (5.28×10^{-3} mL, 0.04×10^{-3} mmol), and $\text{B}(\text{C}_6\text{F}_5)_3$ (2.1 mg, 0.004×10^{-3} mmol) in CH_2Cl_2 (3.96 mL) was stirred for 12 h at 25°C in a glove box under an Ar atmosphere. A small amount of methanol was added to terminate the polymerization. Subsequently, the crude product was purified by dialysis against acetone to obtain a primrose yellow solid product. Yield, 432.8 mg (50%); M_n , 21.8 kg mol $^{-1}$; M_w/M_n , 1.11 (run 4, Table 1).

Synthesis of PDMAm-*b*-PEOEAm

A typical random copolymerization was conducted as follows: Procedure A was performed using Me_2EtSiH (5.28×10^{-3} mL, 0.04×10^{-3} mmol), EOEAm (430.6 mg, 2.0 mmol), DMAm (198.4 mg, 2.0 mmol), $\text{B}(\text{C}_6\text{F}_5)_3$ (10.4 mg, 0.02×10^{-3} mmol), and CH_2Cl_2 (3.96 mL). The polymerization was conducted for 24 h and then terminated by adding methanol to obtain PDMAm-*b*-PEOEAm as a solid polymer. Yield, 413.5 mg (58%); M_n , 15.8 kg mol $^{-1}$; M_w/M_n , 1.09 (run 16, Table 2).

Table 1 Hydrosilylation-promoted group transfer polymerization (GTP) of EOEAm with Me_2EtSiH using $\text{B}(\text{C}_6\text{F}_5)_3$ as the catalyst ^a

Run	Polymer	$[\text{EOEAm}]_0/[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0$	Time/h	M_n,calcd ^b /kg mol $^{-1}$	M_n,MALS ^c (M_w/M_n) ^d /kg mol $^{-1}$	T_{cp} ^e /°C
1	PEOEAm ₂₅	25/1/0.1	6	5.4	5.4 (1.15)	14.5
2	PEOEAm ₅₀	50/1/0.1	12	10.8	10.5 (1.15)	13.9
3	PEOEAm ₇₅	75/1/0.1	12	16.1	15.9 (1.12)	8.9
4	PEOEAm ₁₀₀	100/1/0.1	12	21.5	21.8 (1.11)	8.0
5	PEOEAm ₁₅₀	150/1/0.5	24	32.3	31.8 (1.10)	5.9
6	PEOEAm ₂₀₀	200/1/0.5	24	43.0	42.5 (1.10)	5.0

^a $[\text{EOEAm}]_0$, 1.0 mol L $^{-1}$; solvent, CH_2Cl_2 ; temp., 25°C ; Ar atmosphere; monomer conversion determined by ^1H NMR in CDCl_3 , >99.9%.

^b Calculated using the equation $[\text{EOEAm}]_0/[\text{Me}_2\text{EtSiH}]_0 \times (\text{conv.}) \times (\text{M.W. of the monomer}) + (\text{M.W. of H}) \times 2$. ^c Determined by size-exclusion chromatography (SEC) using an instrument equipped with a multi-angle light scattering (MALS) detector in DMF containing lithium chloride (0.01 mol L $^{-1}$), $dn/dc = 0.0344$. ^d Determined by SEC using an instrument equipped with a refractive index (RI) detector in DMF containing lithium chloride (0.01 mol L $^{-1}$) using PMMA standards. ^e Determined by ultraviolet-visible light (UV-vis) measurements in water (10 g L $^{-1}$).

Synthesis of PEOEAm-*b*-PDMAm

A typical block copolymerization was conducted as follows: Procedure A was performed using Me_2EtSiH (5.28×10^{-3} mL, 0.04×10^{-3} mmol), EOEAm (430.6 mg, 2.0 mmol), $\text{B}(\text{C}_6\text{F}_5)_3$ (20.8 mg, 0.04×10^{-3} mmol), and CH_2Cl_2 (0.36 $\times 10^{-3}$ mL) for 12 h. An appropriate amount of the reaction solution was sampled to confirm the quantitative consumption of EOEAm by ^1H NMR measurements, and then a solution of DMAm (198.4 mg, 2.0 mmol) in CH_2Cl_2 (3.6 mL) was added to the polymerization mixture, and then this mixture was stirred for 24 h. PEOEAm-*b*-PDMAm was obtained as a white solid polymer. Yield, 381.2 mg (53%); M_n , 15.4 kg mol $^{-1}$; M_w/M_n , 1.10 (run 9, Table 2).

Synthesis of PDMAm-*b*-PEOEAm-*b*-PDMAm

A typical random copolymerization was conducted as follows: Procedure A was performed using Me_2EtSiH (5.28×10^{-3} mL, 0.04×10^{-3} mmol), EOEAm (430.6 mg, 2.0 mmol), DMAm (99.8 mg, 1.0 mmol), $\text{B}(\text{C}_6\text{F}_5)_3$ (20.8 mg, 0.04×10^{-3} mmol), and CH_2Cl_2 (2.3 mL) for 24 h. After confirming the quantitative consumption of EOEAm and DMAm by ^1H NMR measurements, a solution of DMAm (99.8 mg, 1.0 mmol) in CH_2Cl_2 (0.9 mL) was added to the polymerization mixture, and then the mixture was stirred for 12 h. PDMAm-*b*-PEOEAm-*b*-PDMAm was obtained as a white solid polymer. Yield, 411.1 mg (57%); M_n , 15.8 kg mol $^{-1}$; M_w/M_n , 1.12 (run 23, Table 3).

Synthesis of PEOEAm-*b*-PDMAm-*b*-PEOEAm

A typical random copolymerization was conducted as follows: Procedure A was performed using Me_2EtSiH (5.28×10^{-3} mL, 0.04×10^{-3} mmol), EOEAm (215.3 mg, 1.0 mmol), and $\text{B}(\text{C}_6\text{F}_5)_3$ (20.8 mg, 0.04×10^{-3} mmol) in CH_2Cl_2 (0.71×10^{-3} mL) for 6 h. After confirming the quantitative consumption of EOEAm by ^1H NMR measurements, a solution of DMAm (199.6 mg, 2.0 mmol) and EOEAm (215.3 mg, 1.0 mmol) in CH_2Cl_2 (2.58 mL) was added to the polymerization mixture, and then the mixture was stirred for 24 h. PEOEAm-*b*-PDMAm-*b*-PEOEAm was obtained as a white solid polymer. Yield, 395.9 mg (55%); M_n , 15.6 kg mol $^{-1}$; M_w/M_n , 1.12 (run 30, Table 3).

Table 2 Thermoresponsive properties of PEOEAm-*b*-PDMAm and PDMAm-*b*-PEOEAm by one-pot synthesis using the hydrosilylation-promoted group transfer copolymerization (GTcoP) of EOEA (M₁) and DMAm (M₂) with Me₂EtSiH (SiH) using B(C₆F₅)₃ as the catalyst^a

Run	Polymer	Block GTcoP		Random GTcoP ^c [M ₁] ₀ /[M ₂] ₀	M _{n,calcd} / kg mol ⁻¹	M _{n,SEC} (M _w /M _n) ^d / kg mol ⁻¹	T _{cp} ^e /°C
		1st GTP ^b [M ₁] ₀ / [SiH] ₀	2nd GTP [M ₂] ₀ / [SiH] ₀				
7	PEOEAm ₃₀ - <i>b</i> -PDMAm ₇₀	30	70		13.4	13.3 (1.11)	33.0
8	PEOEAm ₄₀ - <i>b</i> -PDMAm ₆₀	40	60		14.6	14.5 (1.09)	29.0
9	PEOEAm ₅₀ - <i>b</i> -PDMAm ₅₀	50	50		15.7	15.4 (1.10)	25.0
10	PEOEAm ₆₀ - <i>b</i> -PDMAm ₄₀	60	40		16.9	17.2 (1.11)	23.1
11	PEOEAm ₇₀ - <i>b</i> -PDMAm ₃₀	70	30		18.0	17.9 (1.09)	20.5
12	PEOEAm ₈₀ - <i>b</i> -PDMAm ₂₀	80	20		19.2	19.3 (1.10)	20.0
13	PEOEAm ₉₀ - <i>b</i> -PDMAm ₁₀	90	10		20.4	19.9 (1.12)	13.8
14	PDMAm ₇₀ - <i>b</i> -PEOEAm ₃₀			30/70	13.4	13.4 (1.12)	33.6
15	PDMAm ₆₀ - <i>b</i> -PEOEAm ₄₀			40/60	14.6	14.4 (1.11)	27.0
16	PDMAm ₅₀ - <i>b</i> -PEOEAm ₅₀			50/50	15.7	15.8 (1.09)	25.1
17	PDMAm ₄₀ - <i>b</i> -PEOEAm ₆₀			60/40	16.9	16.7 (1.11)	22.0
18	PDMAm ₃₀ - <i>b</i> -PEOEAm ₇₀			70/30	18.0	18.4 (1.12)	21.5
19	PDMAm ₂₀ - <i>b</i> -PEOEAm ₈₀			80/20	19.2	19.6 (1.10)	19.0
20	PDMAm ₁₀ - <i>b</i> -PEOEAm ₉₀			90/10	20.4	20.1 (1.09)	14.0

^a Solvent, CH₂Cl₂; temp., 25 °C; Ar atmosphere; monomer conversion determined by ¹H NMR spectra in CDCl₃, >99.9%. ^b [EOEA]₀, 1.0 mol L⁻¹; [SiH]₀/[B(C₆F₅)₃]₀, 1.0; time, 12 h (1st GTP) and 24 h (2nd GTP). ^c [M₁ + M₂]₀/[SiH]₀/[B(C₆F₅)₃]₀, 100/1/0.5; time, 24 h. ^d Determined by an SEC instrument equipped with an RI detector in DMF containing lithium chloride (0.01 mol L⁻¹) using PMMA standards. ^e Determined by UV-vis measurements in water (10 g L⁻¹).

Table 3 Thermoresponsive properties of PDMAm-*b*-PEOEAm-*b*-PDMAm and PEOEAm-*b*-PDMAm-*b*-PEOEAm by one-pot synthesis using the hydrosilylation-promoted GTcoP of EOEA (M₁) and DMAm (M₂) with Me₂EtSiH (SiH) using B(C₆F₅)₃ as the catalyst^a

Run	Polymer	1st GTcoP ^b and 2nd GTP		1st GTP ^c and 2nd GTcoP		M _{n,calcd} / kg mol ⁻¹	M _{n,SEC} (M _w /M _n) ^d / kg mol ⁻¹	T _{cp} ^e / °C
		[M ₁ + M ₂] ₀ /[SiH] ₀	[M ₂] ₀ /[SiH] ₀	[M ₁] ₀ /[SiH] ₀	[M ₁ + M ₂] ₀ /[SiH] ₀			
21	PDMAm ₃₅ - <i>b</i> -PEOEAm ₃₀ - <i>b</i> -PDMAm ₃₅	(30 + 35)/1	35/1			13.4	13.5 (1.10)	— ^f
22	PDMAm ₃₀ - <i>b</i> -PEOEAm ₄₀ - <i>b</i> -PDMAm ₃₀	(40 + 30)/1	30/1			14.6	14.6 (1.09)	— ^f
23	PDMAm ₂₅ - <i>b</i> -PEOEAm ₅₀ - <i>b</i> -PDMAm ₂₅	(50 + 25)/1	25/1			15.7	15.8 (1.12)	— ^f
24	PDMAm ₂₀ - <i>b</i> -PEOEAm ₆₀ - <i>b</i> -PDMAm ₂₀	(60 + 20)/1	20/1			16.9	17.1 (1.11)	34.8
25	PDMAm ₁₅ - <i>b</i> -PEOEAm ₇₀ - <i>b</i> -PDMAm ₁₅	(70 + 15)/1	15/1			18.0	17.9 (1.10)	30.5
26	PDMAm ₁₀ - <i>b</i> -PEOEAm ₈₀ - <i>b</i> -PDMAm ₁₀	(80 + 10)/1	10/1			19.2	19.3 (1.11)	24.2
27	PDMAm ₅ - <i>b</i> -PEOEAm ₉₀ - <i>b</i> -PDMAm ₅	(90 + 5)/1	5/1			20.4	20.5 (1.12)	19.5
28	PEOEAm ₁₅ - <i>b</i> -PDMAm ₇₀ - <i>b</i> -PEOEAm ₁₅			15/1	(15 + 70)/1	13.4	13.4 (1.11)	— ^f
29	PEOEAm ₂₀ - <i>b</i> -PDMAm ₆₀ - <i>b</i> -PEOEAm ₂₀			20/1	(20 + 60)/1	14.6	14.5 (1.10)	39.6
30	PEOEAm ₂₅ - <i>b</i> -PDMAm ₅₀ - <i>b</i> -PEOEAm ₂₅			25/1	(25 + 50)/1	15.7	15.6 (1.12)	34.9
31	PEOEAm ₃₀ - <i>b</i> -PDMAm ₄₀ - <i>b</i> -PEOEAm ₃₀			30/1	(30 + 40)/1	16.9	17.0 (1.10)	30.8
32	PEOEAm ₃₅ - <i>b</i> -PDMAm ₃₀ - <i>b</i> -PEOEAm ₃₅			35/1	(35 + 30)/1	18.0	18.1 (1.13)	26.9
33	PEOEAm ₄₀ - <i>b</i> -PDMAm ₂₀ - <i>b</i> -PEOEAm ₄₀			40/1	(40 + 20)/1	19.2	19.5 (1.12)	21.0
34	PEOEAm ₄₅ - <i>b</i> -PDMAm ₁₀ - <i>b</i> -PEOEAm ₄₅			45/1	(45 + 10)/1	20.4	20.4 (1.11)	16.7

^a Solvent, CH₂Cl₂; [SiH]₀/[B(C₆F₅)₃]₀, 1.0; temp., 25 °C; Ar atmosphere; monomer conversion determined by ¹H NMR spectra in CDCl₃, >99.9%.

^b [M₁ + M₂]₀, 1.0 mol L⁻¹; time, 24 h (1st GTcoP), 12 h (2nd GTP, runs 21–24) and 6 h (2nd GTP, runs 25–27). ^c [M₁]₀, 1.0 mol L⁻¹; time, 6 h (1st GTP, runs 28–32), 12 h (1st GTP, runs 33 and 34), and 24 h (2nd GTP). ^d Determined by an SEC instrument equipped with an RI detector in DMF containing lithium chloride (0.01 mol L⁻¹) using PMMA standards. ^e Determined by UV-vis measurements in water (10 g L⁻¹). ^f Insoluble in water.

Results and discussion

Hydrosilylation-promoted GTP of EOEA

EOEA was prepared by the reaction of bis(2-ethoxyethyl) amine and acryloyl chloride. The ¹H and ¹³C NMR spectra of EOEA and its polymer are shown in Fig. S1.† Two signals at 47.32 and 49.09 ppm corresponding to the $-\text{NCH}_2-$ group, at 66.40 and 66.78 ppm corresponding to the $-\text{OCH}_2\text{CH}_3$ group,

and at 68.75 and 68.86 ppm corresponding to the $-\text{NCH}_2\text{CH}_2-$ group are observed in the ¹³C NMR spectrum (Fig. S1c†); these signals are attributed to the *cis* and *trans* conformers of EOEA. However, they exhibited identical polymerization reactivity, as confirmed by the fact that both were consumed equally as polymerization progressed. The conventional GTP method, in which the SKA initiator is used as one of the GTP components, yields polymers with SKA residues attached to

the α -terminus. In contrast, the hydrosilylation-promoted GTP method, in which SKA is generated by the 1,4-hydrosilylation of the monomer and hydrosilane in the polymerization system, yields polymers consisting only of repeating monomer units. Since the thermal properties of the thermoresponsive polymers in aqueous solutions are affected by the molecular mass and terminal substituents, the hydrosilylation-promoted GTP method was used to synthesize PEOEAm. We first described the result for the polymerization of EOEAm using Me_2EtSiH and $\text{B}(\text{C}_6\text{F}_5)_3$ in CH_2Cl_2 with a ratio $[\text{EOEAm}]_0/[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0 = 25/1/0.5$ at 25 °C. The obtained polymer exhibited a unimodal SEC with a relatively low molecular mass distribution (D) of 1.20, as shown in Fig. 1. The number-average molecular mass determined by multi-angle light scattering (MALS) ($M_{n,\text{MALS}}$) was 5.4 kg mol⁻¹; this result agrees with the calculated number-average molecular mass ($M_{n,\text{calcd}}$) of 5.4 kg mol⁻¹.

The ^1H (Fig. S1b†) and ^{13}C NMR spectra (Fig. S1d†) of the obtained polymer showed signals attributed to methylene and methine protons at 1.25–1.90 and 2.15–2.75 ppm, respectively, and those attributed to vinyl methylene and methine carbons at 45–50 and 34–38 ppm, respectively. A more detailed chemical structure of the obtained polymer is revealed in the results of matrix assisted laser desorption/ionization-time of flight mass spectrometry (MALDI-TOF MS). As shown in Fig. 2, only

one series of molecular ion peaks was observed, and the distance between two adjacent molecular ion peaks was 215.3 Da, which agreed with the predicted molecular mass of 215.29 Da for EOEAm as the constitutional repeating unit. In addition, the m/z value of each molecular ion peak clearly indicated the sodium-cationized polymer composition of $[\text{H}-\text{EOEAm}_n-\text{H} + \text{Na}]^+$ (molecular formula: $\text{C}_{11n} + \text{H}_{21n} + \text{N}_n + \text{O}_{3n} + 2\text{Na}$); for instance, an m/z value of 5407.2 Da for a specified peak corresponds to a sodium-cationized 25-mer polymer structure of $[\text{H}-\text{EOEAm}_{25}-\text{H} + \text{Na}]^+$ with a theoretical monoisotopic value of 5407.33 for the molecular formula of $\text{C}_{275}\text{H}_{527}\text{N}_{25}\text{O}_{75}\text{Na}$. This result strongly supported our supposition that the hydrosilylation-promoted GTP of EOEAm was well controlled without any side reactions to produce the polymer with a planned degree of polymerization (DP) of 25, *i.e.*, PEOEAm₂₅. PEOEAm with a higher molecular mass was prepared by the $\text{B}(\text{C}_6\text{F}_5)_3$ -catalyzed GTP of EOEAm and Me_2EtSiH with varying $[\text{EOEAm}]_0/[\text{Me}_2\text{EtSiH}]_0$ ratios of 50, 75, 100, 150, and 200. Table 1 lists the polymerization results. All the SEC traces of the obtained polymers showed a unimodal distribution and shifted to the higher molecular mass region with increasing $[\text{EOEAm}]_0/[\text{Me}_2\text{EtSiH}]_0$ ratios, as shown in Fig. 1. The $M_{n,\text{MALS}}$ values of 10.5, 15.9, 21.8, 31.8, and 42.5 kg mol⁻¹ agreed with the $M_{n,\text{calcd}}$ values of 10.8, 16.1, 21.5, 32.3, and 43.0 kg mol⁻¹, respectively. The D of the obtained polymers decreased from 1.15 to 1.10 with increasing $[\text{EOEAm}]_0/[\text{Me}_2\text{EtSiH}]_0$ ratios. These results indicated that PEOEAm_x with the targeted DPs, *i.e.*, PEOEAm₅₀, PEOEAm₇₅, PEOEAm₁₀₀, PEOEAm₁₅₀ and PEOEAm₂₀₀, can be prepared by the hydrosilylation-promoted GTP of EOEAm by varying the $[\text{EOEAm}]_0/[\text{Me}_2\text{EtSiH}]_0$ ratios (Table 1).

Polymerization reactivity of EOEAm

The GTP method allows the synthesis of a variety of thermo-responsive PEOEAm-based polymers by using EOEAm for synthesizing copolymers and homopolymers. Therefore, we investigated the polymerization reactivity of EOEAm as well as the copolymerization of EOEAm and *N,N*-dimethylacrylamide (DMAm), the simplest *N,N*-disubstituted acrylamide. The resulting poly(*N,N*-dimethylacrylamide) (PDMAm) was water soluble but non-thermally expandable, similar to the reference monomer. First, the difference between the polymerization properties of EOEAm and DMAm was investigated through

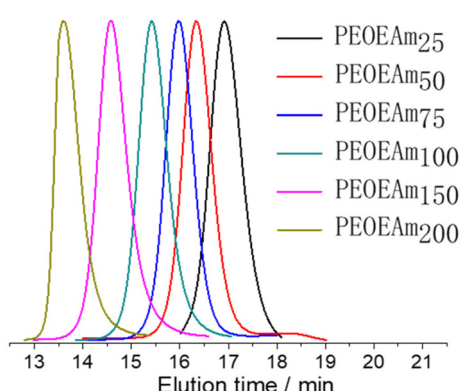


Fig. 1 Size-exclusion chromatography (SEC) traces of PEOEAm obtained by the GTP of EOEAm and Me_2EtSiH with $[\text{EOEAm}]_0/[\text{Me}_2\text{EtSiH}]_0$ ratios of 25, 50, 75, 100, 150, and 200 and with a $[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0$ ratio of 0.5.

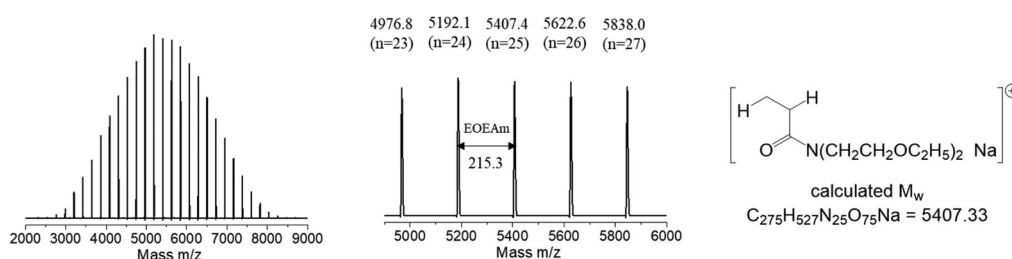


Fig. 2 Matrix assisted laser desorption/ionization-time of flight mass spectrometry (MALDI-TOF MS) spectra of representative PEOEAm with an $M_{n,\text{MALS}}$ of 5.4 kg mol⁻¹ and an M_w/M_n of 1.20 (Table 1, run 1).

kinetic experiments using the hydrosilylation-promoted GTP under the following conditions: $[\text{EOEAm or DMAm}]_0/[\text{Et}_2\text{MeSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0 = 100/1/0.1$ and $[\text{EOEAm or DMAm}]_0 = 1.0 \text{ mol L}^{-1}$. In both GTPs, induction periods were observed as shown in Fig. 3a, and induction times were 30 and 20 min for EOEAm and DMAm, respectively. After the induction time, both monomers were linearly consumed with different polymerization times, and the observed propagation rates ($k_{\text{p,obs}}$) for EOEAm and DMAm were 8.8 and 68.2 min^{-1} , respectively, from the zero-order kinetic plots for monomer conversion (Conv.) *vs.* polymerization time (Time). The molecular mass (M_n) of the obtained polymers increased linearly with increasing monomer conversion, while the dispersity values (D_s) remained below 1.2, as shown in Fig. 3b. These results indicate that the hydrosilylation-promoted GTP of EOEAm with Me_2EtSiH using $\text{B}(\text{C}_6\text{F}_5)_3$ proceeded in a controlled/living manner similar to the case of DMAm, while the polymerization rate of EOEAm was much slower than that of DMAm.^{43–45}

Furthermore, to clarify the copolymerization reactivity of EOEAm toward DMAm, random group transfer copolymerization (GTcoP) of EOEAm and DMAm was performed under the conditions $[\text{EOEAm} + \text{DMAm}]_0/[\text{SKA}^{\text{Et}}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0 = 100/1/0.2$ in CH_2Cl_2 (see the ESI†). The monomer reactivity ratios r_{EOEAm} and r_{DMAm} were determined to be 0.66 and 15.83, respectively, by the Kelen–Tüdös method. The reactivity of vinyl monomers is mainly affected by the electronegativity of the vinyl group and the steric hindrance around the vinyl group, and it is known that there exists a correlation between the electronegativity of the vinyl group and the chemical shift value in the ^{13}C NMR spectrum. The chemical shifts of $\text{CH}_2=$ and $-\text{CH}=$ for EOEAm are 128.14 and 127.57 ppm, respectively, which are very close to the values of 127.69 and 127.41 ppm for DMAm. Since the electronegativity of both monomers is almost equal, the factor that drastically reduced the polymerization properties of EOEAm compared to those of DMAm was mainly the bulky bis(2-ethoxyethyl)amino group in EOEAm.⁴⁶

Block and random GTcoP

To expand the scope of thermoresponsive PEOEAm, block and statistical copolymers consisting of PEOEAm and PDMAM were prepared. The block GTcoP of EOEAm and DMAm was performed using the sequential addition of a monomer with different initial monomer feed ratios of $[\text{EOEAm}]_0$ and $[\text{DMAm}]_0$. The quantitative consumption of EOEAm in the first GTP was confirmed by ^1H NMR; thereafter, a second GTP was initiated by adding DMAm. In addition, the progress of the block GTcoP was confirmed by the shifting of the SEC trace of the polymer obtained from the first GTP of EOEAm to the high molecular mass region by the second GTP of DMAm while maintaining narrow molecular mass dispersions (Fig. S7†). In the case of the GTcoP with a $[\text{EOEAm}]_0/[\text{DMAm}]_0/[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0$ ratio of 50/50/1/1, the M_n (D) of the obtained polymer increased from 10.7 kg mol^{-1} (1.12) for the first GTP to 15.4 kg mol^{-1} (1.10) for the second GTP; this value agrees with $M_{n,\text{calcd}} = 15.7 \text{ kg mol}^{-1}$ (run 9, Table 2 and Fig. S7c†).

The random GTcoP of PEOEAm and PDMAM was expected to produce a gradient copolymer, not a random copolymer because there was an extremely large difference in the monomer reactivity ratio between the r_{EOEAm} of 0.66 and the r_{DMAm} of 15.83. This expectation was supported by the number-average sequence length of EOEAm units (l_{EOEAm}), which could be determined as a parameter reflecting the tendency to isolate the EOEAm–EOEAm diad. For the mole fractions of EOEAm of 0.3 and 0.8 in the monomer feed, l_{EOEAm} was very short, 1.1 and 3.6, respectively (Table S1†).^{47–49} The random GTcoP of EOEAm and DMAm with the $[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0$ ratio of 1/0.2 should produce gradient copolymers (Table S1†), but the formation of block copolymers was achieved by the random GTcoP by increasing the amount of catalyst $[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0$ from 1/0.2 to 1/0.5. The polymerization progress for the random GTcoP of EOEAm and DMAm with the $[\text{EOEAm} + \text{DMAm}]_0/[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0$ ratio of (50 + 50)/1/0.5 in CD_2Cl_2 was followed by measuring the ^1H NMR spectra at polymerization times of 20, 40, and

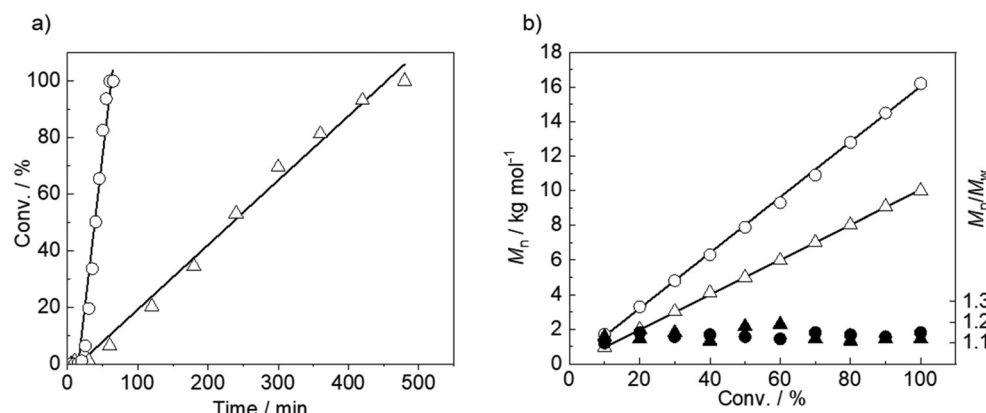


Fig. 3 (a) Zero-order kinetic plots and (b) dependence of molar mass (M_n) and molar mass distribution (M_w/M_n) on monomer conversion (Conv.) in the GTP of (○) EOEAm and (Δ) DMAm ($[\text{EOEAm or DMAm}]_0/[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0, 100/1/0.1; [\text{EOEAm or DMAm}]_0, 1.0 \text{ mol L}^{-1}$).

590 min (Fig. 4). The characteristic proton absorptions attributed to the $=\text{CH-CON-}$ (6.56–6.62 ppm) and $\text{CH}_3\text{N-}$ (3.02–3.09 ppm) groups of DMAm decreased at 20 min and disappeared at 40 min, while those attributed to the $-\text{NCH}_2\text{CH}_2\text{OCH}_2-$ (3.44–3.67 ppm) and $=\text{CH-CON-}$ (6.65–6.71 ppm) groups remained unchanged until 40 min; thereafter, the characteristic proton absorptions attributed to the $-\text{NCH}_2\text{CH}_2\text{OCH}_2-$ and $=\text{CH-CON-}$ groups decreased and finally disappeared completely at 590 min. The ^1H NMR spectrum of the polymer that partially disappeared from the polymerization system after 40 min revealed the characteristic proton absorption attributed to the methylene (1.35–2.07 ppm) and methine (2.26–2.78 ppm) groups in the main chain and $\text{CH}_3\text{N-}$ groups, confirming the formation of PDMAm (Fig. S2a†). Furthermore, the ^1H NMR spectrum of the polymer obtained after 590 min revealed the characteristic proton absorption attributed to the methyl group of the $\text{CH}_3\text{CH}_2\text{O-}$ group and the methylene groups in the $-\text{NCH}_2\text{CH}_2\text{OCH}_2-$ moiety along with the absorption signals of PDMAm, confirming the formation of PDMAm-*b*-PEOEAm (Fig. S2b†). These results indicate that the first GTP of DMAm proceeded preferentially to afford living PDMAm, which functioned as a macro-initiator to continue the second GTP of EOEA, resulting in the formation of PDMAm-*b*-PEOEAm in the random GTcoP of EOEA and DMAm. This means the one-pot synthesis of a diblock copolymer by a random copolymerization. Increasing the amount of catalyst $[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0$ from 1/0.2 to 1/0.5 further increased the difference between the rates of polymerization of DMAm and EOEA. The random GTcoP of EOEA and DMAm was performed by varying the initial molar ratio of EOEA and DMAm under the conditions $[\text{EOEA} + \text{DMAm}]_0/[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0 = 100/1/0.5$. The

copolymerization results of runs 14–20 are summarized in Table 2. For the obtained copolymer $\text{PDMAm}_y\text{-}b\text{-PEOEAm}_x$, the $M_{n,\text{SEC}}$ values were 13.4 kg mol $^{-1}$ for $x/y = 30/70$, 14.4 for 40/60, 15.8 for 50/50, 16.7 for 60/40, 18.4 for 70/30, 19.6 for 80/20, and 20.1 for 90/10, which were in good agreement with the $M_{n,\text{calcd}}$ values. The D values were as small as 1.09–1.12.

The one-pot method was employed to synthesize tri- and tetra-block copolymers. After the first GTcoP of EOEA and DMAm leading to PDMAm-*b*-PEOEAm, DMAm was sequentially added to the first living GTcoP system to afford PDMAm-*b*-PEOEAm-*b*-PDMAm. For extending the system from di- to tri-block segments, the unimodal SEC trace of the diblock copolymer shifted to a higher molar mass region after the second GTP, and the D was as low as 1.09–1.12 (Fig. S9†). The $M_{n,\text{SEC}}$ values of the obtained copolymers were 13.5, 14.6, 15.8, 17.1, 17.9, 19.3, and 20.5 kg mol $^{-1}$, which agreed well with the $M_{n,\text{calcd}}$ values of 13.4, 14.6, 15.7, 16.9, 18.0, 19.2, and 20.4 kg mol $^{-1}$, respectively (Table 3). The characteristic absorptions attributed to PDMAm and PEOEAm were observed in the ^1H NMR spectra of the obtained polymers, confirming the formation of $\text{PDMAm}_{25}\text{-}b\text{-PEOEAm}_{50}\text{-}b\text{-PDMAm}_{25}$ (Fig. S4†). Similarly, the first GTP of EOEA, followed by the second GTcoP of PEOEAm and PDMAm, led to the synthesis of PEOEAm-*b*-PDMAm-*b*-PEOEAm. The extension process from di- to tri-block segments was confirmed from the SEC traces (Fig. S10†). The $M_{n,\text{SEC}}$ of the obtained copolymer agreed with the $M_{n,\text{calcd}}$ (runs 28–34, Table 3), and the D values were as low as 1.10–1.13. The formation of $\text{PEOEAm}_{25}\text{-}b\text{-PDMAm}_{50}\text{-}b\text{-PEOEAm}_{25}$ was confirmed from the ^1H NMR spectrum of the resulting copolymer (Fig. S5†). Finally, a tetra-block copolymer ($\text{PDMAm}\text{-}b\text{-PEOEAm})_2$ was easily synthesized by repeating the random GTcoP of PEOEAm and PDMAm twice (Scheme 2).

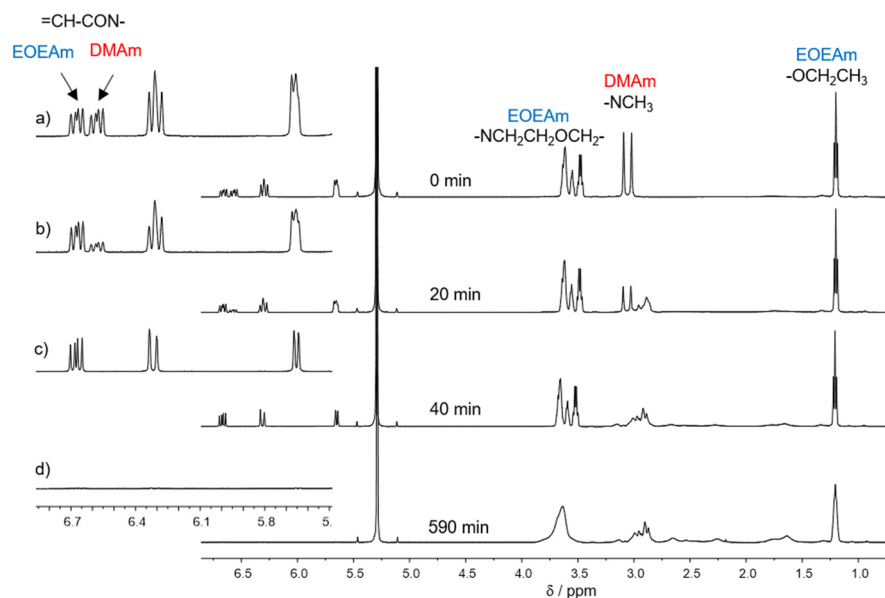
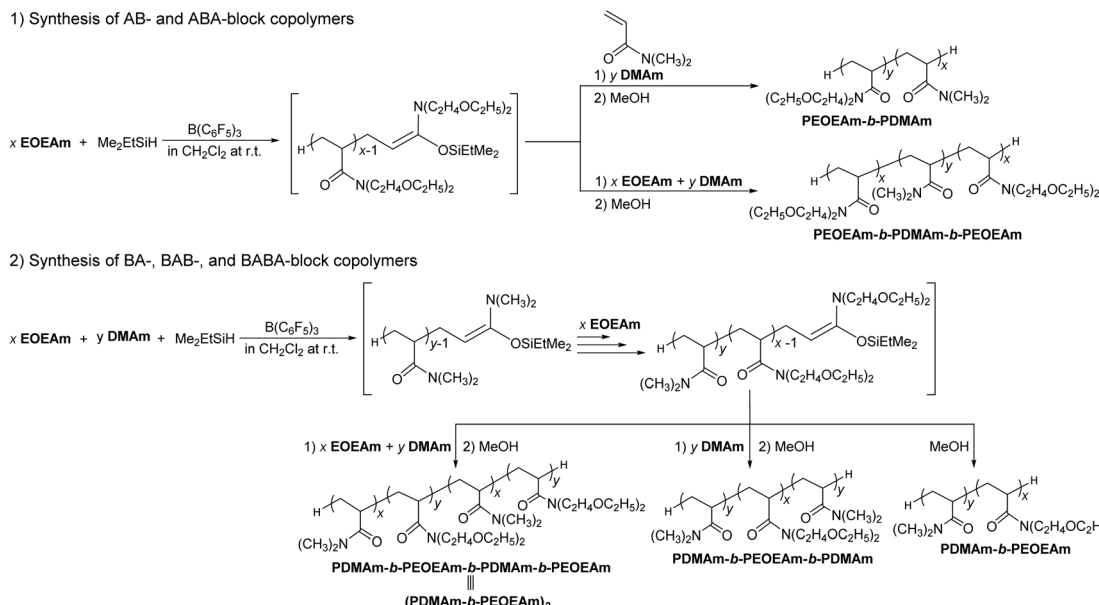


Fig. 4 ^1H nuclear magnetic resonance (NMR) spectra for the GTcoP of EOEA and DMAm at polymerization times of (a) 0, (b) 20, (c) 40, and (d) 590 min and the expanded spectra of the vinyl groups of EOEA and DMAm (solvent, CD_2Cl_2 ; $[\text{EOEA} + \text{DMAm}]_0/[\text{Me}_2\text{EtSiH}]_0/[\text{B}(\text{C}_6\text{F}_5)_3]_0, (50 + 50)/1/0.5$; $[\text{EOEA} + \text{DMAm}]_0, 1.0 \text{ mol L}^{-1}$).



Scheme 2 Synthesis of (1) PEOEA-b-PDMAm and PEOEA-b-PDMAm-b-PEOEA and (2) PDMAm-b-PEOEA, PDMAm-b-PEOEA-b-PDMAm, and (PDMAm-b-PEOEA)₂ by the block and random GTcoP of EOEA and DMAm.

SEC traces revealed that the extension from di- to tetra-block segments proceeded with D values being maintained low (Fig. S11†), and the $M_{n,SEC}$ of the obtained copolymer agreed with the $M_{n,calcd}$ (Table 4). The formation of (PDMAm₂₅-b-PEOEA₂₅)₂ was confirmed from the ¹H NMR spectrum of the resulting copolymer (Fig. S6†).

Thermoresponsive properties of PEOEA and its copolymers with PDMAm

The thermal phase transition behaviors of PEOEA and its copolymers were evaluated at the cloud-point temperature (T_{cp}). At T_{cp} , the transmittance reached 50% of the transmittance-temperature curve that was plotted by monitoring a polymer solution at a concentration of 10 g L⁻¹ with a UV-vis spectrophotometer at a wavelength of 500 nm. The phase-transition properties of PEOEA were compared with those of its analog, namely, poly(*N,N*-bis(2-methoxyethyl)acrylamide) (PMOEAm), to investigate the effect of the side chain of the monomer unit on the thermoresponsive properties. Fig. 5a

shows the dependence of T_{cp} on the degree of polymerization (DP_x) of PEOEA and PMOEAm. For both polymers, with increasing DP_x , T_{cp} decreased from 14.5 to 5.0 °C for PEOEA and from 56.5 to 48.0 °C for PMOEAm. The T_{cp} of PEOEA was drastically reduced because of the increase in hydrophobicity caused by the change in the side chain moiety of the ROCH₂CH₂N- group from methyl to ethyl groups. Such a decrease in T_{cp} (associated with the change from methyl to ethyl groups at the end of the ethylene oxide side chain) was observed for poly(glycidyl ether) and polyisocyanate with ethylene oxide (EO) side chains. Molecular-dynamics (MD) simulations for poly(glycidyl ether) with EO side chains indicated that the hydrophobic hydration shell around the ethyl group at the end of the side chain had a lower hydration shell density than that around the methyl group at the end of the side chain and was more easily broken upon increasing the temperature.^{50–52} The MD simulations also showed that the fluctuations in the polymer chain for the ethyl group at the end of the side chain were smaller than those for the methyl

Table 4 Thermoresponsive properties of (PDMAm-b-PEOEA)₂ by one-pot synthesis using the hydrosilylation-promoted GTcoP of EOEA and DMAm with Me₂EtSiH using B(C₆F₅)₃ as the catalyst^a

Run	Polymer	[EOEA/DMAm] ₀	Time/h	$M_{n,calcd}$ /kg mol ⁻¹	$M_{n,SEC}$ (M_w/M_n) ^b /kg mol ⁻¹	T_{cp} ^c /°C
35	(PDMAm ₃₅ -b-PEOEA ₁₅) ₂	15/35 × 2	8 + 12	13.4	13.9 (1.19)	57.8
36	(PDMAm ₃₀ -b-PEOEA ₂₀) ₂	20/30 × 2	8 + 18	14.6	14.7 (1.20)	54.6
37	(PDMAm ₂₅ -b-PEOEA ₂₅) ₂	25/25 × 2	8 + 18	15.7	15.8 (1.21)	44.9
38	(PDMAm ₂₀ -b-PEOEA ₃₀) ₂	30/20 × 2	8 + 18	16.9	17.1 (1.21)	41.3
39	(PDMAm ₁₅ -b-PEOEA ₃₅) ₂	35/15 × 2	12 + 24	18.0	18.2 (1.18)	38.1
40	(PDMAm ₁₀ -b-PEOEA ₄₀) ₂	40/10 × 2	12 + 24	19.2	19.6 (1.20)	25.5

^a Solvent, CH₂Cl₂; temp., 25 °C; Ar atmosphere; [M]₀, 1.0 mol L⁻¹; [MOEA + EOEA]₀/[Me₂EtSiH]₀/[B(C₆F₅)₃]₀, 50/1/1; monomer conversion determined by ¹H NMR spectra in CDCl₃, >99.9%. ^b Determined by an SEC instrument equipped with an RI detector in DMF containing lithium chloride (0.01 mol L⁻¹) using PMMA standards. ^c Determined by UV-vis measurements in water (10 g L⁻¹).

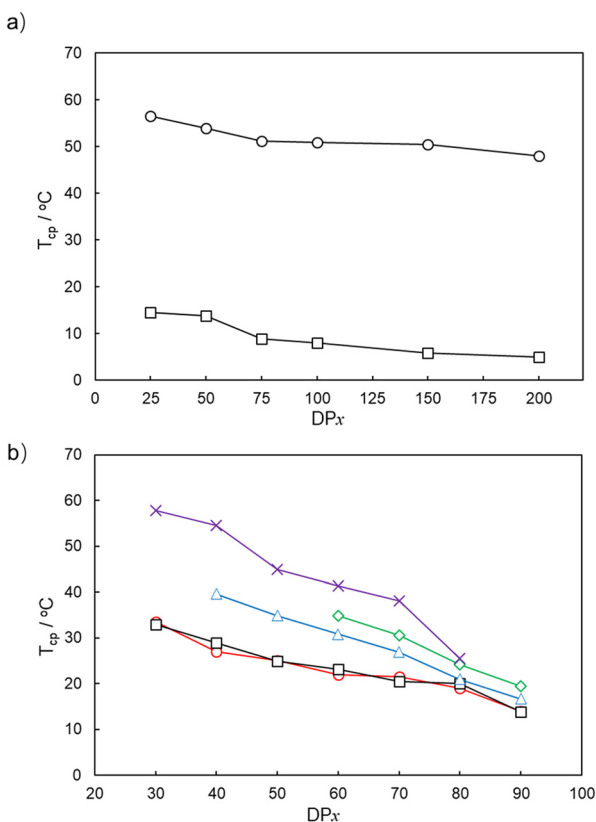


Fig. 5 Dependence of the cloud-point temperature (T_{cp}) on the degree of polymerization (DP_x) of (a) \square PEOEAm_x and \circ PMOEAm_x and (b) (\circ) PDMAm_y-b-PEOEAm_x, (\square) PEOEAm_x-b-PDMAm_y, (\diamond) PDMAm_y-b-PEOEAm_x, (Δ) PEOEAm_x-b-PDMAm_y-b-PEOEAm_x, and (\times) (PDMAm_{y/2}-b-PEOEAm_{x/2})₂.

group at the end of the side chain, making it more difficult for water molecules to penetrate into the gaps between the side chains. This inhibiting effect of the ethyl group on the penetration of water into the gaps between the side chains, together with the hydrophobicity of the ethyl group itself, is thought to contribute to the lower T_{cp} of PEOEAm.

The thermoresponsive properties of the di-, tri-, and tetra-block copolymers are summarized in Fig. 5b. T_{cp} decreased with increasing DP_x of PEOEAm_x for all the block copolymers and increased in the following order: PDMAm-b-PEOEAm \approx PEOEAm-b-PDMAm $<$ PEOEAm-b-PDMAm-b-PEOEAm $<$ PDMAm-b-PEOEAm-b-PDMAm $<$ (PDMAm-b-PEOEAm)₂. In the diblock copolymer system, the dependence of T_{cp} on DP_x was almost the same for both PDMAm-b-PEOEAm (14.0–33.6 °C) and PEOEAm-b-PDMAm (13.8–33.0 °C); this result supports the aforementioned conclusion that the copolymer obtained from the random GTcoP of EOEA and DMAm is not a random structure but a block structure, *i.e.*, PDMAm-b-PEOEAm. In the triblock copolymer system, the T_{cp} values for PEOEAm-b-PDMAm-b-PEOEAm were about 3.5 °C higher than those for PDMAm-b-PEOEAm-b-PDMAm. No T_{cp} was observed for PEOEAm₁₅-b-PDMAm₇₀-b-PEOEAm₁₅ because it was insoluble in water; this result was different from that obtained

for PEOEAm₃₀-b-PDMAm₇₀, which has a T_{cp} of 33.0 °C. Furthermore, PDMAm_y-b-PEOEAm_x-b-PDMAm_y ($x + 2y = 100$) exhibited no thermoresponsive behavior for $x < 50$; this is unlike the case of PEOEAm_x-b-PDMAm_y ($x + y = 100$), which exhibited thermal phase transition for x in the range of 30–90. These results indicated that the sequence of the two block segments in the triblock copolymer system greatly affected the thermoresponsive properties. For the tetra-block copolymer system, the thermoresponsive properties of (PDMAm-b-PEOEAm)₂ can be widely varied (T_{cp} varies from 25.5 to 57.8 °C), which is in contrast to the results obtained for PDMAm-b-PEOEAm, which has a narrow range of T_{cp} (14.0–33.6 °C). The thermoresponsive properties of PEOEAm can be controlled by forming its block copolymers with water soluble, non-thermoresponsive PDMAm, and the T_{cp} increases with an increasing number of blocks in the block copolymer in the following order: PEOEAm₅₀ (13.9 °C) $<$ PDMAm₅₀-b-PEOEAm₅₀ (25.1 °C) $<$ PEOEAm₂₅-b-PDMAm₅₀-b-PEOEAm₂₅ (34.9 °C) $<$ (PDMAm₂₅-b-PEOEAm₂₅)₂ (44.9 °C).

Finally, the thermal phase transition was further discussed using the hydrodynamic radii (R_h s) of block copolymers. Fig. S13[†] shows the distribution of R_h for PDMAm₅₀-b-PEOEAm₅₀, PEOEAm₅₀-b-PDMAm₅₀, PDMAm₂₅-b-PEOEAm₅₀-b-PDMAm₂₅, PEOEAm₂₅-b-PDMAm₅₀-b-PEOEAm₂₅, and (PEOEAm₂₅-b-PDMAm₂₅)₂. Table 5 summarizes the R_h values measured at 20 and 50 °C. For the diblock copolymer system, the R_h values after the phase transition were about 40 times those before the phase transition; R_h values changed from 11.6 to 428.8 nm for PDMAm₅₀-b-PEOEAm₅₀ and from 11.4 to 411.5 nm for PEOEAm₅₀-b-PDMAm₅₀. There was little difference between the two diblock copolymers because the copolymer obtained by the random GTcoP was the diblock copolymer, PDMAm-b-PEOEAm. For the triblock copolymer system, the R_h values increased after the phase transition; they increased from 11.1 to 510.5 nm for PEOEAm₂₅-b-PDMAm₅₀-b-PEOEAm₂₅, whose sizes were similar to those of the diblock copolymers. In contrast, the R_h values of PDMAm₂₅-b-PEOEAm₅₀-b-PDMAm₂₅, which is not thermoresponsive, increased only approximately four times from 12.3 (before the phase transition) to 47.0 nm (after the phase transition). In addition, the R_h value increased from 13.7 (before the phase transition) to 572.9 nm (after the phase transition) for (PEOEAm₂₅-b-PDMAm₂₅)₂; this result is similar to that obtained for the di- and triblock copolymer systems, except that PDMAm₂₅-b-PEOEAm₅₀-b-PDMAm₂₅ was insoluble in water. These results indicated that the di-, tri-, and tetra-block copolymers formed large, disordered aggregates in response to thermal stimuli, resulting in the aqueous copolymer solution changing from a clear to a turbid solution. Moreover, PDMAm_y-b-PEOEAm_x-b-PDMAm_y with $x = 30, 40$, and 50 and PEOEAm₁₅-b-PDMAm₇₀-b-PEOEAm₁₅ could not measure thermal phase transitions because the aqueous solutions of these triblock copolymers were cloudy. The hydrophobicity of these triblock copolymers was unexpected because PEOEAm and PDMAm are soluble in water, but we confirmed that the arrangement of PEOEAm and PDMAm in the multiblock co-

Table 5 Hydrodynamic radii (R_h) of PDMAm₅₀-*b*-PEOEA₅₀, PEOEA₅₀-*b*-PDMAm₅₀, PDMAm₂₅-*b*-PEOEA₅₀-*b*-PDMAm₂₅, PEOEA₂₅-*b*-PDMAm₅₀-*b*-PEOEA₂₅, and (PEOEA₂₅-*b*-PDMAm₂₅)₂ at 20 and 50 °C

Polymer	T_{cp}^a	R_h^b/nm	
		20 °C	50 °C
PDMAm ₅₀ - <i>b</i> -PEOEA ₅₀	25.1	11.6	428.8
PEOEA ₅₀ - <i>b</i> -PDMAm ₅₀	25.0	11.4	411.5
PDMAm ₂₅ - <i>b</i> -PEOEA ₅₀ - <i>b</i> -PDMAm ₂₅	—	12.3	47.0
PEOEA ₂₅ - <i>b</i> -PDMAm ₅₀ - <i>b</i> -PEOEA ₂₅	39.6	11.1	510.5
(PDMAm ₂₅ - <i>b</i> -PEOEA ₂₅) ₂	44.9	13.7	572.9

^a Determined by UV-vis measurements in water (10 g L⁻¹).

^b Determined by dynamic light scattering (DLS) measurements in water (3 g L⁻¹).

polymer has a significant effect on the thermoresponsive properties. Therefore, further studies on the aggregation structure of the multiblock copolymers are needed by measuring the critical micelle concentration (CMC) and by molecular dynamics (MD) simulations.

Conclusions

The hydrosilylation-promoted GTP of EOEAm with Me₂EtSiH using the B(C₆F₅)₃ catalyst can precisely synthesize PEOEA_n, with repeating units of EOEAm only and without initiator residues at the α chain end. The polymerization kinetics of the GTP of EOEAm or DMAm showed that the polymerization rate of EOEAm was much lower than that of DMAm. In the random GTcoP of EOEAm and DMAm, after all DMAm was consumed, the polymerization of EOEAm proceeded to produce not a statistical copolymer but a block copolymer, *i.e.*, the one-pot synthesis of PDMAm-*b*-PEOEA_n. This method was used to synthesize tri- and tetra-block copolymers. The block copolymer of PEOEA_n and PDMAm exhibited a thermal phase transition in which large, disordered aggregates formed in response to thermal stimuli, resulting in the aqueous copolymer solution changing from a clear to a cloudy solution. The T_{cp} values of 14.5–5.0 °C for PEOEA_n were 40 °C lower than those of 56.5–48.0 °C for PMOEAm. Furthermore, the T_{cp} increased as the number of blocks in the block copolymer increased in the following order: PDMAm-*b*-PEOEA_n ≈ PEOEA_n-*b*-PDMAm < PEOEA_n-*b*-PDMAm-*b*-PEOEA_n < PDMAm-*b*-PEOEA_n-*b*-PDMAm < (PDMAm-*b*-PEOEA_n)₂. The exceptions were PEOEA₁₅-*b*-PDMAm₇₀-*b*-PEOEA₁₅ and PDMAm_x-*b*-PEOEA₃₀-*b*-PDMAm_y ($x = 30, 40$, and 50) that did not undergo any phase transition, possibly because these triblock copolymers formed stable micelles consisting of a hydrophobic PEOEA_n core and a hydrophilic PDMAm shell; however, further clarification of this matter is needed. Even for a low-thermoresponsive polymer, by using its block copolymer with a water-soluble polymer, the thermoresponsive temperature could be controlled over a wide range by adjusting the DP ratio of two blocks and the number of blocks in the copolymer.

Conflicts of interest

The authors declare no conflict of interest.

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