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Organocatalytic controlled/living ring-opening polymerization of cyclotrisiloxanes initiated by water with strong organic base catalysts

Ring-opening polymerization of cyclotrisiloxanes using water as an initiator and strong organic bases such as guanidines possessing an R–N=C(N)–NH–R’ unit as catalysts produced a variety of well-defined symmetric polysiloxanes with controlled number-average molecular weights ($M_n = 2.64–102.3$ kg mol$^{-1}$) and narrow polydispersity ($D = 1.03–1.16$). Controlled syntheses of statistical and triblock copolymers were also achieved. Various terminal functionalities were successfully introduced by the end-capping reaction of propagating polysiloxanes using functional chlorosilanes. In this system, even non-dehydrated solvents are usable for the polymerization.

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Organocatalytic controlled/living ring-opening polymerization of cyclotrisiloxanes initiated by water with strong organic base catalysts†

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Organocatalytic controlled/living ring-opening polymerization of cyclotrisiloxanes, such as hexamethyldicyclosiloxane, 1,3,5-trimethyl-1,3,5-triphenylcyclosiloxane, 1,3,5-trimethyl-1,3,5-trivinylcyclosiloxane, and 1,3,5-trimethyl-1,3,5-tris(3,3,3-trifluoropropyl)cyclosiloxane, using water as an initiator and strong organic bases, such as amidines, guanidines, phosphazene bases, and proazaphosphatrane, as catalysts produced a variety of polysiloxanes with controlled number-average molecular weights ($M_n = 2.64–102.3$ kg mol$^{-1}$), narrow polydispersity ($D = 1.03–1.16$), and well-defined symmetric structures. Controlled syntheses of statistical copolymers and triblock copolymers were achieved by copolymerizations of two cyclotrisiloxanes. Various terminal functionalities were successfully introduced by the end-capping reaction of propagating polysiloxanes using functional chlorosilanes. Kinetic investigations demonstrated that the polymerization proceeded through the initiator/chain-end activation mechanism, namely activations of water in the initiation reaction and of terminal silanols in propagating polysiloxanes in the propagation reaction. Catalytic activities of strong organic bases were revealed to depend on their Brønsted basicity and efficiency of the proton transfer in the initiation and propagation reactions. Guanidines possessing an \( R – N = C ( N ) – N H – R' \) unit, in particular 1,3-trimethylene-2-propylguanidine, showed excellent performance as a catalyst. In this system, even non-dehydrated solvents are usable for the polymerization.

Introduction

Polysiloxanes are major components of organosilicon materials that have a wide range of applications including lubricants, insulating and coating materials, sealants, adhesives in the form of oil, rubber, gel, and resin due to their excellent chemical and physical properties.† Precise control of the linear, branched, and cross-linked structure of polysiloxanes is difficult, but would be essential for preparing the next generation of organosilicon materials with much superior properties. The linear polysiloxanes are industrially synthesized by (1) hydrolysis and simultaneous polycondensation of dichlorosilanes or dialkoxy silanes or (2) ring-opening polymerization (ROP) of cyclooligosiloxanes, in particular cyclotetrasiloxanes, using acidic species as catalysts or hydroxide salts as initiators, although the molecular weight distribution of the products becomes broad and the formation of cyclic oligomers as by-products are unavoidable.‡ Anionic ROP of cyclotrisiloxanes initiated by lithium compounds, such as organolithiums and lithium silanolates, has been the sole method to synthesize polysiloxanes and their copolymers with narrow molecular weight distributions and well-defined structures.† However, the biggest drawback of this method is that highly purified and dried starting materials must be used to avoid undesired side reactions, such as termination, backbiting, and chain-transfer reactions. The synthetic difficulty of polysiloxanes with a controlled structure has been a hurdle to actively employ them for developing organosilicon materials.

In the field of polymer science, there has been continuous motivation to develop controlled polymerization that can be conducted with simple starting materials and procedures. Organocatalytic polymerization has been studied over the last decades to realize it.† In particular, organocatalytic ROP of cyclic monomers, such as lactones, cyclic carbonates, epoxides, and cyclic phosphoesters, has been studied using organic acids, such as sulfonic acids, bis(sulfonyl)imides, and phosphoric acids, as well as organic bases, such as amines, amidines, guanidines, phosphazene bases, proazaphosphatranes, cyclopropenimines, and N-heterocyclic carbenes (NHC), as catalysts.‡ Many of them proceed in a controlled/living fashion.
and have been extensively applied to controlled synthesis of various well-defined polymers, including end-functionalized polymers, block copolymers, and star-shaped polymers. The success of the organocatalytic ROP inspired us to apply it to cyclooligosiloxanes and develop a convenient method to synthesize various polysiloxanes with well-defined structures.

ROP of cyclooligosiloxanes and cyclic carboilsiloxanes have been attempted with organic acids and bases as catalysts. Regarding acidic catalysts, HOSO2CF2, HN(SO2CF2)2, HB(C6F5)4, Ph3CB(C6F5)4, and B(C6F5)3 (ref. 27,28) have been used for ROP of hexamethylcyclotrisiloxane (D3(Me2)). None of them produced poly(dimethylsiloxane) (PDMS) with a narrow molecular weight distribution. For the basic catalysts, only those with quite high Bronsted basicity, such as phosphazene bases, NHCs, and bicyclic guanidines, have been employed. Möller and coworkers were the first to employ an electronically neutral strong organic base. They reported a polymerization of octamethylcyclotetrasiloxane (D4(Me2)) using 1-tert-butyl-4,4,4-tris(dimethylamino)-2,2-bis[tris(dimethylamino)phosphoranylidenamino]-2-l,4,5,6-tetramethylimidazol-2-ylidene and 1,3-di-t-butyl-4,4,4-tris(dimethylamino)-2,2-bis[tris(dimethylamino)phosphoranylidenamino]-2-l,4,5,6-tetramethylimidazol-2-ylidene, as catalysts and alcohols as initiators.

Hupfeld and Taylor also reported that a polymerization of D4(Me2) using tBu-P4 as a catalyst almost immediately produced PDMS with Mₙ of up to 4 x 10⁶ kg mol⁻¹ with D of 1.5–1.9. Clarke’s group reported that a polymerization of 1,3,5,7-tetramethyl-1,3,5,7-tetraphenylcyclotetrasiloxane (D₄(Me,Ph)) using tBu-P₄ as a catalyst and methanol as an initiator. The polymerization was very rapid and the resulting PDMS had a very high number-average molecular weight (Mₙ) up to 440 kg mol⁻¹, although polydispersity (D) was broad, 1.7–1.9. Hupfeld and Taylor also reported that a polymerization of D₄(Me2) using tBu-P₄ as a catalyst almost immediately produced PDMS with Mₙ of up to 4 x 10⁶ kg mol⁻¹ with D of 1.5–1.9. Clarke’s group reported that a polymerization of 1,3,5,7-tetramethyl-1,3,5,7-tetraphenylcyclotetrasiloxane (D₄(Me,Ph)) using tBu-P₄ as a catalyst and methanol as an initiator at ambient temperature produced poly[methyl(phenyl)siloxane] (PMPS) with Mₙ of 122–249 kg mol⁻¹ and D of 1.5. Baceiredo and colleagues reported ROP of D₃(Me2) using NHCs, such as 1,3-dicyclohexylimidazol-2-ylidene and 1,3-di-tert-butyl-4,5-dimethylimidazol-2-ylidene, as catalysts and alcohols as initiators to give PDMS with high D of 1.5–1.7. Waymouth and Hedrick and coworkers reported the only example that succeeded in obtaining PDMS with narrow D (<1.2) by organocatalytic ROP of D₃(Me2). They used 1,5,7-triazaabicyclo[4.4.0]dec-5-ene (TBD) as a catalyst and 4-pyrenebutan-1-ol as an initiator, although spectral and chromatographic data of the products were not provided. Thus, controlled synthesis of polysiloxanes by organocatalytic polymerization has not been well established. We considered that an appropriate choice of a catalyst and an initiator is the key to develop a controlled/living ROP of cyclooligosiloxanes. The acidity/basicity of the catalyst should not be too high to avoid side reactions, such as main chain scission and condensation of propagating polysiloxanes, that affect Mₙ, D, and terminal structures of resulting polysiloxanes. Furthermore, an initiation reaction between a cyclooligosiloxane and an initiator should generate propagating polysiloxanes with a sufficiently stable terminal structure at a sufficiently fast rate in comparison with a propagation reaction.

We herein report a controlled/living ROP of cyclotrisiloxanes using water as an initiator, strong organic bases as catalysts, and organochlorosilanes as end-capping agents. As shown in Scheme 1, the developed system is capable of polymerizing various cyclotrisiloxane in a controlled/living fashion and producing a variety of (telechelic) polysiloxanes with controlled Mₙ, narrow molecular weight distributions, and well-defined symmetric structure, which are difficult to obtain by the conventional anionic ROP using lithium compounds as initiators. This new polymerization method has a further advantage that non-dehydrated solvents can be used to give well-defined polysiloxanes.

**Results and discussion**

**Optimization of catalyst for the polymerization of D₃(Me2)**

We found that water initiates a polymerization of D₃(Me2) in the presence of strong organic bases acting as catalysts to give

![Scheme 1](Image)

Scheme 1. Ring-opening polymerization (ROP) of cyclotrisiloxanes using water as an initiator and strong organic bases as catalysts.
PDMS with narrow molecular weight distributions. We first optimized the catalysts for the polymerization by testing the catalytic activity of various strong organic bases with different Bronsted basicity and structural classes. The strong organic bases tested in this study were classified into the following four categories. Amides/guanidines A are amides and guanidines with no hydrogen atoms on their amino groups, such as 1,1,3,3-tetramethylguanidine (TMGa), 1,5-diazabicyclo[4.3.0]non-5-ene (DBN), 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) and 7-methyl-1,5,7-triazabicyclo[4.4.0]dec-5-ene (MTBD). Guanidines B are guanidines with an R–N=C(N)–NH–R’ unit, such as 1,5,7-triazabicyclo[4.3.0]non-6-ene (TBN) and TBD. Phosphazene bases include 2-tert-butylamino-2-diethylamino-1,3-dimethylperhydro-1,3,2-diazaphosphorine (BEMP), tert-butylamino-tri(pyrrolidino) phosphorane (Bu-PP (pyrr)), and 1-ethyl-2,2,4,4,4-pentakis(dimethylamino)-2X3,4X3-catena[n]phosphazene (Et-P2). As proazaphosphatranes, only 2,8,9-triisobutyl-2,5,8,9-tetraaza-1-phosphabicyclo[3.3.3]undecane (TIPB) was employed. The Bronsted basicity of the bases in MeCN (μMeCN)K_BH is shown in Scheme 1 and Table 1.

The catalytic activity of the bases was determined from the apparent rate coefficients of propagation (k_{app} h^{-1}) observed in the polymerizations in tetrahydrofuran (THF) at 30 °C under the conditions of \([\text{D}_{1/2}]_{0}=1.80 \text{ mol L}^{-1}\) and \([\text{D}_{1/2}]_{0}/[\text{H}_2\text{O}]_{0}/[\text{C (catalyst)}]_{0}=10/1/0.005\) for Et-P2 and TiBP, 10/1/0.007 for TBD, and 10/1/0.10 for other bases. The polymerization was initiated by adding a stock solution of water in THF and that of the catalysts in dry THF to a solution of D₃(Me₂) in dry THF in this order. A linear relationship between the polymerization time, \(t\) (h), and \(-\ln(1-c)\), where \(c\) is monomer conversion, was observed in the first-order kinetic plot of each polymerization. According to the eqn (1), the k_{app} values were determined from the slope observed in the plots.

\[
-\ln(1-c) = k_{app} t
\]  

(1)

The observed k_{app} values were normalized to k_{app} [H₂O]₀/[C₃]₀, as listed in Table 1 to directly compare k_{app} values observed in the polymerizations using different amounts of the catalysts. This conversion should be rational, since we confirmed an almost linear relationship of k_{app} with the initial molar ratio of the catalyst and water, [C₃]₀/[H₂O]₀, as discussed in the next section (see Fig. 5).

Fig. 1 shows the dependence of \(\log(k_{app})\) on \(\mu\text{MeCN}K_{BH}\) of strong organic bases. \(\mu\text{MeCN}K_{BH}\) instead of those in THF can be used for the discussion, since Leito’s group has reported that K_BH values in MeCN and THF have a good linear correlation.8,42 The catalytic activity of an organic base per its Bronsted basicity, \(k_{app}/10^{\mu\text{MeCN}K_{BH}}\), increased in the following order: TiBP < phosphazene bases < guanidines/amidines A < guanidines B, which indicated that \(\mu\text{MeCN}K_{BH}\) values of the catalysts were not the only single factor that affected the rates of polymerization. Interestingly, the amidines/guanidines A and the phosphazene bases independently showed linear relationships between \(\log(k_{app})\) and their \(\mu\text{MeCN}K_{BH}\).

Among the phosphazene bases and TiBP with \(\mu\text{MeCN}K_{BH}\) values of 27.5–33.5 (Table 1, entries 11–14), Et-P₂ (1.1 × 10⁷ h⁻¹) and TiBP (5.2 × 10⁷ h⁻¹) showed significantly higher catalytic activity than the other bases due to their very high Bronsted basicity. On the other hand, BEMP (0.14 h⁻¹) and Bu-PP (pyrr) (0.60 h⁻¹) showed catalytic activities that were only comparable to MTBD (0.29 h⁻¹) and TBN (0.16 h⁻¹), which have more than 100 times weaker Bronsted basicity. The very high Bronsted basicity of the phosphazene bases and TiBP caused the frequent occurrence of undesired side reactions (vide infra), in particular, in the polymerizations catalyzed by Et-P₂ and Bu-PP (pyrr), which made these bases unfavorable for controlling the polymerization as can be seen from the \(D\) of the products in each polymerization.

Regarding the amidines/guanidines A with \(\mu\text{MeCN}K_{BH}\) values of 23–25.5 (Table 1, entries 1–4), TMGa (0.0026 h⁻¹), DBN

| Entry | Catalyst | MeCNpK_BH | k_{app} b (h⁻¹) | D of the product (Conv. (%)) |
|-------|----------|-----------|----------------|-----------------------------|
| 1     | TMGa     | 23.3      | 0.0026         | 1.08 (50.7)                 |
| 2     | DBN      | 23.79     | 0.019          | 1.10 (98.5)                 |
| 3     | DBU      | 24.34     | 0.039          | 1.11 (99.1)                 |
| 4     | MTBD     | 25.43     | 0.29           | 1.11 (99.6)                 |
| 5     | TBD      | d         | 0.0036         | n.d.¹                      |
| 6     | TMGb     | d         | 0.093          | 1.13 (99.3)                 |
| 7     | TBN      | 24.55     | 0.16           | 1.11 (99.9)                 |
| 8     | TMIPG    | d         | 1.7            | 1.11 (99.5)                 |
| 9     | TmNPG    | d         | 2.6            | 1.11 (98.0)                 |
| 10    | TBD      | 25.96     | 6.3            | 1.12 (99.1)                 |
| 11    | BEMP     | 27.58     | 0.14           | 1.14 (99.2)                 |
| 12    | tBu-P₂ (pyrr) | 28.35 | 0.60          | 1.23 (99.5)                 |
| 13    | Et-P₂    | 32.94     | 1.1 × 10⁵      | 1.22 (99.8)                 |
| 14    | TiBP     | 33.33     | 5.2 × 10⁵      | 1.13 (99.9)                 |

¹ The polymerizations were carried out under the conditions of \([\text{D}_{1/2}]_{0}=1.80 \text{ mol L}^{-1}\) and \([\text{D}_{1/2}]_{0}/[\text{H}_2\text{O}]_{0}/[\text{C (catalyst)}]_{0}=10/1/0.005\) for Et-P₂ and TiBP, 10/1/0.007 for TBD, and 10/1/0.10 for the other bases.

² Calculated with \(k_{app} = k_{app}[\text{H}_2\text{O}]_{0}/[\text{C}_3]_{0}\). 

D of the obtained PDMS at the indicated conversion of monomer. 

Not reported.

*Not determined.

**Fig. 1. A dependence of the logarithm of the normalized apparent rate coefficients of propagation (log(k_{app})) on \(\mu\text{MeCN}K_{BH}\) of strong organic bases observed in the polymerizations of D₃(Me₂) using water as an initiator and strong organic bases as a catalyst in THF at 30 °C under the conditions of \([\text{D}_{1/2}]_{0}/[\text{H}_2\text{O}]_{0}=10\) and \([\text{D}_{1/2}]_{0}=1.80 \text{ mol L}^{-1}.**
(0.019 h⁻¹), DBU (0.039 h⁻¹), and MTBD (0.29 h⁻¹), showed only low catalytic activity, although undesired side reactions almost never occurred within the observed time range. In contrast, the guanidines B with \(^{13} \text{C} \text{NMR} \) \( p \text{K}_\text{BH} \) values of 24.5–26 (Table 1, entries 7 and 10), such as TBD (6.3 h⁻¹) and TBN (0.16 h⁻¹), showed much higher \( k_{\text{app}}/10 \text{MeC} \text{N} \text{H} \text{OSiMe}_2 \) than the amidines/guanidines A as shown in Fig. 1, even though their structures and Bronsted basicity were not very different. For example, TBD showed 22 times higher catalytic activity than MTBD, although its Bronsted basicity was only around 3.4 times stronger than MTBD. This high catalytic activity of the guanidines B may have originated from the R–N=C(N)–NH–R’ unit that enables high efficiency of the initiation and propagation reactions as described in the section regarding the mechanism (see Scheme 3).

We hence evaluated the catalytic activity of four more guanidines B of which \(^{29} \text{Si} \text{NMR} \) values have not been reported, such as 1,4,6-triazabicyclo[3.3.0]oct-4-ene (TBO), 1,1,2,3-tetramethylguanidine (TMG), 1,3-trimethylene-2-isopropylguanidine (TMiPG), and 1,3-trimethylene-2-propylguanidine (TMnPG), since undesired side reactions rather frequently occurred in the polymerizations catalyzed by TBD and TBN. TMG showed much lower catalytic activity than TBD, showed high catalytic activity next to TBD. The Brønsted basicity of the guanidines B may have originated from the R–N=C(N)–NH–R’ unit that enables high efficiency of the initiation and propagation reactions as described in the section regarding the mechanism (see Scheme 3).

Fig. 2 SEC chromatograms of the PDMS-(OH)\(_2\) (\( M_{n, \text{NMR}} = 2.64 \text{ kg mol}^{-1}, D = 1.14 \)) and the PDMS-(OSiMe\(_3\)Ph)\(_2\) (\( M_{n, \text{NMR}} = 3.17 \text{ kg mol}^{-1}, D = 1.11 \)) synthesized in THF at 30 °C under the conditions of [D\(_3\)Me\(_2\)]\(_0\)/(H\(_2\)O)\(_0\)/[TMnPG]\(_0\) = 10/1/0.10 and [D\(_3\)Me\(_2\)]\(_0\) = 1.80 mol L\(^{-1}\) to give \( \pi, \pi\text{-dihydroxy-}

\[ \text{PDMS-(OH)}_2 \]

\[ \text{PDMS-(OSiMe}_3\text{Ph)}_2 \]

end-capped PDMS (PDMS-(OH)\(_2\)) with \( M_n \) estimated by \(^1\text{H NMR} \) measurement \( (M_{n, \text{NMR}}) \) of 2.64 kg mol\(^{-1}\) and \( D = 1.14 \) (Table 2, entry 1). The observed \( M_{n, \text{NMR}} \) corresponded well to the \( M_n \) calculated of 2.20 kg mol\(^{-1}\). The conversion of monomer reached 98.0% after 90 min of polymerization. The purification of the product was easily achieved by neutralizing TMnPG with an excess amount of benzoic acid and by washing the concentrated oily crude product with acetonitrile (MeCN). Removal of the MeCN layer with a pipette and drying the residue under vacuum gave pure PDMS-(OH)\(_2\) as a colorless viscous liquid. End-capping of the propagating PDMS-(OH)\(_2\) was achieved by directly adding pyridine (8 equiv.) and chlorodimethylphenylsilane (Me\(_2\)PhSiCl, 5 a)-dihydroxy-

\[ \text{PDMS-(OH)}_2 \]

\[ \text{PDMS-(OSiMe}_3\text{Ph)}_2 \]

presumably due to its much lower Brønsted basicity than TBD and TBN.\(^{43}\) TMiPG (0.093 h⁻¹), an acyclic tetramethylguanidine with an R–N=C(N)=NH–R’ unit, showed 36 times higher catalytic activity than TMGa. The replacement of an H–N=C(N)=NH–Me unit certainly contributed to the increase in the catalytic activity. TMiPG (1.7 h⁻¹) and TMnPG (2.6 h⁻¹), monocyclic guanidines with a six-membered ring and the same number of carbon atoms as TBD, showed high catalytic activity next to TBD. The Bronsted basicity of TMnPG has been reported to be at least higher than TBN in CD\(_2\)OD/D\(_2\)O = 8/2 (w/w).\(^{43}\) TMiPG showed a slightly lower catalytic activity than TMnPG presumably because of the higher steric hindrance of the isopropyl group than that of the propyl group. It is noteworthy that the undesired side reactions were much less frequent in the polymerization catalyzed by TMiPG and TMnPG in comparison with that catalyzed by other bases with comparable catalytic activity. Hence, we identified TMnPG as the most suitable catalyst for the polymerization of cyclotrisiloxanes initiated by water.

Fig. 3 \(^1\text{H} \) and \(^{29} \text{Si} \) \(^1\text{H}\) NMR spectra of the synthesized PDMSs (Table 2, entry 1) in CDCl\(_3\): (a) \(^1\text{H} \) NMR spectrum of PDMS-(OH)\(_2\), (b) \(^1\text{H} \) NMR spectrum of PDMS-(OSiMe\(_3\)Ph)\(_2\), (c) \(^{29} \text{Si} \) \(^1\text{H}\) NMR spectrum of PDMS-(OH)\(_2\), (d) \(^{29} \text{Si} \) \(^1\text{H}\) NMR spectrum of PDMS-(OSiMe\(_3\)Ph)\(_2\).
Table 2  Polymerizations of \(D_{3}^{(Me2)}\) initiated by water and catalyzed by TMnPG at 30 °C\(^a\)

| Entry | \([D_{3}^{(Me2)}]_0\) | Solvent\(^b\) | End-capping agent | Time (h) | Conv (%)\(^c\) | \(M_n\) (kg mol\(^{-1}\)) | SEC\(^e\) | NMR\(^f\) | \(D\) | \(k_{p, app}\) (h\(^{-1}\)) |
|-------|-----------------|-----------|-----------------|----------|-----------|----------------|--------|--------|-----|------------|
| 1     | 10/1/0.10       | THF       | None            | 1.5      | 98.0      | 2.20           | 1.66   | 2.64   | 1.14 | 2.6        |
| 2     | 10/1/0.05       | THF       | Me\(_2\)PhSiCl  | 3        | 98.1      | 2.47           | 1.49   | 2.85   | 1.13 | 1.3        |
| 3     | 10/1/0.01       | DMac/THF  | Me\(_2\)PhSiCl  | 0.50     | 98.6      | 2.21           | 0.83   | 2.81   | 1.33 | 8.2        |
| 4     | 10/1/0.10       | CH\(_2\)Cl/THF | None      | 1        | 98.6      | 2.21           | 1.80   | 2.70   | 1.12 | 4.6        |
| 5     | 25/1/0.25       | CH\(_2\)Cl/THF | None      | 2        | 99.5      | 5.54           | 4.09   | 5.70   | 1.06 | 2.4        |
| 6     | 50/1/0.50       | CH\(_2\)Cl/THF | Me\(_2\)PhSiCl | 1.57     | 93.8      | 10.7           | 9.75   | 12.2   | 1.04 | 1.7        |
| 7     | 100/1/1.0       | CH\(_2\)Cl/THF | Me\(_2\)PhSiCl | 3        | 97.3      | 21.9           | 15.6   | 18.9   | 1.04 | 1.2        |
| 8     | 200/1/2.0       | CH\(_2\)Cl/THF | Me\(_2\)PhSiCl | 5.5      | 95.0      | 42.5           | 31.8   | 39.9   | 0.64 | 0.67       |
| 9     | 550/1/5.5       | CH\(_2\)Cl/THF | 97/3 | None      | 18       | 93.0          | 113.8  | 80.0   | 102.3| 0.03       |
| 10    | 10/1/0.10       | CH\(_2\)Cl/THF | Me\(_2\)HSiCl  | 1        | 99.5      | 2.35           | 1.60   | 2.95   | 1.15 | 0.8        |
| 11    | 10/1/0.10       | CH\(_2\)Cl/THF | Me\(_2\)ViSiCl  | 1        | 99.1      | 2.39           | 1.71   | 3.00   | 1.14 | 0.4        |
| 12    | 10/1/0.10       | CH\(_2\)Cl/THF | AllyMe\(_2\)SiCl | 1        | 99.5      | 2.42           | 1.67   | 2.77   | 1.13 | 0.7        |
| 13    | 10/1/0.10       | CH\(_2\)Cl/THF | (CICH\(_2\))Me\(_2\)SiCl | 1 | 99.5 | 2.44 | 1.84 | 3.14 | 1.10 |
| 14    | 10/1/0.10       | CH\(_2\)Cl/THF | (BrCH\(_2\))Me\(_2\)SiCl | 1 | 98.8 | 2.52 | 1.71 | 3.08 | 1.13 |
| 15    | 10/1/0.10       | CH\(_2\)Cl/THF | Me\(_2\)C\(_6\)F\(_5\))SiCl | 1 | 99.3 | 2.68 | 1.72 | 3.38 | 1.13 |
| 16    | 10/1/0.10       | CH\(_2\)Cl/THF | [EtO\(_3\)]Me\(_2\)SiCl | 1 | 99.5 | 2.57 | 1.84 | 3.25 | 1.11 |

\(^a\) \([D_{3}^{(Me2)}]_0 = 1.80 \text{ mol L}^{-1}\). \(^b\) Volume ratios of two solvents are shown for the mixed solvents. \(^c\) Determined by \(^1\)H NMR. \(^d\) Calculated from \(M_n\)calcd = \([D_{3}^{(Me2)}]_0[H_2O]_0[TMnPG]_0 \times \text{Conv.} \times \text{MW. of } [D_{3}^{(Me2)}]_0 = 222.46 + \text{MW. of terminal structures}. \(^e\) Determined by SEC measurements in THF using polystyrene standards.

equiv.) to the reaction mixture at 30 °C and continuing the reaction for 15 min. The purification of the end-capped product (PDMS-(OSiMe\(_2\)Ph)\(_2\)) was also easily achieved by washing the concentrated crude product with methanol or MeCN.\(^{44}\) \(M_n\)NMR and \(D\) of the obtained PDMS-(OSiMe\(_2\)Ph)\(_2\) were determined to be 3.17 kg mol\(^{-1}\) and 1.11, respectively. The end-capping reaction hence did not affect the \(D\) of the original product. Fig. 2 shows the molecular weight distribution of the products measured by the size-exclusion chromatography (SEC) using THF as an eluent and polystyrene as a calibration standard.

The well-defined structures of the obtained PDMS-(OH)\(_2\) and PDMS-(OSiMe\(_2\)Ph)\(_2\) were proven by \(^1\)H and \(^29\)Si\(^{[1]}\)H NMR analysis as shown in Fig. 3. In each spectrum, signals due to the terminal groups and several monomeric units from the termini were separately observed from that due to the inner repeating units. In both \(^1\)H and \(^29\)Si\(^{[1]}\)H NMR spectra of the PDMS-(OSiMe\(_2\)Ph)\(_2\), the signals due to the PDMS-(OH)\(_2\) were not observed, which evidenced the quantitative end-capping of the PDMS-(OH)\(_2\).

Structures of the obtained PDMS-(OH)\(_2\) and PDMS-(OSiMe\(_2\)Ph)\(_2\) were further analyzed by positive ion matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS) using trans-2-[3-(4-tert-butylphenyl)-2-methyl-2-propenylidene]malononitrile (DCTB) as a matrix and sodium trifluoroacetate as a cationization agent. Only one series of peaks was observed in both of the spectra shown in Fig. 4, which indicated that the obtained PDMS-(OH)\(_2\) and PDMS-(OSiMe\(_2\)Ph)\(_2\) consisted of only linear PDMS with two terminal hydroxy groups and two terminal dimethyl(phenyl)siloxy groups, respectively. The \(m/z\) values of the observed molecular ion peaks corresponded to the calculated molar mass of PDMS-(OH)\(_2\) and PDMS-(OSiMe\(_2\)Ph)\(_2\) cationized by a sodium cation. Interestingly, the obtained PDMSs contained PDMSs of which the degree of polymerization was not a multiple of 3, although those PDMSs should not be produced if only a simple ring-opening reaction of \(D_{3}^{(Me2)}\) occurred in the polymerization. This point will be discussed in a later section regarding the mechanism of the polymerization.

As already mentioned in the previous section, a linear relationship was observed in the first-order kinetic plot of the polymizations carried out with \([D_{3}^{(Me2)}]_0[H_2O]_0[TMnPG]_0 = 10/1/0.10 and 10/1/0.05 (Table 2, entries 1 and 2) as shown in Fig. 5a. The \(k_{p, app}\) for the former and the latter polymerizations were determined to be 2.6 and 1.3, which suggested that the \(k_{p, app}\) varies linearly with the initial ratio of the catalyst and water, i.e., \([C]_0/[H_2O]_0\).

Fig. 5b shows the dependence of \(M_n\)NMR and \(D\) on monomer conversion (c). In both of the polymerizations, \(M_n\)NMR increased linearly as \(c\) increased, while \(D\) remained in a range of 1.08–1.14 even until the late stage of polymerization. It was hence found that the propagation reaction was the dominant process in the polymerizations and undesired side reactions were not frequent.
The occurrence of undesired side reactions in this medium proceeded as an inhomogeneous reaction, which was sensitive to the reaction conditions of the conventional anionic ROP using specially synthesized lithium compounds as an initiator. PDMS end-functionalized with dimethylsiloxanes with halogenomethyl, 2,3,4,5,6-pentafluoroethyl, dimethylvinylsiloxanes, and triethoxysilyl groups as shown in Fig. S9† were obtained by the end-capping with the corresponding chlorosilanes at 30 °C for 15 min after the polymerization. The products exhibited narrow $D$ ranging from 1.10 to 1.15. Notably, telechelic polysiloxanes with halogenomethyl, 2,3,4,5,6-pentafluoroethyl, and triethoxysilyl groups with narrow $D$ values were also obtained by our method, although these functional groups are potentially sensitive to the reaction conditions of the conventional anionic ROP of cyclotrisiloxanes using lithium compounds.

It is worth noting that, in contrast to the conventional anionic ROP initiated by lithium compounds, even non-dehydrated solvents were usable for the polymerization. $D_3^{(Me_2)}$ using non-dehydrated THF (purity: >99.5%, stabilizer-free) as a solvent produced PDMS-(OSiMe2Ph)2 with $M_{n,NMR}$ of 2.89 kg mol$^{-1}$ and $D$ of 1.11 under the same conditions as the polymerization shown in entry 1 of Table 2. Non-dehydrated CH$_2$Cl$_2$ (purity: >99.5%, stabilized with 2-methylbut-2-ene) was also usable (see ESIF). The maximum $M$s reachable with non-dehydrated reagents and solvents depend on their water contents.  

**Scope of monomers**

Controlled/living polymerizations of cyclotrisiloxanes other than $D_3^{(Me_2)}$ were also achieved using water as an initiator and TMnPG.
as a catalyst. 1,3,5-Trimethyl-1,3,5-triphenylsiloxane (D(Me,Ph)), cis/trans = 24/76, 1,3,5-trimethyl-1,3,5-trivinylsiloxane (D(Me,Vi)), cis/trans = 23/77, and 1,3,5-trimethyl-1,3,5-tris(3,3,3-trifluoropropyl)cyclosiloxane (D(Me,TfPr)), cis/trans = 20/80 were polymerized under THF at 30 °C under the conditions of [M]0/[H2O]0 = 10/1 and [M]0 = 1.80 mol L−1. Chloro-triethylysilane (Et3SiCl) was used as an end-capping agent for the polymerization of D(Me,Ph), while Me2PhSiCl was used for the polymerizations of D(Me,Vi) and D(Me,TfPr) to give well-defined PMPS, poly[methyl(vinyl)siloxane] (PMVS), and poly[methyl(3,3,3-trifluoropropyl)siloxane] (PMTFPS) with M[n] by 1H NMR spectra and the silicon atoms in the inner repeating units of the three polymers also showed three different signals in the 1H NMR spectra and the silicon atoms in the inner repeating units of the three polymers also showed three different signals in the 29Si{1H} NMR spectra (Fig. S16–18).46 MALDI-TOF MS spectra shown in Fig. S16–18 evidenced that the obtained PMPS, PMVS, and PMTFPS consisted of only linear PMPS with two terminal triethylysilox groups as well as PMVS and PMTFPS with two terminal dimethyl(phenyl)siroxol groups. The polymerizations needed to be terminated when the monomer conversion reached 80–90%, since the condensation of two propagating polymers and the backbiting (vide infra) became noticeable.47 Indeed, the formation of D(Me,Ph) and 1,3,5,7-tetramethyl-1,3,5,7-tetra(nvinyl)cyclosiloxane (D(Me,Vi)) was observed in the polymerizations of D(Me,Ph) and D(Me,Vi) (Table 3, entries 1–4 and Fig. S2 and S3). Mn values of PMPS, PMVS, and PMTFPS were also controllable by changing [M]0/[H2O]0 (Table 3, entries 1 vs. 2, entries 3 vs. 4, and entries 5 vs. 6) as already demonstrated for PDMS.48 The values of k,papp = k,papp_[H2O]0/[10(C1)0] observed in the polymerizations of D(Me,Ph) (17 h–1, Table 3, entry 1), D(Me,Vi) (35 h–1, Table 3, entry 3), and D(Me,TfPr) (99 h–1, Table 3, entry 5) carried out with [M]0/[H2O]0 = 10 were all higher than that observed for D(Me) (4.6 h–1, Table 2, entry 4), presumably due to the effect of electron-withdrawing phenyl, vinyl, and 3,3,3-trifluoropropyl groups, which increased the electrophilicity of the silicon atoms in the three monomers. Interestingly, the polymerizations of D(Me,Ph) showed almost 1 h of an induction period at the initial stage (Table 3, entries 1 and 2), although both of them produced PMPS with narrow D. Considering from the mechanism of the polymerization discussed in a later section, the reaction of water and D(Me,Ph) would proceed slowly while the activation of water by TMnPG was much more dominant than that of a,ω-di-hydroxy-terminated oligomethyl(phenyl)siloxanes (vide infra).

In contrast to cyclosiloxanes, tetrasiloxanes and cocyclotetrasiloxanes, such as D(Me,Vi)49 and D(Me,TfPr), were almost non-reactive by the polymerization of D(Me,Ph) and D(Me,Vi) (Table 3, entries 3–9), even though [Bu4N]49p [(MeCN)2Bu4][HBF4] = 42.7)49 and NHs, such as 1,3-di-alkylimidazol-2-yldiene and 1,3-dialkyl-4,5-dimethylimidazol-2-yldiene (estimated [Me6C6pHBF4] = 32–34 and 34–34, respectively),46 with much higher Brønsted basicity than TMnPG are known to catalyze the polymerization of cyclotetrasiloxanes,29,32

**Characteristics of copolymerization**

Both statistical copolymerization and block copolymerization of two cyclosiloxanes were achieved with the polymerization using water as an initiator and TMnPG as a catalyst. Statistical copolymerization of D(Me,Vi) and D(Me,Vi) was achieved by periodically adding 1 equiv. (to the initial amount of water) of D(Me,Vi)8 times into a reaction mixture of the polymerization of

| Entry | Monomer | [M]0/[H2O]0/[C1]0 | CH2Cl2/THF (v/v) | End-capping agent | Time (min) | Conv. (%) | M[n] (kg mol−1) | k,papp (h−1) | k,papp (h−1) |
|-------|---------|------------------|-----------------|-----------------|------------|-----------|--------------|-------------|-------------|
| 1     | D(Me,Ph) | 10/1/0.01        | 57/43           | Et3SiCl         | 219        | 92.4      | 6.8          | 4.02        | 1.7         |
| 2     | D(Me,Ph) | 30/1/0.03        | 79/21           | Et3SiCl         | 240        | 83.3      | 10.6         | 10.5        | 1.0         |
| 3     | D(Me,Ph) | 10/1/0.01        | 57/43           | Me2PhSiCl       | 72         | 95.5      | 3.1          | 2.76        | 1.1         |
| 4     | D(Me,Vi) | 30/1/0.03        | 79/21           | Me2PhSiCl       | 51         | 80.6      | 5.0          | 6.65        | 1.8         |
| 5     | D(Me,TfPr) | 10/1/0.004  | 70/30           | Me2PhSiCl       | 40         | 92.8      | n.d.         | 4.63        | 1.2         |
| 6     | D(Me,TfPr) | 30/1/0.012       | 88/12           | Me2PhSiCl       | 100        | 94.2      | n.d.         | 4.63        | 1.2         |
| 7     | D(Me,Vi) | 7.5/1/0.10       | 65/35           | None            | 1500       | 10.6      | n.d.         | 0.29        | n.d.        |
| 8     | D(Me,Vi) | 7.5/1/0.01       | 57/43           | None            | 1500       | 1.5       | n.d.         | 0.031       | n.d.        |
| 9     | D(Me,Vi) | 6/1/0.10         | 65/35           | None            | 1500       | 1.5       | n.d.         | 0.031       | n.d.        |
| 10    | D(Me,Vi) + D(Me,Vi) | 25 + 8/1/0.05 | 70/30          | Me2PhSiCl       | 540        | 64.9, >99.9 | n.d.         | 6.02        | 1.3         |
| 11    | D(Me,Vi) | 25/1/0.05        | 67/33           | Me2PhSiCl       | 540        | 84.9      | n.d.         | 4.74        | 1.0         |
| 12    | D(Me,Vi) | 7/1/0.25         | 84/16           | Et3SiCl         | 60         | 92.7      | n.d.         | 5.17        | 1.0         |

a Determined by 1H NMR. b Mol% of the monomer converted to the corresponding polysiloxanes and cyclotetrasiloxanes. c Calculated from [M]0,calculated = [M]0/[H2O]0 × (Conv. to polymer) × (MW. of monomer) × (MW. of terminal structures). d Determined by SEC measurements in THF using polystyrene standards. * Not determined. ** Around 1 h of an induction period was observed.
D$_3^{(Me_2)}$ in CH$_2$Cl$_2$/THF (70/30, v/v) at 30 °C under the conditions of [D$_3^{(Me_2)}$]$_0$/[H$_2$O]$_0$/[TMnPG]$_0$ = 25/1/0.05 (Table 3, entry 10) in a manner similar to the ‘semibatch’ method performed by Cypryk’s group for a copolymerization of D$_3^{(Me_2)}$ and D$_3^{(Me,Ti-Pr)}$, using n-butyllithium as an initiator.$^{28}$ D$_3^{(Me,Vi)}$ was added at 28, 60, 95, 135, 182, 238, 308, and 401 min from the initiation so that a constant amount of D$_3^{(Me_2)}$ was consumed in the time interval between the two additions. The number-average content of the dimethylsiloxane and methyl(vinyl)siloxane monomeric sequences in the obtained copolymer were NMR analysis (Fig. S19) previously observed in a copolymerization of D$_3^{(Me_2)}$D$_3^{(Me_2)}$D$_3^{(Me_2)}$D$_3^{(Me_2)}$/D$_3^{(Me_2)}$D$_3^{(Me,Vi)}$D$_3^{(Me_2)}$D$_3^{(Me,Vi)}$. In contrast, copolymerization of D$_3^{(Me_2)}$D$_3^{(Me,Vi)}$D$_3^{(Me,Vi)}$ units. In contrast, copolymerization of D$_3^{(Me_2)}$ and D$_3^{(Me,Vi)}$ by adding TMnPG to a premixed solution of the two monomers resulted in a rapid and preferential polymerization of D$_3^{(Me_2)}$ as shown in Fig. S20.$^{31}$ The monomeric sequence of the obtained copolymer was visualized based on the population of the triad monomeric sequences, n(D$_3^{(Me_2)})$/n(D$_3^{(Me,Vi)})$, and D as shown in Fig. S21,$^{31}$ which indicated the almost segregated presence of D$_3^{(Me_2)}$D$_3^{(Me_2)}$ and D$_3^{(Me,Vi)}$D$_3^{(Me,Vi)}$ units. In contrast, copolymerization of D$_3^{(Me_2)}$ and D$_3^{(Me,Vi)}$ by adding TMnPG to a premixed solution of the two monomers resulted in a rapid and preferential polymerization of D$_3^{(Me,Vi)}$ as shown in Fig. S22,$^{31}$ although the conversions of D$_3^{(Me_2)}$ in the first polymerization were not quantitative in both of the syntheses. This result indicated that D$_3^{(Me_2)}$ almost did not react in the second stage of the polymerization due to very low reactivity of the propagating end of PMVS and PDPS against D$_3^{(Me_2)}$. The obtained PDPS-b-PDMS-b-PDPS had a much narrower molecular weight distribution (D = 1.06) than those synthesized by the conventional anionic ROP using dithium diphenylsilanediolate as an initiator (D = 1.4–1.8).$^{34}$ These successful block copolymerizations demonstrated the controlled/living nature of theROP of cyclosiloxanes using water and strong organic bases.

**Mechanism of the polymerization**

It is worth discussing the mechanism of the polymerization catalyzed by the strong organic bases, since it has different characteristics compared to that of the conventional anionic ROP of cyclosiloxanes using hydroxide salts, organolithiums, and lithium silanlates as initiators. The structure of the propagating end in the former is a silanol, while that of the latter is a silanolate. Scheme 2 shows the possible elementary reactions in the ROP of cyclosiloxanes using water as an initiator and strong organic bases as catalysts. The polymerization is considered to proceed through the initiator/chain-end activation mechanism. (a) In the initiation reaction (Scheme 2a), a strong organic base first activates a water molecule. The ring-opening reaction of a cyclosiloxane occurs through the nucleophilic attack of the activated water at a silicon atom of a cyclosiloxane, followed by a proton transfer from the water to the cyclosiloxane to form a 1,5-dihydroxytrisiloxane. (b) The propagation reaction (Scheme 2b) proceeds through the activation of a terminal silanol group of a propagating polysiloxane by the strong organic base. The degree of polymerization of a propagating polysiloxane, n, increases by three after one propagation reaction. $K_w$ and $K_p$ can be defined as the equilibrium constants for the activation of water and a terminal silanol, and $k_t$ and $k_p$ can be defined as the rate coefficients for the ring-opening reaction of a monomer with the activated water and terminal silanol. The high catalytic activity of the
guanidines B may have originated from the high efficiency of the proton transfer in the initiation and propagation reactions. The R–N═C(N)–NH–R’ unit in the guanidines B would simultaneously activate water or a silanol on a propagating polymer by the imino group and transfer the hydrogen atom on the amino group to an incoming monomer based on the ‘proton shuttling mechanism’ as depicted in Scheme 3a. In contrast, the bases in the other categories are incapable of ‘proton shuttling’ and need to directly transfer the hydrogen atom as depicted in Scheme 3b. $K_p$ would increase as the acidity of the terminal silanol increases and the steric hindrance of the silanol decreases. $k_p$ would increase as nucleophilicity of the activated silanol increases, electrophilicity of the monomer increases, and the steric hindrance of the silanol and the monomer decrease. (c) ‘Intermolecular transfer’ of a terminal hydroxysilyl group from one propagating polymer to another one (disproportionation) can be considered to occur by a nucleophilic attack of the activated terminal silanol to another terminal silanol (Scheme 2c). This reaction increases and decreases the degree of polymerization of the two propagating polymers by one and causes the formation of polysiloxanes, of which degree of polymerization is not a multiple of 3, as observed in Fig. 4. A model reaction using 1,5-dihydroxy-1,1,3,3,5,5-hexamethyltrisiloxane ($D_3^{(Me_2)_3}(OH)_2$) and TMnP produced a mixture of $\omega,\omega$-dihydroxy-terminated permethyloligosiloxanes with different numbers of siloxy units in 6 min in THF at 30 °C under the conditions of $[D_3^{(Me_2)_3}(OH)_2]_0/[\text{TMnP}]_0=1/0.1$ (Fig. S6†). Similar reactions, ‘intermolecular transfer’ of terminal lithium silanolates on PDMS14,59 as well as ‘disproportionation’ of hydroxy-terminated oligodimethylsiloxanes60–62 have also been reported to occur in the conventional anionic ROP of $D_3^{(Me_2)_3}$ as well as in a transformation of oligosiloxanol in the presence of a strong base. The intermolecular transfer is an allowable side reaction since it does not largely affect $M_n$ and $Đ$ of the resulting polysiloxanes. (d) Condensation (dimerization, Scheme 2d) also occurs during the polymerization by a similar mechanism to (c) as an undesired side reaction. As already mentioned, the condensation of two propagating polysiloxanes becomes pronounced, in particular in the late stage of polymerization, and causes increasing of $Đ$ of the resulting polysiloxanes. It occurs much less frequently than (c). The similar tendency was observed in the disproportionation of oligosiloxanol using phosphazenium hydroxide.62 (e) Backbiting (depolymerization, Scheme 2e) occurs by intramolecular nucleophilic attack of the activated terminal silanol of a propagating polymer to a siloxyl unit in its inner repeating units. The backbiting produces cyclooligosiloxanes, such as
cycloptetrasiloxane and cyclopentasiloxane, and reduces the degree of polymerization and yield of propagating polymers. On the other hand, (f) the intermolecular chain-transfer reaction (Scheme 2f), which is known to occur in a conventional anionic ROP of cyclotrisiloxane,14 did not occur at least in the polymerization of D(Me2)3 catalyzed by TMnPG in THF at 30 °C, although it is a possible undesired side reaction. Trimethylsiloxy-terminated PDMS was not produced in a model polymerization of D(Me2)3 in the presence of dodeca-methylpentasiloxane (see ESI, Fig. S7†). The difference in the chances of side reactions (e) and (f) would be originated from the fact that the silanol and the silicon atom to be attacked always closely present for the former. The terminal silicon atom bearing a hydroxy group would be sterically and electronically more susceptible to a nucleophilic attack than the silicon atoms in the inner repeating units, since a hydroxy group is less bulky and a stronger electron withdrawing group than a dialkylsiloxyl group.65 The frequency of the side reactions increased in the following order: (e) < (d) < (c).

Although only the intramolecular condensation of a propagating polysiloxane can be considered as a possible termination reaction, we did not observe this process in any polymerizations that we conducted. Hence, the polymerization is characterized by the absence of termination reactions, which ensures the controlled/living nature of the polymerization. The total numbers of reactive hydroxy groups in water, which has ‘two’ hydroxyl groups, and propagating polysiloxanes do not change throughout the polymerization even when side reactions (c), (d), and (e) occur. The polymerization can be terminated only by neutralization of the catalyst or end-capping of propagating polymers.

Another possible polymerization mechanism is the ‘nucleophilic monomer activation mechanism’ that is based on the activation of the monomer by the nucleophilic attack of the catalyst.19 However, if so, (1) catalytic activity of the strong organic bases would not be linear to their Bronsted basicity as shown in Fig. 1, since the nucleophilicity (‘silicophilicity’) of the bases and their basicity are independent as it was demonstrated in a condensation of a silanol and an organosilanes with a leaving group catalyzed by an organic base.66 Besides, (2) the first-order kinetic plot would show a convex curve when the interaction of a catalyst and a monomer is high.64

**Effects of reaction conditions on kinetics of the polymerization**

The plausibility of the proposed polymerization mechanism was confirmed by comparing a rate equation derived from the kinetics of the elementary reactions, eqn (2), with the experimentally obtained rate equation, eqn (1). The rate of propagation, \( r \), can be expressed by the product of the rate coefficient of propagation, \( k_p \) (L mol\(^{-1}\) h\(^{-1}\)), the concentration of hydroxy groups activated by catalyst, \([\text{P}^*] \) (mol L\(^{-1}\)), and the concentration of unreacted monomer, \([\text{M}] \) (mol L\(^{-1}\)), when both \( k_w \) and \( k_1 \) are comparable to or greater than \( k_p \).

\[
\frac{d[M]}{dt} = k_p[P^*][M]
\]  

(2)

[\(P^*\)] is expressed by the product of the concentration of hydroxy groups not being activated, \([\text{P}] \) (mol L\(^{-1}\)), the concentration of catalyst not interacting with hydroxy groups, \([\text{C}] \) (mol L\(^{-1}\)), and the equilibrium constant for the activation of silanol groups by the catalyst, \(K_s\) (L mol\(^{-1}\)), i.e., \([\text{P}^*] = K_s[\text{P}][\text{C}]\), since \(K_s\) is defined as \(K_s = [\text{P}^*][\text{P}][\text{C}]\). \([\text{P}^*]\) is hence expressed by eqn (3) using the initial concentration of hydroxy groups, \([\text{P}]_0\) (mol L\(^{-1}\)), the initial concentration of the catalyst, \([\text{C}]_0\) (mol L\(^{-1}\)), since \([\text{P}] = [\text{P}]_0 - [\text{P}^*]\) and \([\text{C}] = [\text{C}]_0 - [\text{P}^*]\).

\[
[\text{P}^*] = K_s([\text{P}]_0 - [\text{P}^*])([\text{C}]_0 - [\text{P}^*])
\]  

(3)

From the quadratic formula, \([\text{P}^*]\) is expressed by eqn (4) as the function of three constants, i.e., \([\text{P}]_0\), \([\text{C}]_0\), and \(K_s\) that varies depending on the catalyst, the temperature, and the solvent employed for a polymerization:

\[
0 = [\text{P}]_0[\text{C}]_0 - \left([\text{P}]_0 + [\text{C}]_0 + \frac{1}{K_s}\right)[\text{P}^*] + [\text{P}^*]^2
\]

\[
[\text{P}^*] = \frac{1}{2}\left([\text{P}]_0 + [\text{C}]_0 + \frac{1}{K_s} - \sqrt{\left([\text{P}]_0 + [\text{C}]_0 + \frac{1}{K_s}\right)^2 - 4[\text{P}]_0[\text{C}]_0}\right)
\]  

(4)

[\(P^*\)] is hence constant, regardless of monomer conversion, \(c\), and frequency of the side reactions (c), (d), and (e). The integration of eqn (2) gives eqn (5):

\[
\ln(1 - c) = k_pl[\text{P}^*]t
\]  

(5)

which corresponds well with eqn (1) when \(k_{p,app}\) is regarded as:

\[
k_{p,app} = k_pl[\text{P}^*]
\]  

(6)

Hence, the proposed mechanism for the polymerization shown in Scheme 2 were proven to be reasonable. The polymerization is considered to proceed through the initiator/chain-end activation mechanism.

Effects of reaction conditions on the kinetics of polymerization can be expected with eqn (4)–(6). Fig. 8a shows the expected dependence of \(k_{p,app}\) on \(K_s\) and \([\text{C}]_0/[\text{P}]_0\) under the conditions of \([\text{M}]_0 = 1.80\) mol L\(^{-1}\) and \([\text{M}]_0/[\text{P}]_0 = 5\). The calculated \(k_{p,app}\) was normalized by dividing it by the \(k_{p,app}\) on \([\text{M}]_0/[\text{P}]_0 = 1\). It was found that \(k_{p,app}\) is almost proportional to \([\text{C}]_0/[\text{P}]_0\) when \([\text{C}]_0/[\text{P}]_0 \leq 1\) regardless of \(K_s\), which corresponded to the results shown in Fig. 5. In contrast, the increase in \(k_{p,app}\) with increasing \([\text{C}]_0/[\text{P}]_0\) depends on \(K_s\) when \([\text{C}]_0/[\text{P}]_0 > 1\) and its rate of change decreases with increasing \(K_s\). Fig. 8b shows the expected dependence of \(k_{p,app}\) on \(K_s\) and \([\text{M}]_0/[\text{P}]_0\) under the conditions of \([\text{M}]_0 = 1.80\) mol L\(^{-1}\) and \([\text{M}]_0/[\text{C}]_0 = 100\). The calculated \(1/k_{p,app}\) was normalized by multiplying it by the \(k_{p,app}\) on \([\text{M}]_0/[\text{P}]_0 = 5\). It was found that \(k_{p,app}\) is almost inversely proportional to \([\text{M}]_0/[\text{P}]_0\) when \(K_s\) is small, while \(k_{p,app}\) only gradually decreases with increasing \([\text{M}]_0/[\text{P}]_0\) when \(K_s\) is large. Considering the values of \(k_{p,app}\) observed in the polymerizations
moisture content of the solvent is taken into account, since it has been considered 'common sense' that intensive dehydration and purification of reagents and solvents is essential for conventional anionic ROP initiated by lithium compounds. This new convenient method to synthesize well-defined polysiloxanes will enable the synthesis of organosilicon and organic/inorganic hybrid materials with diverse architectures and may lead to the development of new advanced materials with improved properties.

Conflicts of interest

There are no conflicts to declare.

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