New Condensation Polymer Precursors Containing Consecutive Silicon Atoms—Decaisopropoxycyclopentasilane and Dodecaethoxyneopentasilane—And Their Sol–Gel Polymerization

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Abstract: The sol–gel polymerization of alkoxysilanes is a convenient and widely used method for the synthesis of silicon polymers and silicon–organic composites. The development of new sol–gel precursors is very important for obtaining new types of sol–gel products. New condensation polymer precursors containing consecutive silicon atoms—decaisopropoxycyclopentasilane (CPS) and dodecaethoxyneopentasilane (NPS)—were synthesized for the preparation of polysilane–polysiloxane material. The CPS and NPS xerogels were prepared by the sol–gel polymerization of CPS and NPS under three reaction conditions (acidic, basic and neutral). The CPS and NPS xerogels were characterized using N₂ physisorption measurements (Brunauer–Emmett–Teller; BET and Brunauer-Joyner-Halenda; BJH), solid-state CP/MAS (cross-polarization/magic angle spinning) NMRs (nuclear magnetic resonances), TEM, and SEM. Their porosity and morphology were strongly affected by the structure of the precursors, and partial oxidative cleavage of Si–Si bonds occurred during the sol–gel process. The new condensation polymer precursors are expected to expand the choice of approaches for new polysilane–polysiloxane.

Keywords: sol–gel precursor; condensation polymer; alkoxycyclopentasilane; alkoxyneopentasilane; porosity; polysilane; polysiloxane; oxidation of Si–Si bond

1. Introduction

Sol–gel polymerization of multialkoxysilanes is a convenient method for the synthesis of silicon polymers and silicon–organic composites under mild conditions [1–9]. A variety of sol–gel silicon polymer precursors such as alkyl or aryl bridged multialkoxysilane ((OR)₃Si-R'-Si(OR)₃, R' = alkyl, and aryl groups) and simple alkoxysilanes (Si(OR)₄ and R'Si(OR)₃) have been investigated [10–13]. Control of the morphologies and textural properties (surface area, pore volume, and pore distribution) of the resulting sol–gel polymer has been achieved by the molecular design of the precursors.

Quite recently, our group found interesting reactivities of peralkoxy-, perchlorocyclopentasilane, and perchloroneopentasilane: (i) Si–Si bond breaking of a soluble polymer synthesized from decaethoxycyclopentasilane with methyltrimethoxysilane occurred even at one minute-baking conditions (200 °C) [14], while a polymer obtained from hexaethoxymethyldicyclohexasilane with methyltrimethoxysilane was thermally stable; (ii) adding a secondary amine into...
decachlorocyclopentasilane and dodecachloroneopentasilane induced Si–Si bond cleavages. The interesting cleavage reactivity of the consecutive silicon bonds in the oligosilanes and the lack of study on the sol–gel process of peralkoxyoligosilanes prompted us to study their sol–gel reaction. To the best of our knowledge, sol–gel reactions of oligosilanes have not been studied. Only commercially available simple alkoxydisilanes—\((\text{OEt})_3\text{SiSi(OEt)}_3\) and \((\text{OME})_3\text{SiSi(OMe)}_3\)—were used for the synthesis of sol–gel silicon oxide materials. We expected that the new core structures and Si–Si bond breaking may affect the morphology and textural property of the resulting sol–gel polymer.

In this report, we describe the synthesis of new condensation polymer precursors containing consecutive silicon atoms—decaisopropoxycyclopentasilane (CPS) and dodecaethoxyneopentasilane (NPS)—as well as their sol–gel reactions and structures of the resulting CPS and NPS xerogels, representing the first sol–gel study of peralkoxyoligosilane.

![Decaisopropoxycyclopentasilane (CPS) and Dodecaethoxyneopentasilane (NPS)](image)

2. Materials and Methods

2.1. Materials

In all of the reactions in which air-sensitive chemicals were used, the reagents and solvents were dried prior to use. Diethyl ether, triethylamine, and \(n\)-hexane were distilled from Na/\(\text{PH}_2\text{CO}\). EtOH and \(\text{PrOH}\) were distilled from calcium hydride. Other starting materials were purchased as reagent grade and were used without further purification. Glassware was flame-dried with nitrogen or argon flushing prior to use. All of the manipulations were performed using the standard Schlenk techniques in nitrogen or argon atmosphere and using a glove box (MBraun, Garching, Germany).

2.2. Instruments and Measurements

\(^1\text{H}, ^{13}\text{C}, \text{and } ^{29}\text{Si NMR spectra were recorded using a Bruker Avance II}\) BBO 400 MHz S1 spectrometer (Bruker, Billerica, MA, USA). The chemical shifts were referenced to internal \(\text{C}_6\text{D}_6\) or \(\text{CDCl}_3\), or external tetramethylsilane. All of the solid-state NMR experiments were conducted with NMR instruments (Varian \text{unity NOVA}, Varian, Palo Alto, CA, USA) using 5 and 2.5 mm double resonance MAS probe heads for \(^{29}\text{Si}\) and \(^1\text{H}\) at 14.1 T \((^1\text{H} \text{resonance frequency } 600 \text{ MHz, wide bore})\) at room temperature. Mass spectra were recorded using a low-resolution (Agilent Technologies GC/MS: 6890N, 5973N mass selective detector) EI (electron ionization) mass spectrometer and a high-resolution (JEOL JMS-600W Agilent 6890 Series, Agilent, Santa Clara, CA, USA) instrument. The morphology of the xerogels was monitored by scanning electron microscopy (SEM, Quanta 250FEG, FEI, Hillsboro, OR, USA). High-resolution transmission electron microscopy (HRTEM) was performed with a JEOL JEM2100F (200 kV) instrument (JEOL, Tokyo, Japan) using a carbon-coated 200-mesh copper grid. Nitrogen adsorption–desorption isotherms were obtained using a surface area analyzer (BELSORP-max and Micromeritics, ASAP 2010, BEL, Osaka, Japan). Thermogravimetric analysis (TGA) curves were recorded using a simultaneous thermal analyzer (STA, STA 8000, Perkin Elmer, Waltham, MA, USA) in \(\text{N}_2\) (50.0 mL/min) atmosphere with heating from 30.0 to 1000.0 °C (10.0 °C/min).

2.3. Synthesis of Decaisopropoxycyclopentasilane (CPS, \(\text{Si}_5(\text{O}^\text{Pr})_{10}\))

\(\text{PrOH}\) (5.8 mL, 0.076 mol) and \(\text{Et}_3\text{N}\) (10.6 mL, 0.076 mol) dissolved in \(\text{Et}_2\text{O}\) (150 mL) were slowly added to decachlorocyclopentasilane \([23,24]\) (3.1 g, 6.3 mmol) dissolved in \(\text{Et}_2\text{O}\) (350 mL) for 1 h at...
were aged for 2 weeks at 30 °C. After gelation, the monoliths were crushed in water (100 mL) and filtered. The solids were washed twice with distilled H2O (100 mL) and washed with n-hexane in argon atmosphere in a glove box, and volatiles were distilled using vacuum distillation. Solid CPS (4.0 g, 5.5 mmol) was obtained at 88% yield. Si5(OiPr)10 (CPS): 1H NMR (C6D6, 400 MHz): δ 4.58 (m, 10H, SiOCH(CH3)2), 1.40 (d, 60H, SiOCH(CH3)2). 13C NMR (C6D6, 100 MHz): δ 26.79 (s, SiOCH(CH3)2), 69.17 (s, SiOCH(CH3)2). 29Si NMR (C6D6, 79 MHz): δ −14.41 (s). HRMS (high resolution mass): C27H60Si5 687.3268 (M+·Pr calcld), 687.3267 (found).

2.4. Synthesis of Dodecaethoxyneopentasilane (NPS, Si5(OEt)12)

EtOH (3.2 mL, 0.055 mol) and Et3N (8.4 mL, 0.060 mol) dissolved in Et2O (150 mL) were slowly added to the dodecaethoxyneopentasilane [25] (3.1 g, 6.3 mmol) dissolved in Et2O (350 mL) for 1 h at −78 °C. After the mixture was stirred for 6 h at −78 °C and slowly warmed to room temperature, the solution was stirred for 12 h at room temperature. The Et3N•Cl− salt was removed by filtration and washed with n-hexane in argon atmosphere in a glove box, and volatiles were distilled using vacuum distillation. NPS (2.3 g, 3.4 mmol) was obtained at 82% yield as an oil. Si5(OEt)12 (NPS): 1H NMR (C6D6, 400 MHz): δ 3.87 (q, 24H, SiOCH3), 58.85 (s, SiOCH3). 13C NMR (C6D6, 100 MHz): δ 18.90 (s, SiOCH2(CH3)2), 1.16 (t, 36H, SiOCH2(CH3)2). 29Si NMR (C6D6, 79 MHz): δ −40.52 (s, Si(Si(OEt)3)4), −152.16 (s, Si(Si(OEt)3)4). HRMS: C24H61O12Si5 681.3009 (M+·Pr calcld), 681.3030 (found).

2.5. Sol–Gel Polymerization of CPS and NPS

The sol–gel polymerization of CPS and NPS were carried out at 0.4 M concentration in EtOH or iPrOH according to the substituent groups of the precursors, respectively. The monomers were dissolved in anhydrous alcohols in 25.0 mL vials. An aqueous catalyst (0.01 wt % HCl or NaOH per water) was added to the vial. The solutions were sealed and shaken at room temperature. Gelation was determined at a point where the solution did not flow as a liquid. After gelation, the monoliths were aged for 2 weeks at 30 °C prior to further processing. The gels were crushed in water (100 mL), and filtered and washed with water (2 × 100 mL) and alcohols (EtOH or iPrOH, 2 × 100 mL). After drying overnight in air, the xerogels were prepared by vacuum drying at 50 °C for 24 h.

2.6. Preparation of CPSH

CPS (1.7 g, 2.32 mmol) was diluted in anhydrous iPrOH (4.5 mL) to adjust to 0.4 M concentration in an oven-dried vial (25 mL). A prepared aqueous acid catalyst, namely, 0.01 wt % HCl dissolved in 10 eq of H2O (0.42 mL, 23 mmol), was added to the vial, and then the vial was sealed and shaken. Gelation occurred within 5 min. After aging for 2 weeks at 30 °C, the wet gel was crushed in distilled H2O (100 mL) and filtered. The solids were washed twice with distilled H2O (100 mL) and anhydrous iPrOH (100 mL). After drying for 12 h in air, a white solid CPSH xerogel (0.54 g) was obtained by vacuum drying for 24 h at 50 °C. CPSH: 1H CP/MAS NMR (600 MHz): δ 0.975, 1.138, 5.126. 13C CP/MAS NMR (150 MHz): δ −1.862, 23.385, 47.661, 65.787. 29Si CP/MAS NMR (120 MHz): δ −29.062 (D1), −70.335 (T1), −100.860 (Q). N2 physisorption surface area (Brunauer–Emmett–Teller; BET) [26,27]: 317 m²/g, N2 absorption mean pore diameter (Brunauer–Joyner–Halenda; BJH): 28, 50, 90 nm. Total weight loss (TGA, 50–1000 °C, N2 atmosphere): 19.3%.

2.7. Preparation of CPSOH

Similarly to the procedure for the preparation of CPSH, CPSOH was prepared using an aqueous base catalyst, namely, 0.01 wt % NaOH dissolved in 10 eq of H2O (0.42 mL, 23 mmol), instead of the acid catalyst. Gelation occurred within 3 min. After aging, washing and drying were carried out in the same manner as described above. A white solid CPSOH xerogel (0.51 g) was obtained. CPSOH: 1H CP/MAS NMR (600 MHz): δ 0.690, 0.975, 1.138, 5.329. 13C CP/MAS NMR (150 MHz): δ 22.738, 65.657. 29Si CP/MAS NMR (120 MHz): δ −26.175 (D), −69.782 (T), −101.229 (Q). N2 physisorption
surface area (BET): 408 m$^2$/g, N$_2$ absorption mean pore diameter (BJH): 24, 43, 76 nm. Total weight loss (TGA, 50–1000 °C, N$_2$ atmosphere): 17.8%.

2.8. Preparation of CPSN

Similarly to the procedure used for the preparation of CPHS, CPSN was prepared using 10 eq of distilled H$_2$O (0.42 mL, 23 mmol) without any acid or base catalyst. Gelation occurred within 10 min. After aging, washing and drying were carried out in the same manner as described above. A white solid CPSN xerogel (0.53 g) was obtained. CPSN: $^1$H CP/MAS NMR (600 MHz): $\delta$ 0.120, 0.527, 4.190. $^{13}$C CP/MAS NMR (150 MHz): $\delta$ 0.534, 24.615, 30.182, 65.978. $^{29}$Si CP/MAS NMR (120 MHz): $\delta$ −32.076 (D), −71.382 (T), −100.984 (Q). N$_2$ physisorption surface area (BET): 150 m$^2$/g, N$_2$ absorption mean pore diameter (BJH): 32, 50, 76 nm. Total weight loss (TGA, 50–1000 °C, N$_2$ atmosphere): 16.2%.

2.9. Preparation of NPSH

The NPS (1.8 g, 2.6 mmol) was diluted in anhydrous EtOH (4.3 mL) to adjust to 0.4 M concentration in an oven-dried vial (25 mL). A prepared aqueous acid catalyst, namely, 0.01 wt % HCl dissolved in 12 eq of H$_2$O (0.56 mL, 31 mmol), was added to the vial and then the vial was sealed and shaken. Gelation occurred within 4 days. After aging for 2 weeks at 30 °C, the wet gel was crushed in distilled H$_2$O (100 mL) and filtered. The solids were washed twice with distilled H$_2$O (100 mL) and anhydrous EtOH (100 mL). After drying for 12 h in air, a white solid NPSH xerogel (0.63 g) was obtained by vacuum drying for 24 h at 50 °C. NPSH: $^1$H CP/MAS NMR (600 MHz): $\delta$ 0.080, 0.487, 3.905. $^{13}$C CP/MAS NMR (150 MHz): $\delta$ 1.052, 17.688, 29.987, 59.117. $^{29}$Si CP/MAS NMR (120 MHz): $\delta$ −80.472 (T), −101.721, −109.951 (Q). Total weight loss (TGA, 50–1000 °C, N$_2$ atmosphere): 14.9%.

2.10. Preparation of NPSOH

Similarly to the procedure used for the preparation of NPSH, NPSOH was prepared using an aqueous base catalyst, namely, 0.01 wt % NaOH dissolved in 12 eq of H$_2$O (0.56 mL, 31 mmol), instead of the acid catalyst. Gelation occurred within 1 day. After aging, washing and drying were carried out in the same manner as described above. A white solid NPSOH xerogel (0.67 g) was obtained. NPSOH: $^1$H CP/MAS NMR (600 MHz): $\delta$ 0.202, 2.806, 6.306. $^{13}$C CP/MAS NMR (150 MHz): $\delta$ 17.365, 30.117, 59.894. $^{29}$Si CP/MAS NMR (120 MHz): $\delta$ −91.158 (T), −101.721, −110.688 (Q). N$_2$ physisorption surface area (BET): 7 m$^2$/g, N$_2$ absorption mean pore diameter (BJH): 66, 90, 119 nm. Total weight loss (TGA, 50–1000 °C, N$_2$ atmosphere): 27.4%.

2.11. Preparation of NPSN

Similarly to the procedure for the preparation of NPSH, NPSOH was prepared using 12 eq of distilled H$_2$O (0.56 mL, 31.0 mmol) without any acid or base catalyst. Gelation occurred within 2 days. After aging, washing and drying were carried out in the same manner as described above. A white solid NPSN xerogel (0.62 g) was obtained. NPSN: $^1$H CP/MAS NMR (600 MHz): $\delta$ 0.527, 3.986. $^{13}$C CP/MAS NMR (150 MHz): $\delta$ 17.000, 17.753, 30.052, 58.793. $^{29}$Si CP/MAS NMR (120 MHz): $\delta$ −80.717 (T), −101.844, −110.442 (Q). N$_2$ physisorption surface area (BET): 28 m$^2$/g, N$_2$ absorption mean pore diameter (BJH): 21, 43, 120 nm. Total weight loss (TGA, 50–1000 °C, N$_2$ atmosphere): 17.9%.

3. Results and Discussion

3.1. Synthesis of the New Condensation Polymer Precursors

To investigate the sol–gel reaction of the oligosilane with consecutive silicon bonds by the molecular designed precursors, cyclic and branched peralkoxyoligosilanes were synthesized. The decaisopropoxycyclopentasilane (CPS)—Si$_5$(OIPr)$_{10}$—and dodecaethoxyneopentasilane (NPS)—Si(Si(OEt)$_3$)$_4$—were synthesized by the alcoholysis of decachlorocyclopentasilane and dodecachloroneopentasilane, respectively (Scheme 1). The alcoholysis reactions were carried out
at −78 °C in Et₂O for 6 h, and then the reaction mixture was slowly warmed to room temperature. After 12 h of stirring at room temperature, the formed white salt (Et₃N⁺Cl⁻) was removed by filtration without contact with air and moisture. The CPS cyclic compound was obtained as a solid at 88% yield. Using a method similar to the method described above, branched precursor NPS was obtained as an oil at 82% yield. The two products were characterized by ¹H NMR (Figures S1 and S4), ¹³C NMR, ²⁹Si NMR (Figures S2 and S5), and HRMS (Figures S3 and S6). The ²⁹Si NMR spectrum of CPS displayed a resonance at −14.4 ppm. In the HRMS data, one ¹Pr group removed fragment of CPS was detected. Two resonances were observed at −40.5 and −152.2 ppm in the ²⁹Si NMR spectrum of NPS and were assigned to the four terminal Si atoms and the central Si atom, respectively.

Scheme 1. Synthesis of the new condensation polymer precursors—decaisopropoxycyclopentasilane (CPS) and dodecaethoxyneopentasilane (NPS).

3.2. Preparation of CPS and NPS Xerogels

To prepare polysilane–polysiloxane materials, sol–gel polymerizations of CPS and NPS (0.4 M) were carried out at room temperature in isopropanol or ethanol according to the substituent groups of the precursors. Hydrolysis and condensation reactions were carried out in neutral, acidic (0.01 wt % HCl per H₂O), and basic (0.01 wt % NaOH per H₂O) conditions. For the hydrolysis, water was used with the number of moles of the equivalent of alkoxy groups of each precursor (10 eq for CPS and 12 eq for NPS). The CPS solution became a viscous gel within a few minutes, while the gelation of NPS required a few days. After gelation, all of the gels were aged for 2 weeks at room temperature to obtain condensation that was as complete as possible prior to further processing. The wet gels were washed with water and the corresponding solvent to remove the catalyst. Xerogels were obtained as opaque white brittle solids by vacuum drying for 24 h at 50 °C. The xerogels were ground into fine powder for analyses (Scheme 2).

Scheme 2. Preparation of CPS and NPS xerogels.
3.3. Characterization of CPS and NPS Xerogels

3.3.1. Surface Area and Porosity

The porosity of the xerogels was determined by \( N_2 \) physiosorption measurements. The textural properties of xerogels are summarized in Table 1. The BET (Brunauer–Emmett–Teller) method was applied to determine the surface area of the xerogels. CPS xerogels showed a larger surface area than the NPS xerogels. CPSOH exhibited the largest surface area of 408 \( \text{m}^2 \cdot \text{g}^{-1} \) and a total pore volume (at \( P/P_0 = 0.99 \)) of 0.74 \( \text{cm}^3 \cdot \text{g}^{-1} \). CPH showed a quite large surface area of 317 \( \text{m}^2 \cdot \text{g}^{-1} \) and the largest total pore volume of 0.83 \( \text{cm}^3 \cdot \text{g}^{-1} \). By contrast, the NPS xerogels were either non-porous or exhibited negligible porosity. These results indicated that the designed structure of the precursors affected the surface area and the porosity of the xerogels. Considering that the core size of CPS is larger than that of NPS, it was assumed that the higher porosity of CPS may be due to the difference in the core size. Usually, bridged polysilsequioxanes, which are large-core silsesquioxanes, are highly porous and have high surface areas [10–12,28–32].

Table 1. Textural properties of the xerogels (BET and BJH).

| Xerogels | Area \( S \) (m\(^2\)/g) | Pore Volume (cm\(^3\)/g) | Mean Pore Diameter (nm) |
|---------|-------------------------|--------------------------|------------------------|
| CPSH    | 317                     | 0.83                     | 28, 50, 90             |
| CP SOH  | 408                     | 0.74                     | 24, 43, 76             |
| CPSN    | 150                     | 0.24                     | 32, 50, 76             |
| NPSH    | No porosity             | -                        | -                      |
| NPSOH   | 7                       | 0.02                     | 66, 90, 119            |
| NPSN    | 28                      | 0.06                     | 21, 43, 120            |

3.3.2. Solid-State CP/MAS NMR Analysis

The xerogels were characterized by solid-state \(^1\)H, \(^{13}\)C, and \(^{29}\)Si CP/MAS NMR spectroscopy. Shorthand notations have been commonly used in the organosilicon literature for the Si–O bonding configurations: The structures of \((\text{SiO})_1\text{Si}, (\text{SiO})_2\text{Si}, (\text{SiO})_3\text{Si},\) and \((\text{SiO})_4\text{Si}\) are referred to as M, D, T, and Q, as this relates to the number of O atoms bonded to a Si atom, respectively. \(^{29}\)Si CP/MAS NMR spectra of the CPS xerogels show the presence of D \((R_2\text{SiO}_2) (\delta \text{ from } -20 \text{ to } -40 \text{ ppm})\), T \((RSiO_3) (\delta \text{ from } -60 \text{ to } -80 \text{ ppm})\), and Q \((\text{SiO}_4)\) units \((\delta \text{ from } -90 \text{ to } -120 \text{ ppm}) [33–37]\). If there was no Si–Si bond cleavage of CPS during the sol–gel process, only the D unit should be observed. However, T and even Q units were observed in Figure 1. The spectra indicate that the oxidative cleavage of Si-Si bonds in cyclopentasilane partially occurred during the sol–gel reaction to give Si–O–Si bonds [38–44]. Generally, when materials with Si–Si bonds on the surface are immersed into water or exposed to dry air, the Si–Si bonds are readily oxidized, and chemically stabilized Si–O–Si bonds are formed. The surface Si–Si bonds of the xerogels may be oxidized since NPS and CPS were exposed to water and air for a long time. Similarly to the \(^{29}\)Si CP/MAS NMR spectrum of CPS xerogels, the Q unit is also found in the spectra of NPS xerogels. Without Si–Si bond breaking of NPS during the sol–gel process, only the T unit should be observed. The resonances at the T and Q units of NPS xerogels observed in Figure 2 imply the partial oxidation of the Si–Si bonds of neopentasilane.
Figure 1. $^{29}$Si CP/MAS NMR spectra of CPS xerogels. (a) CPSH, (b) CPSOH, (c) CPSN.
3.3.3. Microscopy

The morphology and pore texture of xerogels were investigated by scanning electron microscopy (SEM) and high-resolution transmission electron microscopy (HRTEM). For CPS xerogels, Figures 3–5 show disordered meso- and macroporous structures consisting of spherical nanoparticles with other shapes. The sizes of the spherical nanoparticles were measured as 10–20 nm (CPSH), 20–30 nm (CPSOH), and 30–90 nm (CPSN). The different sizes of the particles can be attributed to the different conditions of the sol–gel reactions. While CPSOH consisted of spherical nanoparticles only (Figure 4a,b), aggregated nanorods sized 400–500 nm in CPSH (Figure 3b) and a huge particle sized approximately 1.5 µm in CPSN (Figure 5b) were observed along with spherical nanoparticles. These noncrystalline
composites may have contributed to the reduction of the surface areas of CPSH and CPSN (Table 1). TEM images display the mesoporous textures of the CPS xerogels (Figures 3c,d–5c,d).

Figure 3. Morphology and pore texture of CPSH. SEM images (a,b) and TEM images (c,d).

Figure 4. Morphology and pore texture of CPSOH. SEM images (a,b) and TEM images (c,d).
The morphologies of NPS xerogels were observed to be significantly different from those of the CPS xerogels. NPSH and NPSN are flat plates connected to each other as stacked films, and NPSOH is an irregularly wrinkled plate. In the TEM images, the textures consist of highly crystalline composites with no pores (Figures 6–8).
4. Conclusions

The new condensation polymer precursors containing consecutive silicon atoms—CPS and NPS—were successfully synthesized at high yields. CPS and NPS xerogels were prepared by sol–gel polymerization under three reaction conditions (acidic, basic, and neutral) using the new precursors. The characterizations of CPS and NPS xerogels using BET, BJH, and microscopy showed that their porosity and morphology were strongly affected by the structure of precursors. The CPS xerogels exhibited relatively large surface areas and total pore volumes, while the NPS xerogels exhibited nonporosity or negligible porosity. The relatively high porosity of CPS xerogels was achieved without
any organic substituent, template, or porogen, and they showed 16~19% of weight loss in TGA analysis. Bridged polysilsesquioxanes exhibited higher porosity, however, the weight loss reached around 45% \[10,29\]. 29Si CP/MAS NMR study of the xerogels indicated that a partial oxidative cleavage of the Si–Si bonds in cyclopentasilane and neopentasilane occurred during the sol–gel process. The structural features—a hybrid of Si–Si and Si–O–Si bonds—might provide a new optical property since consecutive silicon bonds absorb UV-visible light \[14\], and porous silica shows photoluminescence. The mixing of different skeletal bonds—Si–O–Si in an insulator and Si–Si in a semiconductor—provided interesting thermal and electrical conductivity. We hope that the new condensation polymer precursors for polysilane–polysiloxane materials will be used in many applications such as core-shell particle, anode materials for Li-ion batteries and non-metal thermal conductors.

**Supplementary Materials:** The following are available online at http://www.mdpi.com/2073-4360/11/5/841/s1, Figure S1: \(^1\)H NMR spectrum of CPS, Figure S2: \(^{29}\)Si NMR spectrum of CPS, Figure S3: HRMS result of CPS, Figure S4: \(^1\)H NMR spectrum of NPS, Figure S5: \(^{29}\)Si NMR spectrum of NPS, Figure S6: HRMS result of NPS, Figure S7: N\(_2\) adsorption–desorption isotherms of the CPS xerogels. (a) CPSH, (b) CPSOH, (c) CPSN. Figure S8: Pore size distribution of the CPS xerogels. (a) CPSH, (b) CPSOH, (c) CPSN.

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