Completely discontinuous organic/inorganic hybrid nanocomposites by self-curing of nanobuilding blocks constructed from reactions of $[\text{HMe}_2\text{SiOSiO}_{1.5}]_8$ with vinylcyclohexene

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Abstract: The reaction of 4-vinyl-1-cyclohexene with $[\text{HMe}_2\text{SiOSiO}_{1.5}]_8$ provides tetra- and octa-2-cyclohexenylethylotcasilsesquioxanes. The tetrabifunctional $[\text{cyclohexenylethylMe}_2\text{SiOSiO}_{1.5}]_4-[\text{Me}_2\text{SiOSiO}_{1.5}]_4$ (average degree of functionalization) melts near 80 °C and can be cast and then cured (by thermal hydrosilylation) into transparent nanocomposite shapes. These materials, while not nanoporous, offer dielectric constants of 2.8–2.9 at 100 kHz to 3 MHz, and are air stable to temperatures ≥400 °C. The resulting materials appear (by X-ray diffraction) to be partially ordered after curing. The octafunctional material also melts at low temperatures (ca 120 °C) and can be copolymerized with hydridosiloxanes to give similar materials with lower thermal stability. The synthesis and characterization of the starting materials is described, as well as thermal curing studies and properties characterization of the resultant nanocomposites.

Keywords: self-curing; bifunctional; silsesquioxanes; oxidation resistance; transparent

INTRODUCTION

We and others have undertaken extensive studies designed to develop routes to organic/inorganic hybrid nanocomposites using nanobuilding blocks based on the cubic silsesquioxanes $[\text{HMe}_2\text{SiOSiO}_{1.5}]_8$, (OHS) and functionalized octaphenyloctasilsesquioxanes $[\text{RPhSiO}_{1.5}]_8$ (R = NH$_2$, Br, I, etc.), both of which are easily prepared from low-cost starting materials.$^{1–35}$ In these systems, the core consists of the smallest single crystal of silica (0.5 nm in diameter) that acts as a rigid, hard particle functionalized at each of eight vertices. The core also provides the heat capacity of silica introducing a robustness not normally found in organic compounds. The functionalization can be used to form organic links or tethers between vertices.

Each functional group occupies one octant in Cartesian space, hence these cage or cubic systems are perfect nanobuilding blocks for constructing nanocomposites with excellent control of uniformity from nanometer to centimeter length scales.$^{31–35}$

The types of tethers and the reaction chemistry used to connect vertices provide access to hybrid materials wherein the periodicity within the nanocomposite, the coupling of the organic phase to the inorganic phase and to itself can be very well defined and tailored.$^{31,32}$ If the rigid core is considered to be a constant, then systematic changes in the nanoarchitecture of the organic components can be used to develop structure–processing–property relationships that allow correlations to be drawn between nanostructure and the macroscopic or global properties of these materials.

Because these nanocomposites closely resemble the ‘interphase’ materials that form at interfaces in macroscopic organic/inorganic composites,$^{36–39}$ they may offer information on property transfer (mechanical, photonic, electronic) across these interfaces, a process

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thought to be critical to developing optimal composite materials. Because they resemble interphase rather than bulk phase materials, they also offer the opportunity to create materials that have properties nonlinearly related to bulk phase properties. The potential exists to create whole new classes of materials.

In the majority of our studies to date, we sought crosslinking chemistry that leads to hybrid nanocomposites wherein single organic tethers connect two cube vertices only. That is, the organic phase interacts with the inorganic phase and vice versa with no chemical bonds forming between tethers. These completely discontinuous nanocomposites allow us to examine the influence of mechanical load transmission across interfaces at the nanoscale rather than through an organic matrix, which would result if the organic tethers were to copolymerize. Furthermore, the tethers used in our studies are typically 1–3 nm in length, such that in most instances, the organic component can be thought to represent a single segment in a wholly organic polymer. This has implications for understanding the behavior of single segments decoupled from polymer chains.

We have demonstrated the success of this approach for sets of epoxy resin nanocomposites. We and others have described extensions to polyimide nanocomposites. However, we return here to our oldest systems wherein simple hydrocarbon units couple cube vertices. These early systems provide the basis for extending the above concepts to systems with more flexible hydrocarbon units.

In our original studies, we systematically characterized products from hydrosilylative crosslinking of [HSiO1.5]8 and [HMe2SiOSiO1.5]8 with [CH2=CHMe2SiOSiO1.5]8 and [CH2=CHSiO1.5]8 giving simple CH2CH2 tethers. The resulting porous materials offered surface areas of 400–600 m2 g−1 and pore volumes typically <0.4 mL g−1. The shortest tethers, formed between [HSiO1.5]8 and [CH2=CHSiO1.5]8, gave the highest surface areas but with crosslink densities ca 45% of theory. Furthermore, the pore size distributions were very broad. In contrast, the longest tethers, obtained from reactions of [HMe2SiOSiO1.5]8 with [CH2=CHMe2SiOSiO1.5]8, gave the lowest surface areas but with crosslink densities 80% of theory and with a narrow pore size distribution of 0.1–0.4 nm. This pore size distribution could be accounted for if each cube was covalently bonded to six neighboring cubes (80% crosslink density) creating intercube pores of 0.1–0.2 nm in diameter and if some cubes were occasionally missing from the periodic structure (defects), accounting for the 0.3–0.4 nm pores. Furthermore, X-ray powder patterns of these materials indicated periodic assemblies as also suggested by the pore size distribution and crosslink densities.

Although the original materials offered potential as thermally stable (to 400 °C) low-k dielectrics and also as porous media for chromatographic applications, for example, they could only be prepared by solution processing that required a catalyst and solvent removal. Consequently, we sought other systems that might be melt processable and thermally crosslinkable but with many of the same advantages. To this end, we explored the use of 4-vinyl-1-cyclohexene as a tether. The vinyl group permits hydrosilylative attachment to the cube, while the cyclohexenyl unit is much less susceptible to hydrosilylation under the same conditions but will thermally hydrosilylate at higher temperatures, offering the potential for melt processing. The results of these efforts are presented below.

EXPERIMENTAL

Analytical methods

Solution NMR

NMR analyses were run in CDCl3 with 1H and 13C data recorded using a Varian INOVA 400 MHz spectrometer. 1H NMR spectra obtained at 400 MHz were collected using a 6000 Hz spectral width, a relaxation delay of 3.5 s, a pulse width of 38° and 30,000 data points. 13C(1H) NMR spectra were obtained at 100.6 MHz using a 25,000 Hz spectral width, a relaxation delay of 1.5 s, a pulse width of 40° and 75,000 data points. CDCl3 was used as internal reference for 1H NMR (7.259 ppm) and 13C NMR (77.23 ppm). 29Si NMR spectra were recorded using a Bruker AM 360 MHz spectrometer operating at 71.5 MHz using 32,000 Hz spectral width, a pulse width of 90°, a relaxation delay of 10.0 s, 32,000 data points and tetramethylsilane as external reference.

Thermal analysis

Dynamic thermal analysis–thermogravimetric analysis (DTA-TGA) of samples was performed using a SDT 2960 simultaneous DTA-TGA instrument (TA Instruments Inc., New Castle, DE). The instrument was calibrated with gold. Measurements were performed using 7–10 mg samples placed in alumina crucibles and heated under a continuous flow of nitrogen or synthetic air (110 mL min−1). The calorimeter was calibrated with indium as a reference. DSC traces were recorded using a 2910 differential scanning calorimeter (TA Instruments Inc., New Castle, DE) under nitrogen (200 mL min−1). The calorimeter was calibrated using indium as a reference. Samples (20–30 mg) were typically heated to 200 °C at 5 °C min−1 and then allowed to cool naturally at 1 °C min−1.

Fourier transform infrared (FTIR) analysis

Spectra were recorded on a Galaxy series FTIR 3000 spectrometer (Mattson Instruments, Madison, WI). Random cuttings of crystalline, optical-grade KBr from International Crystal Laboratories were used to prepare the sample. About 450 mg of KBr were ground in a mortar with a pestle with 10 mg sample

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added to make a 2 wt% mixture. The sample chamber was purged with nitrogen for a minimum of 5 min before data collection. A minimum of 128 scans was collected for each sample at a resolution of 4 cm⁻¹. Peaks were identified manually by selecting the desired region with the cursor.

**Mechanical properties studies**

**Sample preparation**

An aluminium mold (50 × 15 × 5 mm) was made from aluminium foil. Finely ground [cyclohexenylmethylMe₂SiOSiO₁.₅]₄-[HME₂SiOSiO₁.₅]₄ (TCHS) (ca 3 g) was then melted in the mold at 85°C in a vacuum oven (Lindberg Ltd, Blue M, Asheville, NC). The oven was then purged a minimum of three times with nitrogen and the sample was then heated gradually (10°C h⁻¹) to 150°C for 3 days under static vacuum (150 mmHg). The resulting specimen (30 × 10 × 2 mm) was polished with an Econet III polisher/grinder (Buehler Ltd, Lake Bluff, IL) for TMA measurements.

**Dynamic mechanical analysis (DMA)**

DMA was conducted using a 2980 dynamic mechanical analyzer (TA Instrument, New Castle, DE). The sample was mounted on a single cantilever clamp. Mechanical properties were measured under nitrogen in step mode every 10°C from room temperature to 300°C. Prior to each measurement, the furnace was kept at the set temperature for 10 min to ensure thermal equilibration.

**X-ray powder diffraction (XRD) studies**

XRD patterns were obtained from 2θ = 2° to 30° using a Rigaku X-ray diffractometer with Cu Kα radiation (1.542 Å) operating at 40 kV and 100 mA, at a scanning speed of 2° min⁻¹. The powder samples were prepared by milling DMA specimens. The samples were press-packed on a glass specimen holder.

**Dielectric constant measurements**

Dielectric constant measurements were obtained using a Hewlett Packard 419A instrument with sweep mode from 100 Hz to 40 MHz. A specimen was polished to 1.5 mm diameter and with a flat surface. The height of the specimen was measured directly with a caliper. The specimen was set between two copper electrodes to measure the dielectric constant at selected frequencies.

**Synthetic methods**

**Solvents and reagents**

4-Vinyl-1-cyclohexene, dimethylchlorosilane and tetramethylammonium hydroxide (25 wt% in methanol) were purchased from Sigma-Aldrich Co. Hexane, methanol and sodium sulfate were purchased from Fisher-Scientific. Tetraethoxysilane was purchased from Lancaster. Tetramethylsiloxane (TMDS) was purchased from Gelest and polydimethylsiloxane (PDMS) was purchased from Petrarch Systems. All of the above reagents and solvents were used as received. Toluene (Fisher-Scientific) used for hydrosilylation reactions was distilled from sodium/benzophenone under nitrogen.

**Catalyst**

Platinum dicyclopentadiene [Pt(dcpc)] was prepared from hydrogen hexachloroplatinate(IV) hydrate (Sigma-Aldrich) and dicyclopentadiene using published procedures. Sufficient sample (8 mg) was dissolved in distilled toluene (10 mL) to yield a 2.0 mmol L⁻¹ solution, which was stored under nitrogen.

**Octaamion solution**

To a 2 L round-bottomed flask equipped with a magnetic stir bar, 500 mL (4.75 mol) tetramethylammonium hydroxide (25 wt% in methanol), 258 mL (6.55 mol) methanol and 182 mL (10.1 mol) distilled water was added. The flask was cooled in an ice bath and the system was maintained under nitrogen. Then 253 mL (1.13 mol) tetraethoxysilane was added via an addition funnel. The solution turned cloudy and was stirred at room temperature overnight to produce a clear solution of tetramethylammonium octaamion in quantitative yield.

**Octakis(dimethylsiloxy)octasilsesquioxane¹⁻⁶**

To a 5 L round-bottomed flask equipped with a mechanical stirrer were added 2600 mL hexane and 436 mL (4.0 mol) dimethylchlorosilane under nitrogen. The flask was placed in an ice bath, and 1000 mL of octaamion solution ([Si] = 1 mol L⁻¹) was then added via an addition funnel over 2 h. The final mixture was stirred continuously for another 2 h. The two layers were separated with a 2 L separating funnel. The hexane layer was dried with sodium sulfate, gravity filtered through coarse filter paper and the solvent was removed by rotary evaporation to yield a white powder. This powder was rinsed with methanol and collected by vacuum filtration. The product was dried in a vacuum oven at 80°C for 4 h to yield 91.5 g (72% of theoretical) of octakis(dimethylsiloxy)octasilsesquioxane (OHS; octahydrido spacer cube). The product was further purified by subliming at 200°C under vacuum or recrystallization from ether at room temperature.


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0.380, −2.897 (d, H − Si − CH₃), −108.697 (s, Si − O). Mass spectrum (m/z, fragmentation, % intensity): 1016 (M⁺, 36.5), 1003 (M⁺ − CH₃, 100), 957 (M⁺ − Si(CH₃)H), 941 (M⁺ − OSi(CH₃)₂H). TGA (air): ceramic yield = 88.7% actual, 94.3% calculated. TGA (nitrogen): ceramic yield = 0% actual as sample sublimes.

**Tetakis-[2-(3’-cyclohexenyl)ethylidimethylsiloxyl]octaasilsequioxane (TCHS)**

To a 250 mL Schlenk flask equipped with a magnetic stir bar and a condenser was added 21.38 g (0.021 mol) OHS. The reaction vessel was dried with a heat gun under nitrogen. Then 4-vinyl-1-cyclohexene (100 mL) was added by syringe to dissolve the tetakis[2-(3’-cyclohexeny)ethylidimethylsiloxyl]octaasilsequioxane (TCHS) and the reaction was heated at 90 °C for 5 h. Triphenylphosphine (5 mg) was added to deactivate the Pt(dcp) catalyst, and the toluene solvent was removed by rotary evaporation. The remaining solid was rinsed with methanol and collected by vacuum filtration to give a quantitative yield of a white powder.

**Table 1. NMR data for vinylcyclohexene, OHS, TCHS, OCHS**

<table>
<thead>
<tr>
<th>Compound</th>
<th>¹H NMR (ppm)</th>
<th>¹³C NMR (ppm)</th>
<th>²⁹Si NMR (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4-Vinylcyclohexene</td>
<td>5.49 (m, H-5); 5.68 (d, H-5, H-6); 4.92 − 5.05 (m, H-1); 2.26 − 1.39 (m, H-3, H-4, H-7, H-8)</td>
<td>24.7 (C-5); 28.2 (C-6); 30.9 (C-3); 37.4 (C-4); 112.2 (C-8); 125.9 (C-2); 126.7 (C-1); 143.6 (C-7)</td>
<td>2910 (ν CH₃) 1652 (ν C–Si) 1455 (C–H bend) 2137 (ν C–H)</td>
</tr>
<tr>
<td>TCHS</td>
<td>5.65 (s, H-C=C-H); 4.73 (m, Si-H); 2.04 (ddd, H-4); 2.02 (q, H-7); 1.63 − 1.55 (m, H-8); 1.30 (m, H-2); 1.18 (m, H-3); 0.61 (t, H-1); 0.24 (d, H-4)</td>
<td>126.8, 127.0 (C-2), 26.4, 31.6, 29.6, 25.5, 25.4 (sp² C, cyclohexene); 14.6 (Si-CH₃); 0.08 (CH₃-Si-H); −0.46 (CH₃-Si-CH₃)</td>
<td>13.78 (CH₃-Si-CH₃); −0.081, −2.86 (CH₃-Si-H); 108.6 (Si-CH₃); −108.94 (CH₂-Si-O-Si)</td>
</tr>
<tr>
<td>OCHS</td>
<td>5.64, 5.65 (d, H-C=CH₂); 2.08 (ddd, H-4); 2.03 (q, H-7); 1.66, 1.56 (m, H-8); 1.25 − 1.29 (m, H-2); 1.18 (M, H-3)</td>
<td>127.2, 126.9 (C=C); 36.4, 31.9, 29.9, 28.7, 25.6 (sp² C, cyclohexene); 14.86 (Si-CH₃); −0.173 (CH₃-Si-CH₃)</td>
<td>13.49 (CH₃-Si-CH₂); −108.86 (CH₂-Si-O-Si)</td>
</tr>
</tbody>
</table>

**Curing studies**

**Thermal curing**

Typical curing studies were done in circular aluminium pans (44 × 12.5 mm) containing ca 2.0 g of sample. Samples were heated in a vacuum oven (model V0914SA, Lindberg/Blue M, Asheville, NC) at 100 °C under dynamic vacuum to melt the sample and volatilize residual solvent. The vacuum oven was then purged a minimum of three times with nitrogen, heated to 230 °C over 1 h and samples were cured at temperature for a minimum of 5 h under a static vacuum to produce clear solids.

**Radical curing**

In a mortar and pestle, 1 g of TCHS was mixed with 5 mol% 2-azobisisobutyronitrile (AIBN; 0.006 g) and then heated at 100 °C in an aluminium pan under dynamic vacuum to remove any excess solvent. The reaction was then heated at 200 °C for 6 h under nitrogen and static vacuum to produce a transparent, rigid cured polymer.
The resulting product was an opaque, somewhat flexible solid. TGA (air): ceramic yield = 30% actual, 52.3% theoretical. TGA (nitrogen): ceramic yield = 49.7%. FTIR data are listed in Table 3.

Table 3. FTIR data for cured resins

<table>
<thead>
<tr>
<th>Resin Type</th>
<th>OCHS w/OHS (cm⁻¹)</th>
<th>OCHS w/PDMS (cm⁻¹)</th>
<th>OCHS w/TMDS (cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TCHS self-cure</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2916 (ν–C–H alkane)</td>
<td>3023 (ν–C–H alkene)</td>
<td>3023 (ν–C–H alkene)</td>
<td>3023 (ν–C–H alkene)</td>
</tr>
<tr>
<td>2151 (ν–Si–H)</td>
<td>2144 (ν–Si–H)</td>
<td>1652 (ν–C=O)</td>
<td>1652 (ν–C=O)</td>
</tr>
<tr>
<td>1653 (ν–C=O)</td>
<td>1655 (ν–C=O)</td>
<td>1445 (ν–C–H bend)</td>
<td>1445 (ν–C–H bend)</td>
</tr>
<tr>
<td>1259 (ν–Si–CH₃)</td>
<td>1258 (ν–Si–CH₃)</td>
<td>1258 (ν–Si–CH₃)</td>
<td>1258 (ν–Si–CH₃)</td>
</tr>
<tr>
<td>1095 (ν–Si–O)</td>
<td>1096 (ν–Si–O)</td>
<td>1096 (ν–Si–O)</td>
<td>1096 (ν–Si–O)</td>
</tr>
</tbody>
</table>

Reaction of OCHS with OHS

Thermally promoted curing

Finely ground (mortar and pestle) mixtures of 0.73 g (0.0004 mol) OCHS and 0.37 g (0.0004 mol) OHS, both with and without 0.1 mL Pt(dcp) catalyst, were placed in aluminium pans in a vacuum oven and heated to 100 °C under dynamic vacuum to remove any residual solvent. The vacuum oven was purged at 20 °C for 1 h and then ramped at ca 8 °C min⁻¹ to 230 °C, and kept there for a minimum of 5 h under static vacuum (150 mmHg). The mixture never became molten. After 5 h, a brown, brittle solid was produced for both the catalyzed and uncatalyzed experiments. Improved curing was obtained if the triphenylphosphine from the synthesis was carefully removed by column chromatography.

AIBN-promoted curing

OCHS (0.73 g, 0.0004 mol) and 0.37 g (0.0004 mol) OHS were mixed with 5 mol% AIBN (0.0015 g) and ground with a mortar and pestle. The mixture was heated at 110 °C under dynamic vacuum to remove any residual solvent. It was then ramped in nitrogen-purged vacuum oven at a rate of ca 8 °C min⁻¹ to 200 °C for 6 h under static vacuum. The resulting product was an opaque, somewhat flexible solid. TGA (air): ceramic yield = 61.3% actual, 66.2% theoretical. TGA (nitrogen): ceramic yield = 59.9%. FTIR data are listed in Table 3. (See Appendix A for details.)

Reaction of OCHS with PDMS (see Appendix A for details)

In an aluminium pan, 1 g (0.50 mmol) of OCHS was heated at 120 °C under dynamic vacuum to remove excess solvent, and melt the sample. Then, 0.26 mL (0.002 mol) of TMDS and 0.1 mL of Pt(dcp) (2.0 mmol L⁻¹ solution in toluene) was added and ramped in a nitrogen-purged oven under static vacuum at ca 8 °C min⁻¹ to 200 °C for 5 h. An opaque, brittle solid was obtained.

The same reaction was performed by adding 5 mol% AIBN (0.004 g), which was ground together with OCHS with a mortar and pestle. This mixture was heated under nitrogen and static vacuum at 200 °C for 6 h. It produced an opaque, brittle solid. TGA (air): ceramic yield = 41.4% actual, 45.4% theoretical. TGA (nitrogen): ceramic yield = 30.0%. FTIR data are listed in Table 3. (See Appendix A for details.)

RESULTS AND DISCUSSION

In the following sections, we first describe the synthesis and characterization of ethylcyclohexenyl-substituted cubic silsesquioxanes. We then discuss their curing behavior under a variety of conditions followed by an evaluation of their thermomechanical and dielectric properties. These studies lead to some conclusions about their nanocomposite structures and properties.

TCHS/OCHS

TCHS is easily synthesized in >90% yield via platinum-catalyzed hydrosilylation, as per reaction (1).
The resulting solid (actually an average of multiple substitution products and their isomers) is a low-melting (76°C), white crystalline powder that was characterized using 1H NMR, 13C NMR, 29Si NMR (Table 1), FTIR (Table 2), DSC (Figs 1 and 2), XRD (Fig. 3) and DTA-TGA (Figs 4 and 5). Likewise, OCHS is easily synthesized using identical conditions but with excess vinylcyclohexene giving a low-melting (120°C), white crystalline solid.

NMR analysis

The 1H NMR spectrum of the starting cube, OHS, exhibits two peaks, a doublet at 0.252 ppm, corresponding to the methyl groups, and an Si–H peak that appears as a quintet centered at 4.727 ppm. 1H NMR integration of TCHS shows a 1.00:0.96 ratio for CH3–Si–H (0.25 ppm):(CH3)2Si–CH2 (0.13 ppm) providing an average substitution of 4.02. Earlier efforts to synthesize methacrylate derivatives via hydrosilylation when analyzed by mass spectroscopy showed mono-through octasubstitution with an average equivalent to the tetrasubstituted species. 46 We assume this pattern holds true here. OCHS was also characterized by 1H NMR, 13C NMR, and 29Si NMR (Table 1) and XRD (Fig. 3). Following hydrosilylation, only the internal double bond remains, as indicated by a peak at 5.65 ppm (Table 1), implying that reaction occurs solely at the vinyl position.

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These results are further supported by the 13C NMR analysis where 10 different carbon environments are expected and seen (Table 1). The 13C NMR spectrum of OHS shows only one peak at 0.059 ppm for (CH3)2HSi–O. Likewise, the 29Si NMR spectrum of OHS exhibits one peak at −108.7 ppm, corresponding to [HMe2SiO1.5]8, and a doublet centered at −1.26 ppm for [HMe2SiOSiO1.5]8. In contrast, the 29Si NMR spectrum of TCHS shows four peaks (Table 1). The peak at −108.9 ppm is assigned to O–Si–O–Si–CH2 units, while the peak at −108.7 ppm represents unreacted O–Si–O–Si–H units. A doublet centered at −1.39 ppm reveals coupling of (O–Si–O–Si–H) to the proton while the (O–Si–O–Si–CH2) unit gives a singlet at 13.78 ppm.

For OCHS, neither 1H NMR nor 13C NMR spectrum shows any Si–H or CH3–Si–H peaks, indicating that the cube is fully substituted (Table 1). The 13C NMR analysis shows only 9 carbon environments, as expected for the proposed structure. 47–49

FTIR analysis

The FTIR spectrum of TCHS shows an expected νC–Halkene band at 3023 cm−1, a νC=C band at 1652 cm−1 for the internal double bond and a νSi–H peak (2137 cm−1) for unreacted Si–H groups (Table 2). The OCHS spectra are similar, lacking only a νSi–H peak. Both systems exhibit νSi–O bands at 1096 cm−1, characteristic of the cubic core structure, and useful as an internal standard. More detailed FTIR studies on cubic silsesquioxanes can be found in papers by Calzaferri and c-workers. 47–49

DSC analysis

DSC cycling of TCHS to 300°C is shown in Fig. 1. In the first heating cycle, an endotherm due to melting (Tm) is observed beginning near 70°C. An exotherm is then observed with an onset close to 150°C and
a maximum at 220 °C. In the second cycle, neither peak is observed suggesting that thermally promoted hydrosilylation occurs, forming ethylcyclohexyl links (tethers) between cube vertices.

Fig. 2 shows DSC traces of TCHS heated to 150 °C. The intent is to examine its behavior below the curing temperature. In the first cycle, an endotherm appears with a maximum at 76 °C for Tm and on the cooling cycle a weak exotherm appears at 57 °C that may correspond to Tc. In the second cycle, the endotherm for melting broadens, the maximum shifts to 60 °C and the exotherm disappears.

These results can be explained if we assume that on precipitation from solution we produce a partially crystalline material. On heating, the crystalline form melts but if the melt has a limited propensity to crystallize on cooling, expected for a material that consists of a mixture of substituents, then an amorphous phase forms on cooling that can be expected to have a lower melting temperature.

**XRD studies**

Fig. 3 shows powder patterns for OHS, TCHS, OCHS and thermally cured TCHS (resin). The XRD data are best described by using the OHS powder pattern as a reference point. Three primary peaks seen for OHS are at 8.3°, 18.9° and 24.4° indicating a crystalline material. Indeed, the crystal structure is known. These peaks represent periodic distances of 1.1, 0.5 and 0.3 nm and are assigned to the (100) or (010), (120) and (112) planes, respectively. The 0.5 nm distance likely corresponds to the cube (core) body diagonal. The 0.3 nm peak likely corresponds to the separation between the 4–4′ rings of the cubes – the distance between cube faces.

The TCHS material is only partially crystalline as suggested by the broad peak at 7.3°, indicative of a larger unit cell (1.3–1.4 nm), as might be expected for the irregularly substituted cube. The peak at 18.9° shifts only slightly to 18.5° and does not exhibit any broadening indicating a direct relationship to the core structure, as suggested above. Also, the peak at 24.4° disappears. In the OCHS material, we see a similar pattern. The lowest peak shifts further to 6.5° indicating a slight but further increase in unit cell size. A similar shift to slightly lower angles gives a peak at 18.1°. Finally, the crosslinked resin produced by thermal promotion of hydrosilylation also exhibits some ordering. This is discussed further below.

**Thermal properties**

For reference purposes, the TGAs of OHS in nitrogen and air were run (not shown). OHS sublimes completely in nitrogen beginning at about 230 °C giving no ceramic yield. In air, it gives a ceramic yield of 85–90 wt% (94.3 wt% theory) depending on the ramp rates as some of the material sublimes before it oxidizes. Note that if the ramp rates are too fast in air, **OHS will oxidize explosively**.

TGA traces of TCHS in nitrogen and air are shown in Figs 4 and 5, respectively. In both systems, an exotherm is seen at ca 190 °C that corresponds to thermally promoted hydrosilylative crosslinking. In nitrogen, the thermally crosslinked material is stable to >400 °C and then degrades slowly with a final char yield of ca 70 wt%. In air (Fig. 5) the 190 °C exotherm is also accompanied by a slight mass gain as some Si–H oxidation likely occurs concurrent with thermally promoted hydrosilylation. A second exotherm is seen beginning around 350 °C associated with mass loss as the system oxidizes. The final ceramic yield is 66.1 wt% which is close to the calculated ceramic yield of 64 wt% for an average of four ethylcyclohexenyl substituents. These data are in good agreement with the 1H NMR integration.

Related TGA studies of OCHS in nitrogen and air are presented in Figs 6 and 7. The TGA run in nitrogen (to 1000 °C) gives a ceramic yield of...
36.4 wt% (versus 49% theory), with most mass loss between 450 and 550 °C. A small exotherm, seen at ca 200 °C, likely arises from thermally promoted hydrosilylation from residual Si–H groups. A small endotherm is seen centered at about 500 °C. The low ceramic yield and endotherm in the DTA suggest that some OCHS sublimes under nitrogen; however, the material remains thermally stable to about 400 °C. This suggests that it remains a liquid until it sublimes or decomposes. Perhaps, it may be useful as lubricant, especially in a fully hydrogenated form.

In air, the DTA/TGA of OCHS (Fig. 7) shows an exotherm at ca 190 °C with a mass gain due to oxidation of residual Si–H groups whose existence is suggested above. The ceramic yield in air is 46.8% (theory: 49.0%). This ceramic yield is 2% lower than expected and may be due to some competitive sublimation. The DTA trace shows irregular exotherms and endotherms between 400 and 450 °C that are likely a combination of the sublimation process and oxidative degradation processes.

The above DTA/TGA studies provide the basis for conducting bulk polymerization studies to produce larger scale samples for a variety of tests. The following section on curing discusses efforts to make fully crosslinked polymers from TCHS and OCHS.

Curing studies
The studies discussed in this section are divided into self-curing, radical-initiated curing and curing with chain extenders. We begin with the TCHS studies.

TCHS self-curing
Small samples (ca 2 g) of finely ground TCHS powder were cured at 200 °C for 6 h in aluminium pans (44 × 12.5 mm) under nitrogen in a sealed vacuum oven. The cured product gave somewhat flexible, transparent disks with some cracks. Attempts to form tensile specimens produced hard, clear solids at the exposed portions of the mold, but left the interior tacky and incompletely cured. Increasing the cure temperature and time to 230 °C for 8 h gave better
results, but interior sections remained tacky. If the triphenylphosphine used to deactivate the platinum catalyst is not added, the specimen obtained was harder and more brittle.

Fig. 8 shows spectra of TCHS and a self-cured sample. On curing, the intensities of the νC–H alkene (>3000 cm\(^{-1}\)), νSi–H (2151 cm\(^{-1}\)) and νC=\(\text{C}\) (1653 cm\(^{-1}\)) peaks decrease considerably relative to νC–H alkane, but do not completely disappear indicating that the cube is not 100% crosslinked following 8 h of curing. Thus, crosslinking appears to occur by thermally promoted hydrosilylation of the internal double bond. It is likely that 100% crosslinking is not possible due to steric and/or viscosity constraints. Also possible is that the distribution of the functional groups at the cube vertices is not completely regular as discussed below.

Figure 9 provides a schematic model of crosslinked TCHS. Based on this model and our previous studies\(^{27}\) one might anticipate nanoporosity in this system (see below).

The addition of 5 mol% AIBN to TCHS produced a material similar to the uncatalyzed experiment, yet the polymer was more flexible without cracks. FTIR analysis of the AIBN-cured product revealed a spectrum essentially identical to that of the self-cured material. Detailed studies on AIBN-promoted curing are given in Appendix A.

The TMA curve for the TCHS resin (Fig. 10) suggests a possible \(T_g\) near 70 °C. Although we also looked for \(T_g\) in the DMA trace, the samples were too brittle for study. Consequently, it is not clear if the change in slope at 70 °C in the TMA trace should be assigned to \(T_g\) or not. If it is a real \(T_g\) then it can only be assigned to movement of cyclohexyl rings, e.g. between chair and boat configurations, or to unreacted functional groups.

The coefficients of thermal expansion (CTEs) are 125 and 220 ppm °C\(^{-1}\) below and above 70 °C, respectively. As a reference, CTE values for typical silicone rubbers are 200–300 ppm °C\(^{-1}\),\(^{51}\) also in accord with some of the epoxy resin systems we have studied recently.\(^{35}\)

The XRD pattern of the TCHS resin magnified 5 times (Fig. 3) reveals broad weak peaks centered at 7.5° and 17.5° suggesting some periodicity within the resin structure. These peaks are relatively close to the peak positions in TCHS suggesting only modest changes in the cured structure. The fact that there is an XRD pattern at all may be attributed to the flexibility of the resulting ethylcyclohexyl tethers that may allow some ordering in the almost fully crosslinked system.

It is important to note that these systems should be true three-dimensional networks if the attachment of the ethylcyclohexenyl groups around the cage occurs in a random or statistical fashion. Indeed, random attachment seems necessary to obtain high crosslinked densities. The alternative perspective is that attachment is highly ordered, e.g. occurring only on one face of any cube. This would then lead to curing to form a ladder structure as suggested in Fig. 9. These are two extremes that for the moment we cannot differentiate between. But there is one mechanistic argument that can be made for attachment of all functional groups on one face.

The FTIR data in Fig. 8 suggest that only small amounts of Si–H remain. This is also supported by the small (1–2 wt%) mass gains in the TGA trace seen at near 200 °C in Fig. 7 suggesting that very little
Si–H is left to oxidize after curing. One interpretation may be that hydrosilylation is actually more selective than random. We have previously suggested that a platinum catalyst species may react selectively around one cube face simply on the basis of the higher statistical likelihood of reacting with another Si–H unit on that face.\(^{42}\) This would lead on crosslinking to the ladder structure shown in Fig. 9 which might explain the high CTEs observed relative to highly crosslinked cage epoxy resins we have made with CTEs well below 100 ppm °C\(^{-1}\). These issues remain unresolved at present.

Porosimetry studies were undertaken to determine if the resulting materials exhibited well-defined microporosity as in the case with the vinyl cube studies noted above. Surprisingly, after many repeated efforts by several individuals and the BET surface area measurements (Micromeretics ASAP 2000) manufacturer as well, it was determined that these materials were completely non-porous. At first this was quite surprising but these results are identical to those found by Loy and Shea, on studies of silsesquioxanes produced by hydrolysis of (EtO)\(_3\)Si(CH\(_2\))\(_3\)Si(OEt)\(_3\) (and related compounds) in both acid and base, where \(x = 2–8\). Only the materials produced with \(x = 2\) were porous.\(^{4}\)

**Dielectric constant measurements**

The model of Fig. 9 suggests the potential for well-defined porosity as found for materials with shorter tethers. Thus, before learning that these materials are not porous, we began measurements of their dielectric constants. Generally, the dielectric constants of these materials derive primarily from the cube structure.

Discontinuous organic/inorganic hybrid nanocomposites

**OCHS curing with OHS**

Attempts were made to react OCHS with equal molar amounts of OHS by dissolving it in molten OCHS with ca 1 g quantities. Aluminium pans with finely ground (mortar and pestle) OCHS and OHS were heated to 230 °C for 6 h under nitrogen in a sealed vacuum oven with or without Pt(dcp) catalyst. OCHS becomes molten, but OHS appears to be insoluble by visual inspection. Both experiments gave brown, brittle solids.

Samples of cured products, washed with hexane, were insoluble indicating that some crosslinking occurred since both OCHS and OHS are soluble in hexane. Crosslinking is not 100% complete because FTIR (Table 2) bands show \(\nu\)C–H\(_{\text{alkane}}\) (2924 cm\(^{-1}\)), \(\nu\)Si–H (2144 cm\(^{-1}\)), \(\nu\)C–H\(_{\text{alkene}}\) (3023 cm\(^{-1}\)), \(\nu\)C=H (1655 cm\(^{-1}\)) peaks in a 1:0.59:0.81:0.59 ratio, indicating that unreacted Si–H and C=C groups remain.

An improved material was made by mixing a 1:1 molar ratio of OCHS with OHS and 5 mol% AIBN. After curing at 200 °C for 6 h in nitrogen, a clear flexible disk was obtained. Some phase separation was seen in the form of opaque and transparent areas in the sample, most likely due to incomplete dissolution of OHS into the OCHS.

FTIR data (Table 3) reveal identical peaks for both the AIBN-initiated and uninitiated polymerizations. In comparison to FTIR ratios for the uninitiated reaction discussed above, the decreases in Si–H and C=C indicate a greater degree of crosslinking between OCHS and OHS when AIBN is present.

DTA/TGA traces of this polymer in air show a slight mass gain (ca 1.5%) at ca 210 °C (Fig. 11), indicating oxidation of residual Si–H groups. The TGA then shows a steep mass loss at ca 300 °C.

The DTA shows exotherms centered at 280 and 320 °C, corresponding to mass gains first from oxidation of residual Si–H and second from oxidative decomposition of the crosslinked material. The found ceramic yield of 61.3 wt% differs from theory (66.2%).

![Figure 11. TGA/DTA in air of OCHS:OHS 1:1; ramp rate = 10 °C min\(^{-1}\).](image-url)

Discontinuous organic/inorganic hybrid nanocomposites


DOF: 10.1002/pi
likely because of residual unincorporated OHS, which sublimes at ca 230 °C (see above). In nitrogen (Fig. 12), the polymer is stable to ca 400 °C with a ceramic yield of 59.9 wt%. This latter result supports the likely partial sublimation of unreacted material. No exotherms are seen in nitrogen as no oxidative processes are possible. This also indicates that all crosslinking processes from thermally promoted hydrosilylation are complete in the original curing steps.

Similar OCHS coupling studies were run using a hydrido-endcapped PDMS and also TMDS. These studies gave materials of similar low quality and properties and are recorded in Appendix A.

CONCLUSIONS

The use of bifunctional hydrocarbon reactants wherein the reactivity of one functional group is greater than the second allows the development of self-curing silsesquioxanes. The reaction of vinylcyclcohexene with OHS provides tetra- and octacyclohexenylethylcylocasilsesquioxane. The tetra-bifunctional (average degree of functionalization) melts to give low-viscosity liquids that can be cast and subsequently cured to highly crosslinked, transparent flexible nanocomposites. These nanocomposites offer thermal stabilities of ≥400 °C and good low-k dielectric properties without being nanoporous. The absence of UV-absorbing groups suggests that they may also offer promise for deep-UV photoresist applications.

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Figure 12. TGA/DTA in nitrogen of OCHS:OHS 1:1; ramp rate = 10 °C min⁻¹.
APPENDIX A

Detailed studies of AIBN curing

DTA/TGA of AIBN-initiated self-curing shows a decrease in air stability, from ca 400 °C for TCHS to ca 300 °C (Fig. A1). Large exotherms are seen in the DTA trace centered at 335 and 480 °C. Indeed, the exotherm at 335 °C is so great that the temperature recorded by the thermocouple is greater than the furnace temperature as witnessed by the slight drop in temperature below 80 wt%. The ceramic yield for the polymer is 65.5 wt%, in good agreement with the ceramic yield of TCHS monomer (65.7 wt%). DTA-TGA of the polymer run in nitrogen (Fig. A2) shows a gradual mass loss starting at ca 130 °C, which evolves into a much steeper mass loss at ca 400 °C to give a ceramic yield of 61.2%. This is slightly lower than expected and indicates some sublimation, suggesting incomplete curing, as expected from the tacky sample interior.

The DSC trace (not shown) does not show any peaks, which has important implications because it indicates that segmental motions that would account for $T_g$ in typical organic polymers are not possible in cured TCHS. Thus, when the ethylenedichlorohexyloxide (OC) cube derivative is reacted with diaminodiphenyl methane to form single tethers joining two cube vertices, $T_g$ values are seen at ca 150 °C. Clearly, the absence of $T_g$ in these studies indicates that segments of only 5 or 6 atoms are restrained from moving. Alternatively, they occur at below room temperature. These results contrast with our epoxy resin studies. 18–21

Figure A1. TGA/DTA in air of TCHS cured at 230 °C/8 h; ramp rate = 10 °C min⁻¹.

Figure A2. TGA/DTA in nitrogen of TCHS cured at 230 °C/8 h; ramp rate = 10 °C min⁻¹.

Reaction of OCHS with PDMS

OCHS was successfully cured with hydrido-endcapped PDMS (molecular weight ca 400) mixed in a 1:1.25 molar ratio (1.0 g:0.26 g) with 0.1 mL of Pt(dcp). This produced a clear, smooth and flexible solid after curing at 200 °C/5 h in a sealed, nitrogen-purged vacuum oven. The same experiment was run without the Pt(dcp) catalyst. After 200 °C/10 h the sample appeared tacky and incompletely cured.

Attempts to make tensile specimens of OCHS cured with PDMS and 0.2 mL Pt(dcp) using the same method as for self-cured TCHS were unsuccessful. After 200 °C/8 h the sample remained very tacky with many bubbles. These bubbles probably form from release of H₂ during curing.

FTIR analysis (Table 3) of the catalytically cured sample of OCHS and PDMS shows residual νC–Halkene (3023 cm⁻¹) and νC=C (1652 cm⁻¹)
peaks without a corresponding $\nu$Si–H peak. The $\nu$C–H alkane (2917 cm$^{-1}$); $\nu$C–H alkene (3023 cm$^{-1}$); $\nu$Si–H; $\nu$C= =C (1652 cm$^{-1}$) are in a 1:0.38:0.00:0.26 ratio. Vinyl group peaks are expected to remain because a molar ratio of 1:1.25 (OCHS:PDMS) was used. This means that there are 8 cyclohexenyl sites per 2.5 Si–H sites. Assuming that the Si–H groups of PDMS react fully, ca 6 cyclohexenyl sites should remain on OCHS. The decrease in vinyl group intensity along with the absence of $\nu$Si–H indicates that crosslinking proceeds via the same mechanism as for self-cured TCHS.

The same OCHS:PDMS molar ratio was also reacted with 5 mol% AIBN, curing at 200 °C/6 h. The resulting product was opaque and quite brittle. FTIR analysis of this sample showed the same peaks as for the platinum-catalyzed reaction (Table 3), indicating that crosslinking does occur, unfortunately resulting in undesirable properties. The ratio of $\nu$C–H alkane (2910 cm$^{-1}$); $\nu$C–H alkene (3023 cm$^{-1}$); $\nu$Si–H; $\nu$C= =C (1652 cm$^{-1}$) is 1:0.46:0:0.22.

DTA/TGA in air (Fig. A3) shows a slight mass loss at ca 100 °C, and then a much steeper mass loss at ca 300 °C. The ceramic yield is 52.3 wt% versus 59.2% theory, and the DTA trace shows exotherms centered at 373 and 525 °C. In nitrogen (Fig. A4), a slight mass loss is again seen beginning at ca 100 °C, and then a much steeper mass loss at ca 400 °C with a ceramic yield of 49.7%. The DTA trace does not show any exotherms. The above interpretations can also be made as to the thermal decomposition behavior of these materials. The overall poorer thermal stability of this material compared with the above materials suggests that the PDMS linker is the component that degrades first.

**Reaction of OCHS with TMDS**

Attempts were made to react OCHS with TMDS in a 1:4 molar ratio, catalyzed by Pt(dcp). After heating in a sealed vacuum oven at 200 °C/5 h, a transparent, brittle solid was produced.

**Figure A3.** TGA/DTA of OCHS:PDMS copolymer in air.

**Figure A4.** TGA/DTA of OCHS:PDMS copolymer in nitrogen.

**Figure A5.** TGA/DTA of OCHS:TMDS copolymer in air.

**Figure A6.** TGA/DTA of OCHS:TMDS copolymer in nitrogen.
FTIR analysis (Table 3) continues to show \( \nu_{\text{C--H alkene}} \) (3023 cm\(^{-1}\)) and \( \nu_{\text{C=C}} \) (1652 cm\(^{-1}\)) peaks, whereas the \( \nu_{\text{Si--H}} \) peak is absent. Theoretically, there is a 1:1 ratio of cyclohexenyl:Si--H groups. However, because TMDS is a volatile compound and the curing process involves heating in a vacuum oven, some TMDS is likely to evaporate, resulting in excess cyclohexenyl units, explaining the presence of \( \nu_{\text{C--H alkene}} \) in the FTIR spectrum after curing. The \( \nu_{\text{C--H alkane}} \) (2917 cm\(^{-1}\)):\( \nu_{\text{C--H alkene}} \) (3023 cm\(^{-1}\)):\( \nu_{\text{Si--H: C=C}} \) (1652 cm\(^{-1}\)) band ratio was 1:0.60:0:0.22.

The same reaction was run with 5 mol% AIBN, and cured at 200 °C/6 h. The product was an opaque, brittle solid. FTIR analysis shows peaks nearly identical to those in spectra from the platinum-catalyzed reaction, indicating that crosslinking also occurs in the presence of AIBN. The \( \nu_{\text{C--H alkane}} \) (2910 cm\(^{-1}\)):\( \nu_{\text{C--H alkene}} \) (3023 cm\(^{-1}\)):\( \nu_{\text{C=C}} \) (1652 cm\(^{-1}\)) band ratio is 1:0.51:0.27.

The TGA trace in air (Fig. A5) shows a mass loss of ca 12% from 240 to 350 °C likely due to dangling siloxane units that have not successfully crosslinked. A steep mass loss is then seen above 350 °C with a final ceramic yield of 41.4% versus 45.4% theory. Two exotherms are seen in the DTA trace run in air. The first has an onset at 430 °C, and a maximum at 460 °C over a range of ca 60 °C. The second has an onset at 520 °C, and a maximum at 550 °C over a range of 160 °C. Both are ascribed to decomposition of the copolymer.

The TGA run in nitrogen (Fig. A6) shows thermal stability up to ca 400 °C with a ceramic yield of 30.0%. No exotherms are seen in the DTA trace run in nitrogen as expected. The TGA curve seems to be very similar to that of the thermally cured TCHS cube suggesting that short PDMS segments do not affect the overall thermal stability.