Catalytic Synthesis of Polymethylsilsesquioxanes

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**Title**: Catalytic Synthesis of Polymethylsilsesquioxanes

**Author(s)**: Richard M. Laine, Jeffrey A. Rahn, Kay A. Youngdahl, John F. Harrod

**Abstract**: Dimethyltitanocene, Cp₂TiMe₂, where Cp = C₅H₅, can be used to promote the redistribution of -(MeHSiO)₉-cyclomers and linear oligomers to produce a polymethylsilsesquioxane copolymer of the type -(MeHSiO)₉(MeSi(O)₁.₅)₀.₇-. A mechanism for this redistribution is suggested, that involves σ-bond metathesis promoted by a Ti(IV) species. The structural evolution of the polymethylsilsesquioxane copolymer as it is heated to 1000°C is followed using solid state ²⁹Si NMR. TGA experiments and chemical analysis support the NMR results which indicate that most of the starting monomer is either volatilized or undergoes further redistribution, so that by 400°C only polymethylsilsesquioxane remains.

The active catalyst derived from Cp₂TiMe will also catalyze the alcoholysis of the Si-H bonds in the copolymer producing alkoxy derivatives, -(Me(R)O)SiO)₉(MeSi(O)₁.₅)₀.₇-(R = Me, Et, nPr, nBu). This same catalyst system can be used to polymerize polysilazane oligomers such as -(MeHSiNH)₋ or -(H₂SiNMe)₋, but only if some -(MeHSiO)₋ is present. The ceramic yields of 1:1, 1:3, 1:9, and 1:18 ratios of -(MeHSiO)₋:-(MeHSiNH)₋ were much higher than that found for the pure polysilazane.

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Catalytic Synthesis of Polymethylsilsesquioxanes

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Abstract:

Dimethyltitanocene, Cp₂TiMe₂, where Cp = C₅H₅, can be use to promote the
redistribution of -[MeHSiO]ₓ- cyclomers and linear oligomers to produce a
polymethylsilsesquioxane copolymer of the type -[MeHSiO]₀.₃[MeSi(O)₁.₅]₀.₇-. A mechanism for this redistribution is suggested, that involves α-bond
metathesis promoted by a Ti(IV) species. The structural evolution of the
polymethylsilsesquioxane copolymer as it is heated to 1000 °C is followed
using solid state ²⁹Si NMR. TGA experiments and chemical analysis support
the NMR results which indicate that most of the starting monomer is either
volatilized or undergoes further redistribution, so that by 400 °C only
polymethylsilsesquioxane remains.

The active catalyst derived from Cp₂TiMe₂ will also catalyze the
alcoholysis of the Si-H bonds in the copolymer producing alkoxy derivates,
-[Me(RO)SiO]₀.₃[MeSi(O)₁.₅]₀.₇- (R = Me, Et, nPr, nBu). This same catalyst
system can be used to polymerize polysilazane oligomers such as
-[MeHSiNH]ₓ- or [-H₂SiNMe]ₓ-, but only if some -[MeHSiO]ₓ- is present. The
ceramic yields of 1:1, 1:3, 1:9, and 1:18 ratios of -[MeHSiO]ₓ-:[MeHSiNH]ₓ-
were much higher than that found for the pure polysilazane. At higher ratios
of polysilazane, the apparent ceramic composition was the same as that
found for the pure polysilazane.
**Introduction**

Silsesquioxanes, $\text{RSi(O)}_{1.5}$, represent a unique and poorly studied subset of polyalkylsiloxanes. This is despite the fact that they offer many exceptional properties. For example, silsesquioxanes, because of their need to form three Si-O-Si bonds, assume regular polyhedral shapes such as shown for octamethyloctasilsesquioxane:\(^1\)

![Octamethyloctasilsesquioxane](image)

These polyhedral shapes have geometries very similar to those found for silica and its derivatives. Feher et al\(^2\) have used this similarity as the basis for developing molecular models of silica surfaces. The regular geometry also contributes to such properties as high temperature stability\(^1\) and high hardness.\(^3\) Octamethyloctasilsesquioxane is stable in air to 415°C at which temperature it sublimes.

Polysilisesquioxanes appear to have cage rather than ladder structures,\(^4\) wherein the polymer is formed by opening of a polyhedral edge as suggested for polymethylsilsesquioxane:

![Polymethylsilsesquioxane](image)

Polymethylsilsesquioxane, $-\text{[MeSi(O)}_{1.5}\text{]}_x-$, is stable in air to temperatures $>500^\circ\text{C}$ and, as we will show below, to at least 600°C in nitrogen. The phenyl derivative is reported to be stable to temperatures of
800°C. The cage structure has been suggested to be useful for making microporous materials.

Polymethylsilsesquioxanes have been used as protective polymer coatings in the electronics industry and as precursors to silica and SiO$_4$-$x$C$_x$ glasses:

From an engineering standpoint, these materials offer exceptional properties that should lead to widespread applications. Unfortunately, their propensity for forming gels makes it extremely difficult to prepare useful, processable quantities of any given material.

Until recently, the only method of preparing silsesquioxanes was via hydrolysis of the alkyltrichloro or trialkoxysilane:
Separation of the polyhedral or polyalkylsilsesquioxane from the reaction mixture is extremely difficult and the isolable yields for these compounds are quite poor, typically ranging from 15-30%. Consequently, the discovery by one of us that titanium will catalyze the redistribution of alkoxy silanes under extremely mild conditions, reaction (1),8 suggested that

$$3\text{MeHSi(OEt)}_2 < 0.1 \text{ mol } \% \text{ Cp}_2\text{TiMe}_2/\text{RT} > \text{MeSiH}_3 + 2\text{MeSi(OEt)}_3$$

(1)

it might be possible to synthesize polysilsesquioxanes via a similar route which eliminates the need for a hydrolytic synthesis:

$$-[\text{MeHSiO}]_n - 0.1 \text{ mol } \% \text{ Cp}_2\text{TiMe}_2/\text{RT} > \text{MeSiH}_3 + -[\text{MeSi(O)}_{1.5}]_x^-$$

(2)

This reaction works very effectively and we present here an overview of our recent efforts to prepare and characterize the resulting polymers and their properties. The work presented here includes work previously published elsewhere.9,10

Results and Discussion:

Neat mixtures of either cyclic -[MeHSiO]_n^- (n =4,5), or linear oligomeric, Me_3Si-[MeHSiO]_n-H (M_n = 2000 D) with 0.2 mol % (η^5-C_5H_5)_2TiMe_2 will turn royal blue (under N_2 at 20°C) following an induction period of ca 15 min. MeSiH_3 is evolved rapidly with stirring and in 5-7 min the solution becomes extremely viscous and gels in 10-15 min. Solid state $^{29}$Si NMR (see below) indicates that the final gel consists of a copolymer of approximate composition -[MeHSiO]_{0.3}[MeSi(O)]_{1.5}_{0.7}. The induction period that precedes reaction appears to be related to free radical promoted decomposition of (η^5-C_5H_5)_2TiMe_2 which generates the true catalyst.

Soluble product can be obtained by dilution with toluene. Thus, reaction in a five-fold excess (with respect to added -[MeHSiO]_n^-) of toluene gives a stable solution after 72 h of reaction. Reactions attempted with less than a five-fold excess of toluene lead inevitably to the formation of a gel.

Thin films cast from the resulting copolymer/toluene solution exhibit moderate elastomeric properties, excellent adhesion to glass, carbon, and metal surfaces and, can be heated without significant changes in properties to 250°C. Above this temperature, the polymer becomes more and more
brittle; however, no visible degradation occurs up to temperatures of approximately 400°C. However, solid state $^{29}$Si NMR can be used to demonstrate that structural changes do occur on heating.

Because of our continuing interest in polymer precursors to ceramics we followed the structural evolution of the Me$_3$Si-[MeHSiO]$_n$-H derived copolymer during heating to 1000°C using solid state $^{29}$Si NMR (Figure 1). At room temperature, we see the Me$_3$Si (3%) and Me(OH)Si (4%) end caps of the original oligomer together with peaks for -[MeHSiO]$_n$- (30%) and -[MeSi(O)$_{1.5}$]$_x$- (70%). TGA experiments and chemical analysis$^9$ support the NMR results which indicate that most of the starting monomer is either volatilized or undergoes further redistribution such that by 400°C only the pure polymethylsilsesquioxane remains. Note that in the NMR the sharp singlet of the [MeSi(O)$_{1.5}$] silicon is present at 600°C but becomes severely broadened as the polymer is transformed into a glass at 800°C.$^{10}$

Insert Figure 1

Nearly identical copolymer compositions are obtained from the copolymer generated in toluene as determined following solvent removal. The NMR established composition is confirmed by the chemical analysis.$^{11}$ Given the relatively simple $^{29}$Si NMR, the copolymer structure is assumed to be that shown below:

However, it is possible that other polyhedral silsesquioxane structural units also form in the copolymer.

Catalytic redistribution of hydridosiloxanes by transition metals was first reported by Curtis and Epstein.$^{12}$ They discovered that iridium complexes promoted redistribution of H-[Me$_2$SiO]$_x$-SiMe$_2$H and proposed the
following type of mechanism:

\[
\begin{align*}
\text{Ir} + \text{HSiMe}_2\text{OSiMe}_2\text{H} & \rightarrow \text{Ir} \quad \text{HSiMe}_2\text{OSiMe}_2\text{H} \\
-\text{Me}_2\text{SiH}_2 & \rightarrow \quad \text{HSiMe}_2\text{OSiMe}_2\text{H}
\end{align*}
\]

Based on recent work by Tilley et al\textsuperscript{13} on the reactions of d\textsuperscript{9} metals with silanes, we believe that a different mechanism is operating here. This mechanism probably involves \(\sigma\)-bond metathesis promoted by a Ti (IV) species generated by decomposition of \((\eta^5\text{-C}_5\text{H}_5)_2\text{TiMe}_2\). The following mechanism is suggested based on the work of Tilley et al; however, substantiation must await detailed kinetic studies.

We propose a Ti (IV) catalytic process despite the royal blue color of the reaction which is typical Ti (III) compounds and despite our having isolated Ti (III) complexes\textsuperscript{14} in related systems because of the following discovery.
Because the copolymer \([-\text{MeHSiO}]_0.3\text{[MeSi(O)1.5]}_0.7\)- still retains Si-H groups which are susceptible to further modification either by hydrosilylation or alcoholysis, we explored the possibility of changing the copolymer rheological characteristics by reaction with alcohols. We find that addition of alcohols to the toluene reaction solution anytime during the course of reaction leads to very effective alcoholysis of the copolymer:

\[
\text{MeMeI} > \text{I} + \text{MeMeMeSi(O)1.5} \text{[MeSi(O)1.5]}_0.7
\]

The addition of MeOH leads to a rapid color change from royal blue to yellow orange and extremely rapid (almost violent) evolution of hydrogen. The reaction is complete within the time of addition. The other alcohols are less reactive, with the n-BuOH reaction taking 1-2 days at room temperature.

Once solvent is removed, the methoxy derivative will become gel-like in hours to days depending on the temperature of the room. The n-butoxy derivative in contrast is much less susceptible to gelling and will remain as a viscous liquid for periods up to one week. N-propanol solutions of the n-propoxy derivative (25 wt %) will remain stable almost indefinitely. NMR characterization indicates the formation of \([-\text{Me(RO)SiO}]_0.3\text{[MeSi(O)1.5]}_0.7\)- and confirms the initial copolymer composition. These polymers are actually a masked form of \([-\text{MeSi(O)1.5}]_x\)- given that addition of water will lead to hydrolysis of the SiOR bond and water can actually be used to cause thermosetting!

The yellow-orange color is typical of a Ti (IV) catalyst and when coupled with Tilley’s work suggests the above proposed Ti (IV) promoted redistribution reaction.

Polysilazane Polymerizations

The success of this catalyst system suggested that it might also be used
for the catalytic polymerization of polysilazane oligomers such as 
\(-\text{[MeHSiNH]}_x\) or \(-\text{[H}_2\text{SiNMe]}_x\). This would offer an alternative to the 
ruthenium catalysts that we have previously used to form tractable silicon 
nitride preceramic polymers. To our surprise, neither oligomer underwent 
catalytic redistribution when mixed with \((\text{n}^5\text{-C}_5\text{H}_5\text{)}_2\text{TiMe}_2\) despite the fact 
that the catalyst reacted and some small amount of gas (presumably \text{CH}_4) 
evolved coincident with reaction.

This result was disappointing; however, we attempted to catalyze the 
redistribution of \(-\text{[MeHSiO]}_x\) in the presence of \(-\text{[MeHSiNH]}_x\) to explore the 
possibility of trapping the \(-\text{[MeHSiNH]}_x\) oligomer in the resulting 
silsesquioxane gel. We used ceramic yield as a measure of our success. 
Thus, Figure 2 shows TGAs for the copolymer derived from \(-\text{[MeHSiO]}_x\) 
(74-78% ceramic yield at 900°C in \text{N}_2), pure \(-\text{[MeHSiNH]}_x\) (37% ceramic yield 
at 900°C in \text{N}_2) and a 1:1 mixture of \(-\text{[MeHSiO]}_x/-\text{[MeHSiNH]}_x\). If the 1:1 
mixture were to act simply as a physical mixture, then the ceramic yields 
for the combination should be the arithmetic mean or 56%. Instead, we find 
that (see Figure 2) the ceramic yield is ≈ 72%. This would suggest that we 
were successful in trapping the silazane in the interstices of the 
polymethylsilsesquioxane gel. However, an alternate explanation is that in 
the presence of the hydridosiloxane, the catalyst is now capable of 
catalyzing redistribution of polysilazanes. To test this, we changed the 
siloxane:silazane ratio. Table 1 shows both that as the ratio is changed from 
1:1 to 1:3 to 1:18 reaction continues to occur, the ceramic yields stay high 
(relative to pure polysilazane) and the resulting ceramic products begin to 
look as like the ceramic products obtained from pure \(-\text{[MeHSiNH]}_x\). Our 
preliminary conclusion is that titanium catalyzed redistribution of 
\(-\text{[MeHSiNH]}_x\)-requires the presence of some quantity of hydridosiloxane as 
cocatalyst.\textsuperscript{15}

It is important to note here that the apparent compositions reported in 
Table 1 are a form of book-keeping that is not truly indicative of the actual 
nature of the \text{SO}_4\times\text{C}_x glass.\textsuperscript{9} However, these compositions are adequate for 
developing the selectivity to ceramic products, at 900°C, obtained by 
pyrolysis of \(-\text{[MeHSiNH]}_x\) and \(-\text{[H}_2\text{SiNMe]}_x\). The siloxane/silazane mixtures 
are perhaps better treated as mixtures of silicon oxynitride, \text{Si}_2\text{ON}_2 and 
\text{SO}_4\times\text{C}_x or \text{Si}_2\text{ON}_2 and silicon nitride/carbide depending on the percentage of 
initial hydridosiloxane.
<table>
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<th>Compound</th>
<th>Ceramic Yield</th>
<th>Apparent Ceramic Composition</th>
<th>Si₃N₄</th>
<th>SiC</th>
<th>C(x's)</th>
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<tr>
<td>MeHSiNH</td>
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<td></td>
<td>64</td>
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<tr>
<td>H₂SiNMe</td>
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<td></td>
<td>75</td>
<td>--</td>
<td>18</td>
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<tr>
<td>MeHSiO</td>
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<td></td>
<td>70</td>
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<td>10</td>
</tr>
<tr>
<td>1:1 MeHSiO/MeHSiNH</td>
<td>72</td>
<td></td>
<td>31</td>
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<td>10</td>
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<tr>
<td>1:3 MeHSiO/MeHSiNH</td>
<td>64</td>
<td></td>
<td>43</td>
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<td>1:9 MeHSiO/MeHSiNH</td>
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<td>1:18 MeHSiO/MeHSiNH</td>
<td>63</td>
<td></td>
<td>62</td>
<td>19</td>
<td>11</td>
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</table>

Table 1. Apparent Ceramic Compositions for Selected Polysilazanes, Polymethysilsesquioxane and Various Mixtures Pyrolyzed to 900°C in Nitrogen. Heating rate 5°C/min. Apparent ceramic compositions are calculated assuming Si is the limiting element. N is the limiting element when -[H₂SiNMe]ₓ⁻ is the preceramic.

Conclusions

Titanium catalyzed redistribution of -[MeHSiO]ₓ⁻ provides a useful route to tractable, processable methylhydridosiloxane-methylsilsesquioxane copolymers. The "Ti" catalyst active in the redistribution reaction will also promote alcoholysis of the resultant copolymers to produce alkoxy derivatives which display equivalent or slightly better, high temperature stability than the starting copolymer and more controllable rheology.

Although Cp₂TiMe₂ will not catalyze the polymerization of pure -[MeHSiNH]ₓ⁻, in the presence of small amounts of hydridosiloxane, it is an active catalyst precursor and permits catalytic redistribution that leads to a high ceramic yield silicon nitride precursor.
Acknowledgements

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References

3. From data sheet on GE Polyalkylsilsesquioxane (PALS) coatings distributed by General Electric Company.
11. Found C = 19.4%, H = 5.4%, Si = 41.5%, Calc. for [MeHSiO]0.3[MeSi(O)1.5]0.7, C = 18.5%, H = 5.1%, Si = 43.1%
Figure 1. Chemical evolution of Me₃Si-[MeHSiO]ₙ-H derived copolymer during heating to selected temperatures 1000 °C using solid state ²⁹Si NMR. M = Me₃SiO; D = Me₂Si(O)₂; DH = MeHSi(O)₂; DOH = MeSi(OH)(O)₂; T = MeSi(O)₃; and Q = Si(O)₄.

Figure 2. Thermogravimetric Analysis of -[MeSi(O)₁.₅]ₓ⁻ (a), -[MeHSiNH]ₓ⁻ (c), and a 1:1 Copolymer of -[MeHSiNH]ₓ⁻ and -[MeSi(O)₁.₅]ₓ⁻ (b). Pyrolyzed under N₂ at a heating rate of 5 °C/min.