Rheological Behavior of Entangled Polystyrene-Polyhedral Oligosilsesquioxane (POSS) Copolymer (Preprint)

Jian Wu & Patrick T. Mather (Univ. of CT); Timothy S. Haddad (ERC); Gyeong-Man Kim (Martin Luther University, Germany)

Air Force Research Laboratory (AFMC)
AFRL/PRSM
9 Antares Road
Edwards AFB CA 93524-7401

Air Force Research Laboratory (AFMC)
AFRL/PRS
5 Pollux Drive
Edwards AFB CA 93524-7048

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We report on the linear viscoelastic properties of a family of entangled linear thermoplastic non-polar hybrid inorganic-organic polymers: random copolymers of polystyrene (PS) and styryl-based polyhedral oligosilsesquioxane (POSS), R7(Si8O12)(C6H4CH=CH2), with R = isobutyl (iBu). A series of styrene-styryl POSS random copolymers with 0, 6, 15, 30, 50 wt% iBuPOSS were investigated. WAXS and TEM demonstrate that the iBuPOSS disperses in the polymeric matrix at a molecular level. It is observed that the iBuPOSS plays a plasticizer-like effect, yielding a monotonic decrease of the glass transition temperature with increasing iBuPOSS content. Rheological measurements revealed that linear viscoelastic behavior of the copolymers is also profoundly influenced by the presence of iBuPOSS. The incorporation of iBuPOSS dramatically decreases the rubbery plateau modulus (νNG), suggesting a strong dilation effect of isobutyl-POSS on entanglement density. Additionally, the apparent flow activation energy, obtained by fitting the Vogel-Fulcher-Tamman-Hesse equation, monotonically increases with increasing iBuPOSS content, indicating a lower sensitivity of POSS copolymers to changes of temperature. We attribute observations of the microscopic topology of constituent polymer chains to be altered by iBuPOSS comonomers that act as compact volumetric branches. Conversely, intermolecular interactions between iBuPOSS and PS segments do not play an essential role in determining the rheological behavior.

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Rheological Behavior of Entangled Polystyrene-Polyhedral Oligosilsesquioxane (POSS) Copolymers. Preprint

Jian Wu¹, Timothy S. Haddad², Gyeong-Man Kim³, Patrick T. Mather¹*

¹Polymer Program and Department of Chemical Engineering, Univ. Connecticut, Storrs, CT 06269
²ERC, Inc., AFRL/PRSM, Edwards Air Force Base, CA 93524
³Institut für Werkstoffwissenschaft, Martin-Luther-Universität Halle-Wittenberg, Geusaer Straße, D-06217 Merseburg

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Abstract: We report on the linear viscoelastic properties of a family of entangled linear thermoplastic non-polar hybrid inorganic-organic polymers: random copolymers of polystyrene (PS) and styryl-based polyhedral oligosilsesquioxane (POSS), R₇(Si₈O₁₂)(C₆H₄CH=CH₂), with R = isobutyl (iBu). A series of styrene-styryl POSS random copolymers with 0, 6, 15, 30, 50 wt% iBuPOSS were investigated. WAXS and TEM demonstrate that the iBuPOSS disperses in the polymeric matrix at a molecular level. It is observed that the iBuPOSS plays a plasticizer-like effect, yielding a monotonic decrease of the glass transition temperature with increasing iBuPOSS content. Rheological measurements revealed that linear viscoelastic behavior of the copolymers is also profoundly influenced by the presence of iBuPOSS. The incorporation of iBuPOSS dramatically decreases the rubbery plateau modulus (G''), suggesting a strong dilation effect of isobutyl-POSS on entanglement density. Additionally, the apparent flow activation energy, obtained by fitting the Vogel-Fulcher-Tamman-Hesse equation, monotonically increases with increasing iBuPOSS content, indicating a lower sensitivity of POSS copolymers to changes of temperature. We attribute observations of the microscopic topology of constituent polymer chains to be altered by iBuPOSS comonomers that act as compact volumetric branches. Conversely, intermolecular interactions between iBuPOSS and PS segments do not play an essential role in determining the rheological behavior.
I. Introduction

Polymers and polymer-matrix composites are used in numerous high-tech applications, but in many cases replacement of metallic components by polymers in order to lower weight is challenging due to stringent thermo-mechanical requirements. For example, needs exist for polymers and polymer-based composites to serve as cryogenic components of liquid rocket engines, or for polymeric films that survive the space environment. However, such applications demand high strength over a large temperature range and/or survival in harsh environments. With conventional high-performance polymeric materials, largely aromatic-based but with widely varying polymerization chemistry (epoxies, polyimides, etc.), limitations have been encountered and satisfaction of application requirements is only possible in limited cases. Thus, materials engineers have turned to alternative approaches that improve properties by reinforcing the constituent polymer chains at the local, or segmental, level. One particularly successful approach has been the use of hybrid inorganic-organic monomeric units, polyhedral oligosilsesquioxane (POSS), that feature aspects of organic functionality required for intimate dispersion and mechanically rigid silicon-oxygen polyhedral reinforcement allowing molecular reinforcement. POSS monomers are functionalized with not only a polymerizable group or groups, but also with inert organic groups at the other vertices and are within the nano-materials classification. The size of POSS, consisting of an inorganic (silicon-oxygen) core and eight variable organic side groups, is approximately 1.5 nm. Moreover, POSS aggregation to large (~50 nm), but limited scale is commonly observed.

The introduction of POSS moieties pendant to the polymeric chain has a dramatic effect on the physical properties of the polymeric materials, which was detailed in a comprehensive review paper by Li and coworkers.\(^1\) For polystyrene,\(^2\) polynorbornene\(^3\) and poly(methyl methacrylate)\(^4\) tethered by cyclopentyl-POSS (CpPOSS) and cyclohexyl-POSS (CyPOSS), the T\(_g\) monotonically increases with POSS incorporation. The retardation of polymer chain motion has been thought to be ascribed to the interaction or association between POSS groups, and/or a large inertia effect on the segmental mobility
due to the massive POSS groups. This observation is also true for thermosetting networks. For a POSS-epoxy system, Lee and Lichtenhan revealed a retardation effect that hinders the molecular motion of epoxy network junction. Surprisingly, their small-strain stress relaxation experiments revealed monofunctional POSS moieties had no contribution to the deformation process of the epoxy network and it was concluded that multifunctionality would be required for such an effect. Meanwhile, Li et al. extensively studied the influence of multifunctional-POSS on viscoelastic properties of co-polymers, revealing the significant role played by POSS incorporation. However, only few reports on the rheological behavior pertaining to the linear polymers tethered by POSS exist. Romo-Uribe et al. reported the linear viscoelastic properties of unentangled random copolymers from 4-methylstyrene and its POSS derivatives with R = cyclopentyl (Cp) and cyclohexyl (Cy). There, the rheological properties were profoundly altered with increasing POSS incorporation. At higher mole fractions of POSS (CpPOSS > 16 mol% and CyPOSS > 8 mol%), a secondary rubbery plateau with $G' \sim 10^3$ Pa, appeared replacing the expected traditional terminal zone behavior of unfunctionalized poly(4-methylstyrene). This phenomenon was attributed to intermolecular interactions involving two constituents of POSS moieties.

More generally, for polymeric liquids that feature specific interchain interactions, an additional relaxation time may appear in the rheological spectrum, either shorter than or longer than the unmodified terminal (longest) relaxation time. Moreover, such association sites can act the way entanglements do, leading to a decrease of the entanglement molecular weight in an already entangled polymeric liquid as well as a secondary rubbery plateau if the associated relaxation times are sufficiently separated. This mechanism can be described by the “sticky reptation” model developed by Leibler et al. and conceived for the hydrogen-bonded elastomers. Indeed, more recent activity in the field of supramolecular polymers has witnessed similar phenomena. Unfortunately, because of the limited polymerization degree, observations of this type for POSS’s impact on the rubbery plateau was impossible for the system previously studied. However, recent reports from Kopesky et al. found that tethered-POSS incorporated within entangled poly(methyl methacrylate) (PMMA)-based
copolymers featured decreasing plateau modulus ($G_N^0$) relative to PMMA homopolymer, motivating the present work.

How might POSS impact the state of entanglement for a polymer melt? Architecturally, POSS moieties pendant to polymeric backbone are similar to the random branches with spherical shape that are intrinsically incapable of entangling with each other or the backbone chains. Recognizing that chain topology of entangled linear polymeric liquids impacts viscoelastic behavior as explained with the reptation (or tube) model, the rubbery plateau modulus, $G_N^0$, depends on contour properties characterized by contour length $L_{pp}$, tube diameter $d_T$, Kuhn length $a_{PP}$ and packing length $l_p$ that we hypothesize to be influenced by the pendant POSS moieties. Fetters et al. derived the empirical universal relation for $G_N^o$ (Pa) and $l_p$ (Å):

$$G_N^o = 0.00226k_BT/l_p^5$$

where $k_B$ and $T$ are Boltzmann constant ($1.38 \times 10^{-23} \text{m}^2\text{kg}/\text{s}^2\text{K}$) and absolute temperature (K), respectively. This also leads to simple proportionality between tube diameter and packing length: $d_T = 19l_p$. Recently, Everaer et al. made parameter-free and quantitative predictions for $G_N^o$, through the tube model on the basis of purely topological analysis.

In the present study, we report on the thermal and rheological characterization of entangled random copolymers from styrene and its POSS derivatives with R = isobutyl (iBu) with systematical variation of the weight percentage of iBuPOSS in the polymers from 0 to 50wt%. We observe a profound impact of iBu-POSS incorporation on the glass transition temperature, the WLF free-volume parameters, and the plateau modulus, all of which are detailed quantitatively. The substantial modification role of pendant POSS group on the rheological properties of polystyrene (PS) will be emphasized, setting the stage for a subsequent report on the impact of POSS R-group variation on rheological properties.
II. Experimental Section

Synthesis of isobutylPOSS-styrene-co-styrene copolymers. Five polymers containing either 0, 6, 15, 30 or 50 wt % POSS-styrene monomer were all synthesized in the same manner as shown in Figure 1. A 10 molal monomer solution containing a total of 3 grams of monomers was initiated using 0.25 mole % azobis(isobutyronitrile) (AIBN). One example synthesis of 6 wt. % POSS follows: Under a nitrogen atmosphere, a dry O₂-free solution of toluene (2.73 mL), \([((C₈H₇)(i-C₄H₉)₇(Si₈O₁₂)] (180 mg, 0.196 mmole), styrene (2820 mg, 27.08 mmole) and AIBN (11.2 mg, 0.068 mmole) was heated to 60 °C for two days. This was then diluted with 15 mL CHCl₃ and precipitated into 100 mL of methanol. After stirring 1 hour, the copolymer was isolated on fritted glassware and air-dried overnight. \(^1\)H NMR spectroscopy (400 MHz) showed no unreacted monomers and confirmed that the product contained 6 weight % POSS.\(^{20}\) The isolated yield of dry copolymer was 93 % of the theoretical value.

![Synthetic scheme for the preparation of random copolymers yielded from styrene and styryl isobutylPOSS through radical copolymerization initialed by AIBN at 60°C.](image)

**Figure 1.** Synthetic scheme for the preparation of random copolymers yielded from styrene and styryl isobutylPOSS(BuPOSS) through radical copolymerization initialed by AIBN at 60°C.

The random copolymers prepared from styrene and \(^1\)BuPOSS-styryl through radical copolymerization were completely dissolved in the tetrahydrofuran (THF). The polymer solutions with 5wt% concentration were poured into the Teflon™ casting dishes to dry at room temperature for four
days. Next, the cast films are dried at vacuum oven at 50~60°C for two days and then 80~90°C for two days. In order to clean up the residual solvent (THF), the cast films are dried above glass transition (~120°C) for half one day further.

**Characterization**

*Thermal Analysis.* The thermal properties of the random copolymers were investigated by a TA Instrument (DSC 2920) equipped with a mechanical intercooler (temperature to T= –60°C) under a continuous nitrogen purge (50 mL/min). Both calibrations of enthalpy constant and temperature were based on a run in which one standard sample (indium) was heated through its melting point. The samples from solvent-cast films are sealed in aluminum pans with mass in the range of 5~10mg. All measurements were conducted at a scan rate of 10 °C/min following heat-cool-heat procedure from 0 °C to 250 °C. Glass transition temperatures (T_g) were determined by the midpoint of enthalpy change during the second heating.

*Wide Angle X-ray Scattering (WAXS).* In order to assess the microstructure of the random copolymers, Wide-angle X-ray Scattering (WAXS) experiments were performed at room temperature on the solvent-cast films of the samples using a BRUKER D5005 X-ray diffractometer with rotating erode source operated at 40 kV and 40 mA, which were performed at 2θ between 5.0° and 40.0°. Nickel-filtered Cu Kα radiation with wavelength, λ=1.5418Å, was used for the measurements.

*Transmission Electron Microscopy (TEM).* Real space observations of the microstructures adopted by the POSS-PS copolymers were conducted by employing transmission electron microscopy (TEM, JEOL 2010) experiments with an accelerating voltage of 200 kV. The point and line resolution are 0.19nm and 0.14nm, respectively. The thin sections with 50~60 nm thickness were cut at –100°C using a Leica Ultracut E microtome equipped with a diamond knife. The thin slices collected on 300-mesh copper TEM grids were stained under RuO₄ vapor at room temperature. Previous work revealed that this staining agent was POSS-selective.²¹
Rheological Characterization. The investigation of linear viscoelastic properties of the random copolymers was conducted by using the ARES rheometer (TA Instruments). It was equipped with two torque transducers distinguished by their torque capacity: 200 g·cm and 2000 g·cm. The existence and extent of the linear viscoelastic regime was determined by the dynamic storage and loss moduli, $G'(\omega)$ and $G''(\omega)$, as a function of strain (0.1~10%) at the angular frequency with 10 rad/s. All of the measurements were carried out within the linear viscoelastic range, where $G'(\omega)$ and $G''(\omega)$ are independent of strain. The dynamic moduli were measured as a function of frequency, $\omega$, spanning 0.01 – 100 rad/s at temperatures of 120 – 180°C. All of the rheological characterizations are performed under a nitrogen atmosphere using the parallel plates with 8 mm diameters and with the gap between two plates being about 1.0 mm.

III. Results and Discussion

1. Copolymer Synthesis

Free radical polymerization of the PS-\textsuperscript{3}BuPOSS copolymers led to polymers with high molecular weight (> 150 kg/mol) and correspondingly high number-average degrees of polymerization (DP > 1000). Polydispersity indices typical of free radical polymerizations (> 1.4) were achieved. In our earlier work, lower monomer concentrations were employed (4 versus 10 molal) and, consequently, significant lower molecular weights were achieved. Table 1 summarizes the molecular characteristics of the copolymers.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$M_w \cdot 10^3$ (g/mol)</th>
<th>$M_w / M_n$</th>
<th>DP</th>
<th>wt%POSS</th>
<th>mol%POSS</th>
</tr>
</thead>
<tbody>
<tr>
<td>0wt%iBuPOSS</td>
<td>161</td>
<td>1.43</td>
<td>1080</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>6wt%iBuPOSS</td>
<td>186</td>
<td>1.46</td>
<td>1164</td>
<td>5.9</td>
<td>0.70</td>
</tr>
<tr>
<td>15wt%iBuPOSS</td>
<td>195</td>
<td>1.40</td>
<td>1158</td>
<td>15.4</td>
<td>2.02</td>
</tr>
<tr>
<td>30wt%iBuPOSS</td>
<td>300</td>
<td>1.52</td>
<td>1387</td>
<td>33.2</td>
<td>5.34</td>
</tr>
<tr>
<td>50wt%iBuPOSS</td>
<td>419</td>
<td>1.65</td>
<td>1358</td>
<td>52.1</td>
<td>10.95</td>
</tr>
</tbody>
</table>

\textsuperscript{a}
DP’s are calculated from $M_n$ and wt% POSS is derived from mol% POSS, which is directly measured from $^1$H NMR spectroscopy.\textsuperscript{20}

2. Morphological Characterization

Figure 2 shows the wide-angle X-ray scattering (WAXS) patterns of random copolymers with respect to different content of styryl-isobutylPOSS ($^1$BuPOSS). The WAXS pattern of pure PS shows two amorphous halos: one is centered at a $d$-spacing of 4.5Å and the other one is at 8.3Å. In contrast, the pure styryl-isobutylPOSS macromer is a very crystalline substance and features numerous diffraction peaks. In particular, there is a very intense and sharp diffraction peak at 11.0Å associated with the 101$hkl$ reflection of rhombohedral lattice.\textsuperscript{22} By contrast, no such diffraction peak ascribed to the crystal of $^1$BuPOSS macromer appears in the WAXS patterns observed for the copolymers. Copolymerization apparently prevents the crystallization of POSS moieties and leads to a good dispersion with a lack of macroscopic $^1$BuPOSS incorporation. With increasing $^1$BuPOSS loading, the amorphous halo at the $d$-spacing = 8.3Å shifts to the lower angle and the corresponding peak is slightly narrowed. Meanwhile, the amorphous halo at the $d$-spacing = 4.5 Å becomes a little broadened, with scattering from the 50 wt% $^1$BuPOSS sample revealing the origin to be an emergence of an underlying shoulder peak. Furthermore, in contrast with the $^1$BuPOSS macromer, these two peaks are quite distinct from any crystalline reflections of the $^1$BuPOSS macromer. The strongest $^1$BuPOSS reflection peaks appear at $d$-spacing of 11.0 Å and 8.1 Å; however, these regions of the copolymer WAXD patterns are devoid of any features. Thus it clearly indicates that co-polymerization effectively prevents crystallization of $^1$BuPOSS units.
To further elucidate the microstructure of the random copolymers, we made direct observations of selected samples that were microtomed and selectively stained using transmission electron microscopy (TEM). The dark particles revealed in Figure 3(a) directly image the POSS-related particles in the sample with 6wt% 1BuPOSS. The size is only from 1.5 to 3 \( \text{nm} \), approaching the size of a single POSS molecule. Further increase in 1BuPOSS concentration leads to the increase of dispersed particle density shown in Figure 3(b), however, the particle size nearly does not change. We infer that POSS-cages pendant along the PS chain are well-dispersed in the matrix with a molecular level and there is no obvious aggregation of POSS cages, consistent with our WAXS observations described above. The peaks related to the presence of POSS only shift from \( d \)-spacing centered at 8.3 \( \text{Å} \) to 9.7 \( \text{Å} \) implying that the POSS cages are incorporated within the PS chain packing. We conclude that the 1BuPOSS groups grafted on the PS chain were dispersed in the polystyrene matrix at nearly molecular level.
Figure 3. Transmission electron microscopy (TEM) images of as-cast films of $^i$BuPOSS-PS copolymers: (a) 6; (b) 30 wt-% POSS. The samples were stained by RuO₄ vapor at ambient condition.

In contrast, we have noticed POSS aggregation is a commonly observed phenomenon in polynorborene, polyurethane, polyimide, etc, tethered by POSS molecules. Romo-Uribe et al. reported the both cyclopentyl-POSS (CpPOSS) and cyclohexyl-POSS (CyPOSS) pendant on the poly(4-methylstyrene) can aggregate to form nanocrystals when the POSS loading is up to the critical mol% content. Zhang et al. studied the effect of random copolymers of PMMA-POSS on the phase segregation of the typical immiscible polymer blend of PMMA and PS prepared from toluene spin coating. They found that the cyclopentyl functionalized POSS grafted onto the backbone of PMMA plays a compatibilizing role in polymer blend of PMMA and PS. It was concluded that a favorable interaction exists between the POSS functional group and the PS homopolymer. So, POSS aggregation essentially depends on the compatibility between POSS molecules and polymer main chain and there may be a critical concentration, beyond which POSS will tend to form microphase separation from the matrix. Although the POSS used here is functionalized with isobutyl, it is still reasonable for us to expect that $^i$BuPOSS has the potential to be compatible with PS homopolymer. Indeed, our morphological characterizations have revealed that $^i$BuPOSS were dispersed with a molecular level, indicating $^i$BuPOSS/PS interactions.
2. Thermal analysis.

The effects of the presence of tethered POSS cages on the glass transition were studied using
differential scanning calorimetry (DSC). Second heating for the styrene and styryl-\textsuperscript{1}BuPOSS random
copolymers are shown in Figure 4. The glass transition temperatures determined from the midpoint of
enthalpy change are shown in Figure 5, where we can observe that the glass transition temperature of
the random copolymers with \textsuperscript{1}BuPOSS decreases with the increasing POSS content. In addition, we
observe a significant and monotonic decrease in the heat capacity change at T\textsubscript{g} with increasing POSS
loading.

![Figure 4](image)

**Figure 4.** Differential Scanning Calorimetry (DSC) analysis of as-cast \textsuperscript{1}BuPOSS-PS copolymers films
with varying weight percentage of \textsuperscript{1}BuPOSS: (i) 0, (ii) 6, (iii) 15, (iv) 30, and (v) 50 wt-% POSS. The
heating rate was 10 °C/min and the second heating traces are shown. The dash line
represents the T\textsubscript{g} of PS homopolymer determined from the midpoint of enthalpy change.

![Figure 5](image)

**Figure 5.** The variation of glass transition temperature (\textbullet, T\textsubscript{g}) and heat capacity step change (\textbullet, \Delta C\textsubscript{p}) at T\textsubscript{g} as functions of \textsuperscript{1}BuPOSS
content. T\textsubscript{g} is determined from the midpoint of enthalpy change in the DSC trace curves shown
in Figure 4.
The glass transition values of polymers reflect, in part, the rotational energy barrier about $\sigma$ bonds in the backbone chain, lower glass transition temperatures indicating a lower energy barrier. For polymer families with the same backbone structure, the side chain or group and intermolecular interactions greatly influence $T_g$. While rigid side chains restrict torsion about $\sigma$ bonds in the backbone chain, raising $T_g$, flexible side chains, such as the family of poly(alkyl methacrylate)s, exhibit lowering of $T_g$ by “internal plasticization” where the side group increases interchain spacing and free volume, lowering rotation energy barrier and decreasing $T_g$. Thus, considering backbone rotation alone, side chains may either increase or decrease $T_g$, depending on which of the two mentioned factors dominates.

Additionally, if there is strong intermolecular interaction, such as hydrogen bonding, ion-ion interaction, acid-base interaction, among others, the formation of physical cross-linking would restrict polymer chain motion, thus enhancing $T_g$. In POSS-branched polymers, intermolecular interactions consist of both POSS-POSS and POSS-polymer matrix. The styryl-POSS macromer is functionlized with a polymerizable styrenic group, and also decorated with seven inert organic groups (so-called POSS R-group) at other vertices, themselves contributing 80% of the POSS volume and strong R-group sensitivity. For example, POSS incorporation with cyclopentyl(Cp) and cyclohexyl(Cy) R-groups has been uniformly observed to enhance $T_g$ of random copolymers, such as poly(4-methylstyrene-POSS), poly(norbornyl-POSS), poly(methacrylate-POSS), poly(siloxane-POSS), among others. However, Xu et al. reported that styryl-isobutylPOSS incorporation into poly(acetoxy styrene)(PAS) and poly(vinylpyrrolidone)(PVP) first decreases and then increases $T_g$ as the POSS comonomer content is increased.

Here, our copolymers have a polystyrene backbone randomly “branched” by $^1$BuPOSS. The $^1$BuPOSS branches effectively increase the distance between adjacent backbone chains, as shown in Figure 2, and lead to the increase of the mean size of local free volumes. As a result, the chain mobility is increased and $^1$BuPOSS, from this factor alone, may play a plasticizing role. However, the massive volume of POSS cage can undoubtedly induce a steric barrier, increasing the resistance to torsional
rotation about σ bonds in the polystyrene backbone chain. Morphological characterization reveals that 1BuPOSS disperse in the PS matrix with a molecular level. Thus, in our samples, the inter-chain interactions are between molecularly dispersed 1BuPOSS segments and PS segments through relatively weak van der Waals forces. The variation of glass transition temperature in the random copolymers is the net result of all three effects: free volume, steric barrier and POSS—polymeric segment interaction. Our DSC data (Figs. 4, 5) reveal that the glass transition monotonically decreases, though with upward curvature, with increasing 1BuPOSS content, indicating that the dominant factor in determining Tg is the addition of free volume effect by 1BuPOSS. This interpretation will be further supported with rheological measurements employing time-temperature superposition.

3. Rheological Behavior.

A. Time Temperature Superposition

We investigated the applicability of time-temperature superposition (TTS) to the subject PS-POSS copolymers by conducting frequency sweep tests spanning the temperature range 120 °C < T < 180 °C. This limited range was selected in order to avoid thermal degradation and side reactions, yet feature enough sample compliance to avoid exceeding the torque limit of the instrument. All of the experiments were within the linear viscoelastic regime determined by strain sweep tests. Relative to a fixed reference temperature (T=120 °C), we established the time-temperature superposition (TTS) by shifting the logarithmic plots of G'(ω) and G"(ω) along the frequency axis horizontally. Vertical shifts along the modulus axis were not required. Thus, for good superposition,

\[ G'(\omega, T) = G'(a_T \omega, T_r) \]  
\[ G"(\omega, T) = G"(a_T \omega, T_r) \]

where \( T_r \) is the reference temperature and \( a_T \) are the temperature-dependent frequency shift factors. Figure 6 shows the master curves of PS-host random copolymers with different 1BuPOSS content, wherein data extend over ca. 7 decades of reduced frequency.
Figure 6. Master Curve of as-cast films of \( ^1 \text{BuPOSS-PS} \) copolymers with varying weight percentage of \( ^1 \text{BuPOSS} \): (○) 0, (□) 6, (△) 15, (▽) 30, and (◇) 50 wt-% POSS with \( T_r = 120 \, ^\circ \text{C} \): (a) \( G' \); (b) \( G'' \) and (c) \( \tan \delta \). The arrows in (c) indicate the minimum values of \( \tan \delta \) for each sample.
Within the temperature range, time-temperature-superposition was found to be satisfactory and applicable as expected for single phase, amorphous polymers far from any thermally-stimulated morphological transformation. All of the polymers exhibit a rubbery plateau, terminal zone, and the rubber-liquid transition regimes of viscoelastic behavior. This confirms that all of the samples form entangled melts, as anticipated based on the molecular weight values (Table 1).

The time-temperature-superposition shift factor, \( a_T(T) \), can be described by the WLF-equation,\(^{31}\) that is derived on the basis of temperature dependent free volume:

\[
\log a_T = \frac{-C_1^g(T - T_g)}{C_2^g + (T - T_g)}
\]  

(5)

Here, \( C_1^g = B/2.303f_g \) and \( C_2^g = f_g/\alpha_f \), where \( f_g \) is fractional free volume at \( T_g \), \( \alpha_f \) is the temperature coefficient of fractional free volume and \( B \) is a constant, generally assumed to be unity. If the reference temperature, \( T_r \), is different from glass transition temperature, WLF equation will be rewritten as:

\[
\log a_T = \frac{-C_1^r(T - T_r)}{C_2^r + (T - T_r)}
\]  

(6)

where \( C_1^r = B/2.303f_r \), \( C_2^r = f_r/\alpha_f \), and \( f_r \) is the fractional free volume at the reference temperature \( T_r \).

Inspection of Eqn. (6) reveals that plots of \( 1/\log(a_T) \) vs. \( 1/(T-T_r) \) should be linear (if the expression holds) and that the parameters may be determined from the plots’ slopes and intercepts. For all random copolymers examined, such plots (Figure 7) indeed feature a linear relationship, indicating validity of Eqn. 6. From the slopes and intercepts, we have determined \( C_1^r \), \( C_2^r \) and \( f_r \), for each copolymer whose values at the glass transition temperature can be calculated using the following equations.\(^{31}\)

\[
C_1^g = \frac{C_1^r C_2^r}{C_2^r + T_g - T_r}
\]  

(7)

\[
C_2^g = C_2^r + T_g - T_r
\]  

(8)

\[
f_g = \frac{B(C_2^r + T_g - T_r)}{2.303 C_1^r C_2^r}
\]  

(9)
Figure 7. The WLF function plot of shift factor for as-cast films of ^BuPOSS-PS copolymers with varying weight percentage of ^BuPOSS: (●) 0, (○) 6, (■) 15, (□) 30, and (▲) 50 wt-% POSS. The reference temperature is 120°C.

The calculated values of free volume parameters ^BuPOSS-PS random copolymer are tabulated in Table 2. There is not a clear trend in the value of the fractional free volume, \(f_r\), as a function of ^BuPOSS loading at the reference temperature, 120°C, which is not surprising given the variation in \(T_g\) relative to the arbitrary \(T_r\). However, it is very interesting to find that the fractional free volume at \(T_g\) \(f_g\) monotonically decreases with decreasing ^BuPOSS content, where \(f_g\) of PS is almost equal to the numerical value reported elsewhere.\(^{32}\) This result implies that ^BuPOSS molecules can create remarkable void volume in the glass and melts, which leads to associated increases of the segmental mobility. The decrease of glass transition temperature with increasing ^BuPOSS loading observed can be attributed to the additional free volume generated by the tethered POSS molecules.
Table 2. Summary of Viscoelastic Characteristics of Polystyrene-POSS Copolymers with isobutyl R-group at reference temperature 120 °C.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$C_1^f$</th>
<th>$C_2^f$(K)</th>
<th>$f_r/B$</th>
<th>$f_g/B$</th>
<th>$\alpha_f$ (K⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0wt%iBuPOSS</td>
<td>8.18</td>
<td>47.85</td>
<td>0.0531</td>
<td>0.0283</td>
<td>11.09×10⁻⁴</td>
</tr>
<tr>
<td>6wt%iBuPOSS</td>
<td>8.05</td>
<td>60.10</td>
<td>0.0540</td>
<td>0.0298</td>
<td>8.98×10⁻⁴</td>
</tr>
<tr>
<td>15wt%iBuPOSS</td>
<td>9.15</td>
<td>85.55</td>
<td>0.0475</td>
<td>0.0319</td>
<td>5.55×10⁻⁴</td>
</tr>
<tr>
<td>30wt%iBuPOSS</td>
<td>8.79</td>
<td>92.04</td>
<td>0.0494</td>
<td>0.0339</td>
<td>5.36×10⁻⁴</td>
</tr>
<tr>
<td>50wt%iBuPOSS</td>
<td>8.55</td>
<td>96.63</td>
<td>0.0508</td>
<td>0.0345</td>
<td>5.25×10⁻⁴</td>
</tr>
</tbody>
</table>

Meanwhile, the temperature coefficient ($\alpha_f$) of the fractional free volume also decreases with increasing iBuPOSS loading (Table 2) and at the higher wt-% of iBuPOSS, it seems to level off. In other words, the temperature dependence of fractional free volume is lower for increasing iBuPOSS content. According to WLF theory (shown in Fig. 7 to apply) the fractional free volume $f$ can be written approximately as:

$$f = f_g + \alpha_f (T - T_g) \quad T \geq T_g \quad (10)$$

Following this relation and using measured values for all of the parameters, Figure 8 reveals the temperature dependence of free volume fraction for the regime rheologically characterized (120 < $T$ < 180 °C) for the iBuPOSS-PS random copolymers. For the PS homopolymer, despite having the lowest fractional free volume at $T_g$ (Table 2), it shows the largest value at higher temperature due to its largest $\alpha_f$ value. In contrast, for large iBuPOSS-content copolymers the change of fractional free volume with temperature is much smaller. In this sense, the incorporated POSS moieties are somewhat athermal as is the case of inorganic fillers.
Figure 8. The temperature dependence of fractional free volume \( f \) as-cast films of \( \text{BuPOSS-PS} \) copolymers with varying weight percentage of \( \text{BuPOSS} \): (——) 0, (−−−) 6, (−−−−) 15, (−·−·−) 30, and (−−−−−−) 50 wt-% POSS.

B. Rubbery Plateau

Inspection of Figure 6 reveals not only significant thermorheological simplicity for all of the copolymers, but also a strong dependence of the plateau modulus on level of \( \text{BuPOSS} \) incorporation. In characterizing the rubbery plateau regime, the preferred method to estimate the plateau modulus would be to integrate the area under the terminal loss peak,

\[
G_N^0 = \frac{4}{\pi} \int_{-\infty}^{\omega_{\text{max}}} G''(\omega) \ln \omega
\]

where \( \omega_{\text{max}} \) is the frequency at the maximum value of loss modulus \( G''_{\text{max}} \) in the plateau zone. However, due to molecular weight polydispersity (Table 1), our data don’t show a peak in loss modulus. Alternately, there are several semi-empirical ways to estimate the value of the rubbery plateau, \( G_N^0 \).

Wu\textsuperscript{33} proposed an expression to describe the relationship between the crossover modulus, \( G_c \), \([G_c(\omega)=G'(\omega)=G''(\omega)]\) and plateau modulus, \( G_N^0 \). The ratio of \( G_N^c \) and \( G_c \) can be expressed as a function of molecular weight polydispersity, \( p = M_w/M_n \).
\[ \log\left(\frac{G_N^o}{G_c}\right) = 0.380 + \frac{2.63(\log p)}{1 + 2.45(\log p)} \]  

(12)

where \( p \) should be smaller than 3. This equation has been confirmed both theoretically\textsuperscript{33, 34} and experimentally for many homogenous polymer melts. A disadvantage, however, is reliance on arguably inaccurate \( p \) data. Finally, the method ultimately adopted for plateau modulus measurement utilizes the “\( \tan\delta \) minimum criterion”, wherein \( G_N^o \) is set equal to the storage modulus \( G' \) at the frequency where \( \tan\delta \) is at its minimum in the frequency spectrum:\textsuperscript{34, 35}

\[ G_N^o = \left| G' \right|_{\tan\delta \rightarrow \text{min}} \]  

(13)

Indeed, Figure 6(c) revealed that a minimum in \( \tan\delta \) was well-determined for all samples. Figure 9 shows the plateau modulus obtained from these two semi-empirical methods in the random copolymer with \(^1\)BuPOSS. For both cases, the plateau modulus monotonically decreases with \(^1\)BuPOSS content. Moreover the values of \( G_N^o \) calculated from crossover modulus method are higher than those obtained from minimum \( \tan\delta \). The \( G_N^o \) (\( \approx \)2.0\texttimes10\textsuperscript{5} Pa) of PS determined by \( G' \) at minimum \( \tan\delta \) is much closer to the value of plateau modulus obtained by integrating the area under the terminal loss peak of mono-dispersed entangled PS.\textsuperscript{32} Henceforth we adopt the “\( \tan\delta \) minimum criterion” as our preferred method.

**Figure 9.** Rubbery plateau modulus \( (G_N^o) \) as a function of \(^1\)BuPOSS weight fraction: (●) evaluated from the crossover modulus and molecular weight polydispersity, and (▲) determined from \( G' \) at minimum \( \tan\delta \). The dash reference line is plateau modulus of PS reported in ref. 31.
The relationship between the characteristic value of $G_N^o$ and molecular weight between entanglements, $M_e$, is given by the relation\textsuperscript{17, 31, 36}:

$$M_e = \frac{4 \rho RT}{5 G_N^o}$$  \hspace{1cm} (14)

where R is the universal gas constant ($=8.314 \text{ J} \cdot \text{mol}^{-1} \cdot \text{K}^{-1}$), $\rho$ is density, T is the absolute temperature. The density of PS is 1.05 g/cm\textsuperscript{3} at room temperature. Barry\textsuperscript{37} and Larsson\textsuperscript{38, 39} reported crystal densities of POSS-related molecules with different R-groups from CH\textsubscript{3}- (1.51 g/cm\textsuperscript{3}) to n-C\textsubscript{4}H\textsubscript{9}- (1.14 g/cm\textsuperscript{3}) to 1-naphthyl- (1.24 g/cm\textsuperscript{3}). Further, the density of $^i$BuPOSS has been estimated\textsuperscript{15} to be 1.15 g/cm\textsuperscript{3}. All of them have densities higher than PS at room temperature. Qualitatively, it is reasonable that the density values of random copolymers with different R-group are higher than pure PS, although values of density for $^i$BuPOSS random copolymers have not been reported. Regardless, melt densities are not expected to change more than several percent between samples while, in contrast, $G_N^o$ values were observed to change nearly an order of magnitude. The relationship between density ($\rho$, g/cm\textsuperscript{3}) of pure polystyrene and temperature (T, °C) in the range from 100° to 220° can be expressed as follows:\textsuperscript{40}

$$\rho = 1.0865 - 6.19 \times 10^{-4} T + 0.136 \times 10^{-6} T^2$$  \hspace{1cm} (15)

Assuming such temperature-dependent density values, even for our copolymers, the entanglement molecular weight and the number of entanglements per chain of our $^i$BuPOSS-PS copolymers calculated from Eqn. 14 are detailed in Table 3. One can observe that the entanglement molecular weight clearly increases with $^i$BuPOSS loading. In effect, $^i$BuPOSS incorporation along the PS chain dilutes the entanglement density. Meanwhile, the entanglements per chain of our copolymers vary in a range from ~7 to ~13, which further confirms all of the samples effectively entangled, as anticipated based on the molecular weight. The terminal slopes also listed in Table 3 will be discussed later.

From the viewpoint of chemical structure, POSS groups grafted on the PS chain play the same role of branches and essentially change the topology of the PS chain. There have been very extensive studies on the effects of microscopic topology on the rheological behavior in the entangled polyolefin
However, careful studies on randomly branched polystyrene have received less attention. These studies mainly focused on the effect of the long chain branches (LCBs) on the linear and non-linear rheological properties. How spherical cage-like branches, such as POSS molecules, affect the rheological behavior has not been studied.

**Table 3. Rheological Properties of Polystyrene-POSS Copolymers with isobutyl R-group.**

<table>
<thead>
<tr>
<th>Compound</th>
<th>(G_N^0) (kPa)</th>
<th>(M_e) (kg/mol)</th>
<th>(Z=M_w/M_e)</th>
<th>((dG'/d\gamma\omega)_{\omega\rightarrow0})</th>
<th>((dG''/d\gamma\omega)_{\omega\rightarrow0})</th>
</tr>
</thead>
<tbody>
<tr>
<td>0wt%iBuPOSS</td>
<td>205</td>
<td>12.9</td>
<td>12.5</td>
<td>1.65</td>
<td>0.99</td>
</tr>
<tr>
<td>6wt%iBuPOSS</td>
<td>123</td>
<td>21.5</td>
<td>8.6</td>
<td>1.44</td>
<td>0.98</td>
</tr>
<tr>
<td>15wt%iBuPOSS</td>
<td>93</td>
<td>28.5</td>
<td>6.8</td>
<td>1.35</td>
<td>0.97</td>
</tr>
<tr>
<td>30wt%iBuPOSS</td>
<td>83</td>
<td>31.9</td>
<td>9.4</td>
<td>1.33</td>
<td>0.96</td>
</tr>
<tr>
<td>50wt%iBuPOSS</td>
<td>42</td>
<td>61.6</td>
<td>6.8</td>
<td>1.27</td>
<td>0.93</td>
</tr>
</tbody>
</table>

Fetters et al.\textsuperscript{18} developed the relationship between the entanglement density of polymer species and their molecular dimensions in the melt state, resulting in the expressions:

\[
M_e = 218 \times 10^{-6} \rho l_p^3
\]

where \(l_p(\text{Å})\) is the packing length defined as the occupied volume of a chain divided by the mean square end-to-end distance, \(M_e(\text{kg/mol})\) is the entanglement molecular weight, \(\rho(\text{kg/mol})\) is the density and the units of the prefactor, \(218 \times 10^{-6}\), is \(\text{m}^3\cdot\text{mol}^{-1}\cdot\text{Å}^{-3}\). Roughly, this length may be interpreted as the effective chain profile or width. Combining Eqn. 14 and Eqn. 15 allows a density-independent determination of \(l_p\) in terms of \(G_N^0\),

\[
l_p = \left( \frac{3.67 \times 10^3}{RTG_N^0} \right)^{1/3}
\]

where \(R(\text{J}\cdot\text{mol}^{-1}\cdot\text{K}^{-1})\) is the universal gas constant, \(T(\text{K})\) is absolute temperature, \(G_N^0\) (Pa) is the rubbery plateau modulus and prefactor units, \(3.67 \times 10^3\), are \(\text{Å}^3\cdot\text{m}^{-3}\cdot\text{mol}^{-1}\). Thus, smaller plateau moduli imply larger \(l_p\), or chain profiles, as well as larger tube diameters \(d_t\) \((d_t = 19 l_p)\).\textsuperscript{18}
Using Eqn. 17 and the plateau modulus data of Figure 9, we computed values of $l_p$ for $i^\text{BuPOSS}$-PS random copolymers (Figure 10). We find that the packing length monotonically increases from a value of 3.88 Å for PS homopolymer (reported elsewhere as 3.92 Å)\textsuperscript{18} to a value at 6.63 Å for 50wt% (10.95 mol%; $\approx 1$ in 10 repeat units bearing a POSS branch). This is a very large packing length increase (71%). A similar increase (73%) is observed in polymethacrylates when the pendant methyl group of PMMA is uniformly (every repeat unit) substituted by an octyl group.\textsuperscript{18} Thus, substituting one $i^\text{BuPOSS}$ on every tenth repeating unit of a PS chain is topologically equivalent to substituting every repeat unit of a PMA chain with an n-octyl chain. Interestingly enough, ten n-octyl chains features a molecular weight of 1130 g/mol, comparable to (but larger than) the molecular weight of each pendant POSS moiety, 816 g/mol. Certainly, the estimated values of packing length of $i^\text{BuPOSS}$-PS copolymers should be confirmed further by any other accurate experiments, such as small-angle neutron scattering (SANS) and the finer synthetic methods to prepare the narrow dispersed samples would also be beneficial.

![Figure 10](image.png)

**Figure 10.** Plot of packing length ($l_p$) of as-cast films of $i^\text{BuPOSS}$-PS copolymers with varying weight percentage of $i^\text{BuPOSS}$ as a function of POSS weight fraction. The packing length is calculated using Eqn. 17.
From this topological viewpoint, we can explain the relation between the observed dependence of plateau modulus on \(^1\text{BuPOSS}\) content. A schematic drawing to describe the effect of pendant \(^1\text{BuPOSS}\) groups on the dimension of the reptation tube, \(l_p\) or \(d_t\), is shown in Figure 11. Undoubtedly, the pendant \(^1\text{BuPOSS}\) has a great influence on the PS chain topology. Presuming that the dynamics of the present random copolymers follow the tube model, the resultant average tube diameter will increase due to the presence of the pendant \(^1\text{BuPOSS}\) molecules and consequently lower the entanglement density. We envision that the average tube diameter of the constitute PS chains increases with the increasing POSS content. Consequently, the plateau modulus monotonically decreases with increasing POSS content due to the variation of microscopic topology of PS chain.

[Diagram of Polystyrene (PS) and PS-r-POSS]

**Figure 11.** Schematically drawing the effect of the pendant POSS-group on the PS chain topology and the resultant tube dimension.

Possible intermolecular interactions have been ignored in the discussions above. If there exists the strong intermolecular interaction, the plateau modulus will be enhanced and there will be two dominant mechanical relaxations above the glass transition temperature (\(T_g\)): one is for disentanglement and the other is for disassociation of specific intermolecular interaction. This mechanism can be described by “sticky reptation” model developed by Liebler et al.\(^{10}\) conceived for hydrogen-bonded elastomers. Although the interaction due to the individual R-group is weak, each POSS cage contains seven R-groups, resulting in so-called multiple interactions to magnify the total interaction of each POSS cage. Romo-Uribe et al.\(^{9}\) observed the rheological behavior in the unentangled random copolymers with styrene-based POSS cyclopentyl- or cyclohexyl-based and 4-methylstyrene. There,
when POSS loading was increased beyond a critical value, a terminal zone region couldn’t be reached at any high temperature or low frequency and a secondary rubbery plateau appeared. In particular, they observed the critical value for this transformation that depended on R-group: CpPOSS was around 60wt% and while CyPOSS was at 40wt%.

In our case, the iBuPOSS molecules pendant on the PS chain are dispersed at a molecular level and there is not obvious aggregation observed. The primary attractive force of interaction is between iBuPOSS and PS segments. Nevertheless, we observed only a single mechanical relaxation above Tg, this being due to disentanglement. Thus we can propose that the interaction between iBuPOSS and PS matrix is weak and incapable of yielding a secondary rubbery plateau. In our random copolymers from styrene and styryl iBuPOSS, the effect of iBuPOSS on the microscopic chain topology is the dominant effect on the plateau modulus. Meanwhile, it can also be expected that the interaction between POSS and PS matrix is tunable by varying the R-group. Indeed, the R-group dependence of rheological behavior will be discussed in a following paper.

Within the rubbery plateau regime, polymers with linear chain branches normally follow two kinds of relaxation mechanisms below the dynamic glass transition: the branches relax with the rapid Rouse motion (or rapid reptation) at higher frequencies with the effective tube diameter increasing continuously until the branch arms are completely retracted. Then the primary backbone follows the reptation-like relaxation in the widened tubes at the lower frequencies. Thus, there are two separated relaxation scales, a shorter one for branches and a longer one for the backbone. However, all of the master curves shown in Figure 6 do not feature such a relaxation at the higher frequencies, instead featuring only one characteristic relaxation. We attribute this finding to the fact that POSS branches are different from the common polymeric coil branches. In particular, they feature a compact, nanoscale structure incapable of entangling and thus experiencing associated mechanical relaxation.

### C. Behavior in Terminal Zone

At the terminal zone, the polymer melts should be fully relaxed and follow the characteristic power laws for frequency dependence of the shear storage and loss moduli: \( G' \sim \omega^2 \) and \( G'' \sim \omega^1 \). The
slopes of terminal zones were estimated for random copolymers with variation in \textsuperscript{1}BuPOSS content and reported in Table 3. The terminal zone frequency dependence of the storage modulus decreases with the increased \textsuperscript{1}BuPOSS from $\omega^{1.65}$ for pure PS to $\omega^{1.27}$ for 50wt\% \textsuperscript{1}BuPOSS. Meanwhile, there also appears to be a slight decrease in the power-law dependence for $G''$ with the \textsuperscript{1}BuPOSS loading, from $\omega^{0.99}$ for pure PS to $\omega^{0.93}$ for 50wt\% \textsuperscript{1}BuPOSS. According to Doi-Edwards reptation theory,\textsuperscript{17} the relationship between $G'$ and $G''$ for the linear flexible polymer can be expressed as

$$\log G' = 2 \log G'' - \log \left( \frac{\rho R T}{M_c} \right) + \log \left(\frac{\pi^2}{8}\right)$$

where $\rho$ is the density, $R$ is universal gas constant and $T$ is the absolute temperature. We note that such a plot is independent of molecular weight. For the ideal polymeric liquid, the slope of $\log G'$ versus $\log G''$ is 2 at the terminal zone, while the intercept can directly yield $G_N^0$. Figure 12 shows the logarithmic plot of $G'$ and $G''$ (termed the Han plot\textsuperscript{50}) with different \textsuperscript{1}BuPOSS loading.

![Figure 12. Double logarithm plot of $G'$ versus $G''$ for as-cast films of \textsuperscript{1}BuPOSS-PS copolymers with varying weight percentage of \textsuperscript{1}BuPOSS: (○) 0, (▲) 15, and (◇) 50 wt-% POSS.](image)

Like the frequency dependence of $G'$, the slope of $G'$ versus $G''$ also has an \textsuperscript{1}BuPOSS content dependence. In particular, the slope decreases with the \textsuperscript{1}BuPOSS loading. Compared with the tendency
of molecular weight polydispersity with \textsuperscript{i}BuPOSS content, we can’t attribute this deviation from the classical terminal behavior only to the effect of molecular weight polydispersity. Instead, intermolecular interactions may play a role. Morphological characterization showed us that the pendant \textsuperscript{i}BuPOSS groups are dispersed at nearly a molecular level. We argue, then, that the deviation from the classical terminal behavior may be ascribed to the existence of finite interactions between \textsuperscript{i}BuPOSS and PS segments. Although the primary attraction force of interaction between POSS and PS-matrix is weak and therefore, less important in the rubbery plateau frequency regime (where stresses are higher), this interaction could reasonably have influence on the terminal zone behavior as reflected in the Han-plot slope.

We now consider the terminal relaxation time as influenced by \textsuperscript{i}BuPOSS incorporation. Due to significant variation in molecular weight of the copolymer series (Table 1), we focus on the temperature dependence of the terminal relaxation time, rather than the absolute values. The terminal zone can be characterized by several parameters, including the terminal relaxation time, \( \tau \), the zero shear viscosity \( \eta_0 \), and steady state shear compliance \( J_e^0 \). The defining expressions are as follow:

\[
\tau \equiv \eta_0 J_e^0 \tag{19}
\]

\[
\eta_0 = \lim_{\omega \to 0} \frac{G''(\omega)}{\omega} \tag{20}
\]

\[
J_e^0 = \lim_{\omega \to 0} \frac{G'(\omega)}{G''(\omega)^2} \tag{21}
\]

Accordingly, the relaxation time values for each copolymer could be determined from zero-frequency limits of storage and loss modulus spectra obtained at different temperatures.

The temperature dependences of the \textsuperscript{i}BuPOSS-PS copolymer terminal relaxation times were well described by the Vogel-Fulcher-Tamman-Hesse (VFTH) equation:

\[
\tau = A \exp \left( \frac{B}{T - T_0} \right) \tag{22}
\]
where \( T_o \) is the “ideal glass transition temperature” (or “Vogel temperature”) and \( B \) is termed an “apparent activation energy,” physically representing an Arrhenius-like temperature activation energy (though with units of temperature). Figure 13 shows the temperature dependence of terminal relaxation times of the polystyrene and styryl-\(^1\)BuPOSS random copolymers, together with the best-fit VFTH curves. The values of \( T_o \) and \( B \), extracted from the best-fit VFTH-equation curves, are reported as a function of \(^1\)BuPOSS content in Figure 14. We observed that the “apparent activation energy”, \( B \), increases by ca. 50% and \( T_o \) decreases by ca. 40°C when POSS loading is increased from 0 to 50 wt-%. (Note by inspection of Eqn. 22 that increasing \( B \) corresponds to decreasing temperature sensitivity.) Thus, similar to our observations of decreasing \( \alpha_f \) with increasing POSS loading (Table 2), we observed a lowering of the temperature-sensitivity for the terminal relaxation time with \(^1\)BuPOSS copolymerization, as manifested in the VFTH \( B \) “activation energy” parameter.

**Figure 13.** Temperature Dependence of Terminal Zone Relaxation Time (\( \tau \)) for as-cast films of \(^1\)BuPOSS-PS copolymers as a function of POSS weight fraction: (●) 0, (○) 6, (■) 15, (□) 30, and (▲) 50 wt-% POSS. The solid lines represent the best-fit VFTH curves for the copolymers and PS homopolymer.
Figure 14. Plot of Vogel temperature ($T_0$, ⋄) and apparent activation energy (B, ■) of $^i$BuPOSS-PS copolymers as a function of $^i$BuPOSS content, which are extracted from the best-fit curves shown in Figure 14. The symbols of open circle (○) and open square (□) are the values of $T_0$ and B for PS from ref. 46.

There are several theoretical models to explain the effects of branches on the temperature dependence of terminal relaxation time. According to the coupling model,$^{54-56}$ the dynamics of polymer chains can be described by the coupling parameter $n$ ($0 < n < 1$), where the value of $n$ is closely related to the topological constraints of the entangled system. The surrounding environment experienced by a single macromolecular chain is the key factor of this parameter. In particular, the parameter values of the linear and branched polymer can be expressed by their corresponding activation energy, $E$:

$$\frac{E_{\text{branch}}}{E_{\text{linear}}} = \frac{1-n_{\text{linear}}}{1-n_{\text{branch}}}$$ \hspace{1cm} (23)

The coupling model predicts that the temperature dependence of branched polymers will always be weaker (larger $E$) than that of the corresponding linear polymers, as $n_{\text{branch}} > n_{\text{linear}}$. More specifically, branched polymers are expected to have the stronger “coupling” that acts to increase $n$, resulting in broadening the relaxation function and the increase of activation energy of the rheological properties. Usually, this model is valid for branched polymers for which the molecular weight of branches exceeds the entanglement molecular weight, $M_c$.\textsuperscript{54-56} However, only a few studies have been reported for branched polystyrene.\textsuperscript{45-47} Ferri et al.\textsuperscript{47} prepared the randomly branched polystyrene by copolymerization of styrene (St) and divinylbenzene (DVB) and studied the melt rheology of linear and randomly branched polystyrene (LPS ad RBPS), where the RBPS featured entangled branches ($M_{\text{br}}/M_c$...
They found that while the shift factors \( a_T \) of LPS were similar to those of RBPS, the activation energy slightly increased with degree of branching. Our results are qualitatively consistent with these RBPS data, despite architectural dissimilarity.

Semiquantitative comparison of our observations with atomistic molecular dynamics simulations is also possible. In particular, simulations of POSS-polynorbornene random copolymers due to Bharadwaj, Berry and Farmer indicated a weaker temperature-dependence of elastic moduli for CyPOSS- and CpPOSS-polynorbornene copolymers than for the reference PN homopolymer, consistent with our present observations of increasing VFTH apparent activation energy. There, it was reasoned that POSS units act as “anchor” against molecular motion. While it would seem that such anchoring effect would be most important at short time scales, not within the terminal zone, it is reasonable that the nature of polymer reptation with periodic “anchor” requires attention at the level of polymer chain diffusion calculations. To finally elucidate this aspect, the finer synthetic methods to prepare the narrow dispersed random copolymers from styrene and styryl \(^1\)BuPOSS would be beneficial.

**IV. Conclusions**

We investigated the thermal and linear rheological behavior of random copolymers from styrene and styryl-based \(^1\)BuPOSS. The molecularly homogeneous copolymers featured decreasing \( T_g \) with increasing \(^1\)BuPOSS incorporation, a finding attributed to increased free volume. The dynamic moduli were measured within the linear viscoelastic regime over a range of temperatures from 120 °C to 180 °C. Rheological characterization showed that time-temperature superposition (TTS) works well over a range of \(^1\)BuPOSS content up to 50 wt-%. The well-fitted WLF equation revealed that the fractional free volume at \( T_g \), \( f_g \), increases with the \(^1\)BuPOSS loading; however, the corresponding temperature coefficient of free volume was seen to decrease with increasing \(^1\)BuPOSS content. We propose that the \(^1\)BuPOSS groups play a plasticizer-like role in determining the glass transition temperature like linear branches, increasing the glassy free volume fraction to enhance the PS segmental mobility.
Compared with the linear-branched polymers, there is no branch (arm) retraction mechanism at the short times observed in our PS branched by iBuPOSS. The rubbery plateau modulus, defined as the shear storage modulus at the condition of minimum loss tangent, decreases with increasing iBuPOSS incorporation, which we parameterized as an increasing tube diameter and thus packing length due to the presence of the iBuPOSS group along the PS chain, making the polymer chain “bulky”, diluting the entanglement density. Within the terminal zone, the shear storage modulus $G'(\omega)$ featured deviation from an ideal liquid with increasing POSS content; however, no secondary rubbery plateau was observed. This indicated that the interaction between iBuPOSS groups within PS segments is not sufficient to yield “stick reptation”-like behavior. The terminal relaxation times were found to have weaker temperature dependence for random copolymer with iBuPOSS than for the pure linear PS homopolymer, the apparent activation energy increasing with increasing iBuPOSS loading. Like linear polymeric coil branches, the iBuPOSS cage plays a negative effect on the temperature-dependence of terminal relaxation time. This phenomenon seems consistent with the coupling model and with large-scale molecular dynamics simulations.

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