High Pressure Conductivity and NMR Investigation of Siloxane-Based Polymer Electrolytes

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Variable temperature and pressure conductivity and \( ^{23}Na \) nuclear magnetic resonance (NMR) measurements of several siloxane-based polymers complexed with NaCF\(_3\)SO\(_3\) are reported. The conductivity exhibits V.T.F. behavior of the type commonly observed in amorphous polymer electrolytes, and activation volumes (29.9 cm\(^3\)/mol at 313K for one sample) typical of these materials. The NMR linewidths associated with mobile Na\(^+\) ions exhibit motional narrowing above \( T_g \) in a manner similar to other Na-polymer electrolytes. The pressure dependence of the \( ^{23}Na \) linewidth is consistent with a 10-15 K/kbar increase in \( T_g \).
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ABSTRACT

Variable temperature and pressure conductivity and 23Na nuclear magnetic resonance (NMR) measurements of several siloxane-based polymers complexed with NaCF3SO3 are reported. The conductivity exhibits V.T.F. behavior of the type commonly observed in amorphous polymer electrolytes, and activation volumes (29.9 cm³/mol at 313K for one sample) typical of these materials. The NMR linewidths associated with mobile Na+ ions exhibit motional narrowing above \( T_g \) in a manner similar to other Na-polymer electrolytes. The pressure dependence of the 23Na linewidth is consistent with a 10-15 K/kbar increase in \( T_g \).
INTRODUCTION

Ionic mobility in polymer electrolytes depends critically on the ability of the host polymer to execute segmental motions. Consequently materials considered for room temperature applications require a relatively low glass transition temperature ($T_g$). A class of these materials which have recently received considerable attention are those based on copolymers of siloxanes and ethylene oxide.\textsuperscript{1,2} The former constituent provides flexibility to the polymer network which, in addition to enhancing segmental motions, yields desirable mechanical properties. The ethylene oxide (EO) groups provide a polar environment for solvation of the salt.

We have prepared several cross-linked siloxane-EO networks complexed with NaCF$_3$SO$_3$ which, in turn, have been studied by a combination of differential scanning calorimetry (DSC), complex impedance, and $^{23}$Na nuclear magnetic resonance (NMR) techniques. The effects of hydrostatic pressure on electrical conductivity and NMR linewidth have also been investigated.

EXPERIMENTAL

Two kinds of host polymer were employed. The first was based on commercial (Petrarch) dimethylsiloxane - EO copolymer (hereafter denoted as PDMS-EO), the second was based on a monophenylsiloxane-EO copolymer (PMPS-EO). Both materials were complexed with NaCF$_3$SO$_3$ with the following EO/Na ratios: 8:1 for PDMS-EO, and both 8:1 and 12:1 for PMPS-EO. The sample preparation details (including cross-linking and synthesis of the PMPS) are described elsewhere.\textsuperscript{2,3} A description of the conductivity and NMR measurement techniques is also given elsewhere.\textsuperscript{2}
RESULTS AND DISCUSSION

A DSC thermogram for the PDMS-EO complex is shown in Fig. 1. A strong glass transition with a "central" \(T_g\) of about 205K is observed. This is somewhat lower than the value obtained for 8:1 PMPS-EO (233K). The difference may be attributable, in part, to inhomogeneities in the PDMS-EO material as will be discussed later. As expected the 12:1 PMPS-EO has a lower \(T_g\) (225K) than its 8:1 counterpart, although this value is still about 4K higher than \(T_g\) for the uncomplexed host polymer.

The temperature dependence of the electrical conductivity of the PDMS-EO complex is displayed on an Arrhenius plot in Fig. 2. The curvature commonly observed in amorphous polymers is apparent. The data were first analyzed via the V.T.F. equation

\[
\sigma = A T^{-1/2} \exp \left\{ - \frac{E_a}{k(T - T_o)} \right\}
\]

(1)

A non-linear least squares fit (with adjustable parameters \(A\), \(E_a\), and \(T_o\)) was performed and the best-fit parameters are listed in Table 1. It is generally found that \(T_g - T_o\) is about 45-50K. However from Table 1 (\(T_o=187K\)), it is apparent that \(T_g - T_o\) is only 18K. The value of \(T_o\) extracted from the data in Fig. 2 is believed to be quite reliable due to the wide temperature range over which the data were taken and is further validated by the high pressure conductivity results to be discussed later. The PDMS-EO sample was not heated prior to acquiring the DSC data in Fig. 1, whereas the conductivity measurements were obtained after having annealed the sample at 100C (in a roughing vacuum to drive off residual water) for several hours. It is therefore postulated that the salt concentration in the unannealed sample may have been inhomogeneous, yielding a lower \(T_g\) than expected on the basis of the conductivity results.

The high pressure conductivity results are shown in Fig. 3. The isothermal (313K) data were fit to the equation

\[
\log_{10} \sigma = \log_{10} \sigma_0 + aP
\]

(2)
and the best-fit parameters are listed in Table 1. The activation volume can be determined from the data in Fig. 3 from the relation

$$\Delta V^\ddagger = -kT \frac{d(ln \sigma)}{dP}$$  \hspace{1cm} (3)

yielding the result $\Delta V^\ddagger = 29.9 \text{ cm}^3/\text{mol}$. This value is close to that obtained for a PDMS-EO sample complexed with NaCF$_3$COO as opposed to the triflate salt.\textsuperscript{2} The similarity in $\Delta V^\ddagger$ for the two samples (at roughly equal temperatures—that is, for equal values of $T-T_0$) implies that the $T_0$ value in Table 1 is reliable.

For completeness, the conductivity data were also analyzed according to a modified V.T.F. equation\textsuperscript{2}:

$$\sigma = A' \exp \left\{ -\left[ E_a' / k(T-T_o') \right] \right\}$$  \hspace{1cm} (4)

and the W.L.F. equation\textsuperscript{5}

$$\log_{10} \left[ \frac{\sigma(T)}{\sigma(T_3')} \right] = \frac{C_1(T-T_3)}{C_2 + (T-T_3)}$$  \hspace{1cm} (5)

The resultant parameters are also listed in Table 1.

The PMPS-EO samples were not analyzed as extensively, but were shown to have room temperature conductivities comparable to PDMS-EO.
The $^{23}\text{Na}$ NMR results are similar to those obtained for PDMS-EO complexed with NaCF$_3$COO$^2$ as well as other polymer electrolytes including PPO:NaI$^6$, PPO:NaClO$_4$$^7$, and MEEP:NaCF$_3$SO$_3$.$^8$ In particular, two distinct contributions to the NMR absorption are observed. One component has a spin-lattice relaxation time $T_1$ on the order of 1 s and a linewidth that doesn't vary appreciably with temperature. The other component has $T_1$ on the order of 1 ms and exhibits motional line narrowing above $T_g$. The latter contribution has been identified previously with mobile Na$^+$ ions in the complex, while the former component was associated with bound Na.$^{2,6,7}$ The bound Na in PPO$_8$ NaI appeared to reflect the presence of either small amounts of uncomplexed salt (but too small to be detected by powder X-ray diffraction) or clusters of ~10 or more ions.$^6$ Due to the relative complexity of the present materials, it is possible that interactions between Na$^+$ ions and the siloxane chain could lead to additional quantities of bound Na. $^{23}\text{Na}$ free induction decays (FID's) of both lineshape components for 8:1 PMPS-EO (at 273K) obtained by selective saturation and subtraction, are shown in Fig. 4. The top FID corresponds to the mobile Na$^+$ ions and is clearly motionally narrowed (in the frequency domain) with respect to the bound Na response (bottom).

The integrated intensities of the FID's in Fig. 4 can be utilized to compute the ratio of mobile to bound Na (M/B). A plot of M/B vs. T is displayed in Fig. 5., for both 8:1 and 12:1 PMPS-EO. It is clear from the data that increasing T tends to dissociate the bound Na. As expected, the 12:1 concentration exhibits a higher degree of dissociation than 8:1, in that less than 1% of the Na remains bound above room temperature in the 12:1 sample. It should be noted however that even for the 8:1 material at 212K, which is below $T_g$, about 60% of the Na is in the mobile phase. Therefore the additional 40% of carriers generated up to 360K has only a minor
effect on the conductivity which increases by 4-5 orders of magnitude over the same temperature range.

The NMR linewidth is proportional to the reciprocal of the FID time constant ($T_2^*$)$^{-1}$ which is plotted as a function of $T$ in Fig. 6 for both PMPS-EO samples. As observed for all other Na-containing polymer electrolytes,$^{2,6,7,8}$ motional narrowing above $T_g$ is clearly indicated. Another universal feature of the $^{23}$Na results is that the linewidth reaches a minimum at about 45-55K above $T_g$ and then broadens somewhat. The high temperature broadening is consistent with the extremely short ($\sim 300\mu s$) $T_1$ values measured in this region, although a microscopic model for this unusual behavior (i.e. the linewidth normally levels off in a BPP-type treatment$^9$) has not yet been formulated.

The effect of hydrostatic pressure on the linewidth ($T_2^*$)$^{-1}$ is shown in Fig. 7. Only data for PDMS-EO is included in Fig. 7. The two data symbols correspond to previously unpressurized and pressure cycled samples. There is clearly a systematic decrease in linewidth with increasing $P$. This result is in marked contrast to the pressure induced increase in linewidth observed in PPO$_8$NaI,$^6$ however in the latter material the data were obtained at 313K which corresponds to $T-T_g \approx 35K$. The PDMS-EO measurements were performed at 290K or $T-T_g \approx 85K$. From the data in Fig 6, which again is more or less representative of all Na polymer electrolytes, it is clear that $T-T_g=35K$ is within the motional narrowing region while $T-T_g=85K$ is within the motional broadening region. Therefore, the principal effect of pressure appears to be an increase in $T_g$ on the order of 10-15/kbar. This result is consistent with the conductivity pressure dependence which reflects a similar increase in $T_0$. The lack of agreement between the room temperature ambient pressure linewidths from Figures 6 and 7 is attributable partly to magnetic field inhomogeneities associated with the high pressure NMR probe and differences between the PMPS-EO (Fig 6) and PDMS-EO (Fig 7) samples.
Finally, the pressure dependence (at 290K) of M/B in PDMS-EO is displayed in Fig. 8. Comparison with the data in Fig. 5, which clearly shows that M/B decreases as the temperature is lowered toward Tg (a similar result was obtained for PDMS-EO:NaCF$_3$COO$^-$), again implies a pressure induced increase in Tg.

CONCLUSIONS

Several polymer electrolytes consisting of crosslinked siloxane-EO networks complexed with NaCF$_3$SO$_3$ have been investigated by temperature and pressure dependent conductivity and $^{23}$Na NMR. The conductivity exhibits an activation volume and V.T.F. behavior with parameters similar to those reported for PDMS-EO complexed with NaCF$_3$COO$^-$ The mobile ion concentration as probed by $^{23}$Na NMR has been shown to increase with increasing T and decrease with increasing P. The $^{23}$Na linewidth becomes motionally narrowed above Tg, and then broadens due to short T$_1$ values at higher temperatures. The pressure dependence of the linewidth is consistent with an increase in Tg on the order of 10-15K/kbar.

ACKNOWLEDGEMENTS

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REFERENCES


Table 1. Best fit VTF parameters, WLF parameters, and results of the high pressure electrical conductivity studies.

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<th>RMS Deviation</th>
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<th>$E_a$(eV)</th>
<th>$T_o$(K)</th>
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<th>RMS Deviation</th>
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<th>$E'_a$(eV)</th>
<th>$T'_o$(K)</th>
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FIGURE CAPTIONS

Figure 1. DSC plot for the PDMS-EO complex showing a central glass transition at about 205K. The data were taken at 10K/min.

Figure 2. Arrhenius plot of the electrical conductivity data for PDMS-EO. The solid line is the best fit VTF equation utilizing the parameters listed in Table 1.

Figure 3. Electrical conductivity pressure dependence at 313K and best-fit straight line.

Figure 4. Free induction decays for the mobile (top) and bound (bottom) Na⁺ in the 8:1 PMPS-EO complex at 273K. The vertical gain in the bottom figure is a factor of ten higher than in the top.

Figure 5. Mobile to bound (M/B) Na⁺ ratios of both 8:1 and 12:1 PMPS-EO complexes as a function of temperature. Circles, 8:1; Squares, 12:1.

Figure 6. Temperature dependence of ²³Na linewidth (T₂⁻¹) of mobile Na⁺ in 8:1 and 12:1 PMPS-EO complexes. Circles, 8:1; Squares, 12:1.

Figure 7. Pressure dependence of ²³Na linewidth (T₂⁻¹) of mobile Na⁺ in PDMS-EO at 290K. The squares correspond to the sample subjected to pressure for the first time; the circles represent data taken after several 0-0.2GPa cycles.

Figure 8. Mobile to bound (M/B) Na⁺ ratio of PDMS-EO as a function of pressure. Circles - pressure cycled sample; Squares - uncycled (first pressure run).
$\left( T_2^* \right)^{-1} \times 10^3 \text{ sec}^{-1}$ vs. $T (\text{K})$