Ion and polymer dynamics in polymer electrolytes PPO-LiClO$_4$: insights from NMR line-shape analysis

M. Vogel and T. Thorbrügge
Institut für Physikalische Chemie, Westfälische Wilhelms-Universität Münster, Correnstr. 30/36, 48149 Münster, Germany
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We investigate ion and polymer dynamics in polymer electrolytes PPO-LiClO$_4$ performing $^7$H and $^7$Li NMR line-shape analysis. Comparison of temperature dependent $^7$Li and $^2$H NMR spectra gives evidence for a coupling of ion and polymer dynamics. $^2$H NMR spectra for various salt concentrations reveal a strong slow down of the polymer segmental motion when the salt content is increased. The $^2$H NMR line shape further indicates that the segmental motion is governed by dynamical heterogeneities. While the width of the distribution of correlation times $G(\log \tau)$ is moderate for low and high salt content, an extremely broad distribution exists for an intermediate salt concentration 15:1 PPO-LiClO$_4$. For the latter composition, a weighed superposition of two spectral components, reflecting the fast and the slow polymer segments of the distribution, describes the $^2$H NMR line shape over a broad temperature range. Analysis of the temperature dependent relative intensity of both spectral components indicates the existence of a continuous rather than a discontinuous distribution $G(\log \tau)$. Such continuous distribution is consistent with gradual fluctuation of the local salt concentration and, hence, of the local environments of the polymer segments, whereas it is at variance with the existence of large salt-depleted and salt-rich domains featuring fast and slow polymer dynamics, respectively. Finally, for all studied mixtures PPO-LiClO$_4$, the $^2$H NMR line shape strongly depends on the echo delay in the applied echo-pulse sequence, indicating that the structural relaxation of the polymer segments involves successive rotational jumps about small angles $\gamma < 20^\circ$.

I. INTRODUCTION

Many polymers can dissolve salts so as to form mixtures supporting high ionic conductivity. These polymer electrolytes are promising candidates for applications in electrochemical devices such as solid state batteries. However, the use of these materials is limited by the achievable electric conductivities at ambient temperature and, hence, there is considerable interest to accelerate the ionic diffusion. Such efforts require a thorough understanding of the mechanisms for the ion transport. Despite considerable progress over the past decades, a general theory of ion dynamics in polymer electrolytes is still lacking.

The interactions of cations and anions with the polymer matrix are important factors governing the charge transport in polymer electrolytes. It is well established that the segmental motion of the host polymer facilitates the ionic diffusion. However, the understanding of this coupling is incomplete. For example, it was emphasized that couplings for the ionic species may differ since cation-polymer interactions are stronger than anion-polymer interactions. In addition, structural inhomogeneities can affect ionic diffusion in polymer electrolytes. Along these lines, it was stressed that the spatial distribution of the ions, in particular the formation of ionic aggregates, is a relevant factor of the charge transport in polymer electrolytes. In general, the relevance of all these aspects depends on sample composition and temperature, leading to a further complication for a comprehensive theoretical treatment.

Polymer electrolytes based on polypropylene oxide (PPO) have attracted much attention because of their high ionic conductivities together with an amorphous nature. We use $^2$H and $^7$Li NMR to investigate mixtures PPO-LiClO$_4$ with different salt concentrations or, equivalently, different ether oxygen to lithium ratios, O:Li. In the past, various experimental methods were used to characterize structure and dynamics of the PPO-LiClO$_4$ system. The electric conductivity $\sigma_{dc}$ at room temperature shows a broad maximum at intermediate O:Li ratios ranging from about 10:1 to 30:1. For such intermediate salt concentrations, differential scanning calorimetry (DSC) revealed two glass transition steps suggesting a liquid-liquid phase separation into salt-rich and salt-depleted regions. Likewise, an inhomogeneous structure in the intermediate composition range was proposed to explain results from dielectric spectroscopy (DS) which give evidence for the existence of two processes related to fast and slow polymer dynamics. Consistently, for PPO-NaCF$_3$SO$_3$, fast and slow relaxation processes were found using photon correlation spectroscopy (PCS). In contrast, other studies on polymer electrolytes PPO-LiClO$_4$ challenged the existence of salt-rich and salt-depleted microphases. Small angle neutron scattering indicated that there are no structural inhomogeneities on a length scale of several nanometers in harmony with the interpretation of data from coherent quasielastic neutron scattering (QENS). Finally, it was proposed that these apparent discrepancies can be reconciled based on a structure model resulting from reverse Monte-Carlo (RMC) simulations of diffraction data. Specifically, the RMC model of PPO-LiClO$_4$ features salt-rich and salt-depleted regions on a small length scale of about 1 nm. In addition, it reveals that the chain conformation changes as a consequence of coordination to the cations.
In view of the importance of the polymer dynamics for the ionic migration, there is considerable interest to ascertain how presence of salt affects the segmental motion in polymer electrolytes. For PPO as host polymer, it was established that addition of salt slows down the polymer dynamics. This effect was regarded as a consequence of cations acting as transient intra- or intermolecular cross links between different monomers, leading to a reduction of chain flexibility. Furthermore, neutron and light scattering studies reported that the PPO dynamics is more nonexponential when salt is present. This additional stretching was explained by either heterogeneous effects or a broadening of the distribution of correlation times \( G(\log \tau) \), or homogenous effects, i.e., intrinsic stretching. Finally, a DS study on PPO-LiClO\(_4\) reported that, over a broad concentration and temperature range, there is a linear relationship between the ionic diffusion coefficient and the relaxation rate of the polymer segments in contact with salt, suggesting that the segmental motion triggers the elementary steps of the ionic migration.

Concerning NMR work on polymer electrolytes, several studies provided insights into the ion dynamics, whereas applications on the polymer motion are rare. In detail, NMR field-gradient techniques were used to determine cationic and anionic self diffusion coefficients while measurements of the NMR spin-lattice relaxation and the line shape provided information about ion dynamics on a more local scale. On the other hand, investigation of polymer dynamics was limited to analysis of \(^1\)H spin-lattice relaxation at high temperatures. Here, we use \(^7\)Li and \(^2\)H NMR to study both the ion and the polymer dynamics in polymer electrolytes PPO-LiClO\(_4\). In particular, we exploit that \(^2\)H NMR provides straightforward access to the rotational jumps of specific polymer segments. While \(^2\)H NMR line-shape analysis is in the focus of the present contribution, future work will present \(^2\)H NMR multi-time correlation functions. Using this concerted approach, we ascertain in detail to which extent the presence of salt affects the segmental motion in polymer electrolytes.

II. EXPERIMENT

In addition to the neat PPO melt, we investigate polymer electrolytes PPO-LiClO\(_4\) with three salt concentrations, namely, 30:1, 15:1 and 6:1. PPO (M\(_n\) = 6510 g/mol) and LiClO\(_4\) were purchased from Polymer Source Inc. and Aldrich, respectively. Strictly speaking, the used polymer was poly(propylene glycol), having terminal OH groups. To prepare the polymer electrolytes, weighed amounts of PPO and LiClO\(_4\) were dissolved in acetonitrile. The resulting mixtures were stirred at 80°C until a clear homogeneous solution was obtained. Afterwards, solvent and moisture were removed on a vacuum line for several days. The resulting samples were sealed in the NMR tube. \(^1\)H and \(^13\)C NMR spectra confirmed absence of water and solvent.

A Netzsch DSC-200 instrument was used to perform DSC experiments at a heating rate of 10 K/min. We find glass transition temperatures \( T_g = 203 \pm 2 \) K for PPO and 30:1 PPO-LiClO\(_4\), consistent with results reported in the literature. For 6:1 PPO-LiClO\(_4\), we obtain \( T_g = 272 \pm 2 \) K, which is 10–20 K smaller than the literature value, suggesting some differences in the respective compositions. For 15:1 PPO-LiClO\(_4\), a very broad glass transition range extending from about 205 K to 245 K is observed. In previous studies, two glass transition temperatures \( T_g,1 \approx 208 \) K and \( T_g,2 \approx 240 \) K were found for the 16:1 composition.

The \(^2\)H (\(^7\)Li) NMR experiments were performed on Bruker DSX 400 and DSX 500 spectrometers working at Larmor frequencies \( \omega_0/2\pi \) of 61.4 MHz (155.6 MHz) and 76.8 MHz (194.4 MHz), respectively. Two Bruker probes were used to apply the radio frequency pulses resulting in 90° pulse lengths between 2.0 \( \mu s \) and 3.5 \( \mu s \), depending on the nucleus and setup. A flow of nitrogen gas controlled by a Bruker VT 3000 heading unit was utilized to adjust the sample temperature. To remove deviations between set and actual temperature within an uncertainty of \( \pm 1.5 \) K, temperature calibration was done using Pb NMR spectra of lead nitrate. Comparing \(^2\)H NMR spectra for different setups, we determined that neither the magnetic field strength nor the probe affects the results. Moreover, changing the sample position within the coil, we found no evidence for the presence of temperature gradients in the sample. The solid-echo pulse sequence was applied to measure the \(^2\)H and \(^7\)Li NMR spectra. To take into account the respective nuclear spin, the sequences (90°)\(_p\) - \( t_p \) - (90°)\(_y\) and (90°)\(_x\) - \( t_p \) - (64°)\(_y\) were used in \(^2\)H NMR and \(^7\)Li NMR, respectively. The \(^2\)H spin-lattice relaxation times were measured utilizing the saturation-recovery pulse sequence.

III. THEORY

A. Basics of \(^2\)H and \(^7\)Li NMR

In solid-state \(^2\)H NMR and \(^7\)Li NMR, the first order quadrupolar interaction, describing the interaction of the nuclear quadrupole moment with the electric field gradient (EFG) at the nuclear site, is the dominant internal interaction. It results in a frequency shift, \( \omega_Q \), which is detected in the experiment. Specifically, the quadrupolar precession frequency \( \omega_Q \) is given by

\[
\omega_Q(\theta, \phi) = \pm \frac{e}{2} \left( 3 \cos^2 \theta - 1 - \eta \sin^2 \theta \cos 2\phi \right)
\]

Here, \( \theta \) and \( \phi \) specify the orientation of the principal axes system of the EFG tensor with respect to the external static magnetic flux density \( B_0 \) and \( \eta \) is the asymmetry parameter of this tensor. The anisotropy parameter \( \delta \) is related to the quadrupolar coupling constant \( C = e^2 qQ/h \)
according to $\delta = \frac{4}{7} C$ for $^2\text{H}$ ($I = 1$) and $\delta = \frac{4}{7} C$ for $^7\text{Li}$ ($I = 3/2$). For the $^2\text{H}$ nucleus, the two signs in Eq. (1) correspond to the two allowed transitions between the three Zeeman levels. For the $^7\text{Li}$ nucleus, these signs are associated with two satellite transitions $\pm 3/2 \leftrightarrow \pm 1/2$, while the first order quadrupolar interaction does not affect the frequency of the central transition $1/2 \leftrightarrow -1/2$.

In $^2\text{H}$ NMR, the EFG tensor is intimately linked to the molecular frame. In our case of deuterated PPO, [CD$_2$-CD(CD$_3$)-O]$_n$, the monomeric unit features three deuterons in the backbone and three deuterons in the methyl group, which will be denoted as B deuterons and M deuterons, respectively. For the B deuterons, the EFG tensor is axially symmetric, i.e., $\eta = 0$, and its principal z axis points along the direction of the C-D bond. For the M deuterons, fast rotation of the methyl group leads to an averaging of the coupling tensor at the studied temperatures. The averaged tensor is also axially symmetric and its principal z axis is aligned with the three-fold symmetry axis of the methyl group. Altogether, the quadrupolar precession frequency for the B and M deuterons can be written as

$$\omega_Q(\theta) = \frac{\delta_{B,M}}{2} (3 \cos^2 \theta_{B,M} - 1) \quad (2)$$

Here, due to the motional averaging, the anisotropy parameters of the B and the M deuterons are related according to $\delta_B = 3 \delta_{M}$ [28]. For the B deuterons, $\theta_B$ specifies the angle between $\mathbf{B}_0$ and the direction of the C-D bond, whereas for the M deuterons, $\theta_M$ denotes the angle between $\mathbf{B}_0$ and the three-fold symmetry axis of the methyl group. In polymer melts, there is an isotropic distribution of molecular orientations and, hence, a broad spectrum results from the powder average at low temperatures, when molecular dynamics can be neglected. In $^2\text{H}$ NMR, this rigid-lattice spectrum has a specific line shape, called Pake spectrum. For PPO, we expect a superposition of two Pake spectra since the anisotropy parameters for the B and the M deuterons are different.

In $^7\text{Li}$ NMR, rigid-lattice spectra are comprised of two narrow non-overlapping contributions resulting from the central and the satellite transitions, respectively. The central transition, being independent of the quadrupolar interaction, yields a narrow line, while a broad spectral component results from the satellite transitions due to the action of the quadrupolar interaction together with the powder average, see Eq. (1). In polymer electrolytes, the large diversity of lithium ionic environments leads to a broad variety of EFG tensors and, hence, values of $\theta$ and $\eta$. Therefore, one expects a rather unstructured satellite component [28].

B. $^2\text{H}$ and $^7\text{Li}$ NMR line-shape analysis

Molecular dynamics with correlation times $\tau$ on the order of the inverse spectral width, i.e., on the order of microseconds, results in a collapse of the discussed $^2\text{H}$ and $^7\text{Li}$ NMR rigid-lattice spectra. In $^2\text{H}$ NMR, there is a well defined relation between the resonance frequency and the orientation of specific molecular units, see Eq. (2), and, hence, analysis of the time dependence $\omega_Q(\tau)$ provides straightforward access to the polymer segmental motion. In $^7\text{Li}$ NMR, line-narrowing of the satellite component indicates a variation of the EFG tensor at the nuclear site with time, which can result from either lithium ionic motion or, for a static ion, from a rearrangement of the neighboring polymer chains. In polymer electrolytes, both dynamic processes occur on a similar time scale and, hence, it is difficult to disentangle effects due to ion and polymer dynamics. The central line is broadened due to dipolar interactions between the magnetic moments of different nuclei, which are more important in $^7\text{Li}$ than in $^2\text{H}$ NMR. Since homonuclear and heteronuclear dipole-dipole interactions contribute, also the central line narrows as a consequence of both ion and polymer dynamics.

1. $^2\text{H}$ NMR two-phase spectra

To discuss effects of molecular dynamics on the $^2\text{H}$ NMR line shape in more detail, let us assume that there is a single anisotropy parameter $\delta$. Further, we suppose that the polymer segments show isotropic reorientation, as may be expected in the melt. If this dynamics is characterized by a single correlation time $\tau$, a rigid-lattice spectrum and a Lorentzian line will be found at low and high temperatures, respectively, where the limits of slow motion ($\tau \gg 1/\delta$) and fast motion ($\tau \ll 1/\delta$) are valid, while a crossover will occur for $\tau \approx 1/\delta$. For polymer electrolytes, a distribution of correlation times can be expected so that both fast and slow segmental reorientation may occur at the same temperature. Then, the spectrum would be comprised of a weighed superposition of a rigid-lattice spectrum and a narrow line. In general, such “two-phase spectra” can result from two limiting cases for the shape of the distribution of correlation times $G(\log \tau)$. On the one hand, it is possible that the existence of well defined microphases is reflected in a discontinuous distribution of correlation times, which is comprised of two narrow non-overlapping contributions $G_f(\log \tau_f)$ and $G_s(\log \tau_s)$ associated with fast segmental motion in salt-depleted domains ($\tau_f \ll 1/\delta$) and slow segmental motion salt-rich domains ($\tau_s \gg 1/\delta$). On the other hand, one can imagine that gradual fluctuations in the local salt concentration lead to a wide variety of local environments that manifests themselves in a very broad continuous distribution $G(\log \tau)$. Then, the fast and slow segments of this continuous distribution would yield a narrow line and a rigid-lattice spectrum, respectively, whereas contributions from segments with $\tau \approx 1/\delta$ are negligible.

In both limiting cases, the temperature dependent normalized spectral intensity $S(\omega; T)$ can be written as

$$S(\omega; T) = W(T)S_f(\omega) + [1 - W(T)]S_s(\omega) \quad (3)$$
Here, \( S_f(\omega) \) and \( S_s(\omega) \) are the normalized line shapes for the limits of fast and slow motion, respectively. The weighting factor \( W(T) \) of the former spectral pattern (0 \( \leq W \leq 1 \)) is related with the distribution of correlation times according to

\[
W(T) = \int_{-\infty}^{\log 1/\delta} G(\log \tau; T) \, d\log \tau \quad (4)
\]

Based on the temperature dependence of the weighting factor, it is possible to distinguish between both limiting cases for the shape of the distribution \( G(\log \tau) \). For a discontinuous distribution, two rapid rises of \( W(T) \) separate three temperature ranges in which the weighting factor is constant. Specifically, \( W(T) = 0 \) at low temperatures \( (\tau_f, \tau_s \gg 1/\delta) \), \( W(T) = W_f \) at intermediate temperatures \( (\tau_f \ll 1/\delta \ll \tau_s) \), and \( W(T) = 1 \) at high temperatures \( (\tau_f, \tau_s \ll 1/\delta) \). Here, \( W_f \) is the contribution of \( G_f(\log \tau_f) \) to the total distribution of correlation times. In contrast, the temperature dependent shift of a broad continuous \( G(\log \tau) \) leads to a gradual variation of \( W(T) \). For a logarithmic Gaussian distribution, characterized by a temperature independent width,

\[
G(\log \tau; T) = \frac{1}{\sqrt{2\pi}\sigma} \exp\left(-\frac{[\log \tau - \log \tau_m(T)]^2}{2\sigma^2}\right), \quad (5)
\]

one obtains after short calculation

\[
W(T) = \frac{1}{2} + \frac{1}{2} \text{erf}[x(T)], \quad (6)
\]

where \( x = (\log(1/\delta) - \log \tau_m)/(\sqrt{2}\sigma) \) and \( \text{erf}(x) \) is the error function. Therefore, given the temperature dependent shift of \( G(\log \tau) \) is known, analysis of \( W(T) \) provides information about the width of this distribution, as will be used below.

2. \(^2\text{H} \) NMR spectra for various solid-echo delays

Finally, we illustrate that, in \(^2\text{H} \) NMR, variation of the echo delay \( t_p \) in the solid-echo pulse sequence provides access to the reorientational mechanism, as was shown for various types of motion.\(^{30,31,32,33,34,35,36,37,38,39}\) For polymer melts, it was reported that the isotropic re-orientation of the polymer segments during the \( \alpha \) process results from successive rotational jumps about small angles\(^{36,37,40}\). Then, the elementary rotational jumps occur on a much shorter time scale, than the overall loss of correlation. Strictly speaking, the ratio of the jump correlation time \( \tau_j \) and the correlation time \( \tau \) depends on the jump angle \( \gamma \) according to\(^{41}\)

\[
\frac{\tau_j}{\tau} = \frac{3}{2} \sin^2 \gamma. \quad (7)
\]

Thus, for small jump angles \( \gamma < 20^\circ \), say, one may encounter a situation \( \tau_j \approx t_p \ll \tau \) at appropriate temperatures. In the literature, it was shown that, in this case, the \(^2\text{H} \) NMR line shape shows a characteristic dependence on the echo delay \( t_p \).\(^{30,31,33,35,37}\) To illustrate the effects, we calculate \(^2\text{H} \) NMR spectra from random-walk (RW) computer simulations. The methodology of these computation was described in previous work\(^{35,37}\). Here, we assume that C-D bonds perform isotropic rotational jumps with a jump angle \( \gamma = 15^\circ \), which is a typical value for the \( \alpha \) process near the glass transition.\(^{30,40}\) Further, we use a correlation time \( \tau = 1 \text{ ms} \), corresponding to a jump correlation time \( \tau_j \approx 100 \mu \text{s} \). In Fig. 1, we show \(^2\text{H} \) NMR spectra resulting from such dynamics for various values of \( t_p \). We see that the intensity in the center of the solid-echo spectrum strongly decreases with respect to the intensity of the “horns” when the echo delay is extended. The characteristics of this effect depend on the value of the correlation time \( \tau \) and the jump angle \( \gamma \) so that, in principle, the \( t_p \) dependence of the line shape enables determination of these quantities. However, we refrain from such analysis since unambiguous quantification will be difficult if distributions of correlation times and jump angles exist, as expected in our case.

IV. RESULTS

A. \(^2\text{H} \) NMR spin-lattice relaxation

First, we measure the \(^2\text{H} \) NMR spin-lattice relaxation of the mixtures PPO-LiClO\(_4\). The inset of Fig. 2 shows the recovery of the normalized longitudinal magnetization \( M_z(t) \) for PPO at \( T = 222 \text{ K} \). We see a two-step increase as a consequence of diverse spin-lattice relaxation behaviors of the different deuteron species. Specifically,
The spin-lattice relaxation times for PPO and 6:1 PPO-LiClO$_4$ lattice relaxation time of the deuterons in the backbone. Doing so, we consider our finding that the spin-lattice relaxation times. In Fig. 2, we see that the spin-lattice relaxation time decreases upon cooling, indicating $\omega_0\tau \ll 1$ for the methyl group jumps at the studied temperatures. Below 200 K, a somewhat weaker temperature dependence of $T_1^M$ suggests that the $T_1$ minimum ($\omega_0\tau \approx 1$) is approached. Above 200 K, the values $T_1^M$ for PPO and 6:1 PPO-LiClO$_4$ fall on the same line and, hence, the presence of salt does not affect the methyl group motion. Interpolation of these data with an Arrhenius law yields an activation energy $E_a = 13$ kJ/mol. Consistent with our results, QENS work on PPO-LiClO$_4$ found that an exponential correlation function, which is not affected by addition of salt, describes the methyl group jumps. However, this study reported a somewhat higher activation energy of 17 kJ/mol.

In harmony with previous $^2$H NMR work on molecular and polymeric glass formers near $T_g$, we find that the $^2$H spin-lattice relaxation of the B deuterons is nonexponential. Fitting the data for PPO at 222 K to Eq. (S), we obtain a stretching parameter $\beta = 0.48$, see Fig. 2. For the studied polymer electrolytes PPO-LiClO$_4$, including the compositions 15:1 and 30:1, which were measured at a Larmor frequency $\omega_0/(2\pi) = 76.8$ MHz, we find stretching parameters in the range $0.4 \leq \beta \leq 0.6$ at temperatures somewhat above $T_g$. Therefore, we use the $\Gamma$-function to calculate the mean spin-lattice relaxation time according to $\langle T_1^B \rangle = \langle T_1^B \rangle / \beta \Gamma(1/\beta)$. In Fig. 2 it is evident that $\langle T_1^B \rangle$ increases when the temperature is decreased, indicating $\omega_0\tau \gg 1$ for the segmental motion at the studied temperatures. Further, we see that $\langle T_1^B(T) \rangle$ is shifted to higher temperatures for 6:1 PPO-LiClO$_4$, implying that addition of salt slows down the dynamics of the B deuterons. In particular, for both samples, the curves $\langle T_1^B(T) \rangle$ show a kink near the respective glass transition temperature, as was observed for other glass-forming liquids.

In the literature, it was demonstrated that nonexponential $^2$H spin-lattice relaxation is due to a distribution of spin-lattice relaxation times, $V(T_1)$. Since $T_1$ depends on the spectral density $J(\omega)$, which is given by the Fourier transform of the correlation function, such distribution $V(T_1)$ indicates the existence of a distribution of correlation times. In general, fast relaxation processes, e.g., a $\beta$ process, can dominate the spin-lattice relaxation near $T_g$ so that a priori it is not clear whether this distribution of correlation times is associated with the $\alpha$ process. For PPO, findings in DS and PCS give no evidence for the presence of fast processes other than the methyl group motion, consistent with our $^2$H NMR results, see below. Therefore, we conclude that the nonexponential $^2$H spin-lattice relaxation of the B deuterons results from a distribution $G(\log \tau)$ for the $\alpha$ process. Thus, dynamical heterogeneities govern the structural relaxation of the host polymer in polymer electrolytes PPO-LiClO$_4$, as was shown for neat polymer melts and supercooled molecular liquids.

\section{\textsuperscript{2}H NMR line shape}

In Figs. 3 and 4 we present temperature dependent $^2$H NMR spectra of PPO-LiClO$_4$ mixtures. At low temperatures, the rigid-lattice spectra ($\tau \gg 1/\delta$) are comprised of two Pake spectra, reflecting the two deuteron species, see Sec. IIIA. Fitting the line shapes, we obtain anisotropy parameters $\delta_B \approx 118-121$ kHz and $\delta_M \approx 38-40$ kHz. For both deuteron species, the lower values of the specified line-width ranges are found for higher salt contents. At high temperatures, we observe a narrow line, indicative of fast ($\tau \ll 1/\delta$) isotropic reorientation of the polymer segments. When we compare the results for the different compositions, it is evident that the line shape changes at higher temperatures for higher salt concentrations, indicating that the presence of salt slows down the segmental motion. The temperatures, at which $\tau \approx 1/\delta$, are about $400 \pm 75$ K. This is near the glass transition temperature of neat PPO, as determined from DSC measurements. Therefore, we conclude that the $^{13}C$ NMR-derived glass transition temperature, which is about $310 \pm 10$ K, is due to the longitudinal $^{13}C$ spin-lattice relaxation time of the methyl group. This result is consistent with a recent $^{13}C$ NMR study of neat PPO-LiBF$_4$, in which the $^{13}C$ spin-lattice relaxation time of the methyl group is found to be $T_1^M > T_1^0$.

\section{Conclusion}

In summary, we present a comprehensive study of the spin-lattice relaxation of the monomeric backbone deuterons in PPO and its mixtures with LiClO$_4$. Our results show that addition of salt slows down the dynamics of both the B and M deuterons. In particular, for both samples, the curves $\langle T_1^B(T) \rangle$ show a kink near the respective glass transition temperature, as was observed for other glass-forming liquids. The $^2$H NMR spectra reflect the presence of a slow $\alpha$ process, consistent with the existence of a distribution of correlation times. However, the $\beta$ process is not observed at temperatures above $T_g$. This suggests that the $\beta$ process is faster than the $\alpha$ process and is not resolved in our experiments. In addition, we find that the $^2$H NMR line shape changes at higher temperatures for higher salt concentrations, indicating that the presence of salt slows down the segmental motion. The temperatures, at which $\tau \approx 1/\delta$, are about $400 \pm 75$ K. This is near the glass transition temperature of neat PPO, as determined from DSC measurements.
FIG. 3: Temperature dependent $^2$H NMR solid-echo spectra of PPO ($t_p = 20 \mu s$).

FIG. 4: Temperature dependent $^2$H NMR solid-echo spectra of 30:1 PPO-LiClO$_4$ ($t_p = 20 \mu s$).

FIG. 5: Temperature dependent $^2$H NMR solid-echo spectra of 15:1 PPO-LiClO$_4$ ($t_p = 20 \mu s$).

FIG. 6: Temperature dependent $^2$H NMR solid-echo spectra of 6:1 PPO-LiClO$_4$ ($t_p = 20 \mu s$).
85 K higher for 6:1 PPO-LiClO$_4$ than for PPO, showing that the effects are strong, cf. Figs. 4 and 6.

Closer inspection of the $^2$H NMR line-shape changes at intermediate temperatures reveals interesting aspects of the polymer segmental motion in the PPO-LiClO$_4$ mixtures. For PPO, there is a more or less continuous collapse of the rigid-lattice spectrum when the temperature is increased, see Fig. 4, consistent with previous results on glass-forming liquids. Revisiting Figs. 4 and 6, we see some deviations from a continuous line narrowing for the low (30:1) and the high (6:1) salt concentrations. In narrow temperature ranges ($\Delta T < 10$ K), the $^2$H NMR line shape resembles a two-phase spectrum, see Sec. III 14, although the effect is not pronounced. In Fig. 6, it is obvious that the two-phase signature is much better resolved for the intermediate 15:1 salt concentration. Over a broad temperature range, the $^2$H NMR spectrum can be described as a weighed superposition of a narrow line and a rigid-lattice spectrum, indicating the coexistence of fast ($\tau < 1/\delta$) and slow ($\tau \gg 1/\delta$) polymer segmental motion. Hence, for all studied polymer electrolytes PPO-LiClO$_4$, the $^2$H NMR line shape indicates that the segmental motion exhibits dynamical heterogeneities. However, the degree of the nonuniformity of the motion depends on the composition. While the dynamical heterogeneities are very prominent for intermediate salt concentrations, the effect is weaker for both low and high salt content.

For a more detailed analysis of the two-phase spectra for 15:1 PPO-LiClO$_4$, we determine the weighting factor $W(T)$ characterizing the relative contribution of the narrow line, cf. Eq. 4. In Fig. 7, we see that $W(T)$ increases gradually from 0 to 1 when the temperature is increased. As was discussed in Sec. III 14, such behavior indicates the existence of a continuous rather than a discontinuous distribution of correlation times $G(\log \tau)$. Specifically, the latter scenario would manifest itself in a discontinuous variation of the weighting factor.

To estimate the width of the distribution $G(\log \tau)$ for 15:1 PPO-LiClO$_4$, we use input from our $^2$H NMR stimulated-echo study on this material. There, we find that the segmental motion exhibits nonexponential correlation functions, which can be described by a Kohlrausch function, $\exp[-(t/\tau)^\beta]$. The small stretching parameter $\beta \approx 0.2$ shows no systematic variation with temperature, consistent with a temperature independent width of the distribution $G(\log \tau)$. In the temperature range of the two-phase signature, the mean correlation time follows an Arrhenius law, $\log_{10}(\tau(T)) = \log_{10}(\tau_0 + E_a/(T \ln 10))$, where $\log_{10}(\tau_0)/s = -49.3$ and $E_a = 28100$ K. Based on these results, we assume that the segmental motion exhibits a logarithmic Gaussian distribution $G(\log_{10} \tau)$ that shifts with an activation energy $E_a = 28100$ K, where the width parameter $\sigma_{10}$ is temperature independent. For various values of $\sigma_{10}$, we then calculate $W(T)$ according to Eq. 4. In doing so, we also vary the prefactor $\log_{10} \tau_0$ since, for broad logarithmic Gaussian distributions, the mean correlation time $\langle \tau \rangle$ is longer than $\tau_m$. Further, we use $\log_{10} 1/\delta = -5.65$, which is the average of the values resulting from $\delta_M$ and $\delta_B$, respectively.

In Fig. 7, we see that the experimental data is well reproduced for a width parameter $\sigma_{10} = 2.4$ and a prefactor $\log_{10} \tau_0 = -50.3$. This width parameter $\sigma_{10}$ corresponds to a full width at half maximum of more than five orders of magnitude and, hence, the polymer segmental motion in 15:1 PPO-LiClO$_4$ is extremely heterogeneous, see inset of Fig. 7. For this intermediate salt concentration, the two-phase signature is observed in the temperature range 240 K $\lesssim T \lesssim 310$ K. The lower and the higher end of this range correspond roughly to the temperatures where the line-shape changes for PPO and 6:1 PPO-LiClO$_4$ are found, see Figs. 8 and 7, suggesting that the broad distribution of correlation times for the 15:1 composition results from large fluctuations in the local salt concentration. Although the present results rule out the presence of a discontinuous distribution $G(\log \tau)$ for the 15:1 composition, it is possible that the broad continuous distribution is comprised of two distributions, which are associated with polymer dynamics in salt-rich and salt-depleted regions, provided these distributions overlap in large parts.

### C. $^7$Li NMR line shape

Next, we analyze $^7$Li NMR spectra to investigate the lithium ionic dynamics in PPO-LiClO$_4$. For all studied polymer electrolytes, we find that the $^7$Li NMR rigid-lattice spectra at low temperatures are comprised of a

![Fig. 7: Weighting factor $W(T)$ quantifying the relative intensity of the narrow line in the two-phase spectra of 15:1 PPO-LiClO$_4$, see Fig. 6. The data indicates the existence of a broad continuous distribution $G(\log \tau)$. The solid line is an expectation for $W(T)$ resulting from the assumption of a logarithmic Gaussian distribution $G(\log_{10} \tau)$ characterized by a width parameter $\sigma_{10} = 2.4$, see text for details. The inset shows this distribution for $T = 272$ K. The shaded area marks the part of the distribution contributing to the narrow line.](image-url)
FIG. 8: (Color online) Full width at half height of the central transition in $^7$Li NMR spectra of 30:1 PPO-LiClO$_4$, 15:1 PPO-LiClO$_4$ and 6:1 PPO-LiClO$_4$. For the sake of comparison, the data was scaled by the value for the lowest temperatures studied.

FIG. 9: Partially relaxed $^2$H NMR solid-echo spectra of PPO for the indicated echo-delays $t_p$ and $T = 222$ K. The delay between the saturation of the longitudinal magnetization and the start of the solid-echo pulse sequence was 3 ms.

FIG. 10: Partially relaxed $^2$H NMR solid-echo spectra of 15:1 PPO-LiClO$_4$ for the indicated echo-delays $t_p$ and $T = 244$ K. The delay between the saturation of the longitudinal magnetization and the start of the solid-echo pulse sequence was 3 ms.

D. $^2$H NMR solid-echo spectra for various echo-delays

To study the mechanisms for the polymer segmental motion in the PPO-LiClO$_4$ systems, we analyze the dependence of the $^2$H NMR solid-echo spectra on the echo delay $t_p$. In our case, the superposition of two Pake spectra complicates such analysis. Therefore, we exploit that the spin-lattice relaxation of the M deuterons is faster than that of the B deuterons, see Fig. 2, and single out the contribution of the former species in partially relaxed temperatures.
2H NMR spectra. In detail, after saturation of the longitudinal magnetization, we do not wait for its complete recovery, but start the solid-echo pulse sequence at times \( t \approx T_1^M \ll T_1^B \), when the longitudinal magnetization is dominated by the contribution of the M deuterons. In Figs. 9-11 we show partially relaxed 2H NMR solid-echo spectra of PPO-LiClO_4 systems. Evidently, the narrow Pake spectrum of the M deuterons dominates the line shape, whereas contributions from the B deuterons are suppressed in large parts. Only for the 6:1 composition, suppression of the latter contribution is somewhat less effective because of a smaller difference of the spin-lattice relaxation times \( T_1^M \) and \( T_1^P \), see Fig. 4.

The dependence of the partially relaxed 2H NMR spectra on the solid-echo delay \( t_p \) is displayed for PPO at \( T = 222 \) K in Fig. 9 and for 15:1 PPO-LiClO_4 at \( T = 244 \) K in Fig. 10. In both cases, a rigid-lattice spectrum is observed for \( t_p = 20 \) μs, while the relative intensity in the center of the spectrum decreases when the echo delay is extended. These line-shape changes are a fingerprint of small angle rotational jumps, see Fig. 4. Since angular rotations are required until the memory of the initial orientation is lost completely. Thus, for such type of motion one encounters the situation \( \tau_j \approx t_p \ll \tau \) at appropriate temperatures. Then, the elementary jumps lead to line-shape changes for large \( t_p \), i.e., high angular resolution, while a rigid-lattice spectrum results for short \( t_p \), since angular displacements, which are sufficiently large to as to lead to effects for poor spatial resolution, are not achieved on the time scale of the experiment.

According to this argumentation, the \( t_p \) dependence of the 2H NMR solid-echo spectra in Figs. 9 and 10 indicates that \( \tau_j \approx t_p \ll \tau \) holds for the reorientation of the three-fold methyl group axes in PPO and 15:1 PPO-LiClO_4 at the studied temperatures, consistent with the results Figs. 3 and 5. This time scale separation \( \tau_j \ll \tau \) is indicative of reorientation resulting from small angle jumps, see Eq. 7. For PPO, this reorientational mechanism is consistent with results from previous work on polymer melts.\(^{29,31,32}\) Our findings for 15:1 PPO-LiClO_4 together with that for 6:1 PPO-LiClO_4, see Fig. 11, show that the isotropic reorientation of the polymer segments in polymer electrolytes also involves successive small-angle jumps. Hence, on a qualitative level, we find no evidence that addition of salt leads to a change of the mechanism for the reorientation of the polymer segments. In a future publication, we will show that analysis of 2H NMR two-time correlation functions confirms this conclusion on a quantitative level.

Figure 11 displays temperature dependent 2H NMR spectra of PPO for a large solid-echo delay \( t_p = 200 \) μs. We see that the intensity in the center of the spectrum is diminished for \( T = 222 \) K, cf. also Fig. 8 while this effect becomes less prominent when the temperature is decreased, until the line shape is independent of \( t_p \) for \( T \leq 212 \) K. Since the line shape for \( t_p = 200 \) μs is sensitive to small-angle motion, these results indicate that cooling slows down the elementary rotational jumps so that, near \( T_g \), any polymer segmental motion is frozen in on the \( \mu s \) time scale. We further emphasize that the observed temperature dependence of the spectra for a large echo delay together with a \( t_p \) independent line shape at low temperatures rule out that the observed effects are due to experimental imperfections, e.g., due to an inaccurate time origin for the Fourier transformation of the time signal. Moreover, these observations exclude spin diffusion, i.e., a transfer of magnetization as a consequence of flip-flop processes of the spins, as possible origin of the \( t_p \) dependent line shape. Since spin diffusion is not directly related to molecular dynamics, it shows, if at all, a weak temperature dependence and, hence, it cannot account for the pronounced effects in Fig. 11.

Many molecular and polymeric glass formers show a secondary relaxation (\( \beta \) process) that is intrinsic to the glassy state.\(^{33}\) In the literature,\(^{36-38}\) it was demonstrated that, at \( T \lesssim T_g \), the spatially restricted motion associated with the \( \beta \) process leads to a \( t_p \) dependence of the 2H NMR solid-echo spectrum, which resembles that resulting from the elementary steps of the \( \alpha \) process at \( T > T_g \), see Figs. 9-11. In previous NMR work, it was suggested that a \( \beta \) process exists in polymer electrolytes.\(^{33}\) Here, the 2H NMR solid-echo spectra give no evidence that PPO-LiClO_4 mixtures elec-
trolleys exhibit a $\beta$ process. In particular, we determined that the $^2\text{H}$ NMR spectra of 15:1 PPO-LiClO$_4$ are independent of the solid-echo delay in the temperature range $160 \, \text{K} \leq T \leq T_g$. Consistently, no $\beta$ process was observed in DS on polymer electrolytes PPO-LiClO$_4$.\textsuperscript{8}

V. CONCLUSION AND SUMMARY

Previous work on the polymer segmental motion in polymer electrolytes PPO-LiClO$_4$ reported nonexponential two-time correlation functions and broad dielectric loss spectra.\textsuperscript{11,15} Two limiting scenarios can explain such nonexponential relaxation. In the heterogeneous limit, all particles obey exponential correlation functions, however, a distribution of correlation times $G(\log \tau)$ exists. In the homogeneous limit, all particles obey the same correlation function, which is, however, intrinsically non-exponential. For disordered materials, heterogeneous and homogeneous contributions may coexist. Measurements of two-time correlation functions and, hence, the previous approaches do not allow one to discriminate between both scenarios and to quantify the relevance of homogeneous and heterogeneous contributions.\textsuperscript{45} Here, we demonstrated that $^2\text{H}$ NMR spin-lattice relaxation and line shape consistently indicate that pronounced dynamical heterogeneities govern the polymer segmental motion in polymer electrolytes PPO-LiClO$_4$. Specifically, at appropriate temperatures, the existence of a distribution of correlation times $G(\log \tau)$ manifests itself in a nonexponential $^2\text{H}$ spin-lattice relaxation and in a $^2\text{H}$ NMR two-phase line shape, i.e., the spectrum is comprised of a weighed superposition of a narrow line and a rigid-lattice spectrum. While the former spectral component is due to the fast polymer segments of the rate distribution, the latter is due to the slow segments. Hence, in both the $^2\text{H}$ NMR experiments, the existence of dynamical heterogeneities is demonstrated without invoking any model assumptions. However, the present data do not provide information whether or not homogeneous dynamics yields an additional contribution to the nonexponential relaxation. In future work, we will show that quantification of homogeneous and heterogeneous contributions is possible based on analysis of $^2\text{H}$ NMR three-time correlation functions, as was demonstrated for neat molecular and polymeric glass formers.\textsuperscript{50,52}

Analysis of $^2\text{H}$ NMR spin-lattice relaxation and line shape gave further evidence for the existence of a continuous rather than a discontinuous distribution of correlation times for all studied polymer electrolytes PPO-LiClO$_4$. Specifically, such shape of the rate distribution is consistent with a $^2\text{H}$ spin-lattice relaxation following a stretched exponential, as was observed in the experiment. Moreover, in an analysis of the two-phase spectra, a continuous distribution $G(\log \tau)$ was indicated by a continuous variation of the weighting factor $W(T)$, quantifying the relative contribution of the narrow line. However, the width of the distribution of correlation times strongly depends on the salt concentration. While the width is moderate for both low and high salt concentrations, an extremely broad distribution exists for the intermediate salt content 15:1 PPO-LiClO$_4$, leading to the observation of two-phase spectra over a broad temperature range. For 15:1 PPO-LiClO$_4$, we used information about the temperature dependent shift of $G(\log \tau)$ from our concomitant $^2\text{H}$ NMR stimulated-echo study to estimate the width of this distribution from the temperature behavior of the weighting factor $W(T)$. An estimate of more than five orders of magnitude for the full width at half maximum shows that this polymer electrolyte exhibits very pronounced dynamical heterogeneities.

Since it is established that presence of salt slows down the polymer dynamics, it is reasonable to assume that the shape of the distribution of correlation times for the polymer segmental motion reflects features of the salt distribution in polymer electrolytes PPO-LiClO$_4$. Thus, the width of $G(\log \tau)$ implies strong fluctuations in the local salt concentration for the intermediate 15:1 composition, while the structural diversity is reduced for both high and low salt content. In the literature, the possibility of liquid-liquid phase separation into salt-rich and salt-depleted regions was controversially discussed for intermediate salt concentrations.\textsuperscript{8,10,11,12,13,14,15,16} Given the rate of the polymer segmental motion is related to the local salt concentration, the existence of well-defined microphases should manifest itself in the existence of a bimodal distribution $G(\log \tau)$. Our $^2\text{H}$ NMR results for 15:1 PPO-LiClO$_4$ indicate the existence of a continuous $G(\log \tau)$ so that they are at variance with a bimodal distribution of correlation times and, hence, of local salt concentrations unless the contributions from salt-rich and salt-depleted regions overlap in large parts. On the other hand, they are in harmony with short ranged fluctuations of the local salt concentration. In this case, one expects a large variety of local structural motifs leading to a broad continuous $G(\log \tau)$. For example, our $^2\text{H}$ NMR findings for the intermediate salt content are consistent with a RMC model of 16:1 PPO-LiClO$_4$, which features salt-rich and salt-depleted regions on a short length scale of about 10 Å corresponding to about two monomeric units. When we consider the cooperativity of molecular dynamics near the glass transition, the chain connectivity and the effect that salt migration renders the spatial salt distribution time depend on the time scale of the polymer segmental motion, it is highly likely that a continuous distribution of correlation times $G(\log \tau)$ results from this RMC model, consistent with our $^2\text{H}$ NMR results.

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