Understanding Lithium-Ion Dynamics in Single-Ion and Salt-in-Polymer Perfluoropolyethers and Polyethyleneglycol Electrolytes Using Solid-State NMR

Chibueze V. Amanchukwu,*# Anna B. Gunnarsdóttir,# Snehashis Choudhury, Tamsin L. Newlove, Pieter C. M. M. Magusin, Zhenan Bao,* and Clare P. Grey*#

ABSTRACT: Lithium metal batteries with high energy densities can enable a revolution in energy storage and accelerate shifts in electric transportation and electricity generation. However, several morphological and electro-chemo-mechanical challenges impede their development. Solid-state electrolytes such as those based on polymers show great promise in replacing liquid electrolytes in lithium metal batteries. Polyether-based polymer electrolytes are the most investigated but are plagued by low room-temperature ionic conductivity and poor oxidative stability. Hence, there is great need for the development and understanding of ion transport in new classes of polymer electrolytes. Perfluoropolyether (PFPE)-based electrolytes have shown improved oxidative stability, but little is understood about their lithium solvation and transport mechanism. In this work, we use multinuclear solid-state magic angle spinning (MAS) nuclear magnetic resonance (NMR) spectroscopy to investigate the lithium cation environment and mobility in crosslinked single-ion and salt-in-polymer PFPE electrolytes and compare it directly to that of well-known polyethylene glycol (PEG) electrolytes. We show that the interaction of the lithium cation with the polymer backbone is weaker in PFPE systems compared to PEG, likely resulting in stronger ion pairing in the PFPE systems. Line shape analyses show lower lithium mobility in PFPE electrolytes despite lower activation energies being derived from spin-lattice relaxation ($T_1$) measurements as compared to those for the PEG systems. The rapid relaxation is instead ascribed to the local fluctuations caused by polymer backbone mobility. By studying different modes of ion binding (single-ion vs salt-in-polymer), we show that differences across polymer backbones (PFPE vs PEG) have a greater effect on mobility than differences in ion binding modes within each polymer class (especially when single-ion conducting site density is not high). Our ability to use MAS NMR to study polymer electrolytes in their native state opens new opportunities to develop and understand novel polymer or hybrid solid-state electrolytes for next-generation lithium metal batteries.

INTRODUCTION

Developing batteries with higher energy densities is vital for electric transportation and a renewable-powered electric grid.1 Replacing the graphite anode (in current lithium-ion batteries) with lithium metal can at least double the overall battery gravimetric energy density.2 Hence, there has been fervent research to develop rechargeable lithium metal batteries. However, prior efforts to commercialize rechargeable lithium metal batteries failed primarily because of the volatile, flammable liquid electrolytes and the inability to prevent lithium dendrite growth upon metal deposition.3 Designing solid-state lithium metal batteries may alleviate these safety and morphological challenges while maintaining high energy density.3–5 Solid-state electrolytes (SSEs) should support high ion conductivity ($>10^{-3}$ S/cm), be mechanically flexible and thin (<25 μm), and have a wide electrochemical stability window (0–4.5 V vs Li/Li$^+$).3 Currently, inorganic SSEs such as phosphates (e.g., LATP), oxides (LLZO),6 and sulfides7 have been studied heavily.8,4 Although inorganics such as sulfides have ion conductivities on the order of $10^{-2}$ to $10^{-3}$ S/cm,8 no single class of inorganic electrolytes meets all the requirements.3,10

Polymer electrolytes are easy to process as a stand-alone SSE, as part of an inorganic-polymer hybrid electrolyte11,12 or as a binder for the cathode composite.13,14 Furthermore, they can be mechanically flexible and fabricated in thin form factors. Polyethylene oxide or glycol (PEO/PEG) is the archetypal polymer electrolyte and has been studied for over four...
decades. However, PEO still suffers from low-room-temperature ionic conductivity ($\sim 10^{-5}$ S/cm) and poor oxidative stability (<4 V vs Li/Li$^{+}$), preventing its use either as a stand-alone SSE or as part of a cathode composite where voltages cutoffs typically exceed 4 V Li ($e.g.$, with LiCoO$_2$ and LiNi$_x$Mn$_y$Co$_z$O$_2$). We note that recent studies have indicated that PEO may have higher electrochemical stability.

The oxidative challenges facing PEO also plague small ether molecules. New polymer backbone chemistries that do not require the ether group for ion conduction are needed. Polycarbonates such as polyethylene carbonate have been explored, but they still suffer from low-room-temperature ionic conductivity and poor oxidative stability.

Recently, we synthesized a new class of fluorinated-ether small-molecule electrolytes that show high ionic conductivity and high oxidative stability. Previous work has shown that crosslinked PFPE can support ion transport, albeit still with low conductivities ($\sim 10^{-5}$ S/cm). Also, previous solution-state works done by some of us and Balsara and co-workers have shown higher lithium transference number in the perfluoroether backbone, indicating weaker lithium coordination of the polymer backbone compared to that of the traditional PEG. In addition, Lee et al. used density functional theory to show that PFPE has a higher oxidation potential and a higher electrochemical stability window compared to PEO, which was also observed experimentally in small-molecule perfluoroether electrolytes. However, little is known about the lithium-ion solvation structure, mobility, and transport mechanism in solid PFPE electrolytes.

Here, we study two classes of polymer electrolytes with different ion transport mechanisms; “single-ion”/polyelectrolyte and “salt-in-polymer” (see Figure 1a). Salt-in-polymer electrolytes involve a lithium salt dissolved in a polymer and usually suffer from low lithium transference numbers (with  

![Figure 1. Schematic of polymer electrolytes. (a) Illustration of the difference between a polyelectrolyte/single-ion and a “salt-in-polymer” polymer electrolyte. (b) Schematic of a crosslinked single-ion PFPE. S5 and S15 correspond to the percentage of TFSI (5 or 15 wt %) attached to the polymer chain (circled in b), and the “S” indicates it is a single-ion polymer. When the PFPE repeat units in the dashed boxes are replaced by the PEG repeat unit as showcased by the arrows, the single-ion PEG is obtained (PEG-S15). (c) Chemical structures for PFPE-DMA, PEG-DA, and LiTFSI. These are used to fabricate “salt-in-polymer” by dissolving 20 wt % LiTFSI salt in the liquid monomer (either PFPE-DMA or PEG-DA) and photo-crosslinking. Chemical structure of PFPE-DMA was obtained from Devaux et al.](https://doi.org/10.1021/acs.macromol.2c02160)
higher mobility of the anion compared to the cation under an electric field.\textsuperscript{13} To improve the transference number, single-ion/polyelectrolytes have been developed, which immobilize the salt anion within the polymer backbone, making the lithium ion the only mobile species, leading to a lithium transference number close to unity.\textsuperscript{33–35} A third class of polymer electrolytes (not studied here), “polymer-in-salt” electrolytes, has been proposed as a strategy to improve transference number where the proposed ion conduction mechanism includes ion hopping between ion clusters, i.e., decoupling of the ion conduction to the polymer segmental motion.\textsuperscript{36,37}

In this work, we fabricated single-ion and salt-in-polymer PFPE, with the tethered TFSI anion and LiTFSI salt dissolved (Figure 1), respectively. Magic angle spinning (MAS) nuclear magnetic resonance (NMR) was used to study the polymer environment, the lithium-ion solvation structure, and activation barrier for ion transport.\textsuperscript{38} MAS NMR provides high chemical shift resolution for these solid polymers and enables an understanding of ion solvation and mobility in their natural solid state, without the need to dissolve the polymers in solvents.\textsuperscript{39–41} Lithium ($^7$Li) NMR was used to study the cation, while fluorine ($^{19}$F) and proton ($^1$H) NMR was used to characterize the anion and polymer backbone behavior.

We perform equivalent analyses on single-ion PEG and salt-in-polymer PEG to provide insight into the differences between the two classes of polymer backbone. We find that lithium interacts more weakly with the PFPE backbone as compared to PEG, and the lithium-ion mobility in PFPE is less strongly coupled to the polymer mobility as compared to PEG. The lithium-ion mobility is strongly temperature-dependent in PEG, whereas only minor effects are observed in PFPE. Finally, the activation energy for lithium mobility was estimated by spin-lattice relaxation ($T_1$) measurements and reveals a lower energy barrier in the salt-in-polymer PFPE system compared to PEG. The low activation energy derived from relaxation measurements is likely due to the rapid motion of the polymer backbone inducing fluctuations in the local magnetic field of the cation. Here, the use of multinuclear MAS NMR yields mechanistic insights into the ion transport behavior in a relatively new class of PFPE electrolytes and will enable the development of next-generation ion-conducting polymer electrolytes.

## RESULTS AND DISCUSSION

### Synthesis and Physicochemical Characterization

The “single-ion” and the “salt-in-polymer” electrolytes were synthesized according to previously published procedures.\textsuperscript{29,34} The electrolytes were crosslinked to obtain mechanically compliant, stand-alone films. All the precursors and monomers are either liquid or soluble in tetrahydrofuran (THF), with crosslinking leading to insoluble solids (visual confirmation). Figure 1 shows the structures of the polymer precursors and a schematic for the subsequently crosslinked structure. For the single-ion conductor, the TFSI anion was tethered to the backbone using the thiol functionality (Figure 1b). Three different concentrations of the single-ion PFPE were studied: S0 (0 wt % TFSI tethered), S5 (5 wt % TFSI tethered), and S15 (15 wt % TFSI tethered). Note, these are the nominal values because not all the added TFSI will crosslink within the chain (see Experimental Details for a more detailed discussion). By changing the PFPE to a PEG, we can make direct comparisons between a single-ion PFPE (PFPE-S0, S5, S15) and a single-ion PEG (PEG-S15).

The “salt-in-polymer” electrolytes were prepared by completely dissolving 20 wt % LiTFSI in the monomers (PEG-diacylate and PFPE-methacrylate) and photo-crosslinking to obtain solid polymer electrolytes (Table S1). These polymers are conventional in the sense that both cation and anion are expected to be mobile in the crosslinked matrix. By comparing the salt-in-polymer PFPE to PEG, we can draw insights from previously explored crosslinked PEG systems.

MAS NMR and Fourier transform infrared (FTIR) spectroscopies were used to confirm the PFPE polymer structures (Figures S1–S4). FTIR data in Figure S4 show...
the incorporation of the TFSI in the single-ion polymers with a peak at 1625 cm\(^{-1}\), which is present in both PFPE-S5 and PFPE-S15 but is absent in PFPE-S0. In addition, the sets of peaks from 610 to 540 cm\(^{-1}\), associated with a SO\(_2\) scissoring vibration from TFSI, are present in both PFPE-S5 and S15 but not in PFPE-S0.\(^{45}\) For the salt-in-polymer, these sets of peaks are also observed, indicating the presence of LiTFSI. The combination of visual (liquid to solid), FTIR (Figure S4), and solid-state NMR provided evidence of crosslinking in these polymer electrolytes and engendered probing of thermal behavior and ion mobility.

The thermal properties of the polymer electrolytes were studied using thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC). The TGA data in Figure S5 show that the polymer electrolytes are stable until at least 250 °C. The DSC data in Figure S6 show that 20 wt % LiTFSI in PFPE shows no glass transition in the temperature range studied (−50 to 120 °C), consistent with results reported by Devaux et al. for the crosslinked PFPE electrolyte.\(^{\text{35}}\) Previous work by Hu et al. showed two glass-transition temperatures for crosslinked PFPE: \(\sim-116\) °C assigned to the fluorinated backbone and \(\sim0\) °C assigned to the hydrocarbon domains of the PFPE. Based on these results, we assign the weak glass transition for PFPE-S0 and PFPE-S15 at \(\sim-18\) °C (Figure S6) to the methacrylate crosslinking sites.\(^{39,30}\) Meanwhile, there are no glass or melting transitions for PEG-S15 and 20 wt % LiTFSI in PEG in the temperature range studied (−50 to 120 °C). Work by Schönhoff et al.\(^{39}\) showed a \(T_g\) of \(-45\) °C for crosslinked PEG with no salt and a \(T_g\) of \(-32\) °C with 30 mol % LiTFSI. It is possible that the \(-50\) °C DSC temperature cutoff (\(T_g\) could be lower) or different sample preparation techniques contribute to the absence of a \(T_g\) in our crosslinked PEG samples.

**Polymer-Backbone Environments and Dynamics.** The temperature dependence of the polymer backbone was studied using \(^1\)H NMR and \(^19\)F MAS NMR spectroscopy. Within amorphous polymer electrolytes, polymer segmental motion is required for ion transport and thus the ion mobility is often coupled to mobility of the backbone.\(^{17,44}\) Hence, comparing the environments within the different polymer classes is vital.

Figure 2a shows the \(^19\)F MAS NMR spectra of the solid PFPE-S15 polymer as a function of temperature. The main resonances correspond to (−CF\(_2\)-CF\(_2\)-O−) units in the polymer backbone around −89 to −91 ppm and the (−CF\(_2\)-O−) units around −52 to −56 ppm. Table S2 summarizes the chemical shifts for the different groups of the polymer backbone.\(^{45,46}\) Integration of the resonances for (−CF\(_2\)-CF\(_2\)-O−) and (−CF\(_2\)-O−) (see Figure S1) results in a ratio of \(x/y \approx 7/3\), consistent with previous reports.\(^{46}\)

Interestingly, the fluorinated PFPE polymer groups give rise to sharp resonances, whereas the resonance of the TFSI anion (expected at around −81 ppm)\(^{46}\) is not directly observed (we know an anion must be present since the \(^7\)Li MAS NMR in Figure S2 shows the presence of lithium). Integration of the \(^19\)F spectra shows that the TFSI peak is buried underneath the strong signal of the PFPE backbone (Table S2), indicating a mobile fluorinated polymer backbone and a more constrained TFSI anion (Figure S2). The absence of the TFSI peak could also be due to broadening from \(^19\)F−\(^19\)F homonuclear dipolar coupling and the rapid motion in this system affecting the ability of MAS to effectively remove the \(^19\)F−\(^19\)F homonuclear coupling (see discussion for \(^7\)Li below).\(^{46}\)

The resonances of the −CF\(_2\)-CF\(_2\)-O− and −CF\(_2\)-O− units do not shift with salt addition (shown in Figure S7), suggesting that the polymer backbone is weakly affected by the addition of lithium ions. Moreover, the backbone resonances shift only slightly with temperature and the line width remains constant within the temperature regime studied (Figure 2a,c). The lack of sensitivity to temperature and the observation of extremely sharp resonances in a system with strongly coupled \(^19\)F spin systems (CF\(_2\) groups) at the relatively slow MAS frequency of 8 kHz indicates that the PFPE polymer chains are already in the NMR fast motion regime and highly mobile even at \(-20\) °C. The terminal CF\(_2\)-CH\(_3\) groups for PFPE S15 show some

---

**Figure 3.** Lithium-ion local environment as a function of temperature. \(^7\)Li MAS NMR spectra as a function of temperature for (a) PFPE-S15, (b) 20 wt % LiTFSI in PFPE, (c) PEG-S15, and (d) 20 wt % LiTFSI in PEG. The dashed box in (a−d) corresponds to the inset. The chemical shift values indicated in the inset correspond to the maximum peak at 30 °C. The arrow indicates increasing temperature. Temperature range from −20 to 40 °C with a 10 °C increment. The temperature color scales in (a,c) are the same for (b,d). Spinning sidebands are denoted by asterisks (*). MAS spinning speed: 8 kHz.
temperature sensitivity both by \(^1\)H NMR (Figure S8) and the \(^{19}\)F NMR signal around −80 ppm (Figure 2a, shown in more detail for both PFPE S0 and S15 in Figure S9), further supporting that the observed \(T_g\) is related to the methacrylate crosslinking sites rather than the fluorinated backbone.

Figure 2b,d shows the \(^1\)H NMR spectra for the PEG polymer electrolytes, where the resonances are broad at −20 °C and narrow significantly with temperature, indicating increasing mobility of the backbone. In addition, the \(^1\)H peak for the salt-in-polymer 20 wt % LiTFSI in PEG shifts downfield (3.77 ppm) compared to PEG S15 (3.65 ppm) at 30 °C. The differences between the polymer backbone (PFPE vs PEG) as a function of temperature significantly outweigh the differences between ion coordination environment (single-ion/polyelectrolyte vs salt-in-polymer) within the same polymer class.

**Lithium-Ion Environment and Dynamics.** Figure 3 shows the \(^7\)Li MAS NMR spectra, which were used to study the lithium-ion environment in the single-ion and salt-in-polymer electrolytes as a function of temperature. The lithium cations will coordinate to the negatively charged TFSI anions or to the tethered anion backbone of the single-ion polymer, and thus, the coordination environments and mobility are likely to differ. \(^7\)Li is a quadrupolar nucleus \((I = 3/2)\) and here all polymer electrolytes show a single \(^7\)Li peak that consists of a narrow component associated with the \((-1/2 \leftrightarrow 1/2)\) central transition and a broader component or a quadrupolar component associated with the satellite transitions \((\pm 3/2 \leftrightarrow \pm 1/2)\) that result in spinning sidebands under MAS (the line shape shown in more detail in Figures S10 and S11).\(^{47,48}\) For the PEG polymers (Figure 3c,d), a broad line shape is observed at the lower temperatures, which sharpens significantly as the temperature is increased, consistent with the behavior of the polymer backbone. For the PFPE polymers, there are minimal changes to the central peak with temperature, mirroring the behavior of the backbone as observed in Figure 2. The intensities and widths of the \(^7\)Li spinning sidebands do, however, change noticeably with temperature, indicating an onset of motion.

The line width of the central peak was first quantified to probe lithium mobility, where line narrowing is expected with increasing temperature due to motional averaging of dipolar and (to a lesser extent for \(^7\)Li) quadrupolar interactions.\(^{40,41}\) Figure 4a shows a decrease of around 30% in line width for both PFPE-S5 and S15 as the temperature increases between −20 and 40 °C compared to only 19% in the salt-in-polymer PFPE, suggesting poorer ion mobility in the salt-in-polymer. In contrast, a sharp decrease in line width of around 85% is observed for both PEG-S15 and 20 wt % LiTFSI in PEG. The significant changes in line width for PEG-S15 may not be attributed to thermal transitions alone because the DSC data in Figure S6 show no thermal transitions between −20 and 40 °C and are consistent with higher \(^7\)Li mobility.

Motional effects are also observed via the spinning sideband manifold (Figure 3) where a loss of intensity is due to motional averaging of anisotropic interactions. For both PEG electrolytes, the satellite transitions collapse into the central transition with increased temperature, which we mainly attribute to averaging of the quadrupolar interaction. However, in contrast to PEG-S15, a noticeable broadening and then sharpening of the spinning sidebands manifold are seen for 20 wt % LiTFSI PEG (see Figure S11).

To gain more insight into motional effects of the quadrupolar interaction, deconvolutions of the NMR spectra were performed using the dmfit software package.\(^{49}\) The spinning sideband manifold is captured by the broad component, attributed to the satellite transitions. The line widths of the central and satellite peaks from the spectral fittings and the extracted values of the quadrupolar coupling constant \(Q_{\text{ef}}\) are summarized in Figures S11 and S12 over the measured temperature range.

In 20 wt % LiTFSI PEG, an initial broadening is seen for the satellite line width, the satellites then narrowing at temperatures above 0 °C. This broadening is a clear indication of slow motion.\(^{50−53}\) In the limit where the MAS frequency is greater than the quadrupolar coupling, maximum broadening occurs for motion on the order of the spinning frequency (here 8 kHz); when the MAS frequency is less than the static quadrupolar coupling constant, the hopping rate at which maximum broadening occurs is higher than the MAS frequency and now also depends on the quadrupolar frequency. However, for the quadrupolar coupling constants derived for this system (see the Supporting Information), the maximum broadening is seen for hop rates of a similar order of magnitude. Thus, for 20 wt % LiTFSI PEG, the motional broadening indicates motion occurring on the \(\sim 10^{-4}\) s timescale.\(^{53}\) In contrast, PEG-S15 shows negligible spinning sideband intensity, indicating that the quadrupolar interaction is effectively being averaged over the whole temperature range, also shown with continuous decrease in line width of the broad component that is present under the sharper central transition peak. This indicates higher lithium-ion mobility in PEG-S15 that is already faster than the MAS frequency (8 kHz) at −20 °C.

For PFPE-S15, the spinning sideband intensity decreases with temperature and is mirrored in an increase in line width of the broad component (see Figure S11). In comparison, the cations in 20 wt % LiTFSI in PFPE show no onset of motion, with little differences seen in the line widths over the whole temperature range (Figure S11). In conclusion, line shape analysis with temperature suggests that the lithium ions in the
single-ion polymer electrolytes have greater mobility as compared to the salt-in-polymers within each polymer class.

The influence of salt concentration within the single-ion PFPE electrolytes was also studied, and Figure 4b shows that as the salt concentration is increased from PFPE-S5 to PFPE-S15, the $^{7}\text{Li}$ chemical shift remains constant at $-0.46$ ppm, indicating little change in the lithium-ion environment. An upfield shift (negative shift) to $-0.77$ ppm is observed in 20 wt % LiTFSI in PFPE, indicating different chemical environments to the single polymer, possibly due to more ion pairing. The chemical shift reported for pure LiTFSI salt (using LiCl at 0 ppm as a $^{7}\text{Li}$ reference) is $-1.2$ ppm. As Figure 4d shows, the chemical shift is further upfield for both PEG-S15 and 20 wt % LiTFSI in PEG (overlap at $-1.29$ ppm), consistent with significant ether–Li binding and/or interactions with the TFSI units due to ion pairing.

The downfield $^{7}\text{Li}$ shift observed in the PFPE indicates that it cannot be solely bound to multiple TFSI units (as in the salt LiTFSI). However, the lack of an observable change of the polymer backbone in the $^{19}\text{F}$ spectrum between the single-ion and salt-in-polymer systems suggests that the backbone interacts only very weakly, if at all, with the Li$^{+}$ ions (Figure S7). The fluoroether backbone is known for its low polarity and has seen significant use as chemically resistant oils. The higher-frequency shift of Li$^{+}$ ions in PFPE compared to PEG is thus ascribed to a lower coordination environment (small number of ligands) for Li$^{+}$ or possibly the interaction with other ligands such as the methacrylate groups.

$^{19}\text{F}$ MAS NMR of the TFSI anion in PEG (Figure S9) shows a temperature dependence similar to that of the lithium ion in Figure 3 with significant decrease in line width. In 20 wt % LiTFSI PEG, the TFSI appears upfield ($-79.7$ ppm) compared to PEG S15 ($-78.6$ ppm) at 30 °C, which may indicate stronger ion pair formation for the salt-in-polymer. In conclusion, for both PEG and PFPE electrolytes, the temperature dependence of the lithium and anion peak mirrors the polymer behavior, indicating that segmental motion of the polymer backbone and the ion mobility are correlated, which will have implications for ion transport.

**Relaxation Behavior and Activation Energy.** The dependence of the spin-lattice ($T_1$) relaxation with temperature is a sensitive probe of motion on the order of the Larmor frequency ($\sim 10^{-9}$ s). $^7\text{Li}$ can undergo relaxation via both dipole–dipole and quadrupolar mechanisms where the quadrupolar interaction is expected to be dominant in these polymer systems. The relaxation times for $^7\text{Li}$ are shown in Figure 5 and are shorter in the PEG polymer system (Figures S14 and S15) are four times longer than in the PEG system at room temperature, consistent with reports of lower conductivity in PFPE electrolytes. Others have reported that the Arrhenius preexponential factor—a proxy for number of charge carriers—is 3 orders of magnitude lower in liquid PFPE than in PEG.

**Consequences for Lithium-Ion Transport.** In PEG-based polymer electrolytes, lithium ions coordinate with the polymer, and segmental motion of the polymer enables lithium-ion transport. Here, the PFPE backbone is highly mobile at all temperatures studied (on the basis of the $^{19}\text{F}$ NMR in Figure 2). Furthermore, no significant change in the $^{19}\text{F}$ chemical shifts in the PFPE backbone are seen with salt addition (Figure S7), indicating that the lithium ion interacts weakly with the PFPE polymer backbone. Unlike the ether oxygen in PEG, the fluorine atoms in PFPE will withdraw electron density from the fluoroether oxygens, limiting their ability to coordinate to the lithium ions strongly. Note that PFPE electrolytes have both fluoroether oxygen atoms as well as the methacrylate group, but the PFPE alone (without any hydrogenated end groups) does not dissolve salt as we have seen through our own experiments. In contrast, the ether group in PEG has been reported to solvate and support ion transport regardless of the end group selection. Thus, much stronger Li$^{+}$–TFSI$^{-}$ ion pairing likely occurs in the PFPE systems, the Li ions binding more strongly to the negatively

**Figure 5.** Ionic motion and relaxation behavior. (a) $^7\text{Li}$ spin-lattice relaxation times ($T_1$) as a function of temperature. Lines shown are linear fits. (b) Activation energies obtained from the Arrhenius fit to the $T_1$ data in (a). The labeling color scheme in (b) is the same as in (a).
charged TFSI groups and the TFSI groups acting as local traps. The weak interactions between the PFPE and Li\(^+\) ions contribute to the lower conductivity of the PFPE- vs PEG-based systems—despite the high mobility of the PFPE backbone.

Low Li\(^+\) mobility is observed for PFPE-S15, even lower mobility being seen (on the basis of the \(^{7}\)Li sideband manifolds) for the salt-in-polymer. The low Li mobility is consistent with long-range measurements of Li transport (conduction), which are lower for the PFPE vs PEG systems.\(^{40}\)

Despite this, higher activation energy barriers were calculated for the single-ion PFPE compared to salt-in-polymer PFPE (Figure 5). The activation barriers determined by the NMR relaxation measurements are unlikely, however, to be directly related to long-range translational motion. In support of this, the \(^{7}\)Li line shape analysis suggested that Li mobility was in the kHz timescale (i.e., correlation times of the order of \(10^{-4}\) s), while the relaxation analysis corresponds to correlation times of the order of \(10^{-8}\) s (Figure S14). Thus, the dominant relaxation mechanisms must arise from short-range dipolar and quadrupolar interactions, the high mobility of the PFPE polymer chains likely driving the relaxation of the Li cation but not the Li mobility. Even though Li binding to the chains is weak, the overall mobility of the backbone will result in high-frequency fluctuations of the whole system.

With NMR sensitive to localized motion on short time scales, efforts were made to compare the activation energies obtained from \(T_1\) measurements to pulsed field gradient (PFG)-NMR diffusion measurements, which probe long-range motion. However, due to the extremely short \(T_2\) relaxation time (on the order of 0.1 ms), the diffusion coefficient of the cation could not be measured. However, our results compare well with reported conductivity measurements; the line shape analysis shows that the ion motion is faster in the PEG systems that corroborates the higher ionic conductivity data reported for PEG systems.

The interest in PFPE systems is due to their increased oxidative stability as predicted by DFT.\(^{18}\) However, our work shows that the PFPE backbone suffers from poor ionic solvation, which is responsible for the lower ionic conductivity observed in crosslinked PFPE.\(^{29}\) An approach to improve ionic solvation in fluorinated electrolytes was recently explored in a new class of small-molecule electrolytes that eschew the fluoroether oxygen within the polymer backbone and another approach that adds ether groups to the PFPE “end-group” to get combined improved ion solvation/transport coupled to desired high electrochemical stability.\(^{29}\) These approaches may lead to improved polymer—ion interactions within the PFPE polymer class and enable a next generation of highly conductive PFPE electrolytes.

### CONCLUSIONS

The lack of ionically conducting, electrochemically, and mechanically stable solid-state electrolytes has hampered the development of lithium metal batteries. Ether-based polymer electrolytes have been heavily studied but suffer from low-room-temperature ionic conductivity and poor oxidative stability. In contrast, PFPE-based electrolytes are purported to have higher oxidative stability, although little is understood about lithium ion solvation and transport in solid-state PFPE electrolytes. In this work, we use \(^1\)H, \(^{7}\)Li, and \(^{19}\)F MAS NMR to study PFPE- and PEG-based polymer electrolytes and show that while the polymer backbone mobility of PFPE is extremely rapid and even liquid-like at room temperature, only slow Li motion (on the kHz timescale) is observed. By contrast, the motion of the PEG backbone increases gradually with temperature in the relevant battery temperature range of −20 to 40 °C, this motion tracking an increase in Li\(^+\) mobility. The \(^{7}\)Li and \(^{19}\)F chemical shift data show that the lithium ion interacts weakly with PFPE compared with PEG. More ion pairing likely exists in the PFPE systems than in the PEG systems where the Li\(^+\) ions are more effectively dissociated from the negative counter anions (or charge) by binding to the ether groups.

The activation energy barrier obtained using NMR relaxation measurements mirrors that obtained in the literature from conductivity measurements and is lower for the PFPE electrolyte compared to PEG. However, the rapid relaxation times observed by NMR in the PFPE system are associated instead with the mobility of the PFPE chains as opposed to the long-range transport of the Li\(^+\) ions. Despite the stronger ether—Li binding in crosslinked PEG, quantification of the \(^{7}\)Li NMR line shapes shows that ion motion is faster in PEG compared to the PFPE electrolytes. By contrasting single-ion/polyelectrolytes with “salt-in-polymer,” our work indicates that ion mobility increases in single-ion/polyelectrolytes for both crosslinked PEG and PFPE but that the identity of the polymer backbone (PEG vs PFPE) has a stronger influence on ion solvation and mobility. Using MAS NMR to study these polymer electrolytes in their native solid state will allow us to translate these insights into the design of next-generation non-ether polymer electrolytes for lithium-based batteries.

### EXPERIMENTAL DETAILS

**Materials and Synthesis.** Poly(ethylene glycol) diacrylate (PEGDA; \(M_w = 700\)) was obtained from Sigma-Aldrich. Poly(etheretherketone) dimeracrylate (PFPE DMA; \(M_w = 1500\)) is a bifunctional urethane-methacrylate and was obtained from Cornerstone company (Fluorolink MD700). Pentaacryltiol tetraakis(3-mercaptopropanionate) (“Tetrathiol”) and 2,2-dimethoxy-2-phenylacetophenone (DMPA) were obtained from Sigma-Aldrich. Tetrahydrofuran was obtained from Fisher Scientific. LiTFSI was obtained from Solvionic.

“Single-Ion” Conducting Polymers. Lithium 4-styrenesulphonyl-(trifluoromethanesulphonyl) imide (STFSI Li) was synthesized according to a procedure previously reported.\(^{34,35}\) To obtain PFPE S5, S15, and S50, STFSI Li was mixed with the tetra-thiol crosslinker, PFPE, and the DMPA initiator.

PFPE-S0 [0 wt % STFSI, 5 wt % of DMPA initiator with respect to (wt) PFPE + STFSI].

PFPE-S5 [5 wt % STFSI wt PFPE DMA, 1:1 mole ratio (−SH/STFSI + DMA); 5 wt % DMPA].

Example for PFPE-S5: 1 g of PFPE DMA; 0.05 g of STFSI; 0.182 g of tetraeth; 0.052 g of DMPA.

PFPE-S15 [15 wt % STFSI wt PFPE DMA, 1:1 mole ratio (−SH/STFSI + DMA); 5 wt % DMPA].

PFPE S50 [50 wt % STFSI wt PFPE DMA, 1:1 mole ratio (−SH/STFSI + DMA); 5 wt % DMPA].

PEG-S15 [15 wt % STFSI wt PEGDA, 5 wt % DMPA initiator wt PEGDA + STFSI].

These components were added to a vial, and tetrahydrofuran (THF) was added and stirred to dissolve the components. However, for the mixtures containing STFSI, not all STFSI dissolved. Hence, the STFSI ratios reported are nominal values, and the actual STFSI content in the crosslinked network will be lower. The solution was then exposed to ultraviolet light (handheld, 365 nm) for crosslinking. For PFPE-S0 and S5, crosslinking happened even before exposure to UV. For PFPE S50, a 400 W lamp was used for 35 min to induce crosslinking. Once crosslinking occurred for all the polymer samples,
they were washed copiously (at least three times) with excess THF. The polymers were then placed at a 60 °C oven overnight to dry.

“Salt-in-Polymer” Conducting Polymers. PFPE DMA and PEGDA were vacuum-dried at 100 °C before transfer to an Argon glovebox. In an Argon-filled glovebox (MBrAun, O₂ < 0.1 ppm, H₂O < 0.1 ppm), the following measurements were made. In a 20 mL vial, 3 g of PFPE DMA was added, followed by 0.75 g of LiTFSI (to obtain 20 wt %). The vial was stirred at 90 °C to fully dissolve the salt. Then, 30 mg of DMPA was added and stirred until dissolution. As soon as DMPA dissolved, the contents were poured into a Teflon dish and irradiated with a handheld UV lamp (365 nm) for 3 min. The film could be easily peeled off from the dish.

For 20 wt % LiTFSI in PEGDA, a similar procedure was followed. Here, 2 g of PEGDA, 0.5 g of LiTFSI, and 20 mg of DMPA were used. The contents were poured into a glass dish and crosslinked with a handheld UV lamp (365 nm) for 3 min.

Characterization (FTIR, TGA, and DSC). FTIR characterization was performed in an Argon-filled glovebox (MBrAun, O₂ < 0.1 ppm, H₂O < 0.1 ppm) with an Agilent Cary 630 spectrometer. TGA characterization was performed using a Mettler Toledo/DSC 2. A 10 °C/min scan range was used from 30 to 500 °C under a nitrogen environment. DSC characterization was performed using a PerkinElmer DSC 4000. DSC procedure: (1) hold at 30 °C for 2 min. (2) Heat at 20 °C/min from 30 to 120 °C for all the polymers to reset the polymer thermal history. (3) Hold at 120 °C for 2 min. (4) Cool at 15 °C/min from 120 to −50 °C. (5) Hold at −50 °C for 10 min. This was to make sure that the sample actually reached −50 °C since the cooling unit was not great. (6) Heat at 15 °C/min from −50 to 120 °C. This last heating cycle was reported in the main manuscript.

Solid-State NMR Characterization. Experiments were conducted on a Bruker Avance II spectrometer equipped with a 4 mm H/F XY DVT and 2.5 mm X/F/H MAS DVT probe. Variable temperature (cooling) was maintained using an externally connected BCU II unit. Samples were held for at least 15 min at each temperature before measurements were made. Sample preparation was done in an Argon-filled glovebox. The crosslinked polymer samples were cut into fine powders or tiny chunks and packed into either a 2.5 or a 4 mm rotor. The 4 mm rotors were covered using Kel-F caps, which limited our accessible temperature range (−20 to 40 °C). Adamantane was used as an external reference for 1H (1.87 ppm) and 13C (higher frequency peak at 38.6 ppm), 1 M NaF in H₂O was used as an external fluorine reference (−120 ppm), and 1 M LiCl in H₂O was used as the external 7Li reference (0 ppm). 90° pulses were measured, and reference measurements were made before each NMR session. 19F spectra were collected using a spin-synchronized Hahn-echo pulse sequence to remove the background from the probe. Spin-lattice (T₁) relaxation measurements were performed using a saturation-recovery sequence. Spectral fittings of the central peak and spectral linewidths were performed using a dmfit software package.49

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsmacromol.2c02160.

Extended characterization data such as FTIR, DSC, TGA, T₁ relaxation, and ssNMR analyses of the PFPE and PEG polymer electrolytes (PDF)

AUTHOR INFORMATION

Corresponding Authors

Chibueze V. Amanchukwu — Department of Chemical Engineering, Stanford University, Stanford, California 94305, United States; Department of Chemistry, University of Cambridge, Cambridge CB2 1EW, U.K.; Pritzker School of Molecular Engineering, University of Chicago, Chicago, Illinois 60637, United States; orcid.org/0000-0002-6573-1213; Email: chibueze@uchicago.edu

Zhenan Bao — Department of Chemical Engineering, Stanford University, Stanford, California 94305, United States; orcid.org/0000-0002-0972-1715; Email: zbao@stanford.edu

Clare P. Grey — Department of Chemistry, University of Cambridge, Cambridge CB2 1EW, U.K.; orcid.org/0000-0001-5572-192X; Email: cpg27@cam.ac.uk

AUTHORS

Anna B. Gunnarsdottir — Department of Chemistry, University of Cambridge, Cambridge CB2 1EW, U.K.; Faculty of Industrial Engineering, Mechanical Engineering and Computer Science, University of Iceland, Reykjavik 107, Iceland; orcid.org/0000-0001-6593-788X

Snehashis Choudhury — Department of Chemical Engineering, Stanford University, Stanford, California 94305, United States

Tamsin L. Newlove — Department of Chemistry, University of Cambridge, Cambridge CB2 1EW, U.K.

Pieter C. M. Magusin — Department of Chemistry, University of Cambridge, Cambridge CB2 1EW, U.K.; Present Address: Institute for Life Sciences and Chemistry, HU University of Applied Sciences Utrecht, 3501 AA Utrecht, Netherlands; orcid.org/0000-0003-1167-3764

Complete contact information is available at: https://pubs.acs.org/10.1021/acsmacromol.2c02160

Author Contributions
C.V.A. and A.B.G. contributed equally to this work.

Notes
The authors declare no competing financial interest.

ACKNOWLEDGMENTS

C.V.A. acknowledges financial support from the TomKat Center Postdoctoral Fellowship in Sustainable Energy at Stanford and a Visiting Fellowship from Corpus Christi College at the University of Cambridge. S.C. and Z.B. acknowledge support from the Assistant Secretary for Energy Efficiency and Renewable Energy, Office of Vehicle Technologies of the U.S. Department of Energy, under the Battery Materials Research (BMR) program. A.B.G. acknowledges the support from the Royal Society (RP/R1/180147) and EPSRC-EP/M009521/1. We thank Professor Eugene Terentjev for providing us access to his DSC instrument.

REFERENCES


(6) Marbella, L. E.; Zekoll, S.; Kasemchian, J.; Emge, S. P.; Bruce, P. G.; Grey, C. P. 7Li NMR Chemical Shift Imaging To Detect...


