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Exploring the Ion Solvation Environments in Solid-State Polymer Electrolytes through Free-Energy Sampling

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 ABSTRACT: The success of polyethylene oxide (PEO) in solid-state polymer electrolytes for lithium-ion batteries is well established. Recently, in order to understand this success and to explore possible alternatives, we studied polyacetal electrolytes to deepen the understanding of the effect of the local chemical structure on ion transport. Advanced molecular dynamics techniques using newly developed, tailored interaction potentials have helped elucidate the various coordination environments of ions in these systems. In particular, the competition between cation−anion pairing and coordination by the polymer has been explored using free- energy sampling (metadynamics). At equivalent reduced temperatures, with respect to the polymer-specific glass-transition temperature, two-dimensional free-energy plots reveal the existence of multiple coordination environments for the lithium (Li) ions in these systems and their relative stabilities. Furthermore, we observe that the Li-ion movement in PEO follows a serial, stepwise pathway when moving from one coordination state to another, whereas this happens in a more continuous and concerted fashion in 14 a polyacetal such as poly $(1,3$ -dioxalane) $[P(EO-MO)]$. The implication is that interconversion between coordination states of the Li ions may be easier in P(EO-MO). However, the overarching observation from our free-energy analysis is that Li-ion coordination is dominated by the polymer (in either case) and contact-ion pairs are rare. We rationalize the observed higher increase in glass-17 transition temperature (T_g) with salt loading in polyacetals as due to intermolecular Li-ion coordination involving multiple polymer 18 chains, rather than just one chain for PEO-based electrolytes. This interchain coupling in the polyacetals, resulting in the higher T_{ci} works against any gains due to variations in Li-ion coordination that might enhance transport processes over PEO. Further research is required to overcome the interdependence between local coordination and macroscopic properties to compete with PEO electrolytes at the same absolute working temperature.

■ INTRODUCTION

 Lithium-ion batteries are high-energy-density power sources that find uses in the automobile industry for electric cars and 5 various portable devices like smartphones and laptops. $1,2$ There are strong motivations for developing solid-state electrolytes, with solid polymer electrolytes (SPEs) being prime candidates, since they have strong mechanical stability and avoid safety concerns associated with the combination of flammable liquid electrolytes and cell shorting due to lithium dendrite growth in 1 the current technology.^{3−[9](#page-10-0)} Since the early work of Wright^{[10,11](#page-10-0)} 32 and Armand,^{[12](#page-10-0)} poly(ethylene oxide) (PEO) has been studied extensively and has remained one of the more promising 34 SPEs.^{[6](#page-10-0),[13](#page-10-0),[14](#page-10-0)} Polymer electrolytes usually behave as hosts for the ions in the system, with the ions having the ability to move

through the free volume of the polymers assisted by the $_{36}$ segmental motion, with reasonable conductivity possible above the glass-transition temperature. ⁶ Therefore, effective dissolu- $_{38}$ tion of the cations and a low glass-transition temperature are key to good ionic properties in these systems.^{[13](#page-10-0)} Unfortunately, $_{40}$ slow ionic conductivities and low transference numbers in ⁴¹

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⁴² SPEs at battery working temperatures have significantly ⁴³ hindered their performance for practical applications.[6](#page-10-0),[13](#page-10-0)[−][18](#page-10-0)

 Standard molecular dynamics (MD) protocols have been used extensively to explore the ion coordination environments and try to understand the transport mechanism in PEO/ lithium bis(trifluoromethanesulfonyl)imide (LiTFSI) sys-48 tems.^{[19](#page-10-0)-[28](#page-10-0)} Borodin and Smith^{[21](#page-10-0)} used a quantum chemistry- based many-body polarizable force field to look at the local ion coordination environment, ion aggregation, and various contributions to cation transport. They found in their simulations that Li-ion motion is a combination of motion along the polymer chains, along with the segmental motion of PEO, and ion hopping from one segment to another at 55 different timescales. Motivated by experimental findings,² Diddens and Heuer confirmed that at lower salt concen- trations, all Li ions are coordinated by PEO chains and that the improved Li-ion transport with addition of ionic liquids was 59 due to the plasticizing effect of the ionic liquid molecules. 22 60 Recently, Molinari and co-workers^{[24](#page-10-0)} showed that asymmetrical negatively charged clusters at high LiTFSI concentrations are a possible reason for experimental negative transference numbers. Standard MD protocols are however plagued by drawbacks of system size effects, starting configuration dependence, and limited sampling as compared to residence times in coordination environments that can affect the accuracy of the various ensemble averages calculated in spite of the long times that are usually used in these simulations. 69 Recent work by Baskin and Prendergast^{[30,31](#page-10-0)} illustrated the use 70 of free-energy sampling methods like umbrella sampling^{[32,33](#page-10-0)} 71 and metadynamics $34,35$ $34,35$ $34,35$ in the framework of both classical and ab initio MD to overcome some of these deficiencies and gain more rigorous insight into ion solvation environments.

74 In our previous work, $36,37$ $36,37$ $36,37$ the design strategy that was employed was to maintain a high oxygen to carbon ratio in the backbone of the polymer to provide sufficient well-connected solvation sites for the Li cation by incorporating methylene oxide (O−CH2 or MO) repeat units into the polymer 79 backbone along with ethylene oxide (O–CH₂–CH₂ or EO). A series of such polyacetals with varying ratios of EO and MO 81 in the repeat units, $P(nEO-mMO)$, were synthesized³⁶ and 82 systematically studied electrochemically 37 and through pulsed field gradient nuclear magnetic resonance spectroscopy 84 measurements and MD simulations.^{[36](#page-10-0)} The polymers studied previously were P(EO-2MO), P(EO-MO), P(2EO-MO), P(3EO-MO), and P(4EO-MO) compared to PEO with LiTFSI salt. Polyacetal−LiTFSI SPEs were studied at similar 88 reduced temperatures $(T - T_g)$ as the glass-transition temperatures in the presence of LiTFSI varied significantly. It was observed that at a given reduced temperature, the cation self-diffusion coefficient was higher in each polyacetal derivative as compared to PEO. It was also observed that P(EO-MO) and P(EO-2MO) exhibit significantly lower anion self-diffusion coefficients as compared to the polyacetals in the series. Preliminary MD simulations revealed that the MO repeat units containing polymers (i.e., polyacetals) have additional ether oxygens close to the cation coordination environment in a second coordination shell, which distort the primary coordination environment and possibly improve Li- ion transport at a given reduced temperature in these systems. In the present work, we use MD simulations and free-energy sampling (metadynamics) to gather further atomic level insight into the different coordination environments that exist in these systems, their relative stabilities, and how they might affect

transport mechanisms. Our goal is to understand the ¹⁰⁵ underlying structural differences in the lithium coordination ¹⁰⁶ environments in these poly(ether-acetal) systems and how ¹⁰⁷ interchangeable they are. For simplicity, we compare PEO to ¹⁰⁸ P(EO-MO), the most efficacious polymer of the polyacetals ¹⁰⁹ studied previously. 37 We observe from metadynamics simu- 110 lations the existence of multiple stable coordination environ- ¹¹¹ ments with differing numbers of coordinating oxygen atoms ¹¹² from the polymer and TFSI. The importance of running ¹¹³ advanced free-energy calculations as compared to a regular ¹¹⁴ MD protocol is most notable upon examination of the relative ¹¹⁵ stabilities of various coordination environments, which might ¹¹⁶ easily form irreversibly during tractable MD simulations, ¹¹⁷ thereby overestimating their statistical significance. Further- ¹¹⁸ more, PEO exhibits a serial, stepwise minimum free-energy ¹¹⁹ pathway to convert from one coordination state to another, ¹²⁰ while P(EO-MO) has a more direct, continuous pathway with ¹²¹ the transition occurring in a more concerted fashion. However, ¹²² we know that the polyacetals experience a larger increase in $T_{\rm g}$ 123 upon addition of LiTFSI salt, which works against any ¹²⁴ enhanced transport mechanisms at the same absolute temper- ¹²⁵ ature. Our simulations indicate that this is due to enhanced ¹²⁶ interchain connectivity through Li-ion coordination in the ¹²⁷ polyacetals. Overall, these insights provide a fundamental ¹²⁸ understanding of the structural differences between local ¹²⁹ coordination of Li ions in both systems and indicate that ¹³⁰ reducing the T_g increase upon salt loading should be key to 131 defining new chemistries for SPEs and improving performance ¹³² for Li-ion battery applications. Fraction battery applications. 133
■ METHODS 134

The Methods section is divided into three parts. In the first part, we ¹³⁵ describe the MD protocol that we use to prepare the polymer systems 136 with ions at various concentrations. In the second part, we delve into 137 the functional form of the force field and the methods used to ¹³⁸ optimize the existing generalized AMBER force field (GAFF) ¹³⁹ interaction potential^{[38](#page-10-0)} to predict the angles, bonds, dihedrals, and 140 bulk properties, such as density, more reliably over a wide range of 141 compositions. Finally, we elaborate on the free-energy calculations 142 performed using these newly optimized potentials to reveal the 143 prevalent coordination environments of the ions and to explore the 144 mechanism of contact-ion-pair formation. 145

MD Protocol. MD simulations were performed using the large- 146 scale atomic/molecular massively parallel simulator (LAMMPS) 147 code^{[39](#page-11-0)} using an interaction potential with the GAFF^{[38](#page-10-0)} functional form 148

$$
E_{\text{tot}} = \sum_{\text{bonds}} K_{\text{r}} (r - r_0)^2 + \sum_{\text{angles}} K_{\theta} (\theta - \theta_0)^2
$$

+
$$
\sum_{\text{dihedrals}} K_{\phi} [1 + d \cos(n\phi)]
$$

+
$$
\sum_{i < j} \left(4\epsilon_{ij} \left[\frac{\sigma_{ij}^{12}}{r_{ij}^{12}} - \frac{\sigma_{ij}^{6}}{r_{ij}^{6}} \right] + \frac{q_i q_j}{4\pi \epsilon_0 r_{ij}} \right)
$$
(1)

where K_r , K_{θ} , and K_{ϕ} are force constants; r_0 , θ_0 , and ϕ_0 are the 150 equilibrium bond length, bond angle, and dihedral angle, respectively; 151 n is the multiplicity; the σ_{ij} and ϵ_{ij} parameters characterize a Lennard- 152 Jones (LJ) non-bonded interaction; and q_i is the partial charge on the 153 atoms. The total energy E_{tot} is hence expressed as a summation of 154 harmonic bond, angle, and dihedral terms that are primarily intrachain 155 in nature and LJ and electrostatic interactions that largely determine 156 interchain interactions (although they are relevant if the polymer 157 chain is highly curved or forms loops). The partial charges were 158 calculated using the restrained electrostatic potential $(RESP)$ model^{[40](#page-11-0)} 159 after ab initio optimization of isolated molecules with the B3LYP 160 hybrid density functional and the 6-311++gss basis set using 161

 TeraChem.[41](#page-11-0)[−][43](#page-11-0) The charges on the ions were scaled by 0.8 based on 163 ab initio results and suggestions from the previous literature $44,45$ $44,45$ $44,45$ to emulate polarization effects. Geometric mixing rules were used for 165 unlike LJ parameters $(\sigma_{ij}, \epsilon_{ij})$. Intramolecular pairwise LJ and coulombic interactions separated by one and two bonds were set to 0, while those separated by three were set to 0.8 and 0.5, respectively. The short-range cutoff was set to 13 Å, while the electrostatic interactions were cut off at 14 Å. The time step used for all our MD simulations was 1 fs unless otherwise specified.

171 A multi-step annealing protocol was used to create the MD f1 172 structures used in this work (see the top panel of Figure 1). A single

Figure 1. Top: Schematic for ion−polymer system preparation; bottom: saturation of density with annealing cycles.

 chain of the polymer with a chain length of 15−20 monomer units was first created and relaxed in the NVT ensemble [where NVT 175 implies a constant number of particles (N) , volume (V) , and 176 temperature (T)] at 300 K. A total of 30 such relaxed chains were 177 placed in a large box at a very low density (0.1–0.25 g/cm³) to avoid any overlap, and the correct amount of Li and TFSI ions for the specific composition were randomly distributed. Salt concentrations 180 studied in this work ranged from $r = 0$ to 0.16, where $r = \begin{bmatrix} Li \end{bmatrix} / \begin{bmatrix} O \end{bmatrix}$ (the ratio of Li ions to polymer oxygens), with the upper bound defined by involvement of all available O atoms in Li-ion coordination, assuming an average of 6 coordinating O atoms per Li ion. Each annealing cycle consisted of first heating the system from 300 to 900 K in steps of 60 K, relaxing at each step for 20 ps at a nominal heating rate of 3 K/ps, relaxing the system for 100 ps at that temperature and slowly cooling down the system again in steps of 60 K at a nominal cooling rate of 3 K/ps in the NVT ensemble, followed by a room-temperature relaxation in the NPT ensemble [where NPT 190 implies a constant number of particles (N) , pressure (P) , and 191 temperature (T)] at 300 K at atmospheric pressure (1 bar) for 50 ps, allowing for the density to change. The variation of density with each annealing cycle is shown in the bottom panel of Figure 1. Based on these data, we chose to perform 20 annealing cycles for each system to ensure that the system has reached an equilibrium density. The resulting samples were then heated to various temperatures and relaxed for 50−100 ns in the NVT ensemble for further structural analysis.

199 Potential Optimization. As stated above, the first three terms in ²⁰⁰ [eq 1](#page-2-0) that define polymer interactions are intrachain in nature, largely 201 determining the bonds, angles, and dihedrals, respectively, while the interchain interactions are determined by LJ and electrostatic 202 interactions. Hence, we can reasonably separate the optimization of 203 the intrachain parameters to predict the correct bonds, angles, and 204 dihedrals, while holding the charge and the LJ parameters fixed, and ²⁰⁵ modify the latter to optimize bulk properties, such as mass density. 206

Optimizing the existing GAFF interaction potential^{[38](#page-10-0)} first required 207 a reliable set of reference data for bonds, angles, and dihedrals which 208 were produced by relaxing various isolated structures of the monomer 209 or dimer using density functional theory, as described in the previous 210 section. As mentioned previously, the partial charges for the 211 optimization were calculated using the RESP model, 40 with the 212 charges on the ions scaled by 0.8 based on ab initio results and 213 suggestions from the previous literature $44,45$ $44,45$ $44,45$ to emulate polarization 214 effects. Figure 2 shows the density predicted by the new optimized 215 f2

Figure 2. Density predicted by the newly optimized potential for PEO and P(EO-MO) at varying salt loadings compared to available experimental results for PEO.⁴

potential for two systems of interest, PEO and P(EO-MO), compared 216 with experimental data for PEO.^{[46](#page-11-0)} It is observed that our new 217 potential performs well in predicting the measured density of PEO for 218 a wide salt concentration range (see [Tables S1](https://pubs.acs.org/doi/suppl/10.1021/acs.macromol.1c01417/suppl_file/ma1c01417_si_001.pdf)−S4 for details). 219

Furthermore, when we tested the interaction potentials for 220 oligomers of different lengths, we observed that the original ²²¹ (unoptimized) GAFF interaction potential was unable to predict 222 the densities correctly over a range of lengths (see [Figure S1\)](https://pubs.acs.org/doi/suppl/10.1021/acs.macromol.1c01417/suppl_file/ma1c01417_si_001.pdf). Our 223 optimized potential performed well for the longer oligomers 224 (approximating the polymer) but was not able to predict the density 225 correctly at shorter lengths (approximating a molecular liquid). To 226 address this issue, we modified the LJ interaction for terminal carbons ²²⁷ in each chain to be slightly more repulsive (by increasing σ by a factor 228 of 1.25 and reducing ϵ by the same factor)—terminal interactions 229 necessarily begin to dominate for shorter oligomers. With this 230 modification (see [Table S5](https://pubs.acs.org/doi/suppl/10.1021/acs.macromol.1c01417/suppl_file/ma1c01417_si_001.pdf) for modified parameters), we see that the ²³¹ previously observed accuracy in predicting the density of long 232 oligomers (>8 repeat units) remains unaffected, but now, we can also ²³³ reproduce the density for shorter chain lengths. We include this detail 234 for future work. However, the metadynamics simulations for longer 235 oligomers reported below were performed using our first density- ²³⁶ optimized potential and were not repeated to include this 237 modification of termini repulsion. Smaller-scale simulations for ²³⁸ these approximations to polymer systems showed no discernible 239 differences with either potential. 240

Metadynamics. Polymer conformations produced using the 241 aforementioned simulated annealing protocol were used as initial 242 configurations to perform free-energy calculations using the ²⁴³ metadynamics approach. It is important to note that the T_g values 244 predicted by our potential are relatively high as compared to 245 experimentally determined $T_{\rm g}$ values and, hence, for similar 246 phenomenological results, we studied these systems at similar effective 247 temperatures as compared to experiments. The PEO and P(EO-MO) 248 electrolytes studied here had the same salt concentration, $r = 0.08$, 249

Figure 3. Radial distribution function of the Li ions with respect to the oxygen atoms from the (a) polymer and (b) TFSI and the corresponding CNs (c,d), respectively.

Figure 4. Pathways: 2D free-energy analysis of solvation of the Li cation and TFSI anion in PEO and P(EO-MO). Coordination with respect to oxygens of ether/acetyl and oxygens of TFSI anions are used as collective variables. Serial process: PEO shows distinct stepwise pathways to move from one coordination state to another (arrows for guide show the polymer oxygen forming a bond in the first step and then the TFSI oxygen bond breaking in the second step). Concerted process: P(EO-MO) shows a more direct path that implies that bond breaking and forming happen in a more concerted way that allows for easier switching among cation coordinations.

250 and were held at effective temperatures $(T_{\text{eff}} = T - T_g)$ of about 120 K, corresponding to absolute temperatures of 573 and 648 K, respectively. The first collective variable used in this analysis was the coordination number (CN) of a Li ion with respect to oxygen atoms from the anion (TFSI). The other collective variable was the CN of a Li ion with respect to the oxygen atoms of the polymer. These collective variables together represent the enthalpy of solvation of the lithium ion in different coordination environments. The CN is calculated as follows:

$$
CN(type 1, type 2) = \sum_{i \in type1} \sum_{j \in type2} \frac{1 - \left(\frac{|r_i - r_j|}{r_0}\right)^6}{1 - \left(\frac{|r_i - r_j|}{r_0}\right)^{12}}
$$
\n(2)

²⁶⁰ where type 1 is a specific Li ion in our simulations, type 2 is either 261 oxygen of the anion or oxygen of the polymer, and r_0 is the cutoff distance defined for the two groups of atoms. This definition of CN^{47} CN^{47} CN^{47} 262 provides a continuous function, which is why we can have non- 263 integral CNs in our simulations. The various parameters used for the 264 metadynamics simulations such as height (H, kcal/mol) and width 265 (W, unitless) of the Gaussian hills, frequency of hill addition (F, freq, 266 steps), cutoff used for measuring the CN of the Li ion with the ²⁶⁷ polymer and TFSI oxygen atoms $(r_0^{\text{Li}-\text{poly}})$ and $r_0^{\text{Li}-\text{TFSI}}$, Å), and the 268 simulation time of the calculations (t, ns) are all listed in [Table S6](https://pubs.acs.org/doi/suppl/10.1021/acs.macromol.1c01417/suppl_file/ma1c01417_si_001.pdf) for 269 both the initial tests and the production runs. 270

These simulations are performed at a finite salt concentration, with ²⁷¹ a fixed number of Li ions in the simulation cell, but our analysis ²⁷² focuses on a single, randomly chosen Li ion. Therefore, we checked 273 for independence of our results on the specific choice of the Li ion. ²⁷⁴ Metadynamics simulations were performed based on using CNs 275 centered around three different Li ions for each system with different ²⁷⁶ starting coordination environments (see [Table S7\)](https://pubs.acs.org/doi/suppl/10.1021/acs.macromol.1c01417/suppl_file/ma1c01417_si_001.pdf). Irrespective of the 277 Li ion chosen or its initial coordination environment, the resulting 278

Figure 5. 1D cut of the free-energy curves at different numbers of oxygens from TFSI for (a) PEO and (b) P(EO-MO), at different numbers of oxygens from the polymer for (c) PEO and (d) P(EO-MO), and along a diagonal from $(4,1)$ to $(5,0)$ for (e) PEO and $(5,1)$ to $(6,0)$ for (f) P(EO-MO).

279 free-energy landscapes are practically identical (see [Figure S2](https://pubs.acs.org/doi/suppl/10.1021/acs.macromol.1c01417/suppl_file/ma1c01417_si_001.pdf)). Based 280 on this sensitivity analysis, we continued running only one of the 281 simulations for each system to conserve computational time.

282 **RESULTS AND DISCUSSION**

 This section is divided into two parts: first, we investigated the local structural differences in the Li-ion coordination environ- ments within PEO and P(EO-MO) and then performed free-286 energy calculations at the same effective temperature ($T_{\text{eff}} = T$ $287 - T_g$) to reveal the relative stability of distinct ion coordinations and the energy barriers to their interconversion. 289 Next, we delved into the effect of salt loading on T_g in the poly(ether-acetal) series to understand why each polymer derivative responds differently and how this affects the performance of SPEs for battery applications.

²⁹³ Understanding Coordination Environments. MD ²⁹⁴ simulations using our optimized interaction potentials have helped to reveal subtle differences in the Li-ion coordination ²⁹⁵ environment in the poly(ether-acetals) series (see [Figure 3](#page-4-0)a for 296 f3 radial distribution functions and [Figure 3c](#page-4-0) for average CN with ²⁹⁷ respect to polymer oxygen atoms). In our previous work, [36](#page-10-0) we 298 observed that polymers containing MO units exhibit a second ²⁹⁹ oxygen coordination shell around the Li ion, which appears to ³⁰⁰ distort the first coordination shell [see the second peak in the ³⁰¹ P(EO-MO) radial distribution function, between 3 and 4 Å, ³⁰² the inset of [Figure 3a](#page-4-0)]. Our presumption is that this may create ³⁰³ a more open cage structure around the Li ion that facilitates ³⁰⁴ ion transport from one cage to another. The distorted Li-ion ³⁰⁵ coordination environment may also assist in the formation of a ³⁰⁶ lower coordination transition state while the Li ions move from ³⁰⁷ one cage to another. We also see that Li ions in P(EO-MO) ³⁰⁸ have a higher average CN $(CN = 7)$ as compared to PEO (CN) 309 $= 6$), but the increased CN is limited to intermediate distances 310 $(3-4 \text{ Å})$. 311

Table 1. Probability of Finding a Li Ion in Different Coordination Environments in PEO

coordination state	energy (kT)	probability (metadynamics)	probability (regular MD protocol) 50 ns	probability (regular MD protocol) 100 ns
(2,2)	12.480	1.252×10^{-5}	0.0009	0.0016
(3,2)	6.941	0.003185	0.0071	0.0078
(4,2)	8.892	4.527×10^{-4}	0.0073	0.0056
(3,1)	9.717	1.984×10^{-4}	0.0074	0.0074
(4,1)	4.169	0.0510	0.0904	0.0914
(5,1)	5.530	0.0130	0.0639	0.0631
(6,1)	17.04	1.310×10^{-7}	0.0076	0.0072
(4,0)	9.907	1.641×10^{-4}	0.0200	0.0192
(5,0)	2.648	0.2330	0.2946	0.3022
(6,0)	1.550	0.6990	0.4612	0.4578
(7,0)	11.59	3.048×10^{-5}	0.0397	0.0368

Table 2. Probability of Finding a Li Ion in Different Coordination Environments in P(EO-MO)

 Our MD simulations indicate a reduced propensity for cation−anion pairs in P(EO-MO) versus PEO [see the reduced intensity in the first peak of the radial distribution function (see [Figure 3b](#page-4-0)) and the overall reduction in average CN with respect to O atoms in TFSI (see [Figure 3d](#page-4-0))]. We rationalize this observation as due to the more negative partial charge on the MO oxygen atoms as compared to the oxygen atoms of EO moieties. In this sense, P(EO-MO) can compete more effectively than PEO against the TFSI anion for coordinating Li ions, thereby resulting in a lower number of ion pairs.

 Free-energy calculations further elucidate the relative stabilities of the distinct Li−O coordination environments 325 and the possible pathways to move between them (see [Figure](#page-4-0) [4](#page-4-0)). Our results suggest that multiple Li-ion coordination environments coexist in both PEO and P(EO-MO), as evidenced by multiple deep minima.

³²⁹ In PEO, we observe that the deepest free-energy minimum is ³³⁰ close to coordinations (5,0) and (6,0), that is, with 5−6 331 oxygen atoms from the polymer (O_{polymer}) and no oxygen 332 atoms from the TFSI anion (O_{TFSI}) . The second most stable 333 coordination states are at $(5,1)$ and $(4,1)$, that is, with 1 O_{TFSI} 334 and 4−5 O_{polymer}, and further less stable minima are found near ³³⁵ (2,2), (3,2), (4,2), (3,1), (6,1), (4,0), and (7,0). Similarly in ³³⁶ P(EO-MO), we observe that the deepest minimum is close to 337 (6,0) and (7,0), that is, with no O_{TFSI} and 6–7 O_{polymer}. The ³³⁸ second most stable coordination state is at (5,1), and further less stable minima are found near (2,2), (3,2), (4,2), (3,1), ³³⁹ $(4,1)$, $(6,1)$, $(5,0)$, and $(8,0)$. 340

[Figure 5](#page-5-0)a,b shows horizontal one-dimensional (1D) free- 341 f5 energy cuts for PEO and PEOMO through the two- ³⁴² dimensional (2D) landscapes for various O_{TFSI} , [Figure 5](#page-5-0)c,d 343 shows similar vertical cuts for various O_{polymer} , and [Figure 5e](#page-5-0),f 344 shows a cut along a diagonal from $(4,1)$ to $(5,0)$ for PEO and 345 $(5,1)$ to $(6,0)$ for P(EO-MO). 346

Based on the free energies of each minimum, we calculated ³⁴⁷ the approximate probability of a Li ion being in a given ³⁴⁸ coordination state by using an approximate partition function. ³⁴⁹ If ΔG_i is the relative free energy of a specific coordination 350 environment, then the probability of finding a Li ion in that ³⁵¹ environment (p_i) is given by eq 3 352

$$
p_i = \frac{e^{-\beta \Delta G_i}}{\sum_j e^{-\beta \Delta G_j}}
$$
\n(3) 353

where $\beta = 1/k_B T$. The probability of finding Li ions in each of 354 the distinct coordination environments considered is calcu- ³⁵⁵ lated based on this partition function and summarized in Table 356 t1 1 for PEO and Table 2 for P(EO-MO). Direct comparison is 357 t1t2 provided based on MD sampling over 50 and 100 ns from ³⁵⁸ regular (unbiased) trajectories in the NVT ensemble at the ³⁵⁹ same temperature. 360

Our free-energy calculations show that the most likely Li-ion ³⁶¹ coordination environment involves 6 polymer oxygen atoms in ³⁶² both PEO and P(EO-MO), with no coordination to the ³⁶³ oxygen atoms from the TFSI anion. The second most stable ³⁶⁴ coordination state of the Li ion in both cases also excludes the ³⁶⁵ anion, with a higher coordination state, (7,0), preferred in ³⁶⁶ $P(EO-MO)$ but the lower coordination state $(5,0)$ preferred in 367 PEO. In addition, P(EO-MO) exhibits a higher relative ³⁶⁸ probability for the second coordination state versus PEO. ³⁶⁹ We observe that the MD protocol and the free-energy ³⁷⁰ calculations in both systems give the same order of relative ³⁷¹ stability for the various coordination environments. However, ³⁷² we see significant differences between the MD-sampled Li-ion ³⁷³ coordination state probabilities and those estimated based on ³⁷⁴ the relative free energies derived from metadynamics [see ³⁷⁵ especially state (5,0) for P(EO-MO) in Table 2]. Overall, the ³⁷⁶ low-probability coordination environments have higher pop- ³⁷⁷ ulations in the MD trajectories, which we consider to be ³⁷⁸ overestimates. This can be attributed to the relatively high ³⁷⁹ residence times of certain ion coordination environments as ³⁸⁰ compared to the total duration of the MD simulations of 50 ns ³⁸¹ and 100 ns in the regular MD protocol. It is also important to ³⁸² note that the probability of the various coordination states ³⁸³ does not change significantly in the 50 ns between these two ³⁸⁴ regular MD simulations. This would imply either that the ³⁸⁵ regular MD protocol has equilibrated or the more likely ³⁸⁶ scenario that it is stuck in a deep local minimum. Without ³⁸⁷ foresight on the depth of the free-energy landscape, we cannot ³⁸⁸ predict the convergence time. This further establishes the need ³⁸⁹ for free-energy calculations, such as metadynamics, to avoid ³⁹⁰ such pitfalls in regular MD protocols. Metadynamics has an ³⁹¹ inbuilt mechanism to iteratively add bias to the system to ³⁹² permit exploration of the various coordination environments in ³⁹³ a reasonable amount of time. Furthermore, from [Figure 4,](#page-4-0) we ³⁹⁴ also confirm our previous hypothesis that the coordination ³⁹⁵ environment in P(EO-MO) is more diffuse than in PEO as ³⁹⁶ evidenced from the much wider minima in the free-energy ³⁹⁷ curves. 398

Figure 6. Representative snapshots of some of the coordination environments along with the energy barriers to transition from one state to another in (a) PEO at 573 K and (b) P(EO-MO) at 648 K.

 Another important insight gained from these calculations are the minimum free-energy pathways between different coordi- nation environments (see [Figure 4](#page-4-0)). PEO exhibits a serial, stepwise minimum free-energy pathway of bond making and breaking to move from one coordination state to another ([Figure 4\)](#page-4-0). To go from the $(4,1)$ state to the $(5,0)$ state in PEO, the minimum energy pathway for the Li ion is to first form an additional bond with a polymer oxygen atom, that is, (4,1) to (5,1), which requires an activation energy of about 1kT (see [Figure 5a](#page-5-0)), followed by breaking a TFSI oxygen bond, that is, $(5,1)$ to $(5,0)$, which requires an activation energy of about 1.5kT (see [Figure 5c](#page-5-0)). To go directly from the (4,1) to (5,0) state, PEO would require an activation energy of about $3kT$ (see [Figure 5](#page-5-0)e). The energy barriers for some of these transitions along with representative snapshots of the f6 414 coordination states are summarized in Figure 6a.

 P(EO-MO), on the other hand, has a more direct free- energy minimum pathway with the bond breaking and forming happening in a more concerted fashion. To go from $(5,1)$ to (6,0) in P(EO-MO), the minimum energy pathway is a diagonal connecting the two states with the TFSI oxygen bond breaking and polymer oxygen bond forming happening

simultaneously with an energy barrier of about $2kT$ (see 421 [Figure 5](#page-5-0)f). A stepwise pathway of first going from $(5,1)$ to 422 $(6,1)$ would require overcoming an energy barrier of 2.5kT first 423 (see [Figure 5b](#page-5-0)), followed by another 0.5−1kT to move from ⁴²⁴ $(6,1)$ to $(6,0)$ (see [Figure 5d](#page-5-0)), which is energetically 425 unfavorable. The energy barriers for some of these transitions ⁴²⁶ along with representative snapshots of the coordination states ⁴²⁷ are summarized in Figure 6b. Even though the net energy ⁴²⁸ barriers for the minimum energy pathways for both systems are ⁴²⁹ similar, a concerted direct process in P(EO-MO) allows for ⁴³⁰ easier transport from one cage to another as compared to the ⁴³¹ stepwise process in PEO. We also observe that for a Li ion to ⁴³² transition from a fully polymer oxygen coordinated state to one ⁴³³ with at least one oxygen from the TFSI anion has much higher ⁴³⁴ thermal activation energies on average in P(EO-MO) as ⁴³⁵ compared to PEO, which would further elucidate why there is ⁴³⁶ a much lower ion-pair concentration in $P(EO-MO)$. 437

Understanding Li-Polymer Connectivity. In the pre- ⁴³⁸ vious section, we observed that the transport mechanism for ⁴³⁹ the most efficacious^{[37](#page-10-0)} of the MO-containing polymers, P(EO- $_{440}$ MO), is probably more efficient at the same effective ⁴⁴¹ temperature. Unfortunately, even though the neat polymers ⁴⁴²

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Figure 7. (a) Effect of salt loading on glass-transition temperatures from experiments.^{[36](#page-10-0)} (b) Average number of chains for different polymers required to coordinate a Li ion. (c) Representative snapshot of single-chain coordination of a Li atom observed in our simulations in PEO and both single-chain and multi-chain coordination in P(EO-MO). (d) Normalized frequency of finding a Li ion coordinated by different numbers of chains for PEO and (e) P(EO-MO) at 363 K.

⁴⁴³ have similar glass-transition temperatures, it was observed in 444 our previous work 36 that addition of salt to the MO-containing ⁴⁴⁵ polymers results in a larger increase in glass-transition ⁴⁴⁶ temperature as compared to PEO, as seen in the left panel f7 447 of Figure 7a.

⁴⁴⁸ This increase in the glass-transition temperature can be ⁴⁴⁹ directly correlated with the number of polymer chains involved

in coordinating Li ions (see Figure 7b). It is observed that in ⁴⁵⁰ the MO-containing polymers, multiple chains are involved in ⁴⁵¹ coordinating individual Li ions. A single chain of PEO on the ⁴⁵² other hand is more easily able to provide all the required ⁴⁵³ oxygen atoms to fully coordinate the Li ion. Figure 7c shows ⁴⁵⁴ representative snapshots of 6-fold coordinated Li ions in PEO ⁴⁵⁵ coordinated by a single chain and in $P(EO-MO)$ coordinated 456

 by a single chain and two chains. We observe from a 50 ns simulation in the NVT ensemble at 363 K that in PEO, a Li ion on average is 80% coordinated by a single chain but only about 20% by two chains, with a negligible amount of three-chain coordination (see [Figure 7](#page-8-0)d). On the other hand, in P(EO- MO), two-chain coordination predominates (70%), with single-chain and three-chain coordination being comparable to each other at about 15% (see [Figure 7d](#page-8-0)).

 We can speculate from the above data that in PEO, ion transport along polymer chains is much more common, with two-chain coordination occurring when the ion hops from one chain to another. In contrast, in P(EO-MO), Li-ion hopping between polymer chains occurs more frequently due to an inability to move along a single chain or that fully coordinated states are more energetically favorable when they involve two polymer chains. If the former is true, then P(EO-MO) facilitates a more three-dimensional ion transport mechanism, as compared to quasi-1D transport along single chains in PEO. Either way, the binding of multiple chains by the Li ions results 476 in an undesirable increase in the $T_{\rm g}$. The direct consequence of this is that in the polymers we have studied, even though the transport mechanism is possibly more efficient in the MO- containing polyacetals, their higher glass-transition temper- ature leads to lower performance at battery working temper- atures with respect to PEO. Free-energy calculations to elucidate the relative stabilities of the binding motifs in these polymers and the energetics of moving along chains and between chains are currently ongoing, with the aim to help design new polymer electrolytes with favorable glass-transition temperatures and transport mechanisms.

487 CONCLUSIONS

 In this work, we predominantly study two systems from the 489 series of polymer electrolytes studied in our previous work, $36,37$ PEO and P(EO-MO) with LiTFSI, to gain insight into local ion coordination environments. Advanced free-energy sam- pling methods show the existence of multiple distinct coordination environments and their relative stabilities. The ordering of the associated populations of the coordinations is consistent with standard MD sampling; however, we view the MD-sampled populations as overestimates. We observe that 497 the $P(\overline{EO}-MO)$ electrolyte on the whole prefers ion coordination environments with higher CNs as compared to PEO and has much more diffuse coordination environments. Furthermore, PEO has a less efficient, stepwise free-energy minimum pathway for interconversion of coordination states that involves sequentially forming and breaking bonds, while P(EO-MO) has a more direct path between coordination environments and achieves this in a more concerted and efficient way. However, performant electrolytes for battery materials require efficient mechanisms of transport coupled with low glass-transition temperatures. Even with a possibly less efficient transport mechanism at a given effective temperature, PEO still has a lower glass-transition temperature at specific salt loading and hence is still the most suited in this series of polymers for electrolyte materials for battery applications. We observe that a possible reason for this is single-chain coordination in PEO as compared to a higher propensity for multi-chain coordination in polyacetal systems. Further research to understand the energetics of these different binding motifs and transport along and between chains is currently ongoing to understand the correlation between the

local structure and macroscopic properties to guide future ⁵¹⁸

\bullet Supporting Information 521

The Supporting Information is available free of charge at ⁵²² [https://pubs.acs.org/doi/10.1021/acs.macromol.1c01417](https://pubs.acs.org/doi/10.1021/acs.macromol.1c01417?goto=supporting-info). 523

Details of the parameters used in the MD simulations for ⁵²⁴ the LJ, bond, angle, and dihedral interactions for the ⁵²⁵ potential used; details on the modi_cations to LJ ⁵²⁶ parameters of the terminal carbons for fixing the density 527 of shorter-chain-length polymers for future work along ⁵²⁸ with the comparison of densities predicted by these ⁵²⁹ potentials for oligomers of di_erent lengths; details of ⁵³⁰ the parameters used in the metadynamics simulations; ⁵³¹ starting configurations for initial tests; and comparison ⁵³² of these preliminary simulations [\(PDF\)](https://pubs.acs.org/doi/suppl/10.1021/acs.macromol.1c01417/suppl_file/ma1c01417_si_001.pdf) ⁵³³

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585 Notes

⁵⁸⁶ The authors declare no competing financial interest.

⁵⁸⁷ ■ ACKNOWLEDGMENTS

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