Materials Genomics Screens for Adaptive Ion Transport Behavior by Redox-Switchable Microporous Polymer Membranes in Lithium–Sulfur Batteries

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ABSTRACT: Selective ion transport across membranes is critical to the performance of many electrochemical energy storage devices. While design strategies enabling ion-selective transport are well-established, enhancements in membrane selectivity are made at the expense of ionic conductivity. To design membranes with both high selectivity and high ionic conductivity, there are cues to follow from biological systems, where regulated transport of ions across membranes is achieved by transmembrane proteins. The transport functions of these proteins are sensitive to their environment: physical or chemical perturbations to that environment are met with an adaptive response. Here we advance an analogous strategy for achieving adaptive ion transport in microporous polymer membranes. Along the polymer backbone are placed redox-active switches that are activated in situ, at a prescribed electrochemical potential, by the device’s active materials when they enter the membrane’s pore. This transformation has little influence on the membrane’s ionic conductivity; however, the active-material blocking ability of the membrane is enhanced. We show that when used in lithium–sulfur batteries, these membranes offer markedly improved capacity, efficiency, and cycle-life by sequestering polysulfides in the cathode. The origins and implications of this behavior are explored in detail and point to new opportunities for responsive membranes in battery technology development.

INTRODUCTION

Membranes play a critical role in many battery technologies, where they serve to electronically isolate the anode from the cathode and allow the battery’s working ion to diffuse between them.1,2 For battery chemistries that involve active materials that are dissolved, dispersed, or suspended in electrolyte, membranes must also prevent active-material crossover; failure to do so leads to low round-trip energy efficiency and in some cases unacceptable capacity fade.3–5 This is particularly problematic in lithium–sulfur (Li–S) batteries, where inefficiencies and instabilities arise when soluble polysulfides—intermediates in the electrochemical interconversion of S8 and Li2S—cross over and incur a shuttling current or irreversibly react with the lithium–metal anode.6–12 While a number of strategies have been suggested for solving the polysulfide shuttle, including the use of lithiated Naion membranes and polymer14 carbon,15,16 or ceramic-coated separators, none of these approaches were capable of complete blocking of polysulfide crossing without incurring dramatic losses in ionic conductivity.

Here we show that these shortcomings are alleviated in the Li–S battery when its membrane is rationally configured from new redox-switchable polymers of intrinsic microporosity (PIMs) (Figure 1).17–22 Key to our success is the adaptation of the membrane’s transport selectivity for the battery’s working ion in situ. More specifically, we leverage the reducing environment of the sulfur cathode to chemically transform a charge-neutral and size-selective PIM membrane into a lithiated and anionic PIM membrane with enhanced polysulfide-rejecting properties. Our in situ activation strategy sidesteps well-known polymer processing challenges encountered with ionomers, where ion clustering into nonpercolating microphase-separated domains is prevalent and renders the material resistive to ion transport.22–25

The design of adaptive PIM membranes was computationally guided using a materials genome,26–28 where candidate monomer segments were screened for their susceptibility to reduction by polysulfides (i.e., a reduction potential above 2.5 V vs Li/Li+) and ionic conductivity. We experimentally validated these predictions and were further able to demonstrate that progressive reduction and lithiation of the PIM membrane by polysulfides slows...
polysulfide diffusive permeability from $1.7 \times 10^{-10}$ to $9.2 \times 10^{-11}$ cm$^2$ s$^{-1}$—an impressive 570-fold improvement over nonselective Celgard separators$^1$—without significantly impacting the membrane’s ionic conductivity ($\sigma = 5 \times 10^{-3}$ mS cm$^{-1}$ at 298 K). We also showed that by blocking polysulfide crossover, the Coulombic efficiency and cycle-life of Li-S cells greatly improves—most notably in the absence of lithium-anode protecting additives. We also showed that by blocking polysulfides (Li$_2$S$_n$, for $n = 4$–8), which are endogeneous to Li-S batteries. The ability of these new PIM membranes to adapt and sustain their polysulfide-blocking ability in situ is unusual and offers advantages over traditional approaches based on single-ion conducting membranes and other permselective barriers whose beneficial properties are ultimately transient. The origin of this transience is tied to the use of anode-protecting additives in the electrolyte (e.g., LiNO$_3$), which are consumed until exhausted and their stabilizing effects are lost thereafter.29–32

**RESULTS AND DISCUSSION**

PIMs are a compelling and versatile platform to understand structure-transport relationships in microporous polymer membranes. Transport outcomes are rationalized on the basis of membrane porosity and pore architecture and their relation to the species interacting with the membrane.$^{17–21,33}$ The membrane’s structural characteristics are dictated by polymer chain-packing relationships,$^{34–36}$ and these packing relationships are ultimately determined by monomer segments within polymer chains,$^{19–21}$ polymer processing techniques used to cast the membrane,$^{37–39}$ and membrane–electrolyte interactions.$^{40}$ In the past, PIMs have advanced as membranes with passive, nontransformable architectures; these membranes are overwhelmingly used for selective gas transport.$^{17–21}$ In the context of a Li–S battery, however, a myriad of chemical transformations can take place.$^{41–46}$ Therefore, we reasoned that PIM membranes need not be inactive; instead, they might serve as adaptive components whose microporous architectures are switchable, dynamic, and tailored at the molecular level to respond to local chemical cues within the battery’s electrolyte—in this case lithium polysulfides (Li$_2$S$_n$, for $n = 4$–8), which are endogeneous to Li–S batteries. The ability of these new PIM membranes to adapt and sustain their polysulfide-blocking ability in situ is unusual and offers advantages over traditional approaches based on single-ion conducting membranes$^{13,47,48}$ and other permselective barriers$^{49–52}$ whose beneficial properties are ultimately transient. The origin of this transience is tied to the use of anode-protecting additives in the electrolyte (e.g., LiNO$_3$), which are consumed until exhausted and their stabilizing effects are lost thereafter.$^{29–32}$

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**Figure 1.** Directed evolution of a microporous polymer membrane’s ion-transport selectivity. (a) The ion-transport selectivity of membranes cast from polymers of intrinsic microporosity (PIMs) (top right inset) can be enhanced to the benefit of Li–S battery cycle-life when redox-switchable phenazine-containing monomers are activated in situ (bottom left inset) by endogenous reducing polysulfides (Li$_2$S$_n$ for $n = 4$–8). (b) This leads to a feedback loop whereby progressive reduction of the membrane by adventitious polysulfides only serves to further restrict their access to the membrane’s pore voids.

**Figure 2.** Predictive design of redox-switchable monomer segments for adaptive microporous polymer membranes tailored for lithium–sulfur batteries. (a) A library of redox-active compounds was generated and screened computationally using a materials genome, seeking to identify those with reduction potentials ($E_{1/2}$) higher than 2.5 V vs Li/Li$^+$; monomers passing this screen would indicate they are readily reduced by lithium polysulfides present in the battery electrolyte. (b) Atom-by-atom substitutions in various PIM-monomer segments led to a number of hits passing our fitness test for $E_{1/2}$. PIMs incorporating lead compound 1 are known as PIM-7. Battery membranes derived from PIM-7 are thus expected to provide access to a new type of membrane that adapts its ion-transporting behavior by engaging the battery’s intrinsic chemistry for storing and releasing charge.
segments in the delithiated state. Structure relaxation and dipyrrolo[3,4-]
determined value of 9.0 for the battery electrolyte (Figure 2). The library design focused on phenazines (e.g., 1 and 4), 1H-isooindole-1,3(2H)-diones (e.g., 2, 5, and 6), pyrazines (e.g., 3), H-isooindolo[2,1-a]benzimidazol-11-ones (e.g., 7), benzo[g]quinoxalines (e.g., 8 and 10), benzo[1,2-c:4,5-c’]dipyrrrole-1,3,5,7(2H,6H)-tetrones (e.g., 9), and dipyrrroles[3,4-b:3’,4’-c’]pyrazine-1,3,5,7(2H,6H)-tetrones (e.g., 11)—all of which in principle could be reduced and lithiated at oxygen or at nitrogen centers upon interaction with Li2S. For example, members of the library containing 1H-isooindole-1,3(2H)-dione substituents are predicted to be reduced by polysulfides to their lithiated radical anions, while others containing diazaheterocycles were designed to undergo sequential reductions to a closed-shell dianionic (and delithiated) state, in some cases driven by rearomatization (e.g., 1 and 4).

The molecular structure and reduction potential of the PIM membrane segments were predicted using density functional theory (DFT).15,54 As lithium cations can bind to any of the electron-negative heteroatoms in the monomer segments, the most favorable binding site was identified by comparing the DFT-predicted energy of all possible Li-O/N binding configurations. The reduction potential (E1/2) was then predicted by calculating the adiabatic electronic affinity of the segments in the delithiated state. Structure relaxation and energy evaluation were carried out using the M08-SO functional,55 while solvent effects were captured by either the IEF-PCM or SMD implicit solvent models,56 where the dielectric constant value was set to the experimentally determined value of 9.0 for the battery electrolyte (Figure S8). All DFT calculations were performed using the Q-Chem software package.57

Many candidates in the library passed our initial screen (Figure 2b). To discriminate between hits, we noted that PIMs based on 2, 5, 6, and 9 typically form brittle films that preclude use as a flexible membrane. We further hypothesized that closed-shell dianionic outcomes may provide more chemical stability long-term, and thus our focus turned to monomers containing phenazines. Charge-neutral PIMs derived from phenazine-containing monomer segment 1 (calculated E1/2 = 2.90 V for the first peak and 2.28 V for the second peak vs Li/Li+) are known as PIM-1,18,58 however, the redox-active character of these polymers has not been reported previously nor has their ion-transporting ability as a membrane.

Validation of Candidate PIM Membrane Redox Chemistry with Lithium Polysulfides. To validate our predictions, we first synthesized PIM-7 via step-growth polymerization in 78% yield and Mn of 80 kg mol−1. Care was taken to adapt the synthetic methodology to afford PIM-7 with high molecular weight as needed to cast flexible membranes (see Supporting Information). With high molecular weight PIM-7 in hand, we then carried out cyclic voltammetry (CV) on the polymer drop-cast onto a glassy carbon working electrode. PIM-7 exhibited two reversible reduction peaks at E1/2 = 3.05 and 2.85 V vs Li/Li+, consistent with the reduction of the phenazine unit to the radical anion followed by the reduction to the dianionic species (Figure 3a). We noted that while the first reduction was within the range predicted by the genome screen, the second was not. We were able to resolve this incongruity in part by taking into account solvent effects using the SMD solvation model,59 which addresses solute–solvent dispersion interactions that are lacking in the currently available IEF-PCM model. Specifically, the experimentally measured dielectric constant of the electrolyte used in this work was used to simulate the electrostatic interaction for both the IEF-PCM and SMD implicit solvent models. The CDS (cavitation/dispersion/solvent-structure) parameters for the SMD were chosen to simulate the effect of the diglyme solvent: the index of refraction, surface tension, and Abraham’s hydrogen bond basicity were set to 1.4097, 36.83, and 0.859, respectively; all other CDS parameters (aromaticity, electro-negative halogenicity, and Abraham’s hydrogen bond acidity) were set to 0.0. Within the SMD context, we calculated E1/2 = 3.31 V for the first peak and 2.75 V for the second peak (vs Li/Li+) for 1. The implication here is that understanding the second chemical reduction from a lithiated radical anion to a dilithiated dianionic species benefits from the SMD treatment and is likely to best apply to the molecules in the library with multiple redox processes. In parallel, we also demonstrated

![Figure 3](https://example.com/figure3.png)
experimentally that PIM-7 could be chemically reduced when introduced to a dilute solution of Li₂S. The optical signatures of PIM-7 in its charge neutral and dianionic state were readily distinguished by UV–Vis spectroscopy (Figure 3b), with wavelength-shifts in the extinction maxima of 440–330 nm consistent with increased electron density of the polymer in its reduced state. Taken together, these results confirmed that PIM-7 membranes will become negatively charged and lithiated in the reducing environment of the Li–S battery as predicted from the materials genomics screen.

Quantitative Understanding of the Adaptive Ion Transport Behavior of PIM-7 Membranes with Lithium Polysulfides Present. Ion-selective membranes were prepared by casting PIM-7 as a thin layer on a mesoporous Celgard 2325 support using a wire-wound rod coating process. This method afforded uniform, 2 μm-thick coatings of PIM-7 on the flexible polymer support as evidenced by cross-sectional SEM (Figure S10). The packing of polymer chains for PIM-7 in the dry state yields an average pore size of 0.70 nm for the membrane (Figure S11). This size regime is predicted to be ideal for sieving polysulfides by size in battery electrolyte.

In order to confirm that PIM-7 selective layers block polysulfide crossover, we carried out crossover measurements using native supported PIM-7 membranes of a known area and thickness placed between two compartments of a diffusion cell (i.e., an H-cell). The H-cell was configured with dissolved Li₂S₈ (0.8 M S as Li₂S₈ in diglyme containing 0.50 M LiTFSI and 0.15 M LiNO₃) on the retentate side and Li₂S-free electrolyte on the permeate side (Figure 4a, and the cell shown in the inset of Figure 4b). The diffusion of Li₂S₈ to the permeate side was then monitored for up to 15 h using CV, where the concentration of polysulfides could be directly related to the measured peak current in the CV using a calibration curve determined separately for a 1–50 mM concentration regime for Li₂S₈ (Figure S12). We carried out the same experiments on unmodified Celgard separators, which are known to be poorly selective for Li₂S₈ (negative control), and for PIM-1 on Celgard, which has been reported by us to provide selectivity but not adaptability (positive control). From these data, we were able to measure peak current in the CV using a calibration curve concentration of polysulfides. Data obtained at times <3 h were below the limit of quantification and were thus omitted.

Table 1. Performance Metrics Distinguishing Non-Selective, Selective, and Adaptive Polymer Membranes

<table>
<thead>
<tr>
<th>membrane</th>
<th>membrane ionic conductivity (mS cm⁻¹)</th>
<th>polysulfide diffusive permeability (cm² s⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Celgard 2325</td>
<td>1.36 × 10⁻¹</td>
<td>(5.2 ± 0.4) × 10⁻⁶</td>
</tr>
<tr>
<td>PIM-1 on Celgard</td>
<td>5.9 × 10⁻³</td>
<td>(4.3 ± 0.3) × 10⁻⁴</td>
</tr>
<tr>
<td>native PIM-7 on</td>
<td>7 (± 2) × 10⁻³</td>
<td>(1.7 ± 0.1) × 10⁻⁴</td>
</tr>
<tr>
<td>Celgard (0 h)</td>
<td>1.7 (± 0.3) × 10⁻⁴</td>
<td></td>
</tr>
<tr>
<td>activated PIM-7 on</td>
<td>(5 ± 3) × 10⁻⁵</td>
<td>(9.2 ± 0.7) × 10⁻¹</td>
</tr>
</tbody>
</table>

Figure 4. Superior polysulfide-blocking ability by supported PIM-7 membranes and their adaptive transport behaviors in response to Li₂S₈. Time evolution of Li₂S₈ concentration in the permeate (right) of H-cells equipped with (a) Celgard (gray), supported PIM-1 (green) or supported PIM-7 (purple) membranes and (b) supported PIM-7 membranes prereduced for 0, 12, or 24 h. The retentate was charged with an initial concentration of 0.8 M S as Li₂S₈ in electrolyte. Data obtained at times <3 h were below the limit of quantification and were thus omitted.

We next sought to understand the impact of polysulfide-driven reductive chemical transformations on the polysulfide-blocking ability of PIM-7 membranes over time. To do so, PIM-7 membranes were bathed in concentrated solutions of Li₂S₈ (1.0 M S as Li₂S₈ in diglyme containing 0.50 M LiTFSI and 0.15 M LiNO₃) for a prescribed period, either 12 or 24 h, and then rinsed with and soaked in fresh electrolyte. The crossover data showed that PIM-7’s polysulfide-blocking ability is enhanced as the phenazine units are progressively reduced over time by Li₂S₈ (Figure 4b). From these data, we were also able to quantify the evolutionary changes in Li₂S₈ diffusive permeability from the baseline of (1.7 ± 0.1) × 10⁻¹⁰ cm² s⁻¹ for PIM-7 on Celgard in its initial state, to (1.4 ± 0.1) × 10⁻¹⁰ cm² s⁻¹ after 12 h and (9.2 ± 0.7) × 10⁻¹¹ cm² s⁻¹ after 24 h of chemical transformation. Extended application of Li₂S₈ beyond 24 h did not appear to further enhance the membrane’s polysulfide-blocking ability. We attribute this effect to the slow diffusion of polysulfides through the membrane and the feedback loop associated with the reduced form of the membrane further retarding the migration of additional
polysulfides. Thus, the membrane adapts its transport behavior and sustains these functions indefinitely; indeed, supported PIM-7 membranes demonstrated a stable crossover rate for at least 2 days. Advantageously, while the polysulfide-blocking character of supported PIM-7 membranes was enhanced upon increasing reduction of the phenazine subunits, the membrane ionic conductivity remained largely unchanged at $5 \times 10^{-3}$ mS cm$^{-1}$ (Table 1 and Figure S14).

**Implementation of Adaptive PIM-7 Membranes in Lithium–Sulfur Cells.** The superior polysulfide blocking ability of adaptive PIM-7 membranes over nonselective Celgard and passively selective PIM-1 membranes had a profound effect on the sulfur utilization, energy efficiency, and cycle-life of Li–S batteries (Figure 5a–c). Here we assembled Li–S coin cells using a dissolved polysulfide cathode, whereby a semisolid ink containing Li$_2$S$_8$ (1.0 M S as Li$_2$S$_8$ in diglyme containing 0.50 M LiTFSI) and Ketjenblack (5% w/w) was introduced to a high surface-area carbon nanofiber current collector. In this configuration, a high concentration of polysulfides is in direct contact with the membrane; this presents the most aggressive fitness test for the different membrane constructs. All coin cells were tested using electrolytes that were devoid of LiNO$_3$ as a non-anode-protecting additive; in doing so, the Coulombic inefficiencies associated with the polysulfide shuttle can only be improved upon by an ion-selective membrane. All cells were galvanostatically cycled between 1.8–2.8 V at a C/8 rate for up to 200 cycles.

Cells assembled with nonselective Celgard separators (negative control) were prone to Coulombic (and energy) inefficiencies associated with the polysulfide shuttle as has been previously reported.$^{29,32}$ As these cells were cycled, the charging cycle required additional energy with each cycle until the twelfth cycle, at which point the charging cycle continued indefinitely (Figure 5c). On the other hand, Li–S cells assembled with passively selective PIM-1 membranes on Celgard (positive control, Figure 5a) were significantly more effective at arresting the polysulfide shuttle; no infinite-charge regime was observed and the energy required to fully recharge these cells was sustainably low. The sulfur utilization of these cells ($\sim$1100 mA h g$^{-1}$ after the second-cycle discharge) was on par with cells assembled with Celgard, as was the capacity fade in the first few cycles; however, the cycle-life of these cells was significantly extended to 200 cycles. The specific capacity of PIM-1 cells at the end of 200 cycles was 451 mA h g$^{-1}$ with a capacity fade of 0.302% per cycle.

In contrast to cells assembled with either Celgard alone or PIM-1 on Celgard, those assembled with adaptive membranes consisting of PIM-7 on Celgard (Figure 5e) were most effective at preventing the polysulfide shuttle. The initial Coulombic efficiency of these cells was high (92.6%, compared to 87.5% for PIM-1 on Celgard and 72.9% for Celgard alone). We also noted that these cells gave markedly improved sulfur utilization, with a specific capacity of 1407 mA h g$^{-1}$ ($\sim$20% enhancement over both Celgard and PIM-1 on Celgard, and 88% of theoretical); this is consistent with their chemically evolved ability to better sequester the polysulfides to the sulfur cathode. Furthermore, cells assembled with PIM-7 on Celgard were able to sustain capacities of 774 mA h g$^{-1}$ (55% of initial) over 200 cycles, with a capacity fade of 0.225% per cycle. This result is highly unusual given that there are no anode-protecting LiNO$_3$ additives present in the electrolyte, highlighting the important role played by the redox-switchable character of the PIM-7 membrane in enabling and sustaining excellent Li–S cell performance. Notably, the versatile yet powerful aspects noted...
here for an adaptive membrane design are completely overlooked in conventional polymeric and composite ion-transporting materials used in Li–S cells. It is likely that such membranes will find synergistic use in Li–S cells with other advances in sulfur cathodes, electrolytes, and lithium–metal protection schemes.

■ CONCLUSIONS

The emerging view from our work is that macromolecular design strategies for ion-selective polymer membranes are primed for a paradigm-shift. It is now possible to use the redox environment of an electrochemical cell to chemically transform the structure and architecture of the membrane in a manner that enhances the transport selectivity of the membrane. The negative feedback loop associated with polysulfides reacting with PIM-7’s phenazine subunits and then encountering restrictions in their access to deeper pore voids is both unusual and powerful in preventing the polysulfide shuttle. To that point, past work in ion-selective membranes would suggest that it is not possible to enhance the selective transport properties of the membrane without negatively impacting membrane conductivity. Our success in this regard highlights the power of directed evolution in defining new properties in ion-transporting membrane materials. In future schemes, we see the predictive design strategies, led by materials genomics as outlined here, as essential for tailoring the switching ability to any arbitrary battery chemistry. PIMs manifest as a universal platform to address crossover problems across a variety of battery architectures, whether solid-state and solution-based electrolytes are employed. PIM membranes, adaptive and otherwise, therefore stand to significantly advance the field of electrochemical energy storage for aviation, transportation, and the grid.

■ ASSOCIATED CONTENT

* Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscentsci.7b00012.

Materials and methods, synthetic details, electrochemical methods, and other Supporting Information (PDF)

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