Computational Molecular Modeling of Transport Processes in Nanoporous Membranes

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Abstract: In this report we have discussed the important role of molecular modeling, especially the use of the molecular dynamics method, in investigating transport processes in nanoporous materials such as membranes. With the availability of high performance computers, molecular modeling can now be used to study rather complex systems at a fraction of the cost or time requirements of experimental studies. Molecular modeling techniques have the advantage of being able to access spatial and temporal resolution which are difficult to reach in experimental studies. For example, sub-Angstrom level spatial resolution is very accessible as is sub-femtosecond temporal resolution. Due to these advantages, simulation can play two important roles: Firstly because of the increased spatial and temporal resolution, it can help understand phenomena not well understood. As an example, we discuss the study of reverse osmosis processes. Before simulations were used it was thought the separation of water from salt was purely a coulombic phenomenon. However, by applying molecular simulation techniques, it was clearly demonstrated that the solvation of ions made the separation in effect a steric separation and it was the flux which was strongly affected by the coulombic interactions between water and the membrane surface. Additionally, because of their relatively low cost and quick turnaround (by using multiple processor systems now increasingly available) simulations can be a useful screening tool to identify membranes for a potential application. To this end, we have described our studies in determining the most suitable zeolite membrane for redox flow battery applications. As computing facilities become more widely available and new computational methods are developed, we believe molecular modeling will become a key tool in the study of transport processes in nanoporous materials.

Keywords: molecular simulation; membrane separations; ion-transport

1. Introduction

It is estimated that 55% of all energy consumed in chemical processes is spent on separations, of which about 50% is consumed by distillation, 20% by evaporation and 10% by drying, and the remaining 20% by non-thermal separations, include membrane-based separations [1]. Thus, further development of membrane-based separation processes to enable their use in applications currently employing thermal separations can lead to significant energy savings in chemical process industries. One obstacle to the development of membranes for these energy intensive separations is that many membrane-based separation processes are not well understood at the fundamental molecular level, thereby resulting in membrane synthesis becoming an art rather than a science [2–5]. Computational molecular modeling tools such as molecular dynamics [6] can play a crucial role...
in clarifying the molecular forces that result in making a membrane effective for a proposed application. Such molecular level understanding can thus greatly assist in the design of new membranes for a desired separation. In addition, such molecular level tools can also assist in understanding unexpected behavior observed during membrane separation processes by providing access to spatial resolution to a fraction of an Angstrom and temporal resolution to a fraction of a femtosecond. This level of resolution is often difficult to achieve in experimental methods. Selectively permeable membranes perform important roles in a wide range of systems from naturally occurring lipid membranes in biological systems to engineered polymeric membranes in filtration and energy technologies. In order to design technologies that incorporate such membranes, it is crucial to understand the behavior of these systems at the molecular level so that optimal performance and maximum efficiencies can be achieved. Computational molecular modeling tools such as molecular dynamics are ideally suited to provide detail at a level that can aid in the understanding of the transport process.

In this paper we briefly summarize three applications that use molecular dynamics techniques to examine intramembrane transport:

1. Transport of water and ions in reverse osmosis (RO) nanoporous membranes and the role played by ion-solvation in such membranes
2. Ion exchange in zeolite membranes and our finding that the separations were almost completely enthalpically driven rather than entropically
3. Separation of gases using zeolite membranes and the role of membrane loading and diffusion in the observed separation factors achieved in the membranes.

These brief overviews are followed by more detailed discussions of two recent investigations by our group:

1. We describe our investigations of multiple zeolite framework types to determine their transport behavior regarding water, protons, and vanadium ions, and investigate at the molecular level the requirements for their suitability in ion exchange membrane (IEM) applications. In addition to investigating different zeolite frameworks, the effect of composition is also examined by introducing different levels of aluminum substitution into the crystalline structure of a specific zeolite framework. By investigating two characteristics, membrane loading and intramembrane diffusion, it was possible for us to predict the overall ion permeability with the goal of optimizing the amount of aluminum substitution for high proton permeability while maintaining selectivity to undesirable ions. These and similar studies can be instrumental in designing more efficient membranes for important applications such as water purification/desalination and in many proposed applications in energy sustainability.

2. The second application focuses on how molecular simulations can help to understand unexpected or non-intuitive results obtained during experiments. A recent experimental study on the dehydration of alcohols using zeolite membranes showed [7] that the membrane was effective in dehydrating alcohol when 5% water was present but became surprisingly ineffective when the water content was 1%. This was an intriguing observation that molecular simulation was able to explain because of the fine spatial and temporal resolution accessible in such investigations that may not be possible experimentally.
2. Molecular Simulations Applied to Various Membrane Applications

Here we demonstrate the wide range of applications that can be addressed using molecular simulations.

2.1 Transport in Reverse Osmosis Membranes

Reverse osmosis (RO) separations of electrolyte solutions (such as the desalination process), is a rather interesting problem, because it is not an obvious case of separations enabled by differences in molecular sizes. In the case of an aqueous NaCl solution, for example, the size of Na+ (<0.2 nm) is certainly considerably smaller than that of a water molecule (~0.3 nm). Despite this, it is well known that membranes used for desalination, such as those described previously, allow water molecules to permeate quite easily while smaller ions such as Na+ are prevented from such permeation. Since permeation in this case could not be explained due to size differences, it was generally accepted that this was due to surface interactions between the membrane surface and the solute/solvent molecules. Molecular simulations present a useful tool for understanding these molecular forces [8]. This problem has been examined in considerable detail recently using the molecular dynamics method and has led to a significant improvement in our understanding of the forces that play a significant role in the RO-based separation of electrolyte solutions as described below.

The simulations were carried out using previously developed intermolecular potential models for both the water and the ions. For water, the simple point charge (SPC) model was used [9], while for the ions the primitive model was used [10]. These models are known to provide a realistic picture of these systems for a wide range of properties and state conditions [8,9]. Long-range forces were accounted for using the reaction field method [11]. The membrane consisted of a thin ZK4 zeolite layer with pores of 0.42 nm diameter (see Figure 1). The atoms constituting the membrane were not charged so that essentially only steric separation would be possible in these membranes.

![Figure 1. Set up for reverse osmosis separation of brine with zeolite membranes.](image-url)

One of the most remarkable results from the molecular simulation studies of aqueous electrolyte solutions was that no additional molecular forces needed to be introduced to prevent the much smaller ions from permeating the membrane, while permitting the larger water molecules to permeate the membrane. This appeared to be due to the large ionic clusters formed. The ions were...
surrounded by solvating water molecules, thus increasing their effective size quite considerably to almost 1 nm. A typical cluster formed due to the interaction between the ions and a polar solvent is shown in Figure 2. These clusters were found to be quite stable, with a rather high energy of desolvation. In addition to the clusters of the type shown in Figure 2a, larger clusters which involved more than one ion were also observed. Figure 2b shows two ions separated by a layer of water molecules forming one such large cluster. By increasing the temperature of the solution, it was possible to make ions permeate the membrane. This effectively made the larger ionic clusters less stable, and the ions were able to break away from the clusters, and the “bare” ions can then permeate the membrane. This was also found to be the case when an external electric field was included in the simulation system. The electric field also weakened the ionic clusters and again allowed ions to then permeate the membrane. These simulations have showed that the solvation of ions is at least partly responsible for these separations, and in the very least these forces must be taken into account as part of the design in RO-based separations of aqueous electrolyte solutions.

![Figure 2. a) Na+ ion (blue) solvated with water (red/white) to effectively increase its size. b) A hydrated ion pair of Na+ (blue) and Cl- ions (cyan).](image)

The ionic clusters observed are not limited to aqueous electrolyte solutions. In fact, very similar results were obtained for methanolic solutions as well [12]. This shows that sufficiently large and stable ionic clusters are a fairly common occurrence whenever ions are dissolved in polar solvents. The clusters are an essential factor in the facilitation of reverse osmosis purification. Since many industrially important solutions include ions in polar solvents, it is important to account for them in separations involving such solvents.

### 2.2 Transport in Ion-Exchange Membranes

Another example of the use of molecular simulation to examine transport of water and ions in nanoporous membranes is the ion exchange between an aqueous solution and NaA zeolite. Our group carried out such a simulation to understand the molecular basis of such exchanges [13]. The schematic for such a simulation is shown in Figure 3.
Figure 3. A typical system set up for ion exchange. Two reservoirs and two membranes in conjunction with periodic boundary conditions are usually used in simulations as shown here.

Molecular simulations were carried out to study the dynamics and energetics of ion exchanges between monovalent and bivalent cations in supercritical and subcritical (liquid) electrolyte solutions (here Li+ and Ca++ in aqueous solutions of LiCl and CaCl2) and an ion exchange membrane (NaA zeolite) using direct simulations of up to a nanosecond or more. NaA zeolites are widely used in many commercial ion-exchange processes including detergents. Our results showed that with appropriate driving forces, such ion exchange processes can be clearly witnessed and investigated using molecular simulations at these time scales. We also attempted to understand the phenomenon of ion-exchange itself at the molecular level. Our results have shown that the ion-exchange process is primarily energetically driven and entropic forces do not appear to be playing a significant role in the exchanges observed. For supercritical LiCl solutions, we found small differences between the energy of the Li+ inside and outside the membrane. In contrast, for Na+ there was a considerable energetic advantage in being outside the membrane, making the overall exchange process energetically favorable. In subcritical (liquid) LiCl solutions we found exchange to be more favorable energetically than in supercritical solutions. For Ca++ similar trends were observed, except that the differences in the energies were much larger (compared to the corresponding Li+ exchanges), making them more energetically efficient, as has also been observed experimentally [14]. This difference in clearly shown in Figure 4. In addition to clarifying the molecular basis for these ion exchanges, simulations can also potentially be very useful to determine the behavior (e.g., state dependence, etc.) of hydrodynamic parameters commonly used to characterize ion-exchange processes at a
fundamental molecular level, and to determine if the continuum hydrodynamic equations used for ion-exchange processes are applicable to nano-systems.

![Figure 4](image_url)  
**Figure 4.** The differences in configurational energy of, Li+, Ca++, and Na+ ions inside and outside the membrane plotted against the number of time steps. A positive number indicates a preference for being outside the membrane.

### 2.3 Transport in Gas Separations Membranes

Molecular modeling has also been used to study the transport and subsequent separation of gases using nanoporous membranes. A study carried out by our group using a range of zeolite membranes to support gaseous mixtures has shown good agreement with experiments exhibiting the viability of using molecular modeling to study such separations. The simulation setup for such a study is shown in Figure 5.
Figure 5. Initial system setup for gas separation simulations and the structure of FAU, MFI and CHA unit cell.

We performed a rather vigorous test of the reliability of the simulation technique to examine the separation factor of CO₂/N₂ and O₂/N₂ mixtures in zeolite membranes. In separations, the ideal separation factor (ISF) is defined as the ratio of the permeabilities of the two gases in their pure state and the usual separation factor (SF) relates to the permeabilities in mixtures (equimolar in our studies). Data obtained from our simulations can be seen in Figure 6 along with experimental results measured on similar systems.
Figure 6. The permeance of a) O₂/N₂ and b) CO₂/N₂ for both pure systems and equimolar binary mixtures compared with experimental results. The first two sets are experimental results while the last two are simulation results as marked. The numbers above the permeances represent the separation factors (SF). Note the contrasting behavior between pure fluids and mixtures for the two systems that the simulation results correctly predict.

While examining N₂/O₂ mixtures the ISF > SF, in CO₂/N₂ mixtures the opposite trend is observed (SF > ISF). Additional results from our simulations are shown in Figure 7 and confirm this behavior. This phenomenon can be explained as follows, based on observations from our simulations. In the case of N₂/O₂ for pure fluids N₂ has a higher diffusion rate than oxygen which leads to high SF since both have similar loadings. In mixtures this effect is dampened since the narrow pores do not allow N₂ to cross O₂ in mixtures. For pure CO₂ and N₂, CO₂ has a somewhat higher loading than N₂ which leads to CO₂ having a higher selectivity. In mixtures, the loading is exclusively CO₂ because of its high quadrupolar moment, which increases the selectivity in mixtures by almost a factor of 3, correctly predicted by the simulations. In addition, simulations can be a useful tool to determine the type of diffusion in the nanopores. For example, the simulations clearly show surface diffusion of N₂ in the zeolite pores as shown in Figure 7.
Figure 7. Simulation data on permeance of N$_2$ at 322 K in an equimolar N$_2$/O$_2$ mixture as a function of pressure. Note that except for surface diffusion, the two other mechanisms are clearly qualitatively incorrect.

2.4 Membranes for Redox Flow Batteries

Redox flow batteries (RFBs) show significant potential for energy storage because of their safety, capacity, and small environmental footprint [15,16]. However, this technology is not currently widely available due to problems with the ion-exchange membranes needed in RFBs. For an RFB with high storage capacity and high efficiency: (i) the electrode reactions must be reversible; (ii) both the oxidized and reduced species must have a high solubility in the electrolyte solution; (iii) there must be a large difference between the redox potentials. Several ion pairs satisfy these requirements, among them Fe/Cr [17], Zn/Br [18], and Zn/Ce [19]. Vanadium RFBs [20] constitute a special case in that only a single elemental species is present in the ions on both sides of the ion-exchange membrane (IEM). For this case the two half reactions are shown in Equation 1.

\[
\text{Cathode: } VO^{2+} + H_2O - e^- \leftrightarrow VO_2^+ + 2H^+ \\
\text{Anode: } V^{3+} + e^- \leftrightarrow V^{2+}
\] (1)

During charging and discharging, current flows through an external circuit while protons must be transported across an IEM between the two electrode compartments. IEMs thus plays a critical role in the design of an RFB [21]. While being electrically insulating it must also be impermeable to the four reactive, vanadium ions species (see reactions above). The current state-of-the-art is to employ polymeric membranes, more specifically sulfonated fluoropolymer-copolymers (commercially known as Nafion) [22]. While these polymer membranes show favorable behavior as proton exchange membranes in fuel cells (PEMFC) and direct methanol fuel cells (DMFC) [23], in RFBs they have stability problems in the highly reactive environment, resulting in the crossover of reactive ions and thus the reduction of cell efficiency and lifetime [24]. In addition, the highly oxidizing environment in RFBs tends to degrade the polymer membrane [25,26]. While some improvements have been made in these polymeric membranes to increase selectivity and stability...
[22], they have not yet overcome the obstacles for their widespread use in RFBs. New materials must therefore be designed if RFBs are to become a viable means of energy storage. We have focused on using zeolite membranes as an alternative to polymeric membranes. Zeolites are aluminosilicate crystals with ordered pore sizes varying from 0.3 nm to >1.0 nm depending on the framework type. Zeolites with high Si/Al ratios are electrically insulating and are extremely stable in both acidic and basic conditions due to their inert chemical nature [27–29]. Previously, thin zeolite membranes have been used to confirm water/ion separation via the size-exclusion mechanism [8,30,31]. We believe that thin film zeolite membranes could be used successfully as IEMs in RFBs.

To test this possibility, we have carried out molecular dynamics studies of six different zeolite frameworks (ERI, LTA, MFI, BEC, CFI, DON) for vanadium RFBs to determine the transport behavior of ions, protons, and solvent in the nanopores of the membranes [32]. The structure of these membranes were obtained from the IZA-SC’s Database of Zeolite Structures [33] and constructed to have a thickness of a single unit cell. As discussed previously, the hydration of the vanadium(II) [V^{2+}], vanadium(III) [V^{3+}], oxovanadium(IV) [VO^{2+}], and dioxovanadium(V) [VO_{2}^{+}] ions plays a key role in ion transport and this effect was examined in detail. We found that a relatively large pore (~7Å) was necessary for ion transport due to the strongly bound hydration shell that effectively increases the size of the ion. At higher ion concentrations we have observed passive (spontaneous with no external forces) permeation of the pores by the ions. This was observed in our studies when both the ion concentration and the temperature were significantly higher than the normal operating conditions in RFBs (8 mol%, 400K). We note that even at these extreme conditions, we observed permeation and subsequent transport of only H_{3}O^{+} ions in ERI, LTA, and MFI membranes. We did not observe any vanadium ion transport across the zeolite (Figure 8b). Therefore, we propose these membranes would be more suited in RFBs because they exhibit the necessary ion selectivity. This agrees with experimental measurements by Xu et al. [34] who have reported similar selective transport using substrate-supported zeolite membranes with ERI frameworks. In contrast, the larger pores in BEC, CFI, and DON frameworks did allow permeation and transport of all ion species V^{2+}, V^{3+}, VO^{2+}, and VO_{2}^{+} in addition to hydronium ions (Figure 8a), which would thus make them unsuitable for vanadium redox flow batteries as they present no proton selectivity.
This again confirmed that the size of the hydrated ion complex is a key factor in zeolite membrane transport. The only ion transport observed through membranes with smaller pores (< 5 Å) such as ERI, LTA, and MFI zeolite frameworks was that of the hydronium ion (proton). Therefore, these membranes demonstrate the selective transport of hydronium ions over vanadium ions that is an essential requirement for IEMs in vanadium RFBs.

To better understand the mechanism of the transport of the hydrated ions through a range of zeolite pores, we placed a single ion on the pore axis and moved it through the membrane at a constant velocity of 0.75 m/s. This steered molecular dynamics (SMD) technique differs from the passive diffusion observed previously and permits the calculation of the Kirkwood potential of mean force (PMF) [35] across the membrane. The PMF is defined as the potential that yields the average force for all configurations along the “reaction path”. Such methodologies have been employed previously to determine the energy barriers of membrane permeation [36]. The force needed to keep the hydrated ion on the pore axis is assumed to be the opposite of the total pairwise force exerted on the ion:

$$ F_c = -F_p = \nabla U $$ (2)

Where $U$ is the total pairwise intermolecular potential. This force can then be integrated along the path maintained by the SMD which, for the one-dimensional path used here, leads to Equation 3:
The PMFs obtained using this approach are shown in Figure 9.

\[ PMF = \int F_c \cdot dr = \int F_{cx}dx \]  

The jagged behavior of the PMF for the ERI membrane results from the varying number of water molecules in the hydration shell as it moves through the membrane. These pores are rather small, so in order for the ion to be forced through the pores, some of the hydrating waters must be detached so that the cluster is small enough to enter the pore. Once it exits the membrane, the ion is hydrated again resulting in the reappearance of the stable cluster. The ERI profile also shows asymmetric behavior. This is because the rather quick motion of the ion (0.75 m/s) does not allow for the changing number of water molecules in the hydration shell to equilibrate as it continues to move along the pore axis. The local minima observed for LTA, MFI, and BEC membranes indicates the most favorable location for the hydrated complex within the pore. These locations correspond to the cavity at the intersection of the channels running in 3 directions within the zeolite membrane. For CFI and DON membranes the maxima observed is located at the center because their pores are 1-dimensional channels. In Figure 9, it is also evident that the free energy barrier decreases with increasing pore size. While results for other vanadium ions all showed similar behavior, the hydronium ion profiles are quite different in magnitude. In Figure 10 we compare the PMF peak heights for V\textsuperscript{2+} and H\textsubscript{3}O\textsuperscript{+} ions in the six zeolites in Fig. 9. The large differences observed in the free energy barriers for the vanadium ions and hydronium ions for zeolites with pores less than 5 Å shows the effectiveness of selective transport, a desirable characteristic of RFB IEMs.
Further examination of the orientation of the hydrated ion complexes during membrane permeation yields additional insights to the relative size of the zeolite pores and the diffusing species. The \([\text{V(H}_2\text{O)}_6]^{2+/3+}\), \([\text{VO(H}_2\text{O)}_5]^{2+}\), and \([\text{VO}_2(\text{H}_2\text{O)}_4]^+\) complexes have rather stable octahedral structures, so it is interesting to monitor the changes in orientation of the hydrated ions as they traverse the membrane during our directed simulations. We measured the angle between: (1) a single \(\text{V}^{2+/3+}\)-water pair and the pore axis for \(\text{V}^{2+/3+}\) (2) In the case of \(\text{VO}^+\) the V-O bond and the pore axis, and finally for (3) \(\text{VO}^+\) one of the V-O bonds and the pore axis. The results obtained are shown in Figure 11 for the BEC membrane. This membrane was chosen as the cluster size closely matches the pore dimensions (~6.08 Å for the cluster and ~6.23 Å for the pore). Unsurprisingly, this analysis shows that outside the membrane, the \([\text{V(H}_2\text{O)}_6]^+\) complex exhibits random tumbling. However, in the small pore of the BEC membrane the angular motion of the complex is restricted, and it remains in a certain orientation until it leaves the channel.
Figure 11. Orientation of the [V(H$_2$O)$_6$]$^{2+}$ complex with respect to the pore axis during membrane transport in BEC zeolite membrane. The dashed lines indicate the zeolite boundary. Tumbling, followed by alignment and finally tumbling are observed.

In addition to determining how zeolite pore size affects ion transport, we also investigated how zeolite composition could lead to differing behavior. The membranes described in the above analyses all consisted of solely silicon and oxygen representing a Si/Al ratio of infinity. These membranes were shown to display selectivity of the hydronium ion over the heavy vanadium ions in the vanadium-RFB at pore sizes below a certain threshold. The composition of these membranes was then modified to incorporate various levels of aluminum substitution. Seven membranes of the identical MFI framework, but varying aluminum content were constructed representing Si/Al ratios of infinity, 383, 191, 95, 54, 47, and 31. The aluminum atoms were substituted at sites corresponding to the T7, T10, and T12 sites as described in studies which previously modeled the uptake of gas into MFI zeolites [37,38]. The substitutions also followed Lowenstein’s Rule [39,40] which does not allow for the presence of Al-O-Al linkages. In order to ensure that the membrane remained charge neutral, the local negative charges due to the aluminum atoms were spread across the four neighboring oxygens and an extra-framework sodium ion was placed in the channel in the vicinity of the substitution. These cations were allowed to equilibrate and find their lowest energy positions prior to the addition of any water or ions to the system.

The aluminum-substituted MFI membranes were used with the same protocol and system setup to ensure that the local charges on the aluminum atoms did not disrupt the hydrated ion complex or allow vanadium ions to permeate the membrane and alter its selective character. Our results showed that for all levels of Si/Al ratios, we still observed no spontaneous vanadium permeation for the entire simulation time of 10 ns. Once this selective characteristic had again been confirmed, the vanadium cations were replaced with hydronium ions in order to measure the dynamics of proton transport. Since the principle of the RFB mechanism rests on protons being released by the reaction in one half-cell and moving through the membrane to be used in the complimentary reaction in the other half-cell, the membranes not only need to be selective for proton transport, but also must have a high enough permeability that they not limit the reaction. For this reason, it is necessary to observe whether or not the membrane permeability is tunable using the Si/Al
ratio in order that the proton flux be maximized. The flux of any species through a membrane is
proportional to the concentration difference and the permeability \([41]\) (Equation 4).

\[
j_i = \frac{P_m}{L} (c_i - c_o)
\] (4)

Here the permeability, \(P_m\), can be represented as the product of the membrane loading, \(\frac{c_m}{c_o}\), and
the intramembrane diffusion, \(D_m\) (Equation 5). Each of these quantities were measured
independently in order to determine which was more strongly affected by the membrane
composition.

\[
P_m = P_m \frac{c_m}{c_o}
\] (5)

**Membrane Loading:** In order to observe the ionic loading of the membrane, the system consisting of
a single membrane with solution on both sides was studied with the various levels of aluminum
substitution. The solution consisted of a 5 mol\% HCl solution in which the dissociated proton was
modeled as a hydronium ion, or protonated water molecule (H\(_3\)O\(^+\)). For each membrane, the system
was simulated until the number of hydronium ions absorbed reached an equilibrium value, this
process was usually complete after ~2 ns. The number was then averaged over an additional 0.5 ns
to obtain the final values presented in Table 1.

<table>
<thead>
<tr>
<th>No. of Substitutions per Unit Cell</th>
<th>Si/Al Ratio</th>
<th>Hydronium Ion Concentration, mol/L</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00</td>
<td>∞</td>
<td>1.86 ± 0.46</td>
</tr>
<tr>
<td>0.25</td>
<td>383</td>
<td>2.02 ± 0.50</td>
</tr>
<tr>
<td>0.50</td>
<td>191</td>
<td>2.22 ± 0.57</td>
</tr>
<tr>
<td>1.00</td>
<td>95</td>
<td>2.77 ± 0.54</td>
</tr>
<tr>
<td>1.75</td>
<td>53.86</td>
<td>4.36 ± 0.52</td>
</tr>
<tr>
<td>2.00</td>
<td>47</td>
<td>5.42 ± 0.63</td>
</tr>
<tr>
<td>3.00</td>
<td>31</td>
<td>6.37 ± 0.61</td>
</tr>
</tbody>
</table>

The data clearly shows an increase in the number of framework substitutions results in an increase
in the uptake of hydronium ions. This agrees with previous findings regarding water uptake in MFI
membranes \([42]\), however the difference is much more pronounced with the charged species in this
study. This increase in absorption is due to Coulombic attraction between the positively charged
hydronium ions and the local negative charges present on the oxygens where aluminum atoms have
been substituted within the zeolite membrane framework.
Intramembrane Diffusion: In molecular dynamics simulations, the calculation of diffusion coefficients in three dimensional systems is commonly performed using Equation 6, where MSD is the mean square displacement of the species of interest.

\[ D = \frac{1}{6} \lim_{t \to \infty} \frac{d(\text{MSD})}{dt} \]  

Because we are only interested in the rate of diffusion within the membrane and not in the bulk, care must be taken, when finding the MSD, that one accounts only for the intramembrane ions. This method indicates a slight decrease in the diffusion as more framework substitutions are made (Table 2). This decrease can be explained in a similar manner – as the charged framework attracts more hydronium ions and results in an increased loading, this attraction also reduces the mobility of the absorbed ions when compared to those in the bulk solution.

<table>
<thead>
<tr>
<th>No. of Substitutions per Unit Cell</th>
<th>Si/Al Ratio</th>
<th>Hydronium Ion Diffusion, ( \times 10^5 \text{ cm}^2/\text{s} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00</td>
<td>( \infty )</td>
<td>2.66 ± 0.11</td>
</tr>
<tr>
<td>0.25</td>
<td>383</td>
<td>2.56 ± 0.24</td>
</tr>
<tr>
<td>0.50</td>
<td>191</td>
<td>2.40 ± 0.11</td>
</tr>
<tr>
<td>1.00</td>
<td>95</td>
<td>2.35 ± 0.25</td>
</tr>
<tr>
<td>1.75</td>
<td>53.86</td>
<td>2.31 ± 0.23</td>
</tr>
<tr>
<td>2.00</td>
<td>47</td>
<td>2.09 ± 0.18</td>
</tr>
<tr>
<td>3.00</td>
<td>31</td>
<td>1.98 ± 0.31</td>
</tr>
</tbody>
</table>

However, our model for the proton provides an incomplete picture of what occurs in reality. Protons in aqueous solution are known to diffuse by hopping from one water molecule to another adjacent molecule via a process known as the Grothuss mechanism [43]. Work on this topic has been quite extensive [44–47] and modeling of this phenomenon is usually performed using techniques that account for quantum behavior. Here, the model we are using is solely classical in nature and represents a permanently hydrated water molecule [46]. Due to this quantum hopping, the mobility of H+ ions in water is significantly higher than other monoatomic ions. The diffusion measured above only accounts for what is termed “vehicular diffusion”, or the proton riding along with the water molecule. A measure of the “hopping diffusion”, which is the quantum effect, would provide a more complete picture of the hydronium diffusion, but is beyond the scope of this study.

When the membrane loading and intra-membrane diffusion of the various systems are scaled relative to the pure silica membrane and the product is taken to find the relative permeability, an interesting trend emerges (Figure 12). At high Si/Al ratios (low substitution numbers), the relative permeability increases as the higher ion uptake dominates, but as more framework substitutions are made, the decreasing diffusion rate forces the permeability to stabilize at a constant value.
In this case, we observe that a substitution rate of ~2 per unit cell, corresponding to a Si/Al ratio of ~45 there exists a threshold beyond which we no longer see an increasing ionic flux. In this context, this represents the minimum number of aluminum substitutions per unit cell of zeolite that will maximize the proton transport through the membrane. As the values used here are relative to the pure silica membrane, one should not view these values as absolute, but the trends should be reproducible using experimental methods.

2.5 Dehydration of Alcohol

We discuss finally our recent studies using molecular dynamics to understand unexpected results observed in experiments [7]. Again, we emphasize the high temporal and spatial resolution provided by molecular simulations lead to increased insights into certain non-intuitive phenomena. In this particular case, a NaA zeolite membrane was used to dehydrate alcohol via vapor phase pervaporation. While the membrane was quite effective when the water content was 5% by weight, it became ineffective when the water content dropped to 1%. Such an observation certainly is counter-intuitive because in general membranes should be more effective when the undesirable component is at a lower concentration. The schematic of the system designed for this study is shown in Figure 13. The middle compartment of the simulation system contains the vapor phase mixture being investigated. Two layers of NaA zeolite membranes separate this compartment from the two side compartments which are initially empty (vacuum). This provides the driving force for the vapor to permeate the zeolite membranes. The system size was chosen to ensure that no vapor condensation takes place in the bulk phase of the vapor compartment at the system temperature of 423K. By removing some atoms that constitute the membrane we are also able to simulate a zeolite membrane with defects of approximately 1 nm – similar to those observed experimentally [7].
Figure 13. Schematic of the simulation system for vapor permeation through NaA zeolite membranes. Red represents the zeolite framework, water and IPA are between the two zeolite membranes.

Our simulation results, as shown in Fig. 14, confirm the non-intuitive trend observed during experiments [7]. In addition, the simulations also provided significant insight into why this unusual and unexpected phenomenon occurred. Simulations were carried out for pure IPA, as well as with 5 and 10% by weight water in the IPA. As can be clearly observed, the IPA was able to readily permeate the defect in the absence of water. Once water was included in the mixture the IPA permeation completely stopped in our time frame.
Figure 14. Number of IPA molecules permeating the membrane as a function of time for different compositions.

Upon further investigation, our simulations showed that there were two primary reasons that explain the decrease in IPA permeation when water is present. Firstly, we found that water molecules get adsorbed in zeolite pores as well as the defects, thus effectively reducing the effective size of the defects. When water is present at the defect sites, IPA molecules can also get adsorbed (with high adsorption energy) which further contributes to blocking the defect (Figure 15). In addition, we observed another interesting phenomenon in our simulations. In the bulk vapor phase in the presence of water, the IPA molecules tend to form larger IPA clusters which as a result effectively increases the dynamic diameter of the IPA molecules, making it more difficult for them to permeate the zeolite defects. This can be clearly observed in Fig. 16. In addition to this, as can be seen from Fig. 16 (c) and (d), the presence of water also results in fewer IPA molecules at the surface of the zeolite, which further restricts the permeation of IPA into the zeolite. We also observed that when water was present no IPA permeated the membrane.
Figure 15. Contrasting behavior exhibited with pure IPA (left side, a, c, and e) and with 5 wt % water (right side, b, d and f): axial views of the defect showing (a) pure IPA molecules permeating defect and (b) water and IPA molecules blocking the defect (water in blue); (c) as (a) above but IPA molecules not shown; (d) as (b) above but IPA molecules not shown; (e) cross section views showing pure IPA in cavity and (f) cross sectional view of both water and IPA molecules in cavity. The molecular diameters shown as spheres are not to scale.
Figure 16. Changes in the behavior of the vapor phase when water is present: (a) snapshot of the pure IPA simulation system near the zeolite membrane; (b) snapshot of simulation system for 5 wt% water in vapor phase near the zeolite membrane. (c); snapshot of the pure IPA vapor phase for a section of the simulation system (away from membrane) (d): snapshot with 5 wt% water in vapor phase (away from membrane); Key: yellow zeolite membrane; green IPA; and blue water; red: membrane defects

3. Conclusions

Membrane separations have a number of advantages over other more conventional techniques including minimal maintenance due to the lack of moving parts and the fact that unlike other techniques it does not require a phase change. As an increasing amount of membrane-based separations are developed in fields as diverse as gas separation, desalination, dialysis, batteries, biosensing, and drug delivery, it is crucial that these processes be well understood so that their efficiencies can be maximized. This requires a more complete picture of the molecular level phenomena that occur between the membranes and the species they are separating.

In this paper we have discussed the important role molecular simulations (specifically the technique of molecular dynamics) can play towards the realization of this goal. We have described our previous work which has helped provide an improved understanding of fluid and ionic transport in processes as diverse as reverse osmosis, ion exchange, gas separations, and redox flow batteries. The application of these simulations has shown how they can play two important roles. Foremost, they can allow us to understand phenomena not well understood, e.g. why water can permeate membranes in RO separations and ions cannot; or why N₂/O2 and CO₂/N₂ show opposite behavior in their separation factors when switching from pure fluids to mixtures; or why zeolite membranes can be effective in dehydrating alcohols via pervaporation when 5% water is present and become ineffective when only 1% water is present. Secondly, they can be effective screening tools for determining which membranes are suitable for a potential application, e.g. simulations clearly identified the zeolite membranes best suited for redox flow batteries while simultaneously revealing details about the ion transport process and how it can be affected by membrane structure and...
composition. Both of these roles can ultimately help provide a better understanding of the physiochemical behavior of many such membrane-based applications and aid in more efficient process design and optimization.

**Author Contributions:** All authors contributed equally to all aspects of this publication.

**Funding:** This research was supported by grants from the National Science Foundation (CBET 1545560) and State Key Laboratory of Materials-Oriented Chemical Engineering (KL13-05).

**Conflicts of Interest:** The authors declare no conflict of interest.

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