pH-Induced Reversal of Ionic Diode Polarity in 300 nm Thin Membranes Based on a Polymer of Intrinsic Microporosity

Yuanyang Rong ¹, Qilei Song ², Klaus Mathwig ³, Elena Madrid ¹, Daping He ¹, Ralf G. Niemann ¹, Petra J. Cameron ¹, Sara E.C. Dale ⁴, Simon Bending ⁴, Mariolino Carta ⁵, Richard Malpass-Evans ⁵, Neil B. McKeown ⁵, Frank Marken ¹*

¹ Department of Chemistry, University of Bath, Claverton Down, Bath BA2 7AY, UK
² Department of Chemical Engineering, Imperial College London, London SW7 2AZ, UK
³ Pharmaceutical Analysis, Groningen Research Institute of Pharmacy, University of Groningen, P.O. Box 196, 9700 AD Groningen, The Netherlands
⁴ Department of Physics, University of Bath, Claverton Down, Bath BA2 7AY, UK
⁵ School of Chemistry, University of Edinburgh, David Brewster Road, Edinburgh, EH9 3FJ, UK

To be submitted to Electrochemistry Communications

Proofs to F. Marken (f.marken@bath.ca.uk)
Abstract

“Ionic diode” (or current rectification) effects are potentially important for a range of applications including water purification. In this preliminary report, we observe novel ionic diode behaviour of thin (300 nm) membranes based on a polymer of intrinsic microporosity (PIM-EA-TB) supported on a poly-ethylene-terephthalate (PET) film with a 20 μm diameter microhole, and immersed in aqueous electrolyte media. Current rectification effects are observed for half-cells with the same electrolyte solution on both sides of the membrane for cases where cation and anion mobility differ (HCl, other acids, NaOH, etc.) but not for cases where cation and anion mobility are more alike (LiCl, NaCl, KCl, etc.). A pH-dependent reversal of the ionic diode effect is observed and discussed in terms of tentatively assigned mechanisms based on both (i) ion mobility within the PIM-EA-TB nano-membrane and (ii) a possible “mechanical valve effect” linked to membrane potential and electrokinetic movement of the membrane as well as hydrostatic pressure effects.

Keywords: electrokinetic effects; salt separation; membrane potential; nanofluidics; iontronics; electrophysiology
1. Introduction

Interest in “ionic devices” [1,2,3,4,5] has increased over the recent years with the emergence of new nanopore technologies [6,7] and availability of new microporous materials [8]. The potential for applications of these devices ranges from analytical nanofluidic tools [9], natural cell pore mimics [10], to energy harvesting systems, and ionic rectifiers for desalination [11]. Ionic rectification (or “ionic diode” [12]) effects are well-known in particular for gel-junctions [13] and hydrogel materials [14]. Both nano-engineered nano-pores and novel microporous materials can contribute to the development of ionic rectifiers and devices. There are opportunities to exploit properties of new types of microporous materials (such as polymers of intrinsic microporosity or PIMs) for improved ionic diodes. In this preliminary report we explore the effects on ion flow by applying very thin PIM membranes produced by spin-coating followed by lift off and transfer.

Polymers of intrinsic microporosity [15,16] have emerged recently as a novel class of materials based on molecularly highly rigid structures that are highly processible (e.g. soluble in organic solvents for casting of films and membranes [17]) and highly porous with typical BET surface area values \(\approx 1000 \text{ m}^2\text{g}^{-1}[18]\). Initial work focused on the gas-phase, but recently also liquid-phase applications have been explored based on stabilisation of fuel cell catalysts [19], “heterogenisation” of molecular catalysts [20,21], formation of novel microporous heterocarbon structures [22], and current rectification [23]. In previous work we employed solution casting methods to give relatively thick (typically 10 \(\mu\)m or more) PIM membranes. However, it is highly interesting to explore much thinner polymer nano-membranes. It is
demonstrated here, that even very thin films of PIM-EA-TB (with typically 300 nm thickness) can be employed successfully for current rectification in ionic diode devices.

2. Experimental

2.1. Chemical Reagents. Polymer PIM-EA-TB was prepared following a literature procedure [18]. Hydrochloric acid (37%), perchloric acid (70%), sulfuric acid (98%) and phosphoric acid (85%), and electrolyte salts were obtained from Sigma-Aldrich or Fisher Scientific and used without further purification. Solutions were prepared with deionized water of resistivity 18.2MΩ cm (at 20 °C) from a Thermo Scientific water purification system.

2.2. Instrumentation. A potentiostat system (Ivium Compactstat) was employed with Pt wires as working/counter electrodes, and two KCl-saturated calomel reference electrodes (SCE, Radiometer, Copenhagen) as sense and reference electrode in a 4-electrode membrane cell (Figure 1A). The membrane sample was always on the side of the working electrode.

2.3. Preparation of Nano-membranes. A thin PIM-EA-TB film was spin-coated on a cover glass using 2 wt.% PIM-EA-TB in chloroform solution (Laurell WS-650Mz-23NPP, 2000 rpm for 1 min). Next, the PIM-EA-TB film was lifted off in water and then transferred to a PET film with a ca. 20 μm diameter micro-hole (supplied by Laser Micromachining Limited, St. Asaph, Denbighshire LL17 0JG, UK). AFM thickness measurements of the PIM-EA-TB film on a glass slide in dry ambient conditions (Figure 1B,C) reveal typically 300 nm thickness produced with this concentration of PIM-EA-TB.
Figure 1. (A) Schematic drawing of the 4-electrode membrane polarisation experiment based on a 20 µm diameter micro-hole that is covered with a typically 300 nm thin PIM-EA-TB membrane. (B) AFM image of the spin-coated PIM-EA-TB membrane after cutting to reveal the thickness. (C) Cross-sectional profile (inset molecular structure of PIM-EA-TB).
3. Results and Discussion

3.1. Electrolyte and pH Effects. Figure 2A shows cyclic voltammetry data for a 300 nm thin PIM-EA-TB film covering a 20 µm diameter micro-hole in a 6 µm thick PET film. The data for 10 mM HCl (Figure 2Ai) clearly show very low currents in the positive potential range (“closed” diode) and significant currents in the negative potential range (“open” diode). Schematic diagrams in Figure 2E and 2F tentatively (following a mechanism suggested in earlier work [11]) explain this behaviour at least qualitatively based on protonation of the amine sites in PIM-EA-TB (see structure in Figure 1) leading to mixed proton and chloride conductivity. In the “open” diode state, the depletion effect may occur at the open side exposed to the electrolyte solution. In contrast, in the “closed” diode state depletion occurs in a restricted region and therefore causes a significant drop in current and the rectification effect. When extracting currents at -1V and +1V, the rectification ratio for the “nano-membrane” (defined here as \( I_{\text{open}}/I_{\text{closed}} \)) is 180 (compared to 576 reported recently for data obtained for much thicker PIM-EA-TB deposits [23]).
Figure 2. (A) Cyclic voltammograms (50 mV s\(^{-1}\)) for a PIM-EA-TB membrane in 10 mM HCl, 10 mM NaCl, and 10 mM NaOH. (B) Chronoamperometry (+1 and -1 V) for 10 mM HCl, 10 mM NaCl, and 10 mM NaOH. (C) Cyclic voltammograms (50 mV s\(^{-1}\)) for 10 mM NaCl with the pH adjusted with HCl. (D) As before, but with the pH adjusted with NaOH. (E,F) Schematic drawing of the ionic diode in “open” and “closed” states.
Additional chronoamperometry experiments were conducted switching the applied potential between +1 V and -1 V (see Figure 2B) and it can be observed that the current switches rapidly from +0.03 μA to -5.0 μA consistent with a rectification ratio of 180. Similar experiments performed with 10 mM NaCl do not result in significant rectification effects although conductivity through the membrane is observed (see Figure 2Aii). That is, both Na⁺ and Cl⁻ are likely to move through the porous PIM-EA-TB structure, but ion mobilities are similar and significant depletion effects do not arise. Interestingly, for the case of 10 mM NaOH, there is again a current rectification effect (see Figure 2Aiii). Chronoamperometry data in Figure 2Bii show that, when switching the potential from -1 V to +1 V, the current switches from -0.02 μA to 1.6μA, suggesting a difference in mobility for Na⁺ and OH⁻ in PIM-EA-TB causing a reversal of the ionic diode polarity. Data in Figure 2C and 2D show the gradual change in rectification effects with pH consistent with an approximate pKₐ for PIM-EA-TB at pH 4 [23].

3.2. Ionic Strength Effects. When changing the concentration for HCl and for NaOH, currents increase but the rectification ratio at +1 V and -1 V remains approximately constant (Figure 3A-B). However, when surveying different types of acids, a clear trend is observed with HCl giving good rectification effects followed by HClO₄, H₂SO₄, and H₃PO₄ giving only poor rectification effects with rectification ratios of 180, 51, 8.0, and 6.7, respectively (Figure 3C). The poorer rectification effects with H₃PO₄ are likely to be linked to phosphoric acid exhibiting pKₐ values higher than that of the PIM-EA-TB, and therefore providing additional protons to weaken the ion depletion effect. When comparing neutral electrolyte media for a range of different anions (see Figure 3D), significant rectification effects are observed also for BF₄⁻, suggesting that the nature of the anion also can have a strong effect on current rectification phenomena in the PIM-EA-TB membrane.
Figure 3. (A) Cyclic voltammograms (50 mVs$^{-1}$) for PIM-EA-TB membrane in (i) 10 mM, (ii) 50 mM, and (iii) 100 mM HCl. (B) As above, but for (i) 10 mM NaOH, (ii) 50 mM NaOH, and (iii) 100 mM NaOH. (C) Cyclic voltammograms (50 mVs$^{-1}$) for a PIM-EA-TB membrane in 10 mM HCl (i), $\text{H}_2\text{SO}_4$ (ii), $\text{HClO}_4$ (iii), and $\text{H}_3\text{PO}_4$ (iv). (D) As above, but with 10 mM $\text{Na}_2\text{SO}_4$ (i), $\text{NaCl}$ (ii), $\text{NaClO}_4$ (iii), and $\text{NaBF}_4$ (iv). (E) Cyclic voltammograms (50 mVs$^{-1}$) for a PIM-EA-TB membrane in 10 mM HCl with varying hydrostatic pressure (see F inset, the fill-height of one half-cell relative to the second was systematically varied by adding or removing electrolyte to give (i) -10, (ii) -5, (iii) 0, (iv) +5, (v) +10 cm hydrostatic pressure in the counter relative to the working compartment). (F) As before, but for 10 mM NaOH.
3.3. Hydrostatic Pressure Effects. Measurements were performed for 10 mM HCl (Figure 3E) and for 10 mM NaOH (Figure 3F) by varying the hydrostatic pressure across the measurement cell by simply adding or removing electrolyte solution (see inset in Figure 3F, data for 0, +/-5, and +/-10 cm water column). For HCl the ionic diode effect is significantly affected by hydrostatic pressure, whereas the effects for NaOH are small. Strikingly, the “open diode” current is increased when additional pressure is applied. This suggests that further “opening” or “tuning” of the diode could be linked to a “mechanical valve” phenomenon, where “bulging” of the thin PIM-EA-TB nano-membrane over the 20 μm microhole will cause a geometry change and thereby contribute to additional changes in flow of current. Further work, for example revealing how membrane potential (associated to electrokinetic effects) and ion size effects are linked to the observed phenomena, will be necessary to identify, quantify, and exploit the processes behind these PIM nano-membrane-based current rectification effects.

4. Conclusion

It has been shown that a very thin (300 nm) PIM-EA-TB membrane can generate significant “ionic diode” effects and could therefore be employed in ionic rectifier-based devices. The switch in ionic diode polarity is novel and linked to the pH of the electrolyte. This phenomenon is explained here tentatively as based on (i) ion mobility differences within the PIM-EA-TB film and (ii) a valve-like opening of the membrane (driven by electrokinetic or hydrostatic effects) over the supporting PET film microhole. Further work will be required, experimentally and theoretically, to better understand ion mobility and reactivity within PIM films and to further develop and exploit these phenomena in areas such as energy harvesting and desalination.
Acknowledgements

Y.R. thanks the University of Bath for a fee waiver and the China Scholarship Council for a PhD stipend. E.M. acknowledges financial support from the EPSRC (EP/K004956/1). F.M. and N.B.M. gratefully acknowledge the Leverhulme Trust for support (RPG-2014-308).

References


