



In Situ Kinetics Studies in All-Vanadium Redox Flow Batteries

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We report results of polarization measurements resolved for the negative and positive electrodes of vanadium redox batteries (VRBs) using a dynamic hydrogen electrode in an operating battery cell. Electrochemical experiments with symmetric electrolyte feeds were also performed. Greater kinetic polarization is observed at the negative ($V^{3/2+}$) electrode compared to the positive electrode ($V^{5/4+}$), in contrast with previously reported ex situ measurements. For the positive electrode, the polarization in the low-current regime was modest and was not kinetically controlled. The relative rates of reaction are a surprise since it might be expected that the $V^{3/2+}$ redox reaction is a simple outer-sphere electron transfer.

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Increasing demand for renewable energy, focused on intermittent sources such as wind and solar, has resulted in great interest in large-scale energy storage. Redox flow batteries (RFBs) are proposed to allow large-scale, low-cost electrical storage, enabling intermittent energy production technologies to take a greater share of the domestic energy portfolio.¹ To realize this goal, significant improvements are needed in RFB performance and cost.

A variety of processes can contribute to performance limitations in RFBs. At the cell level in the all-vanadium chemistry, reaction kinetics, ionic transport through electrodes and the separator, electronic contact resistance, and mass transport resistance into and out of the electrodes are all expected to reduce the power available from a RFB. Polarization curve analysis has been used in all-vanadium RFBs to aid in separating the above contributions to overpotential at varying operating currents.² It is quite important to assess the relative magnitude of these losses, particularly in the low current regime of high conversion efficiency. Generally, this region can be very strongly affected by kinetic limitations.

VRB reaction kinetics studies in the VRB cell itself are somewhat complicated by the need for a reference electrode. Thus, most investigations of kinetics to date were carried out using ex-situ methods, such as rotating disk electrode (RDE) voltammetry or stationary cyclic voltammetry (CV). Gattrell³ observed Tafel slopes as high as 130–500 mV/decade and exchange currents ranging from ~ 0.2 mA/cm² to ~ 7 mA/cm² for the $V^{5/4+}$ couple on a glassy carbon RDE. Suspicion of kinetic limitations has led some to treat the carbon electrodes to improve reaction rates, but with little appreciable improvement.⁴ Recent work by Manahan⁵ found that improved cell performance resulted from replacing the negative electrode carbon paper with higher surface area carbon paper; similar improvement was not realized when the positive electrode material was replaced. However, two-electrode in-situ tests suffer from the inability to definitively ascribe the observed losses or improvements to either the positive or negative electrode.

Inserting a reference electrode into the VRB single cell allows electrode kinetics measurements to be performed in-situ by separating individual electrode contributions to the observed loss after iR correction. Here we report our implementation of a dynamic hydrogen electrode as a reference electrode for a study focused on the low-current region of the polarization curve to probe the kinetics of the positive and negative sides of a VRB using carbon paper electrodes.

Methods

Cell construction.— The RFB hardware used in this work was a modified direct methanol fuel cell with an active area of 5 cm² described in detail previously.⁶ Single layers of untreated 10 AA carbon paper (SGL group) with a nominal thickness of 410 μ m and 328 cm² BET surface area⁷ were used as electrodes on both sides of the cell. Temperature was maintained at 25°C via cartridge heaters on the cell. Two layers of Nafion 117 (Ion-Power) enabled the incorporation of a dynamic hydrogen electrode (DHE) into the cell,⁸ as illustrated in Figure 1.

The electrolyte was 0.1 M vanadyl sulfate (Alfa Aesar) in 5.0 M sulfuric acid (Alfa Aesar). A state-of-charge (SOC) of 50% was used. Details of electrolyte preparation and charging are given in Aaron et al.⁶ The 0.1 M vanadium concentration was lower than what is used in practical VFRBs but is a more thermally stable solution than obtained at higher concentration.⁹ The electrolyte flow rate was 20 mL/min, corresponding to a maximum utilization of 7.5% at the highest measured current.

Electrochemical measurements.— Potentiostatic polarization curves were performed in the low current region – up to approximately 40 mA/cm² – to focus on kinetic control. All overpotentials were controlled with respect to the stable open circuit potential prior to the beginning of the experiment. To maintain a constant 50% SoC throughout a polarization measurement, 4 s steps of alternating polarity overpotential were applied. Figure 2 shows the applied potential at the $V^{5/4+}$ electrode and the cell current vs. time, illustrating that a 4 s discharging step at, for example, -1 mV overpotential was followed by a 4 s charging step at $+1$ mV overpotential relative to the OCV. It is apparent that the ending OCV was the same as the starting OCV, indicating that the VRB SoC was unchanged throughout the course of the experiment. The current was sampled during the final 25% of each time step to exclude capacitive charging currents. The high frequency resistance (HFR) was measured using AC impedance at 10 mV r.m.s. amplitude at roughly 15 kHz. The areal specific resistance (ASR, or HFR*area) on the positive side was 0.513 Ω -cm² and 0.519 Ω -cm² on the negative side. The whole cell ASR, measured from the positive to the negative electrode, was 1.07 Ω -cm². These values were slightly larger for the symmetric cell experiments, described below.

A second set of experiments was performed to verify the polarization curve studies described above. In these, a symmetric cell was operated in which both sides of the battery were fed a 50% SOC $V^{3/2+}$ or $V^{5/4+}$ electrolyte. The electrolyte for both sides was contained in one external reservoir so that no SOC change could take place. Thus, for the $V^{3/2+}$ symmetric cell, V^{2+} was oxidized to V^{3+} on one side

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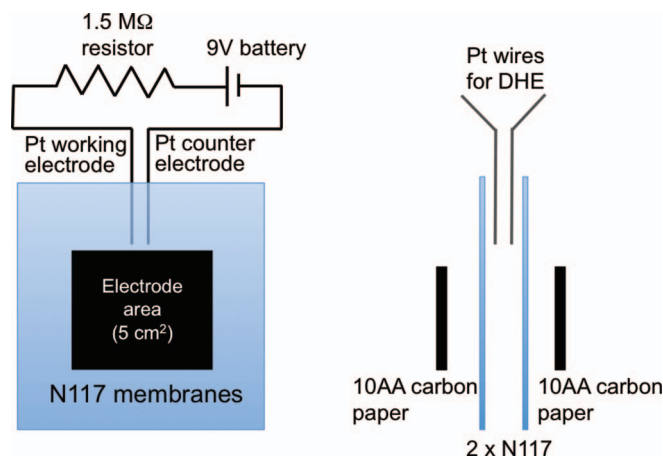


Figure 1. Placement of the DHE in the RFB (face view and side view on left and right, respectively). On the side view, the spacing between the tips of the Pt wires was ~ 1 mm; the tips of the wires were ~ 1 mm outside the active area of the electrodes.

106 while V^{3+} was reduced to V^{2+} on the other side (and similar for the
107 $V^{5/4+}$ symmetric cell).

108 The DHE was inserted between layers of Nafion 117. A steady
109 current of $6 \mu\text{A}$ flowed between the electrodes of the DHE. This pro-
110 vided a stable reference voltage over the course of several months of
111 operation in the VRB. Within experimental error, identical HFR values
112 were observed between the reference and each of the electrodes in
113 the cell, indicating that the DHE provided a plausible reference when
114 placed outside the electrode active area. The accuracy of the potential
115 relative to reference electrode placement, the geometry of the system
116 and the relative rates of reaction (secondary current distribution)
117 were analyzed by He¹⁰ and Liu.¹¹ To meet criteria suggested in these
118 works, we placed the reference electrode far (relative to membrane
119 thickness) outside the edge of either electrode. We also investigated
120 the measured kinetics in both normal and symmetric cells, finding
121 identical current-potential behavior.

122 Results

123 Figure 3 shows charging and discharging polarization data for the
124 positive and negative electrodes of the VRB. The negative electrode
125 exhibited much greater overpotential at all cell current densities than
126 the positive electrode. If the current response to overpotential is ki-
127 netically controlled, a plot of log current against overpotential should
128 achieve a linear trend once the individual electrode reactions have
129 been biased primarily in one direction (i.e. charging or discharging).

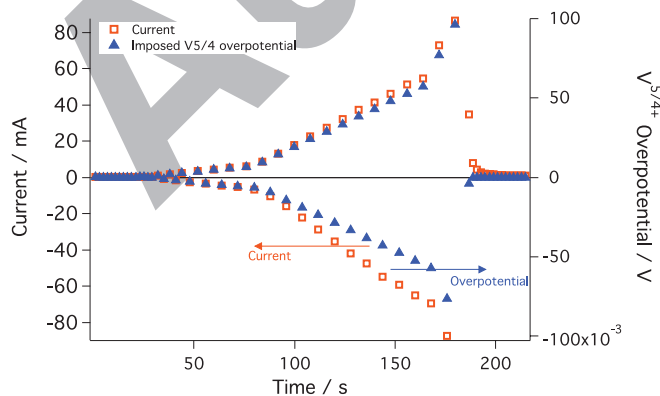


Figure 2. Current and potential profiles over time for the alternating charge-discharge polarization curves. Note that an ending overpotential of 0 V indicates that the VRB SoC was unchanged. The electrolyte was 0.1 M vanadium in 5.0 M H_2SO_4 .

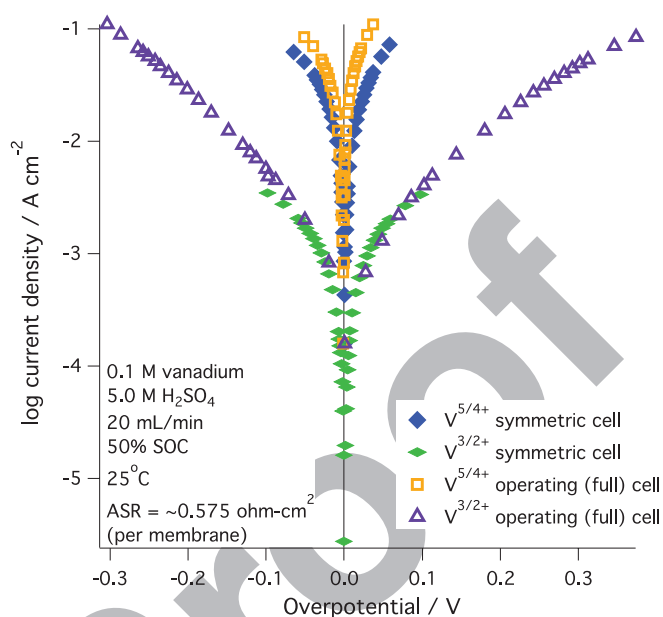


Figure 3. Kinetic region polarization data for VRB charging and discharging behavior. The electrolyte was 0.1 M vanadium in 5.0 M H_2SO_4 .

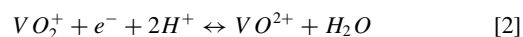
130 Construction of a Tafel plot in this manner allows kinetic parameters
131 such as the Tafel slope and exchange current density to be obtained.
132 The Tafel range spans approximately 1.2 decades of log current den-
133 sity for the full cell experiment. The exchange current density, j_0 , is
134 of particular interest since a higher j_0 indicates faster kinetics.

135 The charging and discharging curves for the negative electrode
136 yielded Tafel slopes of 194 and 204 mV/decade, respectively. From
137 the intercept of the Tafel plot, the geometric area-normalized exchange
138 current density for the $V^{3/2+}$ couple was calculated as 0.148 and
139 0.149 mA/cm^2 (BET-area-normalized 2.25×10^{-3} and 2.28×10^{-3}
140 mA/cm^2) for charging and discharging, respectively. The R^2 values for
141 the linear fits exceeded 0.998, indicating very good correspondence
142 to Tafel behavior for the $V^{3/2+}$ couple.

143 However, the positive electrode did not exhibit linear behavior on
144 the Tafel plot. Indeed, the $V^{5/4+}$ reaction was not sufficiently polarized
145 to reach the Tafel region. In this situation we can estimate the j_0
146 from the linearized Butler-Volmer expression, allowing a plot of current
147 density against overpotential to yield a slope directly proportional to
148 j_0 .¹² With this approach, the geometric area-normalized exchange cur-
149 rent density of 6.48 and 6.65 mA/cm^2 (BET area-normalized 0.0988
150 and 0.101 mA/cm^2) for charging and discharging was obtained for the
151 $V^{5/4+}$ couple (R^2 was 0.9999). Wen et al.¹³ found an exchange
152 current density of 0.218 mA/cm^2 for 0.5 M vanadium at 50% SoC via
153 glassy carbon RDE. The experimental conditions and materials used by
154 Wen et al.¹³ can account for the difference in exchange current. Indeed,
155 it is important to note here that the redox process at different carbon
156 materials could exhibit different kinetic behavior. This expectation is
157 at the heart of our desire to carry out these measurements in situ using
158 the exact battery electrode material.

159 For these materials, we find that the exchange current density for
160 the $V^{5/4+}$ couple is ~ 44 times greater than the $V^{3/2+}$ couple. Clearly,
161 the $V^{5/4+}$ reaction is far more kinetically facile than the $V^{3/2+}$ reaction.
162 The VRB performance loss was dominated by the negative electrode
163 kinetics. The symmetric cell measurements support the operating full
164 cell measurements with very similar current responses to imposed
165 overpotentials..

166 Equations 1 and 2 show the overall reactions that occur at the
167 negative and positive electrodes, respectively.



169 Glancing at equations 1 and 2 can lead to the expectation that
170 the $V^{5/4+}$ couple should have comparatively worse kinetics due to the
171 more complex reaction involving participation of protons and removal
172 of an oxygen from the V^{5+} ion while the $V^{3/2+}$ couple involves a single
173 electron transfer, expected to be a simple outer sphere electron transfer.
174 A more complete understanding of this is outside the scope of this
175 communication. Clearly more detailed work is needed. However, this
176 finding is suggestive relative to catalytic approaches to improving the
177 negative electrode performance.

178 Conclusion

179 Inclusion of a DHE in an operating VRB allowed in-situ kinetics
180 studies to be performed via polarization curve analysis. The negative
181 electrode dominated overpotential at all current densities, including
182 the low-current kinetic region. The $V^{5/4+}$ exchange current density
183 was 44 times greater than that of the $V^{3/2+}$ couple, indicating much
184 faster kinetics for the $V^{5/4+}$ couple. Relatively large kinetic-region
185 losses observed in VRBs can thus be largely attributed to the $V^{3/2+}$
186 couple, which also means that catalysis efforts should be focused on
187 this electrode, rather than the $V^{5/4+}$ couple.

References

1. L. Joerissen, J. Garche, and C. Fabjan, *J. Power Sources*, **127**, 98 (2004). 188
2. D. Aaron, Z. Tang, A. B. Papandrew, and T. A. Zawodzinski Jr., *J. Appl. Electrochem.*, **41**, 1175 (2011). 189
3. M. Gattrell, J. Park, B. MacDougall, J. Apte, S. McCarthy, and C. W. Wu, *J. Electrochem. Soc.*, **151**, A123 (2004). 190
4. K. J. Kim, M. S. Park, J. H. Kim, U. Hwang, N. J. Lee, G. Jeong, and Y. J. Kim, *Chem. Commun.*, **44**, 5455 (2012). 191
5. M. P. Manahan, Q. H. Liu, M. L. Gross, and M. M. Mench, *J. Power Sources*, **222**, 498 (2013). 192
6. D. Aaron, Z. Tang, J. S. Lawton, A. B. Papandrew, and T. A. Zawodzinski, *ECS Trans.*, **41**, 43 (2011). 193
7. S. C. Barton, Y. Sun, B. Chandra, S. White, and J. Hone, *Electrochemical and Solid-State Letters*, **10**, B96 (2007). 194
8. G. Li and P. G. Pickup, *Electrochim. Acta.*, **49**, 4119 (2004). 195
9. J. Zhang, L. Li, Z. Nie, B. Chen, M. Vijayakumar, S. Kim, W. Wang, B. Schwenzer, J. Liu, and Z. Yang, *J. Appl. Electrochem.*, **41**, 1215 (2011). 196
10. W. He and T. V. Nguyen, *J. Electrochem. Soc.*, **151**, A185 (2004). 197
11. Z. Liu, J. S. Wainwright, W. Huang, and R. F. Savinell, *Electrochim. Acta*, **49**, 923 (2004). 198
12. A. Bard and L. Faulkner, *Electrochemical Methods: Fundamentals and Applications*, John Wiley & Sons, New York, 2001. 199
13. Y. H. Wen, H. M. Zhang, P. Qian, H. P. Ma, B. L. Yi, and Y. S. Yang, *Chin. J. Chem.*, **25**, 278 (2007). 200

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