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***In situ* Electrosynthesis of Anthraquinone Electrolytes in Aqueous Flow Batteries**

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Abstract

We demonstrate the electrochemical oxidation of an anthracene derivative to a redox-active anthraquinone at room temperature in a flow cell without the use of hazardous oxidants or noble metal catalysts. The anthraquinone, generated *in situ*, was used as the active species in a flow battery electrolyte without further modification or purification. This potentially scalable, safe, green, and economical electrosynthetic method is also applied to another anthracene-based derivative and may be extended to other redox-active aromatics.

Introduction

Aqueous redox flow batteries (ARFBs) represent a class of devices for storing electrical energy that are especially well suited for large-scale stationary deployment.^{1, 2} Vanadium redox flow batteries, the most developed ARFB technology, have been limited by the high and fluctuating price of vanadium.³

Anthraquinone-based aqueous redox flow batteries are considered as one class of the most promising alternatives to vanadium redox flow batteries because they can be composed of earth-abundant elements such as C, H, O, and N while providing comparable electrochemical performance.⁴⁻⁹ However, reducing the production cost of anthraquinone-based electrolytes and

27 improving their chemical stability are two major challenges preventing them from being cost-
28 competitive.⁹⁻¹⁴ Many factors can influence the synthesis cost of an organic molecule, including
29 the number, duration, complexity, and yields of the reaction steps, the reaction conditions (time,
30 temperature, and pressure), solvent and precursor costs, the cost of waste disposal, and economies
31 of scale. Likewise, a host of factors contributes to the stability, and by extension the long-term
32 viability, of redox-active organics including the chemical structure, solvent conditions, applied
33 potentials, and state of charge. Only through careful consideration of all of these factors can
34 commercial-scale organic ARFBs be viable storage solutions. Therefore, not only is the
35 development of a stable anthraquinone important, but the design of a potentially economical,
36 scalable, and green synthetic route toward targeted molecules is equally significant.^{11, 15}

37 Electrochemically-mediated synthesis (electrosynthesis) enables the replacement of hazardous
38 oxidizing and reducing agents by electric current, or “clean” electrons, through an electrode and
39 has attracted considerable attention for both laboratory and industrial applications in multiple
40 fields of research.¹⁶⁻²¹ Compared to traditional thermochemical synthesis, electrosynthesis can be
41 significantly more environmentally benign due to reduced waste production and alternative
42 chemicals consumed.^{22, 23} However, the necessity of using specific solvents combined with
43 supporting electrolytes, along with their subsequent separations, are some of the primary hurdles
44 limiting the feasibility of electrosynthesis compared to thermochemical processes in many cases.¹⁶

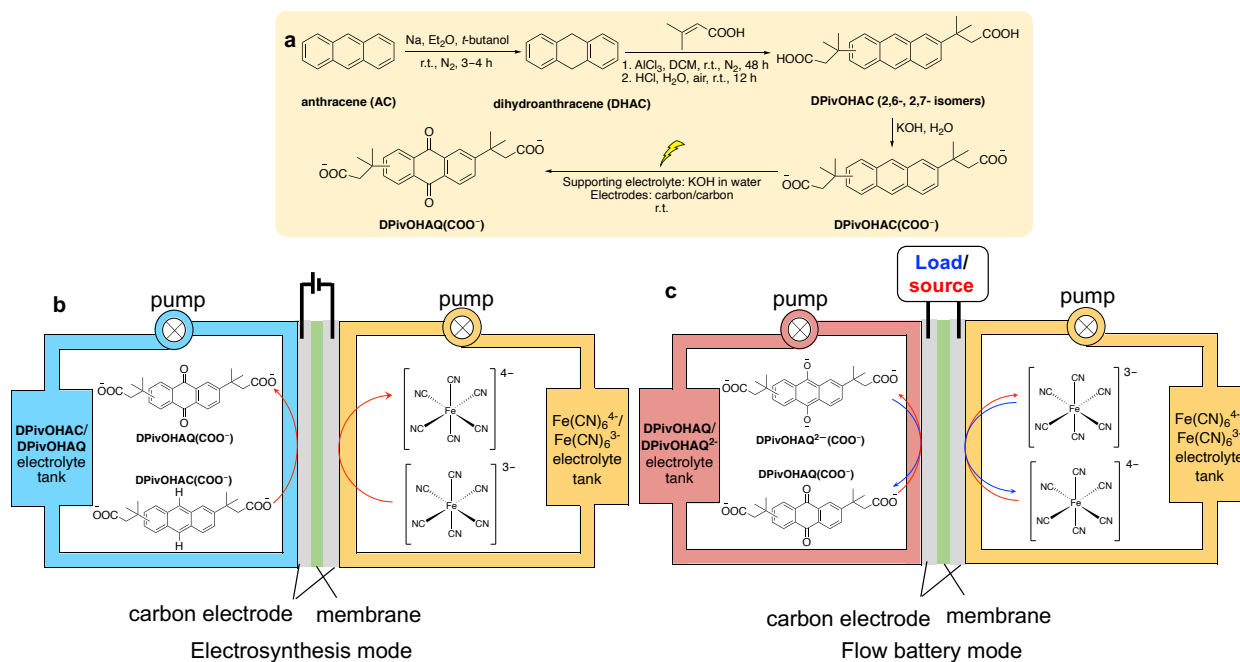
45 As an example, anthraquinone is typically produced from anthracene, an inexpensive and
46 abundant component of coal tar and petroleum.²⁴ Typically, hazardous oxidants such as
47 cerium(IV), chromium(VI), and vanadium(V) compounds dissolved in strong acids, sometimes at
48 elevated temperatures, are used to facilitate this thermochemical conversion.²⁵ To minimize the
49 use of hazardous materials, often these consumed oxidants are electrochemically regenerated and

50 reused for chemical oxidations,²⁵⁻³⁰ that is, a mediated or indirect electrochemical oxidation.
51 However, in both thermochemical conversion and mediated (indirect) electrochemical conversion,
52 isolating anthraquinone from these hazardous solutions can be time- and capital-intensive. Electro-
53 oxidations of anthracene and its derivatives at ~1 mM concentration have been performed
54 previously; however, the low concentrations of anthracene substrates and poor selectivity of the
55 reactions have prevented the method from being synthetically useful.³¹⁻³⁶

56 Using a scalable flow cell setup,³⁷ we demonstrate the capability to electrochemically oxidize
57 water-soluble anthracenes directly to anthraquinones in electrolytes without the use of strong
58 oxidants or catalysts, producing the desired negolyte (negative electrolyte) and ferrocyanide
59 posolyte (positive electrolyte) *in situ*. Compared to conventional thermochemical and
60 electrochemical methods, the new method is safe and potentially inexpensive because it eliminates
61 both the use of hazardous oxidants and the necessity of post-synthesis isolation of the products
62 from the supporting electrolytes. Taking advantage of a flow cell and bulk electrolysis setup, the
63 demonstrated electrosynthetic method is amenable to both continuous and batch processing.
64 Furthermore, we confirmed that the electrosynthetic method can also be extended to other
65 anthracene derivatives.

66 3,3'-(9,10-anthraquinone-diyl)bis(3-methylbutanoic acid) (**DPivOHAQ**) was recently reported
67 as an extremely stable and potentially inexpensive negolyte active species for organic ARFBs.³⁸
68 However, the use of CrO₃ in the synthesis can be highly toxic and explosive if produced in large
69 scale. Figure 1a shows the synthetic route for **DPivOHAQ** in three steps: 1) Through Birch
70 reduction, anthracene (**AC**) is converted to 9,10-dihydroanthracene (**DHAC**) at room temperature
71 (Figure S1). 2) After a Friedel–Crafts reaction and subsequent oxidation by air in one pot, two
72 water-soluble groups are introduced and **DHAC** is re-oxidized to an **AC** derivative (Figure S2),

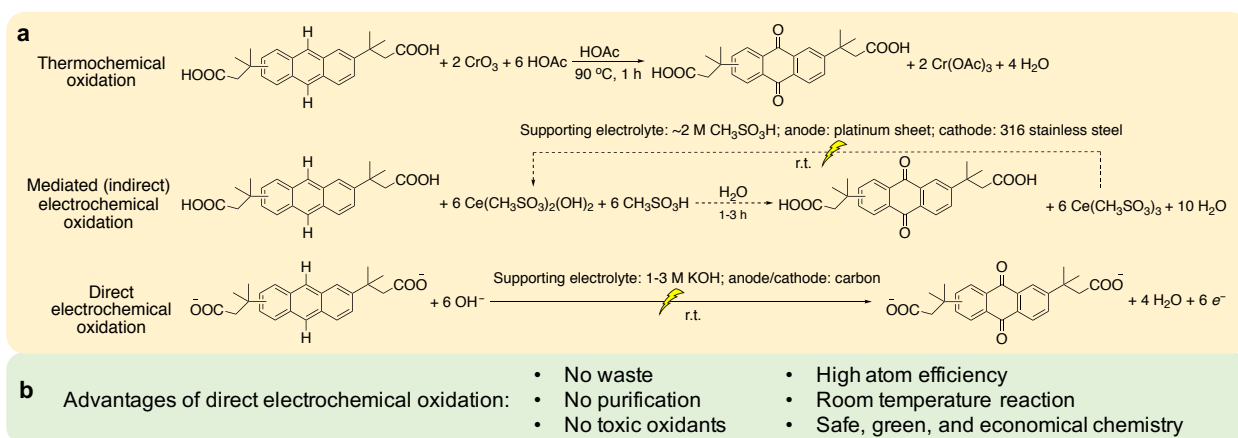
73 forming 3,3'-(anthracene-diyl)bis(3-methylbutanoic acid) (**DPivOHAC**). The **DPivOHAC**
 74 powder was then dissolved in water by adding KOH to deprotonate the carboxylic acid groups. 3)
 75 Lastly, **DPivOHAQ** negolyte active species is produced by electrochemical oxidation in an
 76 aqueous electrolyte without the need for further purification. Figure 1b illustrates how
 77 **DPivOHAQ** and ferrocyanide active species can be produced *in situ* in the flow cell's
 78 electrosynthesis mode. These materials can directly serve as the active species in the negolyte and
 79 the posolyte, respectively, of a flow battery in the same cell as illustrated in Figure 1c.



80
 81 **Figure 1. Preparation of DPivOHAQ and the corresponding flow battery.** (a) The
 82 **DPivOHAQ** synthetic route and conditions starting from anthracene. (b) The setup for
 83 electrosynthesis of **DPivOHAQ** and ferrocyanide. (c) The flow battery setup with **DPivOHAQ**
 84 negolyte (generated *in situ*) and ferrocyanide posolyte (generated *in situ*). **DPivOHAC**: 3,3'-
 85 (anthracene-diyl)bis(3-methylbutanoic acid); **DPivOHAC(COO⁻)** is deprotonated **DPivOHAC**.
 86 **DPivOHAQ**: 3,3'-(9,10-anthraquinone-diyl)bis(3-methylbutanoic acid); **DPivOHAQ(COO⁻)** is
 87 deprotonated **DPivOHAQ**.

88 Figure 2a lists three different oxidation methods for **DPivOHAQ** synthesis. Conventionally,
 89 anthracene derivatives can be chemically oxidized to their anthraquinone forms by oxidants such
 90 as chromium oxide (CrO₃) in strong acidic media at elevated temperature.³⁸ To minimize the use

91 of hazardous oxidants, the strategy of mediated electrochemical oxidation can be performed by
 92 regenerating oxidants such as cerium(IV) compounds.^{26, 29} However, in both of these
 93 thermochemical and indirect electrochemical oxidation processes, tedious and expensive isolation
 94 of anthraquinone-based products from oxidants and acids is required. Taking advantage of the high
 95 solubility of **DPivOHAC** in base, we demonstrate a synthetic route via direct electrochemical
 96 oxidation in alkaline electrolyte with a flow cell. This method allows the complete elimination of
 97 hazardous oxidants and costly separation processes.



98

99 **Figure 2. Comparison of DPivOHAQ synthetic methods.** (a) Thermochemical, mediated
 100 (indirect) electrochemical, and direct electrochemical oxidation reactions to synthesize
 101 **DPivOHAQ**. (b) Advantages of direct electrochemical oxidation *in situ*.

103 Experimental

104 Cell hardware

105 Glassy carbon was used as the working electrode for all three-electrode cyclic voltammetry (CV)
 106 tests with a 5 mm diameter glassy carbon working electrode, an Ag/AgCl reference electrode
 107 (BASi, pre-soaked in 3 M NaCl solution), and a graphite counter electrode. Both undivided cell
 108 and divided cell were built for electrosynthesis. Flow battery experiments were constructed with
 109 cell hardware from Fuel Cell Tech (Albuquerque, NM) assembled into a zero-gap flow cell
 110 configuration. Pyrosealed POCO graphite flow plates with serpentine flow patterns were used for

111 both electrodes. Each electrode comprised a 5 cm² geometric surface area covered by AvCarb
112 HCBA woven carbon fiber without pretreatment, or Pt-coated Toray carbon paper without
113 pretreatment. The membrane is pre-soaked (1 M KOH for 24 hours) Nafion 212.

114 *Undivided electrolytic cell setup (electrochemical oxidation vs. the HER)*

115 Working electrode: carbon felt, where **DPivOHAC(COO⁻)** was oxidized to **DPivOHAQ(COO⁻)**;
116 counter electrode: carbon rod, where water was reduced to hydrogen gas. While the electrolyte
117 was stirred, a constant potential (1.1 V vs. Ag/AgCl) was applied to the divided electrolytic cell
118 until 120% of the required coulombs were extracted from the working electrode.

119 *Divided electrolytic cell setup (electrochemical oxidation vs. the ORR)*

120 Anode: Commercial AvCarb HCBA (woven carbon cloth), where **DPivOHAC(COO⁻)** was
121 oxidized to **DPivOHAQ(COO⁻)**; cathode: platinum coated Toray carbon paper, where humidified
122 air/oxygen was reduced to hydroxide. A constant voltage (1.8 V) was applied to the divided
123 electrolytic cell until the current decreased to 2 mA/cm². The number of extracted electrons was
124 ~1.2 times higher than the theoretical value.

125 *Divided electrolytic cell setup (electrochemical oxidation vs. the reduction of ferricyanide)*

126 Anode: AvCarb HCBA (woven carbon cloth), where **DPivOHAC(COO⁻)** was oxidized to
127 **DPivOHAQ(COO⁻)**; cathode: AvCarb HCBA (woven carbon cloth), where potassium
128 ferricyanide was reduced to potassium ferrocyanide. A constant current density (20 mA/cm²) was
129 applied to the divided cell for at most 1.5 hours with a 1.2 V voltage cutoff; when either time or
130 voltage reached the limit, the potential was held (1.2 V vs. ferro-/ferricyanide) until the current
131 decreased to 2 mA/cm². The number of extracted electrons was ~1.2 times higher than the
132 theoretical value.

133 An aliquot (~250 μL) was transferred from the as-prepared anolyte to an Eppendorf® tube
134 (capacity: 1.5 mL) and acidified by a drop of concentrated HCl to obtain **DPivOHAQ** precipitate.
135 The final **DPivOHAQ** precipitate was re-dissolved in $\text{DMSO-}d_6$ for ^1H NMR measurement. The
136 yield was determined by peak integrations of spectrum. Faradaic efficiency (%) = yield (%) / 1.2.
137 More detailed information can be found in the Supplementary information.

138 **Results and Discussion**

139 In an electrolytic cell, an anodic oxidation half reaction must be accompanied by a cathodic
140 reduction half reaction. As shown in Table 1, we devise three different reduction half reactions to
141 be coupled with direct **DPivOHAC** electrochemical oxidation, *i.e.*, the hydrogen evolution
142 reaction (HER), the oxygen reduction reaction (ORR), and the $\text{Fe}(\text{CN})_6^{3-}$ to $\text{Fe}(\text{CN})_6^{4-}$ reduction
143 reaction. The corresponding oxidation or reduction potentials for these reactions are listed in Table
144 1.

145 For the electrochemical oxidation of **DPivOHAC** to **DPivOHAQ**, two cell types are used, as
146 diagramed and described in Figures S3 and S4. A divided cell uses an ion exchange membrane to
147 separate the two half reactions, resembling the architecture of traditional fuel cells and ARFBs.
148 An undivided cell employs two electrodes suspended in electrolyte without the use of a membrane,
149 reflecting a bulk electrolysis cell.

150 Comparing these three overall reactions, the first one paired with the HER requires the highest
151 voltage; the second one paired with the ORR is known to have slow reaction kinetics and a high
152 overpotential;³⁹ the third one paired with $\text{Fe}(\text{CN})_6^{3-}$ to $\text{Fe}(\text{CN})_6^{4-}$ reduction exhibits the lowest
153 overall reaction cell voltage, suggesting the least amount of energy will be required for
154 electrosynthesis. Another merit of the third reaction is the *in situ* generation of the desired negolyte
155 active species (**DPivOHAQ**) and posolyte active species $\text{Fe}(\text{CN})_6^{4-}$ simultaneously. The

156 disadvantage is that at least six equivalents of ferricyanide and hydroxide are used. Given the
 157 similar reduction potentials of the ORR and of ferricyanide to ferrocyanide, an important direction
 158 for future research is the concurrent reduction of oxygen and ferricyanide in order to achieve high
 159 yields as well as lower ferricyanide usage. By using the same full cell configuration without
 160 changing electrolyte reservoirs, carbon-based electrodes, or ion-exchange membranes, we can
 161 immediately switch from electrosynthesis mode to flow battery mode for electrochemical energy
 162 storage. In this configuration, neither hazardous oxidants nor purification steps are needed, nor is
 163 waste generated. Furthermore, the reaction may proceed at room temperature with high atom
 164 efficiency. The new synthesis is therefore potentially safe, green, economical, and scalable.

165 **Table 1.** Anodic, cathodic, and overall reactions for direct electrochemical oxidation.

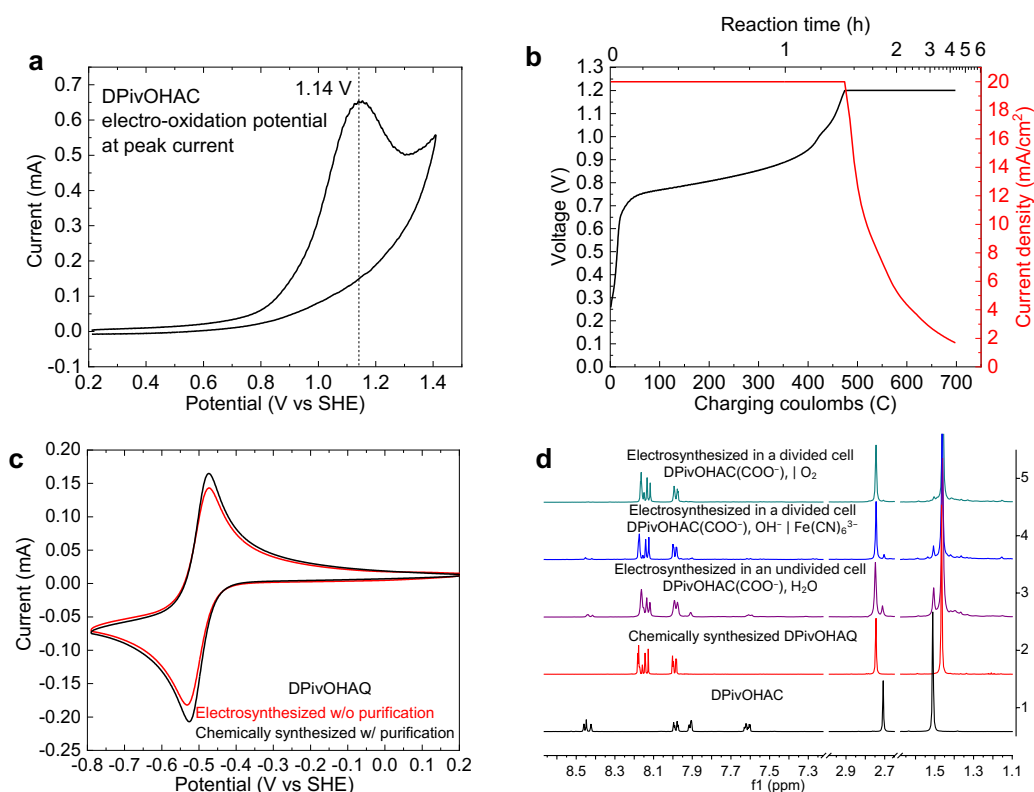
	Reactions	Potential at pH 14 (V vs SHE) / Cell voltage (V)
Anodic	$\text{DPivOHAC}(\text{COO}^-) + 6 \text{OH}^- \longrightarrow \text{DPivOHAQ}(\text{COO}^-) + 4 \text{H}_2\text{O} + 6 e^-$	1.14*
Cathodic	$6 \text{H}_2\text{O} + 6 e^- \longrightarrow 3 \text{H}_2 + 6 \text{OH}^-$ (divided or undivided cell)	-0.83
	$1.5 \text{O}_2 + 6 e^- + 3 \text{H}_2\text{O} \longrightarrow 6 \text{OH}^-$ (divided or undivided cell)	0.40
	$6 \text{Fe}(\text{CN})_6^{3-} + 6 e^- \longrightarrow 6 \text{Fe}(\text{CN})_6^{4-}$ (divided cell)	0.44
Overall	$\text{DPivOHAC}(\text{COO}^-) + 2 \text{H}_2\text{O} \longrightarrow \text{DPivOHAQ}(\text{COO}^-) + 3 \text{H}_2$	1.97
	$\text{DPivOHAC}(\text{COO}^-) + 1.5 \text{O}_2 \longrightarrow \text{DPivOHAQ}(\text{COO}^-) + \text{H}_2\text{O}$	0.74
	$\text{DPivOHAC}(\text{COO}^-) + 6 \text{OH}^- + 6 \text{Fe}(\text{CN})_6^{3-} \longrightarrow \text{DPivOHAQ}(\text{COO}^-) + 6 \text{Fe}(\text{CN})_6^{4-} + 4 \text{H}_2\text{O}$	0.70

166 *: The electro-oxidation potential at peak current

167 The cyclic voltammogram (CV) of **DPivOHAC** at pH 14 (Figure 3a) indicates a peak oxidation
 168 current at 1.14 V vs. SHE. This value is more positive than the standard redox potential of 0.40 V
 169 vs. SHE for the oxygen evolution reaction (OER), and we expect that the OER will be a major side
 170 reaction of electrosynthesis.

171 We then assembled a flow cell with **DPivOHAC** as the anolyte and $\text{K}_3\text{Fe}(\text{CN})_6$ as the catholyte.
 172 Galvanostatic electrolysis with a potentiostatic hold after reaching a potential limit of 1.2 V was
 173 performed for ~4.5 hours to complete the electrosynthesis. The OER side reaction, evidenced by

174 the observation of bubbles generated in the anolyte, precludes a faradaic efficiency of 100%. Thus,
 175 the number of electrons extracted from the anolyte was ~1.2 times higher than the theoretical
 176 number for complete conversion. A plateau appears at ~0.8 V against $K_3Fe(CN)_6$ (0.44 V vs. SHE)
 177 in the voltage profile (Figure 3b).



178

179 **Figure 3. Electrochemical synthesis and characterization of DPivOHAQ.** (a) The cyclic voltammogram
 180 (CV) of 0.1 M DPivOHAC in 1.0 M KCl + 1.0 M KOH aqueous solution. Scan rate: 0.1 V/s. (b)
 181 The electrochemical oxidation was conducted by using a constant current (20 mA/cm²) with a
 182 subsequent potential hold (1.2 V) until the current density decreased to 2 mA/cm². (c) CV of
 183 10 mM electro-synthesized DPivOHAQ (against Fe(CN)₆³⁻) without purification and 10 mM
 184 chemically synthesized DPivOHAQ with purification in 1 M KOH aqueous solutions,
 185 respectively. Scan rate: 0.1 V/s. (d) ¹H NMR spectra of (bottom to top): chemically synthesized
 186 DPivOHAC (black); chemically synthesized DPivOHAQ (red); electro-synthesized DPivOHAQ
 187 in an undivided cell (purple), 17.3% of DPivOHAC remained unreacted according to the
 188 integration, yield: 82.7%; electro-synthesized DPivOHAQ in a divided cell against Fe(CN)₆³⁻
 189 (blue), 7.0% of DPivOHAC remained unreacted according to the integration, yield: 93.0%;
 190 electro-synthesized DPivOHAQ in a divided cell against O₂ (green), 0 % of DPivOHAC remained
 191 unreacted according to the integration, yield: 100%. The deuterated solvent is DMSO-*d*₆, and the
 192 solvent peaks (DMSO and H₂O) were removed to better display the peaks of interest. The

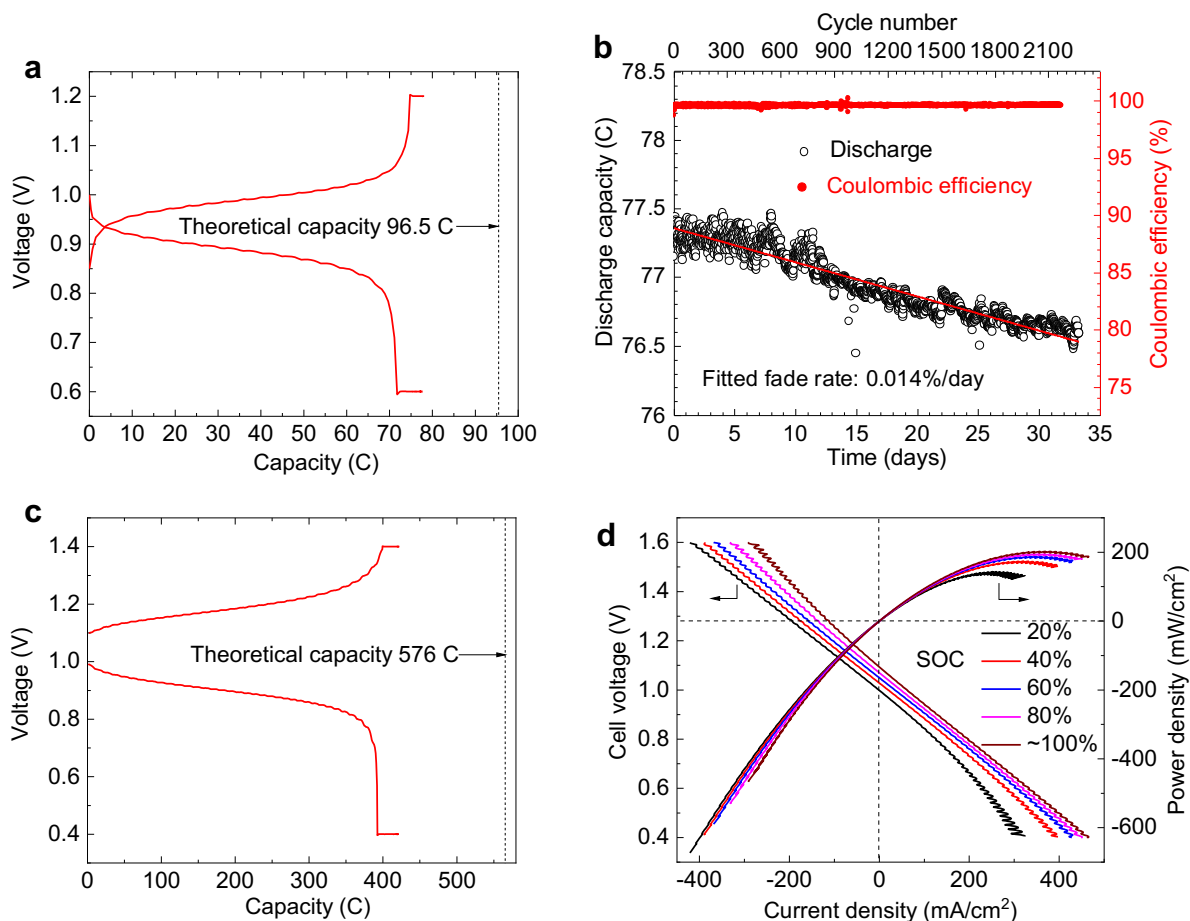
193 electrosynthetic details are described under the headings **Electrosynthesis I, II, and III** in the
194 Supporting Information.

195

196 We compared the CV of **DPivOHAQ** produced by electroynthesis against the reduction of
197 $\text{Fe}(\text{CN})_6^{3-}$ to that of the chemically synthesized product at the same concentration to verify that
198 the reaction products are the same regardless of the synthetic procedure employed (Figure 3c). The
199 two CV curves show identical redox peaks and similar peak currents, indicating a high-yield
200 electroynthesis process. ^1H nuclear magnetic resonance (NMR) spectroscopy was used to further
201 examine the structure of electrothesized **DPivOHAQ** when using either a divided or undivided
202 cell (Figure S3) and to compare the spectra with those of the starting material, **DPivOHAC**, and
203 the chemically synthesized **DPivOHAQ**. The top three spectra in Figure 3d are the ^1H NMR
204 spectra from electrothesized **DPivOHAQ**, in which the dominating peaks have the same
205 chemical shifts as those in the spectrum of chemically synthesized **DPivOHAQ**, further suggesting
206 the desired product was achieved.

207 Slightly different yields of **DPivOHAQ** were obtained when paired with the HER in an
208 undivided cell or with $\text{Fe}(\text{CN})_6^{3-}$ reduction or the ORR in a divided cell (Figure S4). The 82.7%
209 yield when paired with the HER in an undivided cell could be explained by a molecular shuttling
210 effect; *i.e.*, the electrothesized **DPivOHAQ** can first migrate to the cathode where it is reduced,
211 then diffuse back to the anode for re-oxidation. As a result, double counting of electrons can occur.
212 When paired with the $\text{Fe}(\text{CN})_6^{3-}$ reduction half reaction, a yield of 93.0% was obtained. The
213 incomplete yield is likely due to the consumption and therefore decreased concentration of both
214 **DPivOHAC** and OH^- as the electroynthesis continues, making further oxidation increasingly
215 difficult.

216 The use of the ORR half reaction achieved almost 100.0% yield. This exceptional yield may be
 217 attributed to the as-formed OH^- ions on the cathode (ORR) side crossing over to the anolyte and
 218 compensating for any loss of OH^- ions on the anode side. Overall yields in excess of 80.0% for all
 219 three conditions exceed many conventional reactions and are acceptable for direct flow battery use
 220 without purification or separation.



221
 222 **Figure 4. Full cell performance evaluation from Electrosynthesis III and IV.** (a) A
 223 representative charge–discharge profile with 0.1 M **DPivOHAQ**. Negolyte: 5 mL of 0.1 M
 224 **DPivOHAQ** pH = ~13.5. Posolyte: 100 mL of 0.1 M potassium ferro-/ferricyanide solution [~0.06
 225 M $\text{K}_4\text{Fe}(\text{CN})_6$ and ~0.04 M $\text{K}_3\text{Fe}(\text{CN})_6$] pH = ~13.6. (b) Discharge capacity (C) and coulombic
 226 efficiency (%) vs. cycle number and time (days). Negolyte: 4.5 mL of 0.1 M **DPivOHAQ**. Posolyte:
 227 100 mL of 0.1 M ferro-/ferricyanide solution [~0.06 M $\text{K}_4\text{Fe}(\text{CN})_6$ and ~0.04 M $\text{K}_3\text{Fe}(\text{CN})_6$].
 228 Current density: 30 mA/cm^2 with potential hold (cutoffs: 0.6 V, 1.2 V) until current decreased to
 229 2 mA/cm^2 . (c) A representative charge–discharge profile with 0.5 M **DPivOHAQ**. Negolyte: 6
 230 mL of 0.5 M **DPivOHAQ**. Posolyte: 100 mL of 0.5 M potassium ferro-/ferricyanide solution [~0.3

231 M $\text{K}_4\text{Fe}(\text{CN})_6$ and ~ 0.2 M $\text{K}_3\text{Fe}(\text{CN})_6$]. Current density: 100 mA/cm² with potential hold (cutoff:
232 0.4 V, 1.4 V) until current decreased to 2 mA/cm². (d) Polarization curves of the 0.5 M
233 **DPivOHAQ** at the SOC of 20%, 40%, 60%, 80%, and $\sim 100\%$ respectively. Descriptions of
234 **Electrosynthesis III** and **IV** can be found in the Supporting Information.

235
236 To demonstrate the feasibility of switching from the electrosynthesis mode (when paired with
237 $\text{Fe}(\text{CN})_6^{3-}$ reduction) to flow battery mode, we began charge–discharge cycling immediately upon
238 completion of the electrosynthesis, without performing any purification. Because other research
239 has reported that quinones and related compounds can decompose in the presence of light,^{40–42} we
240 wrapped the electrolyte reservoirs with aluminum foil to avoid light-induced decomposition during
241 cell cycling (Figures S13–S15). Figure 4a shows the charge–discharge profile of a single cycle
242 with an open circuit voltage of ~ 1.0 V and a capacity of 84.0 coulombs. Given the 93.0% yield
243 found from the ¹H NMR, the capacity utilization is 93.6%. Long-term cycling was then performed
244 to determine a temporal capacity fade rate of the full cell. Figure 4b demonstrates the discharge
245 capacity and coulombic efficiency over 33.2 days and 2271 cycles with a fitted fade rate of
246 0.014%/day and an average coulombic efficiency of 99.53%. This is consistent with the fade rate
247 of chemically synthesized **DPivOHAQ**.³⁸ The extremely low capacity fade rate is attributed to the
248 chemical stability of the molecular structure. The C–C covalent bond between the anthraquinone
249 core and the functionalizing chains is more robust in strong base and at elevated temperature than
250 the C–O bond demonstrated in previous work.^{4, 5, 38} Furthermore, the two branched methyl groups
251 on the carbon connected to the anthraquinone (AQ) core may increase the stability of the
252 solubilizing chain even when exposed to harsh conditions.¹⁵

253 To examine the feasibility of this method for potential industry use, we further conducted
254 electrosynthesis with a higher concentration (0.5 M) of **DPivOHAC** at a higher current density
255 (100 mA/cm²) (See Figure S5). Figure 4c shows that 0.5 M electrosynthesized negolyte can deliver
256 72.9% of the theoretical capacity. We attribute the discrepancy between the delivered capacity and

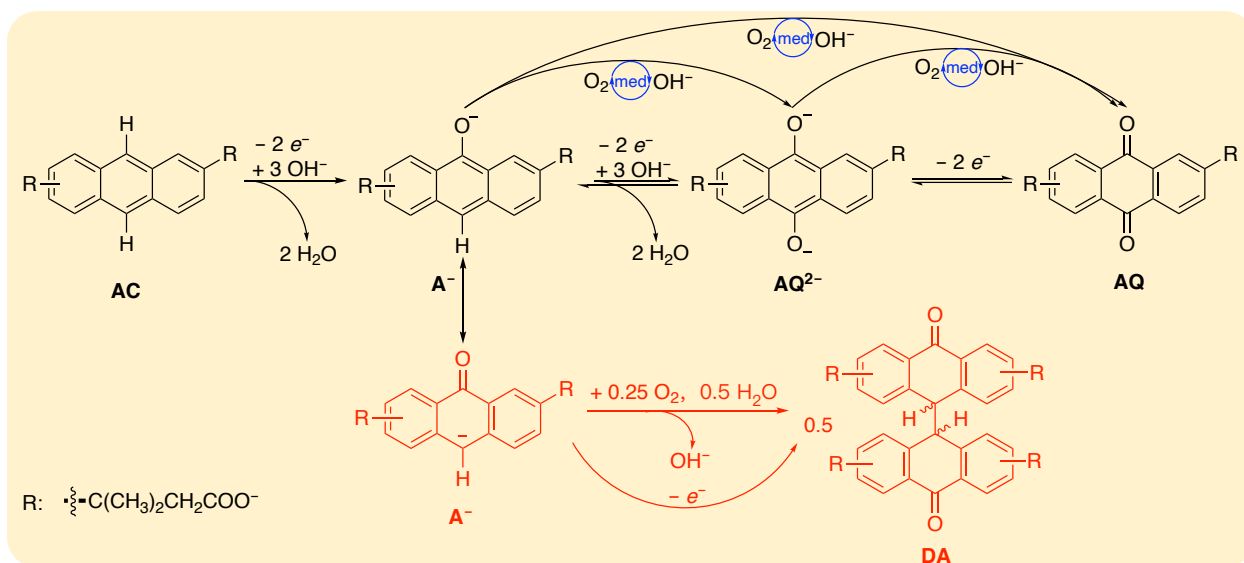
257 the theoretical capacity primarily to incomplete conversion (Figure S6). The capacity utilization is
258 81.9% if we consider that there is 11.0% unreacted **DPivOHAC(COO⁻)** in the negolyte.
259 Additionally, the mass transport of active species at 0.5 M concentration may be another issue
260 limiting the full capacity utilization. The corresponding polarization curve at different states of
261 charge (SOC) is shown in Figure 4d. The peak power density exceeds 0.2 W/cm² when at ~100 %
262 of SOC.

263 Given the total transfer of six electrons during the electrosynthesis of **DPivOHAC** to
264 **DPivOHAQ**, the high yields achieved in this work might be surprising. We hypothesize a three-
265 step successive two-electron transfer mechanism^{34,35}: first, when a potential is applied, anthracene
266 (**AC**) may react with three OH⁻ ions and donate two electrons to produce two water molecules and
267 the anthrone anion (**A⁻**); second, **A⁻** may further react with another three OH⁻ ions and donate
268 another two electrons to generate two water molecules and the deprotonated anthrahydroquinone
269 dianion (**AQ²⁻**); third, **AQ²⁻** may further release two electrons to afford the anthraquinone species
270 (**AQ**). Complete electrochemical conversion in the third step has been well-documented at
271 negative potentials vs. ferro-/ferricyanide^{1,2,43} and should therefore be rapid at positive potentials
272 vs. ferro-/ferricyanide. The reverse reaction of the second step has recently been identified as a
273 side reaction in ARFBs, and the forward reaction is chemically feasible when exposed to O₂ or
274 air.^{14,38} Given the high voltage applied to the cell, it is thus plausible that the forward reactions
275 (**AC** to **A⁻** to **AQ²⁻**/**AQ**) can electrochemically proceed completely and swiftly.

276 Our group has also previously proposed a side reaction pathway for anthraquinones,^{14,44} where
277 the anthrone anion (**A⁻**) can be oxidatively dimerized to dianthrone (**DA**) chemically and/or
278 electrochemically. According to ¹H NMR spectra (Figure 3d) and liquid chromatography–mass
279 spectrometry (LC–MS) results (Figure S7), neither **DA** nor Kolbe electrolysis-related byproducts⁴⁵

280 were detected (Scheme S1), suggesting that **AC/AQ**-related side reactions can be negligible when
281 a sufficient OH^- concentration is present to prevent dianthrone formation and a sufficiently low
282 voltage cutoff is chosen to prevent Kolbe electrolysis dimer formation. The major competing side
283 reaction is the OER, which, along with the reactions of **AC** to A^- to AQ^{2-} , will consume OH^- and
284 may lead to the formation of **DA** as a result of insufficient OH^- ions in the **DPivOHAC** solution
285 (see **Electrosynthesis V** in the SI). Interestingly, the dianthrone (Scheme S2), detected by
286 LC–MS (Figure S11), are surprisingly redox-active when a broad voltage window is applied
287 (Figures S8 and S9 and Scheme S2). On the one hand, the OER can reduce faradaic efficiency; on
288 the other hand, the generated oxygen can serve as a mediator and chemically oxidize intermediates
289 (*i.e.*, A^- , AQ^{2-}) to the final **AQ** form, *i.e.*, mediated (indirect) electrochemical oxidation. Because
290 the entire process involves not only electrochemical oxidations, but also chemical oxidations, it is
291 more appropriate to call it an electrochemical–chemical oxidation process.⁴⁶

292 In the proposed mechanism, the anthrone derivative is an intermediate in the electrochemical
293 oxidation. Anthrone formation has been identified as the major side reaction causing capacity fade
294 in previous work;^{14, 38} therefore, it is plausible that lost capacity of anthraquinone flow battery
295 systems may be recovered and anthraquinone lifetime extended by electrochemically oxidizing
296 anthrone to redox-active anthraquinone derivatives.



297

298 **Scheme 1. Proposed electrochemical oxidation mechanism.** Three-step successive two-electron
 299 transfer process from AC to A⁻, A⁻ to AQ²⁻, and AQ²⁻ to AQ. The generated oxygen from the
 300 OER side reaction may incur chemical oxidation processes including A⁻ to AQ²⁻, AQ²⁻ to AQ,
 301 and oxidative dimerization (A⁻ to DA).

302

303 To demonstrate that the electrochemical oxidation can be applied to other anthracene
 304 derivatives, we performed electrochemical oxidation of 4,4'-(9,10-dihydroanthracene-
 305 diyl)dibutanoic acid (**DBDHAC**), where the molecular core is 9,10-dihydroanthracene.³⁸ The ¹H
 306 NMR results indicate that **DBDHAC** can, like **DPivOHAC**, be electrochemically oxidized to the
 307 final anthraquinone (Figure S12), **DBAQ** (4,4'-(9,10-anthraquinone-diyl)dibutanoic acid), which
 308 has also been shown to be extremely stable.³⁸

309 The shared precursor of **DPivOHAQ** and **DBAQ**, anthracene, is abundant in crude petroleum
 310 and coal tar, and can be synthesized from benzene and benzyl alcohol (Scheme S3).⁴⁷ The
 311 precursor of **DPivOHAQ**, 3,3'-dimethyl acrylic acid, can be industrially produced from malonic
 312 acid, a food acid; the precursor of **DBAQ**, succinic anhydride, can be industrially hydrogenated
 313 from maleic anhydride and used as an important intermediate on an industrial scale. Thus, both
 314 **DPivOHAQ** and **DBAQ** can be readily synthesized from commodity chemicals. Although the
 315 synthetic cost of **DPivOHAQ** or **DBAQ** should be somewhat higher than that of 2,6-

316 dihydroxyanthraquinone (**DHAQ**) due to more steps and more chemicals involved, the capital cost
317 of AORFBs that utilize finite-lifetime electrolytes can be viewed as including the total active cost,
318 which is the sum of the initial cost of redox-active materials and the present value of the future
319 costs of periodic electrolyte replacement.¹³ This can lead to an initial cost—lifetime trade-off in
320 the choice of electrolytes. Over an extended operational lifetime, the total active cost of
321 **DPivOHAQ** or **DBAQ** may be less than that of **DHAQ** due to their much longer lifetimes.¹⁴

322 **Conclusion**

324 This work demonstrates a potentially scalable, safe, green, and economical *in situ*
325 electrosynthetic method for anthraquinone electrolytes in a flow cell without the use of hazardous
326 oxidants or precious metal catalysts. The as-generated electrolytes, which are extremely stable,
327 can be immediately used in a redox flow battery without separation or purification. Other low-cost
328 compounds may also be amenable to this approach, providing a pathway to lower the cost of
329 electrochemical grid storage systems, thereby accelerating the development of a renewable energy
330 economy. The technique extends the opportunities for direct aqueous electrosynthesis to replace
331 thermochemical synthesis of value-added organics.

332 **Supplementary Information**

333 Supplementary Information can be found with this article online at

334 **Acknowledgments**

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338 NSF Graduate Research Fellowship Program, no. DGE1144152 and DGE1745303.

339 **Declaration of Interests**

340 Harvard University has filed a patent application on the materials and the electrosynthetic methods
341 described in this paper.

342

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417

Supplementary Information

***In situ* Electrosynthesis of Anthraquinone Electrolytes in Aqueous Flow Batteries**

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61	hν) and under a 500 W lamp (+ hν) and of (b) DBAQ (0.1 M, pH 12) stored for 1 week in the	
62	absence of light (– hν) and under a 500 W lamp (+ hν). Differences in color were observed	
63	between the two samples of each compound. The formation of a film was also observed in the	
64	DPivOHAQ sample exposed to light.	20
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69	of light (– hν) and under a 500 W lamp (+ hν), each diluted (1:5) in pH 12 D ₂ O or in DMSO- <i>d</i> ₆ . ..	22
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71 **Scheme S3. Complete synthetic routes, conditions, and yields of DPivOHAQ and DBAQ when**
72 **commercially available commodity chemicals are used as starting materials. 23**

73 **Figure S16. ¹H NMR spectra of commercial and synthesized anthracene (AC) in DMSO-*d*₆. The**
74 **peak at 7.37 ppm in the synthesized AC spectrum is from benzene. 23**

75

76 **General information for synthesis and characterization**

77 All reagents were purchased from Sigma-Aldrich or Alfa Aesar and used as received unless
78 otherwise stated. All reactions sensitive to moisture or oxygen were carried out in oven-dried or
79 flame-dried and nitrogen-charged glassware. All anhydrous solvents were saturated with argon
80 and passed through a column of activated alumina immediately prior to use.

81

82 ¹H NMR spectra were recorded on Varian INOVA 500 spectrometers at 500 MHz. NMR spectra
83 were recorded in solutions of deuterated dimethyl sulfoxide (DMSO-*d*₆) with the residual dimethyl
84 sulfoxide (δ 2.25 ppm for ¹H NMR), or deuterated water (D₂O) with the residual H₂O (δ 4.79 ppm
85 for ¹H NMR).

86

87 LC-MS was conducted on a Bruker microTOF-Q II mass spectrometer. The sample was diluted
88 by water/acetonitrile (V/V = 1:1) to the desired concentration (~20 μ M) before LC-MS
89 measurements.

90

91 **Electrochemical characterization**

92 *Cyclic voltammetry measurements*

93 Glassy carbon was used as the working electrode for all three-electrode CV tests with a 5 mm
94 diameter glassy carbon working electrode, an Ag/AgCl reference electrode (BASi, pre-soaked in
95 3 M NaCl solution), and a graphite counter electrode.

96 All electrochemical oxidation and flow cell cycling was conducted with Biologic equipment and
97 corresponding software.

98

99 *Flow cell setup*

100 Flow battery experiments were constructed with cell hardware from Fuel Cell Tech (Albuquerque,
101 NM) assembled into a zero-gap flow cell configuration. Pyrosealed POCO graphite flow plates
102 with serpentine flow patterns were used for both electrodes. Each electrode comprised a 5 cm²
103 geometric surface area covered by a piece of AvCarb HCBA woven carbon fiber. The membrane
104 is pre-soaked (1 M KOH for 24 hours) Nafion 212.

105

106 **Brief description of electrosynthesis**

107 *Undivided electrolytic cell setup*

108 Working electrode: carbon felt, where **DPivOHAC(COO⁻)** was oxidized to **DPivOHAQ(COO⁻)**;
109 counter electrode: carbon rod, where water was reduced to hydrogen gas.

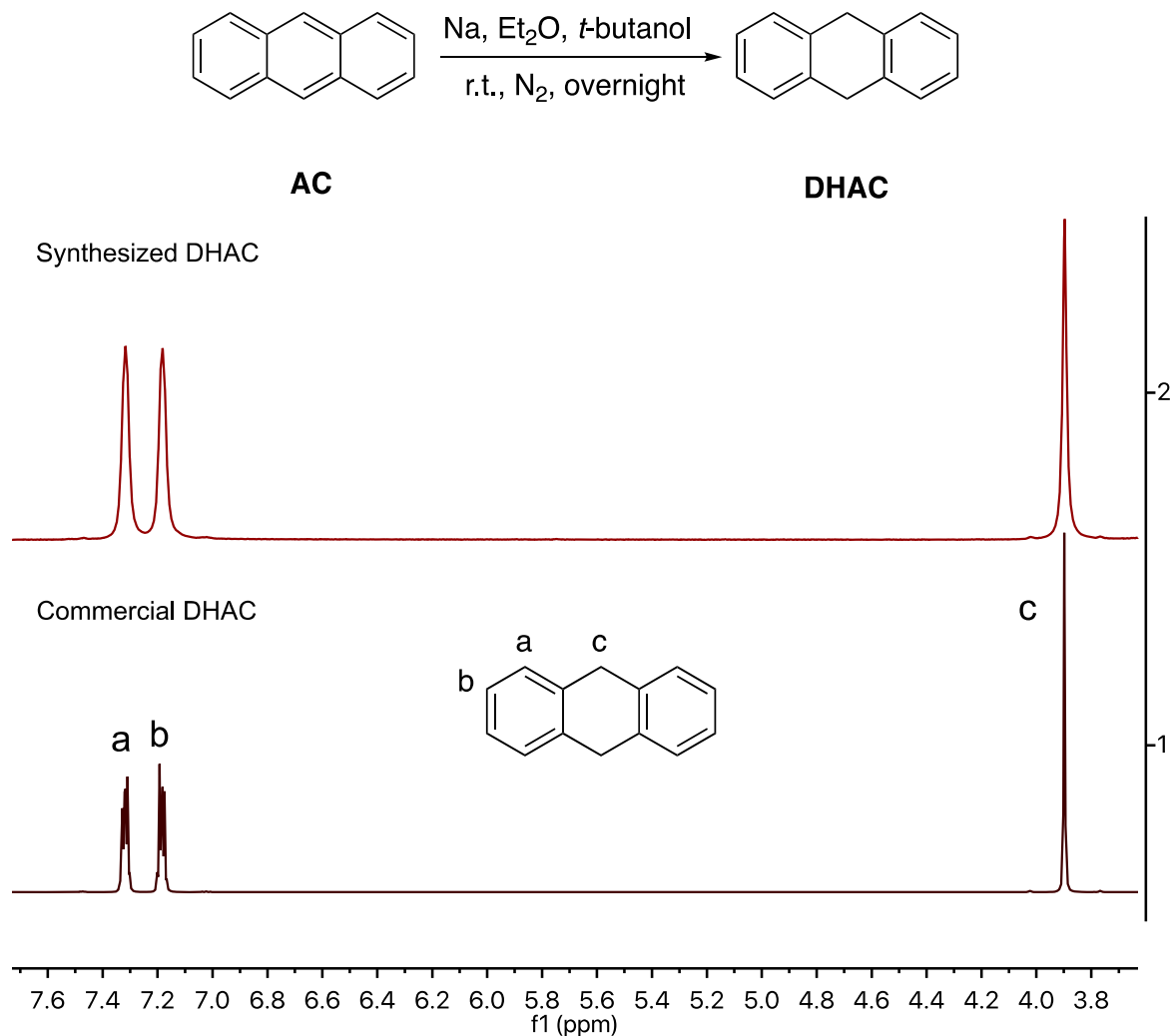
110

111 *Divided electrolytic cell setup vs. the ORR*

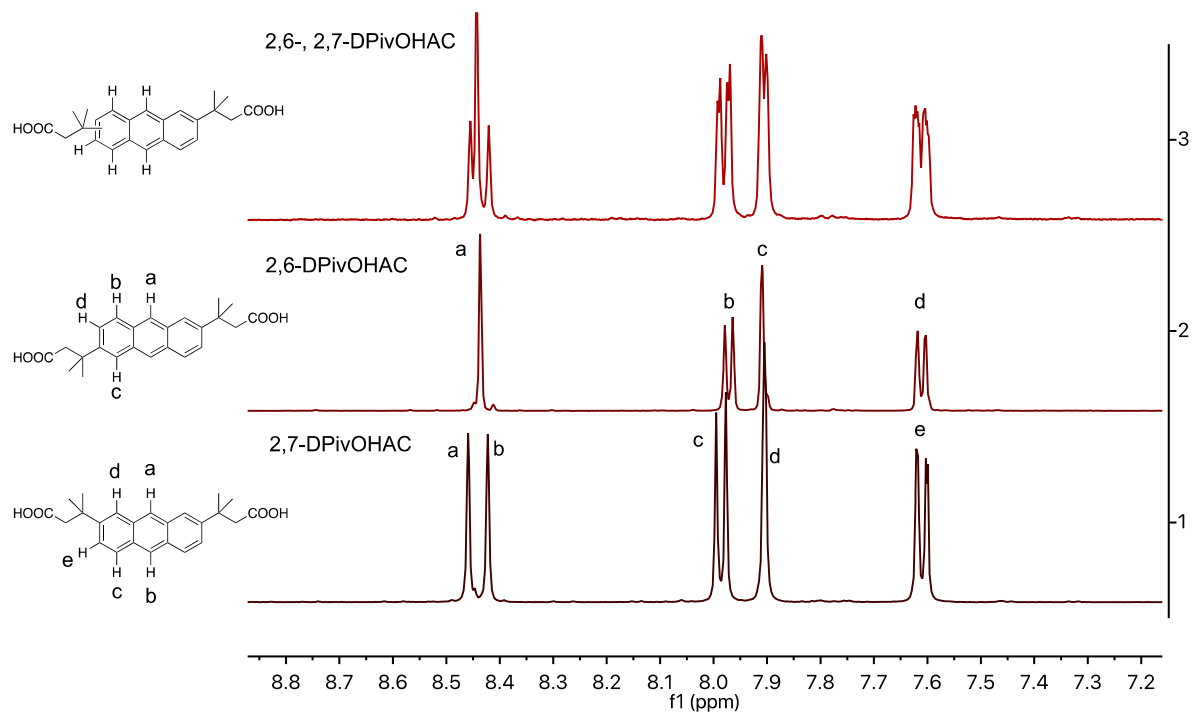
112 Anode: Commercial AvCarb HCBA (woven carbon cloth), where **DPivOHAC(COO⁻)** was
113 oxidized to **DPivOHAQ(COO⁻)**; cathode: platinum coated carbon paper (SGL 39AA), where
114 humidified air/oxygen was reduced to hydroxide.

115

116 *Divided electrolytic cell setup vs. the reduction of ferricyanide*
117 Anode: AvCarb HCBA (woven carbon cloth), where **DPivOHAC(COO⁻)** was oxidized to
118 **DPivOHAQ(COO⁻)**; cathode: AvCarb HCBA (woven carbon cloth), where potassium
119 ferricyanide was reduced to potassium ferrocyanide.
120

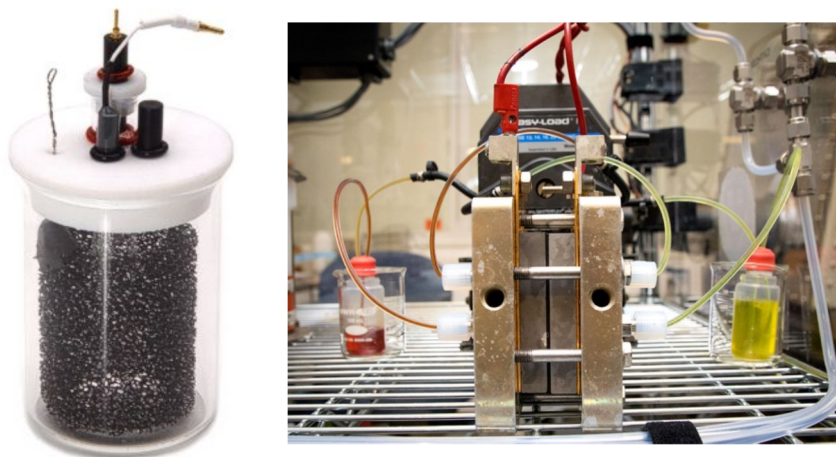


121
122 **Figure S1.** ¹H NMR spectra of commercial and synthesized 9,10-dihydroanthracene (**DHAC**) in
123 DMSO-*d*₆.
124



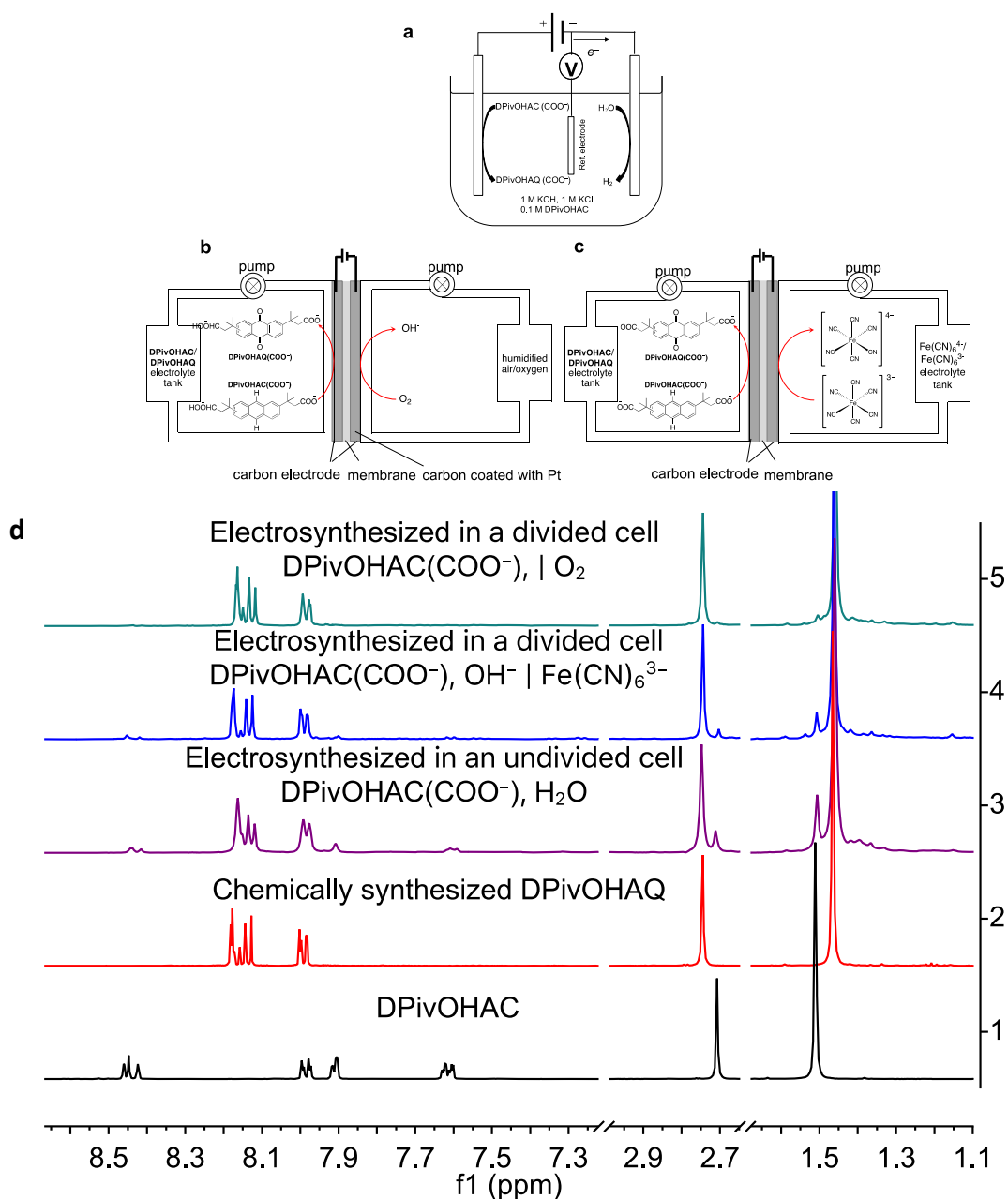
125
 126 **Figure S2.** ^1H NMR spectra of 2,7-, 2,6-DPivOHAC isomers and their mixture (aromatic region)
 127 in $\text{DMSO-}d_6$.
 128

a Undivided cell **b** Divided cell



129
 130 **Figure S3.** Photos of undivided cell (a) and divided cell (b).
 131

132

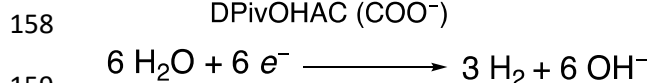
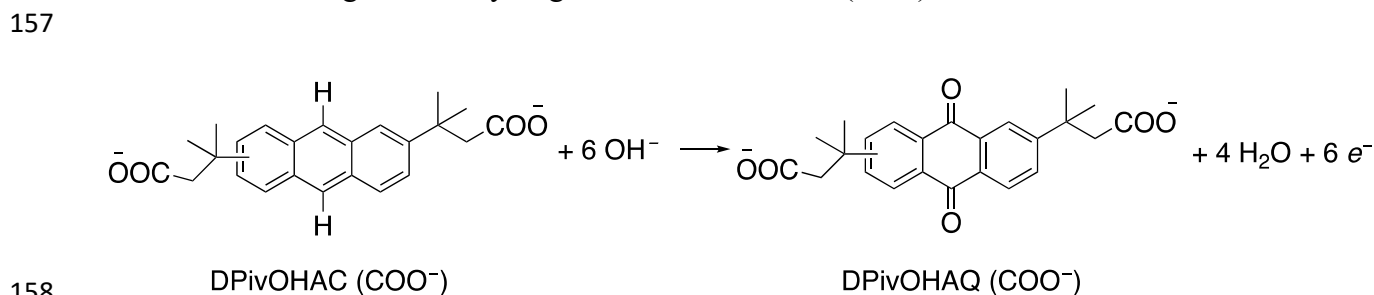


133
 134 **Figure S4.** Schematics of (a) undivided cell against the HER and divided cells against (b) the ORR
 135 and (c) ferricyanide to ferrocyanide, respectively. (d) ¹H NMR spectra of (bottom to top):
 136 chemically synthesized **DPivOHAC** (black); chemically synthesized **DPivOHAQ** (red);
 137 electrosynthesized **DPivOHAQ** in an undivided cell (purple), 17.3% of **DPivOHAC** remained
 138 unreacted according to the integration, yield: 82.7%; electro-synthesized **DPivOHAQ** in a divided
 139 cell against Fe(CN)₆³⁻ (blue), 7.0% of **DPivOHAC** remained unreacted according to the
 140 integration, yield: 93.0%; electro-synthesized **DPivOHAQ** in a divided cell against O₂ (green), 0 %
 141 of **DPivOHAC** remained unreacted according to the integration, yield: 100%. The deuterated
 142 solvent is DMSO-*d*₆, and the solvent peaks (DMSO and H₂O) were removed to better display the
 143 peaks of interest. The electrochemical details are described under the headings **Electrosynthesis**
 144 **I, II, and III.**

145

146 No ion-selective membrane is needed in the undivided cell (against the HER), nor are hydroxides
 147 required theoretically because the HER generates the required number of hydroxides for
 148 **DPivOHAQ** electrosynthesis. Ideally, the divided cell against the ORR will not require hydroxides
 149 either if all generated hydroxides from the ORR can immediately crossover to the **DPivOHAC**
 150 anolyte side. The divided cell against ferri- to ferrocyanide reduction needs six equivalents of
 151 hydroxide for electrosynthesis, the advantage of which is incorporating the electrosynthesis and
 152 flow battery in one setup, and electrosynthesis becomes a part of the on-site setup and takes as
 153 long as the energy/power ratio of the battery.

154
 155 **Electrosynthesis I.** Electrochemical synthesis of **DPivOHAQ(COO⁻)** in an undivided cell at 0.1
 156 M concentration, against the hydrogen evolution reaction (HER).



159
 160
 161 An undivided cell was prepared with carbon felt (XF30A, Toyobo Co., volumetric porosity: 95%)
 162 as the working electrode, a carbon rod as the counter electrode, and Ag/AgCl (3 M NaCl) as the
 163 reference electrode.

164
 165 Electrolyte preparation: 0.378 g **DPivOHAC**, 0.745 g KCl, and 0.561 g KOH were dissolved in
 166 deionized water to obtain a 10 mL solution containing 0.1 M **DPivOHAC**, 1.0 M KCl, and 1.0 M
 167 KOH.

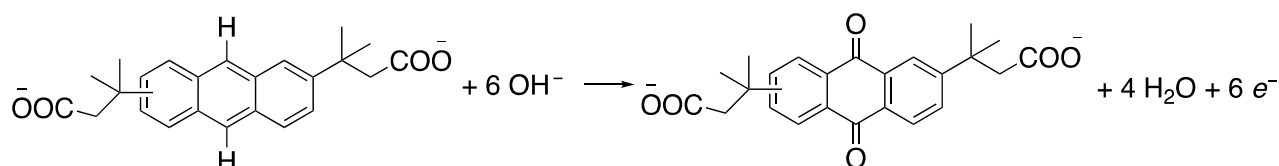
168
 169 On the working electrode: **DPivOHAC(COO⁻)** was oxidized to **DPivOHAQ(COO⁻)**; on the
 170 counter electrode: water was reduced to hydrogen gas.

171
 172 Electrochemical oxidation of **DPivOHAC(COO⁻)**: while the electrolyte was stirred, a constant
 173 potential (1.1 V vs. Ag/AgCl) was applied to the divided electrolytic cell until 120% of the required
 174 coulombs were extracted from the working electrode. [0.1 M * 0.01 L * 96485 C/mol * 6 * 1.2 =
 175 694.7 C, 6 electrons need to be extracted from every **DPivOHAC** molecule].

176
 177 Characterization of anolyte: an aliquot (~250 μL) was transferred from the as-prepared anolyte to
 178 an Eppendorf® tube (capacity: 1.5 mL) and acidified by a drop of concentrated HCl to obtain
 179 **DPivOHAQ** precipitate. The final **DPivOHAQ** precipitate was re-dissolved in DMSO-*d*₆ for ¹H
 180 NMR measurement. According to the integration of the ¹H NMR spectrum (Figure 3d), the yield
 181 is 82.7%. The faradaic efficiency (%) = [yield (%) / 1.2] = 68.9%.

182
 183

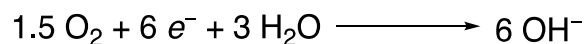
184 **Electrosynthesis II.** Electrochemical synthesis of **DPivOHAQ(COO⁻)** in a divided cell at 0.1 M
 185 concentration, against the oxygen reduction reaction (ORR).
 186



187

188

189



190 In a flow cell setup (divided electrolytic cell), where unbaked AvCarb HCBA was used on the
 191 anode side, the carbon paper was used on the cathode side with coated platinum particles to
 192 catalyze the ORR; Nafion® 212 was used as membrane. The high-frequency area specific
 193 resistance (HF-ASR) was maintained in the range of 1.48–1.54 $\Omega \text{ cm}^2$ before and after
 194 electrosynthesis.

195

196 Anolyte preparation: 0.378 g **DPivOHAC**, 0.745 g KCl, and 0.561 g KOH were dissolved in
 197 deionized water to obtain a 10 mL solution containing 0.1 M **DPivOHAC**, 1.0 M KCl, and 1.0 M
 198 KOH.

199

200 Catholyte preparation: humidified oxygen or air was pumped into the flow cell to participate in
 201 the electrochemical reaction.

202

203 Electrochemical oxidation of **DPivOHAC(COO⁻)**: a constant voltage (1.8 V) was applied to the
 204 divided electrolytic cell until the current decreased to 2 mA/cm². The number of extracted electrons
 205 was ~1.2 times higher than the theoretical value.

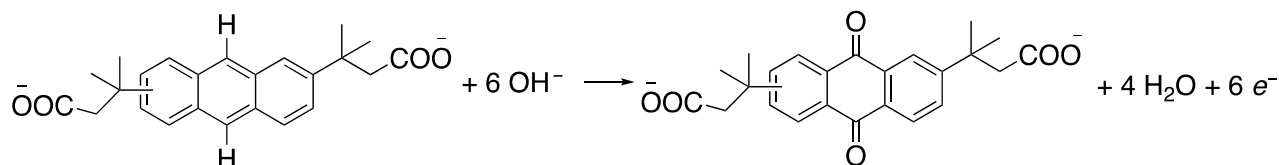
206

207 Characterization of anolyte: an aliquot (~250 μL) was transferred from the as-prepared anolyte to
 208 an Eppendorf® tube (capacity: 1.5 mL) and acidified by concentrated HCl to obtain **DPivOHAQ**
 209 precipitate. The final **DPivOHAQ** precipitate was re-dissolved in DMSO-*d*₆ for ¹H NMR
 210 measurement. According to the integration of the ¹H NMR spectrum (Figure 3d), the yield is 100%.
 211 The faradaic efficiency (%) = [yield (%) / 1.2] = 83.3%.

212

213 **Electrosynthesis III.** Electrochemical synthesis of **DPivOHAQ(COO⁻)** in a divided cell at 0.1 M
 214 concentration, against the reduction of ferricyanide.

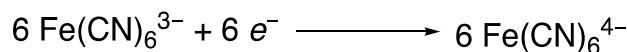
215



216

217

218



219 In a flow cell setup (divided electrolytic cell), unbaked AvCarb HCBA (woven carbon cloth) was
220 used as electrodes for both sides; Nafion® 212 was used as the membrane. The high-frequency
221 area specific resistance (HF-ASR) was maintained at $\sim 1.12 \Omega \text{ cm}^2$ before and after
222 electrosynthesis.

223
224 Anolyte preparation: 0.378 g **DPivOHAC**, 0.745 g KCl, and 0.561 g KOH were dissolved in
225 deionized water to obtain a 10 mL solution containing 0.1 M **DPivOHAC**, 1.0 M KCl, and 1.0 M
226 KOH.

227
228 Catholyte preparation: 3.292 g $\text{K}_3\text{Fe}(\text{CN})_6$, 7.445 g KCl, and 2.805 g KOH were dissolved in
229 deionized water to obtain a 100 mL solution containing 0.1 M $\text{K}_3\text{Fe}(\text{CN})_6$, 1.0 M KCl, and 0.5 M
230 KOH.

231
232 The reason for which 0.5 M KOH was added to the catholyte is to counterbalance the added OH^- in
233 the anolyte, which is required for the electrosynthesis, thereby suppressing the loss of OH^- from
234 the anolyte to the catholyte due to crossover.

235
236 Electrochemical oxidation of **DPivOHAC(COO⁻)**: a constant current density (20 mA/cm^2) was
237 applied to the divided cell for at most 1.5 hours with a 1.2 V voltage cutoff; when either time or
238 voltage reached the limit, the potential was held (1.2 V vs. ferro-/ferricyanide) until the current
239 decreased to 2 mA/cm^2 . The number of extracted electrons was ~ 1.2 times higher than the
240 theoretical value.

241
242 Characterization of anolyte: an aliquot ($\sim 250 \mu\text{L}$) was transferred from the as-prepared anolyte to
243 an Eppendorf® tube (capacity: 1.5 mL) and acidified by a drop of concentrated HCl to obtain
244 **DPivOHAQ** precipitate. The final **DPivOHAQ** precipitate was re-dissolved in $\text{DMSO-}d_6$ for ^1H
245 NMR measurement. According to the integration of the ^1H NMR spectrum (Figure 3d), the yield
246 is 93.0%. The faradaic efficiency (%) = [yield (%) / 1.2] = 77.5%.

247
248 Because a few aliquots were transferred and the volume of as-prepared **DPivOHAQ** changed, 5
249 mL of the **DPivOHAQ** solution was used as the negolyte and 100 mL of the ferro-/ferricyanide
250 solution [$\sim 0.06 \text{ M K}_4\text{Fe}(\text{CN})_6$ and $\sim 0.04 \text{ M K}_3\text{Fe}(\text{CN})_6$] generated from **Electrosynthesis III** was
251 used as the posolyte for charge-discharge cycling. Due to leakage, 4.5 mL of **DPivOHAQ**
252 remained for subsequent cycling.

253
254 **Electrosynthesis IV.** Electrochemical synthesis of **DPivOHAQ(COO⁻)** in a divided cell at 0.5 M
255 concentration with excess hydroxide, against the reduction of ferricyanide.

256
257 In a flow cell setup (divided electrolytic cell), unbaked AvCarb HCBA (woven carbon cloth) was
258 used as electrodes for both sides; Nafion® 212 was used as the membrane. The high-frequency
259 area specific resistance (HF-ASR) was maintained at $\sim 1.1 \Omega \text{ cm}^2$ before and after electrosynthesis.

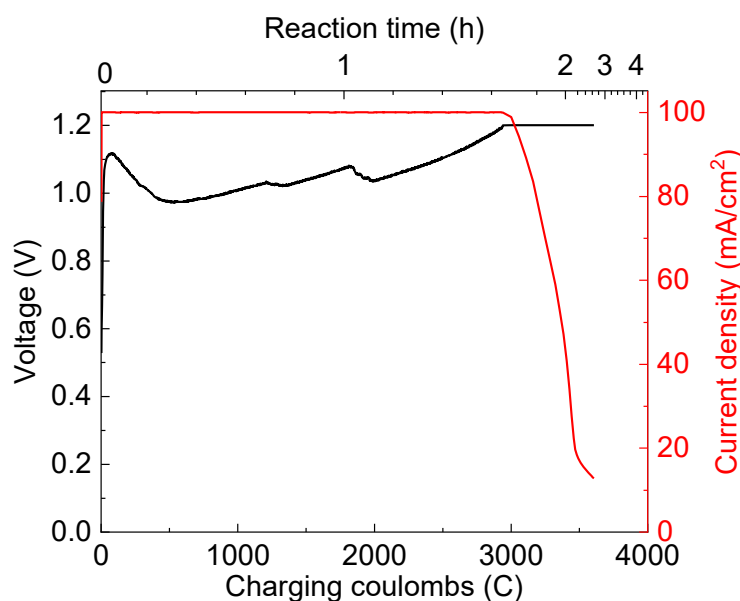
260
261 Anolyte preparation: 1.89 g **DPivOHAC**, 0.745 g KCl, and 0.567 g KOH were dissolved in
262 deionized water to obtain a 10 mL solution containing 0.5 M **DPivOHAC**, 1.0 M KCl, and 1.0 M
263 KOH. Although the **DPivOHAC** electrochemical oxidation requires OH^- ions, we observed that

264 0.5 M **DPivOHAC** tends to crash out of solution when the concentration of KOH exceeds 1.5 M.
265 To circumvent this precipitation issue, we added 1.5 times the required amount of KOH pellets
266 (2.52 g) (*i.e.*, 1.5 times 6 equivalents relative to **DPivOHAC**) into the analyte over the course of
267 constant current charging. According to the Nernst equation, the cell voltage is a function of [OH⁻];
268 thus, the voltage fluctuation reflects the addition of KOH in Figure S5.

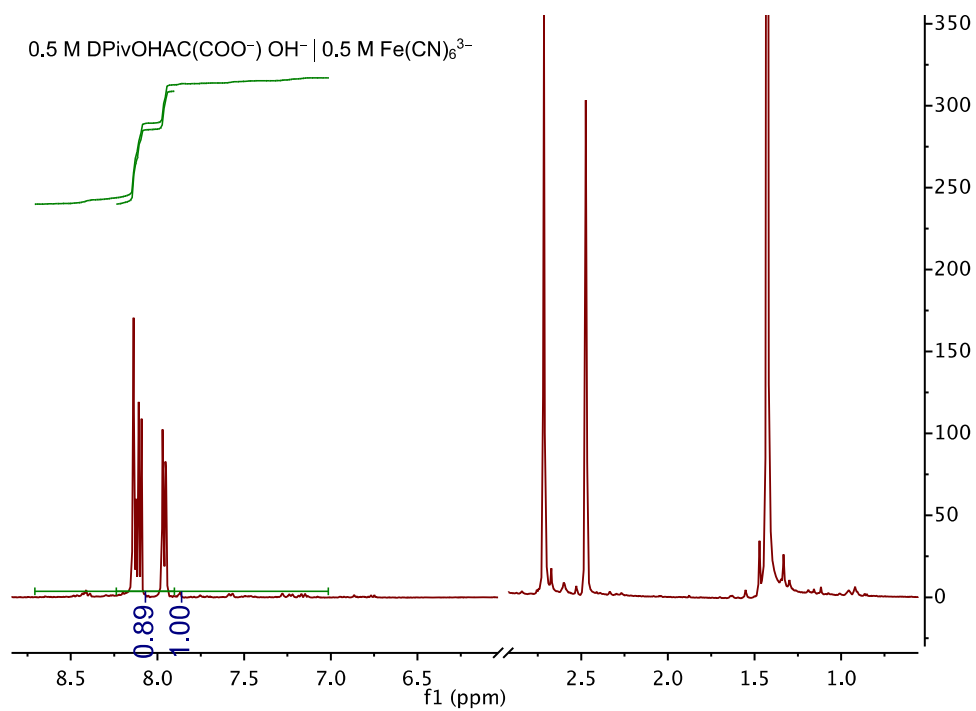
269
270 Catholyte preparation: 16.46 g K₃Fe(CN)₆, 7.445 g KCl, and 2.805 g KOH were dissolved in
271 deionized water to obtain a 100 mL solution containing 0.5 M K₃Fe(CN)₆, 1.0 M KCl, and 0.5 M
272 KOH.

273
274 Electrochemical oxidation of **DPivOHAC(COO⁻)**: a constant current density (100 mA/cm²) was
275 applied to the divided cell for at most 1.7 hours with a 1.2 V voltage cutoff; when either time or
276 voltage reached the limit, the potential was held (1.2 V vs. ferro-/ferricyanide) until the current
277 decreased to 12 mA/cm². The number of extracted electrons was ~1.2 times higher than the
278 theoretical value.

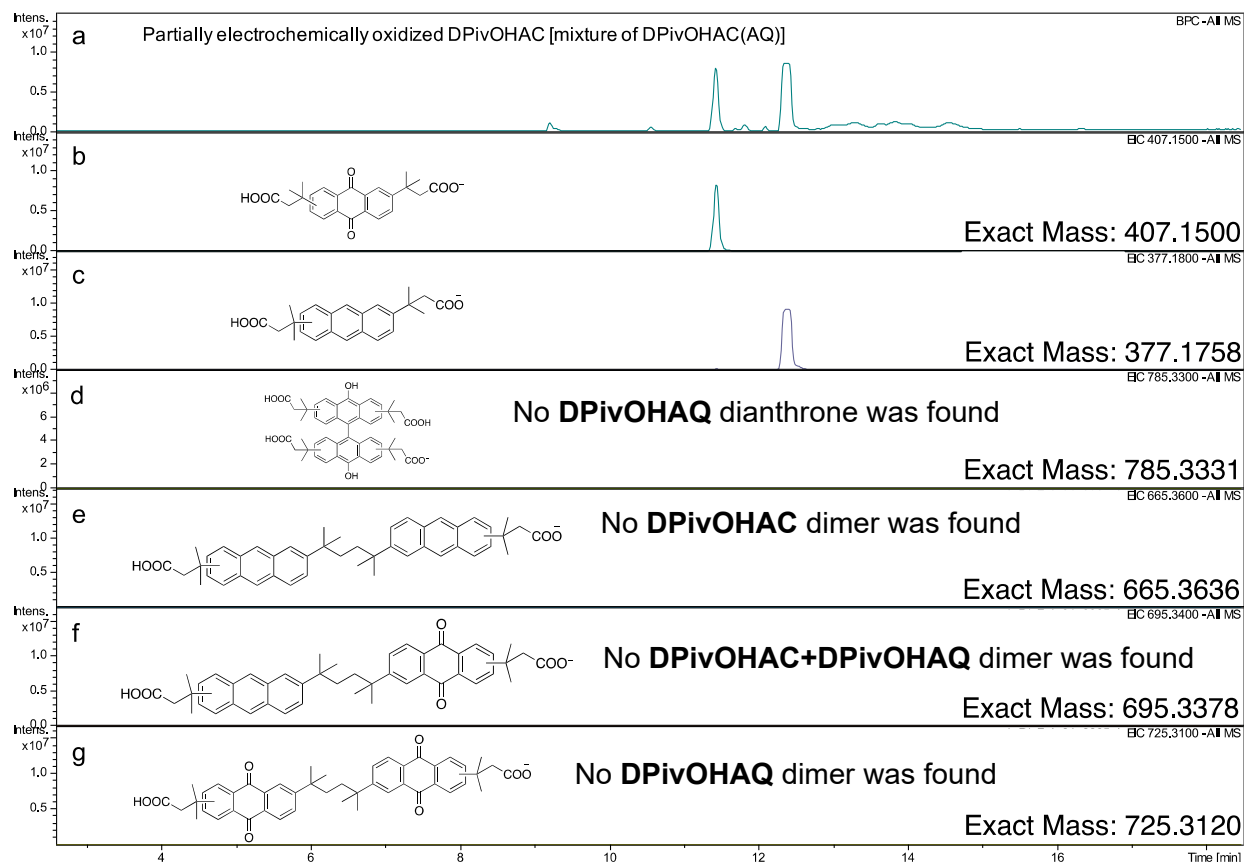
279
280 Characterization of analyte: an aliquot (~250 μL) was transferred from the as-prepared analyte to
281 an Eppendorf® tube (capacity: 1.5 mL) and acidified by a drop of concentrated HCl to obtain
282 **DPivOHAQ** precipitate. The final **DPivOHAQ** precipitate was re-dissolved in DMSO-*d*₆ for ¹H
283 NMR measurement; the yield is 89.0%. The faradaic efficiency (%) = [yield (%) / 1.2] = 74.2%.



284
285 **Figure S5.** The electrochemical oxidation of 0.5 M **DPivOHAC(COO⁻)** (**Electrosynthesis IV**).

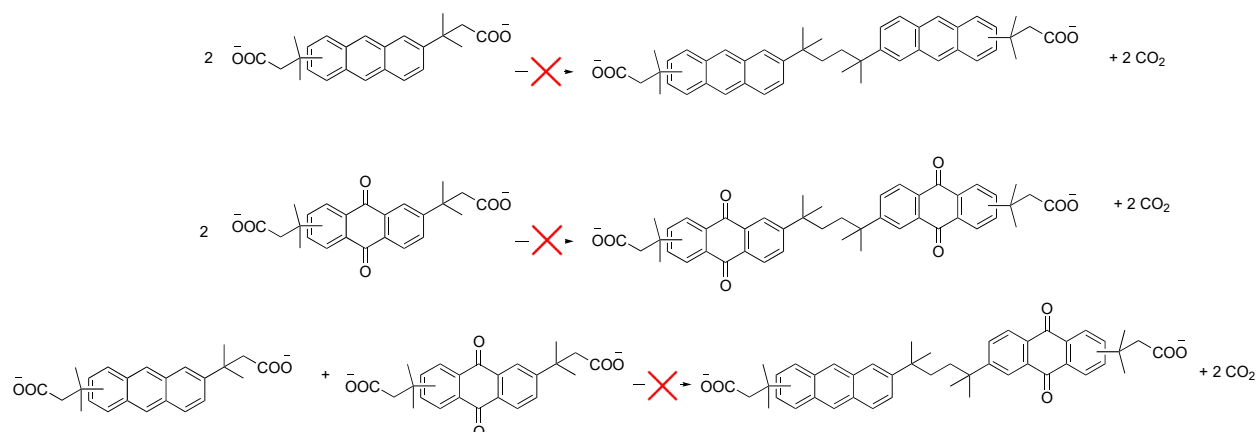


286
 287 **Figure S6.** ¹H NMR spectrum of **DPivOHAQ** in DMSO-*d*₆ synthesized via the procedure
 288 described in **Electrosynthesis IV**. From the aromatic peak integrations, we found that 89.0%
 289 **DPivOHAQ** was generated (when the two set of peaks at chemical shifts of 7.95 and 8.10 ppm
 290 were integrated), 11.0% **DPivOHAC** was remaining.
 291



292
 293 **Figure S7.** Mass spectra of partially electrochemically oxidized **DPivOHAQ** (from **Electrosynthesis IV**)
 294 measured by liquid chromatography–mass spectrometry (LC–MS). (a) The peak intensity and
 295 retention time of partially electrochemically oxidized **DPivOHAQ** under negative mode. (b) The peak
 296 intensity and retention time of **DPivOHAQ** under negative mode. (c) The peak intensity and
 297 retention time of **DPivOHAC** under negative mode. (d) The peak intensity and retention time of
 298 the **DPivOHAQ** dianthrone under negative mode. No peak was found in the given retention time
 299 region, which, in combination with the absence of impurities in the ^1H NMR spectra in Figures 3d
 300 and S6, indicates that no observable **DPivOHAQ** dianthrone was generated during the
 301 electro-synthesis. (e)-(f) The peak intensity and retention time of **DPivOHAQ(AC)**-related Kolbe
 302 electrolysis byproducts under negative mode. No peak was found in the given retention time region,
 303 which, in combination with the absence of impurities in the ^1H NMR spectra in Figures 3d and S6,
 304 indicates that no observable **DPivOHAQ(AC)**-related Kolbe electrolysis byproducts were
 305 generated during the electro-synthesis.

NO Kolbe electrolysis



306
307 **Scheme S1.** Kolbe electrolysis. Kolbe electrolysis-related byproducts are not expected in our cell,
308 as the decarboxylation and dimerization reactions usually require much higher voltages and
309 precious-metal-based electrodes.^{S1} In our cell, we use carbon electrodes and an applied potential
310 of 1.2 V. Additionally, we did not detect any dimer formation from LC–MS measurements.

311
312 **Electrosynthesis V.** Electrochemical synthesis of **DPivOHAQ(COO⁻)** in a divided cell at 0.5 M
313 concentration with a stoichiometric quantity of hydroxide, against the reduction of ferricyanide.

314
315 In a flow cell setup (divided electrolytic cell), unbaked AvCarb HCBA (woven carbon cloth) was
316 used as electrodes for both sides; Nafion® 212 was used as the membrane. The high-frequency
317 area specific resistance (HF–ASR) was maintained at $\sim 1.45 \Omega \text{ cm}^2$ before and after
318 electrosynthesis.

319
320 Anolyte preparation: 1.89 g **DPivOHAC**, 0.745 g KCl, and 0.567 g KOH were dissolved in
321 deionized water to obtain a 10 mL solution containing 0.5 M **DPivOHAC**, 1.0 M KCl, and 1.01
322 M KOH. We added the stoichiometric quantity of KOH pellets (1.68 g) (*i.e.*, 6 equivalents relative
323 to **DPivOHAC**) into the anolyte over the course of constant current charging.

324
325 Catholyte preparation: 16.46 g $\text{K}_3\text{Fe}(\text{CN})_6$, 7.445 g KCl, and 2.805 g KOH were dissolved in
326 deionized water to obtain a 100 mL solution containing 0.5 M $\text{K}_3\text{Fe}(\text{CN})_6$, 1.0 M KCl, and 0.5 M
327 KOH.

328
329 Electrochemical oxidation of **DPivOHAC(COO⁻)**: a constant current density (100 mA/cm^2) was
330 applied to the divided electrolytic cell for at most 1.7 hours with a 1.2 V voltage cutoff; when
331 either time or voltage reached the limit, the potential was held (1.2 V vs. ferro-/ferricyanide) until
332 the current decreased to 12 mA/cm^2 . The number of extracted electrons was ~ 1.2 times higher than
333 the theoretical value.

334
335 Characterization of anolyte: an aliquot ($\sim 250 \mu\text{L}$) was transferred from the as-prepared anolyte to
336 an Eppendorf® tube (capacity: 1.5 mL) and acidified by a drop of concentrated HCl to obtain
337 **DPivOHAQ** precipitate. The final **DPivOHAQ** precipitate was re-dissolved in $\text{DMSO-}d_6$ for ^1H
338 NMR measurement; the yield is 81.8%. The faradaic efficiency (%) = [yield (%) / 1.2] = 68.2%.

339

340 **Formation of dianthrone during electrosynthesis**

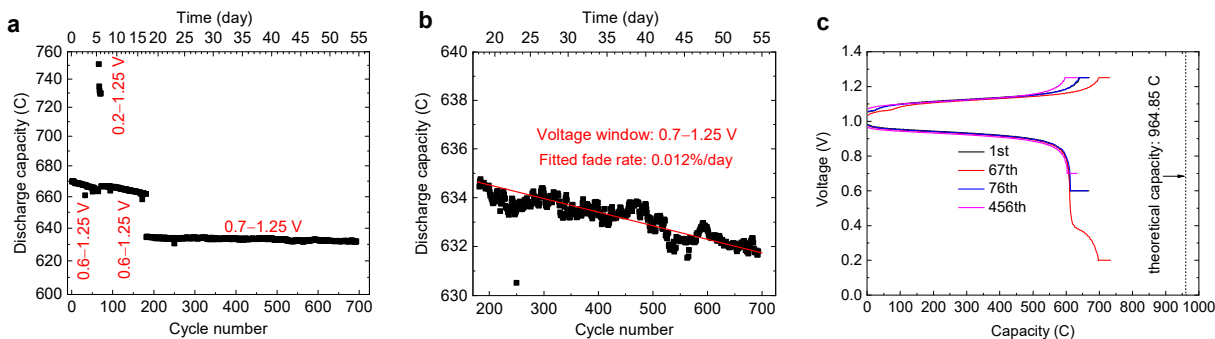
341 Anthrone dimers can be produced during the electrosynthesis when insufficient hydroxide is
342 present.

343
344 When there is excess hydroxide in the solution, although some OH^- ions will be electrochemically
345 oxidized to oxygen via the OER, the remaining OH^- ions are sufficient for the conversion of A^- to
346 AQ^{2-} .

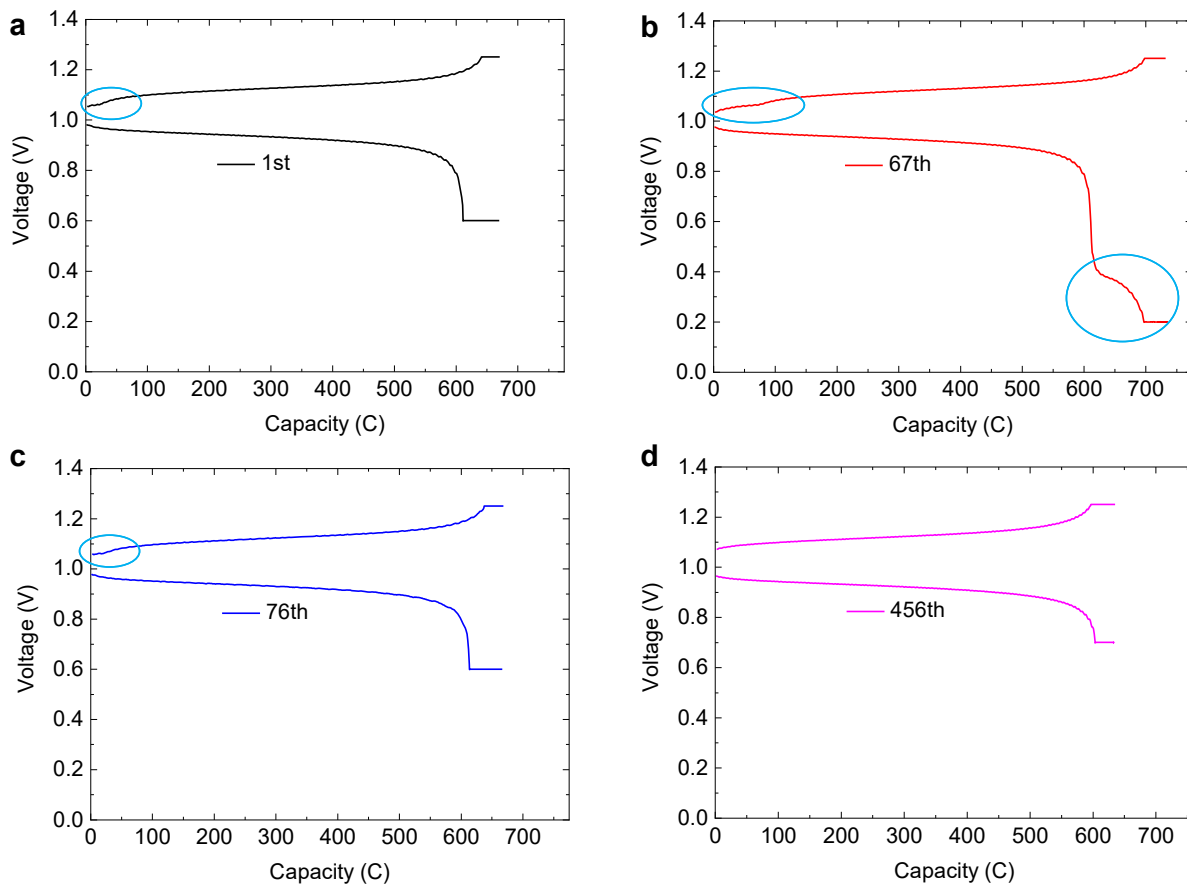
347
348 When there is no excess hydroxide, given that the OER side reaction is an inevitable competing
349 reaction, there will be insufficient OH^- ions for the conversion of A^- to AQ^{2-} ; instead, the
350 anthrone anion A^- may be oxidatively dimerized to the dianthrone **DA**. The following figures and
351 scheme illustrate how **DA** was identified and propose its corresponding electrochemistry.

352
353 During the electrochemical oxidation of the 10 mL 0.5 M **DPivOHAC(COO⁻)** at pH 12, only 1.68
354 g of KOH ($10 \times 0.001 \text{ L} \times 0.5 \text{ M} \times 6 \times 56.1056 \text{ g/mol} = 1.68 \text{ g}$) were added to the solution. Although
355 there is some additional KOH added to the potassium ferricyanide side, hydroxide cannot cross
356 over to the **DPivOHAC** side of the cell sufficiently rapidly to offset its consumption by
357 **DPivOHAC** oxidation and the OER.

358

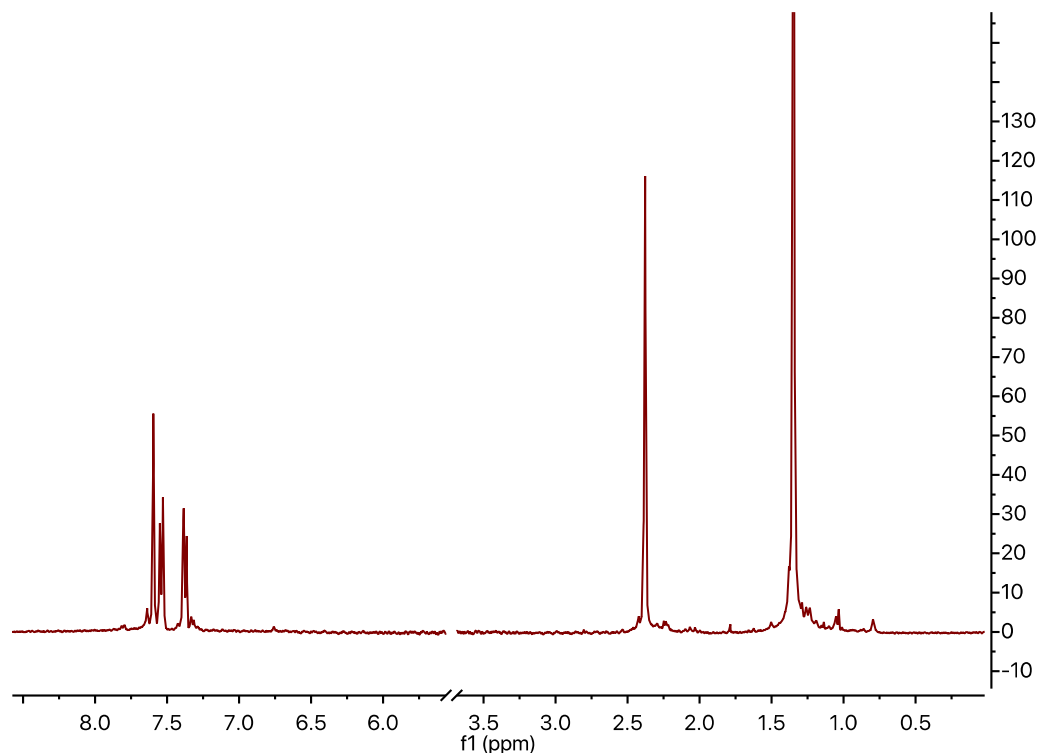


359
360 **Figure S8.** Cell performance of 0.5 M electrosynthesized **DPivOHAQ** when a stoichiometric
361 quantity of hydroxide was added into the **DPivOHAC(COO⁻)** solution (**Electrosynthesis V**). (a)
362 The long-term cycling performance with adjusted lower voltage cutoffs. (b) The zoomed in
363 discharge capacity when 0.7–1.25 V voltage cutoffs were applied; the fitted temporal fade rate
364 was 0.01%/day. (c) The voltage profiles at varying cycle numbers with different lower voltage
365 cutoffs.



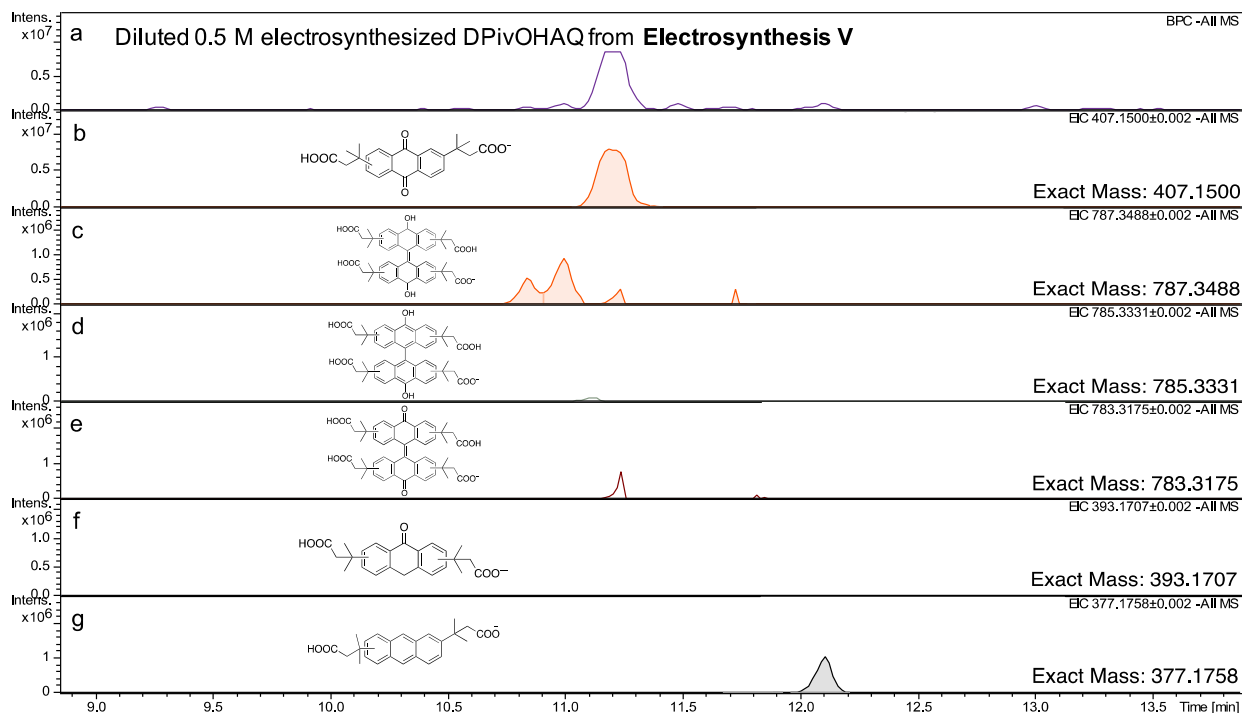
366
 367 **Figure S9.** Voltage profiles of 0.5 M electrosynthesized **DPivOHAQ** when a stoichiometric
 368 quantity of hydroxide was added into the **DPivOHAC** solution (**Electrosynthesis V**) with
 369 different lower voltage cutoffs [(a) 0.6, (b) 0.2, (c) 0.6, and (d) 0.7 V]. The upper voltage cutoff is
 370 kept constant at 1.25 V for the duration of cell cycling.

371
 372 In the 1st cycle, the region in (a) circled in cyan shows a small plateau, indicating some redox-
 373 active byproducts were produced during the electrosynthesis. In the 67th cycle, after lowering the
 374 lower cutoff from 0.6 to 0.2 V, we can clearly see the discharge plateau (in the region of 0.2–0.4
 375 V) attributed to byproducts, and the charge plateau attributed to byproducts is also becoming
 376 longer. In the 76th cycle, after elevating the lower cutoff back to 0.6 V, the shape of the charge
 377 profile becomes nearly the same as the one in the 1st cycle. After the lower voltage cutoff was
 378 further increased to 0.7 V, in the 456th cycle, the small plateau attributed to the byproducts
 379 disappeared.

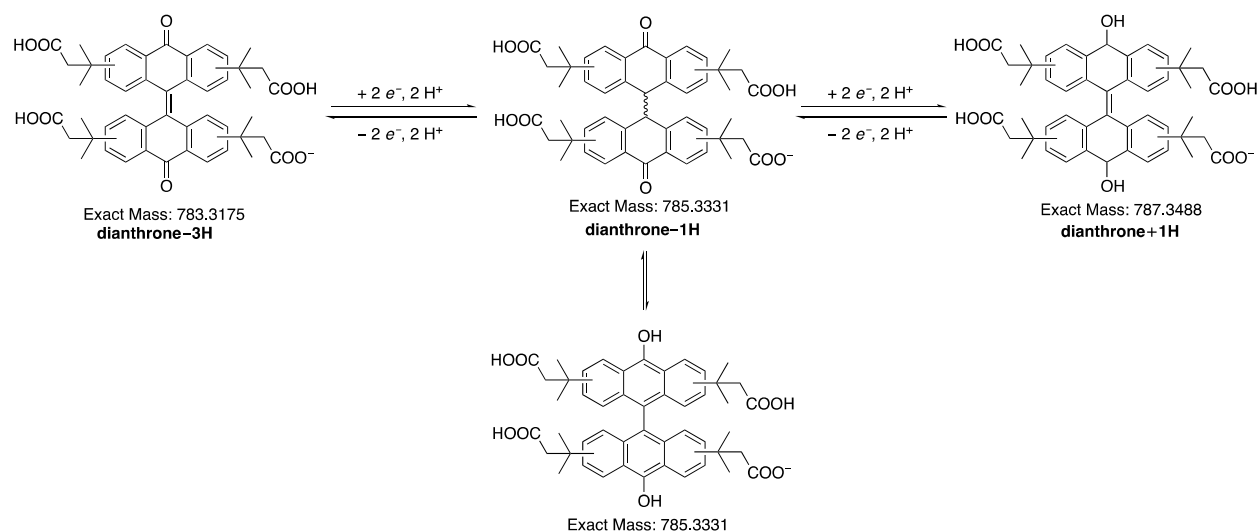


380
 381 **Figure S10.** ^1H NMR spectrum of cycled 0.5 M electrosynthesized **DPivOHAQ** when a
 382 stoichiometric quantity of hydroxide was added into the **DPivOHAC(COO $^-$)** solution
 383 (**Electrosynthesis V**). The solvent peak was removed to clearly show both aromatic and aliphatic
 384 regions of the cycled **DPivOHAQ** solution. The deuterated solvent is D_2O . The dominating peaks
 385 can be assigned to **DPivOHAQ**. Some small impurity peaks were observed, but they are difficult
 386 to identify. The percentages of side products are very close to the detection limit of the ^1H NMR
 387 instrument.

388
 389
 390

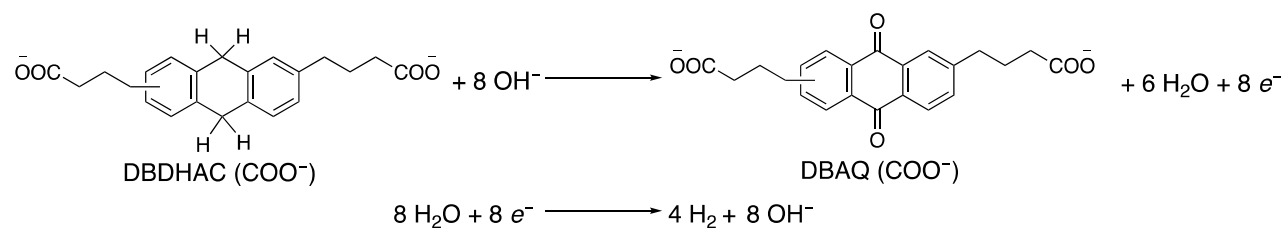


391
 392 **Figure S11.** LC-MS results of cycled 0.5 M electrosynthesized **DPivOHAQ** when a
 393 stoichiometric quantity of hydroxide was added into the **DPivOHAC** solution (**Electrosynthesis**
 394 **V**). (a) The base peak chromatogram of the sample, showing all peaks observed by mass
 395 spectrometry under negative mode. (b) The peak intensity and retention time of **DPivOHAQ-1H**
 396 under negative mode. (c) The peak intensity and retention time of **dianthrone+1H** under negative
 397 mode. (d) The peak intensity and retention time of **dianthrone-1H** under negative mode. (e) The
 398 peak intensity and retention time of **dianthrone-3H** under negative mode. (f) The peak intensity
 399 and retention time of **anthrone-1H** under negative mode (none observed). (g) The peak intensity
 400 and retention time of **DPivOHAC-1H** under negative mode. By integrating the peak areas in (b),
 401 (c), (d), (e), (f) and (g), we found the percentages of **DPivOHAQ** (81.8%), **dianthrone+1H**
 402 (10.2%), **dianthrone-1H** (0.4%), **dianthrone-3H** (1.3%), and **DPivOHAC** (6.3%).
 403



404
405 **Scheme S2.** Proposed possible redox reactions of dianthrones. Because the **dianthrono+1H** (exact
406 mass: 787.3488) and **dianthrono-3H** (exact mass: 783.3175) were detected and plateaus were
407 observed from the voltage profiles, we propose that there are three redox-active states for the
408 dianthrones.

409
410 **Electrosynthesis VI.** Electrochemical synthesis of **DBAQ(COO⁻)** in an undivided electrolytic
411 cell at 0.1 M concentration, against the HER.
412



413
414
415 An undivided electrolytic cell was prepared with carbon felt (XF30A, Toyobo Co., volumetric
416 porosity: 95%) as the working electrode, a carbon rod as the counter electrode, and Ag/AgCl (3 M
417 NaCl) as the reference electrode.

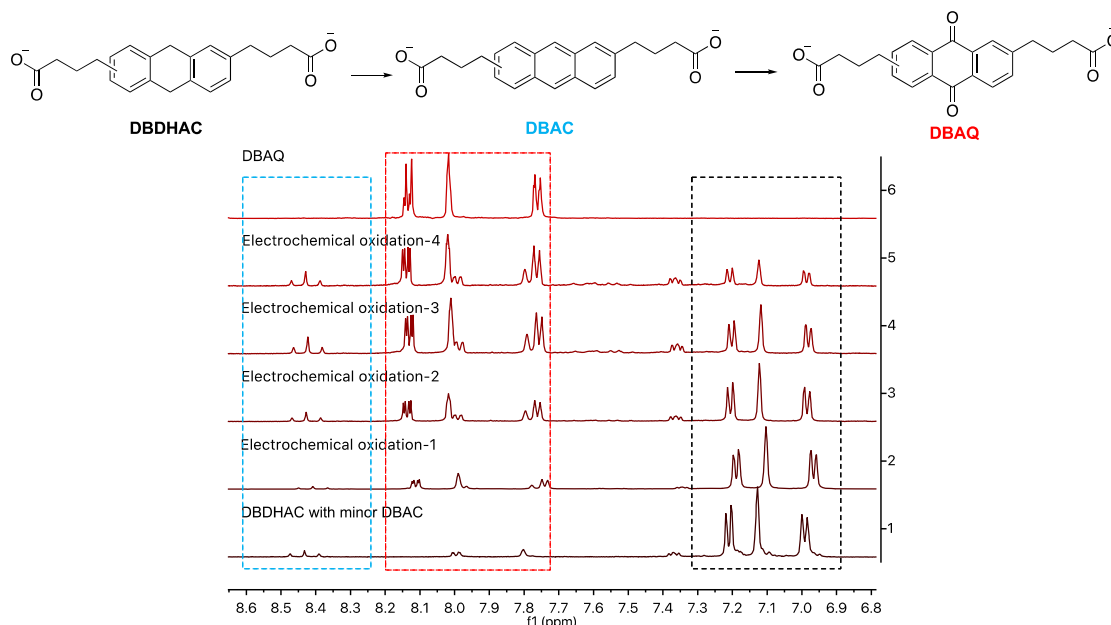
418
419 Electrolyte preparation: 0.35 g **DBDHAC** (synthesized by following our previous work), 0.745 g
420 KCl, and 0.561 g KOH were dissolved in deionized water to obtain a 10 mL solution containing
421 0.1 M **DBDHAC**, 1.0 M KCl, and 1.0 M KOH.

422
423 On the working electrode: **DBDHAC** was oxidized to **DBAQ**; on the counter electrode: water was
424 reduced to hydrogen gas.

425
426 Electrochemical oxidation of **DBDHAC(COO⁻)**: while the electrolyte was stirring, a constant
427 potential (1.1 V vs. Ag/AgCl) was applied to the divided electrolytic cell until 120% of the required
428 coulombs were extracted from the working electrode. [0.1 M * 0.01 L * 96485 C/mol * 8 * 1.2 =
429 926.3 C; 8 electrons need to be extracted from every **DBDHAC** molecule].

430

431 Characterization of analyte: an aliquot (250 μ L) was transferred from the as-prepared analyte to
 432 an Eppendorf® tube (capacity: 1.5 mL) and acidified by a drop of concentrated HCl to obtain
 433 **DBAQ** precipitate. The final **DBAQ** precipitate was re-dissolved in DMSO- d_6 for 1 H NMR
 434 measurement. According to the integration of the 1 H NMR spectrum in the Figure S12, the yield
 435 is 70%. The faradaic efficiency (%) = [yield (%) / 1.2] = 58.3%.
 436



437 **Figure S12.** 1 H NMR spectra of **DBDHAC** (bottom), chemically synthesized **DBAQ** (top), and
 438 electrochemically synthesized **DBAQ** in an undivided cell after varying extents of reaction.
 439 **DBDHAC**: 4,4'-(9,10-dihydroanthracene-diyl)dibutanoic acid; **DBAC**: 4,4'-(anthracene-
 440 diyl)dibutanoic acid; **DBAQ**: 4,4'-(9,10-anthraquinone-diyl)dibutanoic acid. The time interval
 441 between successive measurements labeled electrochemical oxidation-1, 2, 3, and 4 is
 442 approximately one hour. The deuterated solvent is DMSO- d_6 .
 443
 444

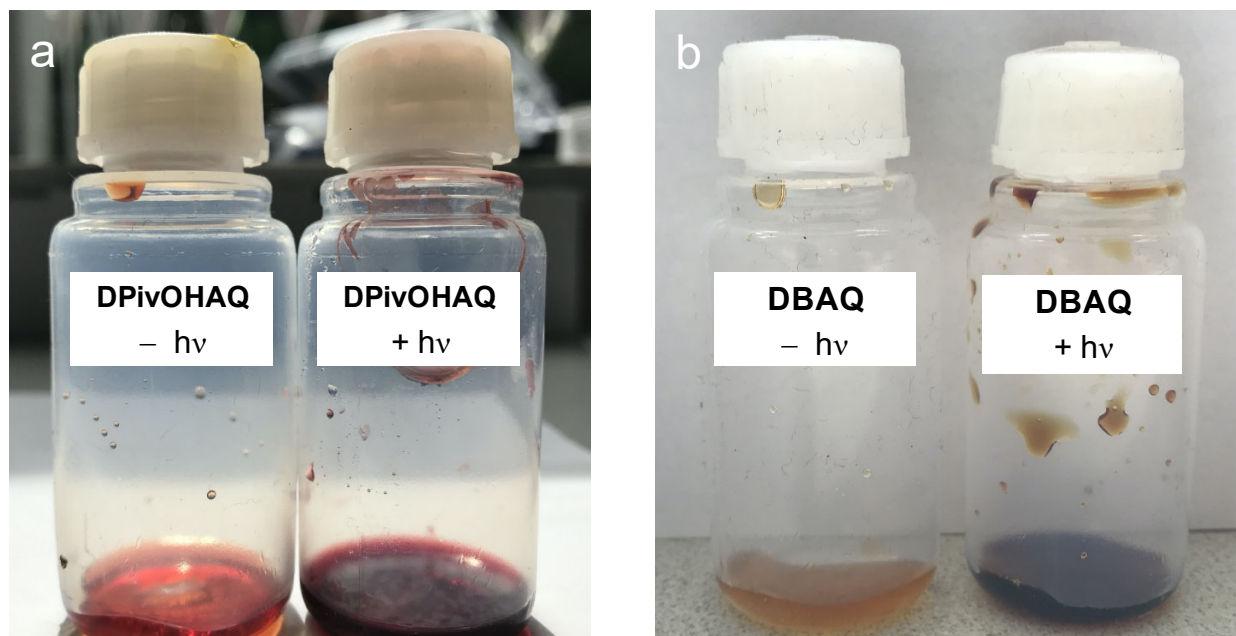
445 Light sensitivity experiments

446 It has been reported that quinones and related compounds can decompose in the presence of
 447 light.^{S2-S6} In order to determine the light sensitivity of **DPivOHAQ** and **DBAQ**, we compared
 448 solutions of each compound held in the presence of and in the absence of light for 1 week. Two
 449 samples of **DPivOHAQ** (0.1 M, pH 12 in water with 1 M KCl, 1.5 mL each) and two samples of
 450 **DBAQ** (0.1 M, pH 12 in water, 1.5 mL each) were prepared in separate FEP bottles (VWR Catalog
 451 No. 16071-008). For each compound, one sample was wrapped in aluminum foil and stored in a
 452 dark drawer for 1 week. The other sample was held for 1 week under a quartz halogen lamp with
 453 a controllable output of 50–1000 W set to 500 W (CowboyStudio QL-1000 W HEAD; ePhotoInc
 454 QL 1000Bulb). The samples exposed to light were allowed to float at the top of a water bath
 455 containing approximately 16 L of water to dissipate excess heat produced by the lamp (the liquid
 456 level decreased gradually due to evaporation and was replenished daily). The liquid level was
 457 maintained at a distance of approximately 20 cm from the light source.
 458

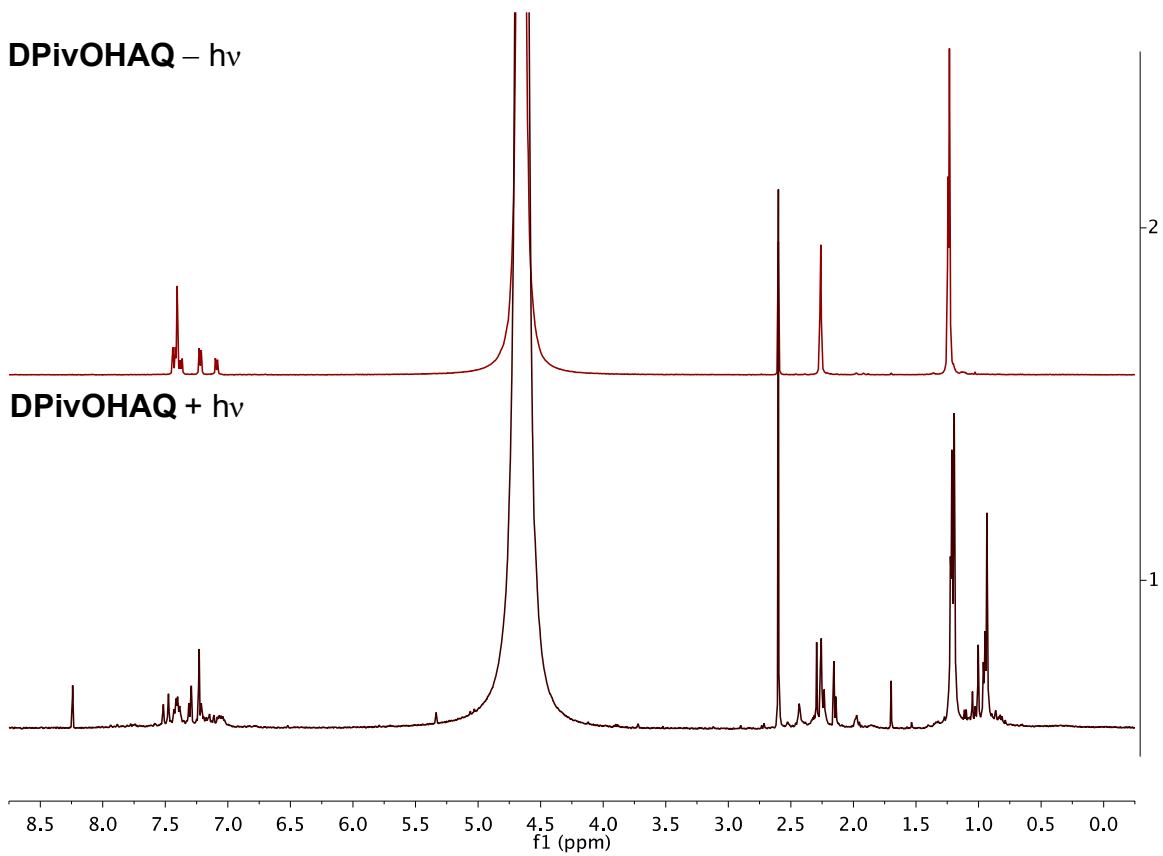
459 After 1 week, differences in color were observed between the samples of each compound stored
 460 in the dark and exposed to light (Figure S13). The formation of a film was also observed in the

461 **DPivOHAQ** sample exposed to light. ^1H NMR spectra of each sample demonstrate decomposition
462 of both compounds stored in the presence of light (Figures S14 and S15).

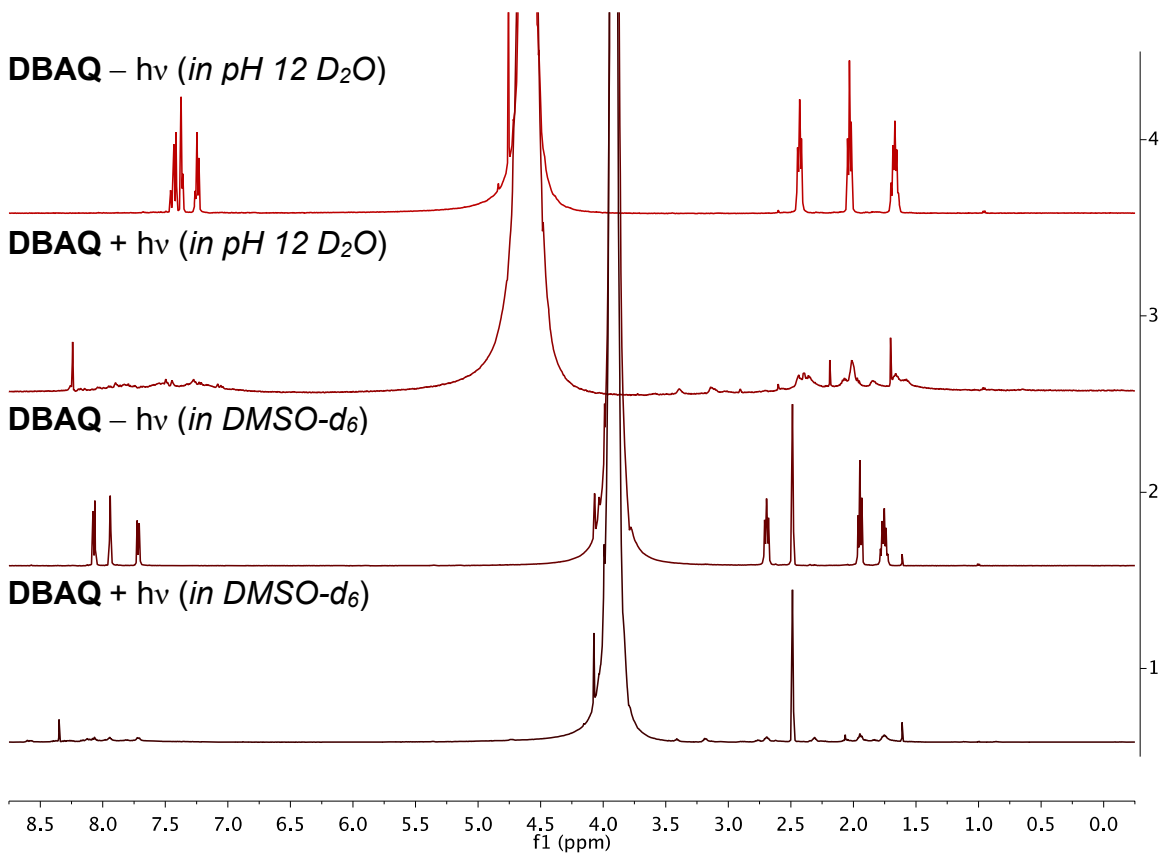
463
464 We therefore wrapped the electrolyte reservoirs with aluminum foil to avoid decomposition due
465 to light exposure during cell cycling.
466



467
468
469 **Figure S13.** Samples of (a) **DPivOHAQ** (0.1 M, pH 12) stored for 1 week in the absence of light
470 (– hv) and under a 500 W lamp (+ hv) and of (b) **DBAQ** (0.1 M, pH 12) stored for 1 week in the
471 absence of light (– hv) and under a 500 W lamp (+ hv). Differences in color were observed between
472 the two samples of each compound. The formation of a film was also observed in the **DPivOHAQ**
473 sample exposed to light.
474



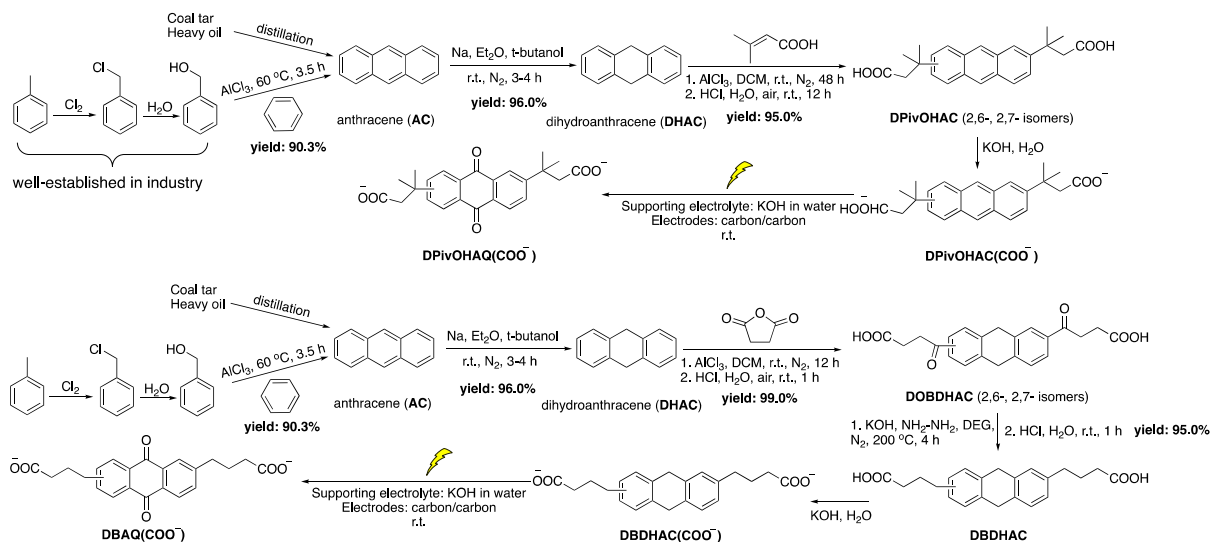
475
 476 **Figure S14.** ^1H NMR spectra of samples of **DPivOHAQ** (0.1 M, pH 12) stored for 1 week in the
 477 absence of light ($-h\nu$) and under a 500 W lamp ($+h\nu$), each diluted (1:5.5) in pH 14 D_2O (1 M
 478 KOD) containing a 9 mM NaCH_3SO_3 internal standard (δ 2.6 ppm).



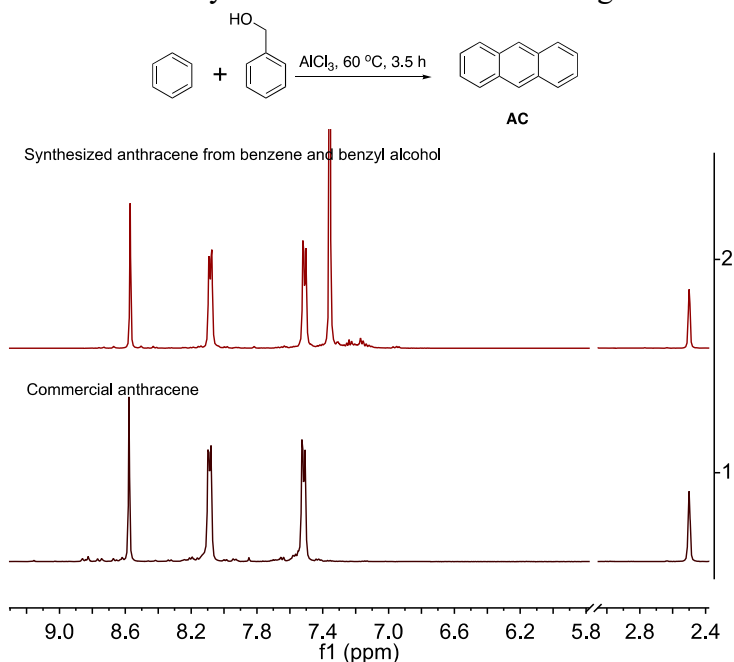
479
 480 **Figure S15.** ^1H NMR spectra of samples of **DBAQ** (0.1 M, pH 12) stored for 1 week in the
 481 absence of light ($-h\nu$) and under a 500 W lamp ($+h\nu$), each diluted (1:5) in pH 12 D_2O or in
 482 $\text{DMSO}-d_6$.

483
 484 **Complete synthesis**
 485

486



487
488 **Scheme S3.** Complete synthetic routes, conditions, and yields of **DPivOHAQ** and **DBAQ** when
489 commercially available commodity chemicals are used as starting materials.



490
491 **Figure S16.** ¹H NMR spectra of commercial and synthesized anthracene (**AC**) in DMSO-*d*₆. The
492 peak at 7.37 ppm in the synthesized **AC** spectrum is from benzene.

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