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Maximization of hydrogen peroxide utilization in a proton exchange membrane H₂O₂ electrolyzer for efficient power-to-hydrogen conversion†

Jie Yang,^{a,b} Ruimin Ding,^{*a} Chang Liu,^{ID a,c} Lifang Chen,^a Qi Wang,^{a,b} Shanshan Liu,^a Qinchao Xu^a and Xi Yin^{ID *a}

A hydrogen peroxide electrolyzer (HPEL) is the workhorse for an energy storage system based on the H₂O₂ electrochemical cycle. The high H₂O₂ utilization towards power-to-hydrogen conversion in the HPEL is essential to ensure the efficiency and cyclability of the system. Unfortunately, the H₂O₂ disproportionation at the anode and its crossover to the cathode in a proton exchange membrane (PEM) HPEL is detrimental to H₂O₂ utilization and must be mitigated. This work investigates the effects of the catalyst type, anode catalyst loading, and PEM thickness on H₂O₂ utilization in a PEM HPEL. The results show that the Co–N–C catalyst exhibits higher H₂O₂ utilization than the Fe–N–C and Pt/C catalysts due to its higher selectivity towards the hydrogen peroxide oxidation reaction (HPOR) and the lesser H₂O₂ disproportionation reaction (HPDR). Increasing the Co–N–C catalyst loading and PEM thickness can effectively inhibit the H₂O₂ crossover and improve the H₂O₂ utilization. On the other hand, the portion of the HPDR and the ohmic loss increase with the catalyst loading and PEM thickness, respectively. A maximum H₂O₂ utilization of over 98% can be achieved by balancing these factors. These results provide valuable guides to the catalyst design and device optimization for efficient energy storage systems based on the electrochemical H₂O₂–H₂ cycle.

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1. Introduction

The percentage of electricity generated by intermittent renewable sources (*e.g.*, wind and solar power) is increasing rapidly. Large-scale, efficient, economical, and zero-emission energy storage systems are needed to integrate renewable-sourced electricity with the traditional power grid and store the electricity in the short or long term.^{1,2} Electrochemical energy storage systems based on the H₂–water cycle,^{3–10} the ammonia–N₂ cycle,^{11–14} the H₂O₂–water cycle,^{15–17} metal-ion batteries,^{18,19} and redox-flow batteries^{20–22} have been proposed for large-scale energy storage and grid balancing.

Recently, we proposed the concept of distributed generation and energy storage systems based on the highly efficient electrochemical cycle of hydrogen peroxide (H₂O₂).^{23–25} In this system, H₂O₂ can be electrolyzed to H₂ and O₂ to store energy in the short term and then be regenerated *via* the two-electron oxygen reduction reaction (2e–ORR) in a fuel cell to generate power. Our proof-of-concept studies have demonstrated a highly efficient and stable PEM H₂O₂ electrolyzer (HPEL) for power-to-hydrogen conversion with a H₂ production Faraday efficiency of 96% and a unitized regenerative hydrogen peroxide cycle cell (UR-HPCC) with a round-trip efficiency (RTE) of above 90% for renewable energy storage.^{23–25} Their unprecedented low system cost and high energy efficiency make the H₂O₂ electrochemical cycle systems highly promising for short- and long-term energy/hydrogen storage, along with other systems (Table S1†). In addition, it is possible to realize long-term energy/hydrogen storage using H₂O₂ as a H₂ carrier and release the hydrogen using a HPEL for various applications.^{23–25}

Achieving a high H₂O₂ utilization towards power-to-hydrogen conversion in the HPEL is essential to ensure the efficiency and cyclability of the H₂O₂–H₂ energy storage system. In an ideal PEM HPEL, the H₂O₂ oxidation reaction (HPOR, H₂O₂ → 2H⁺ + 2e[–] + O₂; E⁰ = 0.695 V *vs.* RHE) takes

^aState Key Laboratory of Coal Conversion, Institute of Coal Chemistry, Chinese Academy of Sciences, Taiyuan, Shanxi 030001, China.

E-mail: dingrm@sxicc.ac.cn, xiyin@sxicc.ac.cn

^bSchool of Chemical Engineering, University of Chinese Academy of Sciences, Beijing 100049, China

^cCenter of Materials Science and Optoelectronics Engineering, University of Chinese Academy of Sciences, Beijing 100049, China

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place and produces protons in the anode and the coupled hydrogen evolution reaction (HER, $2\text{H}^+ + 2\text{e}^- \rightarrow \text{H}_2$; $E^0 = 0 \text{ V vs. RHE}$) occurs in the cathode, converting power to H_2 gas at voltages of 0.7 V and above. In a practical PEM HPEL, as shown in Scheme 1, H_2O_2 in the anode can cross over to the cathode through the PEM and be reduced *via* the hydrogen peroxide reduction reaction (HPRR, $\text{H}_2\text{O}_2 + 2\text{H}^+ + 2\text{e}^- \rightarrow 2\text{H}_2\text{O}$; $E^0 = 1.78 \text{ V vs. RHE}$) at the cathode. This HPRR competes with the HER, decreasing the H_2O_2 utilization towards the power-to- H_2 conversion ($E_{\text{H}_2\text{O}_2\text{-HPOR}}$) and the faradaic efficiency. In addition, the undesired H_2O_2 disproportionation reaction (HPDR, $2\text{H}_2\text{O}_2 \rightarrow 2\text{H}_2\text{O} + \text{O}_2$) may occur at the anode, decreasing the H_2O_2 utilization for H_2 production (Scheme 1).^{24,25} Therefore, it is critical to identify the key factors that affect these processes, mitigate the H_2O_2 crossover, inhibit the competing HPDR, and maximize the H_2O_2 utilization.

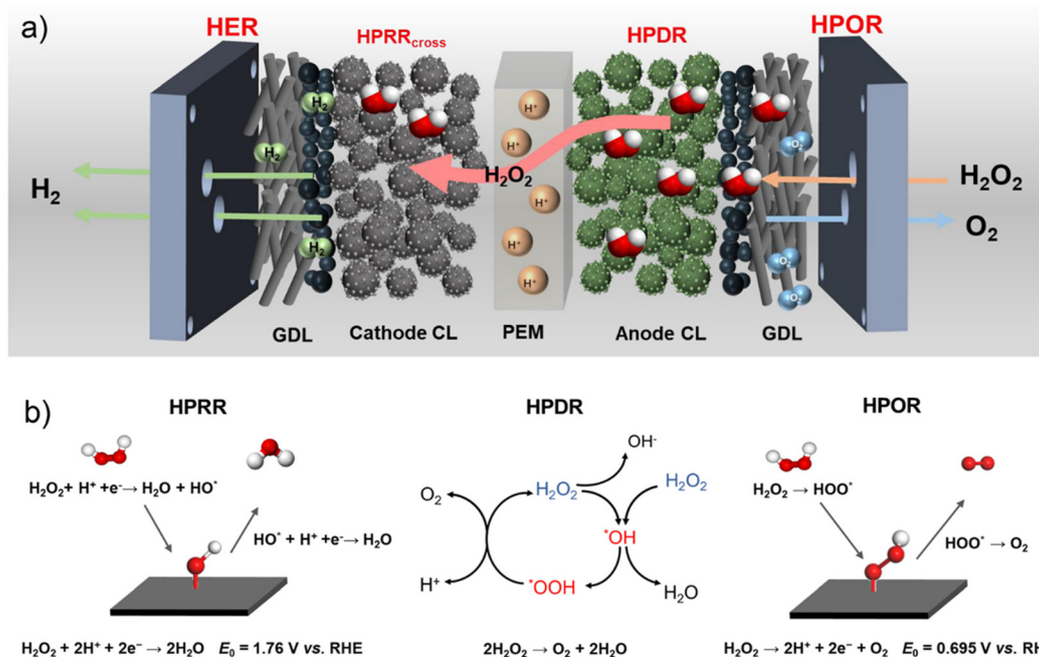
Herein, we focus on the effects of anode catalyst types, anode catalyst loading, and PEM thickness on the H_2O_2 utilization and device performance of the HPEL. By benchmarking the performance of representative platinum group metal-free (PGM-free) catalysts and Pt/C in the electrochemical cell and PEM HPEL, we identify the cobalt- and nitrogen-doped carbon (Co-N-C) catalyst as a better candidate than the Fe-N-C and Pt/C catalysts for its higher selectivity towards the HPOR and lesser HPDR activity. Using Co-N-C, we reveal that increasing the anode catalyst loading and PEM thickness appropriately can effectively inhibit the crossover and thus improve the H_2O_2 utilization. On the other hand, the HPDR intensifies as the anode catalyst loading increases and the internal resistance increases with the PEM thickness. Optimizing the above vital

factors allows us to achieve a maximum H_2O_2 utilization of 98% in the PEM HPEL for power-to-hydrogen conversion. Our results elucidate the interdependences of the competing HPOR, HPDR, H_2O_2 crossover, and H_2O_2 utilization in the PEM HPEL, providing guides for catalyst design and device optimization of efficient energy storage systems based on electrochemical H_2O_2 - H_2 cycles.

2. Experimental

2.1 Membrane electrode assembly (MEA) fabrication

Anode catalyst inks were prepared by dispersing 50 mg of the Co-N-C or Fe-N-C catalyst in a mixture of de-ionized water (DI-water, 30.0 mL), isopropanol (IPA, 30.0 mL), and D521 Nafion dispersion (1070 μL , 5 wt%) in an ultrasonic bath for 2 h or by dispersing 16 mg of Pt/C (Hispec 3000, Johnson Matthey Co.) catalyst in a mixture of DI-water (12.0 mL), IPA (3.0 mL), and D521 Nafion dispersion (165 μL , 5 wt%) in an ultrasonic bath for 30 min. Then, the inks were sprayed onto pieces of 4-cm² hydrophilic carbon cloth (*ca.* 0.33 mm in thickness) at 80 °C to form anode electrodes. Anodes with different catalyst loadings were obtained by controlling the volume of ink sprayed. The catalyst loadings were further confirmed by analyzing the electrochemical surface area (ECSA) of the anode catalysts. Commercial Pt/C gas diffusion electrodes (GDEs) (0.2 mg_{Pt} cm⁻², 20 wt% Pt-C on Sigracet 22BB carbon paper, Fuel Cell Store, USA) were used as cathodes. The Pt/C cathode was hot-pressed to Nafion membranes with different thicknesses (NR211, NR212, NR115, and NR117, Chemours) at



Scheme 1 (a) In a PEM HPEL, H_2 generation at the cathode stems exclusively from the HER. O_2 is generated at the anode *via* the following three processes: HPOR coupled with the HER, HPOR coupled with the HPRR_{cross}, and HPDR at the anode. (b) Schematic representation of the response mechanisms of the HPOR, HPDR,^{26,27} and HPRR.



120 °C and 5.3 MPa for 5 minutes to form the half-MEAs. Then, the anodes and the half-MEAs were assembled in the HPEL hardware to form the full MEAs. A detailed description of the HPEL hardware and the system setup can be found in our previous publications.^{24,25}

2.2 PEM HPEL test and H₂O₂ utilization analysis

The PEM HPEL performance was evaluated at 25 °C and ambient pressure using an established protocol.^{24,25} During the activation process, the Pt/C cathode was filled with pure H₂ gas to form a quasi-reversible hydrogen electrode (RHE). N₂-saturated 0.5 M H₂SO₄ was pumped into the anode and circulated between the anode and a Boro 3.3 glass bottle at a flow rate of 200 mL min⁻¹. The initial cyclic voltammetry (CV) was recorded by polarizing the Co-N-C or Fe-N-C anode from 0 to 1.0 V vs. RHE or the Pt/C anode from 0.05 to 1.2 V vs. RHE at a scan rate of 50 mV s⁻¹ using a CHI 760E bipotentiostat. The double-layer capacitance for the PGM-free catalyst was calculated by integrating the charging current over a potential range of one-volt wide. A capacitance of 30 μF cm⁻² was used to calculate the electrochemical surface area (ECSA), which is used to estimate the loading of the Co-N-C and Fe-N-C catalysts.^{28,29} The loadings of the Pt/C anode catalyst were determined using a Thermo Scientific ARL QUANT'X energy dispersive X-ray fluorescence (EDXRF) spectrometer.

During the PEM HPEL operation, a mixture of 0.5 M H₂SO₄ and 0.5 M H₂O₂ was pumped into the anode and circulated between the anode and a glass flask at 200 mL min⁻¹. Linear sweep AC voltammetry was performed by increasing the voltage from the OCV to 1.0 V at a scan rate of 5 mV s⁻¹, while a 10 kHz AC wave with 5 mV amplitude was applied to measure the real-time high-frequency resistance (HFR). Electrochemical impedance spectroscopy (EIS) was performed to examine the resistance of MEAs at a constant open cell voltage (OCV) with an amplitude of 10 mV. The frequency ranged from 1 Hz to 100 kHz. The *i*R-corrected cell voltage ($E_{iR\text{-free}}$) was obtained using the following equation:

$$E_{iR\text{-free}} = E_{\text{cell}} - E_{\text{ohm}} = E_{\text{cell}} - j \times \text{HFR} \quad (1)$$

where E_{cell} is the measured voltage, E_{ohm} is the ohmic loss, and j is the current density.

The voltage efficiency (VE) was calculated using the following equations:

$$\text{VE} = \frac{E_{\text{rev}}}{E_{\text{cell}}} \times 100\% \quad (2)$$

where E_{rev} is the reversible cell potential of H₂/H⁺/Nafion/O₂/H₂O₂ ($E_{\text{rev}} = 0.704$ V, $T = 298.15$ K).

The H₂ production rate (Y_{H_2}) was calculated according to the following equation:

$$Y_{\text{H}_2} = \frac{(P - P_{\text{H}_2\text{O}})V_{\text{H}_2}}{RTtA} \quad (3)$$

where P is the atmospheric pressure in Taiyuan (92 kPa), $P_{\text{H}_2\text{O}}$ is the saturated vapor pressure of water (kPa) at room temperature, V_{H_2} is the volume of fully humidified H₂ produced, R is the ideal gas constant (8.314 J (mol K)⁻¹), T is the measured

room temperature, t is the collection time, and A is the geometric area of the electrode (cm²).

The efficiencies of H₂O₂ ($E_{\text{H}_2\text{O}_2}$) consumed for the HPOR ($E_{\text{H}_2\text{O}_2\text{-HPOR}}$), crossover ($E_{\text{H}_2\text{O}_2\text{-HPRRCross}}$), and disproportionation ($E_{\text{H}_2\text{O}_2\text{-HPDR}}$) were calculated using the following equation:^{24,25}

$$E_{\text{H}_2\text{O}_2\text{-HPOR}} = \frac{V_{\text{H}_2}}{2V_{\text{O}_2} - V_{\text{H}_2}} \times 100\% \quad (4)$$

$$E_{\text{H}_2\text{O}_2\text{-HPRRCross}} = \frac{2V_{\text{H}_2} \text{FE}_{\text{HPRR}}}{\text{FE}_{\text{H}_2}(2V_{\text{O}_2} - V_{\text{H}_2})} \times 100\% \quad (5)$$

$$E_{\text{H}_2\text{O}_2\text{-HPDR}} = \frac{2 \left(V_{\text{O}_2} - \frac{V_{\text{H}_2}}{\text{FE}_{\text{H}_2}} \right)}{2V_{\text{O}_2} - V_{\text{H}_2}} \times 100\% \quad (6)$$

where V_{O_2} is the volume of fully humidified O₂ produced, FE_{H_2} and FE_{HPRR} are the faradaic efficiencies for the HER and the crossover H₂O₂ reduction reaction ($\text{FE}_{\text{HPRR}} = 100\% - \text{FE}_{\text{H}_2}$), respectively.

3. Results and discussion

3.1 Effects of catalyst selection on the HPEL performance and H₂O₂ utilization

We first benchmarked the performance and H₂O₂ utilization of HPELs constructed using Co-N-C, Fe-N-C, and Pt/C anode catalysts under the same operating conditions (see the ESI for details, Fig. S1 and S2†). These catalysts have been used in the prototypes of PEM HPEL and UR-HPCC devices but lack a systematic comparison.^{24,25} Fig. 1a shows the polarization curves of PEM HPELs with Co-N-C, Fe-N-C, and Pt/C anode catalysts at loadings of 1.0 mg cm⁻², 1.0 mg cm⁻² and 0.06 mg_{Pt} cm⁻², respectively. The Co-N-C catalyst shows a lower open cell voltage (OCV) of ca. 0.71 V than those of Fe-N-C (0.81 V) and Pt/C (0.80 V) catalysts, closer to the theoretical reversible cell voltage ($E_{\text{rev}} = 0.704$ V). The corresponding 8 mV onset overpotential (η_{onset}) yields a 98% maximum voltage efficiency (VE), which is higher than those of Fe-N-C (86.4%) and Pt/C catalysts (87.5%). The result is consistent with that in the RDE system (see the ESI for details, Fig. S3†). When the cell voltage (E_{cell}) increased from the OCV to 1.1 V, the HPOR current density (j) increased monotonically to ca. 380, 397, and 624 mA cm⁻² for the Co-N-C, Fe-N-C, and Pt/C catalysts, respectively. Although the current density of the Pt/C catalyst surpasses that of the Co-N-C catalyst above 0.85 V, the Co-N-C catalyst shows higher HPOR activity than the Fe-N-C and Pt/C catalysts in the low voltage range (0.7–0.8 V). Since the HPOR is a two-electron process and the adsorbed hydroperoxyl (HOO*) is the sole intermediate, the binding free energy of HOO* (ΔG_{HOO^*}) can be used as the thermodynamic activity descriptor for HPOR catalysts.^{24,25} According to the Sabatier principle, the ΔG_{HOO^*} value for an ideal HPOR catalyst should be 4.225 eV at $U = 0$ V. The results of the density functional theory (DFT) calculations show that the ΔG_{HOO^*} values at the Co-N_x active sites in the Co-N-C catalyst, including Co-N₄-pyrrolic (4.16 eV), Co-N₂₊₂-pyridinic (3.66 eV), and Co-N₄-pyridinic (3.50 eV), are closer to the ideal value than those of the



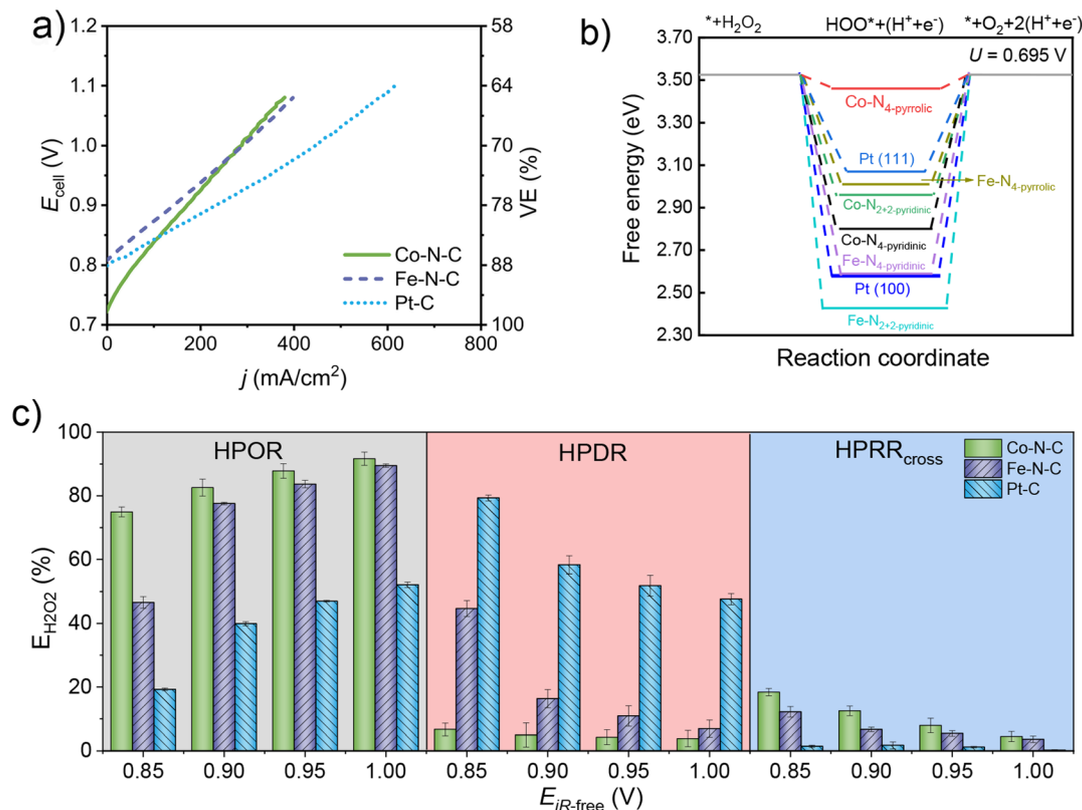


Fig. 1 (a) Polarization curves of PEM HPELs with Co-N-C, Fe-N-C and Pt/C catalysts. (b) Reaction free energy diagram of the HPOR at Co-N₂₊₂, Co-N₄-pyridinic, and Co-N₄-pyrrolic sites in the Co-N-C catalyst, Fe-N₂₊₂, Fe-N₄-pyridinic, and Fe-N₄-pyrrolic sites in the Fe-N-C catalyst, and Pt(111), Pt(100) planes in the Pt/C catalyst. (c) The efficiency of H₂O₂ consumed for the HPOR ($E_{\text{H}_2\text{O}_2\text{-HPOR}}$), disproportionation ($E_{\text{H}_2\text{O}_2\text{-HPDR}}$), and crossover ($E_{\text{H}_2\text{O}_2\text{-HPRRcross}}$) at different $E_{iR\text{-free}}$ values using the Co-N-C, Fe-N-C and Pt/C catalysts.

Fe-N₄-pyridinic active site (3.29 eV), Fe-N₂₊₂-pyridinic (3.13 eV), and Fe-N₄-pyrrolic (3.72 eV) in the Fe-N-C catalyst and the Pt/C catalyst (including Pt (111) (3.77 eV) and Pt (100) (3.28 eV)). And the corresponding free energy diagram of HPOR on these sites at $U = 0.695$ V are shown in Fig. 1b (see the ESI for additional details of DFT calculations and results, Fig. S4–S6 and Table S2†). This more ideal ΔG_{HOO^+} at Co-N_x sites is likely the origin of the higher HPOR activity and lower η_{onset} of the Co-N-C catalyst.^{30–32} Fig. S7a† compares the stability of PEM HPELs with the Fe-N-C, Co-N-C, and Pt-C as anodic catalysts reported elsewhere.^{24,25} In the 20 h stability tests, the voltage at 50 mA cm⁻² changed from 0.852 to 0.840 V, 0.875 to 0.98 V, and 0.86 V to 0.94 V for the Pt-C, Fe-N-C and Co-N-C catalysts, respectively.^{24,25} These results suggest that Pt/C has the highest stability among these catalysts, and the stability of the Co-N-C catalyst is higher than that of the Fe-N-C catalyst. The results from the XPS analysis of Fe-N-C and Co-N-C anodes after the durability test show a decrease of the M-N percentage in the total amount of N (Fig. S7b–e†).²⁴ These results suggest the loss of Fe-N_x or Co-N_x sites. Overall, the Co-N-C catalyst has higher HPOR performance and durability than the Fe-N-C catalyst, and therefore is more suitable for PEM HPEL systems.

Furthermore, we analyzed the effect of catalyst type on H₂O₂ utilization. Fig. 1c shows the efficiencies of H₂O₂ ($E_{\text{H}_2\text{O}_2}$)

consumed for H₂ production ($E_{\text{H}_2\text{O}_2\text{-HPOR}}$), disproportionation ($E_{\text{H}_2\text{O}_2\text{-HPDR}}$), and crossover ($E_{\text{H}_2\text{O}_2\text{-HPRRcross}}$) using the Co-N-C, Fe-N-C, and Pt-C anode catalysts, respectively. Among these catalysts, the Co-N-C catalyst shows the highest $E_{\text{H}_2\text{O}_2\text{-HPOR}}$ and lowest $E_{\text{H}_2\text{O}_2\text{-HPDR}}$ at all iR -corrected cell voltages ($E_{iR\text{-free}}$) from 0.85 to 1.00 V. For all three catalysts, the $E_{\text{H}_2\text{O}_2\text{-HPOR}}$ increases with the $E_{iR\text{-free}}$, while the $E_{\text{H}_2\text{O}_2\text{-HPDR}}$ and the $E_{\text{H}_2\text{O}_2\text{-HPRRcross}}$ decrease. These three variables are interdependent, and their sum is accountable for the total H₂O₂ consumption in the HPEL, hence

$$E_{\text{H}_2\text{O}_2\text{-HPOR}} + E_{\text{H}_2\text{O}_2\text{-HPDR}} + E_{\text{H}_2\text{O}_2\text{-HPRRcross}} = 100\% \quad (7)$$

Their interdependency suggests that by employing an anode catalyst with high activity and selectivity towards the HPOR, the loss of H₂O₂ to the HPDR and crossover can be reduced significantly. Thus, selecting catalysts with high HPOR activity is the key to enhancing H₂O₂ utilization.

More specifically, the Co-N-C catalyst's higher HPOR activity and lower HPDR activity resulted in higher $E_{\text{H}_2\text{O}_2\text{-HPOR}}$ values (consistently above 75%) and lower $E_{\text{H}_2\text{O}_2\text{-HPDR}}$ value (below 3%) throughout the applied voltage range. However, the excess H₂O₂ cannot be completely consumed at the anode, especially at a low cell voltage, so the H₂O₂ crossover can still



occur. The corresponding $E_{\text{H}_2\text{O}_2\text{-HPRR}_{\text{cross}}}$ is consistently above 5%, reaching about 18.3% at an $E_{\text{ir-free}}$ of 0.85 V for the Co-N-C catalyst. For the Pt/C and Fe-N-C anode catalysts, the higher $E_{\text{H}_2\text{O}_2\text{-HPDR}}$ is attributed to their high HPDR activity. The intense HPDR and HPOR within the Pt/C and Fe-N-C anode catalyst layers consume most H_2O_2 and reduce the concentration of H_2O_2 at the interface between the membrane and the catalyst layer, resulting in a low H_2O_2 crossover ($E_{\text{H}_2\text{O}_2\text{-HPRR}_{\text{cross}}}$).

These results show that different catalysts exhibit different behaviors for three competing reactions (HPOR, HPDR, and $\text{HPRR}_{\text{cross}}$), directly affecting the HPEL performance and H_2O_2 utilization. For the Pt/C catalyst, the higher active site density leads to better HPOR activity under high potential operating conditions. However, the high HPDR activity of the Pt/C catalyst significantly reduces the H_2O_2 utilization. Compared to the Pt/C and Fe-N-C catalysts, the Co-N-C catalyst exhibited higher HPOR activity and weaker HPDR activities, which resulted in higher H_2O_2 utilization. High activity and selectivity towards the HPOR and low HPDR activities are essential for anode catalyst selection in PEM HPEL systems.

3.2 Effects of anode catalyst loading on HPEL performance and H_2O_2 utilization

We studied the effects of catalyst loading on the HPEL performance and H_2O_2 utilization using an anode with different loadings of the Co-N-C catalyst. Fig. 2 shows the cross-sectional scanning electron microscopy (SEM) images of the Co-N-C anode catalyst layer (ACL). As the catalyst loading increases from *ca.* 1.0 to 4.5 mg cm^{-2} , the thickness of the ACL increases from *ca.* 6.5 ± 0.5 to $62.5 \pm 10.5 \mu\text{m}$ (Fig. 2f). In the meantime, the coverage of the ACL on the carbon cloth increases with catalyst loading and forms a more continuous ACL, as shown in the top-view SEM micrographs (Fig. S8†).

The texture of the carbon cloth, which is still visible at low catalyst loading, gradually becomes buried by the ACL as the loading increases. However, when the catalyst loading reached *ca.* 3.6 mg cm^{-2} , thin cracks started appearing in the ACL and developed into wider cracks at *ca.* 4.5 mg cm^{-2} (Fig. S8†).

Fig. S9† shows the cyclic voltammograms (CV) of anodes with different Co-N-C catalyst loadings. The double layer capacitance and the ECSA increase as the catalyst loading increases from 1.0 to 4.5 mg cm^{-2} . Fig. 3a and Fig. S10a† show the corresponding PEM HPEL polarization curves, which show the close OCV of *ca.* 0.72–0.73 V. The HPOR current at low voltage (*e.g.* 0.8 V) increased as the loading increased from 1.0 to 3.6 mg cm^{-2} , and then decreased as the loading reached 4.5 mg cm^{-2} (Fig. S10b†). The HPOR current density (j) at high voltage above 0.9 V decreased as the catalyst loading and CL thickness increased (*e.g.* from *ca.* 342 to 263 mA cm^{-2} at 1.00 V). Correspondingly, the internal resistance-corrected cell voltage ($E_{\text{ir-free}}$) increased from 0.80 to 1.00 V and the H_2 production rate (Y_{H_2}) increased at each anode catalyst loading (Fig. 3b). However, Y_{H_2} increased from 4.54 to 5.6 $\text{mmol (cm}^2 \text{ h)}^{-1}$ at 1.0 V as the catalyst loading increased from 1.0 to 3.6 mg cm^{-2} . However, Y_{H_2} decreased to 3.62 $\text{mmol (cm}^2 \text{ h)}^{-1}$ at 1.0 $\text{V}_{\text{ir-free}}$ as the anode catalyst loading increased to 4.5 mg cm^{-2} . Fig. S11† shows the Nyquist plot of electrochemical impedance spectra (EIS) of the HPELs with different anode catalyst loadings measured at the OCV. The high-frequency resistance (HFR, corresponding to the intercept on the x-axis) increased slightly from 0.031 to 0.042 Ω when the anode catalyst loading increased from 1.0 to 4.5 mg cm^{-2} . We believe that the slightly increased HFR values is the result of cracks in the ACL that lead to an apparent increase of electrical contact resistance.^{33,34} The anode polarization resistance ($R_{\text{ct,A}}$) fluctuates slightly from 0.8 ± 0.08 to $0.72 \pm 0.05 \text{ Ohm}$ as the catalyst

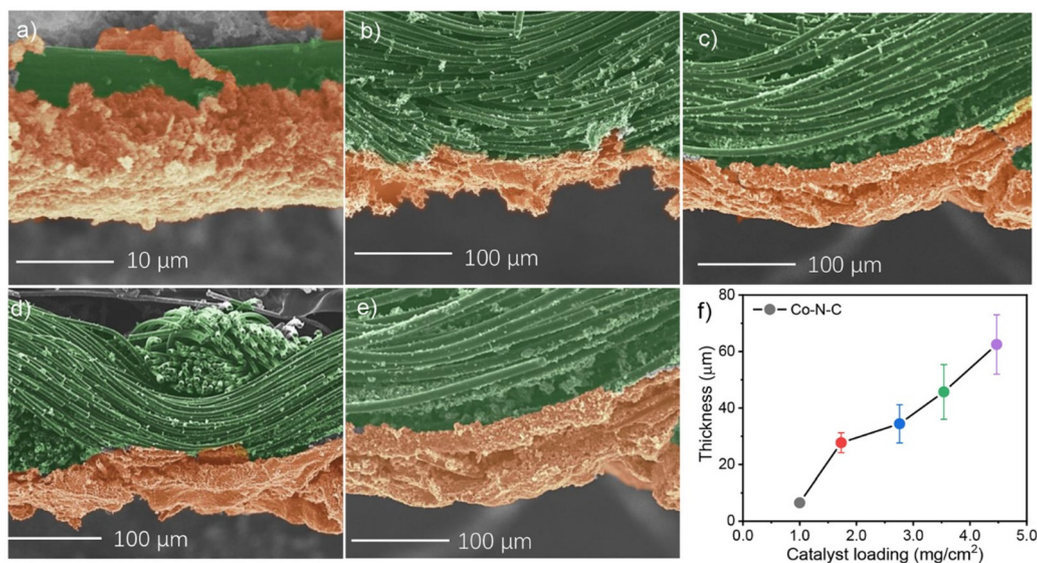


Fig. 2 Cross-sectional SEM micrographs of the Co-N-C ACL with loadings of (a) 1.0, (b) 1.7, (c) 2.8, (d) 3.6 and (e) 4.5 mg cm^{-2} . False colors were applied for clarity. Green: carbon cloth; orange: catalyst layer. (f) Correlation between the catalyst layer thickness and the loading.



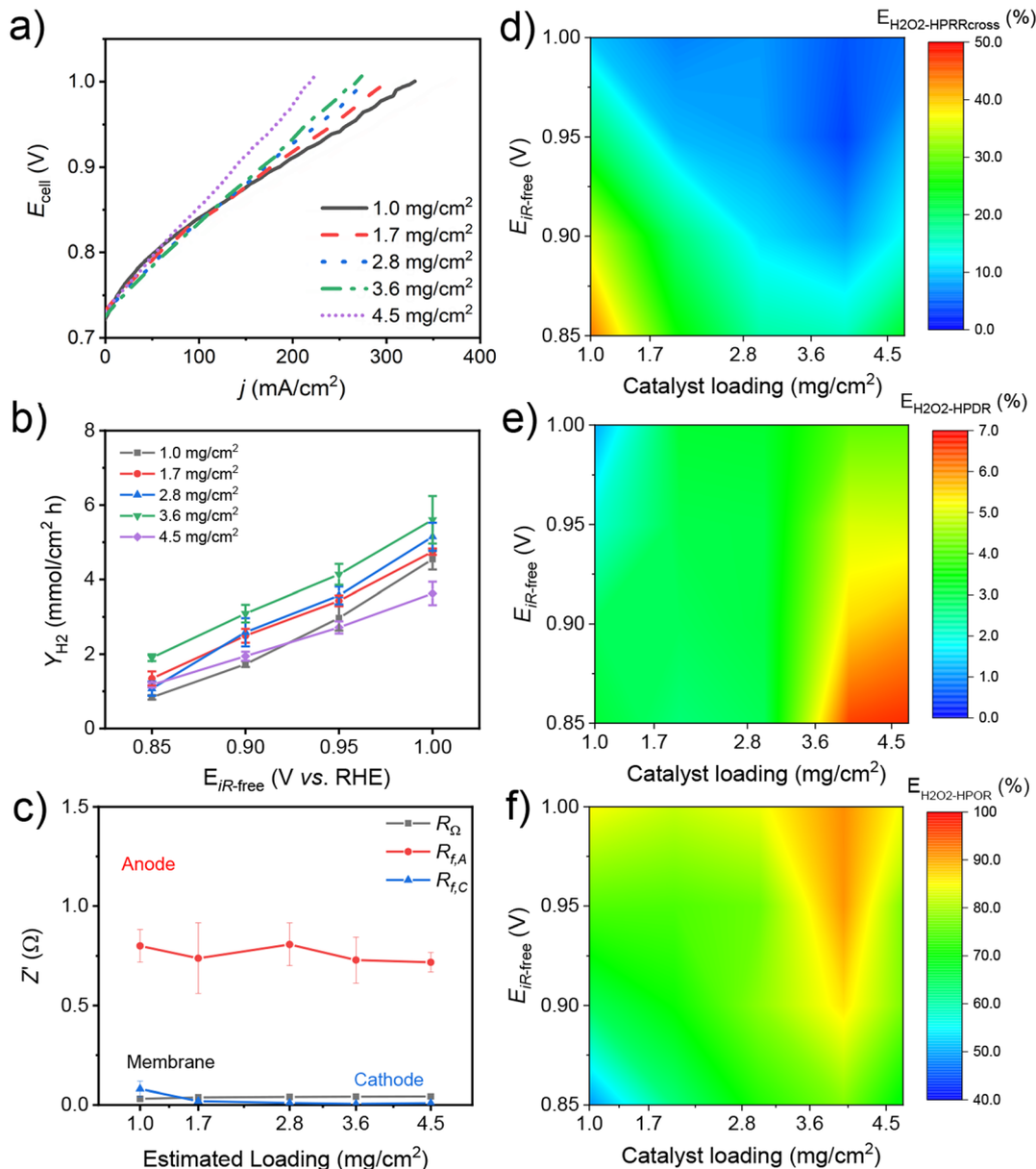


Fig. 3 Performance of PEM HPELs with Co–N–C catalyst loadings of 1.0, 1.7, 2.8, 3.6, and 4.5 mg cm⁻². (a) Polarization curves, (b) Y_{H_2} , (c) variation of R at the OCV with varying catalyst loadings. R_{Ω} refers to the ohmic loss resistance; $R_{ct,A}$ and $R_{ct,C}$ refer to the anode and cathode faradaic resistance, respectively. 2D contour plots of (d) $E_{H_2O_2-HPRRcross}$, (e) $E_{H_2O_2-HPDR}$, and (f) $E_{H_2O_2-HPOR}$ as a function of the catalysts loading and the iR -free cell voltage.

loading increased, while the cathode polarization resistance ($R_{ct,C}$) maintained the same low values (*ca.* 0.01 ± 0.03 Ohm), as shown in Fig. 3c. Clearly, the mass transfer resistance of the anode, as characterized by $R_{ct,A}$, is the main factor affecting the cell performance compared to $R_{ct,C}$ and R_{Ω} . We suggest that at high voltages (> 0.9 V), oxygen generation in large amounts leads to an increase in mass transfer resistance, which is the main reason for the decrease in current density with anode catalyst loading.^{33,35,36}

Fig. 3d–f show the effects of Co–N–C catalyst loading and $E_{iR-free}$ on the H_2O_2 utilization ($E_{H_2O_2-HPOR}$) and the corres-

ponding loss of H_2O_2 ($E_{H_2O_2-HPRRcross}$, $E_{H_2O_2-HPDR}$) in the 2D contour plots (see Fig. S12[†] for the original data). When the catalyst loading increases from 1.0 to 3.6 mg cm⁻², the ACL thickness increased and inhibited the crossover of H_2O_2 . As a result, the $E_{H_2O_2-HPRRcross}$ decreased with the catalyst loading when the loading was below 3.6 mg cm⁻², as shown in Fig. 3d and Fig. S12.[†] However, when the catalyst loading further increased to 4.5 mg cm⁻², wide cracks formed in the ACL and assisted the crossover of H_2O_2 (Fig. S8[†]). On the other hand, as the anode catalyst loading increased, the HPDR intensified due to the decreased HPOR performance, and the $E_{H_2O_2-HPDR}$



increased from 3.1% to 6.7% at an $E_{iR\text{-free}}$ of 0.85 V ($iR\text{-free}$) (Fig. 3e and Fig. S12†). Following the interdependent relationship described in eqn (7), the $E_{\text{H}_2\text{O}_2\text{-HPOR}}$ increases with catalyst loading and peaks at a loading of 3.6 mg cm⁻² (ca. 92% at an $E_{iR\text{-free}}$ of 1.0 V) (Fig. 3f).

We suggest that the increase of anode catalyst loading leads to the increase of ACL thickness, which effectively inhibits the H₂O₂ crossover. However, especially under high voltage operating conditions, the ACL thickness increase significantly increases the mass transport resistance, leading to cell performance degradation. An optimal catalyst loading that balances the interdependent HPOR, HPDR and HPRR_{cross} in the PEM HPEL system is required to maximize the H₂O₂ utilization.

3.3 Effects of PEM thickness on HPEL performance and H₂O₂ utilization

Furthermore, we investigated the effects of PEM thickness on the PEM HPEL performance and H₂O₂ utilization using Nafion membranes with different thicknesses, including Nafion NR 211 (25.4 μm), NR 212 (50.8 μm), NR 115 (127.0 μm), and NR 117 (183.0 μm) (Fig. 4 and Fig. S13†). With the same anode Co-N-C catalyst loading of ca. 1.7 mg cm⁻², these HPELs show consistent anode CV curves (Fig. S14†). While the performance of HPELs decreases with the PEM thickness (Fig. S15†), their $iR\text{-free}$ polarization curves are almost identical (Fig. 4a). When the $E_{iR\text{-free}}$ increased from the OCV to 1.0 V, the j of all HPELs increased monotonically to ca. 360 mA cm⁻². Correspondingly,

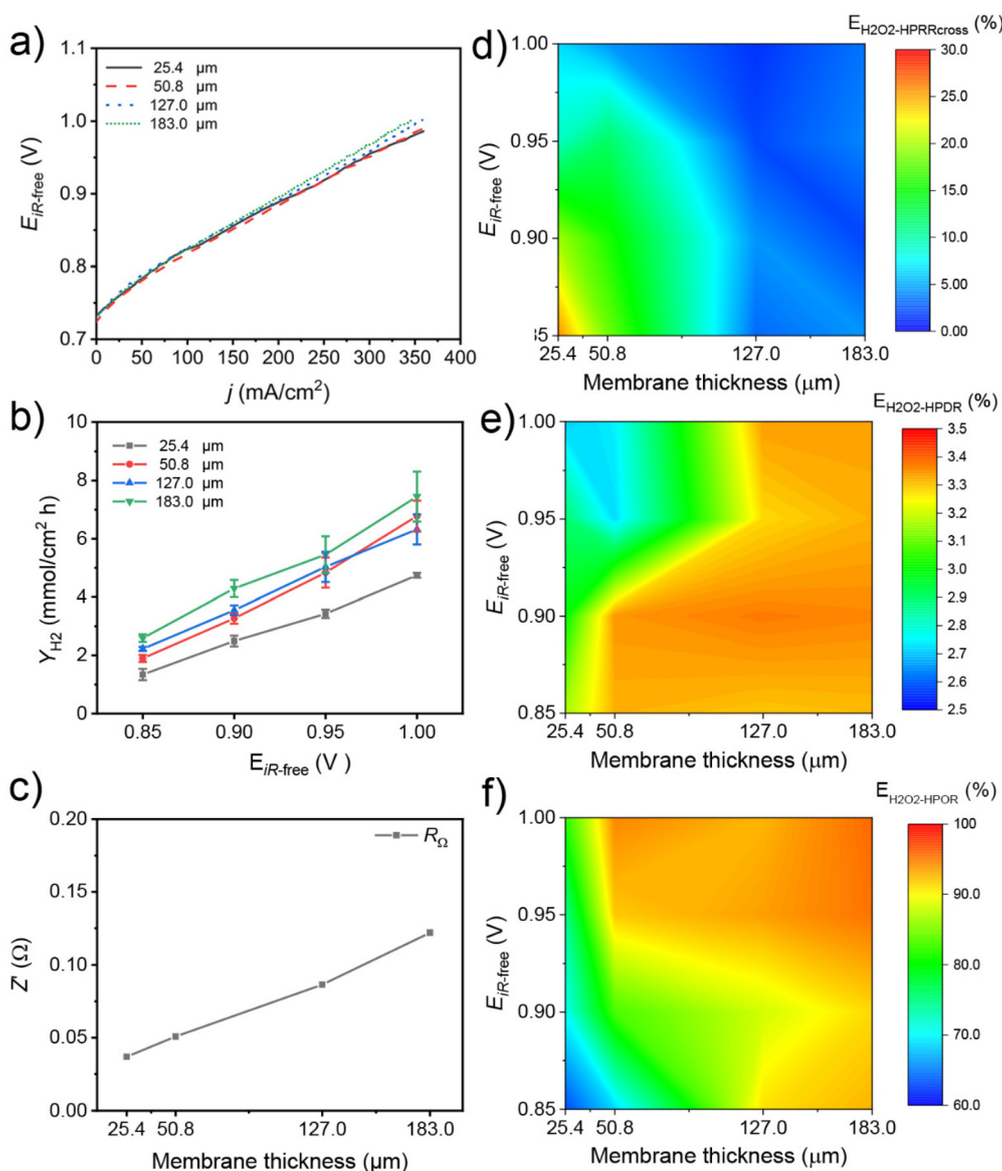


Fig. 4 Performance of PEM HPELs using Nafion membranes with different thicknesses. (a) Polarization curves, (b) Y_{H_2} , and (c) R_{Ω} at the OCV of PEM HPELs with different PEM thicknesses. 2D contour plots of (d) $E_{\text{H}_2\text{O}_2\text{-HPRR}_{\text{cross}}}$, (e) $E_{\text{H}_2\text{O}_2\text{-HPDR}}$, and (f) $E_{\text{H}_2\text{O}_2\text{-HPOR}}$ as a function of the PEM thicknesses and the $iR\text{-free}$ cell voltage.



the H_2 production rate (Y_{H_2}) monotonically increases with the $E_{iR-free}$ and PEM thickness (Fig. 4b). In particular, Y_{H_2} increases from 2.6 to 7.4 mmol (cm² h)⁻¹ as the $E_{iR-free}$ increases from 0.8 to 1.0 V at a PEM thickness of 183 μm. The HFR obtained from the x -intercept of the EIS Nyquist plots increased from 0.038 to 0.12 Ω as the PEM thickness increased from 25.4 to 183.0 μm following the ohmic law (Fig. 4c, and S15[†]). The $R_{ct,A}$ and $R_{ct,C}$ maintained the same level as the PEM thickness increased (Fig. S16[†]). These results suggest that the PEM thickness does not significantly affect the anode polarization behavior within this range but mainly contributes to the ohmic loss of the system.

Fig. 4d–f show the dependence of H_2O_2 utilization on the PEM thicknesses and $E_{iR-free}$. When the PEM thickness increases from 25.4 to 183.0 μm, the $E_{H_2O_2-HPRRcross}$ decreases due to the increased through-plane mass transport resistance for H_2O_2 crossover within the Nafion membrane (Fig. 4d). The $E_{H_2O_2-HPDR}$ decreases with the increase of $E_{iR-free}$ due to the accelerated HPOR rate. On the other hand, the $E_{H_2O_2-HPDR}$ increases with the PEM thickness, which could be attributed to the increased local H_2O_2 concentration within the CL as the H_2O_2 crossover is inhibited (Fig. 4e). The $E_{H_2O_2-HPOR}$ increased with PEM thickness and reached over 90% within the test voltage range (Fig. 4f), following the interdependent relationship described in eqn (6). These results suggest that thicker membranes can effectively inhibit the crossover of H_2O_2 , resulting in lower $E_{H_2O_2-HPRRcross}$ and higher $E_{H_2O_2-HPOR}$, but at the cost of ohmic loss. The tradeoff of $E_{H_2O_2-HPOR}$ and iR loss *via* the optimization of the PEM thickness is needed to maximize the H_2O_2 utilization. Membrane materials with low permeability to H_2O_2 might also improve the H_2O_2 utilization in PEM HPELs.

4. Conclusions and outlook

In summary, we investigated the effects of catalyst type, catalyst loading, and PEM thickness on the performance and H_2O_2 utilization of a PEM HPEL. The results unveiled the interdependency of three fundamental processes, the HPOR, HPDR, and H_2O_2 crossover, which mutually determine the H_2O_2 utilization towards the power-to-hydrogen conversion in the HPEL. Different anode catalysts provide various activities and selectivity towards the HPOR and HPDR. The Co–N–C catalyst is identified as the best anode catalyst for its high activity and selectivity towards the HPOR and low degree of the HPDR. The excellent HPOR activity of the Co–N–C catalyst is attributed to the optimal ΔG_{HOO^*} values at the Co–N₄ sites. In contrast, the Fe–N–C and Pt/C catalysts show higher HPDR activity, lowering H_2O_2 utilization. Increasing the anode catalyst loading and PEM thickness can effectively inhibit the H_2O_2 crossover and improve the H_2O_2 utilization but inevitably lead to higher H_2O_2 loss due to the HPDR and ohmic loss. These findings imply the need for developing highly selective HPOR catalysts and low- H_2O_2 -permeability membranes to boost the performance and H_2O_2 utilization of next-generation PEM HPELs for

renewable energy storage. In addition, the rational design and fabrication of 2e-ORR electrocatalysts with high activity and stability are still necessary to improve the efficiency of H_2 – H_2O_2 cyclic energy storage.

Author contributions

Jie Yang: writing – original draft, methodology, investigation, and visualization. Ruimin Ding: methodology, investigation, visualization, and supervision. Chang Liu: methodology, investigation, and visualization. Lifang Chen: investigation and visualization. Qi Wang: investigation and visualization. Shanshan Liu: investigation. Qinchao Xu: investigation. Xi Yin: conceptualization, methodology, writing – review and editing, supervision, and funding acquisition.

Conflicts of interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Xi Yin, Ruimin Ding, Chang Liu, Shanshan Liu, and Jie Yang have patent CN 202110721698.6 pending to the Institute of Coal Chemistry, Chinese Academy of Sciences. Xi Yin, Jie Yang, Chang Liu, Ruimin Ding, and Shanshan Liu have patent CN202111123206.X pending to the Institute of Coal Chemistry, Chinese Academy of Sciences.

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