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## Dissolution of the Ti porous transport layer in proton exchange membrane water electrolyzers<sup>†</sup>

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The titanium porous transport layer (PTL) is a key component in proton exchange membrane water electrolyzers (PEMWs), facilitating efficient water supply to the catalyst layer while rapidly removing oxygen bubbles. However, in the highly anodic operating environment of PEMWEs, the Ti PTL suffers from degradation, limiting the lifetime of the device. To gain deeper insights into Ti PTL degradation, here we monitor the potential/time-resolved Ti dissolution rates by coupling a PEMWE with an online inductively coupled plasma-mass spectrometer (ICP-MS). The results show that the dissolution of the Ti PTL is a complex and dynamic (electro)chemical event. Initiated by the decreased interfacial pH (even at pH < 1) due to proton accumulation during PEMWE operation, Ti dissolution intensifies with increasing bias potential. However, the dissolved Ti ions are simultaneously hydrolyzed, forming surface Ti oxides that slow down the dissolution rate. Coating the Ti PTL surface with Pt and IrO<sub>2</sub> effectively reduces Ti dissolution, albeit at a higher cost, but they are also susceptible to dissolution during operation. Interestingly, the dissolution profiles of Pt and IrO<sub>2</sub> deposited on the Ti PTL differ significantly from their conventional behavior, which requires further investigation for reliable prediction and optimization of new PTL designs for practical implementation in PEMWEs.

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## Introduction

In the quest for a sustainable, carbon-neutral future, there is a growing energy paradigm shift away from polluting fossil fuels toward a cleaner and more sustainable society.<sup>1</sup> At the heart of this new energy economy is green hydrogen.<sup>2</sup> A critical technology driving this transformation is the proton exchange membrane water electrolyzer (PEMWE),<sup>1</sup> which produces hydrogen gas from abundant water with minimal, or ideally

zero, greenhouse gas emissions. The PEMWE has demonstrated promising performance and rapid response to fluctuations in power supply, making it well-suited for integration with renewable energy sources. However, its harsh operating conditions (>2 V, temperature of 60–80 °C, and acidic environment) limit the choice of materials library that can be used in the PEMWE, as they must have high corrosion resistance.<sup>3,4</sup>

Titanium offers relatively high corrosion resistance under such conditions, with high electrical conductivity and mechanical strength.<sup>5</sup> Therefore, in addition to the bipolar plates, the porous transport layer (PTL), an essential component of the PEMWE for efficiently supplying the water flow to the catalyst layer while facilitating the removal of oxygen bubbles, is typically made of metallic Ti.<sup>6–10</sup> Its remarkable corrosion resistance is due to the formation of a stable protective oxide layer (generally TiO<sub>2</sub>) on its surface when exposed to air or water. This layer is extremely adherent and has the ability to repair itself if damaged, providing robust protection against various corrosive environments. However, the poor electrical conductivity of the protective layer leads to an increase in the ohmic resistance between the PTL and other components of the PEMWE, which can significantly deteriorate the energy efficiency of the PEMWE.<sup>11,12</sup> This drawback has prompted several strategies to ensure durable operation, including optimizing the PTL morphology and coating the metallic Ti surface with thin layers of noble metals (e.g., Pt, Ir, etc.).<sup>13</sup>

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It should be noted, however, that titanium and even noble metals used to protect Ti are not completely immune to corrosion. Using a scanning flow cell (SFC) coupled to an inductively coupled plasma-mass spectrometer (ICP-MS), several research groups have investigated the potential-resolved electrochemical stability of Ti-containing mixed oxide catalysts (*e.g.*, Ir–TiO<sub>x</sub>,<sup>14–16</sup> Ir/CuTiON<sub>x</sub>,<sup>17</sup> and RuIr–TiO<sub>x</sub><sup>18</sup>) in 0.1 M HClO<sub>4</sub> electrolyte. While these studies focused primarily on the dissolution of noble metal catalysts and the subsequent decline in their oxygen evolution reaction (OER) activity — and thus the results may not fully represent the behavior of the Ti PTL in the PEMWE — they showed appreciable Ti dissolution, following a trend similar to that of noble metal dissolution. Meanwhile, Rakousky *et al.* observed the presence of Ti<sup>n+</sup> species at the cathode in an aged catalyst-coated membrane (CCM) after >1000 h of conventional PEMWE operation (2 A cm<sup>-2</sup> and 80 °C).<sup>12</sup> Since the cathode in a pristine CCM was free of Ti and this element was only present in the anode catalyst (a mixture of IrO<sub>2</sub> and TiO<sub>2</sub>) and Ti PTL, Ti dissolution under PEMWE operation was clearly identified. In addition, Ti crossover from the anode PTL to cathode was also observed in the PEMWE for H<sub>2</sub>O<sub>2</sub> synthesis, where Ti felt was used as the PTL but the anode catalyst was only IrO<sub>2</sub>.<sup>19</sup> Thus, the previous findings highlight the need for a deeper understanding of Ti PTL dissolution under PEMWE conditions in order to develop rational PTL designs and operation strategies to meet industrial requirements.

In this study, Ti PTL dissolution in the PEMWE was studied using an online ICP-MS, which allowed potential/time-resolved and quantitative monitoring of the Ti dissolution rate during electrochemical operation. We clearly identified non-negligible Ti dissolution, the rate of which is strongly influenced by the feedstock (or interfacial) pH and applied potential. On the other hand, dissolved Ti ions undergo hydrolysis and form a Ti oxide passivation layer, which suppresses the Ti dissolution rate. We investigated this complex event, during which dissolution and redeposition processes dynamically compete, under various potential profiles on commercialized Ti PTLs with and without Pt and IrO<sub>2</sub> coating layers. Our findings highlight the necessity of similar or further modified diagnostics for better design and evaluation of PTLs to ensure their high durability during PEMWE operation.

## Results and discussion

For the online monitoring of Ti dissolution from the PTL, we coupled a PEMWE cell with an ICP-MS (Fig. 1a). At the anode, the Ti PTL without an OER catalyst, typically Ir(O<sub>x</sub>), was used as the electrode. On the other hand, at the cathode, a Pd/C catalyst rather than conventional Pt/C, which was coated on the gas-diffusion layer (GDL), was employed. It is of note that these distinct configurations of electrodes were due to the elimination of any interferences of Ir or Pt dissolution from the conventional catalytic substances for accurate investigation on Pt/IrO<sub>2</sub>-coated Ti PTLs that will be discussed later, and due to minimization of bubble formation that can disrupt stable operation of the online monitoring. The anode and cathode

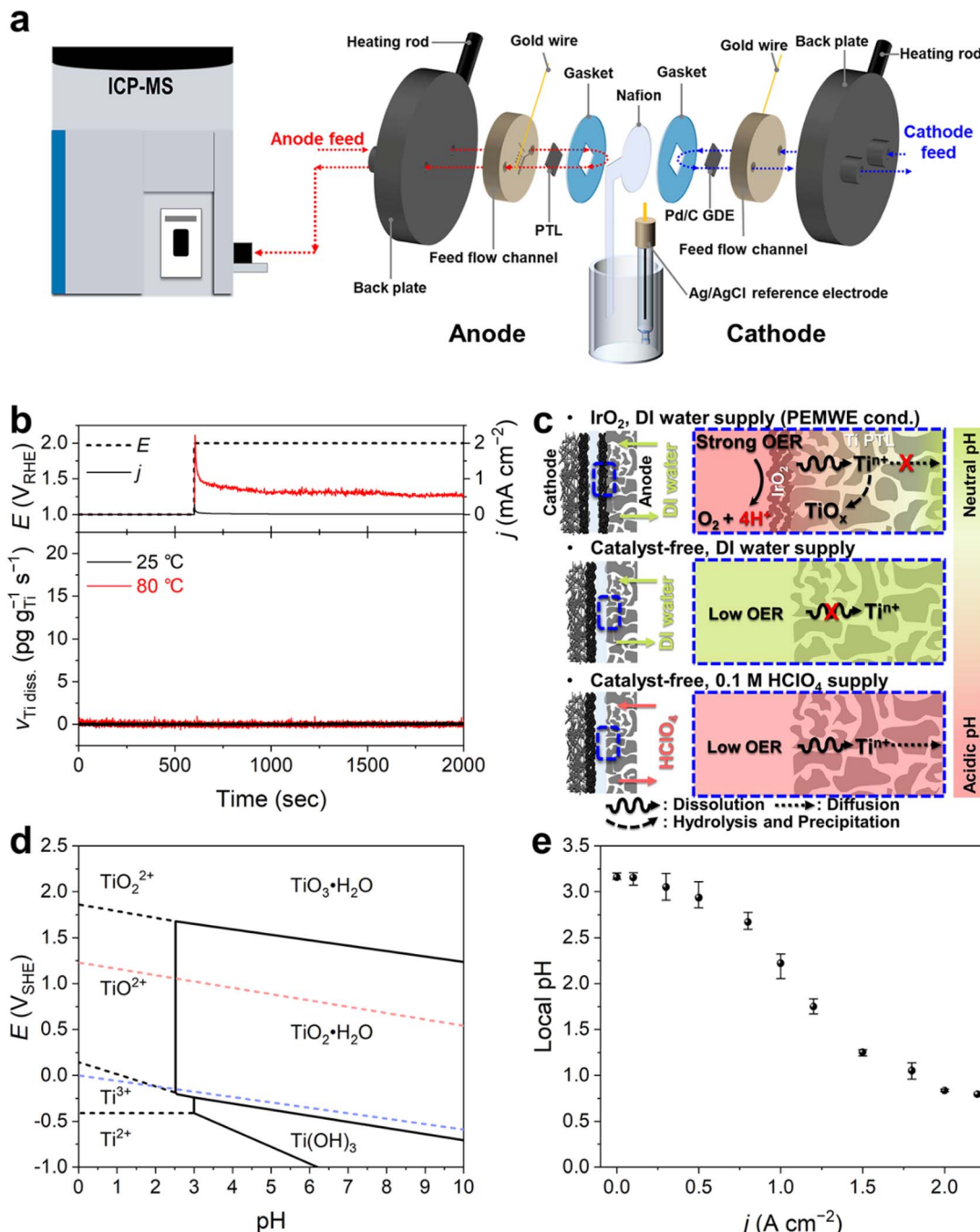
were separated by a Nafion membrane, which was extended out of the electrolyzer cell and connected with a saturated Ag/AgCl reference electrode. This unique cell configuration,<sup>20</sup> beyond the conventional two-electrode system, provided excellent experimental reproducibility and allowed us to accurately control and monitor the anode (or cathode) potential in the electrochemical device (Fig. S1†). Heating rods were inserted into the back plates of the cell to heat the cell. During the measurement, feedstock was supplied at a constant flow rate of 5 mL min<sup>-1</sup> on each electrode side, of which 400 µL min<sup>-1</sup> was introduced into the ICP-MS (see details in the Experimental methods).

Ti dissolution was first investigated at 25 and 80 °C with a continuous flow of deionized (DI) water, a typical feedstock for commercialized PEMWEs, during the anode polarizations at 1 and 2 V *vs.* the reversible hydrogen electrode (RHE; Fig. 1b), which are in the relevant potential range for rest and OER operating potentials, respectively. However, at both temperatures, we found no discernible Ti dissolution during the operation. This result does not agree with previous findings that showed non-negligible Ti dissolution after PEMWE operation,<sup>12,21</sup> thereby implying that our experimental conditions insufficiently reflect those in practical PEMWEs.

We envisioned that this discrepancy primarily originated from different H<sub>2</sub>O consumption and H<sup>+</sup> production rates, leading to consequent differences in local pH at the PTL-electrolyte interface (Fig. 1c). In practical PEMWEs, ampere-order current density (*j*) is typically achieved for efficient evolution of O<sub>2</sub> gas. We also confirmed that our system also reached *j* > 2 A cm<sup>-2</sup> when using the IrO<sub>2</sub> catalyst layer (Fig. S2†). This leads to considerable H<sup>+</sup> production at the anode and its possible accumulation at the interface, lowering the interfacial pH.<sup>22,23</sup> However, in our system, the Ti PTL free from the anode catalyst resulted in a marginal *j* less than 1 mA cm<sup>-2</sup> even at 2 V<sub>RHE</sub> (Fig. 1b), and therefore no considerable changes in the interfacial pH can be reasonably deduced. The importance of feedstock pH on the Ti dissolution can also be understood with the Pourbaix diagram of Ti (Fig. 1d), which shows that pH lower than approximately 2.5 is thermodynamically needed for the exergonic dissolution process of Ti to TiO<sub>2</sub><sup>+</sup> or TiO<sub>2</sub><sup>2+</sup> at potential above 1 V<sub>RHE</sub>. It is of note that, at a given pH 7 of DI water, the most stabilized phase of Ti is solid TiO<sub>2</sub>·H<sub>2</sub>O or TiO<sub>3</sub>·H<sub>2</sub>O depending on the potential applied.<sup>24</sup>

To confirm this hypothesis, *i.e.*, greater accumulation of H<sup>+</sup> at the interface with increasing *j*, we estimated the current-dependent local pH changes using the method recently developed by Sauvé *et al.*,<sup>25</sup> which follows a quasi-equilibrium potential between hydrogen oxidation and reduction reactions on Pt and estimates the interfacial pH (Fig. S3†). The result showed that even at zero current, the interface exhibited a slightly acidic nature (pH 3.2, Fig. 1e), due to the sulfonic acid group-terminated Nafion membrane.<sup>26,27</sup> However, as the current increased, the pH decreased significantly, and it even reached below pH 1 when the *j* exceeded 1.8 A cm<sup>-2</sup>. Therefore, for subsequent online ICP-MS experiments, we opted to use a 0.1 M HClO<sub>4</sub> solution instead of DI water as the feedstock. This choice was made to more accurately simulate the



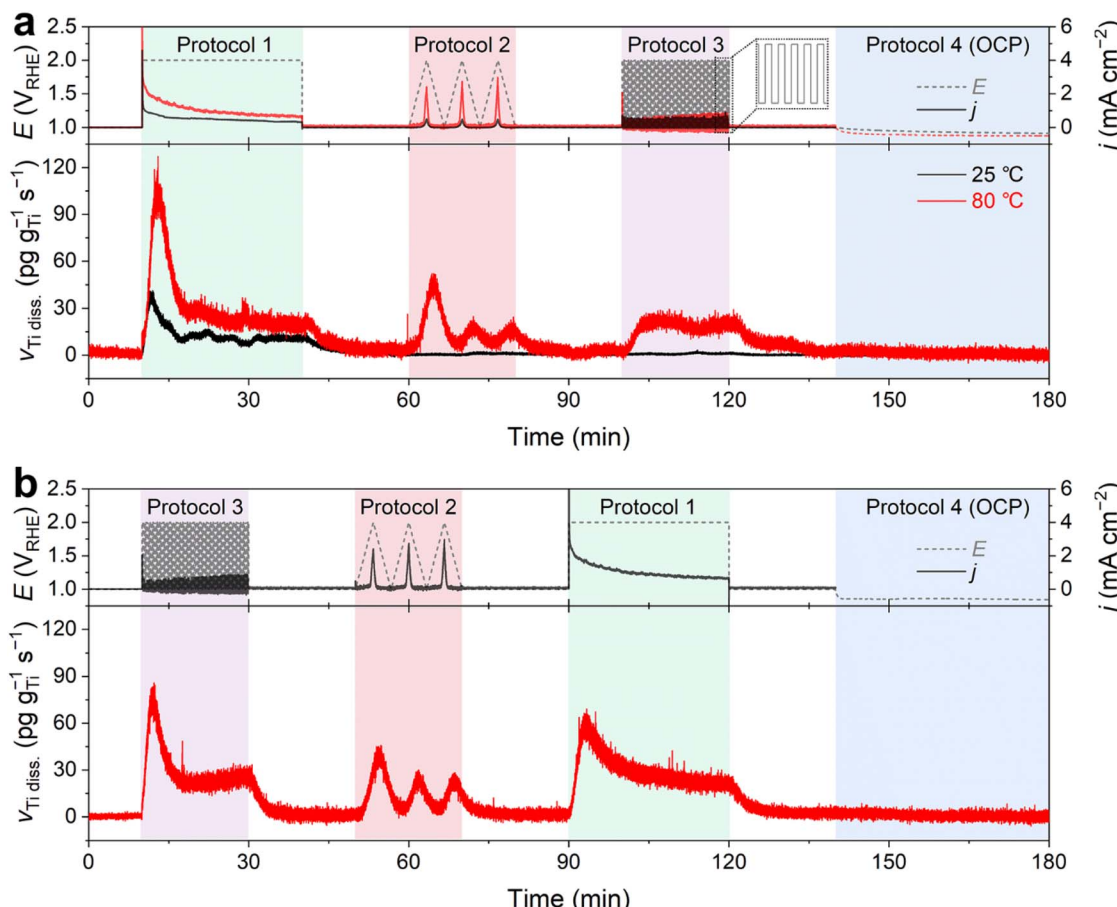


**Fig. 1** (a) Schematic representation of the online PEMWE/ICP-MS setup. (b) Real-time Ti dissolution measured at a static potential of  $2\text{ V}_{\text{RHE}}$  with a supply of  $O_2$ -saturated DI water at  $25\text{ }^\circ\text{C}$  and  $80\text{ }^\circ\text{C}$ . (c) Schematic representation of Ti dissolution behavior under three different operating conditions: PEMWE operation (*i.e.*, Ti PTL with a catalyst layer and DI water supply) and our model systems of catalyst-free Ti PTLs with DI water or  $0.1\text{ M HClO}_4$  supply. (d) Pourbaix diagram of Ti. (e) Estimated local pHs at the electrode–membrane interface measured at different current densities.

interfacial conditions of the real PEMWE without an anode catalyst and to avoid any chemical decomposition and precipitation of dissolved Ti ions into solid  $TiO_x$  as they moved away from the diffusion layer before being analyzed by ICP-MS (Fig. 1c).

In this measurement, we monitored the dissolution rate of Ti in  $O_2$ -saturated  $0.1\text{ M HClO}_4$  during four consecutive potential

protocols (Fig. 2). Protocol 1 entailed the potentiostatic operation of the PEMWE, with a potential hold of the Ti PTL at  $2\text{ V}_{\text{RHE}}$  for 30 min. Conversely, Protocols 2 and 3 involved potentiodynamic conditions, comprising three cyclic voltammograms (CVs) at a scan rate of  $5\text{ mV s}^{-1}$  and 60 cycles of potential pulses of 10 s each within a potential range of  $1\text{--}2\text{ V}_{\text{RHE}}$ , respectively. Typically, the measurement was performed in the order of



**Fig. 2** Real-time dissolution profiles of the Ti PTL measured during potential hold (Protocol 1), 3 cycles of CV (Protocol 2), and potential pulses (Protocol 3). For comparison, Ti dissolution was monitored in the order of (a) Protocols 1 → 2 → 3 at 25 and 80 °C and (b) Protocols 3 → 2 → 1 at 80 °C with a supply of O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> feedstock. At the end of the measurements, the Ti PTL was not externally biased but rested at OCP (Protocol 4).

Protocols 1 → 2 → 3 (Fig. 2a), although the reverse order, Protocols 3 → 2 → 1 (Fig. 2b), was also evaluated. Protocol 4 was then implemented, which corresponded to potential rest, *i.e.*, open circuit potential (OCP), for 40 min. Before and after each protocol, the Ti PTL was polarized at 1 V<sub>RHE</sub> to stabilize the ICP-MS signals.

In contrast to the results observed with DI water (Fig. 1b and S4†), we observed significant Ti dissolution in the HClO<sub>4</sub> feedstock. At 25 °C, significant Ti dissolution rate was found in Protocol 1, but it was almost negligible in the following Protocols 2–4. On the other hand, at 80 °C, a practically more relevant temperature, much greater dissolution of Ti was observed in Protocols 1–3 (Fig. 2a). In addition, it is noteworthy that the Ti dissolution signals were observed in both the anolyte and catholyte (Fig. S5†). This result indicates the crossover of Ti ions from the anode to the cathode, which is consistent with previous results measured after long-term operation of the PEMWE.<sup>12</sup> However, the Ti signal measured in the anolyte was approximately 6 times higher than that in the catholyte, suggesting that the diffusion of dissolved Ti ions from the anode to the cathode was relatively marginal under our experimental

conditions. Therefore, the following investigations focused solely on the dissolution of the Ti PTL in the anolyte at 80 °C.

In the standard protocol sequence (Fig. 2a), no perceptible Ti dissolution was detected at 1 V<sub>RHE</sub>. However, at the initial potential jump to 2 V<sub>RHE</sub> in Protocol 1, the Ti PTL exhibited substantial Ti dissolution of approximately 100 pg g<sub>Ti</sub><sup>-1</sup> s<sup>-1</sup>. However, the intense Ti dissolution rapidly decreased within 500 seconds and stabilized at approximately 25 pg g<sub>Ti</sub><sup>-1</sup> s<sup>-1</sup> during the potential hold. Afterward, in Protocol 3, a consistent Ti dissolution rate was observed, quantitatively similar to the plateau observed in Protocol 1. However, the initial intense Ti dissolution, seen in Protocol 1, was not found in Protocol 3, while the Ti PTL underwent the same potential shift from 1 to 2 V<sub>RHE</sub> and experienced this drastic potential change repeatedly. On the other hand, in the reverse protocol sequence (Fig. 2b), the initially intense but subsequently attenuated Ti dissolution was also recorded in Protocol 3, when it preceded all other protocols. The same conclusion was reached when the potential excursion started with Protocol 2 (Fig. S6†). Therefore, these results implied that the most intense Ti dissolution occurs at the initial potential transition from 1 to 2 V<sub>RHE</sub>, but thereafter

its dissolution rate becomes relatively mitigated regardless of either potentiostatic or pulsed operation at 2 V<sub>RHE</sub>.

It is generally accepted that the Ti surface can be protected against corrosion by forming stable passivation films, *e.g.*, Ti oxide or hydroxide, even in acidic environments.<sup>28,29</sup> However, the intense Ti dissolution at the first potential transition from 1 to 2 V<sub>RHE</sub> allowed us to consider either that the Ti PTL surface is not fully protected by the passivation film, but contains some pits or defective sites, or that the native passivation layer is only quasi-stable, resulting in considerable Ti dissolution from an exposed metallic Ti and/or quasi-stable Ti passivation layer under the highly anodic conditions of 2 V<sub>RHE</sub> (Fig. 3a). Nevertheless, the subsequent exponential decay of the Ti dissolution rate implies the hydrolysis of dissolved Ti ions at the interface and the formation of a relatively stable or thicker passivation layer on the Ti PTL, as also explained in the literature.<sup>28–30</sup> Scanning electron microscopy (SEM) and energy-dispersive

spectroscopy (EDS) of the Ti PTL revealed the increase in oxygen content after its polarization at 2 V<sub>RHE</sub>, corroborating the formation of an additional passivation layer (Fig. 3b and S7†). However, the marginal but consistent Ti dissolution during potential hold at 2 V<sub>RHE</sub> and potential pulses between 1 and 2 V<sub>RHE</sub> suggests that even the newly formed passivation layer may not be completely immune to corrosion, possibly because the thermodynamically most stable phase under such conditions is Ti ions (TiO<sub>2</sub><sup>+</sup> and TiO<sub>2</sub><sup>2+</sup>) rather than solid Ti oxides. Computational modeling based upon the above simple mechanistic scenario toward Ti dissolution also successfully describes the trend of experimental results (Fig. 3c and S8†), *i.e.*, initially intensive but subsequently marginal Ti dissolution, while more complex reactions may need to be further considered for more elaborate prediction. We note that Ti dissolution likely occurs by electrochemical reactions,<sup>31</sup> not chemical reactions<sup>29</sup> since (1) no significant dissolution of Ti was observed at potential hold at 1 V<sub>RHE</sub> before and after the protocols and during potential rest at OCP and (2) the Ti dissolution rate increased with increasing potential as shown in Protocol 2.

After confirmation of possible Ti dissolution from the bare Ti PTL under PEMWE-relevant operating conditions, we investigated the Ti dissolution behavior of the Ti PTL onto which thin protecting layers of Pt and IrO<sub>2</sub> were coated. These modified Ti PTLs are commonly utilized in the PEMWE to mitigate the formation of Ti oxide layers and consequently suppress the increase in contact resistance between the catalyst layer and the PTL.<sup>12,32–35</sup> In this study, we employed commercially available Pt- and IrO<sub>2</sub>-coated Ti PTLs, fabricated on identical Ti PTL substrates, as those investigated in Fig. 2. SEM and EDS studies confirmed the homogeneous dispersion of Pt and IrO<sub>2</sub> over the Ti PTLs (Fig. 4a and b), thickness of which was approximately 100 and 350 nm (Fig. S9†), respectively.

Online ICP-MS results on Pt- and IrO<sub>2</sub>-coated Ti PTLs revealed a considerably reduced Ti dissolution rate compared to that of the bare Ti PTL (Fig. 4c; see S10† for the corresponding current densities). The total Ti dissolution amounts during the protocols were 125 ng g<sub>Ti</sub><sup>-1</sup> for the bare Ti PTL, while Pt- and IrO<sub>2</sub>-coated Ti PTLs recorded only 39 and 4 ng g<sub>Ti</sub><sup>-1</sup>, respectively. Despite the effective protection provided by the IrO<sub>2</sub> layer against Ti dissolution in our case, a direct comparison between Pt- and IrO<sub>2</sub>-coated Ti PTLs and the subsequent discussion on identifying a better protective layer may be less informative, as these modified Ti PTLs exhibit considerable differences in surface morphology (*e.g.*, thickness of the protective layer; Fig. S9†) and noble metal content (Table S1†). More case studies with Pt/IrO<sub>2</sub>/or other protecting layer-coated Ti PTLs will be required to figure out the optimal composition and morphology of the protecting layer. However, at the very least, our results clearly demonstrate that the surface coating strategy is beneficial in mitigating undesirable Ti dissolution from Ti PTLs during PEMWE operation, in addition to the well-known electrical conductivity enhancement.

Our next question is then regarding the stability of the protecting layers during the operation, since even noble Pt and IrO<sub>2</sub> are not immune to corrosion in such highly anodic environments.<sup>14–18,35–39</sup>

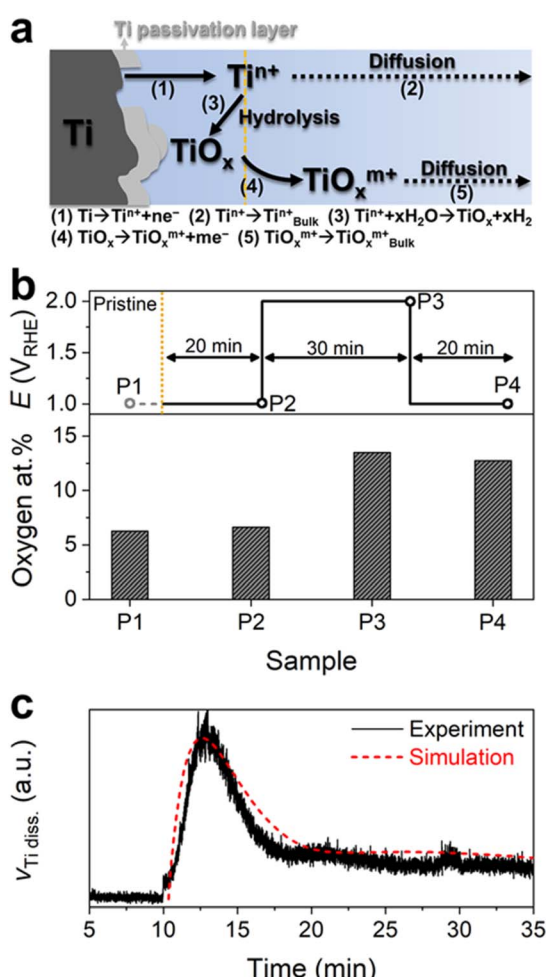


Fig. 3 (a) Schematic representation of the expected Ti PTL dissolution mechanism under highly anodic conditions. The reaction equation of each step is also shown. (b) Oxygen atomic percentages on the Ti PTL before (P1; gray circle) and after specific potential application (P2, P3, and P4; black circles) in O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub>. Oxygen content was estimated by EDS analysis. (c) Comparison of experimental and simulated Ti dissolution profiles at 2 V<sub>RHE</sub>.



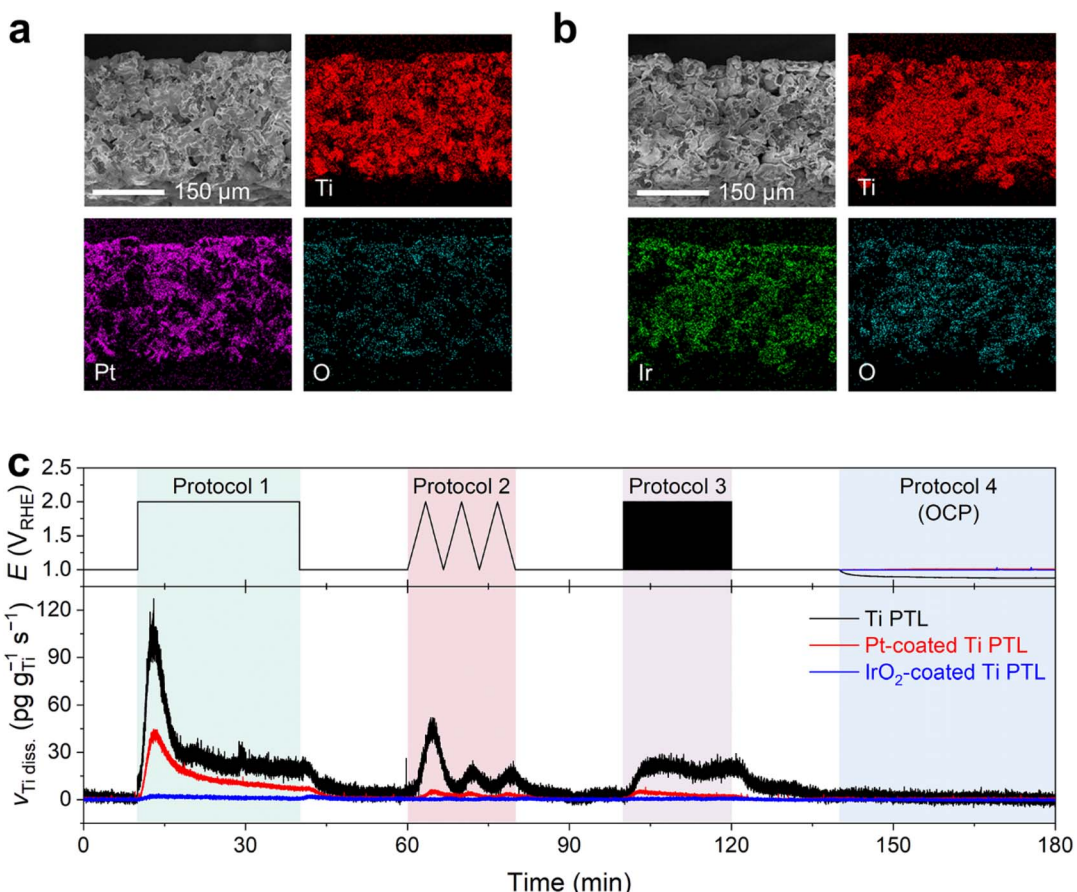


Fig. 4 Cross-sectional SEM and EDS results of (a) Pt- and (b) IrO<sub>2</sub>-coated Ti PTLs. (c) Real-time Ti dissolution profiles of Pt/IrO<sub>2</sub>-coated Ti PTLs measured during potential hold (Protocol 1), 3 cycles of CV (Protocol 2), potential pulses (Protocol 3), and resting at OCP (Protocol 4) with a supply of O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> feedstock.

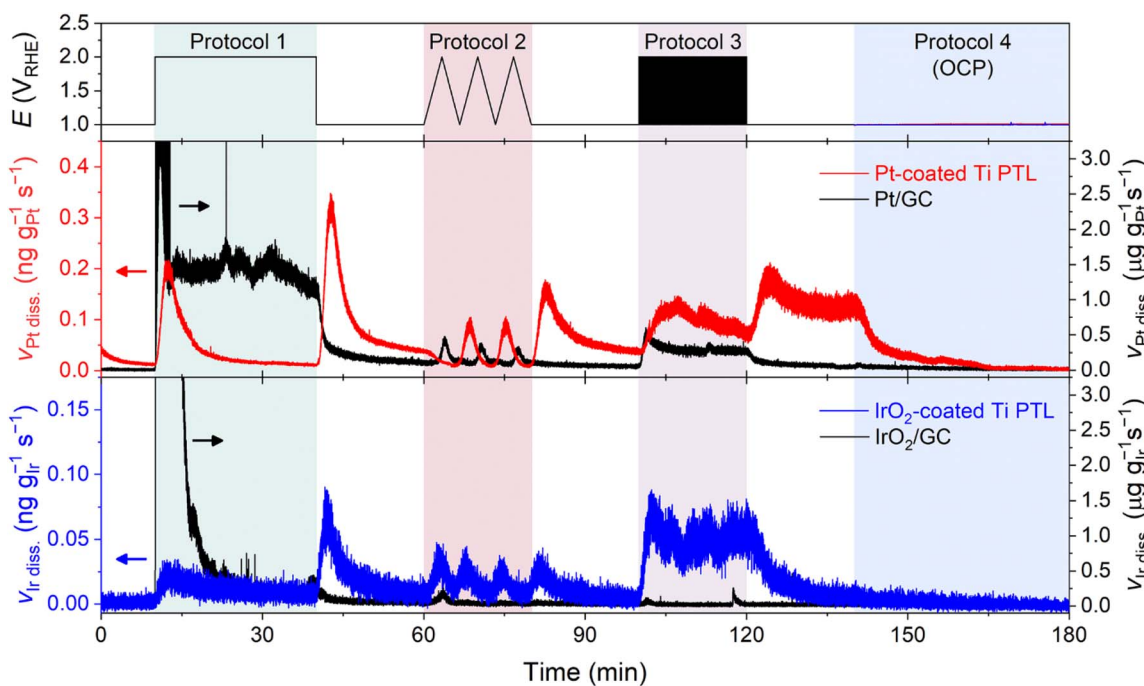


Fig. 5 Real-time dissolution profiles of Pt and Ir from Pt/IrO<sub>2</sub>-coated Ti PTLs measured during potential hold (Protocol 1), 3 cycles of CV (Protocol 2), potential pulses (Protocol 3), and resting at OCP (Protocol 4) with a supply of O<sub>2</sub>-saturated 0.1 M HClO<sub>4</sub> feedstock. For comparison, dissolution profiles of commercially available Pt and IrO<sub>2</sub> nanoparticles deposited on a GC electrode were measured using an electrochemical flow cell at 25 °C.

Fig. 5 displays Pt and Ir dissolution profiles measured upon the modified Ti PTLs during Protocols 1–4 (see Fig. S10† for the corresponding current densities). The results showed non-negligible dissolution of both protecting layers during the potential protocols. For the Pt-coated Ti PTL, Pt dissolution was found during potentiodynamic operation between 1 and 2 V<sub>RHE</sub>, but interestingly a significant Pt dissolution rate was also recorded during the potential hold at 1 V<sub>RHE</sub> that followed the end of each protocol. It is noteworthy that the Pt dissolution profile was different from that of conventional Pt particles deposited on a glassy carbon (GC) electrode, for which no considerable Pt dissolution was identified during the potential hold at 1 V<sub>RHE</sub>. Indeed, for the IrO<sub>2</sub>-coated Ti PTL, Ir dissolution occurred mainly during potentiodynamic operation, *i.e.*, at the beginning and end of Protocol 1 and during Protocols 2 and 3, while conventional IrO<sub>2</sub> showed much less Ir dissolution throughout the measurement except at the beginning of Protocol 1. Such fully different dissolution behaviors of Pt and IrO<sub>2</sub> in Ti PTLs, compared to those of conventional Pt and IrO<sub>2</sub>, thus emphasize that the identity of the supporting electrode or substrate strongly influences the stability of Pt and IrO<sub>2</sub>, as also pointed out recently by Zlatar *et al.*<sup>40</sup>

## Conclusions

In summary, we have investigated the dissolution of Ti PTLs in a PEMWE by integrating it with an online ICP-MS. We have shown that Ti dissolution occurs when the potential exceeds 1 V<sub>RHE</sub> under both potentiostatic and potentiodynamic conditions. It is important to note, however, that our use of acidic 0.1 M HClO<sub>4</sub> feedstock may exaggerate Ti dissolution rates compared to practical PEMWE operations, which typically use neutral DI water. In practical settings, neutral DI water facilitates rapid hydrolysis of dissolved Ti ions, resulting in the precipitation of TiO<sub>x</sub> and preventing the escape of Ti ions into the bulk feedstock (Fig. 1c). On the other hand, the use of 0.1 M HClO<sub>4</sub> also places the entire Ti PTL in an acidic environment, which can cause the entire PTL surface to corrode. Despite this limitation, even limited Ti dissolution at the electrode–membrane interface, a region that can become highly acidic during high current operations, could significantly degrade PEMWE performance. In more detail, the dissolved Ti ions can precipitate on the anode and cathode as TiO<sub>x</sub>, blocking the active surface of catalyst layers and reducing the electrochemically active surface area.<sup>41</sup> Since TiO<sub>x</sub> is an insulator, its accumulation can also lead to loss of electrical contact with active catalytic entities.<sup>11,12</sup> Furthermore, the possible retention of dissolved Ti cations on the membrane through strong electrostatic interaction with anionic functional groups (*e.g.*,  $-\text{SO}_3^-$  in Nafion) can hinder efficient proton transport and reduce cell performance.<sup>42,43</sup> While protective layers on Ti PTLs, *i.e.*, Pt and IrO<sub>2</sub>, can mitigate Ti dissolution to some extent, they can also dissolve under highly anodic conditions, and their dissolution behavior is significantly different from that of conventional Pt and IrO<sub>2</sub>. This discrepancy thus highlights the critical importance of stability measurements directly on practically relevant electrode materials, rather than on well-understood and relatively simple model substances, for

accurate evaluation of their stability. In addition, while we have investigated three commercialized Ti PTLs in this study, a similar approach can be extended to other PTL electrodes with different protective layer loadings, morphologies, fabrication methods, *etc.*, which may affect the dissolution rates of Ti and protective layers.<sup>44,45</sup> Therefore, more comprehensive studies are needed to deepen our understanding and ultimately establish a better design of new Ti PTLs. Our results and the analytical platform used in this study lay the groundwork for such future research.

## Experimental methods

### Online ICP-MS measurements

The anode was a commercial Ti PTL or Pt-/IrO<sub>2</sub>-modified Ti PTL, all provided by Shinsung C&T. The PTLs were tailored to 0.36 cm<sup>2</sup> (0.6 cm × 0.6 cm) and directly introduced into the PEMWE without any OER catalysts. On the other hand, the cathode was a GDL (0.36 cm<sup>2</sup>), on which Pd/C (20 wt% Pd, Sigma-Aldrich) was spray-coated with a target loading of 1 mg<sub>cat.</sub> cm<sup>-2</sup>. To prepare the GDL, a highly hydrophobic carbon mesoporous layer (MPL) was deposited on a carbon paper (MGL280, AvCarb<sup>®</sup>) by spraying an ink, prepared by dispersing 100 mg of Ketjen black EC-300J (KB) and 200 mg of polytetrafluoroethylene (PTFE; 60 wt%, Sigma-Aldrich) in 20 mL of isopropyl alcohol (99.5%, Sigma-Aldrich). The KB content in the MPL was adjusted to 2 mg cm<sup>-2</sup>. The carbon paper was then sequentially heat-treated at 240 and 340 °C under an Ar flow for 30 min each.

Online metal dissolution was analyzed using an ICP-MS (iCAP RQ, Thermo Fisher Scientific) coupled to a homemade PEMWE (Fig. 1a), which verified the comparable performance to other commercial PEMWEs when using a commercial membrane electrode assembly (MEA; Boyaz energy; anode – 4 mg cm<sup>-2</sup> IrO<sub>2</sub> and cathode – 0.5 mg<sub>pt</sub> cm<sup>-2</sup> Pt/C) (Fig. S2†). The anode and cathode were separated by a Nafion 115 membrane (DuPont), which was extended from the PEMWE and connected to a saturated Ag/AgCl reference electrode to control the bias potential using a potentiostat (VMP3, Bio-Logic Science Inc).<sup>20</sup> To avoid unwanted Ti dissolution from a conventional Ti bipolar plate, feed flow channels were fabricated with polyether ether ketone (PEEK). Since the PEEK is an electrical insulator, an Au wire was connected to each electrode to provide electrical contact. All these compartments were assembled in the following order — back plate, flow channel, Au wire, Ti PTL, gasket, Nafion, gasket, Pd/C GDE, Au wire, flow channel, and back plate (Fig. 1a) — and then tightened with six bolts to a torque of 18 Nm. Heating rods were inserted into the back plates of the PEMWE to control the cell temperature, and ICP-MS studies were performed at 25 and 80 °C.

The feedstock was either O<sub>2</sub>-saturated DI water (>18.2 MΩ, Arium<sup>®</sup> Pro, Sartorius) or 0.1 M HClO<sub>4</sub> (diluted from 70% HClO<sub>4</sub>, Sigma-Aldrich), which flowed through each PEEK flow channel at a flow rate of 5 mL min<sup>-1</sup> using a peristaltic pump (LEPP-150L, Labscitech). Prior to introduction of the feedstock into the ICP-MS, 400 μL min<sup>-1</sup> of the outflow was separated and mixed with 0.5 M HNO<sub>3</sub> containing 5 ppb <sup>187</sup>Re as an internal standard at a 1:1 mixing ratio using a Y-connector. Online metal dissolution was estimated from the ratio of metal ions



( $^{48}\text{Ti}$ ,  $^{193}\text{Ir}$ , and  $^{195}\text{Pt}$ ) to  $^{187}\text{Re}$  signals during electrochemical measurements.

In control experiments, the dissolution of Pt and  $\text{IrO}_2$  nanoparticles was investigated at 25 °C using a homemade electrochemical flow cell, which was used in our previous work.<sup>46</sup> Briefly, the cell consisted of a U-shaped channel (1 mm diameter) and two openings (3 mm diameter). On one opening side, a mirror-polished 3 mm GC electrode (002 012, ALS) was introduced as a working electrode, on which 50  $\mu\text{g cm}^{-2}$  of Pt black (HiSPEC 1000, Alfa Aesar) or  $\text{IrO}_2$  powder (Alfa Aesar) was deposited by drop-casting of an ink, prepared by dispersing 10 mg of Pt black or  $\text{IrO}_2$  powder in a mixed solution of DI water (5894  $\mu\text{L}$ ), isopropyl alcohol (1079  $\mu\text{L}$ ), and 5 wt% Nafion solution (100  $\mu\text{L}$ ). On the other opening side, a 3 mm Teflon tube sealed at one end with a PTFE membrane (WP-020-80, Sumitomo Electric Ind., Ltd) was brought up to the working electrode surface to extract the evolved gases by vacuum. The counter electrode was a graphite rod separated from the electrolyte by a Nafion 115 membrane. The reference electrode was a saturated Ag/AgCl electrode connected directly to the electrolyte outlet.

### Local pH measurements

The local pH change at the anode was measured by a method recently developed by Sauvé *et al.*,<sup>25</sup> which monitors the OCP decay transient (potential of the  $\text{H}_2/\text{H}^+$  equilibrium) on Pt. Our homemade PEMWE was used as the electrochemical platform, but a commercial Pt/C MEA (Boyaz energy) was installed. The MEA consisted of a Nafion 115 membrane, and Pt/C was deposited on both sides of the membrane with a loading of 0.5  $\text{mg}_{\text{Pt}} \text{cm}^{-2}$ . A GDL and a Pt-coated Ti PTL were attached to the anode and cathode sides, respectively.  $\text{H}_2$  gas, humidified at 50 °C, was introduced into the anode side at a flow rate of 200 sccm using a mass flow controller (M3030VA with an LTI1000 readout box, LineTech), and Ar-saturated DI water was introduced into the cathode side at a flow rate of 5  $\text{mL min}^{-1}$ . After stabilization of the potential signals at the specified *j*, which ranged from 0.0 to 2.2  $\text{A cm}^{-2}$ , the OCP decay transient was recorded with a data collection interval of 0.2 ms for 1 s. The local pH was calculated using eqn (1) with the OCP value measured at 20 ms after the potential transient to OCP.

$$\text{Local pH} = E_{\text{OCP}@20 \text{ ms}} (\text{V}_{\text{SHE}}) / -0.070 \text{ (@80 } ^\circ\text{C)} \quad (1)$$

### Physical characterization

SEM and EDS analyses were performed at an accelerating voltage of 5 and 15 kV, respectively, using a HITACHI S-4800. For cross-sectional SEM and EDS analyses, PTL samples were prepared by immersion in liquid  $\text{N}_2$  followed by cutting. The noble metal contents on the modified Ti PTLs were measured using an X-ray fluorescence spectrometer (SIMULTIX12; RIGAKU).

### Computational modeling

A computational modeling framework was developed using MATLAB to analyse the dynamic behaviour of Ti PTL dissolution, described by a system of conditional ordinary differential

equations. These equations model the concentration changes of ions and predict the real-time Ti dissolution profile, thereby introducing nonlinear dynamics. The “ode45” solver in MATLAB 2023a was employed for its robust handling of both stiff and non-stiff ordinary differential equation systems, ensuring accurate simulations across varied parameter sets. To refine the model and align it with experimental observations, genetic algorithm optimization was utilized to adjust the kinetic parameters, aiming to minimize the discrepancy between the predictions and experimental data.

### Data availability

Data for this article are available at the Zenodo repository at <https://doi.org/10.5281/zenodo.13117968>.

### Author contributions

Junsic Cho: formal analysis, investigation, visualization, writing – original draft; Dong Hyun Kim: formal analysis, investigation, visualization, writing – original draft; Min Wook Noh: validation; Haesol Kim: methodology; Hong-Gyun Oh: resources; Pilyoung Lee: resources; Soobin Yoon: investigation; Wangyun Won: investigation; Young-June Park: conceptualization, project administration, funding acquisition; Ung Lee: formal analysis, data curation, writing – original draft; Chang Hyuck Choi: writing – original draft, supervision, project administration.

### Conflicts of interest

There are no conflicts to declare.

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### References

- 1 S. Chu, Y. Cui and N. Liu, *Nat. Mater.*, 2017, **16**, 16–22.
- 2 S. E. Hosseini and M. A. Wahid, *Renewable Sustainable Energy Rev.*, 2016, **57**, 850–866.
- 3 H. Jin, B. Ruqia, Y. Park, H. J. Kim, H.-S. Oh, S.-I. Choi and K. Lee, *Adv. Energy Mater.*, 2021, **11**, 2003188.
- 4 M. Bonanno, K. Müller, B. Bensmann, R. Hanke-Rauschenbach, D. Aili, T. Franken, A. Chromik, R. Peach, A. T. S. Freiberg and S. Thiele, *Renewable Sustainable Energy Rev.*, 2024, **9**, 2300281.
- 5 N. D. Tomashov, R. M. Altovsky and G. P. Chernova, *J. Electrochem. Soc.*, 1961, **108**, 113.
- 6 O. Panchenko, E. Borgardt, W. Zwaygardt, F. J. Hackemüller, M. Bram, N. Kardjilov, T. Arlt, I. Manke, M. Müller, D. Stolten and W. Lehnert, *J. Power Sources*, 2018, **390**, 108–115.



7 J. K. Lee, C. H. Lee and A. Bazylak, *J. Power Sources*, 2019, **437**, 226910.

8 T. Schuler, J. M. Ciccone, B. Krentscher, F. Marone, C. Peter, T. J. Schmidt and F. N. Büchi, *Adv. Energy Mater.*, 2020, **10**, 1903216.

9 T. L. Doan, H. E. Lee, S. S. H. Shah, M. Kim, C.-H. Kim, H.-S. Cho and T. Kim, *Int. J. Energy Res.*, 2021, **45**, 14207–14220.

10 D. H. Jeon, S. Kim, M. Kim, C. Lee and H.-S. Cho, *J. Power Sources*, 2023, **553**, 232322.

11 T. Srour, K. Kumar, V. Martin, L. Dubau, F. Maillard, B. Gilles, J. Dillet, S. Didierjean, B. Amoury, T. D. Le and G. Maranzana, *Int. J. Hydrogen Energy*, 2024, **58**, 351–361.

12 C. Rakousky, U. Reimer, K. Wippermann, M. Carmo, W. Lueke and D. Stolten, *J. Power Sources*, 2016, **326**, 120–128.

13 A. Lim, H.-Y. Jeong, Y. Lim, J. Y. Kim, H. Y. Park, J. H. Jang, Y.-E. Sung, J. M. Kim and H. S. Park, *Sci. Adv.*, 2021, **7**, eabf7866.

14 S. Cherevko, T. Reier, A. R. Zeradjanin, Z. Pawolek, P. Strasser and K. J. J. Mayrhofer, *Electrochem. Commun.*, 2014, **48**, 81–85.

15 M. van der Merwe, R. Garcia-Diez, L. Lahn, R. E. Wibowo, J. Frisch, M. Gorgoi, W. Yang, S. Ueda, R. G. Wilks, O. Kasian and M. Bär, *ACS Catal.*, 2023, **13**, 15427–15438.

16 O. Kasian, T. Li, A. M. Mingers, K. Schweinar, A. Savan, A. Ludwig and K. Mayrhofer, *J. Phys.: Energy*, 2021, **3**, 034006.

17 A. Lončar, D. Escalera-López, F. Ruiz-Zepeda, A. Hrnjić, M. Šala, P. Jovanović, M. Bele, S. Cherevko and N. Hodnik, *ACS Catal.*, 2021, **11**, 12510–12519.

18 L. Lahn, A. M. Mingers, A. Savan, A. Ludwig and O. Kasian, *ChemElectroChem*, 2024, **11**, e202300399.

19 J.-P. B. Haraldsted, Z. Révay, R. Frydendal, A. Verdaguér-Casadevall, J. Rossmeisl, J. Kibsgaard and I. Chorkendorff, *Mater. Today Energy*, 2019, **14**, 100352.

20 Q. Xu, S. Z. Oener, G. Lindquist, H. Jiang, C. Li and S. W. Boettcher, *ACS Energy Lett.*, 2021, **6**, 305–312.

21 C. Rakousky, U. Reimer, K. Wippermann, S. Kuhri, M. Carmo, W. Lueke and D. Stolten, *J. Power Sources*, 2017, **342**, 38–47.

22 J. C. Fornaciari, L.-C. Weng, S. M. Alia, C. Zhan, T. A. Pham, A. T. Bell, T. Ogitsu, N. Danilovic and A. Z. Weber, *Electrochim. Acta*, 2022, **405**, 139810.

23 S. Liu, I. Zaharieva, L. D'Amario, S. Mebs, P. Kubella, F. Yang, P. Beyer, M. Haumann and H. Dau, *Adv. Energy Mater.*, 2022, **12**, 2202914.

24 M. Pourbaix, *Atlas of Electrochemical Equilibria in Aqueous Solutions*, National Association of Corrosion Engineers, 1974.

25 E. R. Sauvé, B. Y. Tang, N. K. Razdan, W. L. Toh, S. Weng and Y. Surendranath, *Joule*, 2024, **8**, 728–745.

26 V. Kozlov, N. Bunkin, I. Ps, A. Shkirin, S. Zakharov and Z. Aa, *Water*, 2013, **4**, 129–154.

27 S. A. Berlinger, B. D. McCloskey and A. Z. Weber, *J. Phys. Chem. B*, 2018, **122**, 7790–7796.

28 Y. Fovet, J.-Y. Gal and F. Toumelin-Chemla, *Talanta*, 2001, **53**, 1053–1063.

29 J.-F. Vanhumbeeck and J. Proost, *Corros. Rev.*, 2009, **27**, 117–204.

30 L. Fiedler, T.-C. Ma, B. Fritsch, J. H. Risse, M. Lechner, D. Dworschak, M. Merklein, K. J. J. Mayrhofer and A. Hutzler, *ChemElectroChem*, 2023, **10**, e202300373.

31 R. Srinivasan and F. Fasmin, *An Introduction to Electrochemical Impedance Spectroscopy*, CRC Press, 2021.

32 C. Liu, M. Carmo, G. Bender, A. Everwand, T. Lickert, J. L. Young, T. Smolinka, D. Stolten and W. Lehnert, *Electrochem. Commun.*, 2018, **97**, 96–99.

33 C. Liu, M. Shviro, A. S. Gago, S. F. Zaccarine, G. Bender, P. Gazdzicki, T. Morawietz, I. Biswas, M. Rasinski, A. Everwand, R. Schierholz, J. Pfeilsticker, M. Müller, P. P. Lopes, R.-A. Eichel, B. Pivovar, S. Pylypenko, K. A. Friedrich, W. Lehnert and M. Carmo, *Adv. Energy Mater.*, 2021, **11**, 2002926.

34 T. L. Doan, T. N. Nguyen, Y. S. Jung, C. Lee, M. Kim, S. Lee, H.-S. Cho and T. Kim, *Int. J. Hydrogen Energy*, 2024, **55**, 839–847.

35 C. Rakousky, G. P. Keeley, K. Wippermann, M. Carmo and D. Stolten, *Electrochim. Acta*, 2018, **278**, 324–331.

36 S. Cherevko, S. Geiger, O. Kasian, A. Mingers and K. J. J. Mayrhofer, *J. Electroanal. Chem.*, 2016, **774**, 102–110.

37 O. Kasian, J.-P. Grote, S. Geiger, S. Cherevko and K. J. J. Mayrhofer, *Angew. Chem., Int. Ed.*, 2018, **57**, 2488–2491.

38 S. Cherevko, A. A. Topalov, A. R. Zeradjanin, G. P. Keeley and K. J. J. Mayrhofer, *Electrocatalysis*, 2014, **5**, 235–240.

39 A. A. Topalov, S. Cherevko, A. R. Zeradjanin, J. C. Meier, I. Katsounaros and K. J. J. Mayrhofer, *Chem. Sci.*, 2014, **5**, 631–638.

40 M. Zlatar, D. Escalera-López, M. G. Rodríguez, T. Hrbek, C. Götz, R. Mary Joy, A. Savan, H. P. Tran, H. N. Nong, P. Pobedinskas, V. Briega-Martos, A. Hutzler, T. Böhm, K. Haenen, A. Ludwig, I. Khalakhan, P. Strasser and S. Cherevko, *ACS Catal.*, 2023, **13**, 15375–15392.

41 S. Cherevko, *Curr. Opin. Electrochem.*, 2018, **8**, 118–125.

42 S. A. Grigoriev, D. G. Bessarabov and A. S. Glukhov, *Russ. J. Electrochem.*, 2017, **53**, 808–812.

43 Q. Feng, X. Z. Yuan, G. Liu, B. Wei, Z. Zhang, H. Li and H. Wang, *J. Power Sources*, 2017, **366**, 33–55.

44 C. Liu, K. Wippermann, M. Rasinski, Y. Suo, M. Shviro, M. Carmo and W. Lehnert, *ACS Appl. Mater. Interfaces*, 2021, **13**, 16182–16196.

45 C. Liu, J. A. Wrubel, E. Padgett and G. Bender, *Appl. Energy*, 2024, **356**, 122274.

46 J. Cho, T. Lim, H. Kim, L. Meng, J. Kim, S. Lee, J. H. Lee, G. Y. Jung, K.-S. Lee, F. Viñes, F. Illas, K. S. Exner, S. H. Joo and C. H. Choi, *Nat. Commun.*, 2023, **14**, 3233.

