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# Spectroscopic investigation of the electrochemical growth and stability of am-hydr-IrO<sub>x</sub> grown from IrTi alloys

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Amorphous hydrous iridium oxide thin films (am-hydr-IrO<sub>x</sub>) grown through electrochemical treatment of sputtered iridium thin films are an interesting class of materials for the oxygen evolution reaction (OER). They boast high electrochemical activity and high mass utilisation of the precious metal iridium; however, the application of am-hydr-IrO<sub>x</sub> is limited by poor stability caused by film delamination and electrochemically enhanced corrosion. This study presents a comprehensive investigation of the behaviour of am-hydr-IrO<sub>x</sub> grown from co-sputtered iridium with titanium (IrTi) films. The growth behaviour of IrTi films is investigated in detail by employing X-ray photoelectron spectroscopy, which enables unique insights into the structure and mechanism of the resultant am-hydr-IrO<sub>x</sub> species. Depth-dependent hard X-ray photoelectron spectroscopy (HAXPES) experiments using different excitation energies show that the films separate into an iridium rich hydrous oxide layer supported by an unreacted metallic alloy during growth. Complementary element specific *operando* X-ray absorption spectroscopy studies at the Ir L<sub>3</sub>- and Ti K-edges in combination with electrochemical techniques are further used to ascertain details of the growth mechanisms and develop a hypothesis to connect the electrocatalyst structure to its electrochemical stability under hydrolysis-relevant conditions. Titanium oxide formation at the surfaces acts to impede the formation of am-hydr-IrO<sub>x</sub>, and above a critical concentration, the presence of titanium results in an impressive stability enhancement.

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## 1. Introduction

Proton exchange membrane water electrolyzers (PEM-WEs) are considered to be one of the leading technologies for industrial scale water splitting.<sup>1,2</sup> However, widespread use is limited by the stability of electrocatalyst materials to drive the oxygen

evolution reaction. Due to the harsh, highly oxidizing, and low pH conditions, PEM-WE anodes typically utilise materials containing precious metals such as iridium.<sup>3</sup> The employment of water electrolysis involves coupling of intermittent renewable energy sources with electrolyzers.<sup>4</sup> As a result, electrocatalysts need to be stable also under (to some degree) fluctuating operation schemes. There are several different proposed mechanisms that contribute to the performance loss of iridium-based electrocatalyst materials, each contributing to different extents, based on the electrocatalyst's electronic and crystal structure.<sup>5</sup> For example, potential dependent dissolution mechanisms such as the formation of highly soluble, high valence iridium (Ir<sup>VI</sup>) species<sup>6</sup> or low coordinated iridium sites *via* lattice oxygen evolution have been proposed.<sup>7</sup> Other processes detrimental to the performance of Ir-based anode electrocatalysts are the poorer electronic conductivity of the electrocatalyst after prolonged operation due to discontinuities in the supported active layer<sup>8</sup> or the increased contact resistance with the Ti-based porous transport layer.<sup>9</sup>

Lately, amorphous, hydrous iridium oxides (am-hydr-IrO<sub>x</sub>) have caught attention for acidic water electrolysis due to their high activity, high iridium active site availability and acid

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tolerance.<sup>10–12</sup> The am-hydr-IrO<sub>x</sub> structures can be generated through electrochemical cycling of metallic iridium substrates generating a highly porous, hydrated iridium oxide layer.<sup>13–15</sup> However, compared to crystalline iridium oxide, am-hydr-IrO<sub>x</sub> suffers from poorer stability due to the thermodynamically driven dissolution of the active metal at high overpotentials and low pH.<sup>16–19</sup> In addition, recent work predicts Ir dissolution in am-hydr-IrO<sub>x</sub> by spontaneous formation of iridium vacancies occurring at potentials as low as 0.9 V vs. the reversible hydrogen electrode (RHE).<sup>20</sup>

Mechanical stability of the porous film can be influenced by excessive growth of am-hydr-IrO<sub>x</sub>, where the gas evolution in the porous oxide network can lead to partial or total delamination of the active area from the metallic Ir substrate.<sup>21,22</sup> Mechanical stability is particularly relevant for am-hydr-IrO<sub>x</sub>, as varying voltage from intermittent renewable energy sources coupled to the PEM-WE could lead to further undesirable growth and eventual delamination of the electrocatalyst layer. To improve their stability, many active anode materials are combined with acid-stable oxides either as a support or by partial substitution into the crystal lattice. Oxides of titanium and tin are particularly popular, showing improved stability against dissolution and having positive synergies in combination with iridium.<sup>23–25</sup> However, as they are inactive towards the OER, their combination with active materials often comes at the cost of lower performance.<sup>24,26–28</sup> A more profound understanding of the interaction between, *e.g.* titanium and iridium in its amorphous hydrous form may lead to strategies to stabilize the highly catalytic am-hydr-IrO<sub>x</sub>.

In this work, we investigate the behaviour of hydrous iridium oxide thin films grown from alloys of iridium and titanium of various compositions. The complex growth behaviour of the highly hydroxylated porous electrocatalysts requires characterisation at various stages of the life of the electrocatalyst film. Here, lab-based and synchrotron-based advanced X-ray spectroscopic methods were used to provide depth-resolved information on the chemical composition of the films as they evolve throughout the electrochemically induced growth procedures. In addition to thorough spectroscopic characterisation of the films, the activity and stability of the films were studied and evaluated using a variety of electrochemical techniques. With these results, the role of titanium in the vicinity of iridium atoms during the electrochemical growth of am-hydr-IrO<sub>x</sub> is thoroughly investigated for different Ti concentrations, aiming to reveal the processes triggered by titanium that can improve the stability of iridium-based OER electrocatalysts.

## 2. Results

### 2.1 Characterisation of the pristine IrTi alloy films

Thin films of IrTi alloys were prepared by co-sputtering iridium and titanium onto silicon wafers, yielding surface metal elemental ratios ranging from pure iridium (Ir<sub>100</sub>) to 74 at% Ir (Ir<sub>74</sub>Ti<sub>26</sub>). These elemental surface ratios were derived from the Ti 2p (including metallic and oxidised species) and Ir 4f X-ray photoelectron spectroscopy (XPS) detail spectra shown in Fig. 1a and b. Note that the Ir 4f spectra overlap with the Ti 3s and Ir 5p<sub>1/2</sub> core

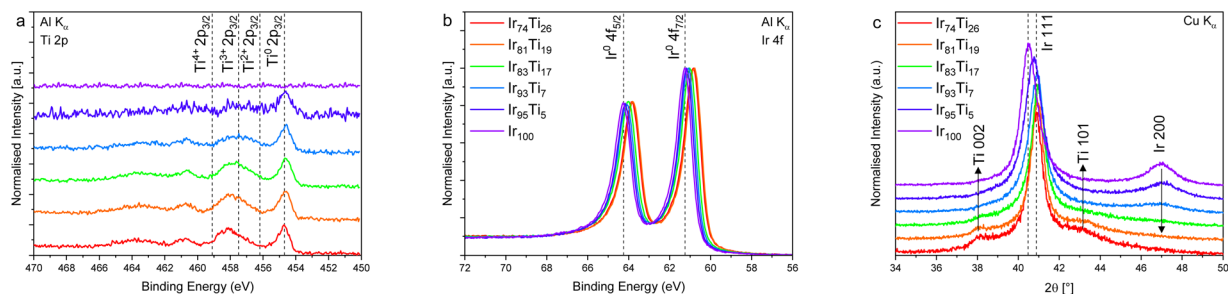
level lines,<sup>23</sup> but their spectral contributions are considered to be negligible when probed with Al K<sub>α</sub>, as in this case their photoionization cross sections are more than one order of magnitude smaller than that of Ir 4f. Details on the quantification of the XPS data are described in the SI (Fig. S4 and Table S1). Fig. S2 shows the survey spectra of the pristine films with the different photoemission lines of the constituent elements labelled. The Ti 2p core level spectra show that titanium exists in multiple chemical states dominated by metallic Ti and oxidised Ti species (Fig. 1a). This observation is consistent with previous reports.<sup>23</sup> The ratio of metallic and oxidised titanium species varies throughout the samples whereby samples with high total titanium content have higher relative proportions of oxidised Ti species. This was quantified by deconvolution of the Ti 2p spectra (Fig. S6) and the relationship with respect to the elemental surface ratio is shown in Fig. S7. The Ir 4f spectra show a shift towards lower binding energies with increasing titanium concentration (Fig. 1b), supporting the argument that a solid solution type alloy is also being formed. Similar effects have been reported with a shift of up to 4 eV to lower binding energies for an equimolar composition.<sup>30</sup> Although, by analogy, a shift up to 2 eV could be expected between Ir<sub>100</sub> and Ir<sub>74</sub>Ti<sub>26</sub>, we only observe a shift of  $\approx -0.5$  eV, consistent with the scenario that parts of the titanium are not completely incorporated into the lattice but are segregated within the iridium matrix, thus, not contributing to the chemical shift of Ir 4f.

X-ray reflectometry indicates a rather uniform film thickness of  $33 \pm 4$  nm for all films independent of Ti content (see Fig. S5 and Table S2). Phase purity and crystal structure of the pristine films were determined with grazing incidence X-ray diffraction (GI-XRD) as shown in Fig. 1c. It is observed that the addition of titanium into the iridium lattice results in a shift of the main Ir (111) reflection to higher angles, from 40.5° (Ir<sub>100</sub>) up to 40.9° (Ir<sub>74</sub>Ti<sub>26</sub>). This is consistent with the successful formation of an alloy, where the introduction of a smaller-radius element leads to a decrease in the associated lattice parameter; however, strain effects in the thin films cannot be excluded. The latter (more specifically directional lattice strain) could also explain why at higher titanium concentrations the more prominent Ti (002) reflex associated with the titanium alpha phase is observed at around 38° (as expected) while the Ti (101) reflex is significantly shifted to higher angles and observed at 43°. The appearance of the peaks in GI-XRD is related to the segregation of a long-range ordered titanium phase (which could be a nanocluster in the alloy, although a segregated phase at the surface could not be excluded). Alloys can be understood as either (a) solid solutions, where there is a single crystal structure with multiple elements occupying the crystallographic sites, or, alternatively, (b) a mixture of two or more metallic phases, where one forms a sub-structure within the parent phase.<sup>30</sup> For these IrTi thin film materials, it appears that both types of alloys are simultaneously present and, therefore, the metallic pristine IrTi films will be referred to as alloys.

### 2.2 Electrochemical growth from the IrTi alloy films

The am-hydr-IrO<sub>x</sub> electrocatalyst layers were grown from their respective metallic IrTi alloy films using methodology reported





**Fig. 1** Characterisation of pristine films. (a) Al  $K_{\alpha}$  (1486.71 eV) excited Ti 2p XPS detail spectra of IrTi films with varying compositions (see the colour code); dotted vertical lines highlight the reference binding energies of the Ti  $2p_{3/2}$  line, consisting of metallic titanium ( $Ti^0$ ) and oxidised titanium species ( $Ti^{2+}$ ,  $Ti^{3+}$ , and  $Ti^{4+}$ ). (b) Corresponding Al  $K_{\alpha}$  excited Ir 4f XPS detail spectra; dotted vertical lines represent the reference binding energies of the spin-orbit doublet of the Ir 4f line for metallic iridium ( $Ir^0$ ). (c) Grazing incident X-ray diffraction patterns (GI-XRD); dotted vertical lines highlight the shift in peak position of the Ir (111) reflex between the  $Ir_{100}$  and the  $Ir_{74}Ti_{26}$  sample. Arrows highlight the decrease of the Ti (002) and (101) reflexes with increasing iridium content and the appearance of a secondary iridium reflection.

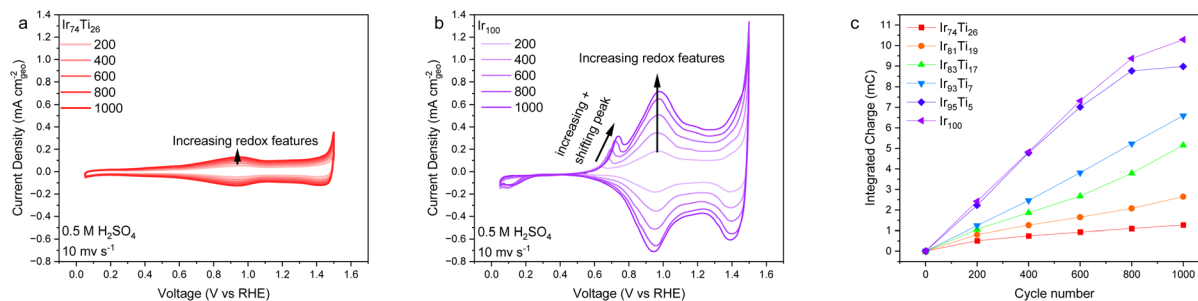
elsewhere.<sup>12</sup> In brief, the IrTi alloys were voltammetrically cycled between reductive and oxidative potentials at  $500 \text{ mV s}^{-1}$  up to 1000 total cycles.<sup>15</sup> After every increment of 200 cycles, a cyclic voltammogram (CV) was measured at a slower rate ( $10 \text{ mV s}^{-1}$ ). The slow CVs of the compositional extremes of the IrTi alloy films,  $Ir_{74}Ti_{26}$  and  $Ir_{100}$ , are shown in Fig. 2a and b respectively, illustrating a significant difference in the growth behaviours, where the pure iridium sample ( $Ir_{100}$ ) exhibits a significantly higher growth rate than its alloy counterpart ( $Ir_{74}Ti_{26}$ , the complete set of electrochemical profiles is shown in Fig. S9). This is seen in the increase in the capacitance and pseudocapacitive-redox with respect to cycle number, highlighted by the arrows in panels (a) and (b) of Fig. 2. This growth reflects an increased surface area with an increased number of electrochemically available iridium sites.

To study this effect further, the areas of the redox features were quantified by integrating the reductive portion of the peak around  $0.95 \text{ V vs. RHE}$  to avoid influence from the small redox feature seen around  $0.7 \text{ V vs. RHE}$ . The number of electrochemically available sites was estimated from the charge, which was calculated from the area of the redox feature and is plotted as a function of cycle number in Fig. 2c. The area of the redox feature was derived using a linear background to remove capacitive effects between  $0.6$  and  $1.2 \text{ V vs. RHE}$ . A clear relationship is observed between the rate of growth with respect to

the number of electrochemical growth cycles and the elemental composition. An increasing titanium content apparently prevents the participation of iridium sites in the electrochemical growth (limiting it), as interpreted from the reduced voltametric charge. This results into a slower growth rate for the films with higher Ti content, as documented by the smallest voltametric charge observed for the  $Ir_{74}Ti_{26}$  thin film. For simplicity, the different numbers of electrochemically available iridium sites, *i.e.* sites within the am-hydr- $IrO_x$  electrocatalysts grown from the different alloy films, will be referred to as “thickness” but this does not necessarily describe a closed layer supported by the metal sublayer. However, it has been shown to linearly correlate with film thickness.<sup>20</sup> Despite titanium having a clear impact on the growth rate of the amorphous hydrous iridium layer, it does not show direct effects on the electrochemical signature of the amorphous hydrous iridium oxide, reflected by the feature around  $0.95 \text{ V vs. RHE}$  observed in all samples. A similar shift of the small anodic peak found around  $0.7 \text{ V vs. RHE}$  as a function of the growth is observed in all samples, as seen in Fig. S9.

### 2.3 Depth-dependent chemical analysis of 1000-cycle grown am-hydr- $IrO_x$ layer films

To probe the chemical composition of the different alloy films grown up to 1000 cycles, they were studied using hard X-ray



**Fig. 2** Electrochemically induced growth behaviour measured by cyclic voltammetry at 200, 400, 600, 800 and 1000 cycles at a scan rate of  $10 \text{ mV s}^{-1}$  in  $0.5 \text{ M H}_2\text{SO}_4$  during the 1000-cycle growth of (a) the  $Ir_{74}Ti_{26}$ , and (b) the  $Ir_{100}$  thin film. (c) Quantified growth behaviour for all studied IrTi films of varying Ti content derived from the integral of the pseudocapacitance of the observed reductive peak at  $\approx 0.95 \text{ V vs. RHE}$  as a function of cycle number.



photoelectron spectroscopy (HAXPES) at the HiKE endstation.<sup>31</sup> Fig. 3a shows the Ir 4f spectra of the samples measured with 2 keV, with pristine Ir<sub>74</sub>Ti<sub>26</sub> as a reference. Dashed lines highlight the positions of the Ir 4f<sub>7/2</sub> contributions for both IrO<sub>2</sub> and metallic iridium. All samples show only evidence of oxidised iridium except for am-hydr-IrO<sub>x</sub> grown from Ir<sub>74</sub>Ti<sub>26</sub>, which also

shows peaks associated with metallic iridium. The Ir 4f region was also measured using a higher excitation energy (6 keV), as shown in Fig. 3b, and, therefore, with a higher probing depth. These less surface-sensitive measurements revealed more unreacted metallic iridium, hinting at a layered structure where the am-hydr-IrO<sub>x</sub> grows on top of the unreacted parts of the

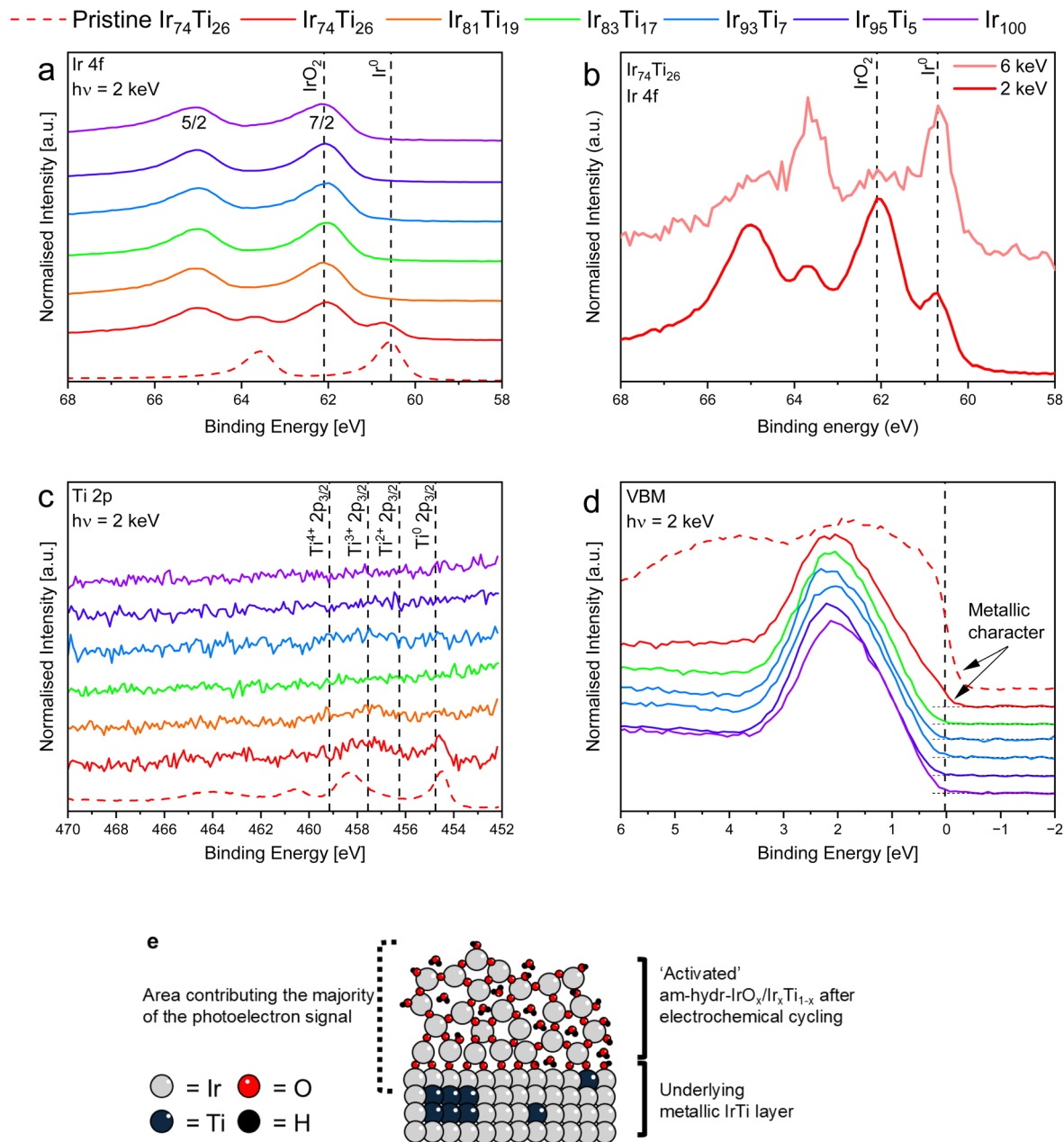


Fig. 3 Hard X-ray photoelectron spectroscopy of 1000-cycle grown am-hydr-IrO<sub>x</sub> (a) Ir 4f HAXPES detail spectra recorded at 2 keV excitation energy. The dotted lines highlight the Ir 4f<sub>7/2</sub> binding energy position of metallic and oxidized iridium. (b) Ir 4f spectra of the am-hydr-IrO<sub>x</sub> layer grown on the Ir<sub>74</sub>Ti<sub>26</sub> alloy film (*i.e.*, the sample with the highest surface titanium content in the pristine alloy film), measured with both 2 and 6 keV. The dotted lines highlight the Ir 4f<sub>7/2</sub> binding energy position of metallic and oxidized iridium. (c) Ti 2p spectra recorded with an excitation energy of 2 keV. The dotted lines highlight the Ti 2p<sub>3/2</sub> binding energies for metallic titanium (Ti<sup>0</sup>) and oxidised species. (d) HAXPES spectra of the valence band maxima. The vertical dotted line indicates the position of the Fermi level (0 eV binding energy). (e) Schematic of the structural model proposed within this work, whereby the am-hydr-IrO<sub>x</sub> layer grows from the IrTi films and is the major contributor to the HAXPES signal, except for the sample with the thinnest am-hydr-IrO<sub>x</sub> top layer (Ir<sub>74</sub>Ti<sub>26</sub>). For this sample, the contribution from the metallic alloy layer is clearly observed in the HAXPES data.



alloy film. Note that the Ir 4f spectra overlap with the Ti 3s and Ir 5p<sub>1/2</sub> core level lines,<sup>23</sup> but their spectral contributions are considered to be negligible when probed with 2 keV (see explanation above). However, for the 6 keV excited Ir 4f spectra, Ti 3s and Ir 5p<sub>1/2</sub> derived spectral contributions have to be considered (especially when quantifying the data), but this does not affect the qualitative statement that more unreacted metallic iridium is present in the less surface-sensitive measurements made above.

Considering that the HAXPES detection limit is between 0.1 and 1 at% (depending on the photoionization cross section of the probed core level), it is worth noting that a Ti 2p HAXPES signal is only detected for the am-hydr-IrO<sub>x</sub> film grown from the Ir<sub>74</sub>Ti<sub>26</sub> film (Fig. 3c). This suggests that Ti is largely absent from the grown am-hydr-IrO<sub>x</sub> layer and the fact that Ti is only detected for the Ir<sub>74</sub>Ti<sub>26</sub> based film can be explained by considering the am-hydr-IrO<sub>x</sub> layer grown from the Ir<sub>74</sub>Ti<sub>26</sub> film as the thinnest one (see Fig. 2c).

To support this hypothesis, HAXPES detail spectra of the valence band maximum (VBM) were also studied to see if there were any significant electronic differences in the am-hydr-IrO<sub>x</sub> grown from IrTi alloy films with different compositions, since small amounts of titanium could influence the electronic structure while not being directly detectable. Fig. 3d shows the measured data of the grown am-hydr-IrO<sub>x</sub> films, with the pristine Ir<sub>74</sub>Ti<sub>26</sub> film also included as a reference.

All am-hydr-IrO<sub>x</sub> layers again show very similar spectral valence band features, except for the one grown from the Ir<sub>74</sub>Ti<sub>26</sub> film having a shallower slope of the leading valence band edge resulting in significant spectral intensity above the Fermi level (highlighted by arrows in Fig. 3d), as can be also observed for the corresponding pristine film, although significantly more pronounced. This further supports the hypothesis that am-hydr-IrO<sub>x</sub> grown from the alloy films presumably contains little to no titanium, which leads to the question of whether the titanium is lost during the electrochemical growth.

In order to study this, X-ray absorption spectroscopy (XAS) at the Ti K-edge with a higher probing depth (0.5 to 1 μm in fluorescence yield<sup>32</sup>) was performed. All 1000-cycle samples show similar spectra, as depicted in Fig. S11a. The spectral fingerprint in these samples, characteristic of metallic Ti<sup>0</sup> mixed with that of titanium oxides in smaller concentrations, is also comparable to the spectra of the pristine IrTi alloy films. This confirms that titanium dissolution presumably does not explain the HAXPES-derived titanium-poor composition of the 1000-cycle grown am-hydr-IrO<sub>x</sub>. This follows chemical intuition as titanium is a very corrosion resistant material, due to the non-porous protective oxide coating formed in the presence of acids.<sup>33</sup> Based on the HAXPES and XAS data, a model illustrated in Fig. 3e is proposed, where am-hydr-IrO<sub>x</sub> grows from the alloy films and covers the entire surface but contains little to no titanium, supported on unreacted metallic alloy, leading to an am-hydr-IrO<sub>x</sub>/Ir<sub>x</sub>Ti<sub>1-x</sub> layer stack. Only for the thinnest film can the underlying alloy be detected by HAXPES, while, for all others, the spectroscopic signals observed originate solely from am-hydr-IrO<sub>x</sub>.

## 2.4 Electrochemical OER activity of the 1000-cycle am-hydr-IrO<sub>x</sub> grown from IrTi alloy films

The cyclic voltammograms corrected for voltage errors caused by solution resistance (*R*) and current flow (*i*) highlighting the electrochemical activity towards the oxygen evolution reaction are compared in Fig. 4a. The series resistances were determined by first growing the am-hydr-IrO<sub>x</sub> layer up to 1000 cycles, measuring the impedance spectrum and then extracting the high frequency response. The series resistance of the 1000-cycle am-hydr-IrO<sub>x</sub> layers is a function of the elemental composition of the alloy films as highlighted in Fig. 4b, whereby the series resistance decreases with higher iridium content in the initial alloy film. After *iR* correction, the similarity in activities between the different samples is somewhat counterintuitive, since thickness, *i.e.* the number of electrochemically active sites derived from the voltametric charge, varies with the metal composition of the film as shown in the inset in Fig. 4a. All evidence up to this point suggest that the catalytic activity of all 1000-cycle am-hydr-IrO<sub>x</sub> layers grown from IrTi alloy films

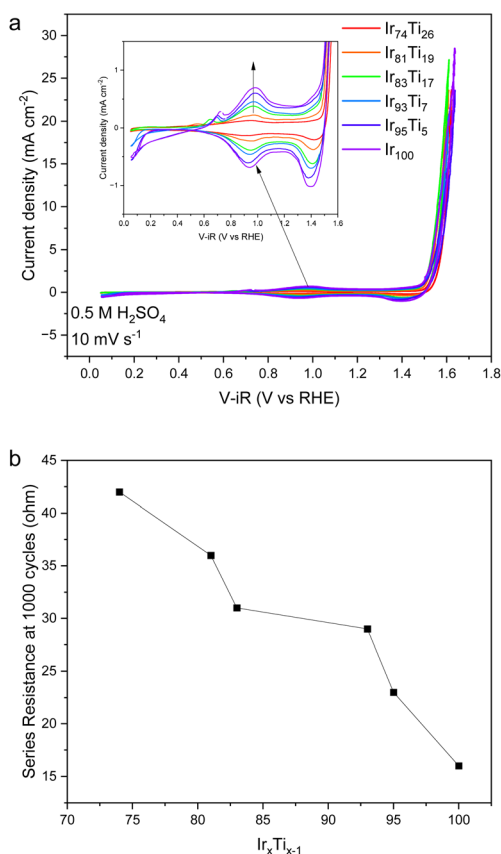


Fig. 4 Electrochemical activity of 1000-cycle am-hydr-IrO<sub>x</sub> layers grown from IrTi alloys with different Ir contents: (a) *iR* compensated cyclic voltammometry. The inset shows an enlarged portion of the cyclic voltammograms highlighting differences in the capacitance between samples. All cyclic voltammograms were measured at 10 mV s<sup>-1</sup> and the current is normalized by the geometric surface area of the sample. (b) Series resistance of the different 1000-cycle am-hydr-IrO<sub>x</sub> layers grown on IrTi alloys with different surface iridium contents. The series resistance was derived from the high frequency response in impedance measurements.



(containing mostly, if not exclusively, iridium as the metal) is very similar, differing only with respect to their thicknesses. In OER electrocatalysis, it is typically assumed that, for the same active site, activity should linearly correlate with the number of active sites (*e.g.*, their total quantity). To probe this relationship for the am-hydr-IrO<sub>x</sub> samples studied, the activity of corresponding layers grown from Ir<sub>100</sub> as a function of cycles was studied and is shown in Fig. S14. A direct correlation between the activity of the sample and its integrated charge (the proxy measurement for electrochemically active sites) should be observed; however, the results are significantly more complex. Fig. S14a shows the cyclic voltammograms of the am-hydr-IrO<sub>x</sub> layers grown on Ir<sub>100</sub> as a function of cycle number, with the inset highlighting the growth of the am-hydr-IrO<sub>x</sub> layer *via* the increase in (pseudo)capacitance of the redox features. The OER activity is measured by the current density at 1.7 V *vs.* RHE and is plotted against the integrated charge of the redox feature in Fig. S14b. From this, it can be derived that the relationship between the number of electrochemically active sites and activity is clearly not direct. Although we see an initial increase in activity between the first 200–400 cycles, this does not follow the steady increase in the redox integral. At high cycle numbers, we see a large drop in activity, coinciding with a small decrease in am-hydr-IrO<sub>x</sub> layer thickness. A detailed interpretation of this observation is developed further in the discussion section below. An alternative explanation for similar *iR* corrected electrochemical activities across the samples, despite the different numbers of exposed catalytic sites, could be due to subtle changes in the electronic structure of the active iridium sites caused by proximal titanium. The electrochemical signature of iridium shown in Fig. 4 and S9 does not appear to reflect any Ti content related changes in the electronic structure of iridium, but rather just shows the extent of am-hydr-IrO<sub>x</sub> growth (thickness). However, this possibility was studied further using *operando* X-ray absorption spectroscopy of the Ir L<sub>3</sub>-edge with details of the experimental procedure found in the SI. After being grown up to 1000 cycles, the am-hydr-IrO<sub>x</sub> layers were held at several voltages where key electrochemical events occur across the pre-catalytic voltage range and the absorption spectra were recorded, as shown in Fig. S15. It was observed that, within the error of the measurement, all samples that were measured exhibited similar white line position shifts as a function of voltage, supporting the conclusions drawn from the electrochemical experiments that electrochemical activity is dominated by the active iridium atoms within the sample and providing evidence that the active species in all samples is am-hydr-IrO<sub>x</sub>, revealed to grow with almost no Ti on top of the remaining unreacted parts of the alloy film.

## 2.5 Electrochemical stability

To study how the initial composition of the IrTi alloy influences the stability of the am-hydr-IrO<sub>x</sub> grown from it, different accelerated testing protocols were utilised to highlight the differences in sensitivity to the different degradation mechanisms between the alloy films. The electrocatalysts were subjected to two different potential regimes. The stability towards cycle-

induced delamination was evaluated by cycling 10 000 times using the typical growth parameters at a scan rate of 500 mV s<sup>-1</sup> with a slower scan of 50 mV s<sup>-1</sup> at increments of 1000 cycles to track changes in the voltametric response. This aims to mimic shut down/switch on protocols, ranging from very high to low potentials within seconds, while limiting potential-induced dissolution by minimising the time spent in a potential range where minimal OER occurs. In contrast, the stability towards potential-induced dissolution was studied potentiostatically at a voltage where OER readily occurs and minimal further growth of am-hydr-IrO<sub>x</sub> is expected. Fig. 5a and b show voltammograms of Ir<sub>100</sub> and Ir<sub>74</sub>Ti<sub>26</sub> films treated to 1000 and 10 000 cycles, respectively. Ir<sub>100</sub> exhibits a dramatic change resulting from this electrochemical procedure; the loss of the capacitive current is consistent with the loss of electrocatalytically active iridium oxide material since electrochemically active iridium contributes to the (pseudo)capacitive current. In addition, the change in voltametric response indicates a chemical change in the electroactive species. In contrast, the Ir<sub>74</sub>Ti<sub>26</sub> alloy film only shows subtle changes in the positions and shape of its electrochemical fingerprint, suggesting maturation of the film but not severe destruction, as seen previously for the pure iridium system. The chemical composition of the am-hydr-IrO<sub>x</sub> films after 10 000 cycles was further studied using XPS/HAXPES and compared to measurements of pristine alloy films and am-hydr-IrO<sub>x</sub> samples grown to 1000 cycles, as shown in Fig. 5c–f. In the pristine Ir<sub>74</sub>Ti<sub>26</sub> film, iridium is in its metallic state. After 1000 cycles, photoemission lines corresponding to oxidized and metallic iridium are observed (see Ir 4f spectra in Fig. 5c). These originate from the surface am-hydr-IrO<sub>x</sub> grown from the alloy film and the unreacted alloy underneath. At the same time, the Ti 2p region (Fig. 5e) shows both oxidized and metallic components in the pristine film. Deviation from expected binding energies was ascribed to localised charging of insulating TiO<sub>2</sub> species. After 1000 cycles, no major changes in the spectral shape of the Ti 2p contributions are detected. However, a loss in the signal to noise ratio is clearly observed which is ascribed to the Ti (constituent of the Ir<sub>74</sub>Ti<sub>26</sub> film) being buried due to the growth of the am-hydr-IrO<sub>x</sub> layer. After 10 000 cycles, titanium is revealed again, suggesting that am-hydr-IrO<sub>x</sub> is now thinner. Therefore, this allows measuring the titanium at the interface between the am-hydr-IrO<sub>x</sub>/Ir<sub>x</sub>Ti<sub>1-x</sub> stack and the underlying metallic alloy. Additionally, the titanium is further oxidized, implicitly indicating a reaction with the electrolyte.

In the pristine Ir<sub>100</sub> film, iridium is in its metallic state. After 1000 cycles, photoemission lines corresponding to oxidized iridium are the only ones observed. This is interpreted as the am-hydr-IrO<sub>x</sub> layer being significantly thicker in this sample and, therefore, no significant signal from the unreacted alloy film is detected. After 10 000 cycles the Ir 4f signal is similar to its pristine counterpart (Fig. 5d). This is interpreted as near-complete delamination of the catalytic layer, where the remaining non-oxidised iridium was electrically isolated due to loss of the conductive catalytic layer and did not participate in the growth. In addition, the strong Si 2p signal arising from the substrate onto which the film was sputtered reveals that the remaining film is very thin or does not cover the support completely (Fig. 5f). In



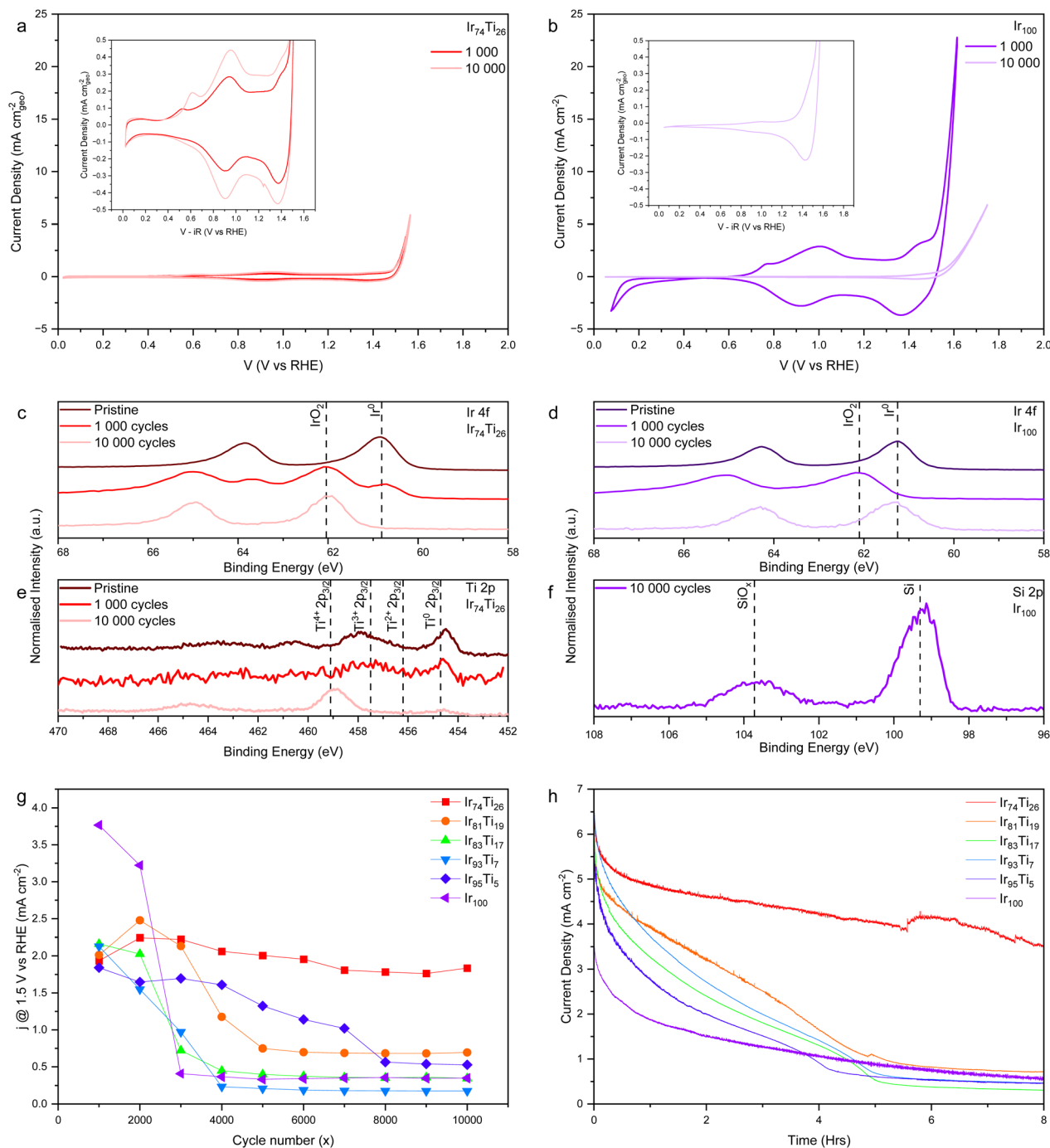
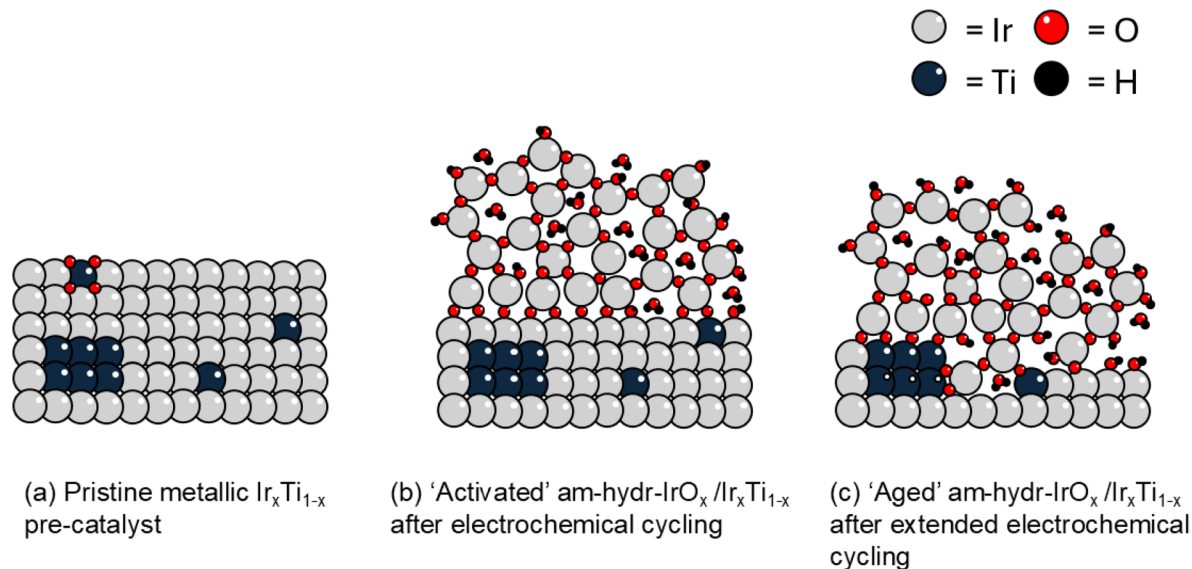


Fig. 5 Electrochemical stability of am-hydr-IrO<sub>x</sub> grown from Ir<sub>x</sub>Ti<sub>100-x</sub>: comparison of cyclic voltammograms measured after 1000 and 10 000 cycles performed on Ir<sub>74</sub>Ti<sub>26</sub> (a) and Ir<sub>100</sub> (b) films. Insets highlight changes in (pseudo)capacitance. (c) and (e) XPS/HAXPES spectra of Ir 4f and Ti 2p, respectively, comparing the pristine Ir<sub>74</sub>Ti<sub>26</sub> alloy film with the sample after 1000 and 10 000 cycles. (d) and (f) XPS/HAXPES spectra of Ir 4f and Si 2p, respectively, comparing the pristine Ir<sub>100</sub> film with the sample after 1000 and 10 000 cycles. The data of the pristine samples and after 10 000 cycles were acquired by XPS with an Al K<sub>α</sub> source, while the data after 1000 cycles were acquired by HAXPES using 2 keV excitation photon energy. (g) Stability towards cycle-induced delamination quantified by the current density, measured at 1.5 V vs. RHE, as a function of cycle number. (h) Potentiostatic stability measured through chronoamperometry at 1.7 V vs. RHE on samples grown to 1000 cycles.

contrast, the sample grown from Ir<sub>74</sub>Ti<sub>26</sub> shows no evidence of delamination (by means of Si peak identification as shown in Fig. S16). However, the elemental ratio of the iridium and titanium species has shifted to Ir<sub>58</sub>Ti<sub>42</sub> (calculated as before, using the Ir 4f and Ti 2p regions). This is consistent with the maturation

of the film through electrochemically driven corrosion into the electrolyte with no significant loss of electroactive area as indicated by a higher pseudo-capacitive response of the redox features. In addition, the Ti 2p spectrum of the 10 000-cycle sample in Fig. 5e shows a lower proportion of metallic titanium,





**Fig. 6** Proposed model for the structural evolution of the studied samples. (a) Pristine metallic alloy film with both isolated crystal domains of titanium and titanium within the iridium matrix. Surface titanium is also partially oxidised through exposure to air. (b) 'Activated' am-hydr- $\text{IrO}_x$  formed on the metallic substrate. (c) 'Aged' am-hydr- $\text{IrO}_x$ , extended cycling has developed the hydrous layer deeper into the metallic substrate allowing more titanium to be accessed spectroscopically due to loss of surface iridium and allowing further oxidation of metallic titanium.

attributed to a more oxidized state, as expected after the extensive electrochemical treatment with the loss of iridium exposing the underlying titanium to the electrolyte.

Fig. 5h shows the potentiostatic stability measured through chronoamperometry at 1.7 V *vs.* RHE after the accelerated degradation tests. Fig. 5g shows the relative rate of degradation of all samples by depicting the loss in current density at 1.5 V *vs.* RHE of a given cycle number. All the high iridium content samples ( $\text{Ir}_{100}$ – $\text{Ir}_{83}$ ) show rapid degradation, likely due to the delamination of the am-hydr- $\text{IrO}_x$  species during the test. However, samples with higher titanium content (*e.g.*  $\text{Ir}_7\text{Ti}_{26}$ ) exhibit significantly slower degradation, with no profound loss of activity (*e.g.* –95% for  $\text{Ir}_7\text{Ti}_{26}$  *vs.* –10% for  $\text{Ir}_{100}$ ).

Fig. 6 illustrates the evolution of the samples through the extended cycling procedure. After 1000 cycles, the pristine metallic pre-catalyst in Fig. 6a transforms into the am-hydr- $\text{IrO}_x / \text{Ir}_x\text{Ti}_{1-x}$  layer stack, in which the underlying alloy is only spectroscopically available for the thinnest am-hydr- $\text{IrO}_x$  layer grown from  $\text{Ir}_7\text{Ti}_{26}$ . After 10 000 cycles, the am-hydr- $\text{IrO}_x$  layer thins, due to electrochemically enhanced corrosion leaching iridium into the electrolyte, revealing titanium that has been partially oxidised due to exposure to the electrolyte.

### 3. Discussion

#### 3.1 Titanium oxide as a growth inhibitor for am-hydr- $\text{IrO}_x$

The complex non-homogenous chemical compositions of the IrTi alloy films were thoroughly characterised and the evolution of the electrochemically grown am-hydr- $\text{IrO}_x$  structure was studied up to layers corresponding to 1000 grow cycles. We show that the presence of the titanium within the pristine alloy is crucial for enhancing the stability (both with respect to Ir dissolution and delamination) of the electrocatalyst by

modifying the growth mechanism of am-hydr- $\text{IrO}_x$ , resulting in a drastically lower growth rate. Additionally, profound improvements in stability are achieved with minimal loss of intrinsic activity.

XPS analysis of the sputtered IrTi films (Fig. S6 and S7) revealed that with higher total titanium content within the film, a higher relative amount of titanium oxide is observed at the surface. Electrochemical cycling induced the formation of am-hydr- $\text{IrO}_x$  layers, whose growth rate was significantly decreased with higher initial Ti content, as evidenced by the pseudo-capacitive currents in Fig. 2.

Hard X-ray photoelectron spectroscopy (HAXPES) measurements (Fig. 3) provided critical insights into the unexpected, layered structure: at 1000 growth cycles, all samples show evidence of a stacked am-hydr- $\text{IrO}_x / \text{Ir}_x\text{Ti}_{1-x}$  layered structure, where at the surface, the electrochemically active am-hydr- $\text{IrO}_x$  interacts with the electrolyte and electrochemically evolves oxygen. This layer is supported by the remaining unreacted IrTi alloy. Evidence from HAXPES (Fig. 3) and *operando* XAS (Fig. S15) shows that am-hydr- $\text{IrO}_x$  grown from all IrTi alloy films differs only in thickness but not composition. Additionally, the amount of titanium in the pristine alloy correlates strongly with the impeded growth rate of am-hydr- $\text{IrO}_x$  (Fig. 2c) and pristine samples with higher titanium content have proportionately higher titanium oxide at the surface (Fig. S6 and S7). By combining these observations, it is evident that the titanium oxide at the sample surface hinders continued growth of am-hydr- $\text{IrO}_x$  by blocking underlying iridium from the electrolyte. After the initial 1000 growth cycles, sufficiently high concentrations of titanium oxide prevent further growth. This is in contrast to am-hydr- $\text{IrO}_x$  grown from pure iridium, which grows continuously until eventual delamination. Although the decreased growth rate of alloys with higher Ti content results in



fewer electrochemically active sites (as revealed by a lower voltametric charge, *i.e.* a thinner film), interestingly it does not negatively affect its OER activity as shown in Fig. 4, even with fewer participating sites. This is likely related to the unique structure of hydrous iridium oxide electrocatalysts. The authors propose that the non-linearity of the thickness–activity relationship could be explained by diffusion limitations of the electrolyte deep within the am-hydr-IrO<sub>x</sub> structure. While the innermost parts of am-hydr-IrO<sub>x</sub> are still electrically active and can still interact with the limited electrolyte that wets the hydrated structure, and hence exhibit measurable electrochemical redox, the local area can be partially depleted of water under operational conditions. This leads to slow H<sub>2</sub>O diffusion, increasing mass transfer resistance and, therefore, slow O<sub>2</sub> generation in these areas.<sup>34</sup> Further work is required to more comprehensively understand these observations but this is beyond the scope of this manuscript.

Now the different chemistries of iridium and titanium can be compared to rationalise this behaviour. As iridium is stable in multiple oxidation states, it can undergo oxidation and partial reduction required to form am-hydr-IrO<sub>x</sub>. Within the cycling potential regime between 0.05–1.5 V, TiO<sub>2</sub> is stable and does not undergo redox reactions.<sup>35</sup> Titanium particles, even on the nano-scale, are resistant to total oxidation due to the formation of a non-porous oxide layer limiting diffusion of electrolyte to underlying metallic titanium.<sup>33</sup> To summarise, during the growth, the oscillating voltage leads to the transformation of iridium to am-hydr-IrO<sub>x</sub> as well as to an opening of the structure, allowing electrolyte to penetrate and react with metallic iridium at later growth cycles. Titanium, in contrast, undergoes limited surface oxidation, hindered by the formation of the non-porous oxide-based passivation layer. It is possible that the presence of the titanium oxide surface layer has the effect of shielding underlying metallic iridium from participating in the growth, leading to a am-hydr-IrO<sub>x</sub> growth rate highly sensitive to the titanium concentration.

### 3.2 Further roles of titanium in stability enhancement

The stabilisation effect of titanium towards iridium dissolution (Fig. 5h) is consistent with previous reports.<sup>24,28,36,37</sup> This raises the question of whether it plays a similar role in the materials studied here or whether all stability improvements can be ascribed to differences in thicknesses of the am-hydr-IrO<sub>x</sub> layer. Thickness is a key factor contributing to the stability of the am-hydr-IrO<sub>x</sub> layer with respect to electrochemical cycling; however alternative factors may also contribute to the potentiostatic stability. Fig. 5h shows that there is a clear critical amount of titanium required in the alloy to achieve high stability in the resultant active am-hydr-IrO<sub>x</sub> layer. This suggests that titanium may influence the stability of iridium with respect to dissolution through a different mechanism, although there is no direct evidence for the presence of titanium within the am-hydr-IrO<sub>x</sub>. In the cases in which titanium could be detected, the am-hydr-IrO<sub>x</sub> is thin enough that it cannot be excluded that the signal corresponds to the metallic alloy film. However, a significant stability increase with titanium content as low as 1.92% has been reported.<sup>38</sup> One possible explanation could be that a small and, therefore, undetected amount of titanium

within the am-hydr-IrO<sub>x</sub> reduces the iridium dissolution rate. Alternatively, since iridium is separated from the alloy when the am-hydr-IrO<sub>x</sub> forms, another possible explanation would be the presence of a titanium-rich interlayer at the interface between the am-hydr-IrO<sub>x</sub>/Ir<sub>x</sub>Ti<sub>1-x</sub> stack. Although there is no direct experimental evidence to support this, it is supported indirectly by the observation of an increase in series resistance with respect to the initial titanium content for films grown up to 1000 cycles (Fig. 4b). Since iridium and its oxides are more electrically conductive, an increasing series resistance could be explained by titanium oxides at this interface.

To further study this, the XPS of the Ti 2p region of am-hydr-IrO<sub>x</sub> grown on Ir<sub>74</sub>Ti<sub>26</sub> cycled 10 000 times was deconvoluted, as shown in Fig. S17, revealing that there is still a significant amount of metallic titanium as well as suboxides of titanium which are still conductive. These results can help to explain the continued electrochemical activity with some loss of activity due to an increased resistance but without total electrical isolation of the electrochemically active layer with respect to the current collector. An analogue observation was found for co-sputtered Ir–Ti films, where Ti suboxides in the interfaces favoured the stability of the electrocatalysts without hindering its activity.<sup>22</sup>

With this information about the surface oxidation state of titanium, the bulk structure of the samples that had undergone 1000 electrochemical cycles was also probed by Ti K-edge XAS measurements, as well as a pristine alloy sample (Ir<sub>74</sub>Ti<sub>26</sub>) for comparison. A linear combination fitting approach of the Ti K-edge XAS spectra was performed using references of metallic titanium as well as several common titanium oxides (TiO, Ti<sub>2</sub>O<sub>3</sub>, Ti<sub>4</sub>O<sub>7</sub> and TiO<sub>2</sub>) as potential candidate structures for titanium within the sample. Attempts to use these structures failed to produce acceptable fits to the measured data, whereby key features of the spectra could not be reproduced as shown in Fig. S11. Additionally, all fits give large contributions of Ti<sub>4</sub>O<sub>7</sub> which does not follow chemical intuition as the samples are prepared under ultra-high vacuum (UHV) conditions and have high metallic contributions at the surface. It has been shown elsewhere that when titanium alloys with other metals, charge-transfer processes can occur between the metals, leading to shifts in the observed spectra.<sup>39</sup> Indeed, this has already been observed for the Ir 4f XPS data of the pristine samples in Fig. 1b, in which a shift towards lower binding energies of Ir 4f is observed. This implies that electron density is moving from titanium to iridium, which corresponds well with the increase in white line position resulting from a decreased electron density around titanium. The fitting procedure was repeated with Ir<sub>91</sub>Ti<sub>9</sub> as a reference (see Fig. S12), since in this sample the titanium is too diluted to form a significant number of isolated structures within the alloy and, therefore, is representative of the IrTi solid solution. The results show that with more titanium in the structure, increasing contributions of Ti and TiO<sub>2</sub> are observed within the fit, which agrees well with previous characterisation. To summarise, after 1000 cycles the unreacted underlying alloy film layer dominates the signal and the electronic state of the layers is best described as a metallic alloy between iridium and titanium, with increasing amounts of isolated titanium at high titanium content.



## 4. Conclusion

This study provides detailed insights into a strategy for improving the stability of am-hydr-IrO<sub>x</sub> electrocatalysts without the typical accompanying loss of activity leading to a promising PEM anode material exhibiting both high activity and stability. By utilizing both lab-based and synchrotron-based spectroscopic techniques, snapshots of the evolving electrocatalyst structure were obtained. Most importantly, it has been shown that the presence of Ti in the initial Ir matrix hinders the growth of am-hydr-IrO<sub>x</sub>. This fact can be used to tune the stability of these film materials by controlling the amount of metallic iridium in the underlying film consumed during the electrochemical growth. The mechanical stability of these films highly depends on the presence of some remaining Ir to anchor the electrocatalyst. Within the pristine samples with higher surface titanium content, there is clear evidence of a higher relative amount of titanium oxide. The am-hydr-IrO<sub>x</sub> growth dynamics are highly sensitive to the amount of titanium oxide and above a critical concentration, a profound improvement in electrochemical stability is observed.

## Conflicts of interest

There are no conflicts to declare.

## Data availability

The raw data utilised in this work have been deposited online in a Zenodo repository that has the following permanent DOI: <https://doi.org/10.5281/zenodo.18164742>.

Supplementary information (SI) is available. See DOI: <https://doi.org/10.1039/d6ta03607d>.

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