Towards maximized utilization of iridium for the acidic oxygen evolution reaction

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ABSTRACT

The reduction in noble metal content for efficient oxygen evolution catalysis is a crucial aspect towards the large scale commercialization of polymer electrolyte membrane electrolyzers. Since catalytic stability and activity are inversely related, long service lifetime still demands large amounts of low-abundant and expensive iridium. In this manuscript we elaborate on the concept of maximizing the utilization of iridium for the oxygen evolution reaction. By combining different thin oxide based support materials with liquid atomic layer deposition of iridium oxide, new possibilities are opened up to grow thin layers of iridium oxide with tunable noble metal amounts. In-situ time- and potential-resolved dissolution experiments reveal how the stability of the substrate and the catalyst layer thickness directly affect the activity and stability of deposited iridium oxide. Based on our results, we elaborate on strategies how to obtain stable and active catalysts with maximized iridium utilisation for the oxygen evolution reaction and demonstrate how the activity and durability can be tailored correspondingly. Our results highlight the potential of utilizing thin noble metal films with earth abundant support materials for future catalytic applications in the energy sector.

KEYWORDS

oxygen evolution reaction, liquid atomic layer deposition, catalysis, iridium, core-shell nanoparticles

1 Introduction

Ruthenium oxide (RuO₂) and iridium oxide (IrO₂) are the most promising catalyst materials for the acidic oxygen evolution reaction (OER), since they combine high intrinsic OER activities with high stability. While RuO₂ is more active, it dissolves to a great extend during OER [1]. IrO₂, in contrast, is typically more stable with slightly higher overpotentials, making it the material of choice in current electrolyzers [2–4]. Different types of Ir-based catalysts exist, ranging from homogeneous molecular catalysts to heterogeneous nanoparticulate, film or bulk catalysts. Heterogeneous iridium catalysts comprise different types of amorphous, hydrous oxide materials to crystalline oxides, perovskites [5, 6], pyrochlores [7, 8] or hollandites [9]. Of all named materials, crystalline IrO₂-oxal unit demonstrates the highest stability and the lowest iridium-dissolution rates due to the formation of a dense, crystalline film, with low electrolyte accessibility to sub-surface iridium oxide and corner shared, so-called “activated oxygen atoms” [3]. The compact catalyst layer results in low overall mass activity since only the outer surface at the solid–liquid interface takes part in the reaction—The measured mass activity is therefore highly dependent on the particle dimensions.

Despite the excellent OER performance of iridium (oxides), their low abundancy, the often-arduous mining conditions and high price remain the major drawbacks for large scale applications of polymer electrolyte membrane electrolyzers. A general trend in electrochemical energy conversion and in the whole field of catalytic applications is the reduction or even the full replacement of noble metals with abundant materials [10–12]. Given the fact that only the outer surface is participating in the catalytic reactions, the bulk could be replaced by less expensive materials—for nanoparticles the principle is known as core-shell concept. The main advantages can be summarized in three points: (i) The noble metal content can be substantially decreased since nearly all active atoms are at the surface and take place in the catalytic reaction; (ii) the underlying substrate could potentially affect the specific activity of the active material positively [13–16]; (iii) a stronger adhesion energy between core and shell material can increase the stability of the active surface elements [17–19]. Exemplarily, for spherical catalyst particles, the electrochemically active surface area (ECSA in m²·g⁻¹) and its dependency on core diameters and shell thicknesses is displayed in Fig. 1. When the noble metal shell is sufficiently thin, the active surface area drops only marginally despite the overall increase in particle size. When keeping the core diameter constant and increasing the shell thickness, however, the ECSA drops drastically, cancelling out the advantage of reduced shell dimensions.

A thin film of IrO₂-oxal in the monolayer range on a cost-efficient support material would secure a high overall mass activity due to...
the high fraction of surface sites that can be utilized. Additionally, certain support materials, such as fluorine doped tin oxide (FTO) [20], antimony doped tin oxide (ATO) [21–23], indium tin oxide (ITO) [24], or Nb-doped TiO₂ [13, 25–27] were reported to positively influence activity and stability of iridium-based catalysts.

In this manuscript, we concentrate on conductive tin oxide support materials and highlight the prerequisites, challenges and limits for OER catalysts with low iridium loading regarding stability and activity. Tin was used as support material due to its relative ease to obtain the necessary conductivity, its reported positive influence on the catalytic activity/stability and its over 10⁴ higher global primary production compared to iridium (data from 2010) [28]. We show that the further developed liquid atomic layer deposition (l-ALD) approach results in low amounts of iridium oxide that can be adapted depending on the amount of cycles applied. The feasibility of the proposed synthesis is shown exemplarily on model thin films allowing for facilitated in- and ex-situ characterisation. We demonstrate that the impact of substrate material, the degree of oxide formation, and the amount of catalyst is crucial for the described design principle which opens up new perspectives for the successful preparation of high-performance catalysts with maximized utilisation of precious iridium.

2 Results and discussion

2.1 Self-adhered iridium complexes

Anchored metal-complexes bridge the gap between heterogeneous and homogeneous catalysis with potential tunability of active sites. The concept seems appealing since the catalyst may be tailored to a specific reaction depending on the used ligands, while still maintaining its molecular identity during catalysis [29–35]. Therefore, we set out with molecular iridium complexes chemisorbed on oxide surfaces (Ir_complex) and tested their performance during OER as an all-surface catalyst [30, 32, 36]. The molecular structure of the used complex can be found in Fig. 2. By exchanging a bound water ligand with surface oxygen functionalities, the Ir_complex Self-adheres to the surface of metal oxides at room temperature. Applying the same synthesis conditions, a (sub)monolayer of the Ir_complex was attached to an FTO and ITO coated glass slide (Fig. 2(a)). A detailed X-ray photoelectron spectroscopy analysis of the adhered complex can be found in the Electronic Supplementary Material (ESM). The as prepared catalyst was tested towards the OER in 0.1 M HClO₄ as shown in Fig. S1(a) in the ESM. Normalized to geometric surface area, the first scan revealed rather low activity in comparison to pristine IrO₂-rutile. However, activation via consecutive potential cycling significantly increased the activity, which originated from the formation of an amorphous IrO_x layer and the loss of organic ligands from the surface, as it was observed before for other electrocatalysts at high potentials [37, 38]. This is in line with cyclic voltammetry curves displayed in Fig. S3 in the ESM which show the characteristic amorphous IrO_x pattern [39]. As stability metric for the OER, the stability number (S-number) was used in order to compare the stability to other OER-catalysts [3]. The S-number describes the number of dissolved metal atoms per one oxygen molecule formed and is independent of surface area, catalyst loading, and the number of active sites. The S-number of Ir_complex on FTO is relatively low and only ~ 150 oxygen atoms are produced per one iridium atom dissolved into the electrolyte (Fig. S1 in the ESM). Note that commonly used iridium-based materials possess S-numbers in the range of 10⁴ to 10⁶ (c.f. Fig. 4) [3].

2.2 Liquid atomic layer deposition

Since the heterogeneously attached iridium-complex fulfilled the prerequisites towards a maximized utilisation of iridium, but did not provide the necessary stability for long-term application, we extended the existing idea to a liquid atomic layer deposition approach [40]. One l-ALD cycle consists of the before described complex adsorption onto oxide surfaces (ITO, FTO) followed by thermal treatment in air. Reiteration of the self-limiting process leads to an adjustment of the iridium oxide loading which depends on the number of cycles. Due to the encountered difficulties in stability of the chemisorbed iridium complex, thermal oxidation of iridium was applied to aim at the formation of stable IrO₂-rutile. Previous studies of iridium oxide on Ti substrates showed the highest OER

![Figure 1](https://example.com/figure1.png)

**Figure 1** Maximizing the utilisation of noble metals in nanoparticles. Dependency of ECSA on (a) varying core diameters with a constant Ir-shell thickness (0.5 nm) and (b) varying shell thicknesses with constant core diameter (1 nm). The calculations result from geometric considerations—The details can be found in the ESM.

![Figure 2](https://example.com/figure2.png)

**Figure 2** Schematic display of liquid atomic layer deposition of iridium oxide on oxide-based substrates (red: oxygen; blue: iridium; yellow: oxide substrate). (a) Consecutive immersion (overnight) of the iridium complex and subsequent annealing at 500 °C leads to the growth of iridium oxide on FTO. (b) STEM images of FTO covered with IrO₂ after 20 cycles at 500 °C. The TEM sample was prepared by focused ion beam (FIB) sectioning. (c) X-ray photoelectron spectra of Sn 3d and Ir 4f levels of the complex adsorbed on the oxide surface (FTO) and after annealing at 500 °C. A detailed analysis can be found in the ESM.
activity after heat treatment between 250 and 350 °C [41]. The low temperatures, however, result in a high level of edge and defects sites with low overall dissolution stability [3, 42]. The formation and crystallization of stable IrO$_2$-rutile was reported to take place between 400 and 500 °C, depending on the used precursors or substrates, which decreases the Ir-dissolution drastically [43, 44]. Therefore, each applied layer was treated for 30 min in air at 500 °C to form oxidized iridium species. For selected samples, the temperature of the last annealing step was increased to 600 °C for 5 h to maximize the transformation from (hydrous) IrO$_x$ to IrO$_2$-rutile [41, 43, 45]. Note that the described Ir$_{\text{complex}}$ contains organic ligands, so that chloride contamination reported for IrCl$_4$ precursors could be excluded [44, 46]. Similar to classical ALD, several cycles are necessary to form one catalytic layer. Here, up to 20 cycles were applied. The iridium loading after one cycle was determined by inductively coupled plasma mass spectrometry (ICP-MS) to be ~ 30 ng·cm$^{-2}$.

In order to inspect the location of Ir with respect to Sn, a lamella of IrO$_x$ on FTO (20 cycles) was prepared by means of focused ion beam-scanning electron microscopy (FIB-SEM, FEI Helios NanoLab 600, Fig S2 in the ESM). The respective scanning transmission electron microscopy (STEM) images and elemental distribution maps are displayed in Fig. 2(b) demonstrating the general feasibility of the synthesis. Depending on the selected temperature, the formation of iridium-rich areas and the formation of an inhomogeneous film is observed (c.f. stability discussion). Especially 600 °C favors the reorganization of IrO$_x$ as shown in Fig. S6 in the ESM. The increase in temperature results in higher Ir/Sn mobility and mixed oxide formation that goes along with surface restructuring and poor homogeneity. Narrow scans of the Ir 4f and Sn 3d core levels determined by X-ray photoelectron spectroscopy of the Ir$_{\text{complex}}$ chemisorbed on the FTO surface before and after thermal treatment (500 °C) are shown in Fig. 2. The shift of the Ir 4f$_{5/2}$ peak from 61.4 eV to 62.1 eV reveals the increased formation of IrO$_x$ [47, 48]. Simultaneously, a shift of the 3d$_{5/2}$ Sn core levels from 485.8 to 486.7 eV indicates the formation of higher (mixed) oxides. The reason lies in the incorporation of Ir(IV) species into the Sn oxide host lattice at this temperature, which decreases the Ir-dissolution drastically [43, 44]. Therefore, each applied layer was treated for 30 min in air at 500 °C to form oxidized iridium species. For selected samples, the temperature of the last annealing step was increased to 600 °C for 5 h to maximize the transformation from (hydrous) IrO$_x$ to IrO$_2$-rutile [41, 43, 45]. Note that the described Ir$_{\text{complex}}$ contains organic ligands, so that chloride contamination reported for IrCl$_4$ precursors could be excluded [44, 46]. Similar to classical ALD, several cycles are necessary to form one catalytic layer. Here, up to 20 cycles were applied. The iridium loading after one cycle was determined by inductively coupled plasma mass spectrometry (ICP-MS) to be ~ 30 ng·cm$^{-2}$.

**2.3 Electrocatalytic activity towards the OER**

The polarisation curves after different adsorption cycles of Ir$_{\text{complex}}$ on FTO and ITO substrates at different temperatures are displayed in Fig. 3(a). The study of various samples of IrO$_x$ on FTO (1, 5, 15, and 20 cycles) reveals that the OER activity scales with the amount of iridium deposited. IrO$_x$ on FTO obtained by 20 consecutive cycles results in high utilization of a stable iridium oxide structure, close to the ideal and theoretical value of mass normalized activity for one monolayer of crystalline IrO$_2$-rutile. In the latter case, the participation of one monolayer was assumed corresponding to 150 ng·cm$^{-2}$ for the 001 facet of IrO$_{2\text{rutile}}$.

ITO was used as support material for comparison. Here, the same amount of cycles (here 5 cycles) result in much higher OER activities which can be explained by several consecutive effects. As stated before, iridium, tin, and indium oxides are susceptible to the formation of mixed oxides at high temperatures (e.g. Ir$_x$Sn$_{1-x}$O$_2$ and Ir$_x$In$_{1-x}$O$_2$ mixed phases) [1, 39]. For ITO, In and Sn dissolve significantly when brought in contact with low pH solutions. The leached In and Sn from the formed Ir$_x$Sn$_{1-x}$O$_2$ and Ir$_x$In$_{1-x}$O$_2$ mixed phases increase the overall surface roughness and result in an “amorphous-like” IrO$_2$ layer with a low amount of interconnected Ir-OIr networks and enhanced edge effects. A similar effect was observed for IrNi-based materials when the non-noble metal leaches during catalysis. The created lattice vacancies generate an unusually large number of d-band holes in the iridium oxide material which results in the observed low kinetic barrier during catalysis [53]. Similarly, the presence of vacancies and defects in cobalt/iron based layered double hydroxides was found to result in higher OER activity in alkaline medium [54]. The role of defects on the OER performance is discussed in detail in Ref. [55]. This observation is also expressed by the varying Tafel slopes obtained for IrO$_x$ on FTO and ITO as discussed in detail in the ESM. While overall this leads to an initial activation of ITO-based samples, long-term performance most likely suffers from the amorphization induced by the leaching process. Figure 3 compares the activity of various catalysts when normalized to the charge associated to the transition from Ir(III) to Ir(IV) and Sn(IV), similar radii, comparable electronegativities, and interfacial energies between IrO$_2$-rutile and SnO$_2$ favour the formation of mixed oxides [49–52]. In the following paragraphs, we will discuss the accompanying consequences of this dependency for (1) the activities and (2) the durability of the prepared catalysts.

**Figure 3** (a) Linear sweep voltammetry (ir corrected) in the OER region recorded for different IrO$_x$ loadings and substrates in 0.1M HClO$_4$ prepared at 500 °C. The displayed numbers represent the applied repetitive deposition cycles on the FTO support. For comparison, a sample with 5 cycles grown on ITO is shown (5$_{\text{ITO}}$). Inset: mass normalized current at 1.65 V$_{\text{RHE}}$. An iridium deposition of 30 ng for each cycle was used for normalisation. *For crystalline IrO$_x$, the participation of one monolayer was assumed corresponding to 150 ng·cm$^{-2}$. (b) Cyclic voltammetry between 0.05 and 1.4 V vs. reversible hydrogen electrode (RHE) with a scan rate of 0.2 V·s$^{-1}$. The displayed shaded area was used to determine the charge associated with the Ir(III) to Ir(IV) transition. (c) Comparison of different catalysts after normalization to the obtained charge from cyclic voltammetry indicated in (b). Ba$_3$PrIrO$_6$ was added as reference catalysts with one of the highest activities reported in Refs. [3, 5, 6].
Ir(IV) (as shown in Fig. 3). The ITO-based catalyst is as active as Ba₅PrIrO₉, a perovskite material with one of the highest reported activities [3, 5, 6]. IrOₓ on FTO possesses a lower specific activity compared to IrO₂ on ITO. The activity is still higher compared to pristine IrO₂, presumably caused by the lower calcination temperature of 500 °C. When annealed to 600 °C, the specific activities of IrOₓ on FTO resemble the specific activity of crystalline IrO₂-rutile (Fig. S4 in the ESM). The determination of the electrochemically active surface area and specific activity, however, still remains challenging for iridium-based materials. In contrast, iridium mass-normalized activities enable a practical comparison between various catalysts as shown in Fig. 1, inset. The loaded iridium content was determined through inductively coupled plasma mass spectrometry (ICP-MS) measurements and compared with the active charge capacity of the Ir³⁺/Ir⁴⁺ transition (c.f. ESM). The hydrous iridium oxide formed on ITO results in a high mass-based activity of ~ 35 A·mg⁻¹ at 1.65 V vs. RHE (5 cycles). This “all surface catalyst” contains low amounts of deposited metal, overall high accessibility to active sites and belongs to the most active catalyst reported in Ref. [6]. For FTO, the highest mass activity was observed for 20 repetitive cycles, resulting in a mass-based activity of ~ 2.5 A·mg⁻¹ at 1.65 V vs. RHE. The specific and mass activities are therefore highly susceptible to the support material employed. In order to shed more light on the ongoing processes during electrocatalysis, in-situ dissolution studies were performed.

2.4 Electrocatalytic stability

The observed activity was correlated with its intrinsic stability and galvanostatic polarization at 1 mA·cm⁻² of all materials was performed. Over the used time interval, the respective elemental dissolution was measured by means of a flow cell (FC) coupled on-line to an ICP-MS until quasi steady state conditions were reached (Fig. 4(a)). A detailed description of the used technique can be found in the ESM and in Refs. [43, 56]. The S-number for each material is estimated from the integrated amount of dissolved metal under quasi steady state conditions and the amount of produced oxygen assuming 100% faradaic efficiency (see Fig. 4(b)). The utilization of different substrates results in very different stability behaviour. Highly active IrO₂ on ITO dissolves to a high extent, while IrO₂ on FTO is significantly more stable, caused by the different extent of amorphous (on ITO) and crystalline character (on FTO) of the respective IrO₂ phases (c.f. amorphous IrO₂ as reference catalyst, Fig. 4(b)). While the choice of stable support material leads to overall higher stability of the catalyst/support system, the temperature influence on the formation of mixed oxides poses substantial challenges to the use of IrO₂/SnO₂-based materials. Indirect corroboration of the previously described mixed oxide formation was found when the dissolution and electrochemical behaviour of pristine FTO was compared to FTO where 15 consecutive cycles of IrO₂ deposition were applied (Fig. S7 in the ESM). For pristine FTO, no dissolution of Sn was observed even up to 2 V RHE while Sn dissolution was observed for 15 cycles that matches the dissolution profile of Ir.

Additional annealing at 600 °C for 5 h increases the stability of IrO₂ on FTO but not to the level of IrO₂-rutile. This indicates that the formation of Ir₅SrSnO₆ mixed oxides has negative effects on the stability of the whole catalytic system. Additionally, annealing at 600 °C has negative consequences for both, the specific and the mass activity (Fig. S4 in the ESM). The predicament lies therefore in the formation of a dense, crystalline IrO₂-rutile film with a high amount of interconnected Ir-O-Ir units during thermal annealing, conditions that, however, also favor increased mobility of surface atoms and the formation of less stable mixed oxides compared to IrO₂-rutile. If the observed dependency can be broken, a mass-based activity close to the most active iridium-based materials can be obtained while keeping the stability in the range of IrO₂-rutile. Our results suggest that an active and stable “all surface” IrO₂-rutile catalyst on an oxide support material has to fulfill the prerequisites of low intermixing and a high number of Ir-O-Ir networks that anchors Ir in place while keeping the total IrO₂-rutile thickness in the monolayer regime.

3 Conclusion

We have shown that post-grafted Ir-organo-complexes are not stable in the harsh environment that the acidic oxygen evolution reaction demands. A further developed synthesis method is proposed that consists of a liquid atomic layer deposition approach. Here, an Ir-organo-complex is attached to an oxide surface and subsequently oxidized. Depending on the amount of repetitive cycles, different catalyst loadings can be obtained. We demonstrate that the stability of the support material has drastic consequences for the activity and stability of the catalyst material. While one of the highest reported specific and mass-based activities were obtained on ITO, the utilization of stabilized surface IrO₂-rutile is at its maximum for FTO as substrate. The higher surface energy of IrO₂-rutile compared to SnO₂-rutile and the driving force for intermixing at high temperatures poses substantial challenges to the synthesis of stable and active OER catalysts based on the combination of iridium- and tin oxide based materials. By breaking the dependency of forming of a dense, crystalline IrO₂-rutile film with a high amount of interconnected Ir-O-Ir units and increased mobility of surface atoms at high temperatures, it is suggested that a mass specific activity close to the most active iridium oxide-based materials can
be obtained, while keeping the dissolution stability high and reducing the overall costs.

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