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Operando XAS Study of the Surface Oxidation State on a Monolayer IrO_x on RuO_x and Ru Oxide Based Nanoparticles for Oxygen Evolution in Acidic Media

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Abstract

Herein we present surface sensitive *operando* XAS L-edge measurements on IrO_x/RuO₂ thin films as well as mass-selected RuO_x and Ru nanoparticles. We observed shifts of the white line XAS peak towards higher energies with applied electrochemical potential. Apart from the case of the metallic Ru nanoparticles, the observed potential dependencies were purely core-level shifts caused by a change in oxidation, which indicates no structural changes. These findings can be explained by different binding energies of oxygenated species on the surface of IrO_x and RuO_x. Simulated XAS spectra show that the average Ir oxidation state change is strongly affected by the coverage of atomic O. The observed shifts in oxidation state suggests that the surface has a high coverage of O at potentials just below the potential where oxygen evolution is exergonic in free energy. This observation is consistent with the notion that the metal-oxygen bond is stronger than ideal.

Introduction

Our current dependence on fossil fuels is being challenged, and many countries are switching to an increasing amount of renewable energy generation due to a combination of economic and political incentives¹. However, most of the new energy sources are producing electricity intermittently, such as wind turbines and solar panels. Today the intermittent power supply is offset by existing fossil fuel power plants, but as the wind- and solar-power fractions increase further this is no longer viable, and new approaches to storing the produced electrical energy are needed. One possibility of storing the energy is in the form of chemical bonds, in which energy conversion devices convert excess electricity into easily storable chemicals, and convert them back to electricity on demand².

For such energy storage technologies to be viable, it is necessary to develop efficient devices that convert electric energy into chemical bonds. This can be in principle obtained in electrochemical cells, but for most relevant electrochemical reactions we do not have catalysts that are efficient enough to avoid large energy-losses. One particularly important chemical reaction is the oxygen evolution reaction (OER: $2H_2O \rightarrow 4H^+ + 4e^- + O_2$), which is one of the half-cell reactions in water splitting. Most synthetic fuels require hydrogen: for instance, dihydrogen is obtained from the hydrogen evolution reaction (HER: $2H^+ + 2e^- \rightarrow H_2$)³, whereas CO₂ can be reduced to hydrocarbons (e.g. $CO_2 + 8H^+ + 8e^- \rightarrow CH_4 + 2H_2O)^4$. However, in both these cases the source of hydrogen is water splitting, and so the OER must be an integral part of the device. Proton exchange membrane (PEM)⁵ electrolyzers, which use acidic electrolyte, have several benefits over traditional alkaline electrolyzers, such as higher current densities and better safety^{6,7}. It is a challenge to find materials that are both active for OER and stable in acidic conditions at the highly oxidative potentials required for water splitting. Most catalysts today are based on the precious metals Ir and Ru^{8–14}. In terms of scaling up energy conversion on a global scale this presents a significant problem, as these metals are extremely scarce¹⁵. RuO_x and IrO_x have shown promising activity for the OER^{10,12,16–24}, albeit most forms of RuO_x present poor stability^{11,22,25}. The sample preparation conditions have a great impact on the activity and stability of RuO_x based OER catalysts²⁶, so even the performance of a single material (RuO_x) can vary greatly^{25,27,28}.

Rossmeisl, Nørskov and coworkers developed a theoretical model to elucidate trends in oxygen evolution activity, using a density functional theory (DFT) based model.^{29–31} They proposed the following mechanism, based on four coupled electron-proton transfers:

* $+H_2O \rightarrow *OH + e^- + H^+$ * $OH \rightarrow *O + e^- + H^+$ * $O + H_2O \rightarrow *OOH + e^- + H^+$ * $OOH \rightarrow *+O_2 + e^- + H^+$

where * represents an active catalytic site. Using the free energies of each step calculated in³⁰, a free energy diagram can be constructed for the entire reaction on a $RuO_2(110)$ surface at different potentials, as seen in Figure 1.



Figure 1. A free energy diagram of the OER mechanism described in the text on a $RuO_2(110)$ surface, based on free energy DFT calculations in reference³⁰. For these calculations, the thermodynamic OER onset is at 1.60 V, the point at which all steps become downhill in energy. The free energy diagram has been calculated at the equilibrium potential 1.23 V, just below the OER onset at 1.50 V, and just above the OER onset at 1.70 V. All potentials are given vs. the reversible hydrogen electrode (RHE).

The ideal catalyst would have a completely flat free energy profile at the equilibrium potential, 1.23 V. However, it turns out that there is a constant energy difference between the binding energies of the two intermediates, *OH and *OOH. This means that even on the best catalyst, there is significant corrugation at the equilibrium potential; overcoming this corrugation requires an overpotential equal in magnitude to the most uphill step. On the most optimal catalyst (as opposed to the aforementioned ideal catalyst), the free energy change between O* formation and OOH* formation would be equal. On a surface which binds *O too weakly, relative to *OH, the overpotential will be due to *O formation. Conversely, a surface which binds *O too strongly, relative to *OH (and by extension *OOH), the overpotential will be due to the *OOH formation. According to Figure 1, the latter is the case for RuO₂(110), where *OOH formation is uphill by 0.37 eV at 1.23 V, while on $IrO_2(110)$ this step is 0.65 eV uphill³¹. These trends result in a Sabatier volcano relationship³¹.

In the current paper, our goal is to probe the proposed reaction mechanism and binding of the intermediates by performing *operando* x-ray measurements. L-edge spectroscopy of the metal atoms using hard X-rays is a bulk sensitive method; even though earlier studies of oxygen reduction on Pt showed that high spectral sensitivity from the interface can be obtained using monolayer catalysts³² and small nanoparticles³³. Herein, we have used a similar approach to investigate the L-edge of Ir and Ru during OER by depositing monolayer amounts of IrO_x on RuO₂ and small nanoparticles of Ru metal and RuO₂. We compared the measured Ir XAS spectra to that of simulated spectra to further connect the reaction mechanism to our measurements. For the simulations, we have chosen the four intermediates mentioned in the proposed reaction mechanism above (empty site, *OH, *O, and *OOH), as well as an *OO-based intermediate. The latter has been predicted theoretically³⁴ and observed experimentally³⁵.

For applications in PEM electrolyzers, nanoparticulate Ir or Ru-based catalysts are often preferred, as they provide a higher surface area per mass of catalyst used. Thus, understanding the catalytic behavior of Ru and RuO₂ in their nanoparticulate form is critical. Therefore, we investigated massselected metallic RuO_x/Ru and thermally oxidized RuO₂ nanoparticles produced by magnetron sputtering and gas aggregation³⁶, which we reported previously¹⁰; this production method ensures

good control of the size and morphology of the catalyst material. As nanoparticles have a high surface-to-bulk ratio they also present a system well suited for studying surface phenomena using X-ray absorption spectroscopy (XAS). Likewise, the ultra-thin IrO_x films³⁷ gave uniquely surface sensitive XAS measurements. To properly understand the reaction mechanism, we investigated all samples under *operando* conditions, having conditions as close as possible to real OER conditions. Catalytic reactions typically take place at the surface; hence their elucidation requires surface sensitive probes. Moreover, as emphasized by Miquel Salmeron and co-workers, measurements under reaction conditions are essential to fully understand surface chemical reactions^{38–40}.

Previously OER has been investigated using *operando* XAS and XPS on IrO_x nanoparticles^{41,42}, as well as on pure Ir or Ru films¹¹ and mixed Ir/Ru compound thin films⁴³. Here we show that the L-edge XAS white line, corresponding to p to d excitations of both Ir and Ru, are sensitive to the applied potential with a shift towards higher photon energies approaching OER conditions. This is most noticeable for the sub-monolayer of IrO_x that gives a maximum surface sensitivity since all Ir atoms are at the interface. The measurements presented here have a higher energy resolution than previously reported XAS spectra, as we are using the Ru L-edge (instead of the K-edge) and a special high energy resolution XAS technique on the Ir L-edge. The adsorbed O atoms cause a change in the oxidation state of the Ir atom, in agreement with previous *operando* x-ray photoelectron spectroscopy studies of IrO₂ nanoparticles⁴². The results give important insights into the reaction mechanism by detecting that the adsorbed O atoms are the stable intermediate providing support for the proposed theoretical mechanism³⁰.

Experimental setup

Sample preparation

The hybrid IrO_x/RuO₂ system was tested by depositing 2 Å of IrO_x on top of a 40 nm RuO₂ thin film. Both films were deposited by magnetron sputtering of a Ru or Ir target at 30 W with the sample substrate being held at 300°C. The oxide was formed during sputtering by a reactive atmosphere at 3 mTorr and a flow of 10 sccm Ar and 4 sccm O₂. The sputter rate was monitored by a quartz crystal microbalance (QCM). Samples of 40 nm RuO₂ thin film were deposited on polycrystalline Au disks, followed by the deposition of 2 Å of IrO_x (sample 2 Å IrO_x/RuO₂/Au). After the deposition, the substrate was held at 300°C for 30 minutes for annealing. Considering a conservative roughness factor of 2 for the RuO₂ thin film, the 2 Å film thickness of IrO_x correspond to sub-monolayer coverage.

The mass-selected ruthenium nanoparticles were prepared under ultra-high vacuum (UHV) using a nanoparticle source equipped with a magnetron sputter gas-aggregation source coupled to a lateral time-of-flight mass filter. The production method can be described in five steps: 1) A ruthenium metal target was sputtered with Ar⁺-ions in a DC magnetron sputter head, 2) the sputtered metal atoms were allowed to condense into nanoparticles in a liquid N₂ cooled "aggregation zone", where He gas was added to facilitate aggregation³⁶. The total pressure in the aggregation zone was in the order of 0.4 mbar. 3) The nanoparticles exited the aggregation zone through an aperture to a differentially pumped region, thus undergoing a supersonic expansion. The nanoparticles then entered an electric lens system, where the negatively charged particles were focused into a narrow beam. 4) The particles were mass-filtered in a lateral time-of-flight mass filter with a resolution of m/ Δ m=20⁴⁴, set to select particles with a mass of 500,000 amu, corresponding to 5 nm in diameter assuming spherical

shape and the bulk density of Ru. 5) The particles were deposited on the glassy carbon substrates supported by steel frames, through an aperture with a diameter of 9 mm, yielding a total loading of about 400 ng. One set of nanoparticles were subsequently oxidized at 400 °C in a flow of pure O_2 for 1 minute, while the other sample got oxidized by being exposed to air for 72 hours.

The used apparatus was a Nano-Beam 2011 nanoparticle source (Birmingham Instruments Inc.), which has a base pressure in the low 10⁻¹⁰ mbar range. For more information about the preparation of mass selected nanoparticles see earlier papers from our laboratory.^{26,45,46}

The nanoparticle source is coupled to another UHV system (Scienta Omicron Multiscan Lab XP), where XPS characterization was conducted using a SPECS XR 50 X-ray gun coupled to an Omicron NanoSAM 7 channel energy analyzer. This chamber has a base pressure in the low 10⁻¹¹ mbar range. The coupling of the two vacuum chambers allows XPS to be carried out after deposition without exposing the nanoparticles to air.

In order to verify that the nanoparticle mass-selection had resulted in a narrow size-distribution, Ru nanoparticles were deposited on Cu TEM grids with a lacey carbon support layer under the same deposition conditions as the samples. One grid was exposed to the same oxidation heat treatment as the glassy carbon windows (400 °C for one minute in a flow of O₂), whereas the other was exposed to air for 72 hours. The TEM grids were then transferred to a Tecnai T20 G2 transmission electron microscope for bright-field imaging. Using the software "ImageJ", the TEM images were subsequently analyzed to obtain size distributions of the deposited nanoparticles. The results were filtered, in order to exclude particles that had landed on top of each other during the deposition.

Synchrotron radiation spectroscopy measurements

All synchrotron measurements were carried out at Stanford Synchrotron Radiation Lightsource (SSRL) at SLAC National Laboratory in California, USA. High energy resolution fluorescence detected X-ray absorption spectroscopy (HERFD-XAS) was performed at beam line 6-2, a hard X-ray wiggler beam line equipped with a Si(111) monochromator in the $\phi=0^{\circ}$ setting. The spectrometer was of the Johann type with 7 bent Si(553) single crystals and equipped with a Vortex fluorescence detector⁴⁷. The intensity of the incoming and transmitted beams was detected using ion chambers. HERFD-XAS was measured at the Ir L₃-edge (11215 eV) on the 2 Å IrO_x/RuO₂/GC sample, focusing on the X-ray absorption near edge spectrum (XANES) only.

Conventional XAS measurements were made at beam line 14-3, a tender X-ray (2 – 5 keV) bending magnet source equipped with a Si(111) monochromator in the ϕ =0° setting. A 4-element Vortex detector was used for fluorescence detection and an ion chamber was used for initial beam intensity detection. Due to the low photon energy, the entire beam path was in a He atmosphere at 1 bar to reduce absorption. At this beam line Ru L₃-edge (2838 eV) XANES were measured on the Ru NP/GC and RuO₂ NP/GC samples.

All experiments were done in an *operando* cell, and two different setups were used at the Ir and Ru L_3 -edges due to the difference in photon energy and thereby different absorbance from the electrolyte. For the Ir L_3 -edge experiments we used a hanging meniscus setup developed at the Nilsson group at SLAC National Laboratory, which is described elsewhere⁴⁸. The cell had a motorized

syringe attached, so that the size of the meniscus could be controlled accurately, forming a cylindrically shaped meniscus 5 mm in diameter. The electrolyte used was 0.05 M H₂SO₄ with a Pt wire counter electrode and a Ag/AgCl reference electrode (Innovative Instruments, Inc. LF-1.6, 3.4M AgCl). To control the potentials a BioLogic potentiostat (BioLogic Scientific Instruments) was used, which was controlled by a separate PC.

For the Ru L₃-edge experiments a back-illumination type cell was used, in which the detected X-rays only penetrate the electrode and not the electrolyte. We fabricated our own electrodes by pyrolyzing 8 µm thick Kapton foil (DuPont) at 1000°C for 1 hour in a flow of N₂. These windows were mounted in stainless steel frames to make handling easier, and a 7 mm diameter hole in the frame defined the active electrode area. Each frame was mounted on a 50 ml high density polyethylene (HD-PE, Nalgene) bottle by epoxy glue, making sure to cover the steel frames completely with epoxy, thus decreasing the active electrode area slightly. Figure S1 in the supporting information (SI) shows pictures of the setup. To avoid the fluorescence from S in the electrolyte, we used 0.1 M HNO₃ electrolyte for these experiments. The electrolyte, a Pt wire counter electrode, and a Ag/AgCl reference electrode (identical to the Ir experiments) were all housed in the HD-PE bottle.

The mass-selected nanoparticles measured at the synchrotron were the same samples that were characterized by XPS in the preparation laboratory, i.e. the as-prepared Ru nanoparticles after exposure to air (5 nm Ru NP/GC) and the thermally oxidized RuO₂ nanoparticles (5 nm RuO₂ NP/GC). We have reported on the preparation and characterization of the IrO_x thin film (2 Å IrO_x/RuO₂/Au) elsewhere³⁷.

Simulating XAS spectra

XAS spectra were simulated using version 9.6 of the FEFF software package⁴⁹. All the simulations were run on the Abisko cluster, which is a part of the Swedish National Infrastructure for Computing (SNIC). We simulated the Ir L₃ edge to compare to the measured spectra on the 2 Å IrO_x/RuO₂/Au sample. These measurements were done using the HERFD XAS technique that reduces the life-time broadening, and to compensate for this in our simulations we reduced the natural broadening by 1.5 eV.

The simulated surface was based on the rutile RuO₂(110) surface, but with the top-most Ru atoms in the surface replaced by Ir. The 2 Å IrO_x/RuO₂/Au sample was polycrystalline, but since the (110) surface is the most stable rutile surface, we chose to use this as the model surface. This surface has two different sites; one with 2 O atoms below the surface, 2 O in the surface, and 2 bridging O above the surface. The other site has 1 O below the surface, 4 O in the surface plane, and 1 O above the surface, see Figure S2. At the potentials used in this experiment the bridge sites are assumed to be occupied by $O^{34,35}$, whereas the different intermediates are found on the coordinatively unsaturated site (cus) on the second Ir site. To get the overall XAS spectra, both sites are simulated and their results averaged. To compare the white line positions of the simulations to those of the measurements, we fitted the simulated spectra using the same procedure as for the measurements.

Results

The combined $IrO_x/RuO_2/Au$ system was structurally and electrochemically characterized previously, with results published in reference³⁷.

To get more insight into the reaction mechanism of OER on IrO_x and RuO_x we fabricated metallic Ru and thermally oxidized RuO₂ nanoparticles. These nanoparticles have been characterized previously in¹⁰, including GA-XRD showing crystallinity of both the as-deposited metallic and thermally oxidized particles. The 5 nm particles in this study are well dispersed and are quite round as seen in the TEM images in Figure S3a and S3b. The as-deposited metallic particles have a narrow size distribution of 4.9±0.3 nm, whereas the oxidized particles have a size of 5.2±0.5 nm but with a tail towards large particle sizes, which could indicate coalescence of a small fraction of particles during the oxidation procedure. The size histograms and fitted size distributions are shown in Figure S3c. The change in particle volume does not correspond to a complete oxidation assuming Ru conservation. However, the as-prepared Ru nanoparticles were transported in air to the TEM, and from the XPS analysis we know that these nanoparticles are partially oxidized (see below). The thermally oxidized RuO_2 nanoparticles show a well-ordered (from TEM) rutile structure (from GA-XRD) based on a previous study¹⁰, and this suggests that these nanoparticles are fully oxidized. Assuming a core-shell structure of the as-prepared Ru nanoparticles, based on the size change they consist of a 2.6 nm metallic core and a 1.15 nm thick oxide shell.

The nanoparticles were further characterized by XPS, and spectra were acquired before being exposed to air, after 72 hours of exposure to air, and the thermally oxidized samples after the oxidation procedure. Only C, O, N and Ru can be found in the survey spectra taken of both the

metallic Ru after air exposure and oxidized RuO₂ after annealing in the tube furnace (see Figure S4). Compared to the ruthenium peak energies after deposition (Ru 3d_{5/2} at 280.4 eV and Ru 3p_{3/2} at 461.8 eV) the peak energies in case of both samples appear to be shifted to higher binding energies (details about peak energies and peak fitting can be found in the supporting information). Figure 2 shows the overlapping C 1s - Ru 3d peaks and the Ru 3p_{3/2} peak for the Ru and RuO₂ samples. As expected the RuO₂ sample has a larger binding energy shift, and thus appears to be more oxidized. The SI contains further details of the XPS characterization, including analysis of the Ru 3p lines.



Figure 2. Detailed Ru 3d and $3p_{3/2}$ XPS spectra of the Ru nanoparticles samples after subtraction of a Shirley-background.

XAS experiments were performed to gain further insight into the chemical state of Ir and Ru during OER. Figure 3 shows XAS spectra of the 2 Å IrO_x on 40 nm RuO_2 thin film on a Au substrate (2 Å $IrO_x/RuO_2/Au$, Figure 3a), 5 nm mass-selected metallic Ru nanoparticles (5 nm Ru NP/GC, Figure 3b),

and 5 nm mass-selected oxidized RuO₂ nanoparticles (5 nm RuO₂ NP/GC, Figure 3c). These spectra are all acquired for the dry as-prepared samples, and they are compared to reference spectra of their respective metals and rutile oxides. The Ir metal and IrO2 references are both sputtered thin films, whereas the metallic Ru reference is a foil from the SSRL reference collection and the RuO₂ reference is a powder (Sigma-Aldrich, 238058-1G, 99.9% purity). The Ir XAS experiment was done using the HERFD XAS technique, and the Ir $L_{\alpha 1}$ emission line was measured prior to the experiment (see Figure S9), and the maximum intensity at 9175 eV was chosen for all XAS spectra. This technique gives an effective life time broadening of 2.1 eV^{50–52}, which is similar to the intrinsic L₃-edge life time broadening of Ru of 2.0 eV⁵¹. The IrO_x sample spectrum is very similar to that of the IrO₂ reference (Figure 3a). The major difference is that the sharp peak just above the absorption edge (the white line) is more intense for the sample, which could be caused by polarization effects in the highly anisotropic IrO_x surface layer. The RuO_2 mass-selected nanoparticles (Figure 3c) are similar to the RuO₂ reference spectrum as expected. Again, the major difference is in the white line intensity, with the sample being more intense than the reference. The RuO_2 reference spectrum reveals a splitting of the white line caused by a crystal field splitting of the Ru 4d states⁵³. But the RuO₂ nanoparticles show this splitting to a much smaller degree, possibly due to a lower degree of crystallinity caused by the small size of the particles. At this point it is also worth noting that the IrO_x does not display any crystal field splitting, consistent with conventional XAS measurements on Ir-compounds^{43,54,55}. Finally, the metallic Ru nanoparticles (Figure 3b) are most similar to the metallic Ru reference as expected. However, the white line is much more intense than the reference, which we expect to be caused by the formation of a surface oxide, as observed by the XPS measurements discussed earlier.



Figure 3. XAS spectra of (a) the 2 Å IrO_x on 40 nm RuO₂ thin film on a Au substrate (2 Å IrO_x/RuO₂/Au), (b) 5 nm mass-selected metallic Ru nanoparticles (5 nm Ru NP/GC), and (c) 5 nm mass-selected oxidized RuO₂ nanoparticles (5 nm RuO₂ NP/GC). They are compared to reference spectra of their respective metal and rutile oxide.

The *operando* XAS measurements of the IrO_x sample were first done at open circuit potential, followed by XAS measurements at various fixed potentials, keeping the potential constant for the duration of each XAS measurement step. All the experimental potentials are given relative to RHE. The potential was first set to 0.5 V and then stepped up to 1.6 V in steps of 0.1 V. The sample potential was increased directly, and there was no open circuit condition in between each step. Due to the bubble formation from the OER the meniscus was broken at 1.6 V, and after restoring it the potential was set to 1.5 V, and then stepped down to 0.6 V in steps of 0.1 V. The potential was then set to 1.0 V and stepped up to 1.7 V in steps of 0.1 V (except between 1.6 V and 1.7 V where the step size was 0.05 V), and this time the electrolyte meniscus stayed intact despite vigorous bubble formation. Figure 4a shows the white line of the 2 Å IrO_x/RuO₂/Au at different potentials, and it is

seen that it shifts with the potential. Compared to the open circuit potential (OCV) the white line has shifted down, remained the same, and shifted up at 0.5 V, 0.9 V, and 1.5 V, respectively. Assuming these changes are caused by core-level shifts due to oxidation state changes, the shifts are equivalent to a reduction, no change, and an increased oxidation, respectively.

The operando XAS of the metallic Ru nanoparticles was performed in a similar way, first at OCV before any potential cycling, followed by OCV after 21 cycles from 0.8 V to 1.2 V. The XAS measurements were then done at fixed potentials from 0.8 V to 1.4 V in steps of 0.1 V (except between 1.3 V and 1.4 V, where the step size was 0.05 V), and the potential was not turned to OCV in between potential steps. The white line at OCV, 0.8 V, and 1.3 V is shown in Figure 4b. At 1.3 V a significant shoulder is seen towards high energy, indicating the growth of an oxide phase. This experiment was stopped after 1.4 V as the Ru had almost completely disappeared, see Figure S10a. In this case the loss of Ru is suspected to be mostly dissolution, as metallic Ru nanoparticles have proven quite unstable in acid¹⁰. Despite the dissolution the measurements are still informative of the OER reaction, as the current is dominated by OER¹⁰. Some Cl contamination was observed, as the Cl K-edge is only ~16 eV below the Ru L₃-edge. It is suspected to be in the form of organic Cl complexes dissolved from the epoxy glue used to mount the sample on the bottle. The amount of Ru and Cl was normalized to the amount of Ru at 0.8 V. Cl evolution has a lower overpotential than OER⁵⁶, and so the Cl we observe is not free Cl ions, as they would get evolved as chlorine gas and disappear immediately. The current during the experiment was not influenced by the Cl concentration, so the Cl did not interfere with the OER.

Finally, the oxidized RuO₂ nanoparticles were measured at OCV both before and after 5 cycles of the potential in the range 0.8 V to 1.2 V. The white line at OCV, 0.8 V, 1.3 V, and 1.7 V is shown in Figure 4c. The changes in the white line position are very small, which is due to a rather low surface to bulk ratio in the 5 nm particles. For this sample XAS was measured at fixed potential from 0.8 V to 1.3 V, back to 0.8 V, and up to 1.7 V. Below 1.2 V the potential step size was 0.2 V, and above it was 0.1 V, and there were no OCV periods in between the potential steps. Figure S10b shows the Ru amount, which was much more stable than the other two Ru samples, but due to the longer running experiment the Cl contamination became guite high.



Figure 4. The white line at different potentials for the (a) 2 Å $IrO_x/RuO_2/Au$, (b) 5 nm Ru NP/GC, and (c) 5 nm RuO₂ NP/GC. For the ultra-thin films of IrO_x the white line moves significantly as a function potential, whereas for the nanoparticles the shift is smaller due to a smaller surface to bulk ratio. The metallic Ru nanoparticles get a shoulder towards high energy at 1.3 V due to the formation of an oxide phase.

Figure 5 shows the white lines of the three samples at each potential, stacked on top of each other so that its position can be tracked from bottom to the top. The white lines of the ultra-thin film of IrO_x follow the potential clearly as discussed above, and as indicated by the red dots. For the metallic Ru nanoparticles, the white line shape changes significantly at the OER and dissolution onset, until the oxide phase becomes dominant at 1.4 V. The positions of the metallic and oxide peaks seen in the white lines are indicated by dashed red lines. The white line position of the oxidized RuO₂ nanoparticles changes very little, but the trend is seen from the fitted (see below) peak position indicated by the red dots. Interestingly the white line shifts to slightly lower energy at the highest potentials.



Figure 5. The white lines at each potential for (a) 2 Å IrO_x/RuO₂/Au, (b) 5 nm Ru NP/GC, and (c) 5 nm RuO₂ NP/GC. The white lines have been offset in the y-direction for clarity, and the beginning of each experiment is on the bottom and the end at the top. For (a) and (c) the red dots indicate the movement of the white line as a function of potential based on peak fitting. In (c) the white line is fitted by two peaks, therefore two sets of red dots are shown. For (b) the two dashed red lines show the positions of the metallic and oxide peaks in the white line.

To interpret the changes in white line position, all the spectra have been quantified by curve fitting. The Ir spectra were fitted with an arctan function as the absorption step and a single Lorentzian for the white line itself. The arctan function was centered at the inflection point for each spectrum, and the Lorentzian peak was fitted by position, width, and height. For the Ru spectra, an arctan function was also used for the absorption step, and two Lorentzian peaks were used to fit the white line due to the crystal field splitting in RuO₂ nanoparticles and coexistence of a metallic and oxide phase in the Ru nanoparticles. The arctan function was again centered at the inflection point of each spectrum, but to get stable fits the two Lorentzian peaks had their widths fixed at 2.8 eV (FWHM), which agrees with the lifetime broadening⁵¹ and gave reasonable fits. The two peaks had their positions and heights fitted freely.

Figure 6 shows the white line Lorentzian peak positions as a function of electrochemical treatment along with the applied potential. For the Ir white lines the position of the single Lorentzian is shown, and for the Ru lines the peak position for the high-energy Lorentzian peak is shown. The error bars represent a confidence interval equivalent to 1 σ obtained from the fitting algorithm. For the Ru spectra additional error due to the Cl contamination was added by simulating the effect of adding a Cl spectrum to a pure Ru spectrum. The peak positions follow what was observed in the raw spectra, but for the IrO_x sample it seems that the white line position does not change any further above 1.5 or 1.6 V. Similarly, for the metallic Ru nanoparticles the peak position saturates at 1.3 V. For the RuO₂ nanoparticles the peak position not only saturates at 1.3 V, it even goes to lower energy. The

inflection point found by center-of-mass analysis of the first derivative is shown in Figure S11 in the SI, and shows the same trends as discussed, just with a smaller movement in energy.



Figure 6. The white line position is shown from fitting the XAS spectra for (a) 2 Å $IrO_x/RuO_2/Au$, (b) 5 nm Ru NP/GC, and (c) 5 nm RuO₂ NP/GC. For the Ir spectra the single Lorentzian peak position is shown, but for the Ru spectra only the high-energy Lorentzian peak position is shown, as it is the most dominant component.

As the XAS measurements here probe transitions from the Ir (Ru) 2p states into empty 5d and 6s (4d and 5s) states, the peak shifts can be caused either by changes in the initial or final state. The initial state energy may change when the absorbing atom changes oxidation state, leading to a core-level shift, whereas changes in the final state energy is tied to changes in the d-band. The XAS features above the white line are linked to the structure due to multiple scattering resonances, and so by comparing this region for different potentials we can see whether the whole spectrum has been shifted by a core-level shift, or if only the d-band has changed.

Figure 7 shows the XAS features above the white line for the three samples. Multiple spectra at low potential and multiple spectra at high potentials have been averaged to provide a better signal-to-noise ratio. Furthermore, the high potential spectrum has been shifted down to align the white lines. The metallic Ru nanoparticles show signs of a changed structure, as the energy of the main feature at around 2890 eV has changed and the peak at ~2868 eV disappears at high potential. This is not surprising, as the metallic Ru undergoes oxidation and forms an oxide phase. For the other two samples, there are no significant changes between the spectra at low and high potential, thus the white line shifts correspond to core-level shift entirely. This means that the shifts we observe are caused by oxidation state changes without any significant structural changes. The change in surface Ir oxidation state by the OER reaction is consistent with previously reported measurements⁴².



Figure 7. The XAS features just above the white line for (a) 2 Å $IrO_x/RuO_2/Au$, (b) 5 nm Ru NP/GC, and (c) 5 nm RuO₂ NP/GC. Any change in this region indicates structural changes, which is only seen in (b), as the metallic Ru nanoparticles oxidize under the measurement conditions. The lack of differences in the other spectra indicates no structural changes. The spectrum taken at high energy has been shifted

to lower energy as indicated in the figure, and the amount is consistent with the core level shifts seen in Figure 6.

We investigated the link between the core level shift and the reaction intermediates further by simulating the XAS spectra obtained on the 2 Å IrO_x/RuO₂/Au system using FEFF. We used Ir-O bond lengths consistent with Ru-O bond lengths reported in references^{35,57,58}, and the position of the white line is shown in Figure 8. Here it is seen that the majority of the core level shift is caused by the *O intermediate. Furthermore, the white line position is almost the same when the cus site is empty or covered by an *OO or *OOH intermediate. In order to be able to compare to other Ir-O bond lengths, we simulated a variety of bond lengths, and their white line positions are shown in Figure S12 in the S1. The error bars shown in Figure 8 are determined from fitting the simulated spectra with an arctan and Lorentzian peak like the measured spectra. The FEFF simulation tool does not provide estimates of the error in the simulation, and the absolute white line position is generally not very accurate in these types of simulations. However, as the atomic system is the same in all simulations, except for the adsorbed reaction intermediate, we believe that the relative error between the white line positions is on the order of the fitting errors.



Figure 8. The figure shows the position of the white line of simulated XAS spectra for surface configurations with different reaction intermediates. The distances indicate the Ir-O bond length of the given intermediate. The error bars are from fitting the simulated spectra using the same procedure as the measured XAS spectra.

The FEFF simulations also show that the measurements agree with a reaction mechanism involving a significant coverage of an *OO based intermediate during OER, as this gives rise to a white line

position almost identical to that of an empty cus site. So spectroscopically it would not be possible to determine whether this intermediate is present, but the results we have are consistent with its presence. The measured shift in the Ir white line is slightly higher than predicted by FEFF, but as the detailed spectroscopic shape around the absorption edge is difficult to calculate the inconsistency is within expected errors.

Discussion

In here we demonstrated that the shift in white line position is primarily cause by a core level binding energy shift, based on a comparison with multiple scattering resonances showing a similar shift as the white line. On this basis we can interpret the data in accordance with the proposed DFT based model of Rossmeisl et al^{29–31}, which shows that the overpotential of $IrO_2(110)$ is due to the conversion of *O to *OOH.³⁰ We observed that the oxidation state of IrO_x , manifested as the Ir white line position, is saturated at high overpotentials. According to the FEFF simulations shown in Figure 8, adsorbed *O causes a markedly more positive shift in the white line position than other adsorbates. Consequently, the white line position is a measure of the *O coverage. In this case the oxidation state increases with increasing potential, as the surface gets further covered by *O. At OER potentials the white line position saturates, which indicates that the surface gets fully covered with adsorbed O that are not turning over to form OO and therefore it would be desirable to destabilize the adsorbed O in future catalyst designs. This core level shift is in agreement with previous *operando* x-ray photoelectron spectroscopy studies of IrO_2 nanoparticles⁴², which also showed from photon energy dependent spectral sensitivity that the change of oxidation state occurs in the topmost Ir layer.

In contrast to IrO_2 , in the case of RuO_2 we observe a decrease in oxidation state beyond 1.5 V. At first glance, this phenomenon could seem counterintuitive. However, it is in fact consistent with the proposed DFT based model shown in Figure 1. In this case the energetically most stable intermediate just below the onset of OER is atomic *O, and a full coverage of *O will bring the Ru in the highest average oxidation state. As the potential is increased somewhat above the potential where all the reaction steps are all downhill in energy, the coverage of *O is now determined by the kinetics of the reaction rather than the thermodynamic stability of each intermediate. As seen in Figure 1 the most difficult step is to go from *O to *OOH, but when the step is overcome the driving force to evolve O_2 gas is very high. Similarly, when the first water molecule is adsorbed to form *OH the further oxidation to *O has a large driving force. Therefore, during OER the two dominant states of Ru are empty sites (*) and covered by atomic *O. In the limit of similar reaction rates for * \rightarrow *O and * $O\rightarrow O_2(g)$ the coverage of *O is 50%. And this lower coverage of *O leads to a lower average oxidation state.

It is interesting to note that we do not observe a decrease in oxidation state on IrO₂, which, according to theoretical calculations, is also limited by *OOH formation³⁰. However, our experimental observation is consistent with the notion that the overpotential required to drive the reaction is higher on IrO₂ than RuO₂. On this basis, we would expect that IrO₂ would also exhibit a decrease in oxidation state at even higher overpotentials.

Conclusion

We measured *operando* XAS spectra on RuO_x based nanoparticles and a IrO_x/RuO₂ thin film, as a function of electrochemical potential. We found that the observed changes in the spectra are primarily due to core level shifts caused by changes in the oxidation state. An exception was metallic Ru nanoparticles, which went through structural changes when forming an oxide phase at high electrochemical potentials. The changes in oxidation state are consistent with an earlier, DFT based reaction mechanism proposed by Rossmeisl, Nørskov and coworkers^{29–31}.

Simulations of the XAS spectra showed that the average oxidation state changes were dominated by the coverage of an atomic O intermediate. The IrO_x showed a saturation of the oxidation state at high electrochemical potentials, meaning that the surface is fully covered by O, consistent with IrO_x binding O too strongly.

The decrease in oxidation state that occurred for the RuO₂ nanoparticles is rationalized by RuO₂ binding the atomic O intermediate too strong compared to the optimum, but weaker than IrO_x. In this case the surface will be covered by atomic O at potentials just below the OER onset, giving the maximum average oxidation state, and during OER the coverage of atomic O will decrease as the active sites are turning over oxygen gas. These observations limit the Ru-O bond strength to be weaker than the Ir-O bond strength, but still stronger than the optimum metal-oxygen bond for OER.

Associated content

Supporting information contains more detailed information on the experimental procedure, sample preparation, and sample characterization.

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TOC Graphic

