Electrochemically active Ir NPs on graphene for OER in acidic aqueous electrolyte investigated by in situ and ex situ spectroscopies

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A B S T R A C T

An electrode for the oxygen evolution reaction based on a conductive bi-layered free standing graphene support functionalized with iridium nanoparticles was fabricated and characterized by means of potentiometric and advanced X-ray spectroscopic techniques. It was found that the electrocatalytic activity of iridium nanoparticles is associated to the formation of Ir 5d electron holes. Strong Ir 5d and O 2p hybridization, however, leads to a concomitant increase O 2p hole character, making oxygen electron deficient and susceptible to nucleophilic attack by water. Consequently, more efficient electrocatalysts can be synthesized by increasing the number of electron-holes shared between the metal d and oxygen 2p.

1. Introduction

The treatment of greenhouse gases and the mitigation of their effects is one of the most important global challenges humanity is now facing [1]. These challenges include the effects of global warming on the climate and their consequences for humans [2], plants, and animals [3], as well as the negative health effects of toxic gases and suspended nanoparticles [4,5]. This issue has to be solved while still providing enough energy to feed a growing population (more than 7 billion) and to operate an infrastructure, which requires bigger and more efficient transportation networks [6]. Renewables (including wind, solar and tidal power) enable opportunities for a large, continuous, and clean energy supply [7]. However, one of the biggest challenges for an economically viable implementation of renewable energy technologies in the current energy infrastructure is caused by the intermittent nature of these energetic resources, which does not match the demand. This mismatch requires strategies to store excess energy for later use. Electrochemical energy conversion and storage plays a crucial role in the overall solution of the global energy challenges [8]. In particular, hydrogen obtained via electrochemical oxidation of water has the potential to replace fossil fuels as a storable and clean fuel [9]; however, this technology requires further development before it can be applied on a large scale [7]. In fact, because of the slow kinetics of the anodic oxygen evolution reaction (OER) [10], it is necessary to apply potentials well beyond the thermodynamically required potential of 1.23 V (vs. SHE) to split water molecules into their principal components (hydrogen and oxygen). For example, to achieve current densities of at least 10 mA/cm\textsuperscript{2} over-potentials of hundreds of millivolts above the OER equilibrium potential have to be applied, with a drastic reduction of the process efficiency [11,12].

Due to their low overpotential and high stability in acidic media iridium-based oxides have been shown to be good candidates for the electrocatalytic oxidation of water [13,14]. Unfortunately, iridium is among the scarcest materials in the earth's crust. Thus, optimized strategies are required for its usage; e.g. by increasing the surface to bulk ratio of the iridium particles [15]. An electronic effect exerted by the active Ir metal species causes the enhanced activity observed, however a detailed chemical speciation is still elusive and is even contradictory in literature in many cases, which requires further investigation to optimize synthesis approaches. Well dispersed...
nanoparticles (NPs) are usually not electrically connected between each other requiring a different strategy to apply the required potential to flow the electrical current necessary to run the reaction. This limitation can be circumvented by using a conductive support material, which drives the current to the catalytic active centers. In this way, the support layer provides electrical contact for the different catalyst NPs yielding high activity and stability under OER conditions, comparable to bulk materials [16]. Graphene was selected as a possible candidate substrate due to its high mechanical strength and flexibility, exceptional thermal and electrical conductivity, and chemical stability under harsh reaction conditions [17–20]. It is well known that defect-free graphene layer is not electro/chemically active [21] and requires functionalization with heteroatoms to manifest chemical reactivity making it as an ideal substrate for electrocatalytic investigations due to its low contribution to the overall reaction rates, allowing the correlation of the catalytic activity to the supported metal.

Photoelectron spectroscopy (PES) is a powerful surface sensitive technique, which provides information about the electronic structure of the core-levels in a non-destructive way. This technique has been applied to a wide range of systems in order to elucidate their chemical composition. The implementation of this technique under reaction conditions is however complicated by the so called “pressure gap” [22] between the high pressure required for realistic reaction conditions and the ultra-high vacuum (UHV) needed in conventional PES chambers for their operation. Recently, holey grids made of Si nitride were combined with photoelectrons transparent bi-layers graphene (BLG) to form windows, which separate the liquid environment from the evacuated (UHV) PES chamber [23,24]. With this approach it was possible to investigate electrochemical electrocatalysts interfaces in the presence of liquid electrolytes and gases [22] under reaction conditions. By taking advantage of this approach, in situ X-ray photoelectron spectroscopy (XPS) was accomplished, allowing the direct correlation between the electronic structure of IrOx (active sites) and its electrocatalytic activity during OER.

In this work, we investigate the variation in the electronic structure of Ir NPs on conductive bi-layers of graphene (BLG) as consequence of applied potential under OER in acidic conditions. We combine in situ with ex situ X-ray photoelectron spectroscopy (PES) and X-ray absorption near-edge structure (XANES) to unveil still undiscovered electronic structural information on the IrOx, active in OER. In particular, valuable information of the occupied (Ir 4f) and partially occupied (Ir 5d, O 2p) orbitals is obtained with these techniques. We show that the formation of electron-holes in the hybridized O 2p/Ir 5d levels is linked to the electrochemical activity shown by iridium oxide. These holes lead to the formation of electron deficient oxygen species, which are the likely centers in the formation of molecular oxygen due to nucleophilic attack by OH or H2O. Furthermore, we report the structural descriptor of the electrocatalytic activity as a consequence of Ir 5d and O 2p hybridization and shared electron-holes and provide guidance for the synthesis of improved electrode materials for OER.

2. Experimental

2.1. Beamsline

In situ synchrotron radiation based experiments were performed at the ISIS beamline of BESSY II in Berlin (Germany). In this facility, the photons are sourced from a bending magnet (D41) and a plane grating monochromator (PGM) yielding an energy range from 80 eV to 2000 eV (soft X-ray range), a flux of $6 \times 10^{19}$ photons/s with 0.1 A ring current using a 111 µm slit and a 80 µm × 200 µm beamspot size. The in situ measurements were accomplished in the ambient pressure X-ray photoelectron spectrometer (AP-XPS) end-station of the ISIS beamline [25].

Hard X-ray absorption measurements at the Ir L$_3$-edge were performed at the beamline BL17C1 [26] of the National Synchrotron Radiation Research Center (NSRRC) in Hsinchu (Taiwan). The photons are sourced from a 25 poles wigglor (W20) with 20 cm period length and a focus spot size of 2 mm × 6 mm. The excitation energy ranges from 4.8 keV up to 14.2 keV. The signal was collected in transmission mode for the powder samples and in fluorescence yield mode for the foil, using an ionization chamber detector.

2.2. Electrode preparation

Graphene was grown by catalytic chemical vapor deposition (CVD) on a 20 µm thick Cu foil (Alfa Aesar 99.8%) using CH$_4$ (diluted in Ar and H$_2$) served as the precursor at 1000 °C in an Aixtron BM Pro (4 inch) reactor. This procedure yielded a continuous polycrystalline film with a grain size in the range of ∼20 µm, which was confirmed by scanning electron microscopy. The graphene layer was fixed to 500 nm of Poly(methyl methacrylate) PMMA (4 wt.% in anisole, 950 k molecular weight) deposited by spin coating. After that, the Cu/graphene/PMMA composite was left floating on a 50 mM aqueous solution of (NH$_4$)$_2$S$_2$O$_8$ in order to detach the copper support. Then the graphene/PMMA layer was floated in deionized water (DI-water) and transferred onto another graphene/copper foil and dried at 50 °C for 5 min. The resulting sample was floated again on the 100 mM (NH$_4$)$_2$S$_2$O$_8$ solution, to remove the copper foil, and rinsed in DI-water. The PMMA/BLG layer was transferred onto a micro-structured Norcada $^+$ Si$_x$N$_4$ TEM grid with a holey area of 200 µm × 200 µm with 500 nm holes and a 1.2 µm pitch distance between them. Finally the PMMA was removed by dissolution in acetone and rinsing in isopropanol (IPA). The adherence between the BLG and the substrate is due to van der Waals interaction ensuring stability for the electrodes. It produces a continuous film that can be used as an electrode in aqueous environments and electrocatalytic applications, among others [21,27]. After the graphene had been transferred to the support, Ir nanoparticles were sputtered onto the surface with a Cressington 208HR sputter coater. The target was Ir 99.99% purity grade (Elektronen-Optik-Service GmbH, Dortmund, Germany), the Ar pressure 0.1 mbar, the current 40 mA, and the sputtering duration 5 seconds. This deposition provides homogeneously distributed Ir NPs on the surface of the graphene, as shown in the TEM measurements (Fig. 1A). A detailed analysis of these measurements revealed a uniform distribution of ∼2 nm NPs covering 22% of the surface on average (Fig. 1A). These particles are composed of metallic Ir which is confirmed by electron energy loss spectroscopy (EELS), where the measured Ir O$_{2.3}$ and Ir N$_{2.7}$ edges correspond to the metallic Iridium spectra [28] (see Fig. 1D). A closer inspection to the morphology shows that the particles are well anchored to the graphene support and exhibit Ir$^0$ (111) face centered cubic (fcc) crystal structure with a lattice plane distance of 0.23 nm, which was confirmed by the Fast Fourier Transformation (FFT) of HRTEM image and is in good agreement with the EELS measurements. The excellent anchorage of the iridium NPs in graphene was confirmed by the ex-situ TEM characterization after the OER (Fig. 1C).

The transmission electron microscopy (TEM) and electron energy loss spectroscopy (EELS) analysis were performed in a Philips CM200 FEG TEM equipped with a GIF Quantum energy filter and Tridiem camera. The EELS spectra were acquired in TEM mode with 1 s acquisition time and energy dispersion of 0.1 eV/channel.

2.3. In situ electrochemical cell

The liquid flow cell was operated inside the main chamber of the AP-XPS endstation of the ISIS beamline [25] at a background pressure of ∼10$^{-7}$ mbar while aqueous solutions circulated on the back side of the free standing graphene membrane described above, which is used as the working electrode. The continuous flow of liquid was assured by a syringe pump 260D Teledyne Isco (Lincoln, USA). The main body of the electrochemical cell is made of polyether ether ketone (PEEK), which is an exceptional electrical insulator and chemically inert. The sealing was
assured with several Kalrez O-rings which have good chemical stability. The cell contains two additional electrodes, the counter- (platinum) and reference-electrode (Ag/AgCl DRIREF-2SH, World Precision Instruments, USA). A cross-section of the cell is depicted in Fig. 2A. A more detailed description was previously reported [24]. Fig. 2B shows the detection scheme through the membrane which requires photoelectrons of at least 500 eV to "escape" through the free standing graphene and reach the vacuum side where the analyzer is placed. The free-standing graphene membrane is used as an electrode, as a support for the Ir NPs, and as a photoelectron transparent window preventing the liquid electrolyte from leaking into the chamber. A scanning electron microscope (SEM) image of the SiNx grid coated with a free standing BLG and decorated with Ir NPs is shown in Fig. 2C. With this approach, it was possible to investigate electrochemical interfaces in presence of liquids. Herein, we use it to investigate the electronic structure of iridium in presence of an aqueous acidic electrolyte under different potentials. This cell was operated in the ISISS beamline of the synchrotron facility BESSY II (Berlin, Germany) [25] establishing a pressure in the main chamber of $\sim 10^{-7}$ mbar while aqueous electrolyte (0.1 M H$_2$SO$_4$) is flowed (1 ml/min) on the back side of the membrane, which removes the bubbles formed during the OER and by water radiolysis. The Ir 4f spectra were collected with a kinetic energy of the photoelectrons of 600 eV (excitation energy of 660 eV) at different potentials. At open circuit potential (OCP) the Ir 4f peak consists of two main peaks ascribed to the spin orbit split Ir 4f$^{7/2}$ and Ir 4f$^{5/2}$ stats.

2.4. Electrolyte preparation

The electrolyte was prepared by diluting 9.8 g of H$_2$SO$_4$ (purity 98%, Alfa Aesar, Massachusetts, USA) in 1 l of Milli-Q water (18.2 MΩ) at room temperature (RT), 25 °C. The electrolyte was continuously saturated by bubbling pure N$_2$ gas, which minimizes the presence of other dissolved gases in the electrolyte. The electrolyte is acidic with a pH equal to 1.

2.5. Potentiostat

Potentiometric control was assured with a Biologic SP-300 (Seyssinet-Pariset, France) allowing different potentiometric and amperometric controls.

2.6. Reference samples

Commercially available iridium powders were purchased from Sigma-Aldrich (99.9% trace metals basis) and AlfaAesar (Premion, 99.99% trace metals basis). In addition, Ir foil was purchased from Mateck (99.99% metals basis). The AlfaAesar powder was used as received and the Sigma-Aldrich powder was washed in Milli-Q water and calcinated at 800 °C in O$_2$ for 50 hours. The AlfaAesar powder is X-ray amorphous (IrO$_x$) while the calcinated sample of Sigma-Aldrich exhibits the rutile-type structure (IrO$_2$). More details of the sample preparation/characterization can be found elsewhere [29]. The Ir foil was subjected to several cycles of sputtering in Ar$^+$ atmosphere and annealing in H$_2$ atmosphere and shows a purely metallic phase.

2.7. DFT calculations

All calculations were performed using the Quantum ESPRESSO package [30] using the Perdew, Burke, and Ernzerhof (PBE) exchange and correlation potential [31] with ultrasoft pseudo potentials from the PSlibrary [32]. A kinetic energy (charge density) cutoff of 30 Ry (300 Ry) was used. The IrO$_x$H$_y$ surface was modeled as a five-layer (110) slab separated from its periodic image by ca. 15 Å of vacuum. Both surfaces of the slab were allowed to relax while the central layer was held fixed.
at its bulk value. The Brillouin zone was sampled with a (12 × 6 × 1) k-point mesh using Marzari–Vanderbilt smearing [33] with a parameter of 0.01 Ry. Integrated charges were determined by a Löwdin population analysis.

3. Results and discussion

The electrocatalytic performance of the graphene electrode described in the experimental section was investigated by means of linear sweep voltammetry (LSV) and cyclic voltammetry (CV). Fig. 1B shows the cyclic voltammogram (CV) of this assembly performed at room temperature with a scan rate of 20 mV/s in 0.1 M H2SO4 versus a Ag/AgCl reference electrode and a Pt counter electrode. The CV shows two oxidation waves, I and II, and two reduction waves, IV and V, in addition to III, which is ascribed to the oxidation of water. It is broadly accepted that I/V and II/IV are redox couples [34]. In addition, the LSV of pristine graphene was compared to a graphene electrode decorated with ∼2 nm Ir NPs, which cover ∼20% of the electrode’s surface (inset Fig. 1B). The electrochemical characterization clearly indicates that the addition of Ir NPs decreases the OER overpotential significantly and increases the current density during OER. As expected, the electrode decorated with Ir NPs is more active, based on the onset potential and current density, than the pristine graphene electrode, proving the correct performance of the experimental methodology used in this work. Notwithstanding the increase in electrocatalytic performance of the graphene electrode when it is decorated with Ir NPs, the nature of the active sites under reaction conditions are still not completely understood. After the OER the electrode was investigated by ex situ transmission electron microscopy (TEM) and electron energy loss spectroscopy (EELS) techniques. Figs. 1C shows the TEM images of the sputtered particles on graphene after the OER, which are homogenous distributed particles on the surface of the graphene with coverage factor of around 20% and particles size of 2 nm in average. The fast Fourier transformation (FFT) pattern proves the existence of reduced iridium particles (induced by the beam) and the two additional rings ascribed to the existence of a non-well defined crystal structure due to the presence of non-stoichiometric iridium oxide (IrₓOᵧ) or/and defects in the lattice. At higher magnification it was found that the iridium nanoparticles are reduced to Ir metallic under vacuum condition in presence of the beam irradiation. It is a well-known issue related to the existence of a non-

Fig. 2. (A) Schematic drawing of the in situ electrochemical cell. (B) Detection scheme. (C) SEM image of the holey SiNₓ structure covered by free standing BLG. (D) In situ characterization of the Ir 4f electronic structure under different applied potentials in presence of 0.1 M H2SO4. (E) Ir L-edge before and after the oxygen evolution reaction of a 20 nm sputtered Ir thin film.
stoichiometric iridium oxide (IrOx) giving Ir\(^0\) (111) cubic structure with a lattice plane distance of 0.23 nm as product of the reduction induced by the beam. Furthermore, EELS performed before and after the OER, just as prepared Ir NPs and after OER, showing that the Ir NPs are metallic after the deposition by sputtering (as shows in Fig. 1D). On the other hand, after the OER, the particles present substantial changes in the electronic structure mostly ascribed to its oxidation state. The EELS spectrum presents a peak at around 51.5 eV and a shoulder at 56.5 eV, as well as Ir N\(_{5,7}\) edges at 62 eV and 65 eV for the N\(_7\) and N\(_6\) transitions respectively, which are ascribed to IrO\(_2\) [35]. The differences in the peak height between Ir as deposited and after OER indicate a change in the electronic structure due to the NPs oxidation. It is generally accepted that the active sites are hydrated Ir-oxohydroxides formed during the OER [36], however the molecular level description of the species responsible for the high catalytic activity is still debated. The results of the in situ characterization of this assembly are shown in Fig. 2D. The Ir 4f spectra were recorded by detecting electrons with a kinetic energy of 600 eV at different potentials. At open circuit potential (OCP), the Ir 4f peak consists of two main peaks ascribed to the spin orbit splitting in Ir 4f\(_{7/2}\) and Ir 4f\(_{5/2}\), respectively. Reference spectra from known Ir oxides are used to fingerprint features in the spectra recorded in situ. For this purpose, three samples were selected: metallic Ir foil, rutile powder, and amorphous powder, as described in the experimental section. Fig. 3A shows the Ir 4f spectra of metallic Ir, rutile IrO\(_2\) and amorphous IrO\(_x\), respectively. Reference spectra from known Ir oxides are used to fingerprint features in the spectra recorded in situ. For this purpose, three samples were selected: metallic Ir foil, rutile powder, and amorphous powder, as described in the experimental section. Fig. 3A shows the Ir 4f spectra of metallic Ir, rutile IrO\(_2\) and amorphous IrO\(_x\), respectively. The metallic main peak (blue) has a binding energy of 60.8 eV and IrO\(_2\) (red) is shifted to higher binding energies (61.7 eV). It is noticeable that for the case of IrO\(_x\) (green) the spectrum is broader than for IrO\(_2\) indicating the existence of another oxidation state at higher binding energy [29]. In addition, the comparison of these measurements with the ex situ Ir L\(_3\)-edge indicate the formation of more 5d electron holes as the consequence of the transition from occupied Ir 2p to unoccupied Ir 5d as shown in Fig. 2E.

In the in situ characterization of the Ir NPs immobilized on the BLG support the two Ir 4f\(_{7/2}\) and Ir 4f\(_{5/2}\) peaks are well described by an asymmetric Doniach-Sunjic (DS) [29] line shape with maximum at a binding energy of 61.7 eV and 63.8 eV, respectively. These binding energies are consistent with a metallic state of Ir in good agreement with the TEM characterization, showing homogeneous dispersed NPs with narrow size distribution of fcc crystals with a Ir\(^0\) (111) lattice plane at a distance of 0.23 nm. Two additional components fitted with a DS line shape appear at a binding energies of 61.8 eV and 64.8 eV, which, together with the shake-up satellites at 1 eV above the main line, can be ascribed to the formation of iridium oxide (IrO\(_x\)(A), like IrO\(_2\) rutile shape). Upon anodic polarization of the electrode at potentials of +0.5 V and +1.0 V the contribution of the IrO\(_x\)(A) species increases. Under OER conditions (+1.6 V), the spectrum evolves into a broader peak shape, which is fitted with two additional DS peaks at 62.3 eV and 65.3 eV and associated satellites 1 eV above the main peaks ascribed to the formation of an additional oxidation state (IrO\(_x\)(B), like amorphous IrO\(_x\) oxide as shown below). Such a broad Ir 4f line shape has been reported under reaction conditions using a proton exchange membrane (PEM) [37,38], which is also present under aqueous electrolyte conditions as it was shown in the in situ experiments in this work. The authors linked the higher binding energies of the Ir 4f level under OER conditions to the formation of an active mixed valence iridium oxide phase. The experiments showed that metallic iridium NPs are transformed into amorphous oxo-hydroxide with increasing potential. Meanwhile, Pfeifer and co-workers have provided for the first time the evidence of the formation of an electron deficient intermediate oxygen species, associated to the active oxide phase [39], providing not only new mechanistic insights into the OER but also confirming further APXPS as a unique tool in surface/interface science and catalysis. Therein, the electron deficient oxygen species were suggested to be susceptible to nucleophilic attack from OH groups or water yielding intermediated hydroperoxo groups, which finally decompose into molecular oxygen. These species are strongly correlated with the existence of an Ir mixed valence state, but unambiguous speciation is still challenging. This complication arises from the fact that the occupancy of the Ir 5d orbitals in the oxide is still unknown whilst its knowledge is of paramount importance as it is directly involved in the hybridization with the O 2p
orbits of the oxygen ligands determines their chemical nature and stability.

To help the assignment of the chemical state of the OER active Ir oxide phase, we complement core level Ir 4f PES with valence band (VB) PES and XANES at the Ir L-edge and O K-edge. The variation in the oxidation state has also a severe effect in the VB spectra (Fig. 3B) where the density of states (DOS) close to the Fermi edge are diminished following the shift to higher BEs observed in the Ir 4f spectra, which is linked to the occupied O 2p and Ir 5d orbitals. Fig. 3C shows the Ir L3-edge which corresponds to the electron transition from the occupied 2p3/2 level to the partially occupied Ir 5d (σ*) and Ir 6s (σ*) orbitals. The orbital splitting into t2g and eg states cannot be resolved in these spectra due the large lifetime broadening [40]. It is obvious that the increase in the main peak intensity of the Ir L3 spectra follows the trend \( \text{IrO}_x > \text{IrO}_2 > \text{Ir}_0 \) revealing that the number of unoccupied states in the 5d band is larger for the amorphous iridium oxide than for the rutile type [41]. Similar trends were revealed under in situ characterization of iridium electrodes [42,43] and in the ex situ characterization shown in Fig. 2E. Assuming the near surface region of the reference samples used as fingerprints is not significantly different than the bulk, the broader Ir 4f (collected at 900 eV KE) line shape for the amorphous iridium shows the same trend for the formation of 5d holes. The O K-edge spectra of the two different oxides are shown in Fig. 3D. The two sharp features at ca. 529–530 eV and 534 eV are related to the electron transition from the occupied O 1s core level to the unoccupied O 2p orbitals hybridized with the Ir 5d orbitals of the Ir, whereas the broader feature at higher excitation energies is due to the excitation of core electrons into the hybridized O 2p-Ir 6sp states [40]. The link between holes on Ir 5d and O 2p can be seen by way of DFT calculations. Fig. 4, for instance, shows how the amount of hole character of Ir and the surface O(H) to which it is bound depends on the oxidation state of a rutile-type IrO2 structure. These charges were obtained for a rutile-type (110) surface with various surface terminations, OH on the bridge and coordinatively unsaturated sites (CUS), O on the bridge and OH on the CUS sites, O on the bridge and CUS sites, and an empty bridge or empty CUS site. In each case the change in hole character (\( \Delta d^- \)) is defined relative to O/ Ir in bulk rutile-type IrO2 such that a positive number indicates an increase in hole character relative to the bulk oxide. Inspection of the figure reveals that as the metal site is oxidized, eliminating H terminations, the number O 2p holes on the O bound to that site also increase in almost linearly fashion. Interestingly, when the number of holes introduced becomes large oxygen acquires more hole density than Ir whereas for a small number of holes the character is shared equally between Ir and O. The O in these latter cases is the electrophilic oxygen species appearing at 529 eV in the O K-edge [39,44]. Thus, oxidation of Ir past the formal IV+ state in rutile-type IrO2 leads to the formation of electron deficient oxygen. Due to the O2p-Ir5d hybridization, the formation of electrons/holes in the Ir 5d leads to the formation of new oxygen species as seen in the pre-edge (\( \sim 529 \) eV) of the O K-edge XANES spectrum. The electron deficiency generated by the formation of 5d holes and the steric factor due to the presence of the other oxygen ligands around the Ir center are considered here as key factors in the formation of the electrophilic oxygen species previously observed, namely the formally \( \text{O}^{5+} \) species [34], which could explain the peculiar reactivity of Ir during the OER. This high degree of covalency leads to the sharing of electron-holes between O 2p and Ir 5 d seen by comparing the O K-edge and the Ir L-edge spectra. Within this picture, when the number of Ir 5d holes increases the strong hybridization with O 2p orbitals leads to a sharing of holes as shown schematically in Fig. 5A, in yellow. Therefore the feature shown in the O K-edge at 529 eV is associated with the amount of hole character introduced to oxygen as iridium is oxidized during the OER. These holes have two roles in OER: (i) as hole on the \( \text{IrO}_2 \) metal site that produces the key energetically non-favorable intermediate metal-oxo/oxyl species and, (ii) the formation of the electrophilic \( \text{O}^{5+} \) oxygen which are susceptible to nucleophilic attack.

Based on the electronic structure revealed by the X-ray spectroscopies used in this work and the fact that metal-oxo-oxyl [45] sites are proposed to be involved in the O–O bond formation leading to the oxygen evolution [46], two different mechanisms are proposed in literature [47]: in the acid base mechanism, the nucleophilic oxygen atom of water or hydroxide attacks the electrophilic oxygen of the metal-oxo-oxyl site and as consequence molecular oxygen evolves; otherwise, in the radical coupling mechanism, two adjacent metal-oxo-oxyl species with radical character couple and evolves molecular oxygen [48]. The acid-base mechanism requires electrophilic oxygen atoms on the surface while radical coupling requires oxygen atoms with radical character, both of which are electron deficient. In latter case, the oxygen radicals must be sufficiently close to each other to form an O–O bond. Hence, electronic, kinetic and steric factors will favor one or the other.
mechanism. The electronic factor is dominated by the electron density at the metal center and the surrounding ligand field, both controlling the metal-oxo-oxyl bond strength. The occupancy of the metal π/π* bonding/antibonding states determine the nature of the oxo-oxyl-sites. In the case of empty d orbitals, the metal is able to accept the px and py electron pairs of the oxo-oxyl ligand forming the bond and making the atomic ensemble extremely electrophilic. The metal-oxide center can therefore act as a Lewis acid site and become susceptible to nucleophilic attack from adsorbed water. A situation with high hole density on Ir would therefore favor the acid-base mechanism. On the other hand, with increasing of the d orbitals’ occupancy, the metal center increasingly acts as an electron donor, resulting in a lower formal Ir-O bond order. The most active amorphous catalyst in this study also exhibited the highest hole density on Ir, as shown by the Ir L3-edge measurements and the cyclic voltammetry. This suggests that the most suitable mechanism describing the intermediate step of O−O bond formation preceding its evolution into molecular oxygen is the nucleophilic attack of water molecules on the metal-oxo-oxyl sites due to the existence in both O 2p and Ir 5d and not so likely only in Ir 5d (oxo bond) or O 2p (oxyl bond) . Thus, the holes on Ir 5d drive the formation of electrochemical active O^{−}\text{•} which was identified as a key factor for catalytic reactivity of iridium oxides, as well as for the reaction path of the OER. Additional hole character on Ir 5d lowers the OER overpotential due to an interaction with the O 2p orbitals, which finally enables the formation of a metal-oxo-oxyl intermediate O^{−}\text{•} species active in OER.

Consequently, the OER on an IrOx catalyst can be tentatively described by its analogy with the oxo-bridge in the photosystem II, where the oxidation of water to molecular oxygen involves the formation of an oxo-oxyl species [49,50]. This active site undergoes nucleophilic attack by water and forms an O−O bond (as shown in Fig. 5B). Similarly, the catalytic oxidation of water on IrOx, happens via an electron deficient oxyl-complex, which is the active species for the formation of the O−O bond as indicated by the formation of shared holes between Ir 5d and O 2p. The oxidation of the resulting hydroperoxy \[\text{[IrO}_2\text{]}^{(6+1)\text{−}}\text{−O−O} \text{−OH}\] intermediate yields the final product dioxygen, and the catalyst recovers by coordination with H\textsubscript{2}O. We postulate that the overall reaction key is the formation of a hydroperoxy bond, through the nucleophilic attack of an oxyl ligand by water or OH. The proposed mechanism is illustrated in Fig. 5B where one cycle fulfills the overall OER reaction formula $\text{IrO}_x + \text{H}_2\text{O}\rightarrow\text{IrO}_x + \text{O}_2 + 4\text{H}^+ + 4\text{e}^−$, with the exact iridium oxidation state remaining as major subject of further investigations requiring accurate preparation of fingerprint samples. The iridium-oxo-oxyl (namely before O^{−}\text{•} [29]) species appear to be the key intermediate, which can activate oxygen, with its reactivity determined by the metal ligand interaction. Higher potentials form holes in the Ir 5d level and the hybridization with the O 2p orbitals leads to an electron deficient oxygen species with electron-holes shared between O and Ir, which is then active in OER due to its double character in the formation of metal-oxo-oxyl and nucleophilic O^{−}\text{•} species yielding hydroperoxy which further oxidize to form dioxygen.

4. Conclusions

In this work, the electronic structure of a free standing graphene electrode decorated with Ir NPs for OER in acidic environment was conducted. In situ X-ray spectroscopy revealed that the OER on the surface of the IrOx NPs is similar to the oxo-oxyl-bridge formation in the photosystem II. The reaction is driven by the formation of shared electron-holes in the O 2p and Ir 5d which activate oxygen by making it electron-deficient. Thus, the oxidation of water into dioxygen involves the formation of an oxo-oxyl-metal-complex and the oxygen in this complex is susceptible to nucleophilic attack by water or OH, which results in hydroperoxy bond formation. The hydroperoxy species is then finally transformed into molecular dioxygen. The proposed reaction pathway was developed around the formation of the O−O bond as the key step. Consequently, the formation of shared holes in the Ir 5d and O 2p is a catalytic activity descriptor and can be used to synthesize more active materials for the OER rather than absorption energy.
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Supplementary materials

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