Pyrochlore iridates (Na,Ca)$_2$Ir$_2$O$_6$·H$_2$O are acid-stable electrocatalysts that are candidates for use in electrolyzers and fuel cells. Ir L$_{III}$-edge X-ray absorption fine structure spectroscopy in 1 M H$_2$SO$_4$ at oxygen evolution conditions suggests the involvement of the electrons from the conduction band of the metallic particles, rather than just surface iridium reacting.

Acid-stable electrocatalysts are sought after for application in polymer electrolyte membrane (PEM) devices, such as electrolyzers for water splitting$^1$ and in fuel cells where they can provide a buffer to counter the corrosion of the carbon support at extremes of potential.$^2$ The acid electrolyte in these PEM devices provides advantages over alkali systems, with high charge density and no detrimental effects of carbonate contamination.$^3$ The dioxides of ruthenium and of iridium, both with the rutile structure, have been proven to be both active and robust catalysts for the oxygen evolution reaction (OER) in these situations.$^4$ The combination of the two precious metals in a ternary mixed oxide has been used to temper the high reactivity of ruthenium, which can be dissolved under operating conditions.$^5$ That work, carried out over a rather limited potential range short of OER conditions, proposed a scenario by which the iridium atoms respond by a two-site reaction, where two types of active sites, which have distinct local structure and electrochemical response, have distinct redox potentials. For (Na,Ce)$_2$Ru$_2$O$_7$, we used in situ XAFS to monitor change of Ir and Ru oxidation state upon application of potential into the OER regime and used the X-ray absorption near edge structure (XANES) to track metal oxidation state, revealing a cooperative response of the two metals under electrocatalytic conditions.$^{15}$ In this communication we report application of this methodology to the pyrochlore system (Na,Ca)$_2$Ir$_2$O$_6$·H$_2$O$^{18}$ and consider the extended X-ray absorption fine structure (EXAFS) to quantify local atomic structure in situ with electrocatalysis at potentials associated with OER. We were interested to study a pure iridate that may offer long-term stability, since the loss of ruthenium remains an issue even in these multinary oxides. In addition, with only a single precious metal to consider we aimed to obtain definitive structural information as a function of applied potential.

The pyrochlore material (Na,Ca)$_2$Ir$_2$O$_6$·H$_2$O was prepared by a hydrothermal crystallisation directly from CaO$_2$, Na$_2$O$_2$.
and IrCl₃·5H₂O in 10 M NaOH solution at 240 °C (ESI†), and was characterised using powder X-ray diffraction (XRD) and scanning transmission electron microscopy (STEM), Fig. 1. Scherrer analysis of the diffraction profile gave an average crystallite domain size of 36.2 ± 2.4 nm, consistent with the STEM imaging, while lowering the synthesis temperature to 170 °C gave a second sample that consisted of smaller crystallite sizes (11.2 ± 1.9 nm by Scherrer analysis, ESI†). Surface areas, measured by nitrogen adsorption isotherms and the BET method for the two samples were 7.2 m² g⁻¹ and 62.7 m² g⁻¹, respectively, consistent with the relative dimensions of crystallites assuming no significant agglomeration (as confirmed by STEM images). The Rietveld method was used to refine chemical composition and this gave empirical formula (Ca₀.70Na₀.24)₂Ir₂O₆·H₂O, for the most crystalline sample with the crystal water content verified using thermogravimetric analysis (ESI†). The Rietveld fit to powder XRD of (Ca₀.70Na₀.24)₂Ir₂O₆·H₂O Fd₃m, a = 10.23978(5) Å, and (b and c) STEM images and EDXA maps of two regions of the material showing homogeneity of elemental distribution.

Prior to in situ EXAFS measurements, ex situ XANES spectra were recorded at the Ir L₃-edge to determine the average oxidation state of Ir in the pyrochlore, by comparison to the reference materials IrCl₃, IrO₂ and BaNa₀.₅Ir₀.₅O₃·d, that contain iridium in oxidation state +3, +4 and 4.9, respectively, Fig. 2. This shows the iridium to be in average oxidation state of +4.5, consistent with the results from crystallography and also earlier work on a material with similar composition but prepared by a different method.¹⁸ The XANES measurements made in situ were similarly analysed (see below) and an important, immediate conclusion from these data is that there is negligible loss of iridium into solution with applied potential in 1 M H₂SO₄ solution, since the white line intensity of the raw data was not seen to diminish during the course of the experiment (see ESI†).

The in situ EXAFS data were analysed to obtain quantitative information about the Ir local environment. A single-shell fit was performed to focus on the Ir–O contribution to the spectrum, and typical fits to the spectra, and their associated Fourier transforms are shown in Fig. 3 for data measured prior to application of potential and at the highest potential, 1.78 V vs. standard hydrogen electrode (SHE). Fitted EXAFS parameters from these spectra are presented in ESI†. For the highly crystalline sample of the pyrochlore, EXAFS spectra were successfully recorded and analysed in a sequence of applied voltage steps, up to 1.78 V, and then upon reversal of potential to a lower than initial starting value. Fig. 4 summarises the structural information obtained from fitting the spectra and full details are provided in the ESI†. These results highlight the...
gradual shortening of Ir–O distance up to 1.48 V, the onset of OER, reaching constant value during OER and then lengthening upon reversal of potential, Fig. 4a. A shorter average Ir–O distance implies oxidation of the iridium. Meanwhile, the Debye–Waller factor essentially shows no change in magnitude over the whole process, Fig. 4b. The iridium oxidation state can be quantified by the shift in edge position, as shown in Fig. 4c, and this measurement was carried out in a separate experiment on materials of two different particle sizes. We can thus conclude that the iridium, on average, increases in oxidation state by 0.5 units under OER conditions and that this is independent of the particle size (and surface area) of the material, despite the greater proportion of available iridium expected for the higher surface area sample.

We also compared the oxidation state shift from XANES with that derived from the Ir–O bond distance using the bond valence sum method, and this showed essentially the same behaviour with applied potential, thus giving independent verification of the changes in average oxidation state seen in situ (ESI†).

To understand the changing local structure of iridium in the pyrochlore during operation as an electrocatalyst we consider the models proposed by Hillman et al. in their study of iridium oxide films. In that work it was observed that the Debye–Waller factor of the Ir–O shell showed an increase upon iridium oxidation (albeit below potentials needed to reach OER conditions), which was reconciled as due to a two-site bond model for redox, in which two iridium species responded differently to applied potential to give a greater static disorder contribution to the Debye–Waller factor. An alternative model is one in which all iridium atoms in the sample respond simultaneously to the applied potential such that the Ir–O bond distance of all are shortened by the same amount with no increase in static disorder. This can be explained by a single-site band model whereby electrons are removed not just from surface sites, but from the conduction band of the metallic particles. The observed behaviour of the pyrochlore is entirely consistent with this idea. Independent verification of this model comes from study of the sample with higher surface area, smaller crystallite...
domain size, which shows no greater extent of iridium oxidation, despite the ten-fold increase in specific surface area. Thus we propose that the iridate pyrochlores show redox behaviour different to the previously studied iridium oxide films: this could be due to the different local structure of the materials since the films were structurally disordered, being hydrated and low density forms of iridium oxide,\textsuperscript{17} rather than the well-defined, crystalline particles that we have studied.

Conclusions

We have shown that \textit{in situ} XAFS (XANES and EXAFS) can be recorded from electrocatalysts operating under realistic conditions to extract quantitative structural information that can be used to probe mechanism of their operation. This method could be applied to other families of oxide materials currently emerging as acid-stable catalysts in important contemporary energy-related applications. For the pyrochlore studied here, we have shown how the bulk metallic character of the oxide particles plays an important role in their mode of operation, with electrons from the conduction band of the oxide being extracted to bring about OER. Complementary cyclic voltammetry (see Fig. S3, ESi\textsuperscript{†}) shows only one redox feature in the potential window of the \textit{in situ} XAFS experiment that would correspond to oxidation/reduction or Ir\textsuperscript{4+}/Ir\textsuperscript{5+}. The surface reactivity must clearly also hold the key to catalysts properties since this is where the water interacts with the catalyst surface, and where the splitting of water and release of oxygen takes place, but the transport of charge from the bulk particle is an important aspect of the overall mechanism. In multinary oxides, such as the pyrochlore studied here, the role of the partner non-precious-metal cations may also be important in providing charge balance at electrocatalysis conditions, which may explain their different mode of operation compared to binary precious-metal oxides. More mechanistic studies on other members of the pyrochlore family would give a greater insight into this idea. Further work is also needed to understand the long-term stability of the pyrochlores under operating conditions, and their application in real devices, which will be topic of forthcoming publications.

Conflicts of interest

There are no conflicts to declare.

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