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EDGE ARTICLE

Anodic deposition of a robust iridium-based water-oxidation catalyst from organometallic precursors†

James D. Blakemore,^a Nathan D. Schley,^a Gerard W. Olack,^b Christopher D. Incarvito,^a Gary W. Brudvig^{*a} and Robert H. Crabtree^{*a}

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Artificial photosynthesis, modeled on natural light-driven oxidation of water in Photosystem II, holds promise as a sustainable source of reducing equivalents for producing fuels. Few robust water-oxidation catalysts capable of mediating this difficult four-electron, four-proton reaction have yet been described. We report a new method for generating an amorphous electrodeposited material, principally consisting of iridium and oxygen, which is a robust and long-lived catalyst for water oxidation, when driven electrochemically. The catalyst material is generated by a simple anodic deposition from Cp*Ir aqua or hydroxo complexes in aqueous solution. This work suggests that organometallic precursors may be useful in electrodeposition of inorganic heterogeneous catalysts.

Introduction

Water-oxidation catalysts are envisioned to play a key role in many proposed schemes for artificial photosynthesis and solar-fuel production.¹ In these proposals, photoelectrochemical cells harvest and store solar energy in chemical form. Although details vary, the strategy typically proposed is the oxidation of water at an anode to produce electrons that in turn reduce H₂O or CO₂ to fuels at the cell cathode.

The high kinetic barrier for oxygen evolution is a key hurdle in each of these proposed systems, since it limits cell power efficiency by introducing a large overpotential.² Additionally, the required oxidizing equivalents must be properly coupled to yield dioxygen, rather than partially oxidized side products (*e.g.*, HO[•], H₂O₂).³ These difficulties hamper the development of water-oxidation catalysts, especially catalysts that can operate under the required harsh oxidizing conditions in a sustained fashion with minimal deactivation.

The dependence of overpotential for oxygen evolution on the nature of the electrode material has been recognized for many years.⁴ Noble metal oxides (*e.g.*, ruthenium, iridium) were observed early to have good catalytic properties for water oxidation.⁵ Among these, iridium materials have proven to be among the most active and stable for oxygen evolution.⁶ The

advent of Dimensionally Stable Anodes® (DSA) in the 1960s introduced iridium oxides as anodes in industrial applications requiring high activity and longevity.⁷ These electrodes are composed of an oxidation resistant titanium/titanium oxide substrate which is coated with an active catalyst layer. This active layer often includes several metal oxides (*e.g.*, tantalum, titanium) in addition to iridium, in order to disperse the active catalyst material, provide high surface area, and reduce corrosion.⁸ However, the catalyst for the oxygen evolution reaction is the iridium material.

Iridium oxides for use as electrode materials have historically been prepared in three ways. In the first, a solution containing iridium chlorides is applied to the electrode surface and then thermally decomposed, giving a nanoparticulate material which is an active catalyst.⁹ In the second method, thermal or chemical decomposition of iridium salts under controlled solution conditions gives rise to free nanoparticles, which can be physically or electrochemically deposited on the electrode surface.^{10,11} Chemical preparations of this second type are used in proton-exchange membrane (PEM) water electrolyzers.¹² Sputtered iridium dioxide has also been shown to be highly active for oxygen evolution.¹³ Additionally, various types of iridium oxide films have been previously shown to be electrochromic, with the most active prepared by the sputtering technique.¹⁴

In more recent work, we have shown that a number of organometallic pentamethylcyclopentadienyl (Cp*) iridium

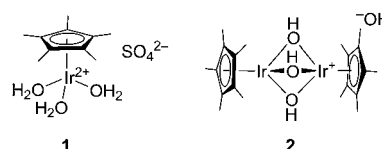


Fig. 1 Structures of iridium(III) complexes **1** and **2**.

^aYale University, Department of Chemistry, PO Box 208107, New Haven, Connecticut, 06520-8107, USA. E-mail: robert.crabtree@yale.edu; gary.brudvig@yale.edu; Fax: +1 203 432-6144; +1 203 432-6144; Tel: +1 203 432-3925; +1 203 432-5202

^bYale University, Department of Geology and Geophysics, PO Box 208109, New Haven, Connecticut, 06520-8109, USA

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complexes are robust and highly active homogeneous catalysts for water oxidation when driven with cerium(IV) as primary oxidant.¹⁵ Complexes **1** and **2** (Fig. 1),¹⁶ having multiple aqua or hydroxo ligands show the fastest rates of oxygen evolution, up to 20–25 turnovers min^{-1} even at pH 0.89 with cerium(IV).¹⁷ These values compare well with recently published work on cobalt catalysis.¹⁸ Kinetic studies, reported elsewhere,¹⁷ show that the order of reaction in iridium for complexes **1** and **2** is greater than one, suggesting dimerization or oligomerization is key to forming the catalyst of highest activity.

We now report that electrochemical oxidation of **1** or **2** induces deposition of a catalytic blue layer of an iridium oxide material (**BL**) capable of catalyzing water oxidation at the anode. Holding the oxidizing potential steady for several minutes is sufficient to form a visible amount of **BL**. Optimal activity is reached after deposition for several hours (see ESI†). **BL** deposits on carbon, platinum, gold, fluorine-doped tin oxide (FTO), and indium tin oxide electrodes. We know of no prior case in which an active material for water oxidation has been deposited from a well-defined organometallic precursor having a metal–carbon bond, although inorganic salts and molecular coordination complexes have been used previously for several metals including cobalt and nickel.¹⁹ Importantly, this route is direct and simple, avoiding harsh conditions and specialized equipment. Furthermore, the deposited **BL** displays interesting electrochromic properties, similar to classical sputtered iridium oxides.

Results and discussion

A cyclic voltammogram of a solution containing **1** (Fig. 2, inset) shows only a simple catalytic wave on the first anodic scan with onset at 1.15 V (all potentials are vs. NHE). On multiple cycling, the **BL** builds up, accompanied by a feature at 0.88 V that grows

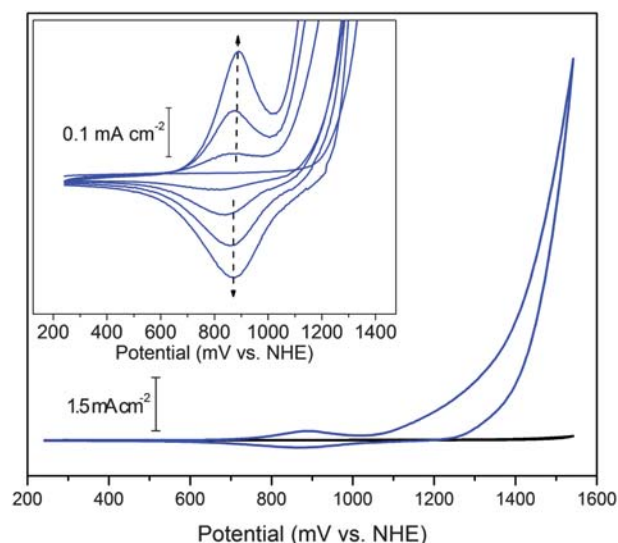


Fig. 2 Cyclic voltammetry of **BL** (blue line) and basal plane graphite electrode background (black line) in 0.1 M KNO_3 at pH 6. The catalytic wave is seen above ca. 1.1 V. Inset: cyclic voltammogram (successive scans) showing deposition of **BL** from a solution containing **1**; reversible peak centered at 0.88 V ($\Delta E_p = 15$ mV). Deposition conditions: 2.3 mM **1**, 0.1 M KNO_3 , pH 2.9; scan rate: 50 mV s^{-1} .

in intensity. As more **BL** deposits, the catalytic wave at 1.1 V also becomes more intense.

Upon rinsing the anode and transferring it to a solution of pure potassium nitrate electrolyte, the reversible feature and catalytic wave from the **BL** are still present at full intensity. The peak separation, ΔE_p , for the reversible feature is ca. 15 mV, consistent with a non-diffusional, surface-bound species (ideally, $\Delta E_p = 0$). Peak currents for the reversible feature show a linear dependence with scan rate, as expected for a surface-bound species (see ESI†).²⁰ Thus, the **BL** is deposited at the electrode surface and is responsible for the catalytic reaction.†

Upon deposition, **BL** functions as a robust and highly active catalyst for water oxidation. Indeed, bubbles of O_2 are rapidly formed at the electrode surface during cyclic voltammetry. Oxygen was confirmed as the product of the catalytic oxidation wave by headspace gas analysis using a fluorescence-probe assay (Fig. 3). An FTO slide, previously prepared with **BL**, was introduced into a closed two-chamber electrochemical cell with 0.1 M KNO_3 (pH 7), and 1.4 V was applied for 1 h. During this time, sufficient charge was passed through the cell to generate 28.9 μmol of dioxygen. 27.7 μmol was detected by the headspace oxygen assay, representing a Faradaic efficiency of 96%. In Fig. 3, it is clear from the parallel behavior of the two signals that nearly all of the oxidizing equivalents contribute to oxygen evolution, rather than to detrimental side reactions. Water was confirmed as the source of oxygen atoms by H_2^{18}O isotope incorporation to give the expected ^{18}O enrichment in product O_2 (see ESI†).

BL is resistant to corrosion and to loss of catalyst during operation. Long time course studies indicate that activity is retained over many hours of continuous use. Specifically, after 9 h of constant electrolysis at an applied potential of 1.4 V (sustained current density: 1.4 mA cm^{-2} , without cell resistance (iR) compensation), the catalyst maintains over 95% of its initial activity (see ESI†).

BL also exhibits electrochromism. If the electrode is polarized at a potential below the reversible wave centered at 0.88 V, the

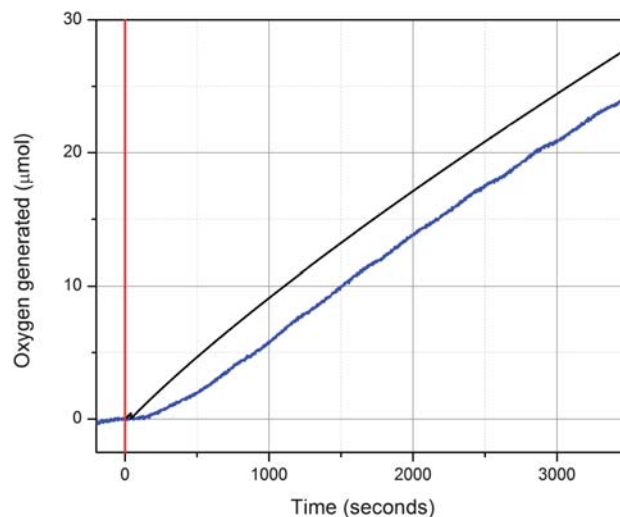


Fig. 3 Oxygen evolution catalyzed by **BL**. Black line: predicted oxygen yield based on current transferred. Blue line: oxygen yield detected by headspace analysis.

film becomes largely translucent, turning a very pale shade of yellow–green. Not surprisingly, polarizing the electrode above the reversible wave causes the striking blue color and opacity to return. This color change also occurs on treatment of **BL** with chemical reductants and oxidants, consistent with an oxidation state change from predominantly iridium(III) to predominantly iridium(IV), similar to results on iridium films prepared by other methods.²¹

The reversible wave at 0.88 V can be used to estimate the number of surface-bound electroactive iridium atoms. In previously studied iridium oxide films, the reversible wave was anticipated to be a one-electron process (iridium(III) to iridium(IV)). However, careful studies by Burke and Whelan led to the observation that the reversible wave is actually a two-electron, three-proton process.²² Based on the similarity of **BL** in voltammetric and catalytic response to the previously described hydrous iridium oxide films, the assumption of a two-electron, three-proton process is used here. After 10 or 20 cycles from 0.3 to 1.5 V at 100 mV s⁻¹, the surface coverage of iridium was estimated to be 4.1 and 8.2 nmol cm⁻², respectively. The former can be estimated at a coverage of less than ten monolayers, in analogy with similar work previously carried out with Au(111) monolayers.²⁰

The turnover frequency on a per-electroactive iridium basis can be estimated with the surface coverage data. At the coverages listed above, the turnover frequency at 1.4 V was estimated to be 1.46 turnovers s⁻¹ for the 4.1 nmol cm⁻² **BL** and 1.11 turnovers s⁻¹ for the 8.2 nmol cm⁻² **BL** (platinum disc rotating disc electrode, $\omega = 1000$ rpm). This is lower than the turnover values reported by Murray, *et al.*, for small iridium dioxide nanoparticles at 1.5 V.^{11b} Furthermore, some surface-bound iridium atoms may not be electroactive, complicating this analysis. In a highly active film which has been deposited for 2 h, approximately 0.25 mg cm⁻² is present on the electrode surface; however, at these loadings, the features in cyclic voltammetry are broadened so significantly that no meaningful analysis can be accomplished. Films of this type have a thickness of between 1 and 2 μm , as estimated by SEM. (see ESI†).

In general, the discrepancy between measured activity and the apparent robustness of catalysis may be explained by a non-Nernstian response from the **BL**. The full width at half maximum measured for the reversible feature at 0.88 V is consistently much larger than predicted by electrochemical theory, even at low catalyst loadings (*vide supra*).¹⁵ Since this is the case, the turnover frequency on a total iridium basis cannot be confidently estimated.

The high activity of the **BL** presumably results from its high surface area. Indeed SEM images show nanoscale features, consistent with this hypothesis (Fig. 4). However, the deposited material is amorphous by X-ray diffraction (only broad, low-angle diffraction is seen; see ESI†).

Elemental analysis on **BL** (after depositing at 1.4 V for 2 h) shows 9% carbon by mass, along with hydrogen, oxygen and iridium. There is negligible nitrogen content.¶ The results are consistent with energy-dispersive X-ray spectroscopy (EDS) measurements, which show the presence of only carbon, oxygen and iridium above background after deposition (see ESI†). **BL** samples deposited from solutions of **1** do not show sulfur incorporation by EDS. As Cp* is the only source of carbon, we

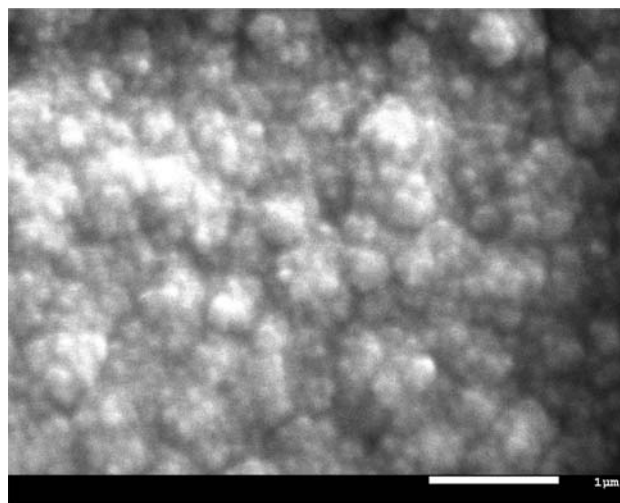


Fig. 4 Top-view SEM image of **BL**. Catalyst was deposited on ITO-coated glass substrate.

conclude that a substoichiometric fraction of the Cp* ligand, or a derivative, is incorporated into **BL**. The Cp* ligand is necessary for deposition, as solutions of iridium trichloride hydrate in potassium nitrate do not form deposits or show catalytic oxidation waves in cyclic voltammetry, nor do less coordinatively unsaturated Cp*Ir complexes such as Cp*Ir(κ^2 -2-phenylpyridine)Cl or Cp*Ir(2,2'-dipyridyl)Cl₂. Oxidation of Cp* methyl groups has been observed previously on treatment with iodine(III) reagents.²³ Similar oxidation could contribute to the deposition process for **BL**, but complexes **1** and **2** are the ones that form a deposit, in contrast to the other half-sandwich iridium compounds we have studied.

IR spectra collected for **BL** also show clear indications of carbon-containing fragments (see ESI†). There are broad, strong stretching frequencies in the region of 1300–1700 cm⁻¹, assigned

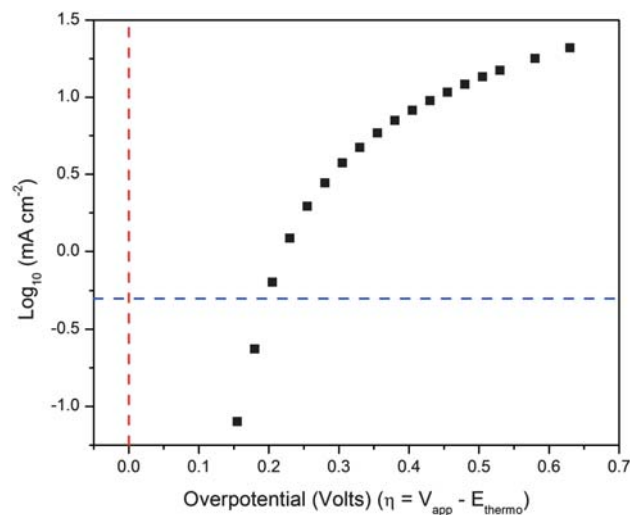


Fig. 5 Tafel plot of catalytic currents. Blue dashed line corresponds to 0.5 mA cm⁻²; red dashed line is at zero overpotential. Conditions: 0.1 M KNO₃, pH 2.9, platinum rotating disc electrode.

to $\nu(\text{C}=\text{O})$ and $\nu(\text{C}-\text{O})$ features, which would arise upon oxidative degradation the Cp^* ligand. However, there are no detectable $\nu(\text{C}-\text{H})$ frequencies, indicating that essentially all of the Cp^* methyl groups have been lost upon formation of **BL**.

We next examined the overpotential for oxygen evolution. Bubble formation and large local pH effects, possible during electrolysis, seem to complicate measurements; therefore, employing a rotating-disc electrode allows us to conduct the experiment under controlled conditions. However, because the Tafel plot (Fig. 5) is not linear, the overpotential cannot be assessed here with complete reliability. Nonetheless, the data show the overpotential required to reach 0.5 mA cm^{-2} is $200 \pm 3 \text{ mV}$. In fact, due to the highly capacitive nature of hydrated noble metal oxides, this overpotential will likely be underestimated (see ESI†). Comparison with very long timecourse data collected without a rotating disc electrode (9 h data from above) at 1.4 mA cm^{-2} shows a near 100 mV higher overpotential, although the rotating disc experiments do not suffer from bubble formation and consequent loss of surface area. Generally, these experiments show there is a low overpotential ($\leq 300 \text{ mV}$) for water oxidation with **BL**, demonstrating the dramatic lowering of the kinetic barrier to $\text{O}=\text{O}$ bond formation achieved by catalysis with **BL**.

Discussion

Iridium oxide materials are known to be highly active catalysts for water oxidation.^{5,6} With this in mind, it is not surprising that formation of **BL** on the electrode surface coincides with the appearance of a catalytic response leading to water oxidation. There are striking similarities in electrocatalytic behavior (*i.e.*, low overpotential) and voltammetric response (*i.e.*, appearance of cyclic voltammograms, electrochromism) between **BL** and other preparations of iridium oxides.²⁴ This strongly suggests that inorganic iridium dioxide is the active catalyst in this case, especially considering the strong effect of preparation method on heterogeneous catalyst performance. However, unlike many other preparations of iridium oxide, **BL** is prepared by an exceptionally mild method. Preparation of the catalyst from organometallic precursors provides a route to explore new solid-state and inorganic catalytic materials for water oxidation. Further, as iridium oxides are one of the most active and industrially-relevant materials available today for water oxidation chemistry, this novel method of preparation will help in elucidating their mechanism of action and perhaps lead to significant improvements in catalyst performance. Specifically, since **BL** is prepared by a mild and facile method, it may be of use in assemblies for achieving light-driven water oxidation.²⁵

Conclusions

In conclusion, we report the iridium oxide **BL** is a highly active and robust electrocatalyst for water oxidation. It is prepared by electrodeposition from aqueous solutions of organometallic Cp^*Ir precursors, allowing careful control of catalyst loading and preparation conditions. The activity of **BL** suggests that it is a promising platform for investigating catalytic water oxidation at electrode surfaces. Furthermore, the method of preparing **BL** from an organometallic precursor suggests that a new, diverse

range of heterogeneous catalysts may be prepared from solutions of organometallic precursors, perhaps eventually with abundant first-row metals.

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Notes and references

- ‡ Turnovers are defined as: $\text{mol O}_2 (\text{mol Ir})^{-1}$.
- § Under the deposition conditions, the local pH at the electrode surface was anticipated to be lower than in the bulk solution, due to proton release during vigorous water oxidation. This would cause **2** to protonate, essentially forming **1** *in situ*. Investigations are now underway to elucidate differences between **BL** deposited from **1** vs. **2**.
- ¶ Elemental analysis results for C, N, H, O and Ir sum to only 74%. This is likely due to a low recovery during analysis, a common occurrence with metal oxide materials. Analysis performed by Columbia Analytical Services, Inc., 3860 S. Palo Verde Rd. Ste. 302, Tucson, Arizona 85714.
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