Open Archive Toulouse Archive Ouverte (OATAO)

OATAO is an open access repository that collects the work of Toulouse researchers and makes it freely available over the web where possible.

This is an author's version published in: http://oatao.univ-toulouse.fr/21775

Official URL: https://doi.org/10.1002/aenm.201702855

To cite this version:


Any correspondence concerning this service should be sent to the repository administrator: tech-oatao@listes-diff.inp-toulouse.fr
Proton Ion Exchange Reaction in Li₃IrO₄: A Way to New H₃₊ₓIrO₄ Phases Electrochemically Active in Both Aqueous and Nonaqueous Electrolytes

Arnaud J. Perez, Robin Beer, Zifeng Lin, Elodie Salager, Pierre-Louis Taberna, Artem M. Abakumov, Patrice Simon, and Jean-Marie Tarascon*

Progress over the past decade in Li-insertion compounds has led to a new class of Li-rich layered oxide electrodes cumulating both cationic and anionic redox processes. Pertaining to this new class of materials are the Li/Na iridate phases, which present a rich crystal chemistry. This work reports on a new protonic iridate phase H₃₊ₓIrO₄ having a layered structure obtained by room temperature acid-leaching of LiIrO₄. This new phase shows reversible charge storage properties of 1.5 e⁻/Ir atom with high rate capabilities in both nonaqueous (vs Li⁺/Li) and aqueous (vs capacitive carbon) media. It is demonstrated that Li-insertion in carbonate LiPF₆-based electrolyte occurs through a classical reduction process (Ir⁵⁺ ↔ Ir³⁺), which is accompanied by a well-defined structural transition. In concentrated H₂SO₄ electrolyte, this work provides evidence that the overall capacity of 1.7 H⁺/Ir results from two additive redox processes with the low potential one showing ohmic limitations. Altogether, the room temperature protonation approach, which can be generalized to various Li-rich phases containing either 3d, 4d or 5d metals, offers great opportunities for the judicious design of attractive electrode materials.

1. Introduction

The field of Li-ion batteries is going through its own paradigm shift with the emergence of the anionic redox process, whereas so far Li-ion batteries were solely relying on cationic redox processes to store energy. The advantages of cumulating cationic and anionic redox processes are exemplified by the numerous studies on the layered Li₁ₓNi₀.₁₂₃M₀.₃₄Sn₀.₂₆O₂, denoted Li-rich NMC, which exhibits capacity as high as 290 mA h g⁻¹, to be compared to the 200 mA h g⁻¹ of the presently used LiNi₀.₆Mn₀.₂Co₀.₂O₂ layered oxide, or NMC 622. Both phases have the same layered structure but solely differ by the presence of Li in the transition metal layers for the Li-rich NMC phases.

In order to understand the scientific background of the anionic redox process in the complex Li-rich NMC materials containing three competing redox centers, scientists have searched for simplified model compounds. Among them, the Li-based ruthenates and iridates Li₂M₀ₓ (M = Ir, Ru), which possess a layered structure and can be described as Li-rich oxides Li[(Li₁ₓM₁₋ₓ)Irₓ]O₂, show a reversible capacity of 270 mA h g⁻¹ and present a less complex redox chemistry because of a sole redox center M. The synthesis of these phases is well documented as they have been the subject of intense fundamental studies on magnetism because of complex spin–orbit coupling effects.

Although these model Ru/Ir-based compounds have limited practical interest for the next generation of Li-ion batteries because of the poor abundance of both Ru and Ir, they have strongly contributed to their fundamental understanding. Studies of the Li₂Ru₁₋ₓSnₓO₂ or Li₂IrₓO₄ phases were, for instance, essential to demonstrate that the extra capacity in Li-rich NMC materials was due to anionic redox,[1] and that the origin of voltage fade in Li-rich electrodes was due to the

Dr. A. J. Perez, R. Beer, Prof. J.-M. Tarascon
Chimie du Solide et Energie – UMR 8260 – Collège de France
11 place Marcelin Berthelot, 75005 Paris, France
E-mail: jean-marie.tarascon@college-de-france.fr
Dr. A. J. Perez, Prof. J.-M. Tarascon
Sorbonne Université – UPMC Univ Paris 06
4 Place Jussieu, 75005 Paris, France
Dr. A. J. Perez, R. Beer, Dr. Z. Lin, Dr. E. Salager, Dr. P.-L. Taberna, Prof. P. Simon, Prof. J.-M. Tarascon
Réseau sur le Stockage Electrochimique de l’Energie (RS2E)
FR CNRS 3459, 33 rue Saint Leu, 80039 Amiens, France

The ORCID identification number(s) for the author(s) of this article can be found under https://doi.org/10.1002/aenm.201702855.

DOI: 10.1002/aenm.201702855
capturing of small cations in tetrahedral sites.\textsuperscript{[4]} Very recently, we expanded this family of model compounds by reporting on a new member Li\textsubscript{1}IrO\textsubscript{4}, which enables to push the limit of insertion reactions to a new record of 3.5 \textit{e}\textsuperscript{−} per transition metal. Aside the field of batteries, it is worth mentioning that both Ru/Ir-based oxides have been heavily studied by electrochemists. First, as highly performant water splitting catalysts, showing a low overpotential and high activity for the oxygen evolution reaction (OER).\textsuperscript{[5,6]} Following the first studies on the formation of an oxide layer at the surface of metallic Ru/Ir electrodes,\textsuperscript{[7,8]} it was proposed that the reversible redox processes happening in this oxide layer could be used to store energy.\textsuperscript{[9,10]} leading to the development of pseudocapacitors, with IrO\textsubscript{2} or RuO\textsubscript{2}•H\textsubscript{2}O being among the stellar materials with capacitances exceeding 800 F g\textsuperscript{−1}.\textsuperscript{[11]} Most of the developments in these fields were done using amorphous, hydrated\textsuperscript{[12]} or high surface area iridium oxides (nanoparticles\textsuperscript{[13]} or nanosheets\textsuperscript{[14]} in order to maximize the shell contribution while often overlooking the bulk contribution to energy storage.

In light of the versatility offered by these Ru(Ir)-based compounds to undergo redox chemical reactions in both aqueous or organic solvents, it is becoming tempting to further extend the rich crystal chemistry of the Li-based iridate phases to proton-based phases and exploit their electrochemistry. Herein we demonstrate, by focusing on Li\textsubscript{1}IrO\textsubscript{4}, the feasibility to prepare a new protonated H\textsubscript{3+}IrO\textsubscript{4} phase by room temperature ion exchange of Li for protons in aqueous media. This protonated phase is shown to present a spectacular reversible electrochemical activity in both nonaqueous media versus metallic Li and in aqueous media versus activated carbon.

### 2. Results

Single phase Li\textsubscript{1}IrO\textsubscript{4} powder was prepared by a previously described ceramic process, which consists in heating a stoichiometric mixture of Ir and Li\textsubscript{2}CO\textsubscript{3} in air at 950 °C for 24 h.\textsuperscript{[15]} The resulting powder was stirred in a 0.1 M H\textsubscript{2}SO\textsubscript{4} solution and, after 1 h of stirring, the powder was recovered by centrifugation and washed with distilled water until its characterization. Comparative scanning electron microscopy (SEM) pictures of the pristine and recovered powders (Figure 1a, inset) indicate the break-up of the initial 10–20 µm aggregates but no impact of the acid treatment on the 1–2 µm size primary particles. The powder composition was indirectly obtained via analysis of the reaction supernatant by complementary inductively coupled plasma mass spectrometry (ICP-MS) for the amount of released lithium, and by acid–base titration to determine the amount of remaining protons in solution. By cross-cutting the results of both measurements we could deduce the composition H\textsubscript{3+}Li\textsubscript{1}IrO\textsubscript{4} for the recovered sample within the accuracy of the analysis. The completion of the exchange reaction was confirmed by the absence of \textsuperscript{7}Li signal in the solid-state nuclear magnetic resonance (NMR) spectrum (Figure S1a, Supporting Information), indicative of the absence of Li in the exchanged sample.

The X-ray powder diffraction pattern of the Li-free sample shows sharp Bragg peaks characteristic of a well crystallized material (Figure 1b). All the peaks could be indexed in a hexagonal \textit{R}$\overline{3}$\textit{m} unit cell, as for the parent phase, with however a main difference in the stacking sequence which is P3 for the H\textsubscript{3+}IrO\textsubscript{4} phase as opposed to O3 for the precursor Li\textsubscript{1}IrO\textsubscript{4} phase (Figure 1c,d). The \textit{a} lattice parameter slightly increases from 2.96 Å for Li\textsubscript{1}IrO\textsubscript{4} to 3.04 Å for H\textsubscript{3+}IrO\textsubscript{4}, while the \textit{c} parameter shows a greater decrease from 14.40 to 14.07 Å as expected since H\textsuperscript{+} is smaller than Li\textsuperscript{+}. Within the O3-Li\textsubscript{1}IrO\textsubscript{4} structure, or Li(Li\textsubscript{1}IrO\textsubscript{3})O\textsubscript{2}, Li\textsuperscript{+} sits in octahedral sites with 1/3 being located in the metal layers and the remaining 2/3 between the layers, but no precise clues can be obtained from the X-ray powder diffraction (XRD) powder pattern regarding the proton positions. To obtain information on the sites occupied by protons, deuterated D\textsubscript{3+}IrO\textsubscript{4} samples were prepared for NMR studies. \textsuperscript{2}H NMR measurements detect a single peak near 15 ppm with a width at half height of 200 ppm (Figure S1, Supporting Information), but this \textsuperscript{2}H peak is broad and without spinning sidebands compared to previous work by Paik et al. on deuterated Li\textsubscript{2}MnO\textsubscript{3}.\textsuperscript{[16]} The \textsuperscript{2}H environments are probably disordered with various distances and geometries relative to the paramagnetic Ir\textsuperscript{3+} centers, resulting in a distribution of Fermi contact shift. If some \textsuperscript{2}H atoms are close to Ir\textsuperscript{4+} centers, they are probably not detected due to the strong paramagnetic interaction with Ir\textsuperscript{4+}.

The thermal stability of the H\textsubscript{3+}IrO\textsubscript{4} phase was investigated by combined thermogravimetric analysis (TGA) and MS measurements by heating a sample under Ar to 1000 °C at a rate of 5 K min\textsuperscript{−1} while analyzing the released gases. The TGA profile for H\textsubscript{3+}IrO\textsubscript{4} (Figure 2a) shows a staircase behavior with three consecutive steps at 190, 700, and 900 °C, corresponding to the departure of H\textsubscript{2}O for the first step and of O\textsubscript{2} for the two remaining ones (Figure 2b), with the products obtained at 500 and 1000 °C being IrO\textsubscript{2} and Ir, respectively, as deduced from XRD of the recovered powders (Figure 2c). From the weight loss, and knowing the final products, we could deduce that the formula of the sample is H\textsubscript{1.4}IrO\textsubscript{4}. The H\textsuperscript{+} content exceeding three cannot be explained by a simple ion exchange reaction which does not involve a redox process. Due to the high redox potential of the H\textsubscript{3+}IrO\textsubscript{4} phase, as we will see later, the material oxidizes water hence leading to a reduced H\textsubscript{1.4}IrO\textsubscript{4} phase with \textit{x} being slightly pH dependent. To further characterize the protons within the sample and their existence as either OH groups or H\textsubscript{2}O, infrared (IR) spectroscopy measurements were done. Note the presence of bands associated to Li carbonate impurities in Li\textsubscript{1}IrO\textsubscript{4} that disappear after acid-leaching (Figure S2, Supporting Information), but to our surprise no bands appear around 1600 and 3600 cm\textsuperscript{−1}; wavelength domains over which the OH groups are IR-active. The absence of such bands could simply be nested in the metallic character of H\textsubscript{1.4}IrO\textsubscript{4} which prevents, for absorption reasons, the detection of OH group, as previously put forward to explain the inability to detect the OH bands in the metallic Li\textsubscript{1.3}H\textsubscript{2}O\textsubscript{2} phase.\textsuperscript{[17]} Having established the structural/physical properties of H\textsubscript{1.4}IrO\textsubscript{4}, we next probe its electrochemical activity versus Li\textsuperscript{+}/Li in nonaqueous media and versus capacitive carbon in aqueous electrolyte.

The electrochemical performances of H\textsubscript{1.4}IrO\textsubscript{4} were measured in organic electrolyte versus Li in Swagelok cells between 1.5 and 4.2 V at a C/10 rate. Remarkably, the cell can reversibly uptake 1.5 Li\textsuperscript{+} with limited polarization (Figure 3a) and shows the feasibility to deliver 75% of its energy density at a rate of 1 C (Figure 3e). Moreover, the voltage profile varies smoothly
with the removal/uptake of Li but also shows a narrow plateau at nearly 3.2 V, which can nicely be visualized on the derivative curve dQ/dV (Figure 3c) and is indicative of a phase transition.

To better understand Li-driven structural changes in (Li,H)$_x$IrO$_4$, in situ XRD measurements were conducted (Figure 4) and XRD patterns were collected with 0.05 step in the lithium stoichiometry. The collected patterns are shown as 2D contour plots in Figure 4a. As an Li$_x$Li$_{1-x}$IrO$_4$ cell is discharged, we observed initially a slight shift of position and then a drastic change in the intensity of the peaks near x = 0.6, with some peaks disappearing at the expense of new ones, which sharpen beyond x = 0.7 to give a well-defined XRD powder pattern that remained till the end of the insertion process at x = 1.5 (Li$_{1.5}$H$_{3-4}$IrO$_4$). Refined XRD patterns demonstrate a Li-driven structural transition, which is accompanied by a change in the stacking sequence from P3 to T1, with the corresponding variation of lattice parameters shown in Figure 4c. There is nearly a continuous increase of the lattice parameter upon reduction, which is consistent with the increase in the Ir ionic radii associated to its sequential Ir$^{5+}$ → Ir$^{4+}$ → Ir$^{3+}$ reduction. By contrast, the interlayer distance (d) remains nearly constant till the P3–T1 transition where it severely drops from 4.68 to 4.52 Å due to the migration of the Li$^+$ ions from prismatic to tetrahedral sites. Further lithiation up to x = 1.5 compensates this contraction with an increase of the interlayer distance d back to 4.71 Å. Such a Li-driven prismatic-tetrahedral

![Figure 1](image1.png)

Figure 1. Refinement of X-ray diffraction data and SEM picture of a) the parent Li$_3$IrO$_4$ and b) acid-leached H$_3$IrO$_4$ phases, with c,d) the corresponding structural models.

![Figure 2](image2.png)

Figure 2. a) TGA analysis under Ar of the acid-leached sample shows three consecutive weight losses, whereas the parent Li$_3$IrO$_4$ is stable up to 1000 °C. b) Mass spectrometry reveals that the first loss up to 500 °C is due to water, followed by losses of O$_2$ at 700 and 900 °C. c) X-ray powder diffraction of the samples taken at 500 and 1000 °C shows diffraction patterns of IrO$_2$ and metallic Ir, respectively.
transition, which is fully reversed upon the removal of one Li⁺ from the Tl phase, is indicative of a reversible Li insertion-deinsertion process. This is fully confirmed by the remarkable similarity between the XRD patterns of the pristine and discharged/charged electrode (Figure S3, Supporting Information). Lastly, both ⁷Li and ²H NMR signal evolutions were monitored during the Li-insertion into D₃.₄IrO₄. Surprisingly, the ²H signal disappears for Li₋₁D₃.₄IrO₄, but can be explained by the strong interaction with the Ir⁴⁺ unpaired electrons. Similarly, this should cause the disappearance of the ⁷Li NMR signal. This is not the case since Li₋₁.₅D₃.₄IrO₄ shows an intense peak near 0 ppm (Figure S1, Supporting Information), quite more intense than in the pristine material, which is characteristic of a diamagnetic material (i.e., without Ir). Altogether, these two observations indirectly suggest that upon reduction of D₃.₄IrO₄, there is in parallel to the insertion process the growth of a lithiated amorphous phase containing little Ir and no deuterium.

One reasonable explanation for this signal is the deposition of nondeuterated electrolyte decomposition products at the surface of the positive electrode.

In a separate experiment, we tested the electrochemical behavior of H₃.₄IrO₄ in aqueous media using three-electrode PFA Swagelok cells with glassy carbon plungers as current collectors. The same polytetrafluoroethylene (PTFE) laminated electrodes, as described before, were used and assembled against negative electrodes containing capacitive carbon (YPS0) in excess with an Hg/Hg₂SO₄ reference electrode, all of which was immersed in 1 M H₂SO₄ water solution. The electrodes were cycled either in galvanostatic mode or by cyclic voltammetry.

Figure 3b shows the variation of the potential as a function of x. Upon reduction (discharge) from the open circuit voltage (OCV) (0.5 V vs Hg/Hg₂SO₄) to ~0.2 V, the material can uptake one H⁺ (H₃.₄IrO₄) while, on the following oxidation (charge) up to 0.8 V, 1.7 H⁺ are removed (H₂.₇IrO₄). ²H NMR performed
Figure 4. In situ XRD study in a,c) nonaqueous (left) and b,d) aqueous (right) electrolytes. Insertion of Li proceeds with a phase transition from P3 to T1 stacking around x = 0.7 and large changes in a parameter (a = 3.04–3.13 Å) and interlayer distance (d = 4.5–4.7 Å), whereas the process in aqueous electrolyte shows limited changes in cell parameters (a = 3.04–3.06 Å, d = 4.7–4.8 Å) that correspond to the low potential region (below 0.4 V vs Hg/HgSO₄). The high potential region cannot be reached at low current because of the concurrent water splitting reaction.

on the D₄₄IrO₄ sample reveals a deuterium signal in the diamagnetic range (0–20 ppm) that we ascribed to the growth of a solid-electrolyte interface (SEI) (Figure S1b, Supporting Information), since as before no signal should be detected due to the paramagnetic influence of Ir⁴⁺. The span of the sidebands, however, suggests a close proximity with the paramagnetic phase or a less symmetric environment compared to D₄₄IrO₄. Once the first cycle is achieved, the electrode can reversibly uptake 1.7 protons with minor polarization between oxidation and reduction curves, suggesting a rapid and reversible H⁺ release/uptake. Worth mentioning is that by starting the electrode on oxidation we can nearly remove 0.7 H⁺ as well. As an indication, from simple charge counting, this implies that on reduction H₃₄IrO₄ transforms to H₄₄IrO₄ which will be oxidized to H₂IrO₄ upon full oxidation.

CVs collected at 1 mV s⁻¹ between -0.1 and 0.8 V versus Hg/HgSO₄ (Figure 3d) show several redox peaks confirming a large electrochemical activity which can be estimated to a reversible uptake of 1.7 protons as determined by galvanostatic cycling. Strikingly, the CV obtained after 5000 cycles at 20 mV s⁻¹ nearly superimposes with the first one, indicative of an amazingly reversible redox process not too common at such level for battery materials (Figure S4, Supporting Information). Finally, the rate capability of the material was measured from 1 to 1000 C. It reveals the high-power performances of the electrode, which can sustain a capacity of 130 mA h g⁻¹ up to 100 mV s⁻¹ (≈400 C) when the electrode thickness is decreased from 150 to 30 µm (Figure 3f). This value competes favorably with pseudocapacitive performances of transition metal carbides (also known as MXene), which can uphold ≈92 mA h g⁻¹ up to 100–200 mV s⁻¹ for electrodes of comparable thickness and loading. However, unlike the MXene or other pseudocapacitive materials, the rate capability of H₃₄IrO₄ is, as expected, largely depending on the electrode design. An even more drastic
To understand the kinetic differences between the low and the high potential processes, Swagelok cells using thin films as positive electrodes were further assembled. Cyclic voltammograms were collected up to scan rates of 50 mV s\(^{-1}\) at first over the full potential range (Figure 5a) and then independently over the low and the high potential range (Figure 5b,c). Both the low and the high potential regions show kinetic limitations as can be seen from the increasing polarization of anodic and cathodic processes (peaks moving to the right or the left in oxidation and reduction, respectively) whatever these potential ranges are swept independently or together. However, noticeable differences can be observed by focusing on the slope of the oxidation and the reduction curves at various rates (Figure 5, bottom) with namely greater rate polarization dependence for the lower potential domain compared to the high potential one. This observation, combined with the disappearance of the low potential domain without carbon, strongly suggests that the kinetic of the low potential process is under ohmic control. Interesting to note is the symmetry of the polarization for the high potential domain as opposed to an asymmetric one for the low potential domain, with the kinetic being more sluggish on oxidation than on reduction. This is indicative of a different ohmic limitation on charge and discharge, which is most likely linked to changes in the electrode conductivity, as will be discussed later.

To get rid of the ohmic limitations at high scan rates and better assess the kinetics of the material itself, we repeated the aforementioned experiments using a cavity microelectrode (CME), which presents several advantages compared to conventional Swagelok set-ups. First, neat active material powders can be used free from any binder. Besides, the feasibility to use only a few micrograms of active material helps to achieve high cycling rates since ohmic drop is negligible for the low current flowing in the electrode. As shown in Figure 6 (Supporting Information), ohmic limitation are effectively negligible from 1 to 200 mV s\(^{-1}\) for all processes, except for the anodic process of the low potential region, whose kinetic seems limited by the (poor) electrical percolation inside the active material.

To fully sort out the origin of the kinetic limitation in the low potential domain, we decided to run step potential-time transient technique (PITT) potentiostatic experiments\(^{19}\) using conventional Swagelok cells in the −0.4 to +0.8 V versus Hg/HgSO\(_4\) range. During each potential step, the current was recorded versus time and then integrated to calculate the electrode capacity at different equivalent potential scan rates (see the Experimental Section). As a result, we could recalculate and replot cyclic voltammetry curves at different potential scan rate. Those CVs calculated from potentiostatic experiments (Figure 6) differ from real CV experiments shown in Figure 5 in the way that ohmic contribution is minimized using a potentiostatic technique. We considered the current at each potential from the recalculated CVs as the sum of a nondiffusion-limited (surface) and a diffusion-limited (bulk) processes\(^{20-22}\)

\[ i(v) = k_1 v + k_2 \sqrt{v} \]  

where the nondiffusion-limited redox contribution is defined by the \( k_1 \) coefficient, associated with electrochemical processes occurring onto accessible surface sites, and the \( k_2 \) coefficient represents bulk diffusion-limited redox processes. Finding the values of \( k_1 \) and \( k_2 \) at each potential allows for the separation of the nondiffusion and diffusion currents (orange and blue in Figure 6, respectively). The area under the CV is the sum of the blue and orange areas. First, the CV envelop is consistent with the experimental CVs plotted in Figure 5 under conventional dynamic conditions, validating the experiments. Furthermore, Figure 6 shows that the electrochemical kinetics of the low and the high potential processes are equivalent since the ratio of their respective area are roughly similar at each scan rates, for both the diffusion limited and the nondiffusion limited contributions. These results unambiguously support the existence of an ohmic limitation in the low potential domain in Figure 6 rather than an electrochemical kinetic limitation.

At this stage a legitimate question regards the mechanistic of such low and high potential processes which, in light of the aforementioned results, are most likely not capacitive. To grasp further insights into this issue, in situ XRD measurements were done using a similar cell as before with the exception that the Be window, which cannot chemically resist acidic electrolyte, was replaced by a Kapton film, hence we reconfigured the positive electrode compartment (see the Experimental Section). We follow a similar experimental protocol as before (XRD pattern taken for every step in \( \Delta E \) of 0.05). The 2D contour plot representation of the collected data (Figure 3b) shows a smooth and continuous displacement of the Bragg peaks toward lower angles upon reduction to −0.2 V, indicative of a proton-driven solid solution insertion process. This process is perfectly reversible as the Bragg peaks move back toward their initial position by switching the current. Refinement of the collected patterns indicates that the P3 stacking is preserved through the process, with however a decrease in both \( a \) and \( c \) lattice parameters when the potential is increased to 0.4 V and a flattening toward the end. Altogether, these results prove that the electrochemical activity at low potential is simply due to a classical insertion-deinsertion reaction.

The upper potential domain turns out to be more difficult to explore as we could not reach oxidation potentials greater than 0.5 V with our cell hardware because of copious water decomposition. To circumvent these difficulties, we returned to ex situ measurements where an electrode was polarized to 0.8 V in Teflon Swagelok cells, the glassy carbon current collector was recovered with the working electrode and directly measured by XRD without further processing (Figure S7, Supporting Information). No change in lattice parameters was detected, suggesting that the high potential storage process does not imply structural changes. To comfort the robustness of such a claim and ensure absence of experimental artefacts, the potential of the sample was measured again in an electrochemical cell after XRD and nearly the same potential was measured at OCV. Lastly, and for sake of completion, it is worth mentioning that identical results were obtained using HCl rather than H\(_2\)SO\(_4\) as electrolyte.
3. Discussion

We have reported a new chemical aspect of the Li-based iridates that is their great aptitude to form protonic phases as demonstrated with the formation of H$_3$IrO$_4$. Such a new phase demonstrates an exciting, but not yet fully understood, electrochemistry with namely an electrochemical activity of 1.5 e$^-$ per Ir metal in both aqueous and nonaqueous media with fast kinetics.

From the structural point of view, this work offers very rich crystal chemistry, summarized in Figure 7, as we start with O3 stacking in the parent Li$_2$IrO$_4$ phase, that changes to P3 stacking in H$_3$IrO$_4$ by acid leaching and eventually transforms to a T1 stacking or maintain the P3 stacking upon additional
Figure 6. Calculated CVs from PITT experiments. The contributions of the diffusion limited and nondiffusion limited electrochemical reaction kinetics are shown in blue and orange colors, respectively, for PITT measurements collected at 1 mV s⁻¹ (top) and 5 mV s⁻¹ (bottom). These contributions were calculated from the deconvolution of the total current (see text).

Figure 7. Overall structural mechanism. Acid leaching of the parent Li₃IrO₄ phase (A) modifies the stacking from O₃ to P₃ for H₃₄IrO₄ (B). Cycling in aqueous media does not modify the stacking (C) whereas insertion of Li in organic electrolyte changes the stacking to T₁ (D).
2.11-2.43 Å to the oxygen atoms of the same layer (Figure 8b). The metal cations and oxygens are shown as gray and red spheres, respectively. The "out-of-plane" and "in-plane" proton positions are colored with dark blue and cyan, respectively. Thick and thin bonds denote short covalent O-H bonds and long hydrogen bonds, respectively.

ordering of the AI02 oxygen atoms from one (Cr02) octahedral layer and a weaker hydrogen bond with the facing oxygen in the adjacent layer. Al(OH)3 shows a honeycomb ordering of the AlO6 octahedra and vacant oxygen octahedra that creates free space to accommodate protons almost in the oxygen close packed planes. The metal cations and oxygens are shown as gray and red spheres, respectively. The "out-of-plane" and "in-plane" proton positions are colored with dark blue and cyan, respectively. Thick and thin bonds denote short covalent O-H bonds and long hydrogen bonds, respectively.

Together, the "out-of-plane" and "in-plane" proton positions can account for the hydrogen atoms in CrOOH.

Turning to the fundamental (de)insertion mechanisms of these protonic phases, they fall in line with the classical cationic insertion mechanism involving the reversible Ir3+ → Ir4+ redox for the storage capacity of Li/H2IrO4 cells. Charging the material above 4.2 V versus Li+/Li results in limited capacity retention indicative of some electrolyte degradation at high potential. Thus, the need to develop a proper electrolyte formulation that resists to hydrolysis to properly evaluate the deprotonation of the material and its long-term cycling stability. In aqueous media there is, besides a low potential process governed by cationic insertion, a high potential process associated to a reversible uptake/release of 0.7 e- for which, based on charge counting, one could invoke some anionic redox process: "H2,7Ir3+O2(OH)1.5". This is quite unlikely as anionic redox affects the structure and usually leads to sluggish kinetic as compared to cationic redox; which is not what we found. Moreover, the participation of lattice oxygen can lead, as discussed by Grimaud et al. to an OER activity which could jeopardize the stability of our electrode, but this is not the case either. Thus, we are left with a small amount of Ir4+, which is not frequent, or, more likely, we underestimated the amount of H+ in the insertion of Li+ or H+, respectively. Therefore, getting insights into the local positioning of the different guest ions, namely H+, is quite complex, owing to their transparency to XRD, as well as to the large incoherent neutron scattering of H+. which can be overcome by using deuterated samples. In absence of such measurements, a valuable alternative consists in looking for possible proton location among the known structures of layered hydroxides such as CrOOH, which adopts a P3 structure (Figure 8a). Typically, protons form strong covalent bond to one oxygen atom, with a typical length of ~0.85–1.0 Å, and one weaker "hydrogen" bond to another oxygen atom, which has more electrostatic nature and a length that can vary between 1.5–2.5 Å. In CrOOH, protons occupy "out-of-plane" position between two successive (CrO2) octahedral layers being closer to one O atom (d(O–H) = 1.05 Å) and more distant from the O atom from the neighboring layer (d(O–H) = 1.53 Å).[21] Thus a linear O–H–O fragment is formed.

Turning to the protons within the metal layers, their position at the Li octahedral sites is quite unlikely. A more likely location could be deduced from the known hydroxide Al(OH)3, which surprisingly has a very similar structure to Li-rich layered oxides with honeycomb ordering.[24] In this structure, the honeycomb ordering of the Al-filled and vacant oxygen octahedra within the (AlO2) layers provides extra "in-plane" proton positions, sitting almost within the oxygen close packed layers, with one short O–H bond of 0.94–1.0 Å and another longer one of 2.11–2.43 Å to the oxygen atoms of the same layer (Figure 8b). Together, the "out-of-plane" and "in-plane" proton positions can account for the hydrogen atoms in Li2H3IrO4.

Now moving to the inserted compound, we should recall that the insertion of Li in Li2H3IrO4 has led to a phase change through which the Li-ions move from octahedral to tetrahedral sites. Along that line it does not come as a surprise that the insertion of 1.5 Li+ in H2H3IrO4 triggers a P3 to T1 structural transition with the extra Li+ occupying the tetrahedral sites. As we suspect that 7Li and 1H NMR signals from the material’s bulk are not visible due to the vicinity of paramagnetic Ir4+ cations, neutron diffraction on deuterated samples is being planned to support most of our hypothesis relying on well-known compounds.

From a material’s prospective we should recall that the iridate family is rich and still largely unexplored, hence the numerous opportunities presented by this simple room temperature one step acid leaching technique. For proof concept generalization, we have already prepared several other members, among which are the α and β-Li2IrO4 phases with either Li or Na, and we could equally confirm the feasibility to obtain the corresponding protonic phases with a reaction kinetics slightly depending upon the pristine material structure. For broader generalization, we could also demonstrate that the ruthenates share similar chemistry versus the leaching technique approaches which, however, compete in some cases with decomposition reactions. So, a multitude of new protonic phases remain to be discovered and studied for their electrochemical properties.

Figure 8. Proton positions in a) layered CrOOH and b) Al(OH)3 bayerite structures. In the CrOOH structure, protons form a strong covalent bond with oxygen atoms from one (CrO2) octahedral layer and a weaker hydrogen bond with the facing oxygen in the adjacent layer. Al(OH)3 shows a honeycomb ordering of the AlO6 octahedra and vacant oxygen octahedra that creates free space to accommodate protons almost in the oxygen close packed planes.
4. Experimental Section

Materials Synthesis and Characterization: Li$_3$IrO$_4$ was prepared by mixing stoichiometric amounts of metallic Ir and Li$_2$CO$_3$ and heating at 950 °C for 24 h in air. The prepared sample was then acid-leached in 0.1 M H$_2$SO$_4$ during 1 h with a fivefold excess of H$^+$ over exchangeable Li$^+$ in the material. The powder and the supernatant were separated by centrifugation and the sample washed three times with distilled water prior to being dried overnight at 55 °C under vacuum. D$_2$SO$_4$ solution and D$_2$O were alternatively used to prepare deuterated samples. The Li$^+$ and H$^+$ contents in the supernatant after reaction were measured by ICP-MS and acid–base titration, respectively. TGA-MS was done under high purity Ar gas (5.0) with 15–20 mg of samples, using a heating ramp of 5 K min$^{-1}$ from room temperature to 1000 °C. X-ray diffraction was performed at each step to confirm the success of the synthesis using a Bruker D8 Advance diffractometer with Cu Kα radiation ($\lambda_{CuKα}$ = 1.54056 Å, $\omega_{CuKα}$ = 1.54439 Å). NMR experiments were performed on a Bruker 4.7 T Avance III spectrometer, using a 1.3 mm double-resonance probehead. The spinning rate was 62.5 kHz except for the D$_2$IrO$_4$ sample that could not be spun above 59 kHz. $^7$Li NMR experiments were performed using a Hahn-echo with a repetition time chosen to ensure full relaxation, and referenced against an aqueous LiCl solution (1 mol L$^{-1}$) at a chemical shift of 0 ppm. $^{1}$H NMR experiments were performed using a quadrupolar echo and a repetition time of 0.1 s, and referenced against liquid D$_2$O at 4.8 ppm.

Electrochemical Characterization: For electrochemical characterization, self-standing PTFE electrodes were prepared. H$_2$IrO$_4$ was mixed with 10 wt% Csp and 5 wt% PTFE, and laminated several times to obtain films of different thickness (10–150 µm) and density of material (4–30 mg cm$^{-2}$). Li half cells were prepared with the thicker films with typical loadings of 10 mg. Metallic lithium was used as the anode material, LiP06 (1 m LiPF$_6$ in ethylene carbonate/propylene carbonate/dimethylcarbonate with 1:1:3 weight ratio) as the electrolyte and Wh atm an GF /D borosilicate glass electrode. The proton release/uptake value was obtained from coulombic titration assuming the formula H$^+$/Li$^+$ for the starting material and that only protons participate to the charge storage mechanism. The cavity microelectrode used here consists in a thin platinum wire (Ø 100 µm) sealed into a low melting glass; a laser beam was then focused on the Pt wire to dig a cavity of 80 µm in diameter and 70 µm in depth. The active material was mixed with 10 wt% Csp and dipped the CME in the powder to fill the cavity. For the in situ XRD experiment, a Swagelok-derived electrochemical cell equipped from a Be window was used in organic electrolyte.$^{57}$ The same measurements were done in acidic media by replacing the Be window with a Kapton film and a flexible graphite film as current collector, the latter being punched with a hole slightly smaller than the working electrode size, in order to ensure a good electronic conductivity without masking the diffraction signal of the material. Prior and after the measurement with the in situ cell, the working and counter electrodes used in the experiment where measured in the three-electrode set up in order to establish the relationship between the cell voltage and the working electrode potential.

PITT Experiments and Kinetic Study: PITT experiments were done in PFA Swagelok cells using thin electrodes. During the experiment, the potential was increased by steps of 25 mV, and hold during 60 s. For each potentiostatic step, the current was recorded versus time. The current was integrated versus time according to $Q = \int i dt$ to calculate the electrode capacity at different equivalent potential scan rates. For instance, the integration of the current during the first $\Delta t = 5$ s for a potential step of 25 mV gives an equivalent scan rate of 5 mV s$^{-1}$ and the integration during the first 25 s gives an equivalent scan rate of 1 mV s$^{-1}$. The mean current during each potential step was calculated according to $I = \frac{Q}{\Delta t}$. As a result, for each potential step, the current and replot cyclic voltammagety curves could be recalculated at different potential scan rates. CVs calculated from potentiostatic experiments minimize the ohmic drop contribution. To estimate the contribution of the diffusion limited and nondiffusion limited processes to the total

material owing to the experimental uncertainty. The kinetic of each process is dictated by the slowest contribution to the electrochemical reaction and we have shown from PITT measurements that the two processes observed at low and high potential have similar electrochemical kinetics (charge transfer and diffusion). They only differ by the existence of a greater ohmic polarization for the low potential domain process which cannot be triggered in absence of carbon additive. Its origin is most likely linked to the increasing amount of Ir$^+$ during the acid leaching treatment. Nevertheless, a question still remains regarding the absence of structural changes detectable by XRD for the high potential process. We thus could envision, in light of previous studies on V$_2$O$_5$ powders showing extra capacities not associated to structural changes, the emergence of a core–shell like process within a single particle, during the uptake-removal of proton, with the shell operating via a pseudocapacitive behavior enabling changes in the Ir oxidation state without affecting the structure. Obviously, further experiments, namely, operando X-ray absorption spectroscopy at Ir-LIII edge to track changes in the Ir oxidation states, have to be done to test our hypothesis and fully elucidate the high potential mechanism observed for the aqueous H$_2$IrO$_4$/Li$_3$IrO$_4$ system.

In summary, we have successfully synthesized a new protolytic iridate phase that shows large capacity storage (1.5 e$^{-}$ per transition metal), while presenting attractive rate capabilities in aqueous systems. Such a storage process was shown to mainly rely on cationic redox reactions with the eventual conivance of a pseudocapacitive behavior to account for the total redox process in aqueous media. Besides the synthesis and fundamental aspects, this work also bears importance application-wise provided that we can implement our protonation strategy to other Li-rich phases, hence deviating from the use of poorly abundant and costly Ir. Lastly, this proton chemistry opens new opportunities within the field of energy bearing in mind that Ir-based materials also present a great interest for water splitting applications.
current, the current at each potential as the sum of the nondiffusion limited and diffusion limited processes was considered as follows: 

\[ i(v) = k_1 v + k_2 v^2 \]

where the nondiffusion limited redox contribution is defined by the \( k_1 \) coefficient, associated with electrochemical processes occurring onto accessible sites, and the \( k_2 \) coefficient representing diffusion limited (bulk) processes. Consequently, it becomes possible to determine the phenomenon governing the reaction kinetics by calculating the ratio \( k_1/k_2 \) and to accurately observe the changes in the kinetic behavior of the material. Considering Equation (1) and plotting \( i/v^2 \) versus \( v \), \( k_1 \) was calculated as the slope of the plot, and \( k_2 \) its y-intercept. Calculation of \( k_1 \) and \( k_2 \) at each potential allows for the separation of the nondiffusion and diffusion currents.

**Supporting Information**

Supporting Information is available from the Wiley Online Library or from the author.

**Acknowledgements**

The authors thank M. Courty for TGA-MS measurements, D. Giaume for ICPMS measurements, C. Yang, A. Grimaud, and G. Rousse for useful discussions. J.-M.T. acknowledges funding from the European Research Council (ERC) (FP/2014)/ERC Grant-Project 670116-ARPEMA.

**Conflict of Interest**

The authors declare no conflict of interest.

**Keywords**

batteries, layered oxides, protons, pseudocapacitors

---


