Evolution of Oxygen Ligands upon Large Redox

Swings of Li₃IrO₄

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Abstract

The limits of intercalation electrochemistry continue to be tested in the quest for ever increasing gains in the storage capability of Li-ion cathodes. The subsequent push for multi-electron reactivity has led to the recognition of the extremely versatile role of oxide ligands in charge compensation when there is a large redox swing. Li₃IrO₄ is a unique model of such activity because it can reversibly cycle between Li₁IrO₄ and Li_{4.7}IrO₄. Here, X-ray spectroscopy, magnetic measurements and computational simulations uncover the evolution of O states in the different steps, compared to the involvement of Ir. While the process between Li₁IrO₄ and Li₃IrO₄ is dominated by the unconventional lattice oxygen redox, the process between Li₃IrO₄ and Li_{4.7}IrO₄ involves a conventional change of the formal oxidation state of Ir, which affects O due to the high covalency. The O states of Li₃IrO₄ exhibit a very high reversibility after the whole 3.7-electron process, completely restoring the pristine state.

1. Introduction

As the world is transitioning into a clean energy paradigm and mobility that is free of emissions, energy storage has grown to be significantly prominent at a societal level. ¹⁻⁵ Among energy storage devices, Li-ion batteries are the most promising due to their high energy density. The manipulation of the redox chemistry of the transition metal (TM)-oxide bond in reactions of electrochemical intercalation in solids was foundational to the advent of this technology. The storage process of the most widely used cathode, α -NaFeO₂-type layered transition metal oxides Li[TM]O₂ (TM = Co, Ni, Mn, among others), involves (de)intercalation concurrent with a conventional redox mechanism: LiTM³⁺O₂ \leftrightarrow xLi⁺ + xe⁻ + Li_(1-x)TM^{(3+x)+}O₂. ⁶ Challenges associated with the structural and chemical damage upon achieving high degrees of oxidation in these phases, ^{1, 7-9} have motivated researchers to define the limits of intercalation reactions to enable the largest possible reversible swings in composition and redox compensation. ³ In particular, a fundamental topic of interest is whether more than 1 mol Li can be stored per mol of total transition metal content.

In comparison with conventional layered oxides Li[TM]O₂, layered oxides with a Li overstoichiometry, Li[Li_xTM_{1-x}]O₂, have recently become the object of scrutiny due to the potential to achieve capacities as high as 300 mAhg⁻¹.¹⁰⁻¹⁹ As a result, they seemingly break the barrier of 1 mol Li extracted per mol TM. This extraordinary activity stems from the accumulative contribution of conventional transition metal redox and O participation. Broadly speaking, unconventional ligand-centered redox (also referred to as "lattice oxygen redox") is triggered in oxides as the stoichiometric ratio of oxide and transition metal increases,²⁰ and non-bonding states are incentivized in O by the existence of Li overstoichiometric in its coordination sphere.²¹ Despite their enhanced electrochemical activity, they still suffer from detrimental chemical behavior, such

as oxygen evolution from the lattice,²²⁻²⁴ extensive structural reorganization,²⁵⁻²⁷ significant voltage hysteresis and fade,²⁸⁻³⁰ and sluggish kinetics.^{16, 31, 32}

In the quest to explore the fundamental limits of redox swing possible in an oxide, Perez et al. discovered that Li₃IrO₄ is capable of reversibly releasing Li at ~4 V vs. Li⁺/Li⁰ to form the nominal composition of Li₁IrO₄, while also uptaking Li down to 1.3 V to form nominally Li_{4.7}IrO₄.³³ The two processes could be combined in a single electrochemical experiment, for a total reversible swing of 3.7 mol Li per mol compound, the largest change observed in an oxide, even when compared to other Li₃TMO₄ phases.³⁴⁻³⁷ This reactivity is thought to be enabled by the ability of oxide ligands to not only compensate changes in the formal charge of Ir, following conventional cation redox, but also to contribute electrons from initially non-bonding states that subsequently hybridize with Ir, such that the metal does not formally change charge.^{33, 38}

Whereas *operando* Ir L_{III}-edge X-ray absorption spectroscopy (XAS) has been employed to study the electrochemical process between Li₁IrO₄ and Li_{4.7}IrO₄,³³ the detailed charge compensation mechanism must be completed by understanding the participation of O in the covalent interactions that enable it. The study of the Li₃IrO₄-Li₁IrO₄ revealed the existence of selective activity of the non-bonding states of the two unique O sites in the structure.³⁸ Here, we evaluate the changes occurring at O states in the complete compositional space, with a focus on whether the most reduced state, Li_{4.7}IrO₄, is the same if produced from Li₁IrO₄, after one initial charge, or directly upon reduction of Li₃IrO₄. To avoid confounding phenomena occurring at the surface of the materials,^{27, 39, 40} we turn to O K-edge XAS with sensitivity to the interior of the material,^{23, 24, 34, 41-46} complemented with magnetic measurements. The results complete the picture of the role of oxygen in the charge compensation mechanism in such a wide redox space.

2. Experimental and Computational Methods

2.1 Sample Preparation

Li₃IrO₄ was synthesized by a solid-state reaction of Li₂CO₃ (Sigma Aldrich, 99%) and metallic Ir (Alfa Aesar, 325 mesh, 99.9%) in stoichiometric proportions, as reported by the previous report.³³ The precursors were ground together using a mortar and pestle and then the homogeneous mixture was heated at 950°C for 24 hours in air atmosphere. After cooling to room temperature naturally, the as-synthesized sample was ground into fine powders and stored in an argon-filled glovebox (H_2O , $O_2 < 0.1$ ppm) for further characterization.

2.2 Working electrode preparation

Samples for *ex situ* XAS were prepared by galvanostatically cycling the positive electrode in typical two-electrode 2032 coin-type cells. The composite positive electrode materials were prepared by homogeneously mixing 80 wt% active material, 10 wt% carbon black (Denka), and 10 wt% polyvinylidene fluoride (PVDF) (Kynar) in 1-methyl-2-pyrrolidone (NMP, Sigma–Aldrich). The slurry were then blade-rolled on an electrochemically grade aluminum foil with the thickness of 6 mm and the typical active material loadings of 3-5 mg/cm², followed by a complete drying under vacuum at 110°C overnight. Dried electrodes were then punched into ½-inch diameter discs and transferred into an argon-filled glovebox (< 0.1 ppm of both H₂O and O₂) for further cell assembling. All the cells were fabricated in an argon-filled glove box with moisture and oxygen levels lower than 0.1 ppm. The cell was composed of the positive electrode and a high-purity lithium foil (Alfa Aesar) as the counter/reference electrode, separated by a 25-μm-thick polypropylene membrane (Celgard 2400), impregnated with 1 M LiPF₆ dissolved in a mixture of ethylene carbonate (EC)/dimethyl carbonate (DMC) (ethyl methyl carbonate (EMC) (3:7, wt %/wt %). The galvanostatic charge–discharge cycling was performed at room temperature using a

BT-Lab tester with a current rate of C/20 (indicating 1 Li is extracted in 20 h) with cutoff voltages of interest. All potentials quoted in this paper were referenced to Li⁺/Li⁰ unless otherwise mentioned. Upon arriving at the state of interest, the cell was immediately stopped and disassembled in the argon-filled glovebox to avoid self-discharging under open-circuit state and the inside cycled electrode was harvested and washed with DMC three times, followed by drying under vacuum in the glovebox antechamber for 30 mins. Cleanly washed and dried electrodes were stored in the argon-filled glovebox for *ex situ* characterization.

Samples for *ex situ* magnetic measurements at different states of charge/discharge were prepared by cycling the pristine material in Swagelok-type cells. The active material was mixed with 10 wt% carbon SP and used as the positive electrode, while metallic lithium was used as the negative electrode. A Whatman GF/D borosilicate glass fiber membrane was used as the separator and was soaked with 1 M LiPF₆ in ethylene carbonate (EC)/propylene carbonate (PC)/dimethyl carbonate (DMC) with a volume ratio of 1:1:3. The cells were assembled in an argon-filled glovebox and galvanostatically cycled using VMP3 cycler at C/20 rate at room temperature. Once the state of interest was reached, the cell was disassembled in the glovebox and the material was harvested and washed using DMC, then dried under vacuum in the glovebox antechamber for 30 mins. Fully dried electrodes were stored in the argon-filled glovebox for characterization.

2.3 Magnetic measurements

Magnetic susceptibility measurements were carried out using a SQUID magnetometer (Quantum design), in zero field cooled (ZFC) mode, with an applied magnetic field of 1 kOe between 300 and 2 K. Samples were prepared by mixing with conductive carbon (Super C65 from Imerys, typical Fe content of 2 ppm), cycled at C/20 in Swagelok cell to the desired Li content and left to equilibrate for a few hours before recovering the powder, which was washed three times

with anhydrous dimethylcarbonate and dried under vacuum. About 10 to 20 mg of the samples were loaded into gel caps for the measurement and sealed in plastic tubes to avoid reaction with air prior to the measurement. The gel cap, plastic tube and conductive carbon were chosen for their negligible contribution to the magnetic susceptibility. The data was fitted between 25 and 250 K using a modified Curie-Weiss law with a temperature independent contribution χ_0 which takes into account the diamagnetic contribution of constituting diamagnetic elements (with an order of magnitude of -10⁻⁵ emu mol⁻¹):

$$\chi = \chi_0 + \frac{C}{T - \theta}$$

2.4 X-ray absorption spectroscopy

O K-edge X-ray absorption spectroscopy (XAS) data was collected *ex situ* at the beamline 4-ID-C of the Advance Photon Source (APS) at the Argonne National Laboratory. Samples of interest were attached to a 7T copper sample holder using conductive carbon tape in an argon-filled glovebox and then transferred into a 3-way transport container and then into an X-ray absorption antechamber through an argon environment to minimize the potential exposure to air. Data were measured simultaneously under both the total electron yield (TEY) mode from the sample photocurrent at $\sim 10^{-9}$ Torr and total fluorescence yield (TFY) mode using a silicon drift diode detector (Vortex). Data was obtained at a spectral resolution of ~ 0.2 eV, with a 2 s dwell time. During the measurement, three scans were performed at each absorption edge, and scans were then averaged to maximize the signal-to-noise ratio. The energy scale of the spectra was calibrated with a Sr₂RuO₄ reference measured simultaneously with samples.

During the measurement, the XAS spectra were recorded over a wide energy range from 520 to 560 eV to cover energies well below and above sample absorptions. Then the normalization and

background removal were performed via established procedures in the literature.⁴⁷ First, the collected data was normalized to I₀ measured from the sample drain current from a freshly coated Au-mesh inserted into the beam path before the X-ray irradiates the sample. Subsequently, a linear, sloping background was removed from all spectra by fitting a linear function to the region from 520 to 524 eV, i.e. at energies below any absorption peaks.

2.5 DFT calculations

XAS simulation

The XAS spectra were computed for all structures using the OCEAN code^{48, 49}OCEAN solves the Bethe-Salpeter equation (BSE)⁵⁰ based on the ground-state charge density and wave function obtained from the DFT-based Quantum Espresso (QE) program package, ⁵¹ using the local-density approximation (LDA or LDA+U)^{52, 53} in conjunction with norm-conserving pseudopotentials.⁵⁴ To solve the Kohn-Sham equation, the cut-off value for the basis plane-waves was set to 952 eV and the k-point grid for the electronic density was converged with increasing k-point grids. As for VASP calculations used to relax all the structures prior to the XAS simulation, Hubbard correction was added to Ir(d) shell and spin-orbit coupling (SOC) was also considered to compute the electronic structures of materials. As shown in Figure S1, very similar electronic structures were obtained for Li₃IrO₄ and Li₄IrO₄ compounds when computed within the DFT+U formalism with or without including SOC. Remarkably, we observed that the splitting of the $t_{\rm 2g}$ and $e_{\rm g}$ states reported as the origin of strong spin-orbit coupling in Ir-based oxides^{55, 56} is already observed in the electronic structures obtained from DFT+U calculations and only slightly modified when SOC is further introduced. Accordingly, the influence of spin-orbit on the O K-edge XAS spectra is expected to be limited as the DFT+U formalism is used for the first-step electronic structure calculation on top of which the XAS simulation is performed.

The main advantage of the BSE approach is to explicitly account for the screened core-hole and photoelectron interaction and to include self-energy correction to the electronic states. In the present calculations, BSE was solved using 500 unoccupied bands and 800 bands were used to build the screened core-hole potential with the default k-point grid. Only dipole-allowed transitions were considered in the XAS calculations. The photon polarization vectors were set at [100], [010], and [001], and the final spectrum of each structure was obtained by averaging the spectra over all polarization vectors and individual oxygen atoms.

Core level shifts (CLS) were included in the calculations to account for the different local environment of the two oxygens occurring in the pristine Li₃IrO₄. The OCEAN code is built on top of one-electron wavefunction pseudopotentials with, in principle, no explicit access to core level states. To overcome this issue, OCEAN uses an optimal projection procedure⁵⁷ to orthogonalize the pseudized valence states to the core-states, alike the projected-augmented-wave (PAW) method developed by Blöchl⁵⁸ and used in VASP calculation. The relative shifts of the core-level energies are then evaluated through the change in the Kohn-Sham potential due to different chemical environments. Then the screening of the core-hole due to the valence electron, which also depends on the local environment of the absorbing atoms and on its charge, is computed through linear response theory. The convergence of the screening procedure was checked in our calculations by increasing the screening radius up to convergence. Absolute excitation energy were then aligned with the experimental data through an ad hoc shift.

3. Results and Discussion

The detailed electrochemical behavior and the corresponding verification of Li content in cycled samples have been reported in our previous publication.³³ Overall, Li can be directly intercalated into Li₃IrO₄ at low potential (black lines in Figure 1a), via a long plateau at \sim 1.7 V,

corresponding to a capacity equivalent to the insertion of 1.7 mol of Li per mol compound at 1.3 V. The subsequent oxidation to 2.5 V proceeds through two processes located at 1.8 and 1.9 V, respectively, leading to almost reversible re-extraction of 1.5 mol of Li. Ir L_{III}-edge XANES reveals significant, yet fully reversible changes in absorption energy associated with a conventional redox mechanism based on formal cationic centers, which reflects population of Ir 5d-O 2p covalent states. Li₃IrO₄ can also be oxidized to 4.5 V, resulting in the amorphization to "a-IrO₃" and an irreversible reduction to 2.0 V with notable voltage hysteresis and capacity loss (gray lines in Figure 1a). By limiting the initial oxidation to Li₁IrO₄, however, Li₃IrO₄ could experience a reversible subsequent reduction to 2.6 V with high reversibility.^{33, 38} The compound can continue to be reduced to 1.3 V to produce an extraordinarily high capacity equivalent to 3.7 mol of Li per mol compound, followed by a reversible oxidation to the pristine state (Figure 1b). In order to study how and in what form O was involved in the electrochemical reactions, electrodes were harvested at representative state points of these electrochemical reactions, as indicated in Figure 1. This article combines data from XAS (Figure 2) and magnetic measurements (Figure 3), all of which probe the electronic structure centered at O 2p and Ir 5d states. Computational analysis probed the changes in electronic structure and quantitatively related them to the XAS.

O K-edge XAS is used to explore the variation of the electronic structure of O. Signals arise from dipole-allowed transition from core O ls to unoccupied O 2p states. In general, the spectra of these transition metal compounds can be divided into two regions (Figures 2a). The pre-edge feature (≤ 535 eV, in general) represents the unoccupied states resulting from O 2p orbitals hybridized with transition metal nd orbitals, and the broad band above 535 eV corresponds to a collection of excitations from O ls orbital to empty states of O 2p orbitals mixed with the transition metal (n+1)s and (n+1)p orbitals, O 3p orbitals, and ultimately, the continuum. The position of the

pre-edge peak is affected by the change in the net electron density of the ligand via donating charge to the surrounding metal ion, the degree of d orbital splitting induced by the ligand field effect, and the overall TM d-manifold orbital energy.⁵⁹ The intensity of these peaks reflects both the density of unoccupied hybridized states and the degree of covalency of the ligand-metal bond.⁶⁰ In addition, the rising edge inflection point in the broad band deduced by the 1st derivative calculation of the XAS spectrum reflects the onset energy of the main edge position,⁵⁹ and thus could indicate the change in the binding energy of the O 1s core level.

The O K-edge XAS of pristine Li₃IrO₄ (Figures 2a) exhibits four distinct features centered at 527.9, 529.1, 531.7, and 533.7 eV, all below 535 eV, with the first three features corresponding to transitions to unoccupied O 2*p*-Ir 5*d* hybridized states, and the broad features above 535 eV arising from O 2*p*-Ir 6*s*, *p* states and the multiple scattering events upon photoionization. It should be noted that the small feature at 533.7 eV can be attributed to Li₂CO₃ impurity which dominates the surface signals (total electron yield, TEY, with a probing depth of 10 nm) (Figure 2a).⁶¹ The result shows that these samples showed a notable sensitivity to air, with batches where contamination was enough to produce these signals, albeit at much lower relative intensity, even in fluorescence yield measurements (total fluorescence yield, TFY, with a penetration depth of 100 nm, probes the interior of the material, Figure 2b). Due to the presence of Li₂CO₃ and the potential for side reactions at the surface, the analysis mainly focused on TFY spectra to probe volumes where the bulk dominates the signal (Figure 2b). It is worthy of notice that spectral intensities in TFY are distorted by the self-absorption of fluorescent photons by the material, which is not present in TEY, so only qualitative trends between samples will be established.

In our previous paper, we reported that Li₃IrO₄ displays extremely reversible oxide-centered redox upon oxidation to Li₁IrO₄.³⁸ The following reduction was also possible, accumulating Li

beyond the initial state (sample F in Figure 1b). 33 O K-edge XAS of sample F (Li_{3.6}IrO₄) presented a shift of the first two pre-edge peaks to higher energy, at 528.1 and 529.1 eV, concurrent with a large decrease of their intensity (Figures 2b and S2), particularly at the lowest energy. These features are confirmed by DFT calculations for which a cationic reduction is predicted, consistent with a sharp decrease in intensity of the two peaks corresponding to the t_{2g} -like peaks supported by the projected density of states (pDOS) of Li₄IrO₄ which shows the decrease of a peak above Fermi level corresponding to the Ir d_{t2g} -O 2p states (Figures 4 and S3). Above 530 eV, the main feature at ~531.7 eV also experienced little shift to position, with an obvious, yet comparably smaller, reduction in intensity. Lastly, the rising absorption edge shifted to lower energy by ~0.2 eV compared to 2.6 V (Figure S4), indicating a relative decrease in the Z_{eff} of O and shallower binding energy shift.

Complete reduction to 1.3 V from Li₁IrO₄ induced a total swing of almost 4 mol of Li, and a nominal composition of Li_{4.7}IrO₄ (sample G in Figure 1b).³³ This sample was marked by an almost complete disappearance of pre-edge signals below 530 eV in the O K-edge XAS data, with the exception of a small, broad peak centered at 529.3 eV and little change above 530 eV compared to sample F (Figures 2b and S2), indicative of the consistency with changes in pDOS (Figure S3). The spectrum significantly broadened at higher energy, around the absorption edge, with a redshift of the inflection point by ~0.6 eV in comparison with the previous state (Figure S4).

The O K-edge XAS of the electrode harvested at 1.3 V after being charged to x = 1 (sample G) was almost perfectly equal to the spectrum of an electrode obtained by direct reduction of pristine Li₃IrO₄ to 1.3 V (sample J, Figures 2b, 2c and S2), further reinforcing the notion that the initial oxidation to Li₁IrO₄ is extremely reversible both from the electrochemical capacity and chemical perspectives. This reversibility is further demonstrated by the highly similar spectra of

pristine Li₃IrO₄ and the sample re-charged to 3 V after a full cycle of oxidation to x = 1 and reduction to 1.3 V (Figures 2b, 2d and S2). These observations emphasize the extremely high reversibility of the redox reaction between Li₁IrO₄ and Li_{4.7}IrO₄.

Significant evolution of the electronic structure between Li₃IrO₄ and Li_{4.7}IrO₄ upon lithiation is also reflected in magnetization data obtained on the samples directly reduced from the pristine state. The reduction process induced an increased paramagnetism from $\mu_{\rm eff} = 0.82 \ \mu_{\rm B}/{\rm Ir}$ in the pristine state (sample A, Li₃IrO₄, Figure 3a), to $\mu_{eff} = 1.46 \,\mu_B/Ir$ for Li₄IrO₄ (sample I, Figure 3b). This value is close to reports for Ir^{4+} (d⁵) compounds such as Li₂IrO₃ or Sr₂IrO₄ (1.7-2.0 μ_B/Ir). ⁶², ⁶³ The magnetization of 5d materials, such as Ir compounds, is affected by the competition between spin-orbit coupling interaction and electron-electron correlation, the main consequence of which is lifting the degeneracy of the t_{2g} manifold into effective total angular momentum $J_{eff} = 3/2$ with low energy and $J_{\text{eff}} = 1/2$ with high energy. This is reflected in the weak paramagnetism ($\mu_{\text{eff}} = 0.2$ - $0.9 \mu_B/Ir$) of Ir^{5+} (d⁴) compounds, corresponding to four paired electrons in the $J_{eff} = 3/2$ levels and a J = 0 ground state, instead of the S = 1 ground state expected for a perfectly octahedral environment in the absence of spin-orbit coupling. In the case of Ir^{4+} (d⁵) compounds, the J_{eff} = 3/2 level is filled, and one electron sits in the $J_{\rm eff} = \frac{1}{2}$ level, leading to a $J = \frac{1}{2}$ ground state, thus explaining the increased moment found for Li₄IrO₄ compared to Li₃IrO₄. In contrast, further lithiation to Li_{4.7}IrO₄, obtained upon complete reduction to 1.3 V, corresponds to a decrease of μ_{eff} to 0.52 µ_B/Ir (Figure 3c). This observation is consistent with a large proportion of Ir³⁺ (d⁶) with filled t_{2g} manifold ($J_{eff} = 3/2$ and $\frac{1}{2}$), corresponding to a J = 0 ground state.^{64, 65}

The changes below 3 V are accompanied by a steady shift of the Ir L_{III}-edge XANES to lower energy reported by Perez et al.³³ Therefore, the data support the injection of electrons into unoccupied O 2p-Ir 5d states following a conventional redox mechanism that can be explained by

changes in the formal oxidation state of Ir as a proxy for the Ir-O covalent bond. The pDOS (Figure S3) from DFT calculations clearly shows the decrease of a peak above Fermi level corresponding to the Ir d_{t2g}-O 2p states upon reduction, reflecting that electron mainly populated states arising from π interactions (t_{2g} in an ideal O_h field). This conclusion is further confirmed by the projection of calculated XAS on the two crystallographic oxygen sites (Figure S5) where the decrease in the two pre-edge peaks below 530 eV is due to the electron population of Ir d_{t2g}-O 2p states. The preedge peaks disappear completely after the composition reaches Li_{4.5}IrO₄ or higher. It should also be noted that after reaching the composition of Li₅IrO₄, the Ir d_{t2g}-O 2p states are filled and no longer observed above the Fermi level which is consistent with decrease in the pre-edge peaks in the experimental O K-edge XAS. It is possible that the smaller reduction of the pre-edge above 530 eV reflected a change in the contribution of O to the σ (e_g) states due to a decreased covalence of the Ir-O bond deduced by the simultaneous increase of Ir-O bond distance resulted from the accommodation of additional charge³³ (Table S1-1) and the calculated atomic Bader charge (Table S1-2). This distinct variation in the intensity reflects a greater role of unoccupied states at the lowest energy, in a behavior reminiscent of a rigid band. The redshift of the main absorption threshold indicates a decrease in the binding energy of the O 1s core level, consistent with repopulation with electrons, and decrease in the $Z_{\text{eff}}\,\text{of}\,O.$

4. Conclusions

Li₃IrO₄ is capable of undergoing consecutive reactions based on unconventional lattice oxygen redox, between Li₁IrO₄ and Li₃IrO₄, followed by a transition to reach Li_{4.7}IrO₄ following a mechanism based on the changes of the formal oxidation state of Ir as a proxy for state with both Ir and O due to the high covalency. All along, O states participate in the redox compensation, with different involvement of Ir and, thus, different changes in electronic structure. Unlike the complex

changes between Li₁IrO₄ and Li₃IrO₄, the filling of states between Li₃IrO₄ and Li_{4.7}IrO₄ is highly reminiscent of a rigid model. Remarkably, the 3.7-electron whole process can be reversed completely to restore the pristine Li₃IrO₄. This report emphasizes the role of ligand oxygen in compensating the charge associated with (de)lithiation via manipulating its covalent interaction with transition metals. Should this manipulation be mimicked for oxides with 3d metals, combined processes of lattice oxygen redox and cationic redox could be achievable in a similarly sequential manner, resulting transformational amounts of charge storage in a single compound.

Acknowledgements

This material is based upon work supported by the National Science Foundation under Grant No. DMR-1809372. This research used resources of the Advanced Photon Source, a U.S. Department of Energy (DOE) Office of Science User Facility operated for the DOE Office of Science by Argonne National Laboratory under Contract No. DE-AC02-06CH11357. Part or all of the research described in this paper was performed at the Canadian Light Source, a national research facility of the University of Saskatchewan, which is supported by the Canada Foundation for Innovation (CFI), the Natural Sciences and Engineering Research Council (NSERC), the National Research Council (NRC), the Canadian Institutes of Health Research (CIHR), the Government of Saskatchewan, and the University of Saskatchewan.

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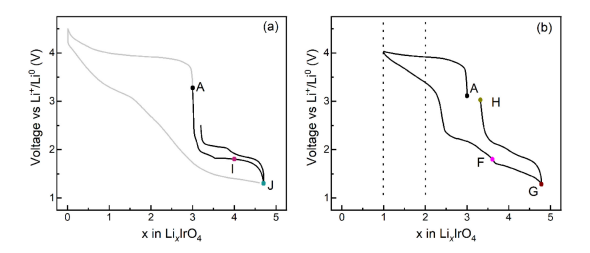


Figure 1. (a) and (b) Voltage-composition profiles of Li₃IrO₄ at different cutoff voltage windows. The electrochemical states of interest are represented with labels of different colors. Samples were harvested from the representative states. A is the pristine state. Cycled samples were obtained after initial oxidation to x = 1 and reduction to 1.8 V (F); reduction to 1.3 V (G); reduction to 1.3 V and re-oxidation to 3.0 V (H). Sample I was obtained by a direct reduction to x = 4 from the pristine state and sample J was obtained by a direction reduction to 1.3 V from the pristine state.

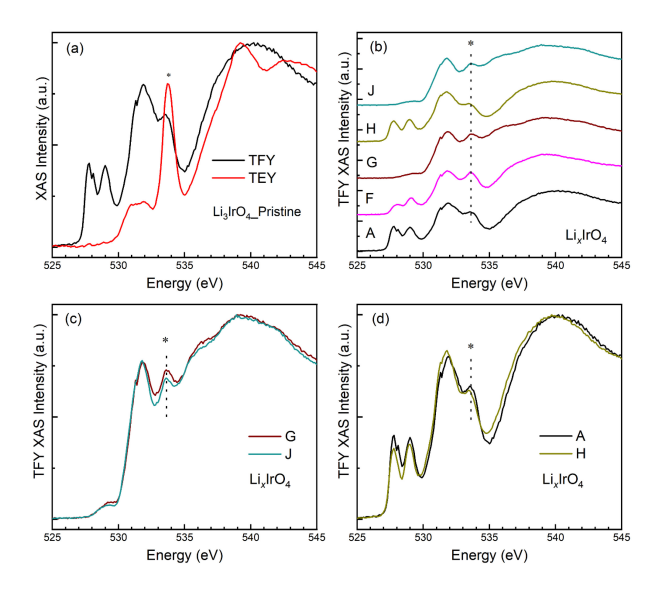


Figure 2. (a) O K-edge XAS spectra of Li₃IrO₄ pristine state measured under different modes. TFY: total fluorescence yield. TEY: total electron yield. The TFY spectrum is dominated by a peak at ~533.7 eV, which is attributed to the transition to π^* (C=O) orbitals in Li₂CO₃ impurity formed mainly in the surface. The minor presence of Li₂CO₃-related peak in TFY also suggests some existence of Li₂CO₃ below the surface. Due to the significant existence of Li₂CO₃ in the surface through all different electrochemical states, O K-edge XAS spectra from TFY were the focus of our work. (b) *Ex situ* O 1*s* XAS spectra of Li_xIrO₄ at difference electrochemical states of charge measured under TFY mode. (c) and (d) Overlaid O 1*s* XAS spectra of sample G and sample

J, sample A and sample H, respectively. The peak at around 533.7 eV (denoted by the asterisk symbol) is related to carbonate-related peak due to the transitions to π^* (C=O) orbitals, which has little influence on the spectral interpretation.

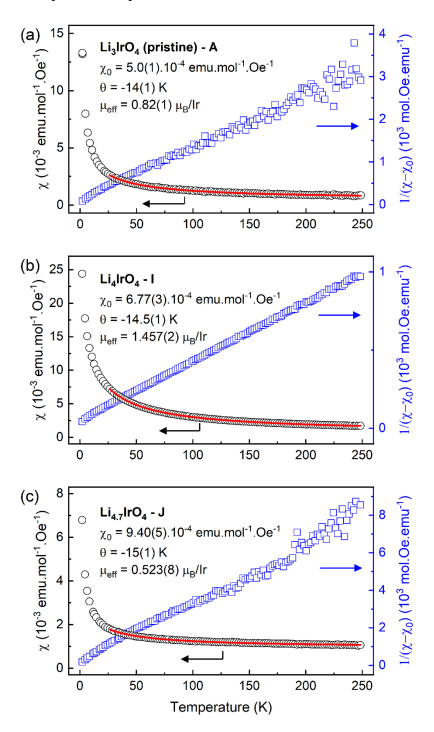


Figure 3. Magnetization measurement of Li_xIrO_4 with different Li contents at x=3 (a), x=4 (b), and x=4.7 (c) was performed *ex situ*. The black circles are zero-field cooled magnetic susceptibility measurements, the red line is obtained from fitting the data with a modified Curie-Weiss law, with the fitting parameters indicated, and the blue circles are the inverse magnetic susceptibility after removing the temperature independent contribution χ_0 .

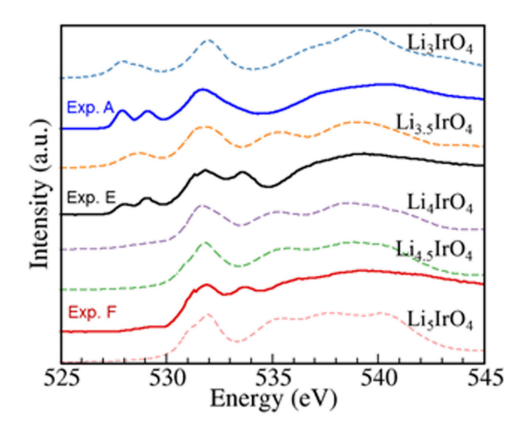


Figure 4. Simulated O K-edge XAS spectra of different electrochemical states and the corresponding experimental results. The XAS projected on the O_{3LP} and O_{2LP} absorbing species are shown in Figure S4. The peak at ~534 eV in the experimental data is ascribed to the presence of Li₂CO₃ impurities.