1 **Metal-Oxygen Decoordination Stabilizes Anion Redox in Li-rich Oxides**

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33 **Abstract**:

34 Reversible high voltage redox chemistry is an essential component of many electrochemical 35 technologies, from (electro)catalysts to lithium-ion batteries. Oxygen anion redox has garnered 36 intense interest for such applications, particularly lithium ion batteries, as it offers substantial 37 redox capacity at > 4 V vs. Li/Li⁺ in a variety of oxide materials. However, oxidation of oxygen 38 is almost universally correlated with irreversible local structural transformations, voltage 39 hysteresis, and voltage fade, which currently preclude its widespread use. By comprehensively 40 studying the Li2-*x*Ir1-*y*Sn*y*O3 model system, which exhibits tunable oxidation state and structural 41 evolution with *y* upon cycling, we reveal that this structure-redox coupling arises from the local 42 stabilization of short ~ 1.8 Å metal-oxygen π bonds and ~ 1.4 Å O–O dimers during oxygen 43 redox, which occurs in $Li_{2-x}Ir_{1-y}Sn_vO_3$ through ligand-to-metal charge transfer. Crucially, 44 formation of these oxidized oxygen species necessitates the decoordination of oxygen to a single 45 covalent bonding partner through formation of vacancies at neighboring cation sites, driving 46 cation disorder. These insights establish a point defect explanation for why anion redox often 47 occurs alongside local structural disordering and voltage hysteresis during cycling. Our findings 48 offer an explanation for the unique electrochemical properties of lithium-rich layered oxides, 49 with implications generally for the design of materials employing oxygen redox chemistry.

51 **Main Text:**

52 Reversible redox chemistry in solids under highly oxidizing conditions (e.g. vs H_2/H^+ , Li/Li⁺, or 53 O₂) is a powerful tool in (electro)chemical systems, increasing the catalytic activity of oxygen-54 evolution and methane-functionalization (electro)catalysts as well as the energy and power 55 densities of lithium-ion batteries $(LIBs)$.¹ In LIBs in particular, employing high-voltage redox 56 has been identified as a promising avenue to meeting the energy density demands of next-57 generation technologies such as plug-in electric vehicles.

58 Recently, anionic oxygen redox has been shown to offer access to substantial high-voltage 59 (de)intercalation capacity in a range of electrode materials,²⁻⁷ spurring an intense research effort 60 to understand this phenomenon. While many oxygen-redox-active materials have been 61 developed, they almost universally exhibit a host of irreversible electrochemical behaviors such 62 as voltage hysteresis and voltage fade.⁸ This is most notable in the anion-redox-active Li-rich 63 layered oxides, $Li_{1+x}M_{1-x}O_2$ (M = a transition metal (TM) or non-transition metal such as Al, Sn, Mg, etc.), which exhibit capacities approaching 300 mAh g^{-1} but have yet to achieve commercial 65 success due to such electrochemical behaviors.^{5, 9} It has been shown both experimentally¹⁰⁻¹² and 66 from first-principles thermodynamics¹³ that the migration of M into empty Li sites⁹ – creating 67 structural disorder in the form of M_{Li}/V_M antisite/cation vacancy point defect pairs – is at the root 68 of voltage profile evolution and depression, particularly during the first cycle. Oxygen redox has 69 separately been shown to trigger voltage hysteresis and sluggish kinetics, $8,14$ with the implication 70 being that oxygen redox and transition metal (TM) migration/vacancy formation are often 71 intrinsically linked. Indeed, the coupling between oxygen redox, cation migration, and voltage 72 hysteresis was recently observed.⁶ However, under the two primary models that have emerged to describe the nature of the lattice oxygen in these materials (either $a \sim 2.5$ Å peroxo-like O_2^{n-1}

74 dimer^{3, 15} or an isolated O⁻ anion^{4, 5, 16}), it is unclear as to why the oxidation of oxygen should 75 lead to the correlated structural transformations and voltage hysteresis. It has been argued 76 qualitatively that the oxidation of oxygen destabilizes the layered structure; $2,15$ however, a robust 77 understanding of the materials design criteria for achieving structurally and electrochemically 78 reversible anion redox remains elusive.

79 In approaching this issue, we turn to layered $Li₂IrO₃$ (equivalently, $Li[Li_{0.33}Ir_{0.67}]O₂$ or LIO)

80 which is reported to exhibit anion redox, yet, unlike other Li-rich layered oxides, exhibits highly

81 reversible structural and electrochemical behavior during cycling.^{15, 17} Upon substituting redox-

82 inactive Sn for Ir, the solid solution material $Li_2Ir_{1-y}Sn_vO_3$ (LISO) forms M_{Li}/V_M defects during

83 charge, accompanied by voltage hysteresis during subsequent cycling that is typical of other 84 anion-redox-active oxides.¹⁵ This tunability makes LIO/LISO an ideal model system for studying 85 the link between anion redox, local structure, and irreversible electrochemistry.

86 In this work, we show that the difference in electrochemical and structural behavior between LIO 87 and LISO holds the key to understanding anion redox. Using a combination of bulk-sensitive 88 spectroscopic and structural probes in conjunction with first principles calculations, we 89 demonstrate that anion redox only occurs simultaneously with structural disordering (i.e. 90 antisite/vacancy formation) in LISO, and is completely absent in LIO up to 4.6 V, further 91 establishing the strong coupling between these two phenomena. We demonstrate that the \sim 2.5 Å 92 O–O dimers that form in LIO do not constitute oxidized oxygen, and that the high degree of 93 covalency and the low voltage of the Ir redox states together prevent oxygen redox from being 94 accessed in LIO. Crucially, we reveal the origin of the coupling between structural evolution and 95 oxygen redox in LISO using density-functional theory (DFT) calculations, wherein the low 96 valence electron count in delithiated LISO promotes charge transfer from O to Ir (essentially,

97 oxygen oxidation *via* ligand to metal charge transfer, or LMCT) through drastic changes in the 98 local bonding configuration. We identify two main bonding configurations that are able to 99 stabilize both the low valence electron count through LMCT and the resulting oxidized oxygen 100 species: short ~ 1.8 Å Ir-O π bonds (equivalently, terminal Ir=O oxo species) and short ~ 1.4 Å 101 O–O dimers. Both configurations require the decoordination of oxygen down to a single covalent 102 bonding partner, which is realized in the layered structure through vacancy formation due to in-103 plane and out-of-plane site disorder (cation migration). Meanwhile, the higher valence electron 104 count in LIO disfavors LMCT and thus LIO exhibits minimal structural disordering, cation redox 105 only, and more reversible electrochemistry. While further investigation is needed to understand 106 how this mechanism depends on the nature of the TM, this point defect model can offer a causal 107 explanation for the nearly universal observation of structural and electrochemical irreversibility 108 in anion-redox-active Li-rich materials and reveals previously unconsidered design parameters 109 through which to improve them, such as the electronic and structural mechanism of LMCT and 110 oxygen decoordination. These principles apply generally to the design of the broad class of 111 highly oxidized oxides where anion redox is likely to play a role.

112

113 **Electrochemistry and phase behavior of Li2IrO3**

114 As previously reported,^{15, 17} Fig. 1a shows that LIO exhibits two well-defined voltage plateaus 115 with average potentials of 3.45 V and 4.15 V versus $Li/L_i⁺$ and a total capacity of 1.5e⁻ per Ir. It 116 was previously shown that LIO exhibits a layered structure with monoclinic *C2/m* symmetry, 117 with the Li and TMs in the TM layers ordered into a honeycomb-like arrangement, and 118 accommodates delithiation through changes in the stacking of the TM layers, changing from 119 initially O3 to $O1^{18}$ at the end of charge. This is further supported here by Rietveld refinement of

120 the synchrotron X-ray diffraction (XRD) patterns (figs. S1 & S2, tables S1 & S2, and

121 Supplementary discussion S1). In addition, we identify the intermediate $Li₁IrO₃$ phase as

122 exhibiting T3 stacking using DFT (Fig. S3). We note that the material studied here exhibits a

123 high degree of initial structural ordering and electrochemical reversibility, and is therefore more

124 similar to that reported by Kobayashi et al.¹⁷ than McCalla et al.¹⁵

125 While it was previously shown that LIO exhibits minimal structural disordering during the first 126 charge, for this study it is important to quantify the cation vacancy/antisite formation during 127 electrochemical cycling such that it can be compared to LISO. We employed an iterative 128 approach to simultaneously refine the in-plane and out-of-plane site disorder as described in 129 Supplementary discussion S1 and Fig. S4. This approach eliminates the difficulty in site 130 occupancy determination resulting from the presence of stacking faults. We confirm the absence 131 of V_{Ir}/Ir_{Ii} defects ($\leq 0.1\pm 0.2\%$ of all Li sites occupied by Ir) in the discharged state both before 132 and after the first electrochemical cycle. In-plane site occupancy refinement of the fully charged 133 O1 structure is challenging due to distortions to the oxygen sublattice¹⁵ and the absence of 134 superstructure peaks. However, out-of-plane refinement shows negligible Ir occupancy in the Li 135 layer in the fully charged state. Consistently, DFT calculations indicate that in-plane and out-of-136 plane Ir antisite/vacancy defect pairs in the fully charged state have a positive formation energy 137 of +0.49 and +0.20 eV (per Li₄Ir₈O₂₄ supercell), respectively. Together, these results show why 138 LIO exhibits reversible electrochemistry: minimal cation disordering occurs during cycling. This 139 is unlike most other Li-rich materials in which anion redox is reported to occur.^{7, 9, 12, 19}

140

141 **Multivalent iridium redox in Li2IrO3**

142 The nature of anion redox in LIO should therefore be examined with further scrutiny. In 143 addition, a strict definition of anion redox is needed. Until now, oxygen redox has been defined 144 as the depopulation of non-bonding (or "weakly/minimally hybridized") oxygen 2p states^{5, 6, 16, 20,} 145 21 . However, this definition does not take into account how materials respond to the depopulation 146 of these states, which as we will show later is a crucial component of oxygen redox. So, we 147 provide a more specific definition here: oxygen redox requires the depopulation of non-bonding 148 (or "weakly/minimally hybridized") oxygen *2p* states wherein the resulting holes reside in 149 predominantly oxygen character orbitals after any structural and electronic reorganization has 150 occurred. A previous study reported anion redox in LIO based on: (a) surface-sensitive X-ray 151 photoelectron spectroscopy (XPS) showing the appearance of an oxygen state with reduced 152 electron density, and (b) neutron diffraction and transmission electron microscopy (TEM) 153 measurements showing the contraction of the octahedral O–O distance from \sim 2.7 Å to \sim 2.5 Å.¹⁵ 154 Here we use bulk-sensitive X-ray spectroscopic techniques to directly determine the nature of the 155 hole states formed during charge and, by induction, the redox processes in LIO. We first show in 156 Fig. 1 analysis of the *operando* transmission X-ray absorption near edge structure (XANES) at 157 the Ir L₃ edge during the first cycle of LIO.^{19, 22, 23} Figure 1b shows that the white line (WL) 158 energy increases linearly during the galvanostatic charge up to $Li_{0.5}IrO₃$ and then decreases 159 linearly on discharge, nearly returning to its original value after the full cycle. Figure 1c, 160 meanwhile, shows that the WL intensity increases up to $Li₁IrO₃$ but then decreases when 161 charging further to $Li_{0.5}IrO₃$, with the trend reversing on discharge. Principal component analysis 162 and non-negative matrix factorization (NMF) of the XANES spectra (Supplementary discussion 163 S2 and Fig. S5) reveal the presence of three spectral end members corresponding to the $Li₂IrO₃$, 164 Li₁IrO₃, and Li_{0.5}IrO₃ compositions, consistent with the presence of two isosbestic points. These

165 three end member spectra (Fig. 1d, inset) reflect the observed changes, with $Li_{0.5}IrO_3$ having the

166 highest WL energy followed by Li_1IrO_3 and Li_2IrO_3 , and with Li_1IrO_3 having the strongest WL

167 peak intensity. A linear combination analysis (LCA) using these end members confirms that the

168 entire spectral evolution is well described by two consecutive two-phase reactions between

 169 Li₂IrO₃ and Li₁IrO₃, and between Li₁IrO₃ and Li_{0.5}IrO₃ (figs. 1d and S5).

170 To understand what these spectral changes indicate, we note that the L₃ WL energy has been

171 shown to vary linearly with formal Ir oxidation state.³⁴ Thus the linear trend during galvanostatic

172 cycling indicates a linear, monotonic change in Ir oxidation state, with a total WL shift of 1.2 eV.

173 The Ir WL energy difference between Ir^{6+} and Ir^{4+} compounds is typically between 1.4-2.0 eV,^{23,}

 2^{24} and the WL shift here is therefore consistent with Ir being oxidized by between 1.2-1.7 e⁻.

175 Although it was argued that a loss in WL intensity indicates Ir reduction in $Li₃IrO₄$,¹⁹ it has been

176 shown that the WL intensity of Ir⁶⁺ is lower than that of Ir⁵⁺,^{23, 24} which may be due to the WL

intensity becoming dominated by relaxation quenching rather than d count for d counts below d^4 -

178 $d^{5.25}$ Thus both the WL energy and intensity suggest a monotonic oxidation of Ir from 4+ to

179 between 5.2+ and 5.7+.

180 Since the linear coefficient between WL energy and oxidation state appears to vary slightly

181 between materials, 2^{3} , 2^{4} it is impossible to precisely quantify the final Ir oxidation state based on

182 XANES studies in other materials. Thus, although the monotonicity of the Ir oxidation is already

183 in striking contrast to the behavior of most anion redox active Li-rich oxides, whose TM ions

184 either stop being oxidized or are even reduced during the high voltage plateau, $4, 6, 16$ and suggests

185 the absence of anion redox in LIO and a final Ir oxidation state of 5.5+, this cannot be proven

186 with XANES alone. To directly assess the contribution of bulk O to the LIO redox mechanism,

187 we perform soft X-ray absorption spectroscopy (sXAS) at the O K edge. Transmission and

188 fluorescence yield (FY) detection modes with probing depths of hundreds of nm were employed 189 (see methods). Figure 2a shows the O K edge sXAS-FY and transmission sXAS spectra of LIO 190 (initially in the $t_{2g}^5 e_g^0$ electronic configuration) during the first electrochemical cycle. The peaks 191 at ~529 eV and 532 eV are ascribed to the unoccupied Ir 5*d*-O 2*p* t_{2g} and e_g^* hybridized states, 192 respectively.^{26, 27} The difference plots taken between different states of charge (Fig. 2b) clearly 193 show that the intensity of the *t2g* peak increases and decreases throughout charge and discharge, 194 mirroring the evolution of the Ir L3 WL (Fig. 1c). Figure 2c shows that the integrated area ratio 195 of the two O K XAS pre-edge peaks follows the same trend in transmission sXAS, which probes 196 the entirety of the particles, confirming that this is a bulk phenomenon. 197 The correlated changes at the O K and Ir L₃ edges indicate redox of a hybridized Ir–O state, and 198 are widely accepted as a fingerprint of conventional "TM–O" cation redox⁶ observed in a variety 199 of compounds such as $\text{LiNi}_{0.5}\text{Mn}_{1.5}\text{O}_4$,²⁸ LiFePO₄,²⁹ 3d layered TM oxides.^{6, 26} While in these 200 systems the oxidation of the TM is accompanied by charge sharing with the anions, this clearly 201 does not qualify as oxygen redox according to the earlier definition since no *additional* 202 unhybridized O *2p* electrons are depopulated beyond the hybridized TM–O states. This is further 203 supported by resonant inelastic X-ray scattering (RIXS) at the O K edge. It has recently been 204 shown that depopulation of unhybridized O *2p* states in some Li-rich materials results in a sharp 205 absorption feature at \sim 530.8 eV excitation energy at the O K edge with a characteristic emission 206 at \sim 523.0 eV, evident in both RIXS maps⁶ and single-energy RIXS spectra, ^{6, 16, 30, 31} which is 207 distinct from the energy of typical hybridization features in most TM oxides. Figure 2d shows, 208 however, that no such feature appears even in the fully charged LIO. Thus, in conjunction with 209 the XANES results, O K edge sXAS and RIXS further support pure multivalent, hybridized Ir–O 210 redox in the absence of oxygen redox in LIO.

211 This is also supported by DFT calculations. The predicted projected density of states (pDOS) of 212 the oxygen *2p* and Ir *5d* states of the three structures formed during cycling are shown in Fig. 2e. 213 Changes in the Ir and O pDOS as a function of Li stoichiometry show that all states depopulated 214 during delithiation contain both Ir and O character. The unhybridized O *2p* states (*i.e.*, the non-215 bonding (NB) O 2p band) lie \sim 2 eV below the Fermi level in Li_{0.5}IrO₃ and are therefore not 216 emptied. Insets in Fig. 2e plot the isosurface of the spin density of the lowest unoccupied (*i.e.*, 217 redox active) states around Ir and O for $Li₁IrO₃$ and $Li_{0.5}IrO₃$. (equivalent to 0.5 electrons per 218 formula unit, shaded in the DOS). In both cases, we observe large spin density around both Ir 219 and O ions in the shape of two distinct hybridized Ir $5d$ -O $2p$ t_{2g} orbitals – likely d_{xy} and mixed 220 $d_{vz} - d_{vz}$. Thus, although all oxygen ions in LIO exhibit the linear Li–O–Li structural motif that 221 was recently shown to raise the energy of the O 2*p* states and promote oxygen redox in many Li-222 rich materials (see Fig. S6),^{5, 6, 32} these results demonstrate that the Ir^{4/5.5+} redox band (where 5.5 223 denotes the average formal oxidation state) is too high in energy for these labile O 2*p* states to 224 compete for redox in LIO, similar to predictions made for V, Cr, and Mo.⁵ 225 Having employed a suite of bulk-sensitive spectroscopic probes and computation, we show that 226 only hybridized TM–O states are accessed in LIO, with Ir reaching the 5.5+ formal oxidation 227 state at the end of charge. We therefore re-classify LIO as anion-redox inactive within the 2.5-4.6 228 V window. In terms of addressing previous conflicting conclusions, it is likely that the oxidized 229 oxygen species observed by XPS are due to near-surface phenomena which do not play a 230 significant role in the bulk redox mechanism. As for the long \sim 2.5 Å O–O dimers that were 231 observed to form in the bulk of charged LIO, we conclude that these species are not sufficient 232 evidence of oxidized oxygen. This is based on the observation that most TM-coordinated

233 oxidized oxygen dimers are between 1.3-1.5 Å in length.³³ We speculate on alternative 234 explanations for the dimerization phenomenon in Supplementary discussion S3 and Fig. S7. 235

Extra capacity beyond Ir5.5+ 236 **in LISO and irreversible electrochemical/structural evolution** 237 When Sn is substituted into Li_2IrO_3 to make solid-solution phase $Li_2Ir_{1\rightarrow}Sn_vO_3$, the 238 electrochemistry changes dramatically and becomes typical of that of anion redox active 239 materials (Fig. 3a). We use the notation LISO## to represent the compound with ## % Sn 240 content, and the term "LISO" to refer to the Sn-substituted materials generally. Regardless of Sn 241 content, when charged beyond Ir^{5.5+}, LISO exhibits a new plateau at \sim 4.35 V, followed by a 242 large voltage drop on the subsequent discharge, a permanently sloped voltage profile, and 243 voltage hysteresis that persists even at open circuit (Fig. S8). Despite the drastic changes to the 244 voltage profile, LISO25 exhibits reasonable capacity retention with cycling (74.5% after 192 245 cycles, Fig. S9). The new high voltage plateau, in conjunction with the additional reversible 246 capacity well beyond 1.5 e- per Ir (Fig. 3a), indicates the presence of a reversible redox partner 247 beyond $Ir^{4+}/Ir^{5.5+}$.

248 In addition to this high-voltage redox partner, the average and local structural behavior of LISO 249 differs from that of LIO. As first demonstrated by McCalla *et al.*,¹⁵ a substantial degree of cation 250 site disorder develops in LISO after first delithiation. Figure 3b quantifies the extent of site 251 disorder in LISO25 before and after electrochemical cycling with either a 4.25 or 4.60 V cutoff. 252 The iterative Rietveld refinement results are shown in Fig. S10 and tables S3 & S4. As shown 253 schematically in Fig. 3b, we assume that each in-plane and out-of-plane antisite defect results in 254 a corresponding vacant M site (V_M) . Since low concentrations of Sn_{Li} and Ir_{Li} have an identical 255 effect on the Bragg peak intensity, XRD cannot be used to differentiate between the migrating

256 species. We therefore perform the refinement assuming either all Ir_{Li} for both in-plane and out-257 of-plane defects or only Sn_{Li} for out-of-plane defects and Ir_{Li} for in-plane defects (assuming all 258 Sn_{Li} for both types of defect leads to full depletion of the Sn content in the M sites in LISO25). 259 We quantify the total amount of disorder by the fraction of V_M , which reaches between 260 8.97 \pm 0.9% (assuming all Ir_{Li} defects) and 12.0 \pm 1.4 % (assuming Sn_{Li} for out-of-plane defects 261 and Ir_{Li} for in-plane defects) after a single cycle between 4.60-2.50 V, approximately evenly split 262 between in-plane and out-of-plane antisite defects. Since the total accessed capacity is 263 approximately the same between LIO and the various LISO compositions, by substituting with 264 Sn we can conclude that the structural disordering is not due to global structural instability 265 caused by low lithium contents. Instead, the disordering appears to be associated with the new 266 redox partner giving rise to the high voltage plateau: the disorder in LISO25 after a full cycle is 267 dramatically greater than in both LIO after the first cycle and LISO25 after a single cycle 268 between 4.25-2.50 V, which are both zero within experimental error. Out-of-plane refinement of 269 the fully charged O1 structure shows that the disorder is increased substantially at 4.60 V (tables 270 S3 & S4), providing a direct connection between the new redox partner, electrochemical 271 irreversibility, and the onset of cation disordering. Increased cation disordering with increased 272 redox capacity beyond Ir^{4+/5.5+} is further supported by pair distribution function (PDF) analysis 273 (figs. S11-S14 and table S5) and EXAFS at the Ir L₃ edge (Fig. S15), which shows that the 274 decrease in scattering intensity of the first and second coordination shells during charge in LISO 275 – indicative of disorder in the atomic distances – is more substantial when the fraction of 276 capacity beyond Ir^{4+/5.5+} is greater (i.e. with increasing Sn content). The changes to the XRD 277 pattern in charged LISO50 (Fig. S16) are complex and make reliable quantification of 278 occupancies with Rietveld refinement challenging. However, refinement in the discharged state

279 (tables S6 & S7) and the PDF and EXAFS behavior are similar to, and show more extensive

280 disordering than, LISO25. As discussed later, DFT also consistently predicts more favorable in-

281 plane and out-of-plane cation migration in fully charged LISO compared to LIO, further

282 supporting cation disordering during charge.

283

284 **O redox switched on by Sn substitution**

285 The electrochemical and structural data confirm that Sn substitution turns on an additional high 286 voltage redox partner that is associated with M_{Li}/V_M defect formation and voltage hysteresis. To 287 further probe the nature of the high voltage LISO redox partner, we perform *operando* 288 transmission XAS at the Ir L_3 and Sn K edges and RIXS at the O K edge. Figures 3c and S17a 289 show that, as with LIO, charging to 4.25 V results in a shift to higher energy of the Ir L_3 WL 290 energy. This is correlated with an increase in intensity of the Ir *5d*-O *2p t2g* peak at the O K edge 291 (Fig. 3d), confirming that this is standard hybridized Ir–O redox. Strikingly, the Ir L_3 WL energy 292 is essentially unchanged when charging beyond 4.25 V, exhibiting only a slight decrease in the 293 peak intensity (Fig. S17c). As argued previously, the invariance of the Ir L_3 WL is indicative of a 294 constant oxidation state and thus we rule out continued hybridized Ir-O redox during the high 295 voltage plateau. Figure S17d shows that the Sn K edge absorption changes minimally during 296 charge in LISO25, ruling out the improbable scenario of Sn oxidation beyond 4+. 297 The constant Ir/Sn oxidation states during the high voltage plateau suggest O redox, and we 298 again turn to the O K edge XAS/RIXS for confirmation. Figure 3d shows that when charging

299 LISO25 from 4.25 to 4.60 V, the *t2g* peak continues to increase in intensity despite the invariance

300 of the Ir L_3 edge WL. This indicates an increased O 2p character in the t_{2g} band without the

301 depopulation of hybridized Ir–O states. Additionally, Fig. 3e shows that when charging to 4.60

302 V, a sharp RIXS feature at 530.7 eV excitation energy and 523.0 eV emission energy (indicated 303 by the white arrows) emerges. This is seen more clearly in the RIXS emission spectra in Fig. 3f. 304 The feature is strongest in LISO50, which exhibits the greatest extent of high-voltage capacity 305 beyond Ir^{4+/5.5+} (Fig. 3a) and antisite/vacancy defects. This RIXS feature has been observed to 306 closely follow oxygen redox activity in several battery electrodes.^{6, 31} Both of these behaviors 307 indicate that, in addition to promoting cation disordering, Sn substitution in LISO also switches 308 on O redox.

309

310 **Cation migration and O redox stabilized by LMCT** *via* **short Ir–O π bonding and O–O**

311 **dimerization**

312 Understanding why Sn substitution gives rise to these phenomena is central to uncovering the 313 origin of the coupling between anion redox and metal migration/vacancy formation. Since the 314 defect concentration is more reliably quantified in LISO25, we focus our DFT analysis on this 315 composition. First, we show in Fig. S18 the evolution of the computed DOS in the control case 316 of ordered, M_{Li}/V_M -free LISO25 as a function of Li stoichiometry. We observe that no 317 unhybridized O *2p* states are depopulated in the fully delithiated O1 structures – only hybridized 318 Ir–O states are accessed, giving Ir a formal oxidation state of at least 6+, which is inconsistent 319 with our XAS results. In fact, the behavior is similar to LIO (albeit more oxidized), with the 320 unhybrdizied O 2*p* states positioned ~1 eV below the Fermi level. Thus, simple delithiation and 321 the O3-O1 stacking change does not explain the high voltage redox process in LISO. 322 Importantly, our DFT calculations suggest that O redox *cannot* occur in LISO after delithiating 323 1.5 Li per formula unit if M_{Li}/V_M defects are not allowed to form.

324 Next, we introduce the experimentally-observed Sn_{Li}/V_M defects and examine their effect on the 325 electronic structure and charge distribution of LISO. Out-of-plane Sn migration into a Li layer 326 site creates a cation vacancy, V_M , and neighboring single-coordinate (dangling) oxygen atoms 327 (Fig. S19). Note that we use the term "single-coordinate" to refer to the number of covalent 328 bonding partners – i.e. counting Ir or Sn, but not Li. We find the formation of Sn_{Li}/V_M defect 329 pairs stabilizes the delithiated structure significantly by 1.34-1.36 eV, whereas the formation of 330 Ir_{Li}/V_M defect pairs is predicted to have a much smaller driving force of 0.02 eV. 331 We attribute the $\text{Sn}_{\text{Li}}/V_{\text{M}}$ stabilization to the response of the resulting M-O dangling bonds, 332 which undergo one of two major transformations. As shown in Fig. 4a, when $M = Ir$, these bonds 333 contract substantially, changing from \sim 2.10 Å to \sim 1.77 Å. Figure 4b shows that the bond 334 contraction accompanies a splitting of the previously unhybridized O *2p* states (shaded, top 335 panel), with some states moving above the Fermi level and becoming oxidized (shaded, bottom 336 panel). The shifted states, now approximately 1.5 eV higher in energy, rehybridize with the Ir *5d* 337 states. Table S8 shows that the bond contraction results in a donation of charge from the 338 dangling oxygens to the Ir bonding partners. Crystal orbital overlap population (COOP) analysis 339 in Supplementary discussion S4 and Fig. S20 shows that the net bond order between the dangling 340 O and Ir ions increases by 45 % as a result. Together, these observations suggest the formation of 341 short Ir–O π bonds (equivalently, terminal oxo ligands) through donation of oxygen lone pair 342 electrons, approaching Ir=O double bonds although an exact bond order is hard to define in an 343 electronically delocalized crystal. Indeed the substantially shortened Ir–O bonds $(1.76 - 1.79 \text{ Å})$ 344 are similar in length to the Ir=O double bond observed in tetrahedral oxotrimesityliridium(V).³⁴ 345 While short Ir–O π bonding in octahedral environments has not been previously observed in

346 molecular compounds, the low effective Ir⁶⁺ *d* count $(d³)$ makes such bonding possible in 347 charged LISO. 35

 348 When M = Sn, on the other hand, the single-coordinate oxygens are predicted to instead pair up 349 to form short (\sim 1.44 Å) O–O dimers that straddle the V_M (Fig. 4c and Fig. S19). Again, this 350 transformation results in a shift of the previously buried NB O 2*p* states from below to above the 351 Fermi level (Fig. 4d) along with donation of electron density from O to a neighboring Ir (table 352 S9). We stress that these localized dimers are distinct from the long (\sim 2.5 Å) dimers previously 353 proposed for this material¹⁵ in that they are well within the range of typical bond lengths for $O_2^{\ n-1}$ 354 species³³ and, crucially, do not form in the absence of cation migration. These behaviors – the 355 stabilization of the charged structure through Sn migration and Ir=O/O–O formation – are also 356 predicted for LISO50 (Fig. S21). We furthermore note that the computational conclusions are 357 independent of the functional employed (table S10).

358 We note that when both oxygens neighboring the V_M are coordinated to an Ir ion, the short Ir–O 359 π bonds are generally predicted to be more favored than O-O dimers (Fig. S21 and table S10). 360 However, as the Sn content is increased, the fraction of Sn-Sn neighbors increases and thus so 361 does the fraction of Sn-coordinated dangling oxygens after TM migration/vacancy formation, 362 which favor O–O formation (Fig. S21 and table S11). This observation may be due to the 363 increased rotational freedom of the Sn–O bond, facilitating O –O dimerization,⁵ as well as the 364 ability of TMs with low d counts (such as $Ir^{5+/6+}$) to adopt terminal oxo ligands in octahedral 365 environments.³⁵ In addition, in-plane M_{Li}/V_M defect pairs can give rise to the same behavior (Fig. 366 S22).

367 Both the V_M-mediated O–O dimer and short Ir–O bond models are consistent with the local 368 structural evolution measured through EXAFS and PDF. EXAFS (Fig. S15) shows a substantial

369 decrease in the Ir–O scattering intensity during charge in LISO, which is exacerbated with

370 increasing oxygen redox capacity, consistent with an increased range of Ir–O bond lengths.

371 Meanwhile PDF (Fig. S14) shows a broadening of the M–O nearest neighbor scattering peak in

372 charged LISO, with an increase in scattering intensity at lower radial distance, consistent with

373 the formation of bonds shorter than the octahedral M–O distance of \sim 2 Å.

374 Additionally, both the O–O dimer and short Ir–O bond mechanisms are supported by the

375 spectroscopy data. Figure 4b predicts the donation of the buried O 2*p* electrons to the over-

376 oxidized Ir ions by raising these buried states above the Fermi level and into the t_{2g} band,

377 preventing Ir from exceeding the 5.5+ oxidation state. This is consistent with the increase in the

378 O K edge t_{2g} peak intensity (Fig. 3d) in the absence of a shift in the Ir L₃ WL energy during

379 oxygen redox (Fig. 3c). Meanwhile, the sharp RIXS feature is associated with specific

380 excitations in highly oxidized states, especially those involving unoccupied O 2*p* states in

381 oxidized oxygen species. Both peroxides³⁶ and O_2 gas³⁷ display the same kind of RIXS feature as

382 do charged Li-rich 3*d* layered oxides.^{4, 6, 16, 31} Consistently, the formation of O–O dimers presents

383 a mechanism through which the buried O 2*p* states may become unoccupied during delithiation

384 in LISO. While our proposal of O-O is consistent with the RIXS result, we cannot rule out the

385 possibility that different types of oxidized oxygen species may also give rise to a similar feature.

386 Under our earlier definition, the formation of O_2^2 dimers qualifies as oxygen redox since, as

387 shown in Fig. 4, O 2*p* states are shifted above the Fermi level (i.e. depopulated) and end up as

388 the peroxide σ^* , which has mostly oxygen character. Thus, this mechanism serves to localize

389 positive charge onto O 2*p* orbitals even in an otherwise highly covalent system. However, in the

390 case of forming short Ir–O π bonds, while the O 2p states are again shifted above the Fermi level,

391 they end up rehybridizing with Ir *t2g* states. In this case, although this results in additional

392 capacity beyond Ir^{4+/5.5+}, the character of these states, and thus the extent to which this counts as 393 oxygen redox, depends on the TM–O hybridization. While the hybridization may be significant 394 in the LISO materials, the nature of such states could be different in less covalent materials, such 395 as 3*d* TM oxides, and could therefore be important for oxygen redox in other systems.

396 To understand why LISO undergoes cation migration and oxygen redox while LIO does not, we 397 show in Supplementary discussion S5 that the driving force for forming Sn_{Li}/V_M defect pairs 398 depends strongly on the electron count in LISO, becoming favorable only when this falls below $1.5 e⁻$ per Ir less than that of the pristine material. Thus, it is clear that the driving force for this 400 structure-redox coupling in LISO is the over-oxidation of Ir beyond an average 5.5+ oxidation 401 state. It is also clear that in the two structural transformations described in Fig. 4 (Ir=O and O–O) 402 substantial donation of charge occurs from O to Ir. It has been shown that in Na-ion battery 403 compounds unusually high oxidation states such as $Fe⁴⁺$ and $Cu³⁺$ are stabilized by LMCT, 404 wherein a substantial donation of charge from oxygen to the metal center occurs through strong 405 $\,$ M–O covalency and low charge transfer energy.³⁸ We therefore propose that when oxidizing 406 beyond Ir^{5.5+} a sufficiently low charge transfer energy is achieved such that substantial LMCT is 407 promoted and, consequently, new structural motifs and bonding configurations that allow for 408 greater LMCT become stabilized. In the case of Ir=O, the Ir–O bond length is substantially 409 shortened such that the covalency and LMCT is increased (as is clear in figs. 4b and S20 and is 410 well known, for example, for the Mn=O bonds in $KMnO₄³⁹$, while in the case of O–O electrons 411 are directly transferred from the $O-O\sigma^*$ to neighboring Ir ions. Figure 5 shows the stepwise 412 process schematically: over-oxidation of Ir through delithiation, followed by Sn migration to 413 create either Ir=O or O–O species with a resulting donation of charge back to Ir, with the net 414 result being oxidation of O. Since the predicted O–O dimer bond length is 1.44 Å, we depict

415 these dimers as peroxide species in Fig. 5. Since the average Ir oxidation state never exceeds 416 5.5+ in LIO, substantial LMCT is not promoted, and therefore LIO exhibits excellent retention of 417 its in-plane and out-of-plane cation order during cycling and no anion redox.

418 Thus, the single-coordinate oxygens simultaneously stabilize both the low valence electron count 419 in charged LISO through LMCT and the resulting oxidized oxygen species, whether they settle 420 as terminal oxo ligands or O–O dimers. Consequently, they promote and stabilize the formation 421 of $\text{Sn}_{\text{Li}}/V_{\text{M}}$ defects. Importantly, this structure-redox coupling quantitatively accounts for the 422 anion redox capacity. The diffraction results give a V_M concentration in LISO25 of 12.0 \pm 1.4% 423 assuming out-of-plane Sn_{Li} defects and in-plane Ir_{Li} defects. This extent of disordering is 424 accompanied by an additional redox capacity of \sim 0.48 Li beyond Ir^{4+/5.5+}. Assuming each 425 decoordinated oxygen contributes one redox electron, this implies that on average \sim 4 oxidized 426 oxygens are generated per migrated atom. Each in-plane and out-of-plane migration pathway 427 studied here generates 4 dangling oxygens, in agreement with this value.

428 We note that the spectroscopic and structural signature of oxidized oxygen in LISO is strikingly 429 similar to that in 3d and 4d Li-rich layered oxides.^{6, 31} Further work is needed to establish the 430 precise commonalities between 3*d*, 4*d*, and 5*d* Li-rich electrode materials, in particular the 431 mechanism of decoordination and the relative driving force for forming short M–O π bonds 432 versus short \sim 1.44 Å O–O dimers. However, this initially suggests that depopulation of the high-433 energy O *2p* states lying along the linear Li-O-Li axes of the honeycomb environments in Li-rich 434 layered oxides is generally stabilized through antisite/vacancy formation, short M–O π bonding, 435 and/or short O–O dimerization. Indeed, ab initio molecular dynamics studies have predicted such 436 mechanisms in $3d$ Li-rich materials.⁴⁰ This also means that the same two-M-coordinate Li-O-Li 437 environments (where M is a covalent, non-alkali cation) that promote oxygen redox by raising

438 the energy of the O 2*p* states also facilitate its stabilization, as only a single M-Li site swap is 439 required to reach single-M-coordinate. In systems such as LISO where these O *2p* states are not 440 sufficiently high lying to be depopulated in the disorder-free state, O redox can still occur if the 441 valence electron count is depleted low enough for substantial LMCT to be promoted via the 442 formation of antisite/vacancy defects, donating the otherwise inaccessible O *2p* electrons to a 443 TM to form the resulting M-O π bonds and O-O dimers.

444 This mechanism contrasts with the conventional thinking that antisite/vacancy formation in Li-445 rich layered oxides is driven by the destabilization of the layered structure at low Li 446 stoichiometries.^{15, 41, 42} We instead propose that this defect behavior is best understood as an 447 overall decoordination of oxygen in response to low valence electron counts and/or oxygen 448 redox. Indeed, this explains why the onset of electrochemical irreversibility for wide a range of 149 Sn content coincides with exceeding the Ir^{5.5+} oxidation state and not a specific Li stoichiometry 450 (Fig. 3a). We also emphasize that this decoordination takes the form of both in-plane and out-of-451 plane cation disorder. This rationalizes early theories of Li-rich layered oxide structural behavior, 452 which proposed a conversion of the honeycomb superstructure to a layered $LiMO₂$ phase via in-453 plane cation migration,⁴³ with later studies observing out-of-plane site disorder creating spinel-454 like structures. 41

455

456 **Conclusion**

457 In summary, by investigating the effect of Sn substitution in the LIO/LISO model system, we 458 have revealed the origin of the strong coupling between anion redox and structural disordering in 459 Li-rich layered oxides that underlies their irreversible electrochemical properties. We find that 460 LIO exhibits no anion redox, and multivalent hybridized Ir–O redox accounts for the entire

461 capacity, reconciling why this material exhibits more reversible electrochemical and structural 462 behavior than most known anion-redox-active Li-ion electrode materials. Thus, the long \sim 2.5 Å 463 O–O "dimers" that form in LIO during charge are not evidence of anion redox. When Sn is 464 substituted, the lower valence electron count towards the end of charge promotes substantial 465 LMCT through the formation of short Ir–O π bonds and \sim 1.4 Å O–O dimers, requiring the 466 presence of neighboring cation vacancies and therefore driving the formation of Sn_{Li}/V_M defect 467 pairs. The LMCT realized by the Ir=O and O–O species maintains Ir at its maximum oxidation 468 state of 5.5+ and results, on balance, in O redox. The spectroscopic and structural similarities 469 between O redox in LISO and a range of other Li-rich layered oxides^{6, 31, 44} suggest that oxygen 470 redox may be generally stabilized in the local coordination environments created through local 471 structural defects. Although further study is needed to establish the commonalities between 472 materials, this explanation rationalizes the widespread coupling of oxygen redox to cation 473 disordering that gives rise to its irreversible electrochemical properties.

474 We expect that the nature of the TM=O and O–O species, along with the kinetics and 475 thermodynamics of forming the M_L/V_M defect pair will determine the materials' voltage, long 476 term cycling stability, and rate capability, offering a new framework in which to optimize the 477 performance of Li-rich electrodes. We further suggest that structures outside the layered $Li_{1+x}M_1$. 478 *x*O2 framework that can accommodate the formation of short O–O and/or TM=O bonds through 479 small distortions rather than substantial cation rearrangement or, alternatively, materials that can 480 achieve substantial LMCT without drastic structural changes (e.g. through enhanced covalency) 481 could exhibit improved electrochemical and structural reversibility during anion redox. Indeed, 482 in Na-ion layered oxides where interlayer cation migration is limited and possibly even 483 prevented by the large interlayer spacing and/or prismatic interlayer site geometry, structurally

484 reversible anion redox can be achieved.^{45, 46} Given the stabilizing effect of cation migration 485 demonstrated here, the relative reactivity of these materials towards the electrolyte and oxygen 486 evolution in the absence of cation migration will be an interesting avenue of further study. Our 487 results reveal a clear strategy for designing materials for applications beyond energy storage 488 where low valence electron counts (high oxidation states) need to be reversibly accessed, such as 489 catalysts for reactions including oxygen evolution,^{20, 47, 48} olefin polymerization,⁴⁹ and methane 490 hydroxylation.⁵⁰

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- 637 638

641 **Methods**

642 Materials

643 Li₂IrO₃ (hereafter, LIO), Li₂Ir_{0.75}Sn_{0.25}O₃ (hereafter, LISO25), and Li₂Ir_{0.5}Sn_{0.5}O₃ (hereafter, 644 LISO50) were synthesized by solid state reactions. Appropriate amounts of $Li₂CO₃$ (Alfa Aesar, 645 99.998% metal basis), IrO₂ (Alfa Aesar, 99.99% metals basis), and SnO₂ (Alfa Aesar 99.9%) 646 metals basis) were mixed using a planetary ball mill. 10% excess amount of $Li₂CO₃$ was used to 647 compensate the lithium evaporation at high temperature. Mixed powder was heat treated at 648 1000 °C for 12 hours twice in a box furnace with an intermittent grinding, and then naturally 649 cooled to room temperature.

650

651 Electrochemical measurements and electrode harvesting

652 For all electrochemistry figures in this study, 80 wt. % active material, 10 wt.% 653 polyvinylidene fluoride (PVDF) binder (MTI Corporation) and 10 wt. % carbon black (Timcal 654 C65) were mixed with N-methyl-2-pyrrolidone (Acros Organics) and the slurry was cast onto 655 carbon-coated Al foil using a doctor blade. The electrode sheet was dried at 110 \degree C in air for 1 h 656 followed by overnight at 60 °C under vacuum. Coin cells (CR2032, Wellcos Corporation) were 657 assembled in an Ar filled glove box with a \sim 11.3 mm diameter LIO/LISO electrode, two 25 μ m 658 thick Celgard separators, a 750 µm thick Li foil counter electrode (Sigma-Aldrich), and 1 M 659 LiPF₆ in 1:1 (v/v) ethylene carbonate (EC)/diethyl carbonate (DEC) electrolyte (Selectilyte LP 660 40, BASF). The coin cells were cycled under a constant current density of a $C/12$ rate (1C = 211) 661 mA g^{-1} _{LIO}, 227.4 mA g^{-1} _{LISO25}, 246.6 mA g^{-1} _{LISO50} corresponding to the (de)lithiation rate of 2Li/h 662 f.u.) between 4.60 V and 2.50 V (LIO and LISO) or 4.25 V and 2.50 V (LISO). Galvanostatic 663 intermittent titration technique (GITT) was applied to measure the open circuit voltage (OCV) at 664 various states of charge and the thermodynamic voltage hysteresis of LIO and LISO. We applied 665 constant current pulses (C/20) for an hour followed by the relaxation for four hours. This was 666 repeated until the GITT cycle was complete.

667 *Ex situ* soft XAS (fluorescence yield), X-ray diffraction (XRD), and resonant inelastic X-668 ray scattering (RIXS) samples were dismantled from coin cells at the indicated states of charge 669 in an Ar filled glove box. Cells were dismantled using a coin cell decrimping instrument (MTI 670 Corporation). The electrodes were then rinsed with excess DEC and dried under vacuum in the 671 glove box antechamber. For XRD, materials were scraped off of the Al foil current collector and 672 then sealed in a glass capillary (0.5 mm diameter, Ted Pella). All the samples were transferred to 673 the desired instrument in a double-sealed Al-coated polypropylene pouch to prevent air exposure.

674

675 X-ray diffraction and Rietveld refinement

676 High resolution powder XRD patterns for Rietveld refinement were measured at beamline 677 2-1 at the Stanford Synchrotron Radiation Lightsource (SSRL, SLAC National Accelerator 678 Laboratory) at 17 keV (0.7293 Å) beam energy for as-synthesized powders and at beamline 11- 679 ID-B at the Advanced Photon Source (APS, Argonne National Laboratory) at 58.4 keV (0.2114 680 Å) beam energy for *ex situ* samples scraped out of electrodes after electrochemical cycling. The 681 size of the X-ray beam was 500 x 1500 µm (SSRL) or 500 x 500 µm (APS). All of the samples 682 were prepared in capillaries to avoid possible preferred orientation of the particles. All of the 683 samples were measured using the transmission geometry.

684 We used the TOPAS software package (Academic v6, Bruker) for Rietveld refinement. 685 Pawley fitting was employed to determine the crystallographic parameters. The pristine and the 686 samples fully charged to 4.60 V were refined with the *C2/m* space group following previous

687 reports.¹⁵ For the T3-Li₁IrO₃, we found the *Cm* space group to give the best results. For LISO, we 688 assumed that Ir and Sn are randomly distributed through the 4h sites in the TM layer, and the 689 migrated Ir/Sn from 4h was assumed to redistribute equally to the 4g and 2c sites in the Li layer. 690 We allowed variations in the Ir occupancy in Li layer of the fully charged (4.60 V) and fully 691 discharged (2.50 V) LIO and confirmed no Ir migration during the first cycle in LIO.

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693 X-ray total scattering and pair-distribution function analysis

694 Atomic pair-distribution function (PDF) was analyzed to gain further insights on the local 695 atomic structures of LIO and LISO of the pristine powder and the samples scraped out of the 696 electrodes charged to 4.60 V and discharged to 2.50 V. X-ray total scattering was collected at 697 beamline 11-ID-B at the APS at 58.4 keV (0.2114 Å) beam energy. All of the samples were 698 packed in Kapton capillaries double sealed in Al-coated pouch in an argon filled glove box, and 699 then transferred to the beamline. The pouch was opened right before the measurement to 700 minimize the air exposure. The data collection time was three minutes per sample. We used 701 PDFgetX2 to obtain the atomic PDF from the total scattering data and PDFGUI to fit the atomic 702 PDF from the starting crystal structures determined by Rietveld refinement described above.

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704 *Operando* Ir L3 edge XAS and EXAFS spectroscopy

705 For *operando* X-ray absorption spectroscopy (XAS), Al-coated poly propylene (PP) pouch 706 cells containing a \sim 11.3 mm diameter LIO/LISO electrode, two 25 μ m thick Celgard separators, 707 a 250 μ m thick Li foil counter electrode (Alfa Aesar), and 1 M LiPF₆ in 1:1 (v/v) ethylene 708 carbonate (EC)/diethyl carbonate (DEC) electrolyte (Selectilyte LP 40, BASF), a Ni negative 709 current collector tab, and an Al positive current collector tab were assembled using a tabletop 710 vacuum impulse sealer (Fuji Impulse) in an argon filled glove box. A stainless steel holder with a 711 pair of Be plates (Ted Pella) as an X-ray window was machined to apply a sufficient pressure on 712 the pouch cells using torque screws. The pressure from the stiff windows significantly improved 713 electronic conduction throughout the micro-porous electrodes ensuring homogeneity. The pouch 714 cells were cycled using a portable potentiostat (SP-150, Biologic) under a constant current 715 density of C/12 (where 1C refers to 2Li/h f.u.) between 4.60 V and 2.50 V (*vs.* Li) with 1 hr rest 716 between charge and discharge.

717 The Ir L3 edge XAS spectra of LIO and LISO was collected at beamline 2-2 of Stanford 718 Synchrotron Radiation Lightsource (SSRL) at SLAC National Accelerator Laboratory and 719 beamline 20-BM of the Advanced Photon Source (APS) at Argonne National Laboratory, 720 respectively. The transmission spectra were measured using a 1 x 7 mm unfocused X-ray beam. 721 A Si (220) (SSRL) or Si (111) (APS) crystal monochromator was detuned to 70% of its original 722 intensity to eliminate high order harmonics. Three ion chambers filled with N_2 gas were used in 723 series to simultaneously measure I_0 , I_1 , and $I_{ref.}$ A Ge reference foil was used to calibrate the 724 photon energy by setting the peak of the first derivative of the K edge absorbance spectrum to be 725 11102 eV. 10 eV, 0.5 eV, and 10 eV energy steps were used in the ranges of 10985-11085 eV, 726 11085.5-11124.5 eV, and 11125-11195 eV, respectively, in order to precisely measure the 727 reference Ge K edge spectra for the calibration followed by the Ir L_3 edge measurement. We 728 used the Athena software package to align and normalize the collected spectra. White line peak 729 was fitted using the sum of an error function and a Lorentzian peak analogous to previous reports. 730 Fourier transformations of k^2 weighted extended X-ray absorption fine-structure (EXAFS) 731 spectra were carried out in the *k* range from 2 \AA^{-1} to 14 \AA^{-1} for all the materials.

734 O K edge Soft X-ray Absorption Spectroscopy: FY, RIXS, and transmission

735 O K edge sXAS of the harvested electrodes were measured in fluorescence yield (FY) mode 736 at SSRL beam line 10-1 equipped with a silicon photodiode detector (AXUV100). Data was 737 acquired under ultrahigh vacuum $(10^{-9}$ Torr) at room temperature with the incident X-ray beam 738 of 500 μm x 500 μm size.

739 RIXS maps were collected at beam line 8.0.1 at the Advanced Light Source (ALS) in 740 Lawrence Berkeley National Laboratory, using the ultrahigh efficiency iRIXS endstation. All the 741 harvested electrodes were sealed in Al pouches in an argon filled glove box and transferred into a 742 specially designed transfer kit and then the experimental vacuum chamber to avoid air exposure. 743 Technical details of the RIXS beam line and data processing can be found in our previous 744 reports.^{6, 51} Two-dimensional emission spectra collected at each excitation energy are aligned 745 using the elastic peak and a reference compound such as $TiO₂$ to generate the full RIXS maps. 746 The color scale has been tuned in the figures to emphasize the contrast of intensity.

747 Transmission O K edge sXAS was measured at ALS beam line 11.0.2 using Scanning 748 Transmission X-ray Microscopy. The samples were prepared by sonicating the harvested 749 electrodes in dimethyl carbonate under argon for two hours at room temperature to separate 750 individual particles out of the composite electrodes. The particle suspension was drop-cast onto 751 copper TEM grids with a carbon film (Ted Pella). The grids were loaded onto a sample holder, 752 sealed in an Al-coated pouch, and then transferred to the beam line. We first collected the STXM 753 images with a 50 nm zone plate, an interferometer-controlled stage, and a point detector. The 754 step size was 50 nm and the dwell time for each pixel was typically 1 ms. STXM spectro-images 755 were aligned in the aXis2000 software package. To obtain average absorption spectra, the 756 aligned absorbance images were filtered at an energy of non-zero intensity, typically 529.5 eV. 757 The pixels below a threshold intensity were set to zero. The remaining un-normalized pixels 758 were then summed to yield the average spectrum. For presentation, normalization of the average 759 spectra was done by subtracting the background intensity and then dividing by the post edge 760 intensity.

761

762 Computational details

763 The calculations for all structures presented were performed with density functional theory 764 (DFT) as implemented in the Vienna *Ab-initio* Simulation Package (VASP).^{52, 53} Valence 765 electrons were described by the plane wave basis set and core electrons were incorporated by the 766 projector augmented-wave method.⁵⁴⁻⁵⁷ Unless otherwise indicated, the perdew-burke-ernzerhof 767 (PBE) functional⁵⁸ with the Hubbard *U* correction⁵⁹ was adopted for the exchange correlation 768 energy. An effective *U* value of 2.75 eV was applied on Ir.⁶⁰ For hybrid calculations, the HSE 769 screened coulomb hybrid density functional⁶¹ was used with a mixing parameter of 0.15 and a 770 range separation parameter of 0.2^{62} . The energy cutoff of the plane wave basis was 520 eV and 771 the *k*-point mesh was $3 \times 3 \times 7$ for a $2 \times 2 \times 1$ supercell of O3-Li_{2-x}MO₃. Van der Waals 1772 interactions were taken into account using the D2 method of Grimme⁶³ for PBE+U and HSE 773 calculations. For SCAN calculations, the rVV10 non-local van der Waals correlation functional 774 was used.⁶⁴ To prepare the structures of $Li_{2-x}IrO_3$, we generated all Li-vacancy orderings within 775 the unit cell of $Li_{2-x}IrO_3$ including 4 formula units using the enumeration technique developed by 776 Hart *et al.*^{65, 66} and 10 configurations at $x = 1.5$ and 0.5 and 100 configurations at $x = 1$ with 777 lowest electrostatic energy were calculated using GGA. The configurations with the lowest 778 DFT/GGA energy at each Li content were selected as most stable structures. Both octahedral and

- 779 tetrahedral sites in Li layer within O3- and O1-Li_xIrO₃ were considered for Li-vacancy orderings
- 780 at x=1. The same Li-vacancy orderings were applied to the $Li_{2-x}Ir_{0.75}Sn_{0.25}O_3$ case. After Sn
- 781 migration to Li layer, the Li-vacancy orderings were re-sampled using the basin hopping 782 algorithm. 67

784 **Data Availability**

785 All experimental data within the article and its Supplementary Information will be made

786 available upon reasonable request to the authors.

787

788 **References**

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867 **Competing Interests:** The authors declare no competing interests.

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870 **Fig. 1. Reversible multivalent iridium redox in Li_{2-x}IrO₃. (a) Capacity-voltage curves of Li₂₋** 871 _xIrO₃ galvanostatically measured at a C/12 rate (17.58 mA g^{-1}) between 4.50-2.50 V for the 872 initial three cycles. The stable phases during the two consecutive two-phase reactions of $Li_{2-x}IrO_3$ 873 occurring at each voltage plateau, obtained by combined XRD Rietveld refinements and DFT 874 calculations, are shown. (**b**) Evolution of the Ir L₃ WL energy (black squares) during the first

886 **Fig. 2**. **Hybridized Ir-O redox in Li2-xIrO3**. (**a**) sXAS fluorescence yield spectra (solid lines) 887 and XAS obtained through scanning transmission X-ray microscopy (STXM-XAS, dashed lines) 888 of the O K edge of $Li_{2-x}IrO_3$ at various voltages throughout the first cycle. From bottom to top, 889 pristine (red, A), charged to 3.9 V (yellow, B), charged to 4.6 V (green, C), discharged to 3.7 V 890 after being charged to 4.6 V (olive, D), discharged to 2.5 V after being charged to 4.6 V (blue, 891 E). (**b**) Difference plot of sXAS obtained from (a) showing the intensity evolution of Ir *5d* -O *2p* t_{2g} and e_g^* peaks. (c) Ratio of the Ir 5d-O 2p t_{2g} peak area to the total Ir 5d-O 2p $t_{2g} + e_g^*$ area 893 measured by RIXS, STXM and sXAS, showing the continuous growth and decay of the t_{2g} area. 894 The O K edge measurements were normalized by the intensity at 545 eV after subtracting the 895 background intensity. (**d**) O K edge RIXS maps of $Li₂IrO₃$, LiIrO₃ and $Li_{0.5}IrO₃$. The right 896 bottom panel shows the RIXS spectra at 530.7 eV excitation energy for each composition where 897 a feature corresponding to anion redox is reported to appear. (**e**) Ir- and O-projected density of 898 states of Li₂IrO₃, LiIrO₃ and Li_{0.5}IrO₃ calculated from first principles, demonstrating no access to 899 the buried O *2p* non-bonding states. Insets show the isosurface of the charge density for the

- 900 lowest unoccupied states corresponding to 0.5 electrons/f.u. in LiIrO₃ and Li_{0.5}IrO₃ (shaded
- 901 region in (e)) visualizing two different Ir 5d-O 2p t_{2g} hybridized states. Yellow and blue show
- 902 negative and positive changes in charge density, respectively. Li ions are omitted for clarity.

923 **Fig. 4. Computational predictions of M-O decoordination and Ir=O/O–O stabilized anion** 924 **redox**. (**a**) Ir–O bond lengths (left) and oxygen coordination environments (right) predicted by 925 DFT in a $Li_{0.5}Ir_{0.75}Sn_{0.25}O_3$ structure before (top) and after (bottom) the formation of a Sn_{Li}/V_M 926 defect when the vacancy neighbors only Ir. Note that the presence of Li neighboring O negligibly 927 affects the O *2p* states due to minimal hybridization between Li and O. (**b**) Projected DOS of the 928 individual oxygen atoms involved in the Ir–O bond contraction before (top) and after (bottom) 929 forming the $\text{Sn}_{\text{Li}}/\text{V}_{\text{M}}$ defect, demonstrating the shift of the O 2p states above the Fermi level, 930 indicating oxygen redox. (**c**) Ir–O bond lengths (left) and oxygen coordination environments 931 (right) before (top) and after (bottom) the formation of a Sn_{Li}/V_M defect when the migrating Sn 932 initially neighbors another Sn, resulting in the formation of a 1.44 Å O–O dimer. (**d**) Projected 933 DOS of the individual oxygen atoms involved in the O–O bond formation before (top) and after

- 934 (bottom) forming the $\text{Sn}_{\text{Li}}/V_M$ defect and O–O dimer, also showing a shift of the O 2p states
- 935 above the Fermi level.

938 **Fig. 5**. **Proposed electronic mechanism of cation migration and LMCT mediated anion** 939 **redox in LISO.** (a) The mechanism of Ir=O formation, wherein Ir is initially oxidized beyond 940 Ir^{5.5+}, promoting LMCT *via* Sn migration, oxygen decoordination, and donation of oxygen lone 941 pair (unhybridized O 2*p*) electrons to forming a short Ir–O π bond. Each black arrow corresponds 942 to the redistribution of an electron pair. **(b)** The mechanism of O–O formation when a dangling 943 O is coordinated to a Sn atom, wherein LMCT is achieved instead through donation of the O–O 944 σ* electrons to Ir. Both cases assume that the number of electrons provided through LMCT 945 mediated oxygen redox is equal to the number of decoordinated oxygens. Note that the 946 calculations in Fig. 4 reflect only the LMCT step in both cases.