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1 Unravelling Solid-State Redox Chemistry in Li $_{1,3}$ **Nb** $_{0,3}$ **Mn** $_{0,4}$ **O₂ Single-**² Crystal Cathode Material

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11 S [Supporting Information](#page-9-0)

 ABSTRACT: Recent reports on high capacities delivered by Li-excess 13 transition-metal oxide cathodes have triggered intense interest in utilizing reversible oxygen redox for high-energy battery applications. To control oxygen electrochemical activities, fundamental understanding of redox chemistry is essential yet has so far proven challenging. In the present 17 study, micrometer-sized $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ single crystals were synthesized for the first time and used as a platform to understand the charge compensation mechanism during Li extraction and insertion. We 20 explicitly demonstrate that the oxidation of O^{2-} to O^{n-} $(0 \lt n \lt 2)$ 21 and O_2 loss from the lattice dominates at 4.5 and 4.7 V, respectively. 22 While both processes occur in the first cycle, only the redox of O^{2-}/O^{n-} participates in the following cycles. The lattice anion redox process triggers irreversible changes in Mn redox, which likely causes the voltage

- 26 activity regions, a single-phase behavior involving only Mn^{3+/4+} and a two-phase behavior involving O^{2−}/O^{n−} (0 ≤ n < 2), were
- 27 found in $Li_xNb_{0.3}Mn_{0.4}O_2$ (0 < x < 1.3). Morphological damage with particle cracking and fracturing was broadly observed when ²⁸ O redox is active, revealing additional challenges in utilizing O redox for high-energy cathode development.
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1. INTRODUCTION

 Lithium-ion batteries (LIBs) have become increasingly important for energy storage in portable electronics, electric vehicles, and stationary application in power grids. Since the discovery of early generations of cathode materials (e.g., 33 LiCoO₂ and LiMn₂O₄) in the 1980s, only a few compounds with other compositions or crystal structures have been 5 reported in the last three decades. $1-3$ $1-3$ The conventional wisdom suggests that in O3-type layered oxides, the 3d orbitals of early transition metals (TMs), are partially overlapped with the 2p orbitals of oxygen.^{[4](#page-9-0)} As such, only about half of the 3d electrons are available to participate in the redox reaction, which largely limits practical capacity (for example, ∼160 mAh/ $_{41}$ g for LiCoO₂) despite their much higher theoretical capacities. To develop high-energy batteries, cathodes with higher specific 43 capacity (>200 mAh/g) and operating voltage (>4.3 V vs $\mathrm{Li}^{+}/$ 44 Li) are needed. $1-3$ $1-3$ $1-3$

 Recently, approaches to enable high-energy cathodes by utilizing redox reactions of both TM cations and oxygen anions have triggered intense interest.[5](#page-9-0)−[7](#page-9-0) One of the most studied examples is the lithium and manganese-rich (LMR) layered oxides with a general formula of $Li_{1+x}Mn_{1-x-y-z}Ni_yCo_zO_2$.^{[8](#page-9-0)−[10](#page-10-0)} 49 Our recent work showed that, contrary to the common notion ⁵⁰ of a nanocomposite structure, the oxide has a single monoclinic ⁵¹ phase $(C2/m)$ with a large number of domains corresponding 52 to different variants. 11 To involve the O 2p electrons in the 53 following electrochemical reactions, the material typically ⁵⁴ undergoes an initial activation process signaled by a unique ⁵⁵ charging voltage profile that is much different from those of the 56 subsequent cycles. Recent studies by Luo et al. suggested the 57 formation of O $^-$ holes in the intermediates, as evidenced by the $\,$ s $\,$ progressive growth of a new peak on the O K-edge X-ray ⁵⁹ absorption spectroscopy (XAS) along with the use of a number 60 of other characterization techniques, including isotopically ⁶¹ labeled differential electrochemical mass spectroscopy 62 (DEMS), X-ray absorption near edge structure (XANES), ⁶³ and resonant inelastic X-ray scattering $(RIXS)$.^{[9](#page-9-0)} However, this 64 remains controversial as experimental evidence is difficult to ⁶⁵

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 obtain, largely due to their high reactivity toward carbonate- based electrolyte and/or self-combine to release carbon dioxide and oxygen gases, respectively. These processes also produce significant amount of oxide-ion vacancies in the unit cell, further destabilizing the crystal structure.[12](#page-10-0) The activation process also triggers unfavorable phase transformations, 72 possibly involving layered \rightarrow spinel \rightarrow rock-salt, as the cycling proceeds, leading to continuous voltage and capacity fades in the cathode.[12](#page-10-0) Various optimization approaches such as doping,[13,14](#page-10-0) surface coating/engineering, and morphology $\frac{1}{26}$ control^{[15](#page-10-0)−[18](#page-10-0)} have been explored as means to mitigate the degradation, but all were met with limited success.

 An alternative approach to address the phase instability is to confine O redox process within a rock-salt structure. This strategy was rarely used as it is generally believed that compounds with rock-salt crystal structure lack lithium-ion conduction pathways and are therefore electrochemically inactive. In 2003, Shigemura et al. first reported that the Li/ Ti/FeO₂ system can deliver a capacity of ca. 200 mAh/g at a 85 slow rate of 10 mAh/ $g^{19,20}$ $g^{19,20}$ $g^{19,20}$ Investigation on lithium-ion 86 conduction pathways in rock-salt $\rm L_{1,211}Mo_{0.467}Cr_{0.3}O_2$ was then reported by Lee et al. in 2014.^{[21](#page-10-0)} Using density functional theory (DFT) calculation, they revealed the percolation of active diffusion channels in disordered Li-excess materials. In 2015, Yabuuchi et al. reported a new rock-salt $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ (LNMO) cathode with an impressive 92 discharge capacity of ca. 300 mAh/g.^{[22](#page-10-0)} With the use of soft/ hard X-ray absorption spectroscopy, the authors proposed the 94 involvement of both $\text{Mn}^{3+/4+}$ and O^{2-}/O^- redox reactions during charge. However, the detailed charge compensation mechanism and role of O, particularly the contribution from 97 oxidation of O^{2-} to O^{n-} $(0 \lt n \lt 2)$ and O loss from the lattice 98 (in the form of O_2 gas, $n = 0$), remain unclear. In this study, 99 well-formed $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ single crystals were synthesized by a molten-salt method for advanced diagnostic studies. The use of single crystals allows us to perform additional single- particle-based studies for systematic comparison of the changes in crystal structure, chemical state, and microstructure as a function of lithium content in the sample. The results provide us important insights on O activities in the oxide sublattice and how to control/tune them through engineering strategies such as chemical composition and surface morphology control.

2. EXPERIMENTAL SECTION

 2.1. Synthesis. In a typical crystal synthesis procedure, 109 stoichiometric amounts of Li_2CO_3 , Nb_2O_5 , and Mn_2O_3 precursors (Sigma-Aldrich, >99%) were milled together with ethanol solvent at 200 rpm for 12 h in a planetary ball mill (RETSCH PM100) using a 112 zirconia jar with zirconia balls. About 10−15% extra $Li₂CO₃$ was used to compensate Li loss that often occurs during high temperature heating. The dried powder was further mixed with KCl flux in a molar ratio of 2.5−5 between the flux and TM precursors (defined as R ratio 116 hereafter). The obtained mixture was then heated at 950 °C for 12 h in 117 an Ar atmosphere. A ramp rate of 4 $^{\circ}$ C/min was used for both heating and cooling. After the reaction, KCl was dissolved in deionized water, and the final product was obtained after filtration and thorough 120 washing with water. Chemical delithiation of $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ was achieved by reacting the pristine powder with various amounts of 0.1 122 M nitronium tetrafluoroborate $(NO₂BF₄)$ in acetonitrile solution in an 123 argon filled glovebox $(O_2 < 1$ ppm and $H_2O < 1$ ppm). The reaction was carried out at room temperature for 1−4 days. The resulting reaction mixtures were filtered, thoroughly washed with acetonitrile, and then dried overnight in a vacuum oven.

127 2.2. Characterization. Chemical composition of the samples was 128 determined by an inductively coupled plasma optical emission

spectrometer (ICP-OES 720 Series). Scanning electron microscopy 129 (SEM) and energy dispersive X-ray spectroscopy (EDS) images were 130 collected on a JEOL JSM-7500F field emission microscope at a 10 kV ¹³¹ accelerating voltage. Prior to the analysis, the powder samples were 132 sputtered with a thin top layer of Au to reduce the charging effect. ¹³³ Samples for transmission electron microscopy (TEM) experiments 134 were prepared by drop casting a sonicated solution of the crystals in 135 anhydrous ethanol onto a carbon coated TEM grid or by sectioning 136 using focused ion beam. Selected area electron diffraction (SAED) was ¹³⁷ performed in TEM mode at 300 kV on the sectioned sample, while 138 high angle annular dark field (HAADF) imaging was performed on ¹³⁹ dispersed particles in scanning transmission electron microscopy 140 (STEM) mode at 120 kV, both using an FEI Titan electron 141 microscope. 142

Phase purity was first analyzed by using laboratory X-ray diffraction ¹⁴³ (XRD) collected on Bruker D2 powder X-ray diffractometer (Cu Kα, ¹⁴⁴ 40 kV, 30 mA). Synchrotron X-ray diffraction patterns were collected ¹⁴⁵ at ambient temperature at 11-BM at the Advanced Photon Source 146 (APS) in Argonne National Laboratory, which operates with a 147 monochromatic X-ray of $\lambda = 0.414$ Å. The scans were collected 148 between 0.5° and 50° (2 θ) at a step size of 0.0001°. The samples were 149 first packed into Kapton capillary tubes before exposing to X-ray for 1 ¹⁵⁰ h. Time-of-flight (TOF) neutron diffraction data were collected at the ¹⁵¹ POWGEN diffractometer at the Spallation Neutron Source, Oak ¹⁵² Ridge National Laboratory. Samples were measured inside the 153 vanadium sample cans, and a single bank wave with center wavelengths 154 of 1.333 Å (2 h data collection) was used. Monochromatic (1.2 Å) 155 neutron diffraction data were obtained using the high resolution ¹⁵⁶ powder diffractometer at BT1 at National Institute of Standards and ¹⁵⁷ Technology (NIST) Center for Neutron Research. The refinement of ¹⁵⁸ the diffraction data was carried out using GSAS/EXPGUI package. 23 23 23 159

Mn K-edges hard X-ray absorption spectroscopy data were collected 160 in transmission mode using a (220) monochromator at SSRL 161 beamline 2-2 and 2-3. The crystals were sandwiched between two 162 Kapton films for data collection. Higher harmonics in the X-ray beam ¹⁶³ were reduced by detuning the Si (220) monochromator by 50% at the 164 Mn edge. Energy calibration was accomplished by using the first ¹⁶⁵ inflection point at 6539 eV in the spectra of Mn metal foil reference. ¹⁶⁶ X-ray absorption near edge structure spectra were analyzed by Sam's 167 Interface for XAS Package or SIXPACK software, with the 168 photoelectron energy origin E_0 determined by the first inflection 169 point of the absorption edge jump. Soft X-ray absorption spectroscopy 170 measurements were carried out at beamlines 8-2 and 10-1 at SSRL. A 171 thin layer of the crystals was spread onto a conductive carbon tape 172 which was attached to an aluminum sample holder inside an Ar filled ¹⁷³ glovebox (O_2 < 1 ppm, H_2O < 1 ppm). Measurements were carried 174 out on the 31-pole wiggler beamline 10-1 at SSRL with a ring current 175 of 350 mA, a 1000 l mm⁻¹ spherical grating monochromator with 20 $\frac{176}{176}$ mm entrance and exit slits, a 0.2 eV energy resolution, and a 1 mm² 177 beam spot. Data were collected at room temperature under ultrahigh 178 vacuum (10^{-9} Torr) in a single load using the total electron yield and 179 fluorescence yield mode detectors.

2.3. Electrochemistry. To prepare the composite electrodes, the 181 active material was first ball-milled with acetylene carbon black ¹⁸² (Denka, 20 wt %) to reduce the particle size and improve its electronic 183 conductivity. The mixture was then mixed with acetylene carbon black 184 and a polyvinylidene fluoride (PVdF) binder (Kynar 2801) in a 8:1:1 ¹⁸⁵ weight ratio in an NMP solvent. The slurry was spread onto aluminum 186 foil and dried overnight at 120 °C under vacuum. Cathode disks with 187 an area of 1.6 cm² and a typical electrode loading of 2-3 mg cm⁻² 188 were cut from the electrode sheets and assembled into 2032-type coin 189 cells in an argon-filled glovebox. Lithium foil (Alfa-Aesar) was used as ¹⁹⁰ counter and reference electrodes, Celgard 2400 polypropylene 191 membrane as separators, and 1 M LiPF₆ in 1:1 (v/v) ethylene 192 carbonate (EC):diethylene carbonate (DEC) (Novolyte Technologies 193 Inc.) as electrolyte. The cells were galvanostatically cycled between 1.5 194 and 4.8 V using a VMP3 multichannel potentiostat/galvanostat 195 controlled by EC-Lab v10.12 software (BioLogic Science Instru- 196 ments). Cyclic voltammetry tests were carried out using the same coin 197 cell configuration with Li foil as counter and reference electrodes. The ¹⁹⁸

199 cells were scanned between 1.5 and 4.8 V with a constant scan rate of 200 5 mV/min. All electrochemical tests were carried out at room 201 temperature.

 Differential electrochemical mass spectroscopy measurements were performed in a customized Swagelok-type cell consisting of a lithium 204 anode, a liquid electrolyte (1.0 M LiPF₆ in EC/DEC 1:1 v/v), and the crystal composite as the working positive electrode. The cell was periodically pulsed with Ar gas at regular intervals to sweep the evolved gases from the headspace to the mass spectrometer for 208 analysis. The experimental setup is described in detail elsewhere. 24 Electrochemical measurements were carried out with a constant current rate of 25 mA/g between 1.5 and 4.8 V with a rest period of 25 min right after the cells reached the upper or lower cutoff voltages. All data collection was controlled by a BioLogic SP-300 potentiostat.

3. RESULTS AND DISCUSSION

 3.1. Synthesis and Characterization of 214 Li_{1.3}Nb_{0.3}Mn_{0.4}O₂ Single Crystals. Synthesis of polycrystal-215 line $L_{1.3}Nb_{0.3}Mn_{0.4}O_2$ cathode material was recently carried out by Yabuuchi et al.[22](#page-10-0) The process involved ball-milling 217 stoichiometric amount of Li_2CO_3 , Mn_2O_3 , and Nb_2O_5 , pressing the resulting mixture into a pellet, and then heating the pellet in Ar atmosphere at 900 °C for 24 h. Synthesis of 220 Li_{1.3}Nb_{0.3}Mn_{0.4}O₂ single crystals, however, has not been reported in the literature. Here, we adopted a molten-salt method and systematically varied the synthesis conditions to 223 obtain phase-pure and discrete $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ single crystals with uniform morphology and size distribution for the first time. The following parameters were carefully optimized in our synthesis: choice of transition-metal precursor salts and flux salt, the molar ratio between the flux and TM precursors (R ratio), choice and amount of lithium precursor, heating temperature and time, and reaction atmosphere. It was found that the best reaction time and temperature were 12 h and 950 231 °C, respectively. Li_3NbO_4 was often obtained as an impurity when the reaction time was too short or the heating temperature was below 950 °C. The loss of lithium at elevated temperature was effectively compensated by using additional 10−15% of the lithium precursor. Various molten salts, 236 including LiCl, NaCl, KCl, CsCl, KOH, and Li_2SO_4 , were 237 used as the flux, but only KCl (mp = 770 $^{\circ}$ C) led to the formation of phase-pure samples. The optimized R ratio was 239 between 2.5 and 5. In addition, the presence of trace O_2 in the 240 synthesis atmosphere can lead to the formation of $Li₂MnO₃$ 241 and Li₃NbO₄ impurities.

 The nominal composition of the as-synthesized oxide was analyzed by an inductively coupled plasma optical emission spectrometer, which confirmed the chemical formula of $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$. As shown in the scanning electron 246 microscopy image in Figure 1a and high angle annular dark field scanning transmission electron microscopy image in Figure 1b, the particles adopted a large spherical shape with an 249 average size of ca. $5-8$ μ m and absence of any grain boundaries. This suggests that all the facets have similar thermodynamic stability and grow at a similar rate under current synthesis conditions. Contrast reversal toward the center of the particles observed on HAADF STEM images indicates that the pristine particles are too thick for detailed analytical electron microscopy. Focused ion beam (FIB) milling was then used to reduce the thickness and a SAED pattern in [110] zone axis was taken over an area covering almost entire single particle (Figure 1c). The pattern matches well with the $_{259}$ one simulated using $Fm\overline{3}m$ structure with a lattice parameter of $260 a = 4.2$ Å (Figure 1d), confirming the single crystal nature of

Figure 1. Morphology and structure of $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ crystals: (a) secondary electron SEM image, (b) HAADF STEM image, (c) SAED pattern taken in [110] zone axis, and (d) simulated electron diffraction pattern corresponding to panel c.

the particle. It is worth noting that the SAED pattern from this ²⁶¹ particle also shows diffuse scattering effects that are character- ²⁶² istic of materials with short-range ordering.^{[25](#page-10-0)} At this point, it is 263 unclear whether this short-range order occurs broadly in the ²⁶⁴ entire sample or if it has any effect on oxide properties and ²⁶⁵ behavior. Both of them require further investigation with ²⁶⁶ complementary analytical techniques, and the results will be ²⁶⁷ reported in a future publication. ²⁶⁸

Bulk phase purity and crystal structure of the synthesized ²⁶⁹ samples were evaluated by using both synchrotron X-ray and ²⁷⁰ neutron diffraction studies. [Figure 2](#page-3-0) shows the Rietveld 271 f2 refinement of the diffraction patterns, while [Table 1](#page-3-0) lists the 272 t1 refined structural parameter. In both sets of diffraction patterns, ²⁷³ the high peak symmetry observed in a wide angular range ²⁷⁴ indicates a single phase with high crystallinity. Rietveld ²⁷⁵ refinement confirms a rock-salt crystal structure with a lattice ²⁷⁶ parameter of 4.1954(1) Å and cell volume of 73.685(1) Å³, 277 , which is in good agreement with the values reported by ²⁷⁸ Yabuuchi et al. on polycrystalline samples. 22 22 22 All cations (Li, Nb, 279 and Mn) were randomly located in the 4a sites, and all M−O ²⁸⁰ bonds had the same distance of $2.0965(1)$ Å, confirming the 281 disordered nature of the oxide crystals. The refined occupancies ²⁸² agree well with the chemical formula of $Li_{1.3}Nb_{0.3}Mn_{0.4}O_2.$ 283

3.2. Electrochemical Studies. The electrochemical ²⁸⁴ performance of $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ crystals was evaluated at 285 room temperature in a half-cell configuration. Due to high ²⁸⁶ resistance and poor utilization of large particles, only limited ²⁸⁷ capacities were obtained on composite electrodes made with ²⁸⁸ the as-synthesized micrometer-sized crystals. In a modified ²⁸⁹ procedure, the crystal size was reduced and electronic ²⁹⁰ conductivity improved by ball-milling the oxide crystals ²⁹¹ together with a carbon black additive in a ²⁹² $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$:carbon weight ratio of 80:20. A SEM image 293 of the mixture is shown in [Figure S1.](http://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.7b05036/suppl_file/cm7b05036_si_001.pdf) The mixture was then ²⁹⁴ added into the standard carbon and PVdF slurry in an N- ²⁹⁵ methyl-2-pyrrolidone (NMP) solvent and cast onto the Al ²⁹⁶

Figure 2. Rietveld refinement of (a) XRD and (b) neutron diffraction patterns collected on pristine $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ single crystals (PWEGEN). Values in parentheses indicate standard deviation.

Table 1. Structural Parameters of Pristine $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ Crystals Determined from Joint Rietveld Refinements of Synchrotron X-ray and TOF Neutron Data^a

| atom | position | Wyck. site | осср. | $d_{(M-O)}$ (Å) | bond valence | $100(U_{\text{iso}})$ (\AA^2) |
|------------|----------|---------------|-------|--|-----------------|------------------------------------|
| Li | 0,0,0 | 4a | 0.65 | 2.0965(1) | 1.092 | 1.13(1) |
| Mn | 0,0,0 | 4a | 0.2 | 2.0965(1) | 2.43 | 1.13(1) |
| Nb | 0,0,0 | 4a | 0.15 | 2.0965(1) | 3.66 | 1.13(1) |
| \circ | 0,0.0.5 | 4b | 1.0 | 2.9650(1) | | 1.72(1) |
| $0.0(0)$. | | | | ${}^aFm\overline{3}m$, a (Å) = 4.1954(1) V (Å ³) = 73.845(1), wR _p = 9.22%, δ = | | |

²⁹⁷ current collector. The final weight ratio of active materi-²⁹⁸ al:carbon additive:PVdF binder in the thus-prepared electrodes 299 was $56:34:10$. Electrode disks with a size of 1.6 cm² and an

active material loading of ca. 2–3 mg/cm² were then cut out 300 and used as cathodes in coin cells, which were cycled between ³⁰¹ 1.5 and 4.8 V vs $\rm Li^+/Li$ at a constant current density of 10 mA/ $_{302}$ g. Figure 3a shows the voltage profiles of the first four cycles. 303 f3 Similar to what was observed in the LMR oxide cathodes, two ³⁰⁴ distinct regions, a sloping region and a plateau region, are ³⁰⁵ present during the first charge. In the sloping region, the ³⁰⁶ voltage increased steadily from the open circuit voltage (OCV) 307 of 3.0 to 4.3 V, which delivered a specific capacity of ∼115 ³⁰⁸ mAh/g. Whereas in the plateau region, the voltage increased ³⁰⁹ only slightly from 4.3 to 4.5 V before it rapidly increased to the 310 cut off voltage of 4.8 V. A specific capacity of \sim 215 mAh/g was 311 obtained above 4.3 V on the first cycle, leading to a total charge 312 capacity of 330 mAh/g. An overall sloping discharge profile was ³¹³ observed, which delivered a total discharge capacity of ∼290 ³¹⁴

Figure 3. (a and b) Voltage profiles of $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ half-cell cycling. (c) Specific capacity and average voltage as a function of cycle number. (d) Cyclic voltammogram of the first 9 scans at a scan rate of 5 mV/min. Solid symbols and open symbols in panel c represent capacity and voltage, respectively.

 mAh/g at the cutoff voltage of 1.5 V. The Coulombic efficiency for the first cycle is therefore ∼88%. Two guidelines, one showing the theoretical capacity originated from the $Mn^{3+}/$ Mn⁴⁺ redox couple at 118 mAh/g and the other showing the theoretical capacity based on Li content at 383 mAh/g are also shown in [Figure 3a](#page-3-0). It is clear that both charge and discharge capacities (330 and 290 mAh/g, respectively) are significantly 322 larger than the theoretical capacity of the Mn^{3+}/Mn^{4+} redox (118 mAh/g). The charge capacity obtained in the sloping 324 region is consistent with the capacity from the Mn^{3+}/Mn^{4+} 325 redox couple, suggesting that Mn^{3+}/Mn^{4+} may be the sole redox process occurring in this region below 4.3 V.

 In the second cycle, there is an overall decrease in the charging voltage of the sloping region, which led to an increased capacity of ∼180 mAh/g upon reaching 4.3 V. The plateau region became somewhat sloping, but voltage increase remained slow. With cycling, the degree of sloping gradually increased, and the capacity obtained from this plateau region continued to decrease. The same trend was also observed on the discharge capacity, which was reduced to 160 mAh/g after 27 cycles, a loss of nearly 45% [\(Figures 3b](#page-3-0) and c). These results obtained from room-temperature cycling are comparable to 337 what was obtained at 60 °C by Yabuuchi et al. Although more carbon was used in our electrode, which likely contributed to the improved performance, the results may also indicate that the crystal samples have a kinetic advantage compared to the polycrystalline samples synthesized using the solid-state method. Further analysis showed that the oxide experienced not only capacity fade but also voltage fade. As shown in [Figure](#page-3-0) [3](#page-3-0)c, the average discharge voltage, obtained by dividing the total 345 cell energy (E) by the cell capacity (Q) at 1.5 V, continues to decrease along with cycling. The extent of decay appears to be sensitive to the discharge cutoff voltage, and much faster decay was observed when the cutoff voltage was reduced from 1.5 to ³⁴⁹ 1.0 V.

350 Further insights on the redox activities of $Li_{1.3}Nb_{0.3}Mn_{0.4}O_2$ crystals were obtained from cyclic voltammetry performed between 1.5 and 4.8 V at a scan rate of 5 mV/min. [Figure 3d](#page-3-0) shows the results obtained from the first nine cycles. A broad peak centered at 3.5 V and a split peak between 4.5 and 4.7 V were observed during the first oxidation, corresponding to the sloping and plateau regions on the charging voltage profile, respectively. In the following scans, the oxidation peak at 3.5 V gradually moves toward high voltage, and its intensity grew while that of the peak at 4.5 V decreased, along with the disappearance of the split peak at 4.7 V after the first cycle. As the peak around 4.5−4.7 V is often attributed to the oxidation 362 of \overline{O}^{2-} to \overline{O}^{n-} (0 $\leq n < 2$), the disappearance of the split peak and the continuous decrease in peak intensity suggest the complex and irreversible nature of the processes. During the first reduction, only a single peak was observed around 3.2 V, which decreased its intensity along with the appearance and gradual growth of a new peak around 2.2 V in the following scans. The pair of peaks centered at 3.5 V during oxidation and 369 3.2 V during reduction is likely associated with the Mn^{3+}/Mn^{4+} redox couple. The growth of the reduction peak at 2.2 V at the expense of the peak at 3.2 V suggests the gradual trans- formation of the Mn species and the increasing contribution from the lower voltage process along with cycling. This transformation was not observed when the upper cutoff voltage was limited to 4.0 V, before the onset of the oxygen oxidation 376 peak. The results suggest that the oxidation of O^{2-} to O^{n-} at high voltage may be responsible for the irreversible conversion

of the Mn^{3+} redox species, which ultimately led to both capacity 378 and voltage fade in this oxide cathode.

Operando differential electrochemical mass spectroscopy was ³⁸⁰ used to further examine the oxidation process of O^{2-} to O^{n-} (0 381 $\leq n < 2$) occurring between 4.5 and 4.7 V. Figure 4 shows the 382 f4

Figure 4. (a) Voltage profile and (b) O_2 and CO_2 gas evolution analyzed by DEMS of a cathode containing 2.66 mg $Li_{1.3}Nb_{0.3}Mn_{0.4}O_2$ active material.

results obtained during the first 2 cycles at a constant current ³⁸³ density of 25 mA/g. The voltage profiles are similar to those ³⁸⁴ obtained from the coin cell tests, validating the excellent quality ³⁸⁵ of the in situ electrochemical cell. During the first cycle, both ³⁸⁶ $CO₂$ and $O₂$ evolution were detected. $CO₂$ evolution began 387 around 3.8 V during the charge (the first blue guideline in the ³⁸⁸ figure) and completed around the mid-discharge cycle (the ³⁸⁹ second blue guideline). There was an increase in intensity ³⁹⁰ around the midplateau after the onset of O_2 evolution, and the 391 peak $CO₂$ evolution was reached at the end of charge at 4.8 V. 392 Oxygen gas was not detected until the midplateau region at ³⁹³ about 4.5 V (the first red guideline in the figure), which also ³⁹⁴ peaked at the upper cutoff voltage of 4.8 V. Oxygen evolution ³⁹⁵ completes at the beginning of first discharge after the resting ³⁹⁶ step (the second red guideline). The cumulative CO_2 and O_2 397 evolved from the first cycle were 1.53 and 0.11 μ mol, 398 respectively, from a cathode that contained 2.66 mg of active ³⁹⁹ materials. Oxygen evolution, therefore, is much reduced ⁴⁰⁰ compared to what was reported on the lithium and ⁴⁰¹ manganese-rich layered oxide system by Luo et al. 9,10 9,10 9,10 9,10 In the 402 second cycle, the amount of $CO₂$ was reduced from 1.53 to 403 0.46 μ mol, while negligible O₂ evolution was detected.

Several sources may contribute to the evolution of $CO₂$ gas. 405 $Li₂CO₃$, a precursor from synthesis or a byproduct formed 406 during air exposure of Li-TM oxide samples, can decompose ⁴⁰⁷ during charging and release $CO₂$ gas. Recent studies confirmed 408 that in LMR oxide cathodes, 26 all CO₂ gas evolved during the 409 first charge in the typical voltage region, results from residual ⁴¹⁰ $Li₂CO₃$ in the samples. Given the similar outgassing behavior of 411 $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ cathode to LMR, we postulate that residual 412 $Li₂CO₃$ likely accounts for most, if not all, $CO₂$ evolution 413 observed in Figure 4. Nevertheless, carbonate-based electrolytes ⁴¹⁴ are usually unstable at voltages above 4.3 V, especially in the ⁴¹⁵ presence of high valence transition metals (such as Mn^{4+}) that 416 may catalyze the process and further lower the decomposition ⁴¹⁷ voltage, and a small percentage of $CO₂$ evolution at high 418 potentials may originate from this degradation process. ⁴¹⁹ Carbonate solvent decomposition due to the attack of oxygen ⁴²⁰ species has also been reported in the context of Li $-O_2$ 421 batteries.^{[27](#page-10-0)-[29](#page-10-0)} This represents another possible mechanism 422

 for lattice oxygen loss in addition to oxygen gas evolution. In the simplest case where we safely assume only O atoms in O₂ 425 gas originate from $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ cathode material, the total lattice O loss from a cathode containing 2.66 mg of active materials was 0.11 μ mol, which corresponds to 0.38 mol % of oxygen ion vacancy in the delithiated cathode and a capacity of 4.48 mAh/g. Although some of the first charge capacity may 430 also come from Li₂CO₃ oxidizes to CO₂ via a 2 e^{$-$}/CO₂ process and side reactions from electrolyte decomposition, it is evident 432 that the redox process of O^{2-} to O^{n-} (0 < n < 2) species accounts for the majority of the first charge capacity above 4.3 434 V (215 mAh/g) and the first discharge capacity (290 mAh/g) 435 in $Li_{13}Nb_{03}Mn_{04}O_{2}$ cathode.

 The DEMS results also reveal that capacity contribution from O extraction from the oxide lattice occurs above ∼4.5 V. This mechanism is active during the first charge which disappears in the following cycles. This is consistent with the observation on the CV studies, where the oxidation peak centered at 4.7 V was observed only during the first scan. The combined results 442 suggest that the peak at 4.7 V is likely related to O^{2-} oxidation 443 to O_2 gas. The peak centering at 4.5 V, therefore, is attributed 444 to the oxidation O^{2-} anion to O^{n-} $(0 < n < 2)$ species. While both $O₂$ gas evolution and lattice oxygen oxidation occur in the 446 first charge, the following cycles involve only the redox of O^{2-} 447 to O^{n-} (0 < *n* < 2) species, signaled by the single oxidation peak at 4.5 V. The continuous decrease in peak intensity along with scanning indicates that the lattice anion redox process is highly irreversible. After merely nine cycles, the O redox process no longer participates.

 3.3. Structural, Chemical, and Morphological Evolu- tion during the First Delithiation. To gain further insights on structural and chemical changes during the first charge, we resorted to chemical delithiation, which allows for the preparation of samples in the large quantity needed for a number of analytical techniques. This approach also allows us to monitor morphological evolution as large single crystals can be used directly without the prior milling process, a necessary step to electrochemically charge and discharge the sample. 461 Pristine $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ crystals were mixed with various amount of nitronium tetrafluoroborate oxidant in acetonitrile to 463 prepare a series of $Li_xNb_{0.3}Mn_{0.4}O_2$ (x = 1.21, 1.1, 1.06, 1.03, 0.92, 0.87, 0.81, 0.76, 0.64, 0.56, 0.49, 0.40, 0.32, 0.22, 0.13, 0.01, and 0) samples at different states of charge, which were subsequently examined by synchrotron X-ray and neutron diffraction, hard and soft X-ray absorption spectroscopy, and SEM. While the Li content was determined by ICP measurements, the Mn/Nb ratio in each sample was verified to be consistent with the calculated value of 1.33 by using 471 energy dispersive X-ray spectroscopy. Figure 5a shows the powder XRD patterns collected at 11-BM at the Advanced Photon Source. As discussed earlier, the pristine $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ is phase pure with a rock-salt crystal structure $(Fm\overline{3}m; a = 4.1954(1)$ Å). The 4a sites are occupied by Li, Nb, and Mn cations, whereas the 4b sites are occupied only by O anions. Oxygen vacancy was not detected in the pristine sample. With Li extraction, a single rock-salt phase remained 479 when the lithium content is above 0.9 ($x > 0.9$). Further removing Li from the structure led to peak broadening and peak shifting toward to high angle, suggesting an overall reduction in lattice dimension. Similar results were also observed in neutron patterns collected on several samples in the series, as shown in Figure 5b. Joint X-ray and neutron t2 485 Rietveld refinement was performed in cases where the same

Figure 5. (a) Synchrotron X-ray diffraction and (b) monochromatic neutron diffraction patterns collected on chemically delithiated $Li_xNb_{0.3}Mn_{0.4}O_2$ crystals. For neutron measurement, Li1.3 and Li0 samples were measured at SNS (POWGEN) using TOF, while Li1.1, Li0.92, Li0.7, Li0.49, and Li0.22 samples were measured at NIST (BT-1) using a single wavelength of $\lambda = 1.19$ Å. Note that the Li contents were determined by ICP measurements where an error range of up to ± 0.06 is possible.

sample was examined by both techniques. [Figure S2](http://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.7b05036/suppl_file/cm7b05036_si_001.pdf) and [Table](#page-6-0) 486 t2 [2](#page-6-0) show the results obtained from Rietveld refinement of these 487 t2 XRD patterns. The structural transformation can be separated ⁴⁸⁸ into two regions: single phase (phase 1) behavior when $x > 0.9$ 489 (region I) and a two-phase (phase 2 and 3) behavior when $0 < 490$ $x < 0.9$ (region II). The unit cell volume and phase fraction as a 491 function of Li content were constructed from the refinement ⁴⁹² and shown in [Figures 6](#page-7-0)a and b, respectively. The Vegard's law 493 f6 is only followed in region I, where the total volume change is ⁴⁹⁴ ca. 0.2%. This is consistent with the participation of simple ⁴⁹⁵ cation redox from Mn^{3+} to Mn^{4+} along with the extraction of 496 0.4 Li⁺ from the Li_{1.3}Nb_{0.3}Mn_{0.4}O₂ structure. Complex mixed 497 oxygen activities are likely involved in region II, where phases 2 ⁴⁹⁸ and 3 with the same rock-salt crystal structure but different ⁴⁹⁹ lattice dimension evolve along with the extraction of Li. Smaller 500 sized phase 3 reaches the maximum fraction when x is \sim 0.5, soi whereas phase 2 becomes the main phase with further Li ⁵⁰² removal. Combining with the results from the electrochemical 503 studies, we propose that lattice O^{2−} oxidation to O^{n−} (0 < n < s04 2) mainly occurs in phase 3, whereas loss of O from the lattice 505 mainly occurs in phase 2. This was further supported by results ⁵⁰⁶ from the joint X-ray and neutron refinements ([Figure S3\)](http://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.7b05036/suppl_file/cm7b05036_si_001.pdf), ⁵⁰⁷ which revealed that ∼2 and 7.8 mol % of oxygen ion vacancies 508 exist in the delithiated Li_xNb_{0.3}Mn_{0.4}O_{2−δ} with $x = 0.2$ and 0, so9 respectively. Compared to the oxygen ion vacancies in the ⁵¹⁰ electrochemically charged samples, the much higher concen- ⁵¹¹ tration is likely due to the fact that chemical delithiation was ⁵¹²

Table 2. Summary of Structural Parameters in Li_xNb_{0.3}Mn_{0.4}O_{2−6} (0 ≤ x ≤ 1.3) Determined from Rietveld Refinement of Synchrotron X-ray Diffraction and Neutron Data⁶

| $\boldsymbol{\mathcal{X}}$ | 1.3 | 1.21 | 1.1 | 0.92 | 0.87 | 0.49 | 0.22 | 0.13 | 0.01 | $\boldsymbol{0}$ |
|---|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|------------------|
| $V_1(\AA^3)$ | 73.845(1) | 73.839(1) | 73.689(1) | 73.544(1) | | | | | | |
| V_2 (\AA^3) | | | | | 73.84(1) | 73.295(1) | 70.838(1) | 68.93(1) | 68.89(1) | |
| $V_3(\AA^3)$ | | | | | 72.877(2) | 72.494(1) | 68.80(2) | 67.07(3) | 66.69(1) | 68.34(1) |
| n_1 | 1.0 | 1.0 | 1.0 | 1.0 | | | | | | |
| n ₂ | | | | | 0.57(1) | 0.24(1) | 0.93(1) | 0.84(1) | 0.93(1) | |
| n_3 | | | | | 0.43(1) | 0.76(1) | 0.07(1) | 0.16(1) | 0.07(1) | 1.0 |
| $d_1(\AA)$ | 2.0965(1) | 2.0976(1) | 2.0962(1) | 2.0948(1) | | | | | | |
| $d_2(\AA)$ | | | | | 2.0976(5) | 2.0736(7) | 2.0690(2) | 2.0500(4) | 2.0495(3) | |
| $d_3(\AA)$ | | | | | 2.0885(4) | 2.0667(4) | 2.0541(3) | 2.0315(3) | 2.0276(7) | 2.0444(1) |
| $100(U_{\rm iso1})~({\rm \AA}^2)$ | 1.13(1) | 1.22(1) | 1.44(1) | 1.47(1) | | | | | | |
| | 1.72(1) | 1.90(1) | 2.18(1) | 2.28(2) | | | | | | |
| $100(U_{\text{iso}})$ (Å ²) | | | | | 1.55(3) | 2.39(1) | 4.11(1) | 4.75(6) | 3.03(4) | |
| | | | | | 2.53(5) | 3.15(1) | 4.85(1) | 6.08(9) | 3.59(5) | |
| $100(U_{\text{iso3}})(\AA^2)$ | | | | | 2.92(4) | 2.39(1) | 4.11(1) | 4.75(6) | 3.03(4) | 5.46(6) |
| | | | | | 2.75(6) | 3.15(1) | 4.85(1) | 6.08(9) | 3.59(5) | 6.15(8) |
| δ | 0(0) | | 0(0) | | | 0(0) | 0.02(1) | | | 0.08(1) |
| $_{\rm w}R_{\rm p}$ (%) | 9.22 | 6.78 | 6.85 | 7.40 | 7.25 | 8.91 | 6.71 | 8.70 | 6.91 | 8.16 |

 aV_1 , V_2 , V_3 , n_1 , n_2 , n_3 , d_1 , d_2 , d_3 , U_{iso1} , U_{iso2} , and U_{iso3} represent the unit cell volume, phase fraction, bond distance of Li/Mn/Nb−O), and thermal factors for Li/Nb/Mn and O in phase 1, 2, and 3, respectively. δ represents oxygen anion vacancy in each phase, while ${}_{w}R_{p}$ represent the residual factor of the refinement.

 performed in an open system that shifts the equilibrium toward more gas evolution. We also like to point out that protons were detected on the chemically delithiated oxides based on the prompt-gamma activation analysis (PGAA). This is likely molecular in nature as delithiated samples generally have higher surface area (as shown in the SEM images in [Figure 9\)](#page-9-0) and tend to absorb water during brief air exposure. However, it is possible that the detected protons may originate from cation exchange between $Li⁺$ and $H⁺$, as previously reported on the 522 classic layered oxides.^{[30](#page-10-0)} Further analysis is needed to fully understand the source of protons in these samples.

 The kinetic properties of delithiation were evaluated by the relationship between the molar ratio of oxidant/oxide used and the residual Li content in the sample [\(Figure 6](#page-7-0)c), as determined by ICP. While a linear relationship is apparent in region I, significant deviation is seen in region II, further confirming the complex nature of mixed anion activities at lower Li contents. Hard X-ray absorption spectroscopy correlates the changes in the absorption edge energy to the variations in the average oxidation state of the absorbing atoms being analyzed in the 533 bulk material.^{[31](#page-10-0),[32](#page-10-0)} The technique also probes the local and electronic structures near the absorbing atoms and has the ability to reveal element-specific structural changes in the samples. Due to the energy limit of the XAS beamline at SSRL, Nb spectra were not collected in this study. A previous report from Yabuuchi et al. indicated that Nb remains at 5+ during the entire charge/discharge process.^{[22](#page-10-0)} The normalized spectra of Mn K-edge and XANES spectra of the chemically delithiated 541 Li_xNb_{0.3}Mn_{0.4}O₂ series are shown in [Figures 7](#page-8-0)a and b, 542 respectively. The spectrum of pristine $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ is 543 consistent with that of Mn³⁺ collected on the standard reference. Substantial blue-shift of the edge energy was observed as the lithium content was reduced from 1.3 to 0.9, 546 consistent with the oxidation of Mn^{3+} to Mn^{4+} . Negligible changes were observed in samples with the Li content below 0.9. The edge position, defined by Photoelectron Energy Origin (E₀), is a commonly used parameter for extracting edge energy 550 level in XANES spectra.^{[33](#page-10-0)} [Figure 7c](#page-8-0) compares the edge position

as a function of lithium content in $Li_xNb_{0.3}Mn_{0.4}O_2$ samples. It 551 is clear that Mn K-edge energy shifts toward higher value in a 552 near linear fashion when $x > 0.9$ (region I) whereas it remains 553 nearly constant when $0 < x < 0.9$ (region II). This is consistent 554 with the results from X-ray and neutron diffraction studies, ⁵⁵⁵ further confirming that Mn is only redox active in region I. 556

Further changes were observed in the intensity of Mn pre- ⁵⁵⁷ edge absorption peaks arising from the dipole forbidden 1s \rightarrow 558 3d electronic transitions, which are typically weak in the ⁵⁵⁹ transition metals. In the presence of 3d and 4p orbital ⁵⁶⁰ hybridization, often resulting from structural distortion in local 561 symmetry or noncentrosymmetric environment between the 562 metal and oxygen coordination, the transitions are made ⁵⁶³ partially allowed, and the peaks become much stronger.^{[34](#page-10-0)–[36](#page-10-0)} As 564 shown in [Figure 7](#page-8-0)d, the intensity of the pre-edge absorption 565 peaks is similar in region I, which becomes noticeably stronger 566 in region II. This suggests that although Mn redox activities are 567 not involved in region II, the nature of Mn−O bond is altered ⁵⁶⁸ in these samples, providing further evidence for oxygen redox ⁵⁶⁹ activities in this region as the processes are likely behind the ⁵⁷⁰ changes in local symmetry and environment of metal−ligand ⁵⁷¹ coordination. Similar to the edge shift observed in [Figures 7a](#page-8-0) ⁵⁷² and b, the absorption energy of the pre-edge peaks also 573 experienced blue shift in region I while maintaining nearly ⁵⁷⁴ constant in region II, corroborating with the observation of Mn ⁵⁷⁵ redox activity in region I only. 576

Oxygen redox activities during delithiation of 577 $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ were investigated by soft XAS that directly 578 probes the chemical changes of the element as a function of ⁵⁷⁹ penetration depth. The depth profiling from the surface to bulk 580 were achieved by using two different detectors in total electron ⁵⁸¹ yield (TEY) and fluorescence yield (FY) modes, which have a ⁵⁸² typical probing depth of 5 and 50 nm, respectively.^{[37](#page-10-0)} [Figures 8a](#page-8-0) 583.68 and b show the evolution of TEY and FY O K-edge XAS ⁵⁸⁴ spectra as a function of Li content in $Li_xNb_{0.3}Mn_{0.4}O_2$. In both s85 cases, the spectra can be separated into the pre-edge region and 586 postedge region at the black dashed line (533 eV) shown in the $_{587}$ figures. Features in the postedge region are often attributed to ⁵⁸⁸

Figure 6. (a and b) Cell volume and phase fraction as a function of x in delithiated $Li_xNb_{0,3}Mn_{0,4}O_2$ samples and (c) relationship between the molar ratio of oxidant/oxide and residue Li content in chemically delithiated $Li_xNb_{0.3}Mn_{0.4}O_2$ samples.

589 O1s \rightarrow O2p-TM4s4p hybridization, while those in the pre-edge 590 region are attributed to O1s \rightarrow O2p-TM3d hybridization.³ Due to the crystal field effect and octahedral coordination of oxygen, the pre-edge peaks are often represented by multiplets between 528 and 533 eV. As the O2p and TM3d band electrons are of most interest to redox activities, we focus on the analysis in the pre-edge region. It is clear that as lithium is 596 extracted from $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$, both TEY and FY spectra exhibit a significant increase in the pre-edge peak intensity. This increase is often used as a qualitative measure for the generated holes left by electron extraction from either oxygen or TM at the O2p-TM3d energy levels.[39](#page-10-0)−[41](#page-10-0) A new shoulder peak at ∼529 eV also appeared and gradually grew along with delithiation, suggesting the involvement additional O-TM 603 hybridization (referred to as p -band) during the process. Due to similarity between the p-band and the pre-edge absorption 605 peak often observed in peroxides and superperoxides, $22,40,42,43$ we speculate this new O-TM hybridization arises from the interaction between oxygen anions as a result of oxygen redox activities. Note that similar results were previously reported on 609 the LMR cathode materials. $22,40$

To better understand the oxygen redox activities as a ⁶¹⁰ function of Li content, quantifications of peak features in both ⁶¹¹ TEY and FY O K-edge XAS spectra were performed, and the ⁶¹² results are shown in [Figures 8](#page-8-0)c and d. The detailed peak fitting ⁶¹³ method is also shown in [Figure S4.](http://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.7b05036/suppl_file/cm7b05036_si_001.pdf) For both TEY and FY ⁶¹⁴ spectra, the ratio between the pre-edge peaks and postedge ⁶¹⁵ peaks continuously increase with lithium extraction, as shown ⁶¹⁶ in [Figure 8c](#page-8-0). This confirms electron extraction from O2p and ⁶¹⁷ TM 3d hybridization and agrees with the previous report by ⁶¹⁸ Yabuuchi et al. 22 22 22 It is, however, unclear on the contribution 619 from TM and O due to the hybridization nature between them. ⁶²⁰ The quantification on the p -band intensity, on the other hand, 621 provides an opportunity to directly probe oxygen redox ⁶²² activities. As shown in [Figure 8d](#page-8-0), the initial p -band intensity 623 is fairly low in the pristine $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ and remains nearly 624 constant below Li content of ∼0.9, suggesting negligible ⁶²⁵ interactions between oxygen anions or oxygen redox activities. ⁶²⁶ As x decreases to below 0.9, there is an increase in the p -band 627 intensity, and the trend continues to full delithiation $(x = 0)$. 628 The results suggest a continuous increase in oxygen valence 629 states in this region and the participation of oxygen redox for ⁶³⁰ charge compensation. Combined with the results from the Mn ⁶³¹ K edge XAS spectra which shows that Mn is active when $x > 632$ 0.9 and inactive when $x < 0.9$, the quantification of O K-edge 633 XAS spectra enables the understanding of complete charge ⁶³⁴ compensation mechanism during delithiation of ⁶³⁵ $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ crystals. Note that the p-band intensity in 636 the $x < 0.9$ region is generally higher in the FY mode than that 637 in the TEY mode, consistent with a difference in oxygen ⁶³⁸ activities, likely enhanced lattice O loss on the surface. 639

The evolution of particle morphology during the first ⁶⁴⁰ delithiation is revealed by the SEM images collected on the ⁶⁴¹ series of delithiated crystal samples [\(Figure 9](#page-9-0)). Some surface 642 f9 roughing was observed but the particles remain intact when $x > 643$ 0.9. With further lithium extraction, large cracks begin to ⁶⁴⁴ appear, which tend to propagate across the entire particle, ⁶⁴⁵ leading to the eventual fracturing of the crystals. The number of ⁶⁴⁶ cracks on each crystal increases along with the decrease in x but 647 appears to reach the maximum at a Li content of 0.5−0.6. This ⁶⁴⁸ coincides with the occurrence of maximum fraction of phase 3, ⁶⁴⁹ as shown in Figure 6b. The change in morphology is also ⁶⁵⁰ consistent with the fact that Mn is redox active when $x > 0.9$ 651 while O is redox active when $x < 0.9$. The participation of 652 oxygen redox appears to cause significant morphological ⁶⁵³ damage. Our results also suggest that lattice oxidation of O^{2-} 654 to O^{n-} (0 < n < 2) is more damaging than O_2 gas evolution, 655 although it is possible that particles become more stable when x 656 is less than 0.5−0.6 as mechanical stress is released by certain ⁶⁵⁷ cracks and fractures at that point. Further evaluation on oxygen ⁶⁵⁸ oxidation and mechanical damage in Li-excess oxides is ⁶⁵⁹ warranted, but we emphasize that particle cracking and ⁶⁶⁰ fracturing create fresh surfaces which can contribute to ⁶⁶¹ increased O loss from the lattice and side reactions with the ⁶⁶² electrolyte. Successful surface stabilization strategies such as ⁶⁶³ elemental segregation and coating treatment therefore need to ⁶⁶⁴ take consideration in particle morphology evolution during ⁶⁶⁵ cycling. ⁶⁶⁶

4. CONCLUSION

Disordered, Li-excess transition-metal oxide cathodes are ⁶⁶⁷ capable of delivering high capacities and are promising next- ⁶⁶⁸ generation LIB cathodes. In this study, uniform and phase-pure ⁶⁶⁹ $Li_{1,3}Nb_{0,3}Mn_{0,4}O_2$ single crystals were synthesized by a molten- 670

Figure 7. (a and b) Mn K-edge hard XAS and XANES spectra, (c) XANES energy edge (E_0) as a function of Li content in Li_nNb_{0.3}Mn_{0.4}O₂ samples, and (d) Mn K-edge pre-edge spectra.

Figure 8. (a and b) O K-edge soft XAS spectra obtained in TEY and FY modes, (c) intensity ratio between the pre-edge and postedge absorption peaks as a function of lithium content, and (d) intensity of the p-band as a function of lithium content. In panels a and b, black dashed line divides the pre- and postedge regions, and the blue dashed arrow follows the evolution of the p-band intensity.

 salt method for the first time and used as a diagnostic tool to understand the charge compensation mechanism during Li extraction and insertion. Electrochemical evaluation suggests structural irreversibility after the first cycle, leading to capacity and voltage fade of the cathode. Both $CO₂$ and $O₂$ gas evolution was found during the first cycle, with the latter only accounting for ∼1% of the charge capacity, in addition to the 678 contribution of ~35% capacity from the Mn^{3+/4+} redox couple. A large fraction of capacity therefore originates from the redox 680 process of Q^{2-}/Q^{n-} (0 < n < 2). Joint refinement on

synchrotron X-ray and neutron diffraction patterns of $_{681}$ $Li_xNb_{0.3}Mn_{0.4}O_2$ series showed that the oxide undergoes a $_{682}$ single-phase reaction involving $Mn^{3+/4+}$ redox reaction when 683 0.9 < x < 1.3 and a two-phase reaction involving mixed O^{2-}/e^{84} O^{n-} (0 $\leq n < 2$) reactions when 0 < x < 0.9. For the fully ₆₈₅ delithiated sample, \sim 7.8 mol % of oxide-ion vacancies were 686 found, resulting from O loss from the lattice. Cracks and $_{687}$ fractures initiated and intensified with deep lithium extraction, 688 which serves as a means to release the mechanical stress ⁶⁸⁹

Figure 9. SEM images of chemically delithiated $\text{Li}_x\text{Nb}_{0.3}\text{Mn}_0\text{4O}_2$ crystals: $x = (a)$ 1.3, (b) 1.1, (c) 0.87, (d) 0.76, (e) 0.56, and (f) 0.13.

⁶⁹⁰ endured by the large crystals during the oxygen redox ⁶⁹¹ processes.

⁶⁹² ■ ASSOCIATED CONTENT

693 Supporting Information

⁶⁹⁴ The Supporting Information is available free of charge on the ⁶⁹⁵ [ACS Publications website](http://pubs.acs.org) at DOI: [10.1021/acs.chemma-](http://pubs.acs.org/doi/abs/10.1021/acs.chemmater.7b05036)⁶⁹⁶ [ter.7b05036.](http://pubs.acs.org/doi/abs/10.1021/acs.chemmater.7b05036)

697 SEM image of carbon mixed $LiNb_{0.3}Mn_{0.4}O₂$ crystals after ball milling, Rietveld refinement of synchrotron X- ray diffraction patterns collected on chemically delithi-700 ated $Li_xNb_{0.3}Mn_{0.4}O_2$ crystals, Rietveld refinement of synchrotron X-ray and neutron diffraction patterns collected on chemically delithiated $Li_0Nb_{0.3}Mn_{0.4}O_2$ crystals, and process used to quantify O K-edge XAS data ([PDF](http://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.7b05036/suppl_file/cm7b05036_si_001.pdf))

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- 713 Notes

⁷¹⁴ The authors declare no competing financial interest.

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(1) Goodenough, J. B.; Manthiram, A. A perspective on electrical 736 energy storage. MRS Commun. 2014, 4, 135-142. 737

(2) Li, W.; Song, B.; Manthiram, A. High-voltage positive electrode 738 materials for lithium-ion batteries. Chem. Soc. Rev. 2017, 46, 3006− 739 3059. 740

(3) Manthiram, A.; Knight, J. C.; Myung, S.-T.; Oh, S.-M.; Sun, Y.-K. 741 Nickel-Rich and Lithium-Rich Layered Oxide Cathodes: Progress and 742 Perspectives. Adv. Energy Mater. 2016, 6, 1501010. 743

(4) Aydinol, M. K.; Kohan, A. F.; Ceder, G.; Cho, K.; Joannopoulos, 744 J. Ab initio study of lithium intercalation in metal oxides and metal 745 dichalcogenides. Phys. Rev. B: Condens. Matter Mater. Phys. 1997, 56, 746 1354−1365. 747

(5) McCalla, E.; Abakumov, A. M.; Saubanere, M.; Foix, D.; Berg, E. 748 J.; Rousse, G.; Doublet, M.-L.; Gonbeau, D.; Novak, P.; Van Tendeloo, 749 G.; Dominko, R.; Tarascon, J.-M. Visualization of O-O peroxo-like 750 dimers in high-capacity layered oxides for Li-ion batteries. Science 751 (Washington, DC, U. S.) 2015, 350, 1516−1521. 752

(6) Pearce, P. E.; Perez, A. J.; Rousse, G.; Saubanere, M.; Batuk, D.; 753 Foix, D.; McCalla, E.; Abakumov, A. M.; Van Tendeloo, G.; Doublet, 754 M.-L.; Tarascon, J.-M. Evidence for anionic redox activity in a 755 tridimensional-ordered Li-rich positive electrode $β$ -Li₂IrO₃. Nat. Mater. 756 2017, 16, 580–586. 757

(7) Sathiya, M.; Rousse, G.; Ramesha, K.; Laisa, C. P.; Vezin, H.; 758 Sougrati, M. T.; Doublet, M. L.; Foix, D.; Gonbeau, D.; Walker, W.; 759 Prakash, A. S.; Ben Hassine, M.; Dupont, L.; Tarascon, J. M. Reversible 760 anionic redox chemistry in high-capacity layered-oxide electrodes. Nat. 761 Mater. 2013, 12, 827−835. 762

(8) Armstrong, A. R.; Holzapfel, M.; Novak, P.; Johnson, C. S.; Kang, 763 S.-H.; Thackeray, M. M.; Bruce, P. G. Demonstrating Oxygen Loss 764 and Associated Structural Reorganization in the Lithium Battery 765 Cathode Li $[Ni_{0.2}Li_{0.2}Mn_{0.6}]O_2$. J. Am. Chem. Soc. 2006, 128, 8694− 766
8698. 8698. 767

(9) Luo, K.; Roberts, M. R.; Guerrini, N.; Tapia-Ruiz, N.; Hao, R.; 768 Massel, F.; Pickup, D. M.; Ramos, S.; Liu, Y.-S.; Guo, J.; Chadwick, A. 769 V.; Duda, L. C.; Bruce, P. G. Anion Redox Chemistry in the Cobalt 770

771 Free 3d Transition Metal Oxide Intercalation Electrode Li-772 [Li_{0.2}Ni_{0.2}Mn_{0.6}]O₂. J. Am. Chem. Soc. 2016, 138, 11211−11218.

773 (10) Luo, K.; Roberts, M. R.; Hao, R.; Guerrini, N.; Pickup, D. M.;

 Liu, Y.-S.; Edstrom, K.; Guo, J.; Chadwick, A. V.; Duda, L. C.; Bruce, P. G. Charge-compensation in 3d-transition-metal-oxide intercalation cathodes through the generation of localized electron holes on oxygen. Nat. Chem. 2016, 8, 684−691.

 (11) Shukla, A. K.; Ramasse, Q. M.; Ophus, C.; Duncan, H.; Hage, F.; Chen, G. Unravelling structural ambiguities in lithium- and manganese-rich transition metal oxides. Nat. Commun. 2015, 6, 8711. (12) Mohanty, D.; Li, J.; Abraham, D. P.; Huq, A.; Payzant, E. A.;

782 Wood, D. L.; Daniel, C. Unraveling the Voltage-Fade Mechanism in 783 High-Energy-Density Lithium-Ion Batteries: Origin of the Tetrahedral

 Cations for Spinel Conversion. Chem. Mater. 2014, 26, 6272−6280. (13) Wang, D.; Huang, Y.; Huo, Z.; Chen, L. Synthesize and electrochemical characterization of Mg-doped Li-rich layered Li- $\left[\text{Li}_{0.2}\text{Ni}_{0.2}\text{Mn}_{0.6}\right]$ O₂ cathode material. Electrochim. Acta 2013, 107, 788 461−466.

 (14) Song, B.; Zhou, C.; Wang, H.; Liu, H.; Liu, Z.; Lai, M. O.; Lu, L. Advances in sustain stable voltage of Cr-doped Li-rich layered cathodes for lithium ion batteries. J. Electrochem. Soc. 2014, 161, A1723−A1730.

793 (15) Li, G. R.; Feng, X.; Ding, Y.; Ye, S. H.; Gao, X. P. AlF3-coated 794 Li(Li0.17Ni0.25Mn0.58)O2 as cathode material for Li-ion batteries. 795 Electrochim. Acta 2012, 78, 308−315.

 (16) Shi, S. J.; Tu, J. P.; Tang, Y. Y.; Liu, X. Y.; Zhang, Y. Q.; Wang, X. L.; Gu, C. D. Enhanced cycling stability of Li- $\left[\text{Li}_{0.2}\text{Mn}_{0.54}\text{Ni}_{0.13}\text{Co}_{0.13}\right]\text{O}_2$ by surface modification of MgO with melting impregnation method. Electrochim. Acta 2013, 88, 671−679. (17) Zhang, X.; Belharouak, I.; Li, L.; Lei, Y.; Elam, J. W.; Nie, A.; Chen, X.; Yassar, R. S.; Axelbaum, R. L. Structural and Electrochemical 802 Study of Al_2O_3 and TiO₂ Coated $Li_{1,2}Ni_{0,13}Mn_{0,54}Co_{0,13}O_2$ Cathode Material Using ALD. Adv. Energy Mater. 2013, 3, 1299−1307.

804 (18) Kim, K. J.; Jo, Y. N.; Lee, W. J.; Subburaj, T.; Prasanna, K.; Lee, 805 C. W. Effects of inorganic salts on the morphological, structural, and 806 electrochemical properties of prepared nickel-rich $Li[Ni_{0.6}Co_{0.2}Mn_{0.2}]$ -807 O₂. J. Power Sources 2014, 268, 349–355.

 (19) Shigemura, H.; Tabuchi, M.; Sakaebe, H.; Kobayashi, H.; Kageyama, H. Lithium Extraction and Insertion Behavior of 810 Nanocrystalline Li_2TiO_3 -LiFe O_2 Solid Solution with Cubic Rock Salt Structure. J. Electrochem. Soc. 2003, 150, A638−A644.

 (20) Tabuchi, M.; Nakashima, A.; Shigemura, H.; Ado, K.; Kobayashi, H.; Sakaebe, H.; Tatsumi, K.; Kageyama, H.; Nakamura, 814 T.; Kanno, R. Fine $Li_{(4-x)/3}Ti_{(2-2x)/3}Fe_xO_2$ (0.18 $\leq x \leq 0.67$) powder with cubic rock-salt structure as a positive electrode material for rechargeable lithium batteries. J. Mater. Chem. 2003, 13, 1747−1757. (21) Lee, J.; Urban, A.; Li, X.; Su, D.; Hautier, G.; Ceder, G. Unlocking the Potential of Cation-Disordered Oxides for Rechargeable Lithium Batteries. Science (Washington, DC, U. S.) 2014, 343, 519− 820 522.

 (22) Yabuuchi, N.; Takeuchi, M.; Nakayama, M.; Shiiba, H.; Ogawa, M.; Nakayama, K.; Ohta, T.; Endo, D.; Ozaki, T.; Inamasu, T.; Sato, K.; Komaba, S. High-capacity electrode materials for rechargeable 824 lithium batteries: Li_3NbO_4 -based system with cation-disordered 825 rocksalt structure. Proc. Natl. Acad. Sci. U. S. A. 2015, 112, 7650-7655. (23) Toby, B. H. EXPGUI, a graphical user interface for GSAS. J. Appl. Crystallogr. 2001, 34, 210−213.

 (24) Xu, J.; Renfrew, S.; Marcus, M. A.; Sun, M.; McCloskey, B. D.; 829 Tong, W. Investigating $Li_2NiO_2-Li_2CuO_2$ Solid Solutions as High- Capacity Cathode Materials for Li-Ion Batteries. J. Phys. Chem. C 2017, 121, 11100−11107.

832 (25) Edington, J. W. Practical Electron Microscopy in Materials Science; 833 Van Nostrand Reinhold Co., 1976.

 (26) Renfrew, S.; McCloskey, B. D. Residual lithium carbonate 835 predominantly accounts for first cycle $CO₂$ and CO outgassing of Li- stoichiometric and Li-rich layered transition metal oxides. J. Am. Chem. Soc. 2017, 139, 17853−17860.

838 (27) Freunberger, S. A.; Chen, Y.; Peng, Z.; Griffin, J. M.; Hardwick, 839 L. J.; Barde, F.; Novak, P.; Bruce, P. G. Reactions in the Rechargeable Lithium-O₂ Battery with Alkyl Carbonate Electrolytes. J. Am. Chem. 840 Soc. 2011, 133, 8040-8047.

(28) Aurbach, D.; Daroux, M.; Faguy, P.; Yeager, E. The 842 electrochemistry of noble metal electrodes in aprotic organic solvents 843 containing lithium salts. J. Electroanal. Chem. Interfacial Electrochem. 844 1991, 297, 225−44. 845

(29) Aurbach, D.; Gofer, Y.; Langzam, J. The correlation between 846 surface chemistry, surface morphology, and cycling efficiency of 847 lithium electrodes in a few polar aprotic systems. J. Electrochem. Soc. 848 1989, 136, 3198. 849

(30) Choi, J.; Alvarez, E.; Arunkumar, T. A.; Manthiram, A. Proton 850 insertion into oxide cathodes during chemical delithiation. Electrochem. 851 Solid-State Lett. 2006, 9, A241−A244. 852

(31) Terada, Y.; Yasaka, K.; Nishikawa, F.; Konishi, T.; Yoshio, M.; 853 Nakai, I. In Situ XAFS Analysis of $Li(Mn,M)$, O_4 (M = Cr, Co, Ni) 5 V 854 Cathode Materials for Lithium-Ion Secondary Batteries. J. Solid State 855 Chem. 2001, 156, 286–291. 856

(32) Nam, K.-W.; Bak, S.-M.; Hu, E.; Yu, X.; Zhou, Y.; Wang, X.; Wu, 857 L.; Zhu, Y.; Chung, K.-Y.; Yang, X.-Q. Combining In Situ Synchrotron 858 X-Ray Diffraction and Absorption Techniques with Transmission 859 Electron Microscopy to Study the Origin of Thermal Instability in 860 Overcharged Cathode Materials for Lithium-Ion Batteries. Adv. Funct. 861 Mater. 2013, 23, 1047−1063. 862

(33) Hu, E.; Bak, S.-M.; Liu, J.; Yu, X.; Zhou, Y.; Ehrlich, S. N.; Yang, 863 X.-Q.; Nam, K.-W. Oxygen-Release-Related Thermal Stability and 864 Decomposition Pathways of $Li_xNi_{0.5}Mn_{1.5}O_4$ Cathode Materials. Chem. 865 Mater. 2014, 26, 1108−1118. 866

(34) Kim, M. G.; Shin, H. J.; Kim, J.-H.; Park, S.-H.; Sun, Y.-K. XAS 867 Investigation of Inhomogeneous Metal-Oxygen Bond Covalency in 868 Bulk and Surface for Charge Compensation in Li-Ion Battery Cathode 869 $Li[Ni_{1/3}Co_{1/3}Mn_{1/3}]O_2$ Material. J. Electrochem. Soc. 2005, 152, 870 A1320-A1328.

(35) Kim, M. G.; Yo, C. H. X-ray Absorption Spectroscopic Study of 872 Chemically and Electrochemically Li Ion Extracted $Li_yCo_{0.85}Al_{0.15}O_2$ 873
Compounds, J. Phys. Chem. B 1999. 103, 6457–6465. Compounds. J. Phys. Chem. B 1999, 103, 6457–6465.

(36) Ignatov, A. Y.; Ali, N.; Khalid, S. Mn K-edge XANES study of 875 the $La_{1-x} CaxMnO₃$ colossal magnetoresistive manganites. *Phys. Rev. B: 876*
Condens. Matter Mater. *Phys.* **2001**. 64. 014413/1–014413/16. 877 Condens. Matter Mater. Phys. 2001, 64, 014413/1-014413/16. (37) Stö hr, J. NEXAFS Spectroscopy; Springer, 1992. 878

(38) Yoon, W.-S.; Balasubramanian, M.; Chung, K. Y.; Yang, X.-Q.; 879 McBreen, J.; Grey, C. P.; Fischer, D. A. Investigation of the Charge 880 Compensation Mechanism on the Electrochemically Li-Ion Dein- 881 tercalated $Li_{1-x}Co_{1/3}Ni_{1/3}Mn_{1/3}O_2$ Electrode System by Combination 882 of Soft and Hard X-ray Absorption Spectroscopy. J. Am. Chem. Soc. 883 2005, 127, 17479−17487. 884

(39) de Groot, F. M. F.; Grioni, M.; Fuggle, J. C.; Ghijsen, J.; 885 Sawatzky, G. A.; Petersen, H. Oxygen 1s x-ray-absorption edges of 886 transition-metal oxides. Phys. Rev. B: Condens. Matter Mater. Phys. 887 1989, 40, 5715−5723. 888

(40) Oishi, M.; Yamanaka, K.; Watanabe, I.; Shimoda, K.; Matsunaga, 889 T.; Arai, H.; Ukyo, Y.; Uchimoto, Y.; Ogumi, Z.; Ohta, T. Direct 890 observation of reversible oxygen anion redox reaction in Li-rich 891 manganese oxide, $Li₂MnO₃$, studied by soft X-ray absorption 892 spectroscopy. J. Mater. Chem. A 2016, 4, 9293-9302. 893

(41) Ma, C.; Alvarado, J.; Xu, J.; Clément, R. J.; Kodur, M.; Tong, 894 W.; Grey, C. P.; Meng, Y. S. Exploring Oxygen Activity in the High 895 Energy P2-Type $\text{Na}_{0.78}\text{Ni}_{0.23}\text{Mn}_{0.69}\text{O}_2$ Cathode Material for Na-Ion 896 Batteries. J. Am. Chem. Soc. 2017, 139, 4835–4845. 897

(42) Kang, J. S.; Kim, D. H.; Hwang, J. H.; Baik, J.; Shin, H. J.; Kim, 898 M.; Jeong, Y. H.; Min, B. I. Soft x-ray absorption and photoemission 899 spectroscopy study of superoxide KO₂. Phys. Rev. B: Condens. Matter 900
Mater. Phys. **2010**, 82, 193102. Mater. Phys. 2010, 82, 193102.

(43) Yilmaz, E.; Yogi, C.; Yamanaka, K.; Ohta, T.; Byon, H. R. 902 Promoting Formation of Noncrystalline Li₂O₂ in the Li−O₂ Battery 903 with RuO₂ Nanoparticles. Nano Lett. 2013, 13, 4679–4684. 904