Gradient-morph LiCoO₂ single crystals with stabilized energy density above 3400 W h L⁻¹

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The cycling stability of LiCoO₂ under high voltages (>4.5 V) was plagued by hybrid anion- and cation-redox (HACR) induced oxygen escape and uncontrolled phase collapse. With DEMS and in situ XANES mapping at the NSLS-II, we demonstrate that oxygen escape triggers irreversible transformations into "bad" surface phases that rapidly propagate inward. Enabling HACR but stopping global oxygen migration is key to a stable high-energy cathode. Therefore, we developed ~10 μm single crystals with LiCoO₂ in the bulk smoothly transitioning to Co-free LiMn₀.７₅Ni₀.２₅O₂ at the surface. By means of initial electrochemical formation, a semi-coherent LiMn₁.₅Ni₀.₅O₄ spinel-like shell was established in operando with little oxygen loss to integrity wrap the LiCoO₂ bulk. Then we obtained gradient-morph LiCoO₂ single crystals to prevent the percolating migration of oxygen out of the particle and achieved enhanced HACR reversibility at high voltages. The gradient-morph HACR cathode undergoes substantially stabilized cycling when charged to above 4.6 V, and hence a stable cyclic volumetric energy density of >3400 W h L⁻¹ has been achieved in a pouch full-cell coupled with a commercial graphite anode and lean electrolyte (2 g A h⁻¹), exhibiting up to 2906 W h L⁻¹ even after 300 cycles.

Though diverse transition-metal (TM) oxides have been investigated as cathode materials for Li-ion batteries (LIB), LiCoO₂ (LCO) still takes the dominant position in the current LIB market for consumer electronics even after Goodenough’s¹ and SONY’s² developments, due to the good rate performance, simple scalability and high compressed density of above 4.1 g cm⁻³. However, LCO is only working within a limited voltage range (<4.35 V) or capacity (<165 mAh g⁻¹) at present in order to maintain the cycling stability,³ which is far from the theoretical capacity of 274 mA h g⁻¹ when Li is fully extracted, so it still provides great potential to achieve much higher energy density.⁴ The cycling stability of LCO to high charging voltages (>4.5 V) has been investigated for decades.⁴⁻⁶ Some literature studies reported⁷⁻¹⁰ that bulk phase transformations (including O³⁻→H₁⁻→O¹) occurring above 4.5 V caused rapid degradation of LCO, while others reported that the highly delithiated Li₁-xCoO₂ (x > 0.5) aggravated the interfacial side reactions with the electrolyte.¹¹⁻¹³ However, we have argued that the exact sequence of events in bulk vs. surface degradation initiators need careful analysis.¹⁴

Because of the heavy Co₃d–O₂p hybridization (Fig. 1a, note that its energy resonance is much more than that of Ni₃d–O₂p or Mn₃d–O₂p, indicating stronger covalency), deep delithiation from LCO must extract electrons from both Co and O ions. Manthiram et al.¹⁵ reported that when L₁₋ₓCoO₂ was delithiated beyond x = 0.5, further delithiation requires the oxidation of O²⁻ (O³⁻→O⁵⁻, x < 2), which has lately been verified by soft X-ray absorption spectroscopy (sXAS).¹⁶,¹⁷ That means hybrid anion- and cation-redox (HACR) reactions must be involved when charging LCO to above ~150 mA h g⁻¹. Although utilizing the last one-half of LCO's theoretical capacity (from 150 mA h g⁻¹ to 274 mA h g⁻¹) in the high-voltage range is extremely enticing, the involvement of oxygen anion-redox and the associated oxygen mobility (O²⁻ has a much lower migration energy barrier than O³⁻¹⁸⁻²⁰ and escape,²⁰ causing lattice collapse and side reactions with the necessarily lean electrolyte in industrial batteries, is a killer problem.

The significant HACR activities entail O²⁻ oxidation, which would make oxygen ions more mobile¹⁸⁻²⁰ and enable oxygen migration (OM). OM can be local or global: global OM (GOM) means the oxygen ions can move long-range in the particle all the way to the surface and be released from the particle surface, possibly reacting with the liquid electrolyte and causing oxygen depletion with a particularly pernicious irreversible phase...
that "infect" the interior, producing flaws and micro-cracks, further driving GOM and IPT propagating deep into the interior. In this work, we will demonstrate the oxygen release at high voltages to promote fast electrochemical kinetics. (3) The shell should not participate in oxygen anion-redox and exclude oxygen vacancy in cycling to eliminate percolation paths for GOM. According to the band diagram in Fig. 1a, the shell should be Co-free (e.g. M can be Mn, Ni etc.), which is not highly hybridized with O) to avoid the oxygen oxidation at high voltages. These requirements point to a high-voltage spinel LiMn1.5Ni0.5O4 phase as a likely candidate for the shell. The LiMn1.5Ni0.5O4 shell is well-known to be very stable up to 5 V, so one should expect a stable high-voltage cycling, when the spinel shell completely encloses and protects the LCO core.

However, so far researchers have never achieved a complete-wetting and lattice-coherent LiMn1.5Ni0.5O4 shell to fully wrap around the LCO lattice. With most of the coating methods, one tends to get islands-coating instead of an integral shell. The islands-coating cannot shut down GOM in cycling. So in this work, we developed a special way to exactly establish such an integral lattice-coherent LiMn1.5Ni0.5O4 spinel shell to fully enclose LCO and shut down GOM in the high-voltage cycling.

In this work, we first developed lattice-coherent LiCoO2 and LiMn0.75Ni0.25O2 single crystals. Because LiMn0.75Ni0.25O2 is isostructural with LCO (both layered), it is easily able to fully

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**Fig. 1** Electronic bands, lattice structures and preparation process of LiCoO2 and the gradient XyLiCoO2-(1-Xy)LiMn0.75Ni0.25O2 single-crystal. (a) The band structures in LiCoO2 and Co-free LMO (M = Mn, Ni etc.). (b) The surface reconstruction of the gradient XyLiCoO2-(1-Xy)LiMn0.75Ni0.25O2 crystal in the initial electrochemical formation. In this process, the oxygen sublattice maintains lattice-coherency during the layered LiMn0.75Ni0.25O2 → spinel LiMn1.5Ni0.5O4 transformation, with preserved Mn : Ni : O ratio and no oxygen loss. The blue balls represent Co ions, while the violet, green and red balls represent Mn/Ni, Li and O ions, respectively. (c) Schematic showing the creation of the gradient XyLiCoO2-(1-Xy)LiMn0.75Ni0.25O2 single-crystal with an abnormal grain growth at high temperature.
and integrally enclose the LCO bulk by high-temperature annealing. As is known, such a Mn-rich LiMn$_{0.75}$Ni$_{0.25}$O$_2$ layered phase is not electrochemically stable and thus rarely prepared for batteries, but here it is just coated as a precursor for the eventual integral LiMn$_{1.5}$Ni$_{0.5}$O$_4$ shell. As shown in Fig. 1b and c, the pre-prepared particles are composed of χ$_0$/LiCoO$_2$-(1 – χ$_0$)LiMn$_{0.75}$Ni$_{0.25}$O$_2$. The core, with χ(χ$_0$ core) = 1, is pure LCO, and χ$_0$ gradually decreases to 0 (χ(χ$_0$ surface) = 0) from the bulk to the surface. That is, the composition in the particle changes from LCO to LiMn$_{0.75}$Ni$_{0.25}$O$_2$ from the core to the shell, but all M and O ions share a coherent layered crystal lattice (Fig. 1b), constituting a gradient LiMn$_{1.5}$Ni$_{0.5}$O$_4$ shell with epitaxial layer crystal structure that can transform into the desirable fully wetting LiMn$_{1.5}$Ni$_{0.5}$O$_4$ shell with spinel crystal structure, as illustrated in Fig. 1b. The LiMn$_{1.5}$Ni$_{0.5}$O$_4$ spinel-like shell that in operando transformed from the lattice-coherent LiMn$_{0.75}$Ni$_{0.25}$O$_2$ by Jahn-Teller distortion in room-temperature cycling will still keep oxygen sublattice-coherency with the LCO bulk and fully enclose LCO. Crucially, although the final LiMn$_{1.5}$Ni$_{0.5}$O$_4$ has a cubic crystal structure while the LiMn$_{0.75}$Ni$_{0.25}$O$_2$ precursor has a layered structure, unlike LiCoO$_2$ → Co$_2$O$_4$, this LiMn$_{0.75}$Ni$_{0.25}$O$_2$ → LiMn$_{1.5}$Ni$_{0.5}$O$_4$ transformation preserves the O: Mn: Ni ratios and requires neither oxygen loss nor V$_0$ injection into the particle that would enhance OM, as only the O: Li ratio changes from 2 to 4. This is the secret for the orderly and controlled LiMn$_{0.75}$Ni$_{0.25}$O$_2$ → LiMn$_{1.5}$Ni$_{0.5}$O$_4$ surface transformation (TM/O all in place and only displacive Jahn-Teller distortion is needed) upon formation by electrochemical delithiation, and the destination of this surface transformation, LiMn$_{1.5}$Ni$_{0.5}$O$_4$, is unarguably the best high-voltage coating phase as it enables quick Li$^+$ conduction but prevents GOM. Though LiMn$_{1.5}$Ni$_{0.5}$O$_4$ is cubic spinel while LCO has a layered structure, the spinel LiMn$_{1.5}$Ni$_{0.5}$O$_4$ shell established in this way still shares the same oxygen-sublattice with the LCO bulk, so we would like to call the particle after the battery-formation process a gradient-morph LCO single crystal (GM-LCO, Fig. 1b). GM-LCO can achieve the following unique performances in prolonged high-voltage cycling: (1) with ~10 μm single crystals, 4.1 g cm$^{-3}$ compressed density and high electronic conductivity, the cathode can cycle to ~190 mA h g$^{-1}$, ~230 mA h g$^{-1}$ and ~270 mA h g$^{-1}$ when charged to 4.5 V, 4.6 V and 4.7 V vs. Li, respectively. While the spinel-like shell contributes little capacity, it is very stable and can always keep crystal dense in cycling. Though the O$^2-$ ions in the LCO bulk participate in HACR and intend to migrate when charged to above 4.5 V, they are completely enclosed and can hardly trespass the shell. GOM and IPT can thus be efficiently prevented, which “tames” the oxygen ions in the LCO core to be a highly-reversible anion-redox (“solid oxygen”) HACR concept like the Li–sulfur chemistry in the high-voltage cycling. In this work, we will show that the cycling stability of G-LCO is highly respectable for consumer electronics applications even when cycled to 4.7 V. (2) Substantially different from the surface passivation, the LiMn$_{1.5}$Ni$_{0.5}$O$_4$ shell conducts Li$^+$ and polarons exceptionally well, preserving the facile kinetics of LCO in cycling, despite the large single crystals for high compressed density. (3) By keeping the Mn valence at +4 in cycling, the LiMn$_{1.5}$Ni$_{0.5}$O$_4$ shell prevents both Co and Mn dissolutions, which can further stabilize the cycling of the practical full-cells versus graphite anodes. Furthermore, the new particles also have better compatibility with the commercial carbonate electrolyte at high voltages, so we will show that G-LCO has ultra-stable high-voltage cycling in a pouch full-cell with a commercial graphite anode and very lean electrolyte (2 g A h$^{-1}$).

Preparation of χ($\rho$)LiCoO$_2$-(1 – χ($\rho$))-LiMn$_{0.75}$Ni$_{0.25}$O$_2$ gradient single crystals

As the Mn valence in the LiMn$_{0.75}$Ni$_{0.25}$O$_2$ precursor is below +4, such a layered crystal was rarely prepared in air as bulk material, as it is thermodynamically unstable. However, we will demonstrate that a thin coherent LiMn$_{0.75}$Ni$_{0.25}$O$_2$ shell can be created on the surface of a huge LCO particle, which acts as a giant seed for the rapid grain coarsening that consumes the original Li/Mn/Ni oxide nanograins, with a composition-gradient region between the LCO core and the LiMn$_{0.75}$Ni$_{0.25}$O$_2$ shell. Such epitaxial substrate has a stabilizing effect on the surface layer phase, as the elimination of phase boundary energies and grain boundary energies compensates for the increased bulk energy, as long as the LiMn$_{0.75}$Ni$_{0.25}$O$_2$-like shell is thinner than the coherency limit. In this work, we created a lattice-coherent LiMn$_{0.75}$Ni$_{0.25}$O$_2$ shell on the pristine LCO (P-LCO) particle with a wet-coating process followed by high-temperature annealing (Method). The deposited Li/Mn/Ni has an opportunity to self-nucleate to other crystals (e.g. spinel) and grow into islands on the LCO particle surface. However, when we deposit only tens of nanometers thick layers of Li/Mn/Ni precursors on the surface of a ~10 μm primary LCO particle, the huge LCO crystal will easily coarsen away the thin Li/Mn/Ni oxide nanocrystals in the calculation process due to grain coarsening, as shown in Fig. 1c. This means that the coated Li/Mn/Ni oxides can directly grow into the same layered lattice with the LCO substrate following the path of the abnormal grain growth (AGG). Because LiMn$_{0.75}$Ni$_{0.25}$O$_2$ is isostructural with LCO, the two components can mutually diffuse and form a lattice-coherent solid solution single crystal when annealed at high temperature. Therefore, the new particle can remain single-crystalline, but one can expect pure LCO in the core and Mn/Ni/Co gradient concentration in the shell.

We characterize the G-LCO particles before the electrochemical cycling, which was produced with 94 wt% LCO and 6 wt% LiMn$_{0.75}$Ni$_{0.25}$O$_2$. The X-ray diffraction (XRD) pattern in Fig. 2a indicated that G-LCO particles were still composed of a single phase, as there was no other peak observed than the R3m phase peak. Fig. 2b and c show the scanning electron microscopy (SEM) images of P-LCO and G-LCO particles. It was clear that P-LCO particles had a well-crystallized morphology and smooth surface with a particle size of 10–15 μm in diameter.
but the particles became round and were completely enclosed after the formation of the lattice-coherent shell. The STEM image of the particle cross-section (after FIB) in Fig. 2d indicated that there was no phase boundary in G-LCO particles. The STEM-HAADF images in Fig. 2e and f further proved the single-crystallinity of the particle, as the (001) lattice plane arranged well from the deep core to the outermost surface. EDS mapping under SEM in the ESI,† Fig. S2, verified the uniform and full conformal coverage of the Mn shell on the LCO particle, and that under STEM in Fig. 2d accurately indicated that the TM element in the core was only Co and the content gradually decreased in the ~150 nm surface region and to 0% (Co-free) in the ~10 nm outermost surface, where Mn and Ni were concentrated. There was not a sharp edge between the Mn/Ni and Co distributions, indicating that the two compositions of LiMn$_{0.75}$Ni$_{0.25}$O$_2$ and LCO formed a solid solution with a smooth gradient transition that promoted the lattice-coherency between the LiMn$_{0.75}$Ni$_{0.25}$O$_2$ shell and LCO core. The Mn/Co/Ni distribution in G-LCO was quantified with EELS line-scan from the shell to core (Fig. 2g and h), and the result showed that Co was ~33% (consistent with LCO) in the bulk but gradually decreased in the ~150 nm surface region and to 0% (Co-free) in the ~10 nm outermost surface, where Mn and Ni gradually concentrated from 0% to ~25% and ~8% (consistent with LiMn$_{0.75}$Ni$_{0.25}$O$_2$) while O remained constant at ~67% across the whole particle.

The above measurements indicated the successful preparation of the gradient X$_{r}$LiCoO$_2$/(1-X$_{r}$)LiMn$_{0.75}$Ni$_{0.25}$O$_2$ single crystals, with X$_{r}$core = 1 and X$_{r}$surface = 0. Additionally, according to the sXAS analysis in the ESI,† Fig. S3, the Mn valence can be quantified as +3.4 at the surface, quite close to the theoretical value in LiMn$_{0.75}^{(+3.33)}$Ni$_{0.25}^{(+2)}$O$_2$, which further verified the composition of LiMn$_{0.75}$Ni$_{0.25}$O$_2$ at the particle surface. Generally, the composition profile of G-LCO can be further optimized by adjusting the ratio of Li/Mn/Ni, M elements and annealing temperatures etc. to get better electrochemical performances.

Again we want to emphasize that the layered LiMn$_{0.75}^{(+3.33)}$Ni$_{0.25}^{(+2)}$O$_2$ is not a typical compound that people can easily synthesize or use electrochemically in the bulk. But here by virtue of the large
LCO single crystal as the seed and substrate, we stabilized this LiMn$_{0.75}$Ni$_{0.25}$O$_2$ phase as a thin epitaxial surface layer, that serves as the precursor for the future LiMn$_{0.75}$Ni$_{0.25}$O$_2$ transformation by O and TM preserving displacive Jahn–Teller distortion. Because of the pre-positioning of Mn : Ni : O in the right ratio, and because the good spinel can still share the same oxygen sublattice with the LCO bulk, only minimum structural disruption is needed to achieve a conformal protection with the “good spinel”, in a highly organized and well-controlled surface transformation during the initial cycles.

**Stabilized high cyclic energy density up to 3400 W h L$^{-1}$ in pouch full-cells**

We prepared the cathode electrodes with a standard industrial procedure, comprising 96 wt% active sample, 2 wt% carbon black and 2 wt% PVDF, with $\sim 17$ mg cm$^{-2}$ loading of active material on an Al foil. The P-LCO and G-LCO cathodes were firstly cycled in half-cells between 3.0–4.6 V. As shown in Fig. 3a, P-LCO and G-LCO had similar capacity in the 1st cycle: P-LCO was charged to 252.1 mA h g$^{-1}$ and discharged to 224.5 mA h g$^{-1}$, whereas G-LCO was charged to 251.2 mA h g$^{-1}$ and discharged to 226.9 mA h g$^{-1}$. However, Fig. 3a shows that the charge/discharge profile of P-LCO totally faded after 100 cycles, while that of G-LCO kept a similar shape to the initial cycle after 100 cycles. The cycling performance in Fig. 3b shows that the capacity of P-LCO decayed rapidly in the initial 30 cycles, and only 45% of the initial capacity and 34% of the initial energy density were retained after 40 cycles. However, G-LCO retained more than 80% of the initial capacity and energy density after 100 cycles. The cyclic voltammetry of P-LCO and G-LCO after different cycles can be found in the ESI.$^\dagger$ Fig. S4.

We use the coulombic inefficiency (CI $\equiv 100\% - \text{CE}$, which is defined as the difference between coulombic efficiency and 100%) in half-cells to compare the capacity loss from the cathode in each cycle.$^{10}$ As shown in Fig. 3b, the CI of G-LCO was only about half that of P-LCO in each cycle, indicating a $\sim 50\%$ suppressed capacity loss in each discharge from the G-LCO cathode. The suppressed CI of G-LCO not only indicated thermodynamic stability, but could also indicate a maintained kinetics of G-LCO during each charging and discharging,$^{41}$ while the higher CI of P-LCO reflected a serious kinetic retardation in each cycle when the growth of “bad” spinel Co$_3$O$_4$ in charging immediately added more impedance for the subsequent discharging. Therefore the difference in cyclic capacity between P-LCO and G-LCO is enlarged at higher rates. As shown in Fig. 3c, G-LCO had a similar capacity to P-LCO in the initial cycles at 20 mA g$^{-1}$, but the capacity retention of P-LCO decreased considerably at high rates, while that of G-LCO decreased little. In particular, at 400 mA g$^{-1}$ after 40 cycles, G-LCO can still discharge 175 mA h g$^{-1}$, three-times higher than...
that of P-LCO (41 mA h g⁻¹). When reducing the rate to 20 mA g⁻¹, the capacity of G-LCO reverted to 210 mA h g⁻¹ whereas that of P-LCO restored to 163 mA h g⁻¹. That is, the kinetic capacity loss of G-LCO, when increasing the rate from 20 mA g⁻¹ to 400 mA g⁻¹, was only about 1/4 that of P-LCO after 50 cycles. This result clearly indicated that the impedance of G-LCO was stabilized in the high-voltage cycling.

We then tested P-LCO and G-LCO cathodes in pouch full-cells with matched commercial graphite anodes (Method). As shown in Fig. 3d, though both P-LCO and G-LCO can discharge to about 205 mA h g⁻¹ and 780 W h kg⁻¹ in the initial cycles when cycled within 3.0–4.50 V, either the capacity or energy density of P-LCO suffered rapid decay in 60 cycles and totally died within 200 cycles. However, G-LCO had very little decay in the cycling, as the capacity remained stable above 191.2 mA h g⁻¹ with an energy density of 736.3 W h kg⁻¹ after 300 cycles. When cycled within 3.0–4.55 V, G-LCO discharged to 216 mA h g⁻¹ and 844 W h kg⁻¹ in the initial cycles, and retained 176.3 mA h g⁻¹ with 664.7 W h kg⁻¹ after 300 cycles. The previous highest energy density of LCO obtained in pouch full-cells vs. graphite anode was also around 800 W h kg⁻¹, reported by Zhang et al.⁵ but the cyclic retention decayed to below 700 W h kg⁻¹ in 70 cycles. The G-LCO in this work has achieved energy-density retention of 736.3 W h kg⁻¹ after 300 cycles. Additionally, as we used only a very lean electrolyte (2 g A h⁻¹) in the full-cell, the good cycling also proved that G-LCO had much better compatibility with the regular carbonate electrolyte at high voltages.⁴²

The stabilized high energy density significantly improved the volumetric energy density of the cathode. Both the compressed densities of the P-LCO and G-LCO electrodes were measured as 4.1 ± 0.1 g cm⁻³ (ESI† Table S1), the same as that provided by the current commercial manufacturers. Then the volumetric energy density of P-LCO and G-LCO cathodes can be compared in Fig. 3e, calculated from the cycling performance in the full-cell (Fig. 3d). Though the initial energy densities of the P-LCO and G-LCO cathodes were similar when charged to 4.50 V, both around 3100 W h L⁻¹, P-LCO decayed to almost 0 after 300 cycles. However, the volumetric energy density of the LCO cathode can remain as high as 2906 W h L⁻¹ after 300 cycles. When cycled to 4.55 V, G-LCO can even achieve 3404 W h L⁻¹ and retain 2793 W h L⁻¹ after 300 cycles. As far as we know, this is the first time that one achieved such a high cyclic volumetric energy density of cathode after 300 cycles, which is ~30% higher than that of the commercial LCO on the market cycled within 4.35 V.

**Prevention of GOM and Co₃O₄ in the full-delithiated G-LCO particles**

The rapid degradation of P-LCO in the high-voltage cycling will be shown to be caused by GOM and the accumulated IPT. Firstly, we used differential electrochemical mass spectroscopy (DEMS) to monitor the O₂ and CO₂ release while charging the cathodes to 4.60 V vs. Li. As shown in Fig. 4a, O₂ started to evolve from P-LCO at 4.35 V (~175 mA h g⁻¹), and the amount linearly increased afterward and especially sped up when holding the cathode at 4.60 V. The accompanying CO₂ release also verified the rampant decomposition of the precious electrolyte when oxygen was released from the cathode. In contrast, GOM was substantially prevented in G-LCO particles at high voltages. As shown in Fig. 4b, little O₂ was released from G-LCO even when holding the cathode at 4.60 V for more than 1 hour. The elimination of CO₂ release also indicated a stabilized interface between the G-LCO particles and the carbonate electrolyte when OM was prevented, which will not only favor the stabilization of the particle surface, but also prevent the depletion of the precious electrolyte to stabilize the cycling, as according to the industrial practice, the total weight of the liquid electrolyte injected is only ~40% of the cathode weight and must be able to wet the anode, cathode and separator even after thousands of cycles.

The occurrence of OM would immediately result in IPT in the P-LCO particle, producing the bad spinel Co₃O₄, where the average Co valence is below +3. Hereby, we utilized an in situ nano-XANES mapping with synchrotron-based full-field X-ray imaging at the NSLS-II of the Brookhaven National Laboratory, which can monitor the spatial distribution of Co valence when charging the cathode (Methods). Thanks to its high spatial resolution (~21 nm) and high temporal resolution,⁴³⁻⁴⁵ we can track the detailed information of IPT from the *operando* spatial Co valence change during charging. Fig. 4c–e showed the in situ Co valence distribution in P-LCO at different states of charges (SOCs). Firstly, we can verify that the particle surface always delithiated more than the bulk while charging, as Fig. 4d shows that the Co valence is higher (in green) at the surface than in the bulk (in yellow). When only focusing on site S at the particle surface, we can note that Co was firstly oxidized when charging P-LCO from 3.0 V (Fig. 4e) to 4.2 V (Fig. 4d), indicating the routine delithiation of LiCoO₂ (ESI† Fig. S5a), it can be inferred that Co₃O₄ was produced at 4.6 V at site S. This is the first time that, using an in situ characterization method, one confirmed that GOM and IPT (CoO₂ → Co₃O₄) started from some specific patches at the LCO particle surface at high voltages. Meanwhile, we also obtained a gradient Co-valence distribution near site S at 4.6 V (Fig. 4e and ESI† Fig. S5a), which showed that the Co ions in the core were able to be oxidized to around +4, but those near the surface were gradually reduced to +2.7. The gradient distribution of Co valence indicated the growth of Co₃O₄ inward in the bulk. When Co₃O₄ wrapped the P-LCO particles in the prolonged cycles, both the capacity and voltage were seen to be severely degraded, as shown in Fig. 3. Very differently, the stopped GOM prevented catastrophic uncontrolled changes to “bad” surface phases in the G-LCO particle. As shown in Fig. 4g–j, all Co ions in the particle were continuously oxidized from +3 to about +4 while charging G-LCO from 3 V to 4.6 V. There was no patch showing reduced Co at the particle surface (ESI† Fig. S5b) in the whole charging process, indicating the
prevention of $\text{Co}_3\text{O}_4$ formation at high voltages. The prevented GOM and Co reduction in G-LCO while charging maintained high kinetics, thus promoting a well suppressed CI within 0.5% and enhanced capacity retention in the prolonged high-voltage cycling as shown in Fig. 3.

To fundamentally understand the prevention of GOM in G-LCO particles, it is essential to investigate the shell reconstruction in the high-voltage cycling. Because of Jahn–Teller distortion,23,24 the layered Mn-based LMO (like LiMnO$_2$) would prefer transforming into a spinel-like phase when sufficiently delithiated by charging to above 4.0 V. In this work, the STEM-HAADF images and FFT patterns in Fig. 5a–f clearly indicated that the Co-free LiMn$_{0.75}$Ni$_{0.25}$O$_2$ shell transformed into a LiMn$_{1.35}$Ni$_{0.25}$O$_4$ spinel-like phase when sufficiently delithiated by charging to above 4.0 V. In this work, the STEM-HAADF images and FFT patterns in Fig. 5a–f clearly indicated that the Co-free LiMn$_{0.75}$Ni$_{0.25}$O$_2$ shell transformed into a LiMn$_{1.35}$Ni$_{0.25}$O$_4$ spinel-like phase when sufficiently delithiated by charging to above 4.0 V. In this work, the STEM-HAADF images and FFT patterns in Fig. 5a–f clearly indicated that the Co-free LiMn$_{0.75}$Ni$_{0.25}$O$_2$ shell transformed into a LiMn$_{1.35}$Ni$_{0.25}$O$_4$ spinel-like phase when sufficiently delithiated by charging to above 4.0 V. In this work, the STEM-HAADF images and FFT patterns in Fig. 5a–f clearly indicated that the Co-free LiMn$_{0.75}$Ni$_{0.25}$O$_2$ shell transformed into a LiMn$_{1.35}$Ni$_{0.25}$O$_4$ spinel-like shell (G-LCO → GM-LCO) after the initial cycles, whereas the LCO core still kept a layered structure.47 Though this LiMn$_{0.75}$Ni$_{0.25}$O$_2$ → LiMn$_{1.35}$Ni$_{0.25}$O$_4$ transformation that occurred in the shell was also irreversible, the operando-created LiMn$_{1.3}$Ni$_{0.5}$O$_4$ shell kept a semi-coherent lattice with the LCO core instead of producing phase boundaries or microcracks (Fig. 5d), as both the LCO core and LiMn$_{1.3}$Ni$_{0.5}$O$_4$ shell kept a coherent oxygen sublattice (ESI† Fig. S6, Table S2 and Discussion S1). Additionally, neither this phase transformation nor the subsequent high-voltage cycling involves oxygen loss (no O$^2_-$ was oxidized or released) in the spinel-like shell, so neither VO nor flaws were pumped into the particle. The quantitative sXAS analysis in Fig. 5g, showing that the Mn valence at the discharge state at the G-LCO particle surface changed from +3.4 to +4 after a few initial cycles, was absolutely consistent with the LiMn(+3.33)$^0.75$Ni(+2)$^0.25$O$_2$ → LiMn(+4)$^0.75$Ni(+2)$^0.25$O$_4$ transformation, and verified the formation of a VO-free shell (if there was VO in the LiMn$_{1.3}$Ni$_{0.5}$O$_4$ spinel, it must contain Mn$^{3+}$ ions$^{48,49}$). Therefore, the robust coherent VO-free LiMn$_{1.3}$Ni$_{0.5}$O$_4$ shell can stop GOM in the LCO lattice, even though HACR and O$^2_-$ oxidation still occur very actively inside the LCO core. Unlike the spinel...
Co$_3$O$_4$ in P-LCO, the well-known high-voltage spinel LiMn$_{1.5}$Ni$_{0.5}$O$_4$ shell does not block Li$^+$ diffusion or electron conductivity, but supplies a 3D 8a-16c-8a path to promote fast Li$^+$ diffusion in cycling.\textsuperscript{24,25}  

sXAS was performed to directly investigate HACR by tracking the oxidation states of O at high-voltages. Both P-LCO and G-LCO cathodes were charged to 4.7 V and then held for 48 h, and the FY sXAS O K-edges were recorded in this charge process. As shown in Fig. 5h and i, both the FY sXAS O K-edges of P-LCO and G-LCO were broadened toward higher energy when charged to 4.7 V, indicating that both O$^{2-}$ ions were oxidized to O$^{n-}$ ($x < 2$)\textsuperscript{5,17} in the two materials. However, after holding at 4.7 V for 48 h, the peak intensity due to O$^{2-}$ on the FY O K-edge of P-LCO decreased a lot, indicating that partial O$^{2-}$ ions were reduced to O$^{n-}$ due to GOM (e.g. O$^{2-}$ $\rightarrow$ O$^{n-}$ + O$_2$). But the O K-edge of G-LCO remained nearly unchanged after 48 h, confirming that the oxidized O$^{n-}$ was significantly stabilized in the G-LCO particle. The sXAS analysis strongly proved the prevention of GOM in the G-LCO particles at high voltages, which agreed well with the DEMS and \textit{in situ} XANES mapping results in Fig. 4. The cycling stability of the fully-charged G-LCO to 4.7 V was also much improved compared to that of P-LCO (ESI,\textsuperscript{†} Fig. S7).

When GOM was prevented in the G-LCO particles by the \textit{operando}-created semi-coherent spinel-like nanoshell, the layered single crystal structure of the LCO core could be maintained in the high-voltage cycling, while IPT (Co$_3$O$_4$) continuously grew into the

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**Fig. 5** Surface reconstruction in the battery formation process and prevented oxygen migration in G-LCO particles. (a) STEM image of the G-LCO particle, showing the phase transformation into a spinel-like phase in the shell, while the core still keeping a layered structure in the G-LCO particle. (b and c) FFT patterns transformed from the spinel surface (b) and LCO bulk (c), as marked in (a) with yellow circles. (d) STEM-HAADF image of the intersection of the shell and the core, showing the semi-coherency between the \textit{operando}-created spinel-like shell and the LCO core. Inserted are the simulated spinel LiMn$_{1.5}$Ni$_{0.5}$O$_4$ structure (left) and LiCoO$_2$ structure (right), where the green and red balls represent Co ions and oxygen ions, while the navy-blue, violet and grey balls represent Mn and Ni ions, respectively. (e and f) FFT patterns transformed from the spinel surface (e) and LCO bulk (f), picked up from the left region and right region in (d). (g) sXAS TEY Mn L$_3$-edge of G-LCO at the discharge state before cycling and after 2 cycles; the Mn valence was quantitatively analyzed by linear-fitting from the standard MnO, Mn$_2$O$_3$ and MnO$_2$ references. (h and i) sXAS FY O K-edge spectra of P-LCO (h) and G-LCO (i) at different states of charge in the initial cycle.
bulk of P-LCO. Fig. 6a shows the SEM image of P-LCO after 200 cycles between 3.0–4.60 V in half cells, which indicated that severely cracked fragments can be found everywhere at the particle surface. The collapsed surface phase (IPT) can grow thick enough towards the bulk and seriously block the charge transfer reaction. As shown in the HRTEM image of the cross-section of P-LCO after FIB preparation (Fig. 6b and c), the spinel Co$_3$O$_4$ almost spread across the whole particle from the surface (site A) to the center (site B). There were also lots of oxygen voids (in red circles in Fig. 6c) observed in the particle bulk because of the continuous oxygen migration and escape in the high-voltage cycling.\textsuperscript{18} The selected area electron diffraction (SAED) pattern in Fig. 6d clearly verified that lots of the layered structure had transformed into spinel Co$_3$O$_4$ in P-LCO after 200 cycles. However, very differently, from the SEM image in Fig. 6e, G-LCO still kept robust single-crystal morphology after 200 cycles between 3.0–4.6 V, without crack or fragment at the particle surface. The HRTEM image of G-LCO after FIB in Fig. 6f and g further indicated the well maintained layered structure throughout the whole LCO core after cycling, as the (001) lattice plane was well arranged from the deep core (site D) up to the shell (site C). The SAED pattern in Fig. 6h additionally proved that most of the layered structure was maintained in the LCO core.

The stabilized phase in G-LCO would maintain good kinetics in the high-voltage cycling, which can be better understood by comparing the Li$^+$ diffusivity ($D_{Li}$) between P-LCO and G-LCO in the full-cells. The potentiostatic intermittent titration technique (PITT) was used to quantify $D_{Li}$ in the charging process. As shown in Fig. 7a, the $D_{Li}$ values of P-LCO and G-LCO were similar in the 3$^{rd}$ charging, both between $10^{-12}$–$10^{-11}$ cm$^2$ s$^{-1}$; however $D_{Li}$ of P-LCO decreased to around $10^{-14}$ cm$^2$ s$^{-1}$ while that of G-LCO still remained around $10^{-12}$ cm$^2$ s$^{-1}$ after 200 cycles. Note that $D_{Li}$ here refers to the diffusion of Li$^+$ and polaron in the full-cell, so $D_{Li}$ here can significantly indicate the practical electrochemical kinetics of the full-cell.

Besides the cathode, the detailed condition of the graphite anode can also affect the kinetics when cycling the full-cell. The solid electrolyte interface (SEI) at the anode was analyzed by EDX mapping, as shown in Fig. 7b–f. It clearly indicated that while a similar amount of F was present at the graphite anode by comparing Fig. 7b and d, there was a massive amount of Co deposited on the surface of the graphite anode after cycling with P-LCO (Fig. 7c), which was greatly suppressed at the anode after cycling with G-LCO (Fig. 7e). The suppression of Co dissolution from the cathode and deposition at the anode significantly maintained the stability of the SEI and interfacial Li$^+$ transfer.\textsuperscript{13,50} Additionally, as we expected, there was also very little Mn detected on the surface of the graphite anode after cycling with G-LCO (Fig. 7f), because the Mn in the shell always kept +4 in the high voltage cycling (Fig. 5g) in LiMn$_{1.5}$Ni$_{0.5}$O$_4$, which have little dissolution in the carbonate electrolyte\textsuperscript{31,52} unlike the Mn$^{3+}$ or Mn$^{2+}$ cations.
Conclusions

We have investigated the global oxygen migration (GOM) and irreversible phase transformation (IPT) while charging LCO to high voltages by DEMS and in situ XANES mapping, and demonstrated that the reduction of transition metal cations started from some patches at the pristine LCO particle surface upon charging to above 4.3 V, which immediately triggered IPT into a bad spinel (CoO$_2$-Co$_3$O$_4$) to block the Li-ion transport, resulting in a rapid cycling fading. In order to prevent the harmful GOM and uncontrollable phase transformations at high voltages, we intentionally created a thin LiMn($^{+3.33}$)$_{0.75}$Ni($^{+2}$)$_{0.25}$O$_2$ layer epitaxially wrapping the LCO single crystal seed by high-temperature processing and abnormal grain growth, which templates a future O-and-TM-preserving surface transformation into the good spinel LiMn$_{1.5}$Ni$_{0.5}$O$_4$ that shares the same oxygen sublattice with LCO. We have proved that the oxidized oxygen ions were significantly stabilized in the LCO core when it was even fully-charged. The elimination of GOM stabilizes hybrid anion- and cation-redox (HACR) reactions in the high-voltage cycling, and preserves the good kinetics of LCO single crystals. The test in pouch full-cells with a commercial graphite anode further showed an extremely stable cycling performance when cycled to 4.55 V. Therefore, a cathode electrode with a super high volumetric energy density of 2906 W h L$^{-1}$ has been achieved after 300 cycles in a full-cell.

For generalization, this gradient-morph (GM-LMO) single-crystal structure can also be applied to other cathode materials as a general strategy to establish a desirable specific shell that cannot be directly created on a particle. We can firstly coat the particle with an intermediate layer that is isostructural with the bulk to realize the integral wrapping by a regular high-temperature reaction (prepare G-LMO first), and then it can transform in operando into the eventual desirable shell (e.g. spinel, G-LMO $\rightarrow$ GM-LMO) to fully enclose and protect the particle in prolonged cycling.

Methods

Preparation of the LiMn$_{0.75}$Ni$_{0.25}$O$_2$ $\rightarrow$ LiCoO$_2$ gradient morph single crystals

The pristine LiCoO$_2$ single crystals were synthesized by a solid-reaction method. Firstly, Co$_3$O$_4$ (≥99%, Sigma-Aldrich) and Li$_2$CO$_3$ (ACS Reagent, ≥99%, Sigma-Aldrich) were sufficiently mixed with a mole ratio of 1:1.5 (with 5% excess of Li$_2$CO$_3$); then the mixture was heated at 1000–1100 °C for 10 hours with a heating and cooling rate of 5 °C min$^{-1}$ to get the pristine LCO particles (P-LCO).

The pristine LCO particles were centrifuged with water to remove the small particles with size below micro-meters, and then ultrasonicated in an ethanol solution with LiOOCH$_3$ (Reagent Plus$^\text{®}$, ≥99%, Sigma-Aldrich), Mn(COOCH$_3$)$_2$ (Reagent Plus$^\text{®}$, ≥99%, Sigma-Aldrich) and Ni(COOCH$_3$)$_2$ (Reagent Plus$^\text{®}$, ≥99%, Sigma-Aldrich) dissolved with a mole ratio of 1.05:0.75:0.25. After that, the mixture was dried in a 60–80 °C water bath with stirring. Finally, the obtained powder was heated at 900 °C for 8 hours with a heating and cooling rate of 5 °C min$^{-1}$ to get the pristine LCO particles (P-LCO).

For generalization, this gradient-morph (GM-LMO) single-crystal structure can also be applied to other cathode materials as a general strategy to establish a desirable specific shell that...
Electrochemical test

The cathode electrode was made of 96 wt% active material, 2 wt% carbon black, and 2 wt% polyvinylene fluoride (PVDF) binder, which was pasted on an Al current collector and compressed under 20 MPa. R2032 coin cells were fabricated with the above cathode, Li metal anode, a Celgard 2400 polymeric separator and a commercial electrolyte solution of 1.2 M LiPF₆ dissolved in a mixture of EC and DEC with a volume ratio of 1:1, and 2 wt% vinylene carbonate additive. Pouch cell-fulls were fabricated with the above cathodes, a commercial graphite anode (double-side coated), a separator of Celgard 2400 polymer, and a commercial electrolyte solution of 1.2 M LiPF₆ dissolved in a mixture of EC and DEC with a volume ratio of 1:1, and 2 wt% vinylene carbonate additive. The loading density of LCO cathode was about ~17 mg cm⁻² with ~3.5 mA h cm⁻² (tested in half-cell under 0.1C); the commercial graphite anode had a loading density of ~12 mg cm⁻² with ~3.8 mA g cm⁻² (tested in half-cell under 0.1C). The pouch cell fulls were fabricated with double layers of electrodes (one graphite anode foil with both sides coated and two LCO cathode foils with a single side coated). The amount of electrolyte added was about 2 g A h⁻¹ in the pouch cell. A LAND CT2001A 8-channel automatic battery test system (Wuhan Lanhe Electronics) was used for charging/discharging of the cells. An electrochemical workstation (Gamry Instr, Reference 3000) was used for the cyclic voltammetry (CV) test between 3.0 V and 5.0 V with 0.05 mV s⁻¹ scanning rate, and for the potentiostatic intermittent titration technique (PITT) with constant potential for 200 seconds followed by 1800 seconds of relaxation with a voltage-step of 40 mV from 3.8 V to 4.6 V. The electrochemical tests were all carried out at room temperature. A self-made quantitative DEMS connected to a Thermo Scientific mass spectrometer (MS). High-purity Ar gas was used as the carrier gas with a flow rate of 3 mL min⁻¹ during the cycling process. In the constant current charging process, the current was 50 mA g⁻¹, and MS spectra were collected every 30 seconds.

Characterizations

X-ray diffraction was performed on a PANalytical X’Pert PRO X-ray diffractor using a Cu target under 45 kV and 40 mA. The data were collected with 2° min⁻¹ and analyzed with the HighScore Plus software. Scanning electron microscopy was performed on a Zeiss Merlin high-resolution scanning electron microscope with X-ray spectroscopy (EDX), High-angle annular dark field (HAADF) images, energy dispersive X-ray spectroscopy (EDS) and electron energy loss spectroscopy (EELS) spectra were acquired from a JEOL ARM 200F scanning transmission electron microscope (STEM) operated at 200 keV, equipped with a cold field emission gun and integrated aberration (Cs) corrector. Dual EELS spectrometers were used to collect both the low-loss and high-loss spectra, and the low-loss spectra were utilized to correct the drift of the zero-loss peak (ZLP). Based on the full-width at half-maximum (FWHM) of the zero-loss peak (ZLP), the optimal energy-resolution of EELS was about 1 eV. The STEM sample was prepared using an FEI Helios NanoLab 600 DualBeam FIB/SEM equipped with a Ga ion source. A Pt layer was deposited on top to protect the cathode particle. High-resolution transmission electron microscopy (HRTEM) images and selected area electron diffraction (SAED) patterns were taken on a JEOL 2010F transmission electron microscope operated at 200 kV.

In situ nano-XANES mapping

In situ nano-XANES mapping was performed at the FXI beamline (18-ID) at the National Synchrotron Light Source II (NSLS-II), Brookhaven National Laboratory. Pouch-cell configurations with P-LCO and G-LCO as cathodes were used for measurements. Each pouch cell was charged at a constant current of 8 mA and stopped at different voltages (4.2 V, 4.4 V and 4.6 V) while collecting XANES spectra. A volume in the middle of the cell was randomly chosen for imaging. XANES images were taken at different energies across the Co absorption edge (7.588–8.153 keV, 1 eV interval). The effective pixel size of each image was 21 nm. Standard samples (CoO and LiCoO₂) were used to extract the reference absorption spectra for Co²⁺ and Co³⁺ oxidation states.

Soft X-ray absorption spectroscopy (sXAS) measurements

Soft X-ray absorption spectroscopy (sXAS) measurements were carried out at the IOS beamline (23-ID-2) at the National Synchrotron Light Source II (NSLS-II), Brookhaven National Laboratory. Spectra were acquired in total electron yield (TEY) and partial fluorescence yield (PFY) modes. The estimated incident X-ray energy resolution was ~0.05 eV at the O K-edge. The monochromator absorption features and beam instabilities were normalized out by dividing the detected PFY and TEY signals by the drain current of a clean gold 10 mesh placed in the incident beam. TEY spectra were recorded from the drain current of the sample and PFY data were acquired using a Vortex EM silicon drift detector. The sXAS spectra for the O K-edge were recorded over a wide energy range from 520 to 565 eV covering energies well below and above sample absorptions. The normalization of the O K-edge was performed. The sample signal was divided by the incident intensity measured from the sample drain current from a freshly coated Au mesh inserted into the beam path before the X-rays could impinge on the sample. (2) The linear, sloping background was removed by fitting a line to the flat low energy region (520–524 eV) of the sXAS spectrum. (3) The spectrum was normalized by setting the flat low energy region to zero and the post-edge to unity (unit edge jump). The photon energy selected for the post edge was 560 eV, beyond the region of any absorption (peaks).

List of abbreviations

| Abbreviation | Description |
|--------------|-------------|
| TM           | Transition metal |
| LIB          | Li ion batteries |
| LCO          | LiCoO₂ |
HACR Hybrid anion- and cation-redox
OM Oxygen migration
GOM Global oxygen migration
sXAS Soft X-ray absorption spectroscopy
IPT Irreversible phase transformation
SCC Stress-corrosion cracking
G-LCO Gradient LiMn0.75Ni0.25O2 → LiCoO2 single crystal
GM-LCO Gradient-morph LCO single crystal
P-LCO Pristine LCO
AGG Abnormal grain growth
XRD X-ray diffraction
SEM Scanning electron microscopy
HRTEM High resolution transmission electron microscopy
STEM-HAADF Scanning transmission electron microscopy-high angle annular dark field
CI Coulombic inefficiency
FIB Focused ion beam
DEMDS Differential electrochemical mass spectroscopy
SOCs States of charges
XANES X-ray absorption near edge structure
SAED Selected area electron diffraction
$\bar{\Delta}_{Li}$ Li$^+$ diffusivity
$V_{O}^{**}$ Oxygen vacancy
FFT Fast Fourier transform

Author contributions

Z. Z. and J. L. conceived and designed the experiments. Z. Z. synthesized the materials, did the material characterization and performed the electrochemical measurements. D. Y. performed the aberration corrected STEM imaging, STEM-EDS mapping, EELS line scan, HRTEM imaging and in situ nano-XANES mapping. R. G. did the FIB sample preparation for STEM characterizations. Z. S. performed the lattice coherency calculation and analysis. I. W. measured the sXAS experiments. Z. Z., D. Y. and J. L. wrote the paper. All authors analyzed the data, discussed the results and commented on the manuscript.

Data availability

The data that support the plots in this paper and other findings of this study are available from the corresponding author upon reasonable request.

Conflicts of interest

The authors declare no competing interests.

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