

Imitating Architectural Mortise-Tenon Structure for Stable Ni-Rich Layered Cathodes

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Abstract: Ni-rich layered oxides are the most promising cathodes for Li-ion batteries, but chemomechanical failures during cycling and large first-cycle capacity loss hinder their applications in highenergy batteries. Herein, by introducing spinel-like mortise-tenon structures into the layered phase of LiNi_{0.8}Co_{0.1}Mn_{0.1}O₂, the adverse volume variations in cathode materials can be significantly suppressed. Meanwhile, this mortise-tenon structures play the role of the expressway for fast lithiumion transport, which is substantiated by experiments and calculations. Moreover, the particles with mortise-tenon structures usually terminate with the most stable (003) facet. The new cathode exhibits a discharge capacity of 215 mAh g⁻¹ at 0.1 C with an initial Coulombic efficiency of 97.5%, and capacity retention of 82.2% after 1200 cycles at 1 C. This work offers a viable lattice engineering to address the stability and low initial Coulombic efficiency of the Ni-rich layered oxides, and facilitates the implementation of Li-ion batteries with high-energy density and long durability.

1. Introduction

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Anisotropic volume variation is an intrinsic disadvantage of layered cathodic oxides because of the insertion and extraction of interlaminar ions during cycling.^[1-4] In detail, the lattice changes along axis a(b) are small, while those along axis c are large. This heterogeneous volume change (especially during the H2-H3 phase change process) in Ni-rich cathode materials leads to crystal structure fatigue (including distorted lattice and formation of cracks) that renders the active material ineffective.^[5-8] Thus, suppressing the volume change in Ni-rich materials is vital importance for durable batteries. Modified strategies such as doping or coating can partially conquer this issue,^[7-10] but the evolution of intrinsically anisotropic lattice parameters in layered oxides is inevitable. It has been recently proven that perovskite phase rivet and high-entropy oxides can significantly alleviate the volume change and

thereby stabilized the Ni-rich cathode.^[11, 12] Despite these advances, it remains challenging and daunting to effectively regulate the detrimental anisotropic volume change and simultaneously achieve a high capacity.

In addition to volume variations during battery operation, another universal drawback of Ni-rich layered materials is that they exhibit a large first-cycle specific capacity loss (12-30%).^[13-17] Such a large capacity loss leads to a low initial Coulombic efficiency, which was raised by the Nobel Laureate M. Stanley Whittingham recently.^[18] The main reason for the first-cycle issue can be attributed to the sluggish Li⁺ diffusion in the bulk materials, especially at the end of the discharge process.^[18-20] It is widely recognized that a high Ni-content or high voltage can enhance the specific capacity, but accompanied by a trade-off of stability.^[21-23] Thus, eliminating the large first-cycle capacity loss is an effective way to resolve the stability-capacity dilemma of Ni-contained cathodes. However, up to date, there is no effective strategy to avoid the large first cycle capacity loss. The above-mentioned volume variation and sluggish ion diffusion can all be traced to structural drawbacks; thus, structural engineering is expected to be the most effective and straightforward way to improve the electrochemical performance of the Ni-rich layered materials.

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Inspired by the robust mortise-tenon joints in traditional Chinese buildings, we design and fabricate layered LiNi_{0.8}C_{0.1}Mn_{0.1}O₂ with spinel-like mortise-tenon structures (NCM811-MT). Like logs in the architectural MT joints for a perfect connection, the layers of NCM811 are wedged into each other through the jagged ends at the spinel-like MT zones. The advantages of layered materials with such structure are summarized as follows. Firstly, the MT structure is strain-retardant to prevent the volume variations. Secondly, the MT structure can provide an expressway for fast lithium-ion transport. Thirdly, the MT structure in layered materials results in most of the particle surfaces being terminated by the stable (003) lattice planes. These superiorities are evidenced by extensive experimental and simulation data, and contribute to the significantly enhanced cycling stability and

initial Coulombic efficiency of NCM811-MT. This architectural MT design broadens the insights for stable crystal structures, and paves the way for developing advanced and durable batteries with high energy densities through structural engineering.

2. Results and discussion

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Figure 1a shows a typical MT structure in a traditional Chinese building, which can survive more than hundreds of years.^[24, 25] The crystallographic schematics of coupled MT structures and a plain layered structure are displayed in Figure 1b and 1c, respectively, which are also magnified in Figure S1 for better clarity. Both high resolution transmission electron microscopy (HRTEM) images and selected area electron diffraction (SAED) patterns reveal that the NCM811 exhibits typical layered crystalline structure in space group $R\overline{3}m$ (Figure S2). The aberration-corrected high-angle annular dark-field scanning transmission electron microscope (HAADF-STEM) images of a NCM811-MT particle are presented in Figure 1d-i. The MT joints separate the whole particle into four domains with the adjacent areas are [100] and [$\overline{1}$ 00] zone axis alternately. The adjacent (003) planes tilt about 109.5° or 70.5°, which are coherent with the spinel-like MT structure. Consequently, rational merging these domains together creates a spinel-like MT structure and a stable (003) surface-dominated particle (Figure S3-S5). For better understanding, the complex SAED pattern in Figure S3 was elaborated in Figure S6. While the average particle size of NCM811-MT reduced to 93 nm (155 nm for NCM811) after introducing of Li₂SO₄, the particles were more angular (Figure S7).



Figure 1. Illustration and characterizations of MT structures. (a) MT structure in a traditional Chinese building (Cherry Garden, Wuhan University). (b-c) Schematic of particles with coupled MT structures (b) and with a plain layered structure (c). Blue, pink and orange octahedra represent spinel MT structure in [110] axis, layered structure in [$\overline{100}$] axis and layered structure in [100] axis, respectively. (d) HAADF image of a NCM811-MT particle. (e-i) Atomically resolved HAADF STEM images at the five locations marked in (d). (j) XRD patterns of NCM811 and NCM811-MT.

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In addition to the local morphology investigation, the X-ray powder diffraction (XRD) was conducted to study the ensemble-averaged crystal structure of the particles (Figure 1j and Figure S8). All the evident peaks can be indexed to a typical α -NaFeO₂ layered structure in the space group $R\overline{3}m$. Rietveld refinements were conducted and the results were listed in Table S1. The lattice parameter *c* (*a*) is decreased (increased) from 14.191 (2.869) Å for NCM811 to 14.181 (2.872) Å for NCM811-MT. Meanwhile, the anti-site rate is similar for the two samples (3.8% for NCM811 and 4.6% for NCM811-MT), thus its impact on the lattice is insignificant.^[26, 27] The (003) d-spacing of the layered structure is

larger than the (111) d-spacing of the spinel structure, and this lattice change indicates the formation of spinel structures in NCM811-MT.^[28] The grain size of NCM811 and NCM811-MT is calculated to be 117.2 (1.7) nm and 24.3 (0.2) nm, respectively. Taking the particle size into account, the ratio of grain size/average particle size is 75% and 26% for NCM811 and NCM811-MT, which implies that the Accepted Articl NCM811-MT particle consists of multiple grains.

It is worth noting that trace Li₂SO₄ can still be detected in NCM811-MT (Figure S8c), which is also confirmed by the spectra of X-ray photoelectron spectroscopy (XPS, Figure S9), suggesting that Li₂SO₄ is very stable and does not react with the materials at high temperatures.^[29] The Li₂SO₄ was quantified to be 2.14 wt.% (molar ratio is 1.91%) in the NCM811-MT product by a carbon-sulfur analyzer. The spinel-like MT structures are formed during the process of 'welding' small particles into a larger one (Figure S10), and Li_2SO_4 is uniformly distributed in the material during synthesis (Figure S11). It is speculated that Li₂SO₄ can transport ions to build the spinel-like MT joint zone at high temperatures.^[30] To further investigate the possible formation mechanism of MT structures, we synthesized NCM811 with Na_2SO_4 or Li_3PO_4 as the additive. As results, spinel-like MT structures are observed in the final products (Figure S12). Indeed, spinel structures can form on the surface of lithium-rich layered oxides when polyanions are introduced during synthesis.^[31] Thus, polyanions such as SO₄²⁻ and PO₄³⁻ play a key role in the formation of spinel-like structures. The formation process of the spinel-like MT structures is summarized and presented in Figure S13.

The fine crystalline structures determine the ion storage properties of the layered materials. The initial charge-discharge curves of NCM811-MT and NCM811 at 0.1 C (1 C = 280 mA g^{-1}) are compared in **Figure 2**a. With a similar charge capacity of \sim 220 mAh g⁻¹, NCM811-MT and NCM811 presents a discharge capacity of 215 mAh g⁻¹ and 195 mAh g⁻¹, respectively. The initial Coulombic efficiency of NCM811-MT reaches 97.5%, which is much higher than that of NCM811 (88.4%). The dQ/dV curves in Figure 2b reveal that the detrimental H2-H3 phase transition was suppressed in NCM811-MT, which

is consistence with the cyclic voltammetry (CV) results (Figure S14). The pair of redox peaks at ~3.1 V may be attributed to the capacity contribution of spinel structure in NCM811-MT.^[12] The much lower initial capacity loss at different cut-off voltages (Figure S15) confirms that the long-standing inherent kinetics limitation (in the low voltage range during discharging) of Ni-rich layered material was overcome in NCM811-MT.^[14, 18, 19]



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Figure 2. Electrochemical performance of the NCM811 and NCM811-MT. (a) Initial charge-discharge profiles at the current of 0.1 C. (b) The corresponding differential capacity (dQ/dV) curves. (c) Rate performance. (d-e) Cycling performance.

NCM811-MT exhibits higher capacities than that of NCM811 at various rates (Figure 2c), implying that the MT structures can enable superior rate capabilities. The capacity retention after 1200 cycles with a voltage range of 2.8-4.3 V for NCM811-MT is 82.2%, which is considerably higher than that of NCM811 (26.5% of 1000 cycles) (Figure 2d). When the cut-off voltage was further increased to 4.5 V, the NCM811-MT still exhibits capacity retention of 81.5% after 800 cycles, which obviously outranges

NCM811 cell (Figure 2e). The charge/discharge and dQ/dV profiles during cycling reveal a considerably suppressed internal resistance drop^[32] and improved structural reversibility in NCM811-MT (Figure S16 and S17). It should be noted that the cycling stability of NCM811-MT (state-of-the-art) is much higher than recently reported single crystal^[33] and core-shell microstructured Ni-rich materials,^[34] and the details of performance comparison was shown in Table S2.

To reveal the function of the spinel-like MT structure in the cathode, we conducted in situ XRD to study the volume change and phase evolution during the charge/discharge process. The cells were tested at 2.8-4.3 V at a current of 0.1 C. In the XRD pattern of hexagonal layered structure NCM811, the position changes of (003) peak indicate the variation of c-value, while (110) represents the variation of *a* value.^[35] In general, during de-lithiation/lithiation of NCM811, the *c* value changes significantly, while the a value changes slightly. The contour maps of (003) and (110) reflections of the two samples are summarized in Figure 3, according to the recorded patterns in Figure S18. While the (003) peak initial mild shift to low angle followed by a sudden shift to high angle during charging for both of the samples, and vice versa during discharging (Figure 3a, 3b). The (003) peak variation of NCM811-MT was 0.32°, which is much lower than 0.57° for NCM811. The variation of (110) during charging for NCM811-MT and NCM811 was 1.34° and 1.53°, respectively (Figure 3c, 3d). The calculated c-axis lattice parameters variation is less pronounced in NCM811-MT compared to that of NCM811, with a maximum Δc of 1.91% (vs 2.84% for NCM811) (Figure 3e). The maximum Δa is similar but the difference tends to be slighter, which was 2.05% and 2.31% for NCM811-MT and NCM811, respectively (Figure 3f). As a result, the volume change was significantly limited in NCM811-MT. The less significant volume change of NCM811-MT relative to NCM811 can be associated with the spinellike MT structure, which constructs a robust connection between the unconsolidated layers along c and the rigid lattice along *a*.

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Figure 3. Structural evolution during the first charge-discharge process. Contour plot of (a) the (003) and (c) the (110) reflection of NCM811. Contour plot of (b) the (003) and (d) the (110) reflection of NCM811-MT. (e-f) Lattice parameter of c and a as functions of voltage.

We further investigated the structural and interfacial evolutions of the cathodic materials after long-term cycles using TEM. The emerging of microcracks and surface degradation were observed in the cycled NCM811 (**Figure 4**a-c and Figure S19), which is similar to previous results.^[36] As confirmed by the SEAD pattern (Figure 4b) and HRTEM images (Figure 4c), large area distortion was observed in the NCM811 after cycling. In sharp contrast, no structural distortion was observed in cycled NCM811-MT, and the surface degradation was quite limited (Figure 4d-f and Figure S20). More severe structure

degradations are expected when further increasing the cut-off voltage to 4.5 V,^[37] and obvious cracks and severe lattice distortions were observed in NCM811 (Figure 4g-i and Figure S21). No rock-salt phases formed in NCM811 at 4.5 V could be related to the rapidly growing overpotential (Figure S16, S17), which blocks lithium de-lithiation. The residual Li acts as an immense obstacle to the phase transition toward rock-salt.^[38] However, the NCM811-MT was still stable, without any distortions and cracks formed in the lattice (Figure 4j-l and Figure S22). Besides, the spinel-like MT and layered structures in NCM811-MT were well preserved and have not transformed into rock-salt phase, as proved by the HRTEM images (Figure 4I and Figure S22). It can be inferred that, the spinel-like MT structure presents a strain-retardant function, which is beneficial to enhance structural stability of NCM811-MT.^[11] The improved surface stability of NCM811-MT can be ascribed to its dominated (003) surface.^[39-41]



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Figure 4. Morphological and structural evolution of the cathode materials after 500 cycles with cutoff voltages of (a-f) 4.3 V and (g-l) 4.5 V at 1 C. (a-c) and (g-i) TEM images, corresponding SAED patterns and HRTEM images of NCM811. (d-f) and (j-l) TEM images, corresponding SAED patterns and HRTEM images of NCM811-MT. The selected areas for electron diffraction are marked by dash circles in a, d, g and j. The subscripts L and S indicate layered and spinel structures, respectively.

To further understand the diffusion kinetics of lithium ions, we conducted electrochemical impedance spectroscopy (EIS) of the two cells at different cycles (Figure S23). The equivalent circuit consists of solid electrolyte interface resistance (R_{sEl} , high-frequency range semicircle), charge transfer resistance (R_{ct} , medium-frequency range semicircle) and Warburg-type element reflecting the diffusion of Li⁺ in cathode materials (low-frequency tail).^[42] Considerable R_{ct} increment was observed upon cycling for NCM811, which increased from 14.8 Ω (1st) to 486.7 Ω (500th). In contrast, the R_{ct} of NCM811-MT increased slightly from 2.4 Ω to 44.4 Ω after 500 cycles (Figure S23c), implying fewer side reactions happen. Furthermore, the Li-ion diffusion coefficient (D_{Li+}) of NCM811-MT remains almost constant during cycling (~3×10⁻¹⁰ cm² s⁻¹), while such parameter in NCM811 is reduced by an order of magnitude after only 300 cycles (from ~3×10⁻¹⁰ to ~3×10⁻¹¹ cm² s⁻¹) (Figure S23d). Due to the spinel-like MT structure, NCM811, indicating a well-maintained structure, which is consistent with the TEM.

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The enhanced initial Coulombic efficiency and rate performance of the NCM811-MT, is considered to be closely related to the inborn spinel-like MT structure. To elaborate on how the MT structure impacts the Li ions transport, lithium-ion diffusions barriers at the grain boundary (GB), in the spinel domain and the layered domain have been calculated. The model of highly symmetrical layered structures connected by a spinel-like MT is displayed in **Figure 5**a. The lithium-ion diffusion energy barriers oblique to the GB and within the spinel structure is 0.18 eV and 0.19 eV (Figure 5c and 5d), respectively. Both of the values are much lower than that in the layered structure (0.62 eV, Figure 5e).

According to the Arrhenius relationship, the decrease of the energy barrier can significantly facilitate the ionic diffusion.^[43] Therefore, a spinel-like MT structure implanted in the layered structure is an expressway for lithium-ion transport in the bulk materials. To verify the calculation results, the lithium-ion diffusion coefficient based on the galvanostatic intermittent titration technique (GITT) of the cathodic materials with and without spinel-like MT was conducted (Figure 5b). It should be noted that, the ion diffusion coefficient values are slightly different in GITT and EIS due to the different measurement principles. The measured D_{Ur} of NCM811-MT is obviously higher than that of NCM811, especially with high lithium content (Figure 5b). Therefore, it can be concluded that due to the implantation of spinel-like MT structures, the intrinsic slow lithium kinetics plague of NCM811 at the high lithium contents is overcome.

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Figure 5. Fast lithium-ion diffusion in NCM811-MT. (a) Schematic of the spinel, layered domain and their boundaries. (b) Lithium-ion diffusion coefficient by GITT. (c-e) Lithium-ion diffusion energy barrier (c) at the GB, (d) within the spinel and (e) in the layered domain.

3. Conclusion

In summary, we propose and synthesize Ni-rich layered LiNi_{0.8}Co_{0.1}Mn_{0.1}O₂ materials with implanted spinel-like MT structures. The MT structures not only improve the mechanochemical stability by suppressing the intrinsic anisotropic volume change, but also provide an expressway for fast Li-ion transport. In addition, due to the formation of MT structures, the NCM particles are extensively

terminated with the most stable (003) planes, which can mitigate the cathode-electrolyte interfacial reactions. As a consequence, NCM811-MT exhibits quite excellent electrochemical properties with a capacity retention of 82.2% after 1200 cycles at 1 C and a high initial Coulombic efficiency of 97.5% at 0.1 C. This new lattice engineering strategy, introducing robust spinel-like MT structures into the layered oxides, may shed light on the design of Ni-rich layered cathodes with high structural stability and negligible first-cycle capacity loss.

4. Experimental section

*Synthesis of LiNi*_{0.8}*Co*_{0.1}*Mn*_{0.1}*O*₂ *with spinel-like mortise-tenon structure (NCM811-MT):* The materials were prepared by a sol-gel method. Stoichiometric amounts of lithium nitrate (Macklin, 99.9%), nickel nitrate hexahydrate (Aladdin, AR, 98%), cobalt nitrate hexahydrate (Aladdin, AR, 99%) and manganese nitrate (Macklin, AR, 50 wt.% in H₂O) were mixed with citric acid (Macklin, 99.5%) and lithium sulfate (Macklin, 99%) in 200 mL deionized water (total transition metals: 0.08 mol). The molar ratios of citric acid to the total metal ions and sulfate to transition metals were 1:4 and 2:100, respectively. After magnetic stirring overnight, the solution was dried at 80 °C to form gel, which was subsequently heated at 230 °C for 5 h and then ground well to form the precursor. The precursor (5 g each time) was calcined at 550 °C for 4 h and 750 °C for 15 h in oxygen. In comparison, LiNi_{0.8}Co_{0.1}Mn_{0.1}O₂ (NCM811) was also synthesized by the same method without adding lithium sulfate.

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Materials characterizations: The crystallographic structure of samples was identified by power X-ray diffraction (XRD, Bruker D8-Advance diffractometer) at a step mode with 0.02° per step and a sampling time of 1 s, using Cu-K α radiation at 40 kV and 80 mA. The *in situ* XRD tests were performed on the same instrument using an *in situ* XRD cell in the 2 ϑ angular range of 15-70° at a scan rate of 5° min⁻¹. FullProf program was employed for the XRD Rietveld refinements. The morphology and microstructure

of the samples were characterized by scanning electron microscope (SEM, ZEISS Supra 55 field emission SEM) and high-resolution field-emission transmission electron microscopy (FETEM, JEOL-3200FS, 300 kV). The cycled cell (fully discharged) was disassembled to get the cathode, which was washed by DMC for 5 times and immersed in NMP (sealed in a tube) in an argon filled glove box. To obtain the dispersive particles for TEM observation, the tube with a cathode inside was ultrasonically treated for 30 min. The air exposure time of the cycled particles is less than 10 min, before TEM observation. TEM coupled with energy dispersive X-ray spectrometry (EDS) was used to detect the elemental distribution. XPS with the excitation energy of 1486.68 eV (Al Kα source) was used to analysize the chemical state and composition. The atomic structure of the samples was acquired on an high-angle annular dark-field scanning transmission electron microscope (HAADF-STEM, JEM-AEM300F, Japan) operated at 300 kV and equipped with double spherical aberration correctors. The element analysis and etching profiles were acquired on an ESCALab 250Xi electron spectrometer. Carbon-sulfur analyzer (Horiba EMIA-8100, Contrinex, Germany) was used to determine the content of S in the final product. 521 4095, ja, Downloaded from https://onlinelibrary.wiley.com/doi/10.1002/adma.202301096 by University Town Of Shenzhen, Wiley Online Library on [15/05/2023]. See the Terms and Conditions (https://onlinelibrary.wiley.com/dems-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons License

Electrochemical measurements: Electrochemical properties were evaluated using coin cells (CR2032) assembled in an argon filled glove box. To fabricate the working electrodes, 80 wt% active materials, 10 wt% polyvinylidene fluoride (PVDF) and 10 wt% acetylene black were mixed in N-methyl pyrrolidone (NMP) solvent, followed by cast the slurry onto the aluminum foil and dried at 100 °C overnight in a vacuum oven. The mass loading of the active materials is about 2 mg cm⁻². Metal Li and Celgard 2316 were used as the anode and separator, respectively. The electrolyte was 1 M LiPF₆ dissolved in a 1:1:1 volume ratio of ethylene carbonate: ethyl methyl carbonate: dimethyl carbonate, with a 5% vinylene carbonate additive. Galvanostatic charge-discharge and galvanostatic intermittent titration technique (GITT, charging/discharging and waiting time is 10 min and 1 hour, respectively) measurements were carried out on a Neware battery test system (CT-4008T-5V10mA-164, Shenzhen,

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China) at room temperature (~25 °C). Cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS) were conducted on a Solartron electrochemical workstation. The cells were charged to 4.3 V before the EIS tests (100 kHz to 0.01 Hz with an AC oscillation amplitude of 5 mV for measurement).

First principles calculations: The Vienna *Ab initio* Simulation Package (VASP) with a plane-wave basis set^[44] and the projector-augmented wave pseudopotentials^[45] were used to perform density functional theory (DFT) calculations with a plane-wave cutoff energy of 520 eV. The spin-polarized generalized gradient approximation (GGA) with Perdew-Burke-Ernzerhof (PBE) function was used to treat the exchange-correlation effects^[46]. To correctly characterize the localized electronic states of the transition metal oxide materials, the GGA+U method was used.^[47] The U value for the Ni 3d state was chosen to be 6.4 eV.^[48] The Monkhorst-Pack *k*-point grid^[49] with a total number of at least 1000/(the number of atoms per cell) points for all directions, was used for the integration in the Brillouin zone. The convergence tolerance for residual force on each atom during structure relaxation was 0.02 eV/Å. The convergence criterion of force was 0.05 eV/Å.^[50]

Supporting Information

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

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Conflicts of interest

The authors declare no conflict of interest.

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A layered LiNi_{0.8}Co_{0.1}Mn_{0.1}O₂ material with implanted architectural mortise-tenon (MT) structures is developed. These MT structures not only improve the mechanochemical stability by suppressing the intrinsic anisotropic volume change but also provide an expressway for fast Li-ion transport. This layered LiNi_{0.8}C_{0.1}Mn_{0.1}O₂ exhibits a capacity retention of 82% over 1200 cycles (1C) and a high initial Coulombic efficiency of 97.5%.

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Imitating Architectural Mortise-Tenon Structure for Stable Ni-Rich Layered Cathodes

