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A study of room-temperature Li_xMn_{1.5}Ni_{0.5}O₄ solid solutions

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Understanding the kinetic implication of solid-solution vs. biphasic reaction pathways is critical for the development of advanced intercalation electrode materials. Yet this has been a long-standing challenge in materials science due to the elusive metastable nature of solid solution phases. The present study reports the synthesis, isolation, and characterization of room-temperature $\text{Li}_x \text{Mn}_{1.5} \text{Ni}_{0.5} \text{O}_4$ solid solutions. *In situ* XRD studies performed on pristine and chemically-delithiated, micron-sized single crystals reveal the thermal behavior of $\text{Li}_x \text{Mn}_{1.5} \text{Ni}_{0.5} \text{O}_4$ ($0 \le x \le 1$) cathode material consisting of three cubic phases: $\text{LiMn}_{1.5} \text{Ni}_{0.5} \text{O}_4$ (Phase I), $\text{Li}_{0.5} \text{Mn}_{1.5} \text{Ni}_{0.5} \text{O}_4$ (Phase II) and $\text{Mn}_{1.5} \text{Ni}_{0.5} \text{O}_4$ (Phase III). A phase diagram capturing the structural changes as functions of both temperature and Li content was established. The work not only demonstrates the possibility of synthesizing alternative electrode materials that are metastable in nature, but also enables in-depth evaluation on the physical, electrochemical and kinetic properties of transient intermediate phases and their role in battery electrode performance.

significant number of Li-ion batteries (LIBs) positive electrode materials operate in a first-order phase transition mechanism where the equilibrium electrochemical potential (E) of the electrode does not vary with the ratio of the phases present in the system¹⁻⁴. Two-phase transformation has traditionally been associated with severe limitations, including slow kinetics and poor stability of the materials. The large volume change between the involved phases leads to the accumulation of strain and mechanical degradation of the active particles, and the creation of Li ions transport barriers at the phase boundaries⁵. Various approaches, including downsizing the particles⁶ and creating anti-site defects in the lattice⁷, were proposed to reduce the miscibility gap and promote single-phase transformation in LiFePO₄. A recent report describes the use of *in situ* XRD with high temporal resolution to capture the appearance of transient solid solutions during high-rate cycling of LiFePO₄ nanoparticle electrodes⁸. The formation of Li_xFePO₄ (0 < x < 1) solid solutions was also accomplished by heating various mixtures of LiFePO₄ and FePO₄ to about $350^{\circ}C^{9,10}$. However, room-temperature single phases needed for detailed characterization were not attainable due to their high metastability.

LiMn_{1.5}Ni_{0.5}O₄ (LMNO) is a promising positive electrode material for high-energy density LIBs due to its 3D lithium-ion diffusion paths, a high operating voltage of 4.7 V, and a theoretical capacity of 147 mAh/g (energy density of 700 Wh/kg)¹¹⁻¹⁴. The spinel transforms through two topotactic two-phase reactions involving three cubic phases: LiMn_{1.5}Ni_{0.5}O₄ is first oxidized to $\Box_{0.5}$ Li_{0.5}Mn_{1.5}Ni_{0.5}O₄ and then \Box Mn_{1.5}Ni_{0.5}O₄ during charge, and vice versa during discharge^{15,16}. Here \Box denotes the vacant octahedral sites in the structure. The volume changes of Phase I/II and Phase II/III are approximately 3% and 3.3%, respectively. Several parameters, including particle surface facets¹⁷, particle size^{18–20}, substitution^{16,21,22}, and structural ordering^{11,23,24}, have been shown to have a large influence on the phase transformation pathway. First-principles calculation suggested that in perfectly disordered LMNO, single phase transformation throughout the entire Li composition range is possible at room temperature²⁵. Experimental studies, on the other hand, reported the presence of a large solid solution region followed by one or two two-phase transformations in disordered LMNO^{17,26}. For ordered LMNO, two two-phase transitions between three cubic phases were typically observed, with the Li content range of Phase I solid solution varying significantly among reports^{27,28}. Although single-phase transformation is considered kinetically advantageous over the two-phase process, owing to the observation that disordered LMNO with a larger solid solution region is often associated with better performance²⁹, the kinetic role of solid solution is largely unknown.

Encouraged by previous studies on temperature-dependent solid solution behavior in the LiFePO₄/FePO₄ system, we performed comprehensive thermal studies on a series of chemically delithiated $\text{Li}_x \text{Mn}_{1.5} \text{Ni}_{0.5} \text{O}_4$ (Li_xMNO, $0 \le x < 1$) prepared from structurally ordered LMNO crystals with exclusive (111) surface facets. The thermal treatment resulted in a reduction in the miscibility gap between the cubic phases I and II, and a series of single-phase solid solutions were attainable even at room temperature. Combining various techniques such as *in situ* temperature-controlled powder XRD (TXRD), Raman, Fourier transform infrared (FTIR) and X-ray



Figure 1 | (a) Room-temperature XRD patterns and (b) phase composition of the as-prepared $\text{Li}_x \text{Mn}_{1.5} \text{Ni}_{0.5} \text{O}_4$ crystal samples. Global Li content as indicated which has a standard deviation of less than 0.8%.

absorption (XAS) spectroscopies, a phase diagram capturing the changes as functions of both temperature and Li content was established, and the physical properties of Li_xMNO solid solutions were investigated.

Results

In situ XRD studies on the thermal behavior of Li_x MNO ($0 \le x \le$ 1). Uniform, micron-sized, and octahedron-shaped LiMn_{1.5}Ni_{0.5}O₄ crystals were synthesized by a molten salt method, as reported in our previous publication¹⁷. Extensive characterization by high-resolution transmission electron microscopy (HRTEM), selected area electron diffraction (SAED), FTIR, nuclear magnetic resonance (NMR) and electrochemical techniques revealed that the sample adopted near perfectly-ordered structure with a space group of P4₃32, and the particles are enclosed by 100% of (111) surface facets. A series of delithiated Li_x MNO, with x = 0.90, 0.82, 0.71, 0.51, 0.40, 0.25, 0.11,0.06 and 0, were prepared by chemical oxidation with varying amounts of a 0.1 M solution of nitronium tetrafluoroborate (NO_2BF_4) in acetonitrile. The lithium content x, solely controlled by the ratio between the spinel and the oxidizing agent, was determined by ICP analysis. The reactions resulted in a variety of mixtures among three cubic phases: LiMn_{1.5}Ni_{0.5}O₄ (Phase I), Li_{0.5}Ni_{0.5}Mn_{1.5}O₄ (Phase II) and Ni_{0.5}Mn_{1.5}O₄ (Phase III). The

XRD patterns and the weight fraction of the phases in the samples were obtained through full-pattern Rietveld refinements and the results are shown in Fig. 1a and 1b, respectively. Low level of lithium extraction (x > 0.71) led to the formation of cubic Phase II at the expense of Phase I. Phase III appeared on further Li removal, which rendered the coexistence of three cubic phases in the samples with an intermediate Li content of $0.25 < x \le 0.71$, although only about 3% of Phase III was present in Li_{0.71}MNO. At low Li content ($x \le 0.25$), the samples were composed of Phase II and III only. The lattice parameters of the Phases I, II and III were refined to be 8.1687(2), 8.0910(6) and 8.0005(3) Å, respectively. The smaller lattice dimensions in Phase II and III are directly related to the reduced Li content as well as a higher average Ni oxidation state in the spinel lattice.

It is worth noting that previous *in situ* XRD studies on the ordered LMNO octahedrons have revealed the existence of solid solution at lower state of charge, to an estimated Li content of $0.59 < x < 1^{17}$. In the literature, the reported *in situ* experiments so far showed significant variation in the lithium content range where solid solution behavior was observed. Aside from the differences in the step size used in the individual experiments, suble variation in sample crystal structure is believed to play a major role^{17,26,30}. Our effort to isolate the single phases from the electrochemically charged electrodes, how-





Figure 2 | Temperature-controlled XRD patterns of selected $\text{Li}_x \text{Mn}_{1.5} \text{Ni}_{0.5} \text{O}_4$ crystal samples: (a) x = 0.82, (b) x = 0.51, (c) x = 0.25, and (d) x = 0.



Figure 3 | XRD patterns of the Li_xMn_{1.5}Ni_{0.5}O₄ crystal samples collected at (a) 250°C and (b) after cooling to 25°C.

ever, was unsuccessful. After removing the same amount of Li in the x > 0.71 region, nearly identical phase mixtures were obtained from both electrochemically and chemically prepared Li_xMNO, as confirmed by ex situ XRD patterns collected on the cathodes recovered from partially charged cells (Supplementary Figure 1). This suggests that the solid solutions appearing during electrochemical charge and discharge are metastable transient intermediates that bear kinetic significance but thermodynamically unstable. Currently, no viable approach for the synthesis of phase-pure Li_xMNO intermediates has been reported.

a)

b

In situ temperature-controlled XRD studies were carried out to systematically investigate the thermal behavior of Li_x MNO ($0 \le x \le$ 1). Chemically delithiated samples were used due to its simplicity in preparing large quantities needed for this study as well as the purity in the absence of binder and carbon additives. Fig. 2 shows the TXRD patterns of selected Li_x MNO compositions (x = 0.82, 0.51, 0.25 and 0) collected during both heating and cooling. For comparison, the patterns of pristine LMNO are also shown in the supplemental information (Supplementary Figure 2), which displayed straightforward thermal expansion and contraction behavior resulting from heating and cooling, respectively. In all cases, no significant changes were detected below 150°C. Between 150 and 225°C, the peaks from the initial cubic phases broadened and started to merge in the samples with x = 0.82 (Fig. 2a) and 0.51 (Fig. 2b), and the complete transformation into a single cubic phase was observed at 250°C. The single phase formation temperature increases along with the decrease of Li content. The reduction in the miscibility gap at the high

temperature was attributed to the increase in Li ion diffusion from the lithium-rich to the lithium-poor phase, similar to the phenomena observed in the LiFePO₄/FePO₄ system. The lowered formation temperature of Li_xMNO is consistent with the fact that Li mobility is generally higher in LMNO than the LiFePO4 system. The cubic framework of the thermal-driven solid solution maintained during the cooling, with no phase separation observed even at room temperature. In the fully delithiated sample composed of Phase III only (x = 0, Fig. 2d), a new spinel-type phase with expanded lattice dimensions appeared above 200°C, suggesting that Phase III is susceptible to thermal-induced phase conversion. Here, the use of "a new spinel-type phase" is to denote the decomposition products, although it is possible that this phase consists of several structurally similar phases and/or compounds. At 250°C, the sample is composed of the decomposition products and Phase III in an approximately 1:1 weight ratio. Further conversion occurred during the early stage of cooling which led to much larger presence of the spinel-type phase in the cooled MNO at room temperature. For x = 0.25 consisting of Phase II and III in the as-prepared sample, three components, Phase II, the spinel-type phase and Phase III, were observed at 200°C (Fig. 2c). Upon further heating to 250°C, Phase II disappeared and the mixture was composed of the spinel-type phase and Phase III only. Fig. 3 compares the TXRD patterns of the Li_x MNO series (x =1, 0.90, 0.82, 0.71, 0.51, 0.40, 0.25, 0.11, 0.06 and 0) collect at 250°C (Fig. 3a) and after cooling to RT (Fig. 3b). At 250°C, single phases were obtained in samples with $x \ge 0.51$ which maintained phase pure after cooling to RT. Samples with Li content below 0.51, however,





Figure 4 | (a) Changes in the lattice parameters upon heating the $Li_xMn_{1.5}Ni_{0.5}O_4$ crystal samples ($0.51 \le x \le 1$), (b) the change in solid-solution lattice parameter on cooling, (c) changes in the lattice parameters upon heating the $Li_xMn_{1.5}Ni_{0.5}O_4$ crystal samples ($0 \le x \le 0.40$), and (d) refined domain size of the main phase (as indicated) in Li_xMNO at 250°C. Error range in the lattice parameters is less than ± 0.001 . Phase I: square; Phase II: circle; Phase III: triangle; Spinel-type phase: star.

were composed of mixtures at both 250°C and RT. The cooled samples with $x \le 0.25$ consisted mostly of the spinel-type phase which has significantly broader peaks compared to the initial spinel phases in the as-prepared samples.

The relationship between the lattice parameters of the phases present and the heating temperature is shown in Fig. 4, established from the fullpattern Rietveld refinements of the TXRD patterns. For samples with $0.51 \le x < 1$ that have minimal presence of Phase III, the cubic lattice parameters grew closer to each other when the temperature is raised above 150°C which eventually became a single lattice parameter around 250°C (Fig. 4a), confirming the formation of a single-phase solid solution in these samples. All solid solutions can be indexed using the same space group of $Fd\bar{3}m$, with the refinements produced similar lattice parameters in the range of 8.1873(9) to 8.1907(8) Å. This is consistent with the previous observation that in ordered LMNO, the lattice dimension is relatively insensitive to lithium content at high x^{17} . Upon cooling, the changes in lattice parameters followed a near linear relationship as a response to thermal contraction, reaching the range of 8.1687(8) to 8.1720(2) Å (Fig. 4b). The results confirm the absence of structural decomposition or phase demixing during the cooling process, further suggesting that, compared to the much discussed LixFePO4 solid solutions, thermal-driven Li_xMNO solid solution phases possess higher stability. For samples with $0 < x \le 0.40$ that have a predominant presence of Phase III (\geq 22 wt%), the merge of all three lattice parameters was not observed during heating (Fig. 4c). At 250°C, the lattice parameters of Phase II merged into the new spinel-type phase at about 8.1961(4) Å, but Phase III remained in all samples.

One possible compound represented by the new spinel-type phase is NiMn₂O₄-type spinel, which was previously reported as a common thermal decomposition product in Ni and Mn containing ternary oxides. Consistent with our results, Hu et al. have shown that fully delithiated and structurally ordered MNO was thermally unstable which released oxygen and transformed into a NiMn₂O₄-type phase at a temperature as low as $250^{\circ}C^{31}$. The transition occurs when the tetragonal Li is fully or partially replaced by Mn which renders a much large lattice parameter at room-temperature, up to 8.40 Å with full replacement. Our effort to obtain room-temperature lattice parameter of the newly formed spinel-type phase was not successful as cooling of the samples with $x \leq 0.25$ led to significant peak broadening (Supplementary Figure 3) and inaccurate refinement of the XRD patterns.

Fig. 4d compares the refined domain size of the main phase in Li_x MNO at 250°C, namely the solid solutions in the 0.40 < $x \le 1$ group and the new spinel-type phase in the $0 \le x < 0.40$ group. The domains size of ~50 nm in the former was reduced to half (~25 nm) in the latter case, consistent with major structural rearrangement and phase transformation associated with the thermal instability of Phase III in the Li-poor samples.



Figure 5 | (a) Raman and (b) FTIR spectra of the Li_xMNO samples collected at room temperature. Left: as-prepared and right: heat-treated.

Vibrational spectroscopy of Li_xMNO ($0 \le x \le 1$). Raman and FTIR spectroscopies probe the vibration modes of the bonds in a single molecule which are powerful in elucidating crystal symmetry of oxide samples. Unlike the powder XRD measurements, both techniques are able to detect small amounts of impurities as well as short-range ordering in amorphous phases that are typically transparent to XRD^{26,32}. Fig. 5a shows the Raman spectra of as-prepared L_xMNO samples (left) and their heat-treated counterparts that were recovered from the TXRD studies (right). In the as-prepared samples, typical Raman bands associated with cubic spinel lattice vibrations are displayed in the 400-700 cm⁻¹ region²⁴. The peaks at 404 and 496 cm⁻¹ on the pristine LMNO spectrum are commonly attributed to E_g and $F_{2g}^{(2)}$ Ni²⁺-O stretching modes. With decreasing x, the disappearance of the peaks at 404 cm^{-1} and the appearance of the peaks around 485 and 545 cm⁻¹ indicate oxidation of Ni^{2+} to Ni^{4+33} . The peaks between 570 and 650 cm⁻¹ are assigned as A_{1g} and $F_{2g}^{(1)}$ Mn-O modes in the MnO₆ octahedron, with their

position and relative intensity ratio closely related to Mn-O bond length, MnO₆ distortion, and Mn oxidation state. It has been shown that an increase in Mn³⁺ content, often associated with a loss of translational symmetry due to the Jahn teller effect, leads to the broadening of the bands from both A_{1g} and $F_{2g}^{(1)}$ modes. This broadening may be further enhanced by an increase in electrical conductivity induced by charge transfer via electron hopping between Mn³⁺ and Mn⁴⁺. On the pristine LMNO spectrum, the well-resolved split of the $F_{2g}^{(1)}$ bands at 593 and 612 cm⁻¹, which mainly originate from the vibration modes of the Mn4+-O bonds, is consistent with the ordering of the transition-metal cations (Ni²⁺ and Mn⁴⁺) where supplementary Raman modes are expected to be active. The A_{1g} band at 635 cm⁻¹ is related to both Mn⁴⁺-O and Mn³⁺-O vibrations. The intensity ratio of 593/635, therefore, is traditionally used as an indicator for Mn4+ concentration in the lattice as well as the average Mn oxidation state in the sample³⁴. The dominance of the peak at 593 cm⁻¹ and its continuous shift towards lower frequency



Figure 6 | XANES spectra of Ni K-edge: (a) as-prepared, (b) heat-treated, and (c) the relationship between the Ni energy edge at the first inflection point (E_0) and Li content in the samples.

along with decreasing lithium content can be attributed to a combination of slight increase in Mn^{4+}/Mn^{3+} ratio and a decrease in Li ion concentration at 8a sites. As determined by the electrochemical studies in our previous report, the estimated Mn^{3+} concentration in the pristine octahedron sample is close to $1.5\%^{17}$.

No significant differences were observed when comparing the spectra of the as-prepared and heat-treated samples with $0.71 \le x \le 1$, all of which resembled that of the LMNO pristine sample. This further confirms the retention of spinel local symmetry in the thermally-induced solid solutions and phase purity of the samples. On the other hand, substantial peak broadening and reduced scattering intensities were observed in the heated-treated samples with $0 \le x \le 0.51$, suggesting thermal induced structural changes, formation of secondary phases with reduced domain size, and possibly, enhanced electrical conductivity³⁵. The dominance of the broad bands at 486 and 588 cm⁻¹ on the spectra suggests enhanced distortion of MnO₆ octahedron and the presence of a new spinel phase with reduced symmetry, such as λ -MnO₂ or NiMn₂O₄-type spinel^{36,37}.

The corresponding FTIR spectra of the Li_xMNO samples before and after heat treatment are shown in Fig. 5b. The presence of the eight well-defined bands at 430, 468, 480, 501, 557, 594, 621, and 650 cm⁻¹ is consistent with the ordered structure in the pristine LMNO sample. Similar to the XRD and Raman results, the cubic spinel structure in the pristine sample was retained and no significant change in symmetry resulted from either delithiation or heat treatment of the samples with $0.71 \le x \le 1$. In the thermally-treated samples, peak broadening and changes in intensity ratios between the bands were first observed in $L_{0.51}$ MNO, which exacerbated as *x* decreased. Since Li_{0.51}MNO is the first sample in the series where the presence of Phase III becomes significant enough (13 wt% in x =0.51 as opposed to <3 wt% in x = 0.71, the previous sample in the series) to enable the detection of its decomposition products by the vibrational techniques, the results provide further evidence that the formation of the new spinel-type phase in Li_xMNO ($0 \le x \le 0.51$) is directly related to the thermal decomposition of Phase III. The higher sensitivity of the vibrational spectroscopy in detecting phases in small amounts and with short-range ordering complements the results from the bulk XRD study where the technique limits the detection of impurities in x = 0.51.

X-ray absorption spectroscopy of Li_x MNO ($0 \le x \le 1$). 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 bulk material^{38,39}. The technique also probes the local and electronic structures near the absorbing atoms and it has the ability to reveal element-specific structural changes in the samples. Fig. 6 shows the Ni K-edge Xray absorption near edge structure (XANES) spectra from the asprepared (Fig. 6a) and heat-treated (Fig. 6b) LixMNO series, respectively. In the as-prepared samples, chemical oxidation led to a monotonous edge-shift to higher energy, consistent with a continuous increase in average oxidation state of nickel as it transformed from the divalent state in LiMn_{1.5}Ni_{0.5}O₄ to tetravalent state in Mn_{1.5}Ni_{0.5}O₄. Two isosbestic points, indicative of a three-component system, were observed at 8353.8 and 8355 eV on the XANES spectra, which is consistent with the presence of Phase I, II and III in the as-prepared samples. The results are also in good agreement with the in situ XANES data recently reported by Arai et al.40 For comparison, the room-temperature spectra of the LixMNO samples subjected to the heat treatment are shown in Fig. 6b. Negligible changes were observed in samples with high Li content of $0.71 \le x$ \leq 1. The samples with lower lithium content of $0 \leq x \leq 0.51$, on the other hand, experienced substantial edge shift towards lower energy, with the absorption edges nearly overlapping with those of lithium-



Figure 7 | XANES spectra of Mn K-edge: (a) as-prepared, (b) heat-treated, and (c) the relationship between the Ni energy edge at the first inflection point (E_0) and Li content in the samples.

rich samples (0.71 $\leq x \leq$ 1). Fig. 6c compares the edge position, defined by Photoelectron Energy Origin (E₀) that is a commonly used parameter for extracting edge energy level in XANES spectra^{31,39}, as a function of lithium content in the samples before and after thermal treatment. Clearly, heating led to significant reduction in Ni oxidation state in the samples with x < 0.71, suggesting that the formation of the new spinel-type phase is related to Ni reduction. Fig. 7 shows the Mn *K*-edge XANES spectra from the as-prepared and heat-treated samples. Only slight energy increase resulted from chemical oxidation (Fig. 7a), corresponding to the low concentration of Mn³⁺ in the pristine sample (1.5%) and limited Mn³⁺/Mn⁴⁺ redox activity. Unlike the

Ni case, heating led to negligible edge shift in the entire series (Fig. 7b), suggesting the absence of significant Mn redox activities during thermal decomposition of Phase III. The edge position as a function of lithium content before and after the heat treatment is compared in Fig. 7c.

Further changes were observed in the intensity of Mn pre-edge absorption peaks arising from the dipole forbidden $1s \rightarrow 3d$ electronic transitions, which are typically week in the transition metals. In the presence of 3d and 4p orbital hybridization, often resulting from structural distortion in local symmetry or non-centrosymmetric environment between the metal and oxygen coordination, the transitions are made partially allowed and the peaks become



Figure 8 | XANES spectra of Mn pre-edge: (a) $0.71 \le x \le 1$ and (b) $0 \le x \le 0.25$. Dotted lines: as-prepared and solid lines: heated-treated samples.



Figure 9 | Phase diagram of $\rm Li_xMn_{1.5}Ni_{0.5}O_4$ on heating. \times indicates the temperature where significant structural changes were observed in the $\rm Li_xMn_{1.5}Ni_{0.5}O_4$ TXRD patterns. The phase compositions in each region as labeled.

much stronger⁴¹⁻⁴³. Also, in an event where a transition metal occupies the tetrahedral instead of the octahedral site, much stronger preedge peaks are observed owing to the lack of inversion center in tetrahedral geometry. As shown in Fig. 8, heat-treatment led to negligible changes in samples with $0.71 \le x \le 1$ (Fig. 8a). The treated samples with low lithium contents, particularly those with $0 \le x \le 0.25$, on the other hand, have noticeably increased Mn pre-edge absorption peaks (Fig. 8b). This suggests that although no significant Mn redox activities are involved in the thermal decomposition of Phase III, some Mn cations may migrate from the octahedral to the tetrahedral sites. This is consistent with the possible formation of NiMn₂O₄-type phase through partial replacement of the tetragonal Li by Mn. Further analysis is necessary in order to determine the chemical nature of the thermal decomposition products.

Phase diagram of Li_{*x*}**MNO** ($0 \le x \le 1$). On the basis of the results obtained from TXRD, Raman, FTIR and X-ray absorption spectroscopies, the phase diagram of LixMNO on heating was constructed (Fig. 9). Because of the nature of chemical delithiation, the Li content step size in this study is relatively coarse which limits the resolution of our phase diagram. Overall, the phase behavior of Li_xMNO as a function of Li content can be divided into three regions at room temperature. In the first region where Li content is between 0.71 and 1, the initial cubic phases I and II merge at high temperature and phase-pure solid solutions form when the temperature is raised above 200°C. The formation temperature increases along with the decrease of Li content. This is the only domain where single phases are attainable in the Li_xMNO system at both elevated and room temperatures. In the second domain where samples are composed of Phase I, II and III (0.25 < x < 0.71), the merge of Phase I and II led to the formation of solid solution but Phase III remains and a new spinel-type impurity phase also appears at high temperature. Further increasing temperature to above 265°C led to the complete conversion of Phase III to the spinel-type phase (Supplementary Figure 4). For $0 < x \le 0.25$, no solid solution formation was observed during the heating process. Phase III decomposed to the spinel-type phase and the final treated samples are composed of both phases.

While Phase I and II in LMNO system are able to merge into a single solid solution at high temperature, Phase III has very low solubility in the lithium containing spinel Phase I and/or II. Phase III also has the lowest thermal stability among the three cubic phases, which decomposes into a spinel-type phase upon heating at a temperature as low as 150°C. The decomposition temperature is largely influenced by the presence of Phase I, which decreases along with the reduction in Phase I content in the sample. It appears that in the absence of Phase I, Phase II undergoes thermal decomposition as well but at a temperature much higher than that of Phase III.

Discussion

The study sheds light on the thermal behavior of the Li_xMNO system and reports the isolation of phase pure room-temperature Li_xMNO solid solutions for the first time. A series of delithiated Li_xMNO samples were prepared by chemical delithiation. While the samples with a Li content between 0.71 and 1 were composed of Phase I and II only, the ones with a low Li content between 0 and 0.25 consisted of Phase II and III, with the latter being the dominant phase in the mixture. The samples with an intermediate Li content between 0.25 and 0.71 were mixtures of all three cubic phases. TXRD studies successfully revealed structural evolution of LixMNO as functions of both temperature and Li content. At high temperature, single-phase Li_xMNO solid solutions formed in Li-rich samples as the miscibility gap between Phase I and II was reduced. The intermediates remained phase pure after cooling to room temperature, and their structural similarity to the original spinel Phase I was further confirmed by the vibrational and XAS studies. A new spinel-type impurity phase was detected in samples with x < 0.71, which was attributed to the thermal decomposition products of Phase III. It was speculated that the process releases oxygen from the spinel lattice and triggers nickel reduction to a lower valence state as a compensation mechanism. The chemical nature of the new spinel-type phase, however, is unclear at this point, and it is possible that it represents several structurally similar phases and/or compounds. Further characterization is currently underway and the results will be reported in a future publication.

The study reveals that the solid solution behavior in LMNO system bears a great deal of similarity to that of LiFePO₄, such as: 1) they are metastable in nature, 2) they can appear as transient intermediates during electrochemical charge and/or discharge, and 3) their formation is promoted by temperature. However, Li_xMNO solid solutions appear to possess higher stability compared to Li_xFePO₄, despite metastable in nature. They are easily detectable by in situ techniques such as XRD, and the thermally-driven single phases can be preserved at room temperature. Considering the higher Li mobility in LMNO, the reason for the enhanced stability is unclear and it requires further investigation. Our findings provide pathways for detailed evaluation on the properties of solid solution intermediates, which may further lead to the understanding of their role in the performance of intercalation electrode materials. This study also sets an example for the synthesis of difficult metastable phases that may be used as new electrode materials.

Methods

Unless otherwise specified, all chemicals were obtained from Aldrich with a purity of 97% or higher. Well-formed octahedral-shaped LiMn1.5Ni0.5O4 crystals were prepared according to the procedure described in our previous report¹⁷. Various levels of chemical delithiation were achieved by reacting the pristine powder with 0.1 M nitronium tetrafluoroborate in acetonitrile solution in an Argon filled glove box (O2 < 1 ppm and H_2O < 1 ppm) at room temperature. The resulting reaction mixtures were filtered, thoroughly washed with acetonitrile, and then dried overnight in a vacuum oven. Chemical composition of the samples was determined by an inductively coupled plasma optical emission spectrometer (ICP-OES, Perkin-Elmer Optima 5400). X-ray diffraction patterns were collected using a Panalytical X'Pert Pro diffractometer with monochromatized Cu K_{α} radiation. The scans were collected between 32 and 80° (2 θ) at a rate of 0.0001°/s and a step size of 0.022°. Lattice parameters and phase ratios were determined by full-pattern Rietveld refinements using Riqas software (Materials Data, Inc.). Temperature-controlled XRD studies were carried out in the same diffractometer equipped with an Anton Parr HTK 1200 hot stage. The samples were heated in air at a rate of 5°C/min, and the XRD patterns were recorded at a temperature step size of 25°C, with each temperature holding for 5 min before data collection. The same procedure was used for the cooling process. Lattice parameters and phase ratios were determined by full-pattern Rietveld refinements using a Pearson VII function in the RIQAS software (Materials Data,

Inc.). The structural model used is $Fd\bar{3}m$ space group with the following Wyckoff position: Li (8a), Ni/Mn (16d) and O (32e). The refined scale factor was used to obtain the phase fractions. The goodness of the fitting is shown in Table 1 in the supporting information. FTIR measurements were performed on KBr pellets using a Nicolet 6700 spectrometer in transmission mode with a spectral resolution of 4 cm⁻¹. Raman spectra of the powder samples were recorded on a "Labram" Raman confocal microscope system (ISA Groupe Horiba) in the confocal backscattering configuration with a 488 nm Argon ion laser (Coherent Inc. Innova 70), a plan olympus $10 \times$ magnification, a 0.25 numerical aperture objective lense, and a 22 focal length. Hard XAS data (Mn and Ni K-edges) were collected in transmission mode using a (220) monochromator at SSRL beamline 4-1. Fine powders of as-prepared and heat-treated Li_xMNO samples were sandwiched between a kapton tap for data collection. Higher harmonics in the X-ray beam were reduced by detuning the Si (220) monochromator by 35% at the Ni edge and 50% at the Mn edge. Energy calibration was accomplished by using the first inflection points in the spectra of Ni and Mn metal foil references, which were 8333 and 6539 eV, respectively. XANES data were analyzed by Sam's Interface for XAS Package or SIXPACK software, with the Photoelectron Energy Origin E₀ determined by the first inflection point of the absorption edge jump.

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Author contributions

S.K. prepared the samples, performed the experiemental studies, and prepared the figures under the supervision of G.C. S.K. and G.C. wrote the main manuscript text. A.J. and R.K. performed Raman characterization of the samples and contributed to writing the manuscript. All authors reviewed the manuscript.

Additional information

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