Entropy and crystal-facet modulation of P2-type layered cathodes for long-lasting sodium-based batteries

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P2-type sodium manganese-rich layered oxides are promising cathode candidates for sodium-based batteries because of their appealing cost-effective and capacity features. However, the structural distortion and cationic rearrangement induced by irreversible phase transition and anionic redox reaction at high cell voltage (i.e., >4.0 V) cause sluggish Na-ion kinetics and severe capacity decay. To circumvent these issues, here, we report a strategy to develop P2-type layered cathodes via configurational entropy and ion-diffusion structural tuning. In situ synchrotron X-ray diffraction combined with electrochemical kinetic tests and microstructural characterizations reveal that the entropy-tuned Na0.62Mn0.67Ni0.23-Cu0.05Mg0.07Ti0.01O2 (CuMgTi-571) cathode possesses more {010} active facet, improved structural and thermal stability and faster anionic redox kinetics compared to Na0.62Mn0.67Ni0.37O2. When tested in combination with a Na metal anode and a non-aqueous NaClO4-based electrolyte solution in coin cell configuration, the CuMgTi-571-based positive electrode enables an 87% capacity retention after 500 cycles at 120 mA g\textsuperscript{-1} and about 75% capacity retention after 2000 cycles at 1.2 A g\textsuperscript{-1}.

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a-ion batteries (NIBs) have attracted significant interest as a low-cost alternative to Li-ion systems for large-scale energy storage applications due to the natural abundance of Na resources and the similar chemical properties of Na and Li\(^{1-3}\). The foremost challenge in advancing NIBs lies in developing high-performance and low-cost electrode materials. Among the available cathode materials, P2-type layered oxides, especially sodium manganese-rich oxides (Na\(_{x}\)Mn\(_2\)O\(_4\)) have been deemed as one of the most fascinating cathode candidates due to their high specific capacities and the natural abundance of Mn\(^{4-5}\). Moreover, oxygen anionic redox activity has been confirmed in these cathodes when charging beyond 4.0 V (vs. Na/Na\(^+\)), which can increase the reversible capacity to even over 200 mAh g\(^{-1}\) at low charge/discharge specific current (e.g., 10 mA g\(^{-1}\))\(^{6,7,8}\). However, compared with Li\(^+\), the insertion/extraction of Na\(^+\) in the reported P2 cathodes generally shows poor reversibility and sluggish kinetics owing to its larger ionic radius (1.02 Å for Na\(^+\) vs. 0.76 Å for Li\(^+\))\(^{9}\), Supplementary Table 1). Moreover, shifting the large and heavy Na\(^+\) leads to the gliding of MO\(_2\) layers and the formation of the irreversible O\(_2\) or OP\(_4\) phase accompanied with the pronounced volume variation during cycling, leading to fast capacity decay and short cycle life\(^{5,8,9}\). On the other hand, the irreversible lattice oxygen due to the structural instability results in detrimental cationic migration and oxygen network distortion, leading to voltage decay and poor kinetics\(^{10,11}\).

Over the past decades, extensive research has been conducted to stabilize anionic redox in the Mn-based P2-type layered cathodes\(^{12-14}\). However, the reported cycle stability by surface coating or alloys doping, especially under fast-charge conditions, remain far less satisfied\(^{15-19}\). Instead, rather less attention has been paid to improving ion transport kinetics. Ion transport kinetics within electrode materials play a pivotal role in the battery performance, which depends highly on the host structure and ion-diffusion tunnels. A stable host structure can avoid the collapse of lattice and diffusion channels, which is the precondition for reversible ion insertion/extraction. The construction of more diffusion-favorable tunnels is beneficial for the rapid migration of large amounts of ions. Thus, the key to addressing the slow ion kinetics lies in constructing a stable host structure with more migration tunnels.

Recently, high-entropy oxides have shown significantly improved structural stability during charge/discharge\(^{20,21}\). Hu et al. has revealed that high-entropy O3-type Na\(_{0.62}Mn_{0.67}Cu_{0.05}Mg_{0.09-2y}Ti_{0.02}O_2\) cathode could enable reversible O3-P3 phase transition and cationic redox within a narrower voltage range of 2.0–3.9 V\(^{20}\). In addition to structural stability, [010] planes with an open structure and ion-migration tunnels are favorable active planes for ion transport of layered cathode materials. Our previous work has demonstrated that increasing [010] active planes could significantly improve the rate performance of lithium layered oxide cathodes\(^{22,23}\). P2 cathodes with increased exposure of [010] facets have also been reported, which however were evaluated only within a low-voltage region (2.0–4.0 V).

High-voltage operation (>4.2 V) is an effective way to increase the specific capacity of battery materials. However, it is well-known in the battery community that high-voltage operation often leads to undesired structural degradation of the cathode. To date, despite numerous attempts through various strategies, the stabilization of layered cathodes at high voltage remains a formidable challenge. To the best of our knowledge, the correlation between entropy and anionic redox kinetics/reversibility of layered cathodes at high voltage remains elusive.

Based on the above discussions, simultaneously increasing the entropy and active facets would be a powerful gateway to enable P2-type layered cathodes with ultrafast charging capability and ultralong cycle life (Fig. 1), which however has not been reported so far. In this work, we have designed and synthesized a series of multi-element P2-type Na\(_{0.62}Mn_{0.67}Ni_{0.23}Cu_{0.05}Mg_{0.09-2y}TiO_2\) (CuMgTi-571, y = 0.01; CuMgTi-552, y = 0.02; CuMgTi-533, y = 0.03) layered cathodes with different configurational entropy and [010] active facets. Through systematically comparing their electrochemical performance and investigating their phase transition and Na\(^+\) diffusion coefficient during charge/discharge, we have identified the correlation between entropy and active crystal facet, cationic/anionic kinetics as well as an electrochemical performance of P2-type cathodes. In comparison with Na\(_{0.62}Mn_{0.67}Ni_{0.37}O_2\), the optimal Na\(_{0.62}Mn_{0.67}Ni_{0.23}Cu_{0.05}Mg_{0.07}Ti_{0.01}O_2\) cathode with higher entropy and increased active [010] facet not only effectively suppressed destructive structural variation and extends cycle life, but also significantly promoted the anionic redox and fast charging capability during high-voltage cycling (i.e., 2.0–4.3 V cell voltage range).

**Results**

**Structural and morphological characterization.** The crystalline structures and phase purities of the entropy-tuned Na\(_{0.62}Mn_{0.67}Ni_{0.23}Cu_{0.05}Mg_{0.09-2y}TiO_2\) and the non-entropy-tuned NaNiO\(_2\) were characterized by X-ray diffraction (XRD) together with the Rietveld refinement. As shown in Fig. 2, all the characteristic peaks from each sample are sharp, clear, and well fitted with the typical hexagonal P2\(_1\)-structure with P6\(_3\)/mmc space group, indicating the high purity and good crystallinity of the samples\(^{24,25}\). The pure P2 phase of Na\(_{0.62}Mn_{0.67}Ni_{0.23}Cu_{0.05}Mg_{0.09-2y}TiO_2\) suggests uniform incorporation of Cu\(^{2+}\), Mg\(^{2+}\), and Ti\(^{4+}\) into the lattice of the materials. Compared to Mn\(^{4+}\) (0.53 Å) and Ni\(^{2+}\) (0.69 Å), the introduction of larger-size cations such as Cu\(^{2+}\) (0.73 Å) and Mg\(^{2+}\) (0.72 Å) may lead to a debate on the size variation of interlayer spacing. However, previous literature has shown that interlayer spacing doesn’t always increase with an increased substitution amount of large-sized cations\(^{16}\). Meanwhile, a high entropy sample Na\(_{0.62}Mn_{0.67}Ni_{0.23}Co_{0.03}Al_{0.03}Ti_{0.02}O_2\) consisting of small-sized cations (Co\(^{3+}\): 0.61 Å, Al\(^{3+}\): 0.53 Å, Ti\(^{4+}\): 0.60 Å) was synthesized and showed that introduction of cations with smaller ionic radii into...
P2-typed structure can lead to increased interlayer spacing (Supplementary Fig. 1).

Therefore, to obtain the lattice parameters and interlayer spacing of Na$_{0.62}$Mn$_{0.67}$Ni$_{0.23}$Cu$_{0.05}$Mg$_{0.09}$-2yTi$_{y}$O$_{2}$ and NaMNO$_{2}$ samples, the synthesized materials were refined by the Rietveld method as displayed in Fig. 2a–d, respectively. According to the refined crystallographic data presented in Fig. 2e and Supplementary Tables 2–5, the Mn$^{4+}$, Ni$^{2+}$, Cu$^{2+}$, Mg$^{2+}$, and Ti$^{4+}$ ions are located in the octahedral 2a Wyckoff sites. Na$^{+}$ ions are located in two prismatic sites (2b and 2d, respectively). The interlayer spacings (d_{o-o}) of CuMgTi-533, CuMgTi-552, CuMgTi-571, and NaMNO$_{2}$ were 4.09, 4.03, 4.09, and 3.98 Å, respectively. The maximum changes of interlayer distance (Δd_{o-o}) are 0.11 Å. Compared to previous report work by increasing the Na$^{+}$ layer distance from 3.6 to 5.8 Å to improve the electrochemical performance$^{26}$, the interlayer distance change here is much smaller.

The morphologies and detailed crystal structures of the entropy-tuned Na$_{0.62}$Mn$_{0.67}$Ni$_{0.23}$Cu$_{0.05}$Mg$_{0.09}$-2yTi$_{y}$O$_{2}$ and the non-entropy-tuned NaMNO$_{2}$ were characterized by scanning electron microscopy (SEM) and transmission electron microscopy (TEM). Low-magnification SEM images (Supplementary Fig. 2) showed that all the as-prepared samples are composed of micrometer-sized particles with predominantly 1–2 μm in diameter. High-magnification SEM images (Fig. 3a–d) revealed that these particles have rather smooth surfaces over the whole particle. TEM images from the individual structure (Supplementary Fig. 3a–d) further confirmed the smooth and clean surfaces of these micrometric particles. The atomic compositions of Na$_{0.62}$Mn$_{0.67}$Ni$_{0.23}$Cu$_{0.05}$Mg$_{0.09}$-2yTi$_{y}$O$_{2}$ and NaMNO$_{2}$ samples were confirmed by inductively coupled plasma-mass spectrometry (ICP-MS), as displayed in Supplementary Table 6. The ICP results revealed that the atomic ratios of metals in all samples are highly consistent with the designated atomic ratio. Moreover, as shown in Fig. 3e and Supplementary Fig. 4, the elemental mappings by high angle annular dark field scanning transmission electron microscopy (HAADF-STEM) confirmed the existence and uniform distribution of Na, Mn, Ni, Cu, Mg, and Ti in the Na$_{0.62}$Mn$_{0.67}$Ni$_{0.23}$Cu$_{0.05}$Mg$_{0.09}$-2yTi$_{y}$O$_{2}$ and Na, Mn, Ni in the NaMNO$_{2}$ samples.

The selected area electron diffraction (SAED) patterns of CuMgTi-571 (Fig. 3f) exhibited hexagonal spot patterns, which is an indication of the single-crystalline feature and hexagonal crystalline structure of the microparticles. Based on the hexagonal phase system of P2-type layered oxides, it can be inferred that the front facets of microparticles are {001} planes and the side facets are {010} planes (Fig. 3g). The atomic arrangement of {010} planes presents an open structure, which facilitates Na$^{+}$ transport.

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**Fig. 2 Interlayer spacing of synthesized cathodes.** XRD patterns and Rietveld refinement plots of a CuMgTi-533, b CuMgTi-552, c CuMgTi-571, and d NaMNO$_{2}$ samples. e The detailed atomic model of interlayer spacing (d_{o-o}) in P2-type layered structure is presented on the left. The values of lattice parameter c and interlayer spacing for four samples are presented on the right.
The percentage of {010} facets and conformational entropy variation may reasonably alter the surface energy of crystal planes, influencing the growth rate of different facets. To the best of our knowledge, the effect of conformational entropy variation may reasonably alter the surface energy of crystal planes, influencing the growth rate of different facets. Therefore, high conformational entropy is the impetus for the growth of a high percentage {010} active facets in this work, while the temperature is the driving force for the growth of [010] active facets in the case of O3 lithium compounds (LiNi1/3Co1/3Mn1/3O2 and Li(Li0.17Ni0.23Mn0.60)O2) with low entropy. These results indicate that (conformational) entropy variation may reasonably alter the surface energy of crystal planes influencing the growth rate of different facets. To the best of our knowledge, the effect of configurational entropy on the surface structure of electrode materials has been barely reported and requires more comprehensive study.

\[
\gamma(T) = Y_0 - (T/A)\left(S_{\text{vib}} + S_{\text{config}}\right)
\]

where \(\gamma(T)\) and \(Y_0\) are surface energy at any temperature and 0 K, respectively. \(T\) is temperature, \(A\) is Helmholtz free energy, \(S_{\text{vib}}\) and \(S_{\text{config}}\) are the system’s vibrational and configurational entropy, respectively. It can be found that the surface energy of a system with simple composition and low entropy is mainly influenced by temperature, while the surface energy of polycomponent system with high entropy is mainly influenced by configurational entropy \(S_{\text{config}}\). The P2-type layered cathode materials in our current work contain six cations, which are high-entropy systems. The surface energy of these P2-type entropy-tuned materials is mainly affected by configurational entropy \(S_{\text{config}}\) at a given temperature. Thus, configurational entropy variation may reasonably alter the surface energy of the crystal plane, influencing the growth rate of different facets and promoting the formation of the layered cathode with more {010} facets. Therefore, high configurational entropy is the impetus for the growth of a high percentage {010} active facets in this work, while the temperature is the driving force for the growth of {010} active facets in the case of O3 lithium compounds (LiNi1/3Co1/3Mn1/3O2 and Li(Li0.17Ni0.23Mn0.60)O2) with low entropy. These results indicate that (conformational) entropy variation may reasonably alter the surface energy of crystal planes influencing the growth rate of different facets. To the best of our knowledge, the effect of configurational entropy on the surface structure of electrode materials has been barely reported and requires more comprehensive study.
Electrochemical performances. To clearly unravel the effect of entropy and crystal-facet modulation on the structural stability and performance of P2-structured cathode material, a series of electrochemical characterizations were performed. Cyclic voltammetry (CV) measurements were conducted to compare the redox behaviors of the entropy-tuned Na0.62Mn0.67Ni0.23Cu0.05Mg0.09-2yTi2O5 and the non-entropy-tuned NaMNO2. As shown in Supplementary Fig. 5, similar to most P2-type cathode materials, both Na0.62Mn0.67Ni0.23Cu0.05Mg0.09-2yTi2O5 and NaMNO2 cathodes exhibit multiple pairs of redox peaks, corresponding to a series of phase transition reactions occurring during charge/discharge. According to the previously reported results, the low-voltage reduction peak located at ca. 2.26 V is associated with the reduction reaction of Mn3+/Mn2+ (Supplementary Fig. 5)30,31. Other weak peaks in 2.0–3.0 V are associated with the Na+ vacancy ordering process. The redox peaks at ca. 3.47/3.30 and 3.70/3.52 V could be attributed to the redox processes of Ni2+/Ni3+ and Ni3+/Ni4+, respectively (Supplementary Fig. 5)31. The high-voltage redox peaks at ca. 4.29/3.96 V are related to oxygen anionic redox, which can be demonstrated by the results of X-ray photoelectron spectroscopy (XPS) analysis (Supplementary Fig. 6). Compared with the CV plots of the NaMNO2 cathode, a new pair of extraction/insertion peaks at ca. 3.95/3.81 V appears in all the Na0.62Mn0.67Ni0.23Cu0.05Mg0.09-2yTi2O5 cathodes and could be attributed to the redox behavior of Cu2+/Cu3+ (Supplementary Fig. 5)32. The CV results suggest that Ni2+/Ni3+ and Cu2+/Cu3+ as well as (O2−/O2n−) (1 ≤ n ≤ 3) are the redox-active couples within the potential range applied and will contribute to the capacity of the as-prepared cathodes.

The anionic redox reaction (O2−/O2n−) (1 ≤ n ≤ 3) is reversible in the first CV curves of CuMgTi-571 and NaMNO2 (Supplementary Fig. 5). However, they show a significant difference in the voltage hysteresis (e.g., the oxidation and reduction peak of anionic redox locates at different potentials). The hysteresis phenomenon is often observed in the cathode materials with anionic redox33–36. The peak potential differences (ΔEp) of anionic redox reaction in NaMNO2 is 0.59 V (4.30 and 3.71 V for oxidation and reduction peak, respectively). In contrast, the (ΔEp) of anionic redox reaction in CuMgTi-571 is decreased to 0.33 V (4.29 and 3.96 V for oxidation and reduction peak, respectively).

Figure 4a, b shows the charge/discharge profiles of CuMgTi-571 and NaMNO2 at 0.1 C of different cycles (both materials were tested using a Na metal anode in coin cell configuration), respectively. It can be seen that the NaMNO2 undergoes a severe voltage fade and further specific capacity decay during cycling, which are characteristic features of Li-excess cathodes that also involve oxygen anionic redox37,38. By contrast, the CuMgTi-571 cathode exhbits less voltage fading during the cycling test in comparison with the NaMNO2 cathode, indicating that the voltage fade issue in CuMgTi-571 is alleviated. This result can be seen more clearly in the normalized voltage capacity curve in Supplementary Fig. 7a, b. Figure 4c compares the cycle performance of CuMgTi-571 with higher entropy and NaMNO2 with much lower entropy at 0.1 C (12 mA g−1). As shown, CuMgTi-571 could deliver an initial discharge capacity of 148.2 mAh g−1, and can still maintain 132.9 mAh g−1 after 100 cycles, resulting in capacity retention of 89.6%. Although NaMNO2 delivers a comparable first discharge capacity (150.5 mAh g−1 with CuMgTi-571), the discharge capacity of NaMNO2 decreases fast and remains only 60.2% (90.6 mAh g−1) of its initial capacity after 50 cycles. Moreover, the average coulombic efficiency of CuMgTi-571 achieves 98.0% compared to 95.4% of NaMNO2, suggesting the fast dynamics and high reversibility of CuMgTi-571. To investigate the reversibility of electrochemical reaction in CuMgTi-571 and NaMNO2, dQ/dV curves after 50 cycles at 0.1 C (12 mA g−1) rate are displayed in Supplementary Fig. 8. It can be seen that the anionic redox peaks at high voltage have been well maintained in the CuMgTi-571 in comparison with NaMNO2. These sharp contrasts confirmed that entropy modulation can significantly stabilize the structures and anionic redox of P2 layered cathodes during high-voltage charge/discharge.

The rate capability of the entropy-tuned Na0.62Mn0.67Ni0.23Cu0.05Mg0.09-2yTi2O5 and the non-entropy-tuned NaMNO2 were further evaluated at various charge/discharge rates from 0.1 C to 10 C (both materials are tested using a Na metal anode in coin cell configuration, 1 C = 120 mA g−1). The CuMgTi-571 cathode shows better rate performance than NaMNO2. Among them, CuMgTi-571 demonstrates the best rate capability. When the rates are increased successively from 0.1 to 0.2, 0.5, 1, 2, 5 and 10 C (1 C = 120 mA g−1), the CuMgTi-571 cathode enables cell-specific capacities of 143.5, 128.3, 122.9, 120.1, 115.1, 104.4, and 82.6 mAh g−1, respectively. When the 0.1 C-rate switches back to 0.1 C (12 mA g−1), a discharge capacity of 151.8 mAh g−1 can be immediately recovered, indicating the tolerance for fast Na+ transport and stable structure of CuMgTi-571 cathode. The changes in the interlayer distance (d) between NaMNO2 and CuMgTi-571 are only 0.11 Å. Such a small variation of interlayer distance is not sufficient to account for the improvement in the rate capability. In contrast, the relatively larger entropy difference (e.g., 0.35 R between CuMgTi-571 and NaMNO2) and active-facet percentage difference (e.g., 16.3% between CuMgTi-571 and NaMNO2) should play a dominant role in the observed improvement on rate capability. With an increase in the entropy and active facet, the reversible capacities of Na0.62Mn0.67Ni0.23Cu0.05Mg0.09-2yTi2O5 under various rates were increased accordingly.

A similar trend can be found in their long-term cyclability, which is an important parameter for battery materials when used for practical applications. In this regard, their long cycling performances were further evaluated by charging/discharging at 1 C (120 mA g−1) for 500 cycles (The materials are tested using a Na metal anode in coin cell configuration, Fig. 5). Figure 5a, b show the voltage profiles of CuMgTi-571 and NaMNO2 for the 100th, 200th, 300th, 400th, and 500th cycles during charge/discharge at 1 C (120 mA g−1) rate, respectively. CuMgTi-571 cathode again displays less voltage fading during the long cycling compared with the NaMNO2 cathode. Figure 5c compares the cycling performances of Na0.62Mn0.67Ni0.23Cu0.05Mg0.09-2yTi2O5 and NaMNO2 at 1 C (120 mA g−1) for 500 cycles. It can be clearly seen that the CuMgTi-571 cathode with the highest entropy and [010] active facet percentage also displays better long-term stability with the highest reversible capacity (103.3 mAh g−1) and capacity retention (87%) after 500 cycles. While the reversible capacities and retention of CuMgTi-552 (89.8 mAh g−1), CuMgTi-533 (65.9 mAh g−1, 62.7%) were decreased accordingly. In sharp contrast, the NaMNO2 with the lowest entropy and least [010] active facet exhibits rapid capacity decay, which can only maintain a reversible capacity of 30.9 mAh g−1 after 500 cycles. In addition, the NaMNO2 cathode also exhibits a lower Coulombic efficiency than the CuMgTi-571 cathode during charge/discharge (Supplementary Fig. 10a).

Furthermore, we tested the long-term cycle stability of CuMgTi-571 and NaMNO2 cathode under extreme fast charging/discharging of 10 C (ca. 6 min charge/discharge, 10 C = 1200 mA g−1, both materials are tested using a Na metal anode in coin cell configuration). The discharge capacity of the NaMNO2 cathode faded rapidly during cycling and dropped to 22.9 mAh g−1 at the 1000th cycle (Fig. 5d). The CuMgTi-571 cathode can still display a fast-charging performance with a reversible capacity of 59.3 mAh g−1.
and reasonable capacity retention of 75.4% after 2000 cycles (Fig. 5d). The Coulombic efficiency of CuMgTi-571 cathode is closer to 100% in comparison with that of NaMNO2 cathode, indicating better sodiation/de-sodiation reversibility (Supplementary Fig. 10b). We compared the electrochemical performance of our entropy and active-facet tuned cathode with other reported P2-type layered cathodes tested at different conditions (Supplementary Table 7 tested at a high rate and Supplementary Table 8 tested at the same cut-off voltage). The comparison indicates a significant advance of the CuMgTi-571 in terms of capacity, cycle number, capacity retention even high-rate capability.

To further verify the benefit of entropy-driven structural stability on the Na⁺ kinetics of the P2-type layered cathodes, Naₐ₀.₆₂Mₐ₀.₆₇Nⁱ₀.₂₃Cₐ₀.₀₆Mₜ₀.ₙ₉₋₂ₚ₂ₜ₂O₂ and NaMNO₂ cathodes. Charge/discharge profiles of a CuMgTi-571 and b NaMNO₂ at 0.1 C (12 mA g⁻¹) between 2.0 and 4.3 V. c Cycling performances of CuMgTi-571 and NaMNO₂ at 0.1 C (12 mA g⁻¹) for 100 cycles. d Rate performances of Naₐ₀.₆₂Mₐ₀.₆₇Nⁱ₀.₂₃Cₐ₀.₀₆Mₜ₀.ₙ₉₋₂ₚ₂ₜ₂O₂ and NaMNO₂ from 0.1 C to 10 C (1 C = 120 mA g⁻¹). Cycling and rate tests were conducted in a coin cell using Na metal as an anode at 25 °C. Hollow and solid circles in (c, d) represent charge and discharge capacity, respectively. Triangles represent coulombic efficiency.

Fig. 4 Cycling and rate performances of Naₐ₀.₆₂Mₐ₀.₆₇Nⁱ₀.₂₃Cₐ₀.₀₆Mₜ₀.ₙ₉₋₂ₚ₂ₜ₂O₂ and NaMNO₂ cathodes. Charge/discharge profiles of a CuMgTi-571 and b NaMNO₂ at 0.1 C (12 mA g⁻¹) between 2.0 and 4.3 V. c Cycling performances of CuMgTi-571 and NaMNO₂ at 0.1 C (12 mA g⁻¹) for 100 cycles. d Rate performances of Naₐ₀.₆₂Mₐ₀.₆₇Nⁱ₀.₂₃Cₐ₀.₀₆Mₜ₀.ₙ₉₋₂ₚ₂ₜ₂O₂ and NaMNO₂ from 0.1 C to 10 C (1 C = 120 mA g⁻¹). Cycling and rate tests were conducted in a coin cell using Na metal as an anode at 25 °C. Hollow and solid circles in (c, d) represent charge and discharge capacity, respectively. Triangles represent coulombic efficiency.

To examine the effect of the conductivity on the anionic redox reaction of CuMgTi-571 and NaMNO₂ at high voltage, in situ electrochemical impedance spectroscopy (EIS) experiments were conducted using Na metal as a counter/reference electrode. Supplementary Fig. 13 shows the Nyquist plots of the CuMgTi-571 and NaMNO₂ at different potentials of the first charge and discharge process. Upon Na⁺ insertion/extraction, both samples experience a series of processes in the bulk and at the surface, leading to different suppressed semicircles in Nyquist plots of the first charge and discharge process. At the high voltage region (4.0–4.3 V), the Nyquist plots of both samples exhibit two distinct parts including a depressed semicircle in the high-frequency region and an inclined line in the low-frequency zone. The high-frequency semicircle should be related to the charge-transfer resistance. By comparing the diameters of the semicircles in the high-voltage region (Supplementary Table 9), the impedances of CuMgTi-571 and NaMNO₂ are similar, implying similar electronic conductivity of the two de-sodiated samples. Therefore, the significantly improved kinetics of the anionic redox in the
CuMgTi-571 is mainly contributed by its structure with high active facets and high entropy, which can provide more stable migration tunnels and is helpful for accelerating Na\(^{+}\) diffusion.

### Phase transition and structural stability

Na\(^{+}\) kinetics is strongly associated with the structural stability of the material and anionic redox usually results in the generation of oxygen vacancies and migration of transition metal (TM) ions, which could cause phase transition and new phase formation. To check the impact of entropy and crystal-facet tuning on the structural stability of the material, in situ synchrotron high energy X-ray diffraction (HEXRD) was conducted on CuMgTi-571 during charge/discharge (tested in coin cell using a Na metal anode), and the results are displayed in Fig. 7 and Supplementary Fig. 14. Figure 7a, b shows the waterfall and contour plot of HEXRD patterns of the CuMgTi-571 electrode charged/discharged at 0.1 C (120 mA g\(^{-1}\)) between 2.0 and 4.3 V, respectively. Upon Na\(^{+}\) extraction (charge), the (002) and (004) peaks shift toward a lower 2\(\theta\) angle due to the increase in the repulsive electrostatic interaction between adjacent MO\(_2\) sheets; while (100), (102), (108), and (112) peaks consecutively peaks shift toward a higher 2\(\theta\) angle with the contraction of a-, b-axis\(^{42-44}\). During the subsequent discharge, the XRD pattern of CuMgTi-571 exhibited an exactly opposite evolution. During the whole charge/discharge process, the well-defined peaks corresponding to the P2 phase are well maintained and returned to the initial states after one full cycle. No new peaks belonging to OP\(_{445}\), O\(_{28}\), or Z\(_6\) phase are detected when charged to 4.3 V, and no existing peaks vanish or split, indicating that new phase formation and phase transition do not occur in this material. The changes in lattice parameters upon cycling based on the refinement results are displayed in Fig. 7c. It is clear that the lattice parameter evolution is highly reversible during the charge and discharge process. The largest changes of the \(a\), \(c\), and \(V\) in the first cycle are only 0.74, 0.91, and 0.57%, respectively, which are much smaller than those of the state-of-the-art reported layered cathodes (Supplementary Table 10 and Supplementary Note 3). In addition, the structural stability of CuMgTi-571 at different cycles was also investigated and the results are shown in Supplementary Fig. 15. The XRD patterns of the cycled samples and pristine are almost identical, indicating that the P2-type structure can be well maintained after extensive cycling. The high reversibility of lattice parameter evolution and structural evolution upon Na\(^{+}\) (de-) intercalation manifests the unusual structure stability of entropy and crystal facet tuned cathode.

In situ HEXRD during heating of de-sodiated cathode powder was also performed to investigate the phase stability of entropy and crystal-facet tuned cathodes at high temperatures (heating from 30 to 450 \(^{\circ}\)C) to investigate their thermal tolerance.
The tested CuMgTi-571 and NaMNO2 materials were scraped from 4.2 V-charged cathodes. The (002) peak, which is most sensitive to the Na content during charging/discharging, is usually used to monitor the phase transformation of P2-type layered cathodes during heating. The (002) reflection of charged CuMgTi-571 continuously moves to a lower angle with increasing the temperature due to the lattice expansion caused by heating (Fig. 8a). Distinct from CuMgTi-571, the (002) reflection of charged NaMNO2 first shifts to a lower angle upon heating, and then moves to a higher angle when the heating temperature reached above 222 °C (Fig. 8b). Such phenomenon indicates that new phases such as spinel or rock-salt were formed in the charged NaMNO2 during the heating. Moreover, the (002) peak fading starts at 350 and 322 °C for the CuMgTi-571 and NaMNO2 cathodes, respectively, further confirming the inferior structural stability of the NaMNO2. The formation of a new phase and inferior structural stability would lead to irregular migration channels and hence sluggish Na$^+$ diffusion.

To further unravel the robust structural stability of the entropy and crystal-facet tuned cathodes, the morphologies of CuMgTi-571 and NaMNO2 after 500 cycles at 1 C (1 C = 120 mA g$^{-1}$) were further compared (both materials using a Na metal anode are tested in coin cell configuration). SEM images in Fig. 8c and Supplementary Fig. 16a show a well-preserved morphology of the CuMgTi-571 particles without noticeable structural degradation, displaying a stable inner structure robust enough to endure the long-term insertion/extraction of Na$^+$. In contrast, the morphology of the NaMNO2 particles changed after long-term cycling (Fig. 8d and Supplementary Fig. 16b). The smooth side surfaces before the cycles split into many thin sheets after 500 cycles, indicating that the repeated Na$^+$ de/intercalation leads to volumetric expansion and serious structural degradation, which could be associated with the fast capacity fade of the NaMNO2.

**Discussion**

The above results clearly demonstrate that fast and reversible (de-) intercalation of large Na$^+$ have been achieved simultaneously in the Na$_{0.62}$Mn$_{0.67}$Ni$_{0.23}$Cu$_{0.05}$Mg$_{0.09}$Ti$_{2}$O$_{2}$. The Na$^+$ extraction/insertion kinetics are mainly attributed to the increased structural stability and ion-diffusion channels. Configurational entropy contribution to the decreased Gibbs free energy in materials retards
dislocation movement and stabilizes crystal structure. Such entropy-driven high structural stability is robust enough to restrain volume variation, structure distortion, and irreversible phase transition, thus safeguarding a stable migration path and reversible Na\(^{+}\) (de-)intercalation. Moreover, the increased \{010\} electrochemically active planes due to the increased entropy create more favorable and efficient tunnels for the migration of Na\(^{+}\), which facilitate rapid Na\(^{+}\) (de-)intercalation and promote electrochemical kinetics.

In summary, configurational entropy could result in a thermodynamically stable structure through a local minimization of Gibbs free energy, and a high percentage of \{010\} facets in the layered cathodes could provide more channels for ion transport. Based on the advantages of high entropy and high active facets, a combinatorial strategy of entropy modulation and active-facet modulation was developed to stabilize the structure and optimize ion-migration pathways of P2-type layered cathodes. The correlation of entropy, active facet, Na\(^{+}\) kinetics, and electrochemical performance of the Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)Cu\(_{0.05}\)Mg\(_{0.09}\)-2\(_y\)Ti\(_y\)O\(_2\), \(y = 0.01, 0.02, \text{and } 0.03\) were prepared using a stirring hydrothermal method. In a typical synthesis, stoichiometric amounts (with 7% excess sodium) of NaCH\(_3\)COO·3H\(_2\)O, Mn(CH\(_3\)COO)\(_2\)·4H\(_2\)O, Ni(CH\(_3\)COO)\(_2\)·4H\(_2\)O, Cu(CH\(_3\)COO)\(_2\)·H\(_2\)O, Mg(CH\(_3\)COO)\(_2\)·4H\(_2\)O, and TiO\(_2\) were mixed in 60 mL of deionized water. An appropriate amount of H\(_2\)C\(_2\)O\(_4\) (molar ratio of oxalic acid to metal ions is 1:3:1) was used as a precipitant agent and also dissolved in deionized water. H\(_2\)C\(_2\)O\(_4\) solution was added dropwise into the above mixture under stirring. Afterward, the reaction mixture was put into 100 mL Teflon-lined autoclaves and aged at 130 °C for 12 h under stirring. The green precipitates were collected by solvent evaporation at 80 °C. The resulting precipitates were annealed in air at 500 °C for 6 h and then at 900 °C for 12 h, and P2-Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)-2\(_y\)Ti\(_y\)O\(_2\) samples were finally obtained. In addition, Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.21}\)Cu\(_{0.05}\)Mg\(_{0.015}\)Zn\(_{0.015}\)Sn\(_{0.015}\)Y\(_{0.015}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.37}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)Cu\(_{0.05}\)Mg\(_{0.07}\)Ti\(_{0.01}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.37}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)Cu\(_{0.05}\)Mg\(_{0.05}\)Ti\(_{0.02}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)Cu\(_{0.05}\)Mg\(_{0.07}\)Ti\(_{0.01}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)Cu\(_{0.05}\)Mg\(_{0.03}\)Ti\(_{0.03}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.21}\)Cu\(_{0.05}\)Mg\(_{0.01}\)Zn\(_{0.015}\)Sn\(_{0.015}\)Y\(_{0.015}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.21}\)Cu\(_{0.05}\)Mg\(_{0.01}\)Zn\(_{0.015}\)Sn\(_{0.015}\)Zr\(_{0.015}\)O\(_2\) were also prepared by same method. The obtained P2-type layered Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)-2\(_y\)Ti\(_y\)O\(_2\) samples were labeled as CuMgTi-533, CuMgti-552, CuMgTi-571, and NaMNO2, respectively.

**Methods**

**Materials synthesis.** A series of multi-element P2-structured layered oxides (Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)Cu\(_{0.05}\)Mg\(_{0.09}\)-2\(_y\)Ti\(_y\)O\(_2\), \(y = 0.01, 0.02, \text{and } 0.03\) were prepared using a stirring hydrothermal method. In a typical synthesis, stoichiometric amounts (with 7% excess sodium) of NaCH\(_3\)COO·3H\(_2\)O, Mn(CH\(_3\)COO)\(_2\)·4H\(_2\)O, Ni(CH\(_3\)COO)\(_2\)·4H\(_2\)O, Cu(CH\(_3\)COO)\(_2\)·H\(_2\)O, Mg(CH\(_3\)COO)\(_2\)·4H\(_2\)O, and TiO\(_2\) were mixed in 60 mL of deionized water. An appropriate amount of H\(_2\)C\(_2\)O\(_4\) (molar ratio of oxalic acid to metal ions is 1:3:1) was used as a precipitant agent and also dissolved in deionized water. H\(_2\)C\(_2\)O\(_4\) solution was added dropwise into the above mixture under stirring. Afterward, the reaction mixture was put into 100 mL Teflon-lined autoclaves and aged at 130 °C for 12 h under stirring. The green precipitates were collected by solvent evaporation at 80 °C. The resulting precipitates were annealed in air at 500 °C for 6 h and then at 900 °C for 12 h, and P2-Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)-2\(_y\)Ti\(_y\)O\(_2\) samples were finally obtained. In addition, Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.21}\)Cu\(_{0.05}\)Mg\(_{0.015}\)Zn\(_{0.015}\)Sn\(_{0.015}\)Y\(_{0.015}\)O\(_2\), and Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.21}\)Cu\(_{0.05}\)Mg\(_{0.01}\)Zn\(_{0.015}\)Sn\(_{0.015}\)Zr\(_{0.015}\)O\(_2\) were also prepared by same method. The obtained P2-type layered Na\(_{0.62}\)Mn\(_{0.67}\)Ni\(_{0.23}\)-2\(_y\)Ti\(_y\)O\(_2\) samples were labeled as CuMgTi-533, CuMgti-552, CuMgTi-571, and NaMNO2, respectively.

**Physicochemical characterizations.** All the samples were characterized by scanning electron microscopy (SEM, Hitachi S-4800), transmission electron microscopy (TEM, JEOLJEM-2100), inductively coupled plasma-mass spectrometry (ICP-MS, Thermo X2), X-ray diffraction (XRD, Bruker D8 Discover X-ray analytical systems with Cu K\(\alpha\) radiation), energy dispersive X-ray (EDX, an EDX detector system attached to FEI Talos-S), and X-ray photoelectron spectroscopy (XPS, ESCALAB 250Xi).

The cycled electrode samples for ex situ XPS and XRD characterization were obtained by disassembling the cycled half-cells using Na metal anode in Ar-filled glovebox (H\(_2\)O < 0.1 ppm, O\(_2\) < 0.1 ppm). The electrodes were rinsed using dimethyl carbonate (DMC, 99%, Sigma Aldrich) repeatedly, then dried in glovebox to remove the solvents. For ex situ XPS measurements, the dried electrodes were first placed into a vacuum transfer vessel in glovebox and then transferred into the
XPS ultrahigh vacuum chamber. For ex situ XRD measurements, the dried electrodes were transferred from glovebox and then tested in air.

HEXRD measurements. The in situ synchrotron high energy XRD characterizations of the CuMgTi-571 electrode were carried out at the beamline 11-ID-C of the Advanced Photon Source at Argonne National Laboratory with a wavelength of 0.1173 Å. The Na|CuMgTi-571 coin cells (type CR2032) with two identical holes on the top and bottom cases were assembled in the argon-filled glovebox and then sealed with Kapton tape after filling with the electrolyte. The electrolyte was 1 M NaPF₆ (≥ 98%, Sigma Aldrich) dissolved in propylene carbonate (PC, 99%, Sigma Aldrich) with 2 vol% fluoroethylene carbonate (FEC, 99%, Sigma Aldrich). A constant charge/discharge current of 20 mA g⁻¹ was applied by using a MACCOr system between 2.0 and 4.3 V at 25 °C. Each 2D diffraction pattern was obtained by a PerkinElmer amorphous silicon 2D detector during the charge/discharge process and then were converted to 1D patterns using fit2D software with a calibrated using a standard CeO₂ sample with the same experimental setup. The postmortem cathode HEXRD measurements during the heating was also conducted at 11-ID-C of Advanced Photon Source. Both the NaNMO₂ and NiMgTi-571 cathode were charged to 4.2 V and then disassembled in the glovebox and scratch off the active materials carefully. The charged cathode materials were then loaded into a stainless-steel DSC autoclave and sealed in the glovebox to prevent any humid/air contamination during the in situ heating experiment. A Linkam TS1500 furnace was used to heat the charged samples from 30 to 450 °C at 5 °C min⁻¹ and a PerkinElmer 2D detector was continuously used to acquire the 2D diffraction data during the heating process.

Electrochemical characterization. All the electrochemical measurements were performed using CR2016 coin cells assembled in an Ar-filled glovebox. Cathodes were prepared by casting a slurry composed of 70 wt% active material, 20 wt% acetylene black, and 10 wt% polyvinylidene difluoride (PVDF, HSV900, Arkema) binder dissolved in N-methyl-2-pyrrolidone (NMP, 99.9%, Aladdin) on an Al foil (99%, 16 μm in thickness, Hefei Kejing, China) and drying the slurry under vacuum at 120 °C for 12 h. The areal loading of the active material was about 3–4 mg cm⁻². A high-purity Na metal foil was used as anode and glass fiber as separator (675 μm in thickness, 2.7 μm average pore size, Whatman). Na metal foil (1.5 mm in thickness, 99.5%, Sinopharm) was made from Na chunks under mineral oil, which was removed from the surface of Na by dried paper inside the glovebox. Na chunks was then pressed into a thin sheet under 50 Pa inside the glovebox, which was further punched into 16 mm diameter plates and used as an anode. The electrolyte was 1 M NaClO₄ dissolved in a mixed solvent of propylene carbonate (PC) and fluoroethylene carbonate (FEC) (49:1, v/v). The amount of electrolyte used in the coin cells was controlled at around 280 µL. Charge-discharge experiments were performed galvanostatically between 2.0 and 4.3 V at different C-rates (1 C = 120 mA g⁻¹) on a LANHE-CT2001A test system (Wuhan, China). Cyclic voltammetry (CV) measurements were carried out on a CHI760E electrochemical workstation (Chenhua, Shanghai China) at a scanning rate of 0.2 mV s⁻¹ within the potential of 2.0–4.3 V. Galvanostatic intermittent titration technique (GITT) tests were conducted in the voltage range of 2.0–4.3 V by applying repeated current pulses at a current rate of 0.1 C for 15 min followed by a 3 h rest. The in situ electrochemical impedance spectroscopy (EIS) experiments were executed using a PARSTAT2273 electrochemical workstation in a frequency range from 0.1 Hz to 100 KHz with an AC amplitude of 5 mV. EIS tests of Na||CuMgTi-571 and Na||NaNMO₂ coin cells
were performed after each potentiostatic intermittent titration technique (PITT) program completed. The step potential difference was 0.1 V, and the step elapsed time was 5000 s. Each EIS test contains 50 data points. All the electrochemical tests were made at 25 °C.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

The data that support the findings of this study are available from the corresponding authors F.F., K.A., S.-G.S and G.-L.X. upon reasonable request.

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**Author contributions**

F.F. and G.-L.X. conceived the idea and designed the experiments; X.F. prepared the P2-layered cathode for highly reversible oxygen redox chemistry. F.F. and G.-L.X. wrote and revised the manuscript.

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