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Li₄NiTeO₆ as a positive electrode for Li-ion batteries[†]

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Layered Li₄NiTeO₆ was shown to reversibly release/uptake ~2 lithium ions per formula unit with fair capacity retention upon long cycling. The Li electrochemical reactivity mechanism differs from that of Li₂MO₃ and is rooted in the Ni⁴⁺/Ni²⁺ redox couple, that takes place at a higher potential than conventional LiNi_{1-x}Mn_xO₂ compounds. We explain this in terms of inductive effect due to Te⁶⁺ ions (or the TeO₆⁶⁻ moiety).

At their early stages (1991), commercial Li-ion batteries were using two intercalation compounds LiCoO2 and graphite as positive and negative electrodes, respectively.¹ Since then, layered oxides have garnered increasing interest with numerous compositional attempts to alleviate the high cost and toxicity associated with cobalt, while preserving the large energy densities resulting from cobalt's high redox voltage and low molecular weight. Through such an exploration, nickel based compounds such as LiNi_{0.33}Mn_{0.33}Co_{0.33}O₂ and LiNi_{0.5}Mn_{0.5}O₂ have been identified to be prime alternatives to LiCoO₂ in the battery market, owing to the accessibility of two-electron Ni⁴⁺/Ni²⁺ redox process that enables capacities of ~ 200 mA h g⁻¹ to be achieved.^{2,3} Additional improvements, leading to capacities higher than 220 mA h g^{-1} , have been made via the introduction of lithium-rich layered oxides $(Li_{1+x}Ni_{\nu}Co_{z}Mn_{1-x-\nu-z}O_{2})$, which are basically considered to be composed of two different layered oxides LiMO2 and Li2MO3.4 Nevertheless, a poor understanding of both their Li-insertion mechanism and continuous voltage decay upon cycling has plagued the commercial use of these Li-rich phases. Addressing these issues has drained humongous research efforts worldwide, but no consensus had been reached until our group explained, via a simple chemical approach combining XPS and EPR experiments, that: (i) the extra measured capacity within these materials is rooted in cumulative reversible cationic $(M^{n+} \rightarrow M^{(n+1)+})$ and anionic $(O^{2-} \rightarrow O_2^{2-})$ redox processes,

and (ii) that the voltage decay upon cycling can be minimized through Sn-substitution for Ru in $\text{Li}_2\text{Ru}_{1-y}\text{Sn}_y\text{O}_3$.⁵ We explained the overall mechanism in terms of Ru(4d)–O(2p) hybridization, implying the proximity of Ru-4d and O-2p levels within the material band structure.

Because of the rarity and high cost of Ru, one part of this continuing work is the exploration of Ru-free compounds showing *n*d–sp hybridization and belonging to the Li₄MM'O₆ family, which contains a huge number of members, as the only restriction requires that the sum of M and M' formal oxidation states be equal to 8 to ensure electro-neutrality.⁶ Based on the fact that most of the studies addressing holes on oxygen deal with Ni-based oxides,³ we have selected Ni-based Li₂Ni_{0.5}Te_{0.5}O₃ (hereafter represented as Li₄Ni²⁺⁻Te⁶⁺O₆) with the hope of achieving high capacity through (i) the 2e⁻ redox process associated with Ni⁴⁺/Ni²⁺ as observed for the aforementioned layered oxides LiNi_{0.33}Mn_{0.33}Co_{0.33}O₂ and LiNi_{0.5}Mn_{0.5}O₂,^{2,3} and (ii) oxygen redox activity, provided that a high degree of Ni(3d)–O(2p) hybridization can be achieved.

 Li_4NiTeO_6 was prepared by a conventional ceramic method from a stoichiometric mixture of Li_2CO_3 (Aldrich), $NiC_4H_6O_4$ · $4H_2O$ and TeO_2 (across organics) (2:1:1) annealed at about 1000 °C for 18 h, followed by heating at 1050 °C for 6 h with intermediate grinding. 10% Li_2CO_3 was added in excess to compensate lithium loss due to high temperature heat treatment. The structure of the compound was analyzed by Rietveld refinement⁷ using the Fullprof program⁸ against high-resolution synchrotron powder diffraction data, collected on the 11-BM beamline at the Advanced Photon Source (APS, Argonne National Laboratory) with a wavelength of 0.4138 Å.

Bragg peaks unambiguously indicate that the sample crystallizes in the *C2/m* space group with cell parameters a = 5.1584(1) Å, b = 8.8806(1) Å, c = 5.1366(1) Å and $\beta = 110.241(1)^{\circ}$ (V = 220.777(3) Å³), *i.e.* a cell similar to the one reported for Li₂MnO₃.⁹ Refinements using different distributions of atoms within the layers led to a structure with Li layers alternating with honeycomb Li/Ni/Te layers (Fig. 1). In the latter, Li and Ni are statistically distributed in the same octahedral site (colored in yellow/green), surrounding TeO₆ octahedra (colored in blue). The final refinement which includes a strain analysis is shown in Fig. 1, and the deduced structural parameters are detailed in ESI[†] (Fig. S1, Tables S1 and S2).

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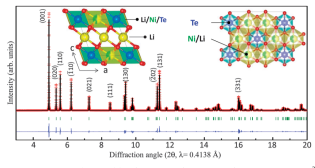


Fig. 1 Structure and Rietveld refinement of Li₄NiTeO₆ ($R_{Bragg} = 3.17\%$, $\chi^2 = 10.9$). Inset: Stacking of Li/Ni/Te layers and Li layers (left), view of one honeycomb Li/Ni/Te layer, for which Li and Ni are equally distributed on the same site (right). Li is yellow, Ni is green, Te is blue and O is red.

Electrochemical behaviour of Li₄NiTeO₆ as a cathode material has been explored in lithium half cells (Swagelok[™] cells) using 1 M $LiPF_6$ in EC:PC:DMC (1:1:3 wt ratio) as an electrolyte. The working electrode was made by milling the sample with 20% carbon black (Super P, Timcal) using a high energy SPEX 800 miller for 20 minutes. The voltage-composition trace is shown in Fig. 2a. Upon charging, the cell rapidly reaches a voltage plateau located at 4.2 V vs. Li^+/Li , which lasts until the removal of ~1.5 Li^+ ions, prior to smoothly and continuously increasing to 4.6 V through the end of charging process which corresponds to total removal of $\sim 2 \text{ Li}^+$ ions solely based on electron counting. The following discharge shows an identical profile (smooth decay + plateau), but is shorter since solely 1.5 Li⁺ ions are reinserted upon discharge. Such a large irreversibility between the first charge and discharge is reminiscent of some electrolyte decomposition at high potential, thus leading to uncertainty in the amount of Li reinserted, which is below 2, as we will show later on using XPS. This leads to an overall reversible capacity of 110 mA h g^{-1} for Li₄NiTeO₆ which is sustained upon cycling since the cell retains nearly 90% of its initial capacity after 100 cycles (Fig. 2b). The derivative plot shown in the inset of Fig. 2a confirms a single redox process upon cycling. Increasing the charging potential to 5 V showed a small broad second oxidation peak (left inset in Fig. 2b) centred at 4.8 V and a rapid decay of cell performance upon cycling (right inset in Fig. 2b). Such capacity decay is mainly rooted in a voltage-driven transformation/ decomposition of Li₄NiTeO₆, as suggested by high degree of amorphisation of the 5 V oxidized sample.

Overall, the voltage profile of Li/Li₄NiTeO₆ cells drastically differs from those measured previously for Li/Li₂Ru_{1-y}M_yO₃ (M = Sn, Mn)^{5,10} cells, as they present a single voltage plateau upon consecutive cycles instead of a stair-case charge profile followed by a S-type curve upon discharge. This indicates a different lithiation–delithiation mechanism that we have explored by both *in situ* XRD and XPS measurements.

Fig. 2c shows the XRD patterns collected during the charge of a home-made $\text{Li}/\text{Li}_4\text{NiTeO}_6$ electrochemical cell equipped with a Be window. Upon removal of Li, there is appearance of a set of weak Bragg peaks which grow at the expense of the peaks of the pristine phase, and become single phase after the removal of 2 lithium ions. This new $\text{Li}_{4-x}\text{NiTeO}_6$ phase, formed at the end of charge (4.6 V), with *x* being not accurately defined due to competing electrolyte oxidation, can also be indexed to a C2/m monoclinic cell

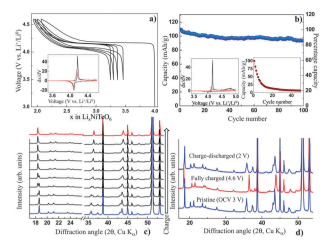
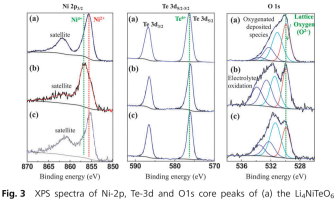


Fig. 2 Electrochemical performance of Li₄NiTeO₆. (a) Voltage *vs.* composition profile for Li₄NiTeO₆, with (as inset) the derivative plot for the first (black) and second (red) cycles. (b) Capacity retention for a Li/Li₄NiTeO₆ cell together with (lower left inset) a derivative plot for the first (black) and second (red) cycles on cycling to a higher voltage (2 to 5 V); (lower right inset) a capacity retention plot when the cell is cycled over 2 to 5 V. (c) *In situ* XRD patterns (λ_{cu}) while Li₄NiTeO₆ is charged at a C/20 rate, (d) patterns collected for the pristine electrode, charged to 4.6 V and fully discharged issued from (c).

with lattice parameters a = 5.037(1) Å, b = 8.716(1) Å, c = 5.163(1) Å; $\beta = 108.79(1)^\circ$, *i.e.* a volume of 214.61(5) Å³ which is less than 3% smaller than that of pristine phase. The *c*-lattice parameter increase upon Li removal arises from increasing repulsion between MO₂ layers which become less screened by Li-ions, while the decrease in both *a* and *b* lattice parameters originates from the cation size reduction linked to the Ni²⁺ \rightarrow Ni⁴⁺ oxidation process. Upon discharge, the XRD pattern for the fully discharged material being similar to that of the pristine phase (Fig. 2d) indicates the structural reversibility of the electrochemical process. A closer examination of the lattice parameter values indicated a slightly lower unit cell volume for the charged–discharged sample (V = 219.28 Å³ vs. 220.78 Å³), suggesting the presence of tiny amounts of smaller Ni⁴⁺ (lower ionic radii than Ni²⁺) in the reduced sample as confirmed by XPS.

XPS analyses (Fig. 3) were carried out for pristine Li₄NiTeO₆ electrode mixed with 20% carbon (a), and similar electrodes charged to 4.6 V (b), and charged to 4.6 V and then discharged to 2 V (c). The Ni- $2p_{3/2}$ core spectra for the pristine sample show a main peak at 855.6 eV and a satellite peak at 861.7 eV, confirming the presence of Ni²⁺ in the pristine phase.¹¹ Upon charging to 4.6 V, the Ni-2p_{3/2} peak shifts to a higher binding energy (856.7 eV) which corresponds to Ni4+ with a small shoulder indicative of remaining part of Ni²⁺ which we could estimate to be $\sim 20\%$ by XPS analysis. This agrees with our electrochemical measurements, indicating minor electrolyte decomposition so that all charging current is not fully used to oxidize Ni²⁺ to Ni⁴⁺. Upon discharge, the Ni-2p_{3/2} binding energy shifts back to that of pristine phase, confirming mainly the reversibility of the process with a very small amount of residual Ni⁴⁺ (tiny shoulder in the left of the Ni²⁺ peak) in agreement with the electrochemical data. Turning to Te-3d core spectra, the Te-3d_{5/2} peak at 576.3 eV is characteristic of Te⁶⁺ referenced to Li4TeO5 and Li2TeO3 (Te-3d5/2 respectively at 576.3 eV and 575.9 eV). It remains unaltered for the entire charge-discharge process indicating that Te does not participate



pristine electrode, (b) charged to 4.6 V, (c) further discharged to 2.0 V.

in the redox process.¹² Lastly, the O-1s core spectra of the aforementioned samples were collected to check for eventual observation of peroxo-like species as seen with $\text{Li}_2\text{Ru}_{1-y}M_yO_3$ compounds reported earlier.^{5,10} The peaks characteristic of O²⁻ anions belonging to the crystalline network at 529.9 eV and weakly adsorbed surface species are visible in the pristine sample spectra. The O-1s spectra of the charged sample show an increase of the high binding energy components (~ 532.5 eV and ~ 534.0 eV) ascribed to some oxidation of the electrolyte,¹³ however no significant change in the O-lattice/ (Ni + Te) atomic ratio was observed between the pristine and charged samples, in line with the absence of an additional peak which could be due to formation of peroxo-like species. However, caution should be exercised here, owing to the broadness of the collected spectra for the charged samples.

Conclusively, the electrochemical behaviour of Li_4NiTeO_6 differs from other reported Li-rich layered oxides and Mn/Sn-substituted Li_2RuO_3 systems, although they all belong to the same structural family. The main difference lies in the shape of cycling profile, which shows a classical two-phase insertion–deinsertion process. This indirectly implies the non-participation of anionic species in contrast to $Li_2Ru_{1-y}M_yO_3$ (M = Mn, Sn) system, suggesting that the total capacity of Li_4NiTeO_6 can simply be explained on the basis of a classical insertion process involving the Ni⁴⁺/Ni²⁺ redox couple. This is further confirmed by full preservation of the host framework throughout cycling as deduced by XRD.

Overall, the insertion mechanism in $\text{Li}_4\text{NiTeO}_6$ can be compared with typical intercalation seen in layered $\text{LiNi}_{0.5}\text{Mn}_{0.5}\text{O}_2$ and not with that of $\text{Li}_2\text{Ru}_{1-y}\text{M}_y\text{O}_3$ (M = Mn, Sn), the reason most likely being that the 3d energy levels associated with Ni^{2+} are higher in energy than the 4d levels of Ru^{4+} , therefore diminishing the chances of undergoing pronounced M(nd)–X(np) metal–ligand hybridization. The absence of two distinct steps corresponding to the $\text{Ni}^{2+}/\text{Ni}^{3+}$ and $\text{Ni}^{3+}/\text{Ni}^{4+}$ redox processes as Ni^{2+} is oxidized to Ni^{4+} is most likely due, as explained for $\text{LiNi}_{0.5}\text{Mn}_{0.5}\text{O}_2$, to the overlapping of two redox couples.³

Lastly, the higher Ni²⁺/Ni⁴⁺ redox voltage observed for Li₄NiTeO₆ (4.2 V) as compared to other Ni²⁺-based layered oxides (<4 V) can simply be explained by considering Te⁶⁺ in Li₄NiTeO₆ as a TeO₆⁶⁻ moiety. This TeO₆⁶⁻ anion – being more electronegative than O²⁻ – increases the redox potential through its inductive effect, as observed in polyanionic compounds when SO₄²⁻

units are replaced by PO_4^{3-} for instance. The significant increase in potential through this inductive effect is related to the closeness in energy of the Te(5p) orbitals and O(2p) orbitals, as depicted in Fig. S2, ESI.[†] Specifically, the strong Te(5p)– O(2p) hybridization lowers the electron density on the Ni–O bonds, consequently increasing their ionic character and therefore the energy required to oxidise Ni²⁺ to Ni⁴⁺. This is confirmed by XPS as the (O-1s lattice – Ni-2p_{3/2}) binding energy difference of 326.7 eV is higher (by 0.3 – 0.4 eV) than values reported for Ni²⁺ based layered oxides.^{11,14} This chemical approach combined with simple concepts of orbital interactions could provide an interesting means for tuning the potential of Ni⁴⁺/Ni²⁺ redox couple in other layered oxides. A variety of other Li₄NiMO₆ compounds are being presently tested to fully validate this point.

To summarize, we have reported that the layered oxide Li₄NiTeO₆ reversibly reacts with 1.5 Li⁺ at a potential of 4.2 V vs. Li⁺/Li⁰ through a classical insertion mechanism involving the Ni⁴⁺/Ni²⁺ redox couple. With a capacity of only 110 mA h g^{-1} and the use of a toxic element such as Te, the practical interest in this phase is limited clearly. Still, this could be made useful for practical applications provided a less heavy and toxic initiator can be found to replace Te⁶⁺. From a fundamental standpoint, however, this study stresses that the design of compounds showing both anionic and cationic redox activities is not trivial in the absence of theoretical support. Ni was obviously not the right choice, presumably due to large separation between Ni⁴⁺-3d levels and O-2p levels. This is an open call to theorists for evaluating the M(nd)-X(n'p) metal-ligand hybridization strength in these layered Li₄MM'O₆ oxides for various M and M', and additionally for testing our findings about the feasibility of tuning the redox voltage of M via the proper choice of an inactive M' cation. We await fruitful experimentalist-theorist interactions.

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