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Structural and electrochemical insights into novel Wadsley Roth $Nb_7Ti_{1.5}Mo_{1.5}O_{25}$ and $Ta_7Ti_{1.5}Mo_{1.5}O_{25}$ anodes for Li-ion battery application†

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Niobium based anodes are gaining increasing popularity for application in high-power lithium-ion batteries, due to their high theoretical capacities, inherent safety at high current densities, and long-term stability. Here, we report the discovery and characterisation of a new Wadsley Roth niobate system, $Nb_7Ti_15Mo_{1.5}O_{25}$, showing that it is isostructural with known systems: Nb_9PO_{25} and Nb_9VO_{25} . To evaluate the material's electrochemical performance, including performance at high current densities (for potential high power applications), and long-term stability, Li half-coin cells were prepared. The material showed an initial capacity of 268(9) mA h g⁻¹ at 0.01 A g⁻¹ (voltage range of 2.5–1.0 V). However, in subsequent cycles, some of this initial capacity is lost, which is attributed to Li trapping associated with the presence of reducible MoO₄ units, similar to the situation observed for isostructural Nb₉VO₂₅. After this initial irreversible capacity loss, the material showed good performance at high current density rates, such that at 2 A g⁻¹ and 4 A g⁻¹ respective capacities of 132(10) mA h g⁻¹ and 115(14) mA g⁻¹ were delivered. Moreover, the material showed respectable capacity retention (97%) after being cycled for 100 cycles at 0.2 A g^{−1}. In order to identify the different Nb, Ti, Mo redox couples involved in this system, a Ta analogue was also synthesized (Ta₇Ti_{1.5}Mo_{1.5}O₂₅) and the electrochemical performance for this phase is also reported. This phase shows a lower initial capacity at 0.01 A g^{-1} (140(3) mA h g^{-1}) than the Nb analogue in the same voltage range, which can be increased (225 mA h g⁻¹) if a lower cutoff voltage (0.5 V) is applied. The capacity retention for this Ta system after 100 cycles at 0.2 A g^{-1} is similar to the Nb analogue (97%). Further work has explored whether the Nb–Ti–Mo contents could be varied, and these results showed that single phase Nb_{10−2x}Ti_xMo_xO₂₅ samples could be prepared for 1.5 \leq x \leq 1.75, and electrochemical testing results for the $x = 1.75$ endmember are also reported. Overall, this research highlights the synthesis and electrochemical characterisation of two new Wadsley Roth phases, and further highlights the challenges associated with the presence of reducible cations in tetrahedral sites in such structures with respect to minimising initial irreversible capacity loss. PAPER
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Introduction

With the target of meeting Net Zero in the coming decade, interest in Li-ion battery (LiB) technology has increased substantially, due to their applications in technologies to limit emissions with respect to energy infrastructure. As a prerequisite for commercialization, LiBs need to be safe, cheap, have long lifetimes, and operate at high capacities. $1-5$ To date,

graphite (Fig. 1) is the typical anode of choice and fulfils a large proportion of commercialization requirements (cheap, long-life and a high theoretical capacity of 372 mA h g^{-1}),⁶ but is limited to low (dis)charging rates due to inherent safety implications associated with the formation of lithium dendrites at high charge/discharge rates.⁷ Fast charging and discharging are essential in high-power applications such as power tools, drones, battery powered fork lifts, mining trucks, and rail transport. $Li_4Ti_5O_{12}$ (175 mA h g^{-1} ; LTO)⁸ spinel (Fig. 1) is a suitable alternative to graphite, which offers a fastcharging solution. However, while this material has favourable attributes in terms of safety and rate capability (the higher operating voltage $(1.5 \text{ V} \text{ vs. Li/Li}^+)$ removes the risk of Li dendrite formation), 9 it leads to a lower energy density for the overall battery compared to a graphite anode cell.

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Fig. 1 Crystal structures of graphite (left), LTO (middle), and Wadsley–Roth Nb₉PO₂₅ block (3 × 3 block highlighted) structure (right).

A strategy to improve the energy density for high power application can be found in the use of Wadsley–Roth block structure materials, such as the titanium-doped niobium oxide materials, *e.g.* TiNb₂O₇ (TNO), Ti₂Nb₁₀O₂₉ and TiNb₂₄O₆₂.^{10–15} The dual electron redox ability of the niobium cation, with not only the Nb^{5+}/Nb^{4+} redox couple but a further reduction to Nb^{3+} , coupled with the single redox of Ti^{4+} , allows the theoretical gravimetric capacities to be greatly enhanced, while the operating voltage window of the niobium-based systems is comparable to LTO, thus avoiding Li dendrite formation issues.¹⁸ However, these systems can suffer from low electronic conductivity prior to lithiation. Increased initial conductivity is seen if oxygen vacancies are introduced into these systems along with carbon encapsulation as identified by Liu et al. in a study of a $Nb_{12}O_{29-x}/C$ system.¹⁶ These systems have attracted considerable recent interest, with a wide range of compositions studied, with additional recent examples reported for W-doped niobate and Mo-doped niobate systems including: $Nb_{12}WO_{33}$,¹⁷ $Nb_{16}W_5O_{55}$,¹⁸ $MoNb_{12}O_{33}$,^{19,20} and $(W_{0.2}V_{0.8})_3O_7$.²¹ In addition to oxide materials work has also been done on heterogeneous transition metal selenide systems (FeSe₂/CoSe₂) and $\cos\left(\frac{S_2}{S_1S_2}\right)$ which can provide high capacities and cycle stability at high rates $(4 \text{ A } g^{-1})$.^{22,23}

These Wadsley–Roth structures can be classed as oxygendeficient derivatives of the ReO_3 crystal structure, containing blocks of corner-linked octahedra, where the oxygen deficiency is accommodated by crystallographic shear planes, in addition to tetrahedral cations in some systems. The corner linked blocks of distorted octahedra are translated in one dimension making $(n \times m)$ networks of MO₆ octahedra with *n* and *m* denoting the length and width of the block.²⁴ At the end of these blocks a crystallographic shear plane is created by edgesharing octahedra, $25,26$ and these crystallographic shear planes help to stabilise the structure by preventing excessive volume expansion on lithiation, thus limiting the degradation of the anode material. $27-29$ The block size in these structures is highly dependent on the metal-to-oxygen ratio, and a higher number of oxygens per metal allows more connections between the corner-sharing octahedra resulting in more substantial block sizes.²⁹ As noted above, the Wadsley–Roth structures are not limited to solely metal octahedra geometry, and in some cases tetrahedral metal sites are also observed. 24 Examples of compounds with such tetrahedral sites include H–Nb₂O₅ and Nb₉PO₂₅.^{30–33} The former Nb₂O₅ material has multiple polymorphs, which are not limited to the Wadsley– Roth structure, and have been discussed in detail from a structural and electrochemical perspective by Griffith et al.³⁴

Of direct relevance to the work reported in this paper is $Nb₉PO₂₅$. This phase was initially investigated by Roth and co-workers in 1965 and shown to be a (3×3) block structure consisting of NbO₆ octahedra and PO₄ tetrahedra (Fig. 1).^{30,31} In 2002 Patoux et al. conducted lithium insertion studies and concluded that in Nb_9PO_{25} there was an approximate 10% volume expansion during lithiation, with a reported intercalation of 13.5 Li ions per formula unit – and a capacity of 210 mA h g^{-1} .³⁵ The related compound Nb₉VO₂₅ is isostructural to Nb_9PO_{25} , with the tetrahedral sites being VO₄.³⁶⁻³⁸ The main difference with respect to their electrochemical performance, resides with the significant first cycle loss of the V-analogue, attributed to irreversible reduction of V^{5+} to V^{3+} which limits the subsequent lithium intercalation. $35,39,40$ In 1983 Cava et al. investigated the lithium insertion mechanisms associated with these Wadsley-Roth type structures.²⁶ The Li inserts into cavity sites within the block structure, and it has been proposed that the crystallographic shear planes in these systems help to stabilize the structure and prevent octahedral tilt distortions from occurring, thus maintaining open pathways for fast lithium diffusion. Cava and co-workers identified clear trends between 12 Wadsley–Roth structures and their electronic structure and phase evolution during (de) lithiation.²⁶ The voltage profiles for these Wadsley–Roth shear structures can be split up into 3 sections. The first region is a solid solution with a sloping profile, the second being a two-phase like region with a flatter voltage profile. Depending on the structure this region can be almost flat, like in $H-Nb₂O₅$, or have a slight slope $(TiNb₂O₇)³⁴$ The third region is a solid solution similar to the first region. 29

As part of an investigation into new $Nb₂O₅$ based anode materials, we investigated the substitution of equal amounts of Ti and Mo into $Nb₂O₅$ leading to the identification of a novel Nb–Ti–Mo–O material, which is isostructural to Nb_9PO_{25} and Nb₉VO₂₅. Initial experiments showed single phase samples for the composition $Nb_7Ti_{1.5}Mo_{1.5}O_{25}$ (NTMO71515), and here we report the characterisation of this material and the related Ta-analogue (Ta₇Ti_{1.5}Mo_{1.5}O₂₅; TTMO71515).

Through subsequent work on the more technologically interesting Nb system, we illustrate that the phase has a small solid solution range, which can be written as $Nb_{10-2x}Ti_xMo_xO_{25}$ (1.5 $\leq x \leq 1.75$).

Experimental

Synthesis and characterisation

 $Nb_7Ti_{1.5}Mo_{1.5}O_{25}$ (NTMO71515) and Ta₇Ti_{1.5}Mo_{1.5}O₂₅ (TTMO71515) were prepared using the standard solid-sate method. Stoichiometric amounts of $Nb₂O₅$ (Alfa Aesar, 99.5%), Ta₂O₅ (Alfa Aesar, 99%), MoO₃ (Alfa Aesar, 99.95%), and TiO₂ (Anatase; Alfa Aesar) were weighed out and ground together in an agate pestle and mortar, before heating in an alumina crucible at 900 °C for 12 hours under air. The samples were reground and reheated for a further 12 hours at 900 °C under air. Further experiments investigated whether a solid solution range, Nb_{10−2x}Ti_xMo_xO₂₅, was possible, and x values between $1.25 \le x \le 2$ were investigated.

Phase purity was confirmed by powder X-ray diffraction (PXRD) using a Bruker D2 diffractometer between 15-70° 2θ (Co K_{α}). For structure determination, data were collected between 5–120° 2θ on a PANalytical Empyrean diffractometer (Cu K_{α}) for 4 hours, with Rietveld refinements conducted using GSAS II.

Raman spectra were collected using a Renishaw inVia Raman microscope with a 532 nm laser line at 1% power, involving 10 seconds of exposure and data accumulated over five times, in the range of 50–1500 cm^{-1} .⁴¹

Scanning electron microscopy (SEM) images were collected on a Hitachi TM4000 plus tabletop microscope using back scattering electrons at a magnification of ×1500 and 5 kV.

Coating and cell fabrication

The electrode ink coating was prepared with an Active Material (AM): Carbon Black (CB): Binder (B) ratio of $80:10:10$ using a Thinky Mixer. The binder – polyvinylidene difluoride (PVDF) – was mixed with N-methyl pyrrolidone (NMP) initially for 5 min/1300 rpm, before additions of the SuperP carbon black and the active material, with subsequent additions of NMP to produce a slurry (10 min/1300 rpm for each step). To degas the mixture, a final mix of 3 min/1800 rpm was performed. The resulting slurry was cast onto copper foil using a draw-down coater, where the bar height was set to 200 μ m. The resulting coating was dried for up to 2 hours at 80 °C before being transferred for overnight drying in a vacuum oven pre-set at 110 °C. The coatings were then calendered between 2 stainless steel

rollers at 80 °C until the porosity of the coated material was approximately between 30–40%.

The Li-half cells were fabricated in an Ar filled glovebox. Initially, circular disks of the coated active material were cut out (12 mm) and weighed outside the glovebox before being transferred inside. Steel 2032 cases were used, with a single 1 mm stainless steel spacer for compression. The lithium metal electrode was prepared from a dispensed fraction of lithium ribbon, where the surface was scratched using a stainless steel spatula (to remove the tarnished surface and to leave a shiny and rough texture) before being cut to size (12.7 mm) ready for assembly. The electrolyte was 1.0 M LiPF₆ in $50:50$ (v/v) ethylene carbonate and dimethyl carbonate – two 50 μ L additions were made during assembly. The glass fibre separator was cut to size (14.3 mm).

Electrochemical testing

The electrochemical performances of the materials were evaluated using a BioLogic BCS805 battery tester, with the cells cycled between 1.0–2.5 V in galvanostatic mode. A lower voltage limit of 1.0 V is used instead of 0.01 V (as applied for graphite electrodes) to avoid the potential for Li dendrite formation, as well excessive side reactions caused by the formation of an SEI layer which will lead to lithium inventory loss. Furthermore, the use of such a larger voltage (2.5–0.01 V) window is undesirable for cell manufacturers, as it would lead to a consequent wider full cell voltage range. All cells had formation cycling applied with a current density of 0.01 A g^{-1} between the two voltage limits. A constant voltage step was applied on lithiation, where the lower voltage limit (1 V) was held for 3 hours or until the current rate decayed to 40% of the value originally applied. A rest of 10 minutes between each cycle was also incorporated. Paper Matrix of an investigation into new NbAc, based anode sollers at 80 V until the Porosity of the constantion or the common paper of the state is likely denoted the state is a set allows a state of the state is the st

The rate studies for these materials were performed asymmetrically such that on lithiation a current density of 0.1 A g^{-1} was consistently applied with the constant voltage step (with 40% current decay limit). The delithiation rate was gradually increased from 0.1 to 4 A g^{-1} with 5 cycles at each step. Asymmetric cycling was introduced to reduce the current rate limitations created by the use of Li metal in an organic electrolyte.²⁰ After the rate study, five further cycles were conducted at 0.1 A g^{-1} . Following the observation of a higher capacity on returning to 0.1 A g^{-1} , a further test with a modified formation cycle was conducted to see if we could improve the initial capacities of these systems. This formation cycle involved 3 cycles at 0.01 A g^{-1} , 5 cycles at 0.1 A g^{-1} and finally 10 cycles at 2 A g^{-1} .

For each measurement, three cells were tested under the same conditions, to deduce the average of the (de)lithiation capacities and the associated error in the values (given in parentheses in the text).

A further lower voltage test was performed for TTMO71515 using a BioLogic BCS805 battery tester, with the cell being cycled between 0.5–2.5 V in galvanostatic mode. 3 formation cycles were done at a current density of 0.01 A g^{-1} for this test.

Long-term cycling for NTMO71515 and TTMO71515 was performed using a BioLogic BCS805 battery tester, with the cell being cycled between 1–2.5 V in galvanostatic mode. 100 cycles at 0.2 A g^{-1} were performed for each cell.

Results and discussion

Both materials $Nb_7Ti_{1.5}Mo_{1.5}O_{25}$ (NTMO71515) and $Ta_7Ti_{1.5}Mo_{1.5}O_{25}$ (TTMO71515) were initially characterised with PXRD. The results showed an XRD pattern similar to the related systems, Nb_9PO_{25} and Nb_9VO_{25} (tetragonal; space group $I\bar{4}$) as shown in Fig. 2. The structures were refined using the structure of Nb_9PO_{25} as a starting model, with the Rietveld refinement fit shown in Fig. 3 for NTMO71515 and the resulting cell parameters given in Table 1.

Fig. 2 PXRD patterns of NTMO71515 (black) and TTMO71515 (red) Co Kα. The peak positions for the isostructural Nb_9PO_{25} phase are also shown (black dashes).

Fig. 3 Observed, calculated and difference XRD profiles for NTMO71515 ($R_{\rm wp}$ – 5.72%, $R_{\rm p}$ – 3.59%) (Cu Ka) from Rietveld refinement.

Table 1 Unit cell parameters of $Nb_7Ti_{1.5}Mo_{1.5}O_{25}$ (NTMO71515) and $Ta_7Ti_{1.5}Mo_{1.5}O_{25}$ (TTMO71515)

Material	a(A)	c(A)	Volume (A^3)	$R_{\rm WD}$ (%)
NTMO71515	15.6879(3)	3.8031(1)	936.00(4)	5.72
TTMO71515	15.6587(3)	3.8043(7)	932.80(3)	9.51

To aid understanding of the doping strategy from the structural perspective, the original crystal model used in this refinement (Nb₉PO₂₅) has been included (Fig. 4). This consists of a (3×3) block structure along with tetrahedral sites, with the 4 different crystallographic sites labelled sites 1–4. Site 1 is the corner-sharing tetrahedra which is occupied by P^{5+} for Nb_9PO_{25} , while sites 2–4 represent different octahedral sites: site 2, the corner-sharing octahedra in the centre of the 3×3 block, and sites 3 and 4 the edge-sharing octahedra.

Due to the presence of multiple cations, including isoelectronic Nb^{5+} and Mo^{6+} , and low X-ray scattering of the oxygen, the refinement was conducted in stages – with the starting point involving all octahedral cation sites (site $2-4$) set to Nb^{5+} , with Mo^{6+} occupying the tetrahedral site (due to the known greater preference of $Mo⁶⁺$ for tetrahedral coordination). Sites which showed low $Nb⁵⁺$ content (site 4) were then assumed to be occupied by some Ti^{4+} . Ti^{4+} was then added to all the octahedral and tetrahedral sites and refined. Due to $Nb⁵⁺$ and $Mo⁶⁺$ being isoelectronic it was not possible to refine individual Nb and Mo occupancies and it is assumed that the $Nb⁵⁺$ octahedral sites also contain some $Mo⁶⁺$ to match the formula of the material. In order, to confirm the preferred occupancy of $Mo⁶⁺$ in the tetrahedral sites, the structure of the equivalent Ta system was also refined (Rietveld refinement profile shown in the ESI (ESI Fig. 1†)). Here the significantly different scattering factors of Mo^{6+} and Ta^{5+} allowed the confirmation of the preference for Mo^{6+} in the tetrahedral site. The structural parameters for both systems are tabulated in the ESI (ESI Tables 1 and 2†). Oalton Transactions Werelaston sections (and the comparison of the comparison of the section of the s

The morphology of both systems was investigated by SEM (ESI Fig. 2 and 3†) and found to exhibit rod-like character.

Raman spectra were recorded for both samples, as shown in Fig. 5, to further evaluate the bonding in these materials. The spectra show that at wavenumbers below 150 cm^{-1} , the peaks for the Ta-analogue are shifted to lower wavenumbers compared to the Nb material, with the most intense Raman

Fig. 4 Crystal structure of the Nb_9PO_{25} system. There are 4 crystallographic sites in the structure. Site 1 is the tetrahedral cation site (orange). Site 2 is the centre octahedra of the 3×3 block (dark blue). Site 3 (blue) and 4 (light blue) are the edge-sharing octahedra.

Fig. 5 Raman spectra of NTMO71515 (black) and TTMO71515 (red).

mode being at 128 cm⁻¹ and 112 cm⁻¹ for the Nb and Ta phases, respectively. At wavenumbers greater than 150 cm^{-1} , the opposite occurs, as the Raman peaks for the Ta-analogue are shifted to higher wavenumbers. Previous work by Perfler et al., comparing TiNb₂O₇ and TiTa₂O₇, observed similar trends.42 By comparison with this past literature, the symmetric metal–oxygen stretching modes for the corner and edge-sharing TaO $_6$ /NbO₆ octahedra can be assigned to the 1008 cm⁻¹ peak (Ta phase), and 992 cm^{-1} peak (Nb phase). The peaks at 932 cm⁻¹ (Ta) and 918 cm⁻¹ (Nb) are likely due to a combination metal–oxygen stretching modes from the TaO_6/NbO_6 octahedra and $MoO₄$ tetrahedra.⁴³ The shifts in the Raman bands to higher wavenumbers for the Ta phase can be attributed to the higher bond strengths and force constants of the Ta– O bonds compared to the Nb–O bonds, with the vibrational modes predominantly involving vibrations of the O atoms, with the metal at the centre of the octahedra not moving. 42 For the lower wavenumber region (below 150 cm−¹) the bands can be attributed to O–(Nb/Ta)–O symmetric and antisymmetric bending vibrations, that predominantly involve vibrations of the Nb/Ta atoms (the increase in mass for Ta at the centre of the $MO₆$ octahedra leading to a shift to lower wavenumber).⁴⁴ Similar to this previous study comparing TiNb₂O₇ and TiTa₂O₇, the bands situated between the wavenumbers 550–700 cm−¹ can be attributed to the Ti–O stretching modes. 42

Electrochemical performance

The compounds have 3 redox active elements Nb/Ta, Ti, Mo where Nb^{5+}/Ta^{5+} might be expected to reduce to $+4/+3$, Ti^{4+} to

 Ti^{3+} and Mo⁶⁺ to Mo⁴⁺. Assuming complete reduction to Nb³⁺/ Ta^{3+} , Ti^{3+} , Mo^{4+} , 18.5 Li would be intercalated into the systems which would give theoretical capacities for NTMO71515 $(RMM: 1266 g mol⁻¹)$ and TTMO71515 $(RMM: 1882 g mol⁻¹)$ of 391 and 263 mA h g^{-1} , respectively. The charge-discharge voltage profiles for NTMO71515 and TTMO71515 on formation cycling (at current density of 0.01 A g^{-1}) are presented in Fig. 6. The initial discharge capacity for NTMO71515 was found to be 268(9) mA h g^{-1} which correlates to *ca*. 12.7 moles of electrons (lithium insertion) in the process, with the lower capacity attributed with only partial reduction of Nb to 3+, similarly to observed previously for other Wadsley Roth phases. The second cycle shows a 37 mA h g^{-1} capacity loss $(1.7 \text{ Li}^+$ irreversibly trapped) with a returned discharge capacity of 231(17) mA h g^{-1} . For the TTMO71515 system the capacity for the first discharge was found to be 140(3) mA h g^{-1} corresponding to lower level of Li incorporation under the same conditions, ca. 9.8 moles of electrons (lithium insertion), suggesting a significantly reduced level of Ta reduction compared to Nb. The second cycle shows a discharge capacity of 98 (4) mA h g^{-1} , corresponding to a loss of 2.9 Li⁺, which is greater than observed for NTMO71515. A breakdown of all the capacities associated with different voltage windows for all 3 formation cycles is provided in the ESI (ESI Tables 3 and 4†) along with voltage profile comparisons between NTMO71515 and TTMO71515 (ESI Fig. 4–6†). **Paper** Context Common Co

To understand the chemistry and the redox couples at play during (de)lithiation, differential scanning calorimetry (dQ/dV) data were plotted (Fig. 7). For the NTMO71515 system, three reversible peaks and one irreversible peak can be observed in the dQ/dV plot. The irreversible peak located at 2.20 V is attributed to the Mo reduction process in the tetrahedral sites. The irreversible redox contributes to a drop in capacity, which is attributed to Li being trapped by this irreversible Mo reduction. The isostructural phase, $Nb₉VO₂₅$, has a similar significant decrease in capacity due to the irreversible reduction of V in the tetrahedral sites.^{45,46} At 1.90 V a reversible peak is observed corresponding to the overlapping redox couples of $Mo^{6+}-Mo^{4+}$ (oct sites), $Ti^{4+}-Ti^{3+}$ and $Nb^{5+}-Nb^{4+}$ and the peak at 1.42 V can be attributed as the partial $Nb^{4+}-Nb^{3+}$ redox peak.^{20,47-49} The dQ/dV plot for NTMO71515 shows no clear shifting of the peaks between the first and third cycle. However, the intensities do decrease slightly between the first

Fig. 6 Galvanostatic discharge–charge curves of NTMO71515 (a) and TTMO71515 (b), vs. Li metal, with a current density of 0.01 A g^{-1} applied.

Fig. 7 Differential scanning calorimetry (dQ/dV) plot derived from the galvanostatic discharge/charge profile of NTMO71515 (a) and TTMO71515 (b), vs. Li metal at a rate corresponding to 0.01 A g^{-1} .

and second cycle highlighting the loss of Li during the first cycle.²⁰ ESI Fig. 9† shows a large decrease in peak intensities and shifting of the peaks in the dQ/dV plot between the formation cycles at 0.01 A g^{-1} and the start of the rate test at 0.1 A g^{-1} , which correlates with the further significant decrease in capacity we observe between these 2 rates.

In order to provide further support for the above assignments, the dQ/dV plots for the Ta-analogue were measured, which show similarities to NTMO71515. For TTMO71515 three reversible and one unreversible redox peak can be observed. Like NTMO71515 the irreversible redox peak at 1.94 V is assigned to the tetrahedral Mo site redox process (shifted to lower V for the Ta analogue) whilst the $Mo^{6+}-Mo^{4+}$ (oct site) shows one peak at 1.62 V.^{50,51} The dQ/dV plot shows a reversible peak at 1.42 V which is attributed as the Ti^{4+} – Ti^{3+} redox process. The third reversible peak at 1.12 V is attributed to the Ta^{5+} –Ta⁴⁺ partial redox process. Comparisons between the dQ / dV plots for NTMO71515 and TTMO71515 can be seen in ESI Fig. 7-9.† The dQ/dV plot for TTMO71515 shows a similar decrease in peak intensities to the NTMO71515 system which we also accredit to the irreversible loss in Li between the first and second cycle due to Li trapping. There is a clear shift in all the redox peak positions to lower voltages compared to the Nb analogue, illustrating a significant influence on the overall redox processes on changing from Nb to Ta. Given the observed shift in the redox peaks to lower voltages for the Ta analogue, a further analysis of the Ta sample was performed to a lower voltage, 0.5–2.5 V, at a current density of 0.01 A g^{-1} for 3 cycles (ESI Fig. 10†). A breakdown of all the capacities associated with different voltage windows for all 3 cycles is provided in the ESI (ESI Table 5†). This study confirmed that there is Ta redox below 1 V as the capacity increased from 140(3) mA h g^{-1} when the voltage limit was 1 V to 225 mA h g^{-1} for the 0.5 V cut-off. The resulting dQ/dV plot also highlights further Ta reversible redox around 1 V (ESI Fig. 11†).

To test the electrochemical rate performance and capacity retention of NTMO71515 and TTMO71515, additional measurements were performed at varying current densities (0.1–4 A g^{-1}) using asymmetric cycling conditions (1–2.5 V window), where the lithiation rate was kept constant at 0.1 A g^{-1} whilst the delithiation rate was gradually increased. The resulting gravimetric capacity at each rate for NTMO71515 is

shown in Fig. 8. While there is an initial drop in capacity on cycling from 0.01 to 0.1 A g^{-1} (values of 220(19) and 151(12) mA h g^{-1} obtained respectively), subsequent retention at higher rates is good, with the average capacity found to be 132 (10) and 115(14) mA h g^{-1} , for the rates of 2 A g^{-1} and 4 A g^{-1} respectively (corresponding to ca. 7.5C and 15C respectively). The capacity retention over the 5 cycles for each rate does not drop below 98% and there is only a 13% loss in capacity between current densities of 0.1 and 2 A g^{-1} (151(12)–132(10) mA h g⁻¹). Even between 0.1 and 4 A g⁻¹ (151(12)-115(14) mA h g−¹) there is only a 27% decrease in capacity highlighting the good rate capability of this system. ESI Table 6† shows the average capacity retention at different current densities and the associated C-rate for NTMO71515. Upon completing the rate study with a final 5 cycles at 0.1 A g^{-1} , interestingly the capacity recovered to 160(11) mA h g^{-1} , which is higher than its original value at the start of the rate study (151(12) A h g^{-1}) – thus suggesting the original formation process needs to be further optimised in these systems, and thus may offer further potential to enhance performance, and that the high delithiation rates applied did not negatively impact the material. Long-term cycling was performed on this NTMO71515 material which confirmed good capacity retention on extended cycling (ESI Fig. 12†). For this study, 100 cycles at a current density of **Calco Transactions**
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Fig. 8 Average specific capacities of three NTMO71515 cells undergoing asymmetric cycling – such that the lithiation current density is maintained at 0.1 A g^{-1} , while the delithiation gradually increases after 5 cycles at each rate step. The red error bars are for lithiation, the black ones are for the delithiation.

0.2 A g^{-1} were performed and a 97% retention in capacity was delivered emphasising the good cycle-life of this system.

To investigate the increase in capacity at lower current density rates when fast cycling is performed, we prepared additional cells where we added extra formation cycling at higher rates – 3 cycles at 0.01 A $\mathrm{g}^{-1},$ 5 cycles at 0.1 A g^{-1} and finally 10 cycles at 2 A g^{-1} . This revised formation cycling was tested, and the results can be seen in ESI Fig. 15.† The results show that the capacity of the material at 0.1 A g^{-1} can be slightly enhanced from 149(1) mA h g⁻¹ to 156(1) mA h g⁻¹ if a faster rate $(2 \text{ A } g^{-1})$ is applied in the formation cycle. This increase in capacity suggests that further study of formation protocols is warranted for Wadsley Roth phases, such as these.

TTMO71515 was also shown to display good capacity retention, after the initial decrease seen for the current densities between 0.01 and 0.1 A g^{-1} (95(2)–79(3) mA h g^{-1}), corresponding to a 18% reduction in capacity. Subsequently the capacity only decreased from 79(2) mA h g^{-1} to 60(1) mA h g^{-1} for the wide range of current densities from 0.1–4 A g^{-1} (27% drop in capacity) with the capacity retention over 5 cycles at each rate staying above 99% (Fig. 9). ESI Table 7† shows the average capacity retention at different current densities and the associated C-rate for TTMO71515. However, the Ta-analogue exhibits lower gravimetric capacities, due to its high relative molecular mass and lack of Ta redox activity above 1 V – thus making it a poorer anode material compared to the Nb analogue in the range 2.5–1.0 V. Moreover Ta is significantly higher in cost than Nb, which is another negative attribute. Long-term cycling was also performed for this system for 100 cycles at 0.2 A g^{-1} . The capacity dropped from 89 mA h g^{-1} to 86 mA h g−¹ over 100 cycles showcasing a similar 97% capacity retention observed as for NTMO71515 (ESI Fig. 13†). Paper Matter published on 31% where the published on 1978 and 1978 a

Investigation of solid solution range for the Nb–Ti–Mo–O system

In order to investigate whether these were line phases, or exhibited a solid solution range, the Nb system was investigated in more detail, with varying Ti/Mo contents: $Nb_{10-2x}Ti_xMo_xO₂₅$, x varied between 1.25 and 2. These experiments showed single

Fig. 9 Average specific capacities of three TTMO71515 cells undergoing asymmetric cycling – such that the lithiation current density is maintained at 0.1 A g $^{-1}$, while the delithiation gradually increases after 5 cycles at each rate step. The red error bars are for lithiation, black are for the delithiation.

phase samples were observed for $1.5 \le x \le 1.75$, indicating a narrow solid solution range, with impurities observed for higher and lower Ti/Mo contents (see Fig. 10, with cell parameters given in Table 2), albeit the impurities were small for $x =$ 2 (ESI Fig. 14†).

The electrochemical performance of the $x = 1.75$ endmember of the solid solution series was also investigated (Fig. 11). The electrochemical performance showed a higher initial capacity (290(4) mA h g^{-1}) along with an accompanying higher first cycle loss (19%) than NTMO71515, with a decrease to 234(4) mA h g^{-1} in the second cycle, which is most likely related to the higher Mo content. After the initial losses, NTMO65175175 has good capacity retention for rates between 0.1–2 A g^{-1} with the capacity retention over 5 cycles at each rate staying above 99%. However, at faster rates $(4 \text{ A } g^{-1})$ the reten-

Fig. 10 XRD patterns of the solid solution $Nb_{10-2x}Ti_xMo_xO₂₅$ (1.25 ≤ x ≤ 2). Nb_9PO_{25} phase tick marks (black dashes) are also shown.

Table 2 Unit cell parameters of NTMO71515 and NTMO65175175

Material	a(A)	c(A)	Volume (A^3)
NTMO71515	15.6879(3)	3.8031(1)	936.00(4)
NTMO65175175	15.6853(2)	3.8015(3)	935.28(3)

Fig. 11 Average specific capacities of two NTMO65175175 cells undergoing asymmetric cycling – such that the lithiation current density is maintained at 0.1 A g^{-1} , while the delithiation gradually increases after 5 cycles at each rate step. The red error bars are for lithiation, black are for the delithiation.

tion falls off to 89%, with the capacity fading from 118(2) to 106 (1) mA h g^{-1} . Upon completing the rate study with a final 5 cycles at 0.1 A $\rm g^{-1}$, the lithiation capacity recovered to 158(7) mA h g⁻¹ compared to 170(5) mA h g⁻¹ for the first set of cycles at 0.1 A g^{-1} . This decrease in capacity suggests this higher Mo content material does not recover as well as NTMO71515 after being cycled at higher current densities. ESI Table 8† shows the average capacity retention at different current densities and the associated C-rate for NTMO65175175.

Conclusions

In conclusion, we have reported the synthesis and characterisation of the new Wadsley Roth phases $Nb_7Ti_{1.5}Mo_{1.5}O_{25}$ (NTMO71515) and Ta₇Ti_{1.5}Mo_{1.5}O₂₅ (TTMO71515), which are isostructural with Nb_9PO_{25}/Nb_9VO_{25} . From electrochemical testing within a Li-half coin cell, NTMO71515 shows high capacity for first cycle of 268(9) mA h g^{-1} (2.5–1.0 V), which decreases significantly over the initial few cycles, attributed to Li trapping mediated by irreversible reduction of Mo in the tetrahedral sites (similar to prior results for $Nb₉VO₂₅$). Despite the initial capacity lost, this material shows subsequent good capacity retention at increasing rates between 0.1 and 2 A g^{-1} with the capacity only dropping from 152 mA h g^{-1} to 133 mA h g^{-1} over this range– equivalent to a 13% drop. Furthermore, the capacity recovers to above its original values when the lower current density 0.1 A g^{-1} is re-applied, suggesting further optimisation is required for the formation cycling to maximise values, while also showing that the material is unaffected by high rate cycling. Further studies showed that this phase can also be prepared containing higher Ti/Mo contents, with a solid solution range observed; $Nb_{10-2x}Ti_xMo_xO_{25}$ (1.5 ≤ $x \le 1.75$). The Ta analogue phase, TTMO71515, showed lower capacities, attributed both to the heavier mass and the reduced Ta redox in the 1.0–2.5 V voltage window. However, the comparison of the data for this phase and NTMO71515 allowed confirmation of the Mo preference for the tetrahedral site and helped in the assignment of the redox couples at play in both materials. **Open Access Articles** Articles. The morphological state and the state of 2023. Downloaded in the more and the more state and the state of the

Author contributions

Alex J. Green: Investigation, data curation, formal analysis, visualization, writing – original draft, writing – review and editing. Elizabeth H. Driscoll: Methodology, visualization, writing – original draft, and writing– review and editing. Yazid Lakhdar: Methodology – review and editing. Emma Kendrick: Supervision, resources, and writing – review and editing. Peter R. Slater: Conceptualization, methodology, supervision, resources, writing – original draft, and writing – review and editing.

Conflicts of interest

There are no conflicts to declare.

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