

Highlighting a study on the battery cycling and atomic structure stability of the metastable 'R' polymorph of Nb₂O₅ from Professor Megan M. Butala's laboratory in the Department of Materials Science and Engineering at the University of Florida (United States).

R-Nb₂O₅ has an 'idealized' V₂O₅ structure and Wadsley-Roth-like structural stability during Li-ion battery cycling

With Wadsley-Roth-like 'slabs' and an 'idealized' V_2O_5 structure, metastable R-Nb₂O₅ bridges two well-studied families of Li-ion battery electrodes. Inter- and intrapolyhedral connectivity stabilize the structure against polyhedral rotations during lithiation and delithiation, providing excellent reversibility with negligible structure changes during cycling.



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1 Introduction

Early transition metal oxides featuring crystallographic shear planes are a technologically and fundamentally compelling family of Li-ion battery electrode candidates. These materials, sometimes referred to as Wadsley–Roth or Magnéli phases,¹⁻⁵ have demonstrated excellent performance as high potential anodes. With operating potentials between 1 V and 1.5 V and excellent rate capabilities, they offer a pathway to safe, highpower Li-ion batteries as alternatives to graphite and $Li_4Ti_5O_{12}$.⁶⁻⁸ Wadsley–Roth (WR) materials can also have multielectron redox, in which more than one redox reaction occurs per redox active transition metal, contributing to high energy densities, *e.g.*, reduction of Nb⁵⁺ to Nb³⁺, allowing for storage of >1 mol Li per transition metal, M.^{5,9-12}

In addition to being technologically relevant, WR materials provide a foundation for identifying atomic and electronic structure features that support effective transport of ions and electrons.^{9,13,14} WR materials have ReO₃-like blocks, in which [MO₆] octahedra are corner-sharing (Fig. 1a and c) with square

R-Nb₂O₅ has an 'idealized' V₂O₅ structure and Wadsley–Roth-like structural stability during Li-ion battery cycling[†]

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The adoption of batteries across diverse applications requires electrode materials with a wider range of performance metrics, such as cost, safety, and material availability. Along the path to discovering new commercially viable materials, a fundamental understanding of chemical and atomic structure features that provide structural stability and effective ion transport is essential. In support of new understanding, we report the cycling behavior of metastable $R-Nb_2O_5$. $R-Nb_2O_5$ adopts an 'idealized' V_2O_5 structure, in which $[NbO_6]$ octahedra alternate in edge- and corner-sharing resulting in ReO_3 -like slabs, whereas Wadsley–Roth materials have ReO_3 -like blocks, linked through edge-sharing octahedra at intersecting crystallographic shear planes. We find that this slab structure is stable during cycling, with minor atomic structure changes and cycling curves that are symmetric on discharge and charge, resembling the behavior of Wadsley–Roth materials more than other related materials, such as ReO_3 , V_2O_5 , or Nb_3O_7F . Based on our findings, $R-Nb_2O_5$ can serve as a 'structural bridge' between Wadsley–Roth block structures and V_2O_5 , through which we can relate inter- and intra-polyhedral structures to cycling behavior and structural stability during cycling.

channels that provide fast Li transport. These blocks are connected across crystallographic shear planes, at which [MO₆] octahedra are edge-sharing (Fig. 1b and c). During lithiation of ReO₃, [ReO₆] octahedra rotate around corner-sharing vertices.^{4,15} In contrast, in WR materials, the edge-sharing octahedra at crystallographic shear planes stabilize against this rotation, keeping square channels accessible for fast Li⁺ transport.^{9,11} In addition to structural stability, edge-sharing octahedra have a favorable overlap of d-electron orbitals, which results in high electronic conductivity upon even minor reduction, contributing to their high-rate capabilities.^{11,16,17}

Among WR and related materials studied as battery electrodes, there are many niobates,^{16,18} including those with the chemical formula Nb₂O₅.^{17,19-24} We report here, for the first time, the cycling behavior and charge storage mechanism of yet another polymorph *R*-Nb₂O₅. '*R*' represents a "neutral designation",^{19,25,26} and is not related to the crystal system or preparation conditions. This metastable polymorph adopts an idealized V₂O₅ structure (Fig. 1f and 2)² and was first identified by Gruehn.²⁵ *R*-Nb₂O₅ can be described relative to WR block structures as having ReO₃-like slabs (rather than blocks) connected by parallel (rather than perpendicular) shear planes (Fig. 2a).^{2,27}

WR block structures can be described by the connectivity of $[MO_6]$ octahedra, specifically in terms of the number of cornersharing octahedra in a given direction. For example, a block that is *m* corner-sharing octahedra wide, *n* high, and infinite in the third direction would be represented as $(m \times n) \propto .^3$ For *H*-Nb₂O₅, this representation would be $(3 \times 5)_{\infty}$ (Fig. 1c).

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[†] Electronic supplementary information (ESI) available: Additional details and results from Rietveld refinements of atomic structure models as well as cycling data in Swagelok cells, representative of cycling from which *ex situ* discharged and charged powders were prepared, and coin cells cycled at C/10 and 1C. See DOI: https://doi.org/10.1039/d2ta08653k

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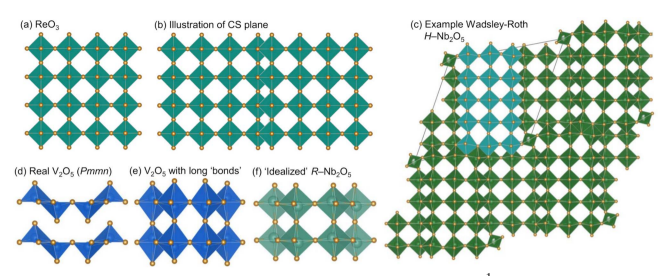


Fig. 1 (a) ReO₃ has only corner-sharing [MO₆] octahedra. (b) Adjacent ReO₃-like blocks offset by $\frac{1}{2}$ an octahedron in 2 directions make a crystallographic shear (CS) plane with edge-sharing [MO₆] octahedra. (c) Wadsley–Roth structures are comprised of ReO₃-like blocks connected by intersecting CS planes; *H*-Nb₂O₅ is shown as an example. (d) V₂O₅ has layers of edge- and corner-sharing [VO₅] square pyramids. (e) Connecting the long V–O distance across the layers, a relationship between (d) real and (f) 'idealized' V₂O₅ structures can be visualized. (f) Metastable polymorph *R*-Nb₂O₅ adopts an idealized V₂O₅ structure with edge- and corner-sharing octahedra.

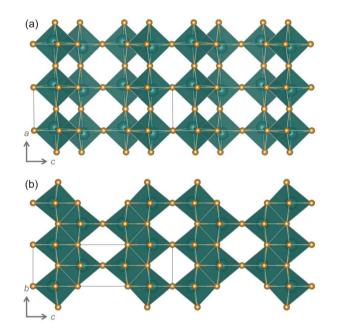


Fig. 2 Visualizations of R-Nb₂O₅ based on the original report (ICSD collection code 25765)²⁵ viewed along the (a) [010] and (b) [100] directions.

Borrowing the WR structure notation,³ the structure of *R*-Nb₂O₅ can be represented as $(2 \times \infty)_{\infty}$.

Both α -V₂O₅ and ReO₃ can be considered as 'parent structures' for *R*-Nb₂O₅.^{27,28} Like ReO₃, V₂O₅ also undergoes polyhedral rotations during lithiation that result in extensive atomic structure changes.^{29,30} V₂O₅ has layers composed of edge- and cornersharing [VO₅] square pyramids (Fig. 1d). Across the layers, there is a long V–O distance (≈ 2.8 Å); when a 'bond' is drawn between these atoms, V₂O₅ can be visualized as a three-dimensionally-connected structure (Fig. 1e). The larger ionic radius of Nb

relative to that of V results in the octahedral, rather than squarepyramidal, coordination of Nb in *R*-Nb₂O₅, approaching a relatively idealized structure.^{2,31} Given the different atomic structure evolutions and stabilizations of ReO₃ and WR block structures, we analogously find that *R*-Nb₂O₅ has improved structural stability upon lithiation relative to V₂O₅. In addition to being stable against polyhedral rotation during lithiation, we also find that *R*-Nb₂O₅ has minimal atomic structure evolution during cycling, akin to the relatively small changes in WR materials, for example, unit cell volume changes $\leq 6\%$.^{10,11}

The 'ideality' of *R*-Nb₂O₅ relative to that of α -V₂O₅ can be illustrated by comparing their distributions of M–O distances. For *R*-Nb₂O₅, they range from 1.7 Å to 2.3 Å (this work); this is narrower than the range for V₂O₅, which is from 1.58 Å to 2.8 Å,³² and broader than that for ReO₃, in which there is a single Re–O bond length of 1.9 Å.³³ Nb₃O₇F is another slab structure, denoted as $(3 \times \infty)_{\infty}$, that has also been studied as a Li-ion electrode¹⁵ and has a similar distribution as *R*-Nb₂O₅, from 1.85 Å to 2.2 Å.³⁴

Overall, we find that the cycling behavior and structural stability of R-Nb₂O₅ are more similar to WRs than to ReO₃,^{4,35} α -V₂O₅,²⁹ or Nb₃O₇F.¹⁵ R-Nb₂O₅ has symmetric discharge and charge profiles that do not significantly change between the first and later cycles. In addition, atomic structure changes are minimal, with no evidence of the polyhedral rotation that ReO₃ or V₂O₅ undergo during lithiation. We observed unit cell changes of less than 5% and a reduced off-centering of Nb in cycled products.⁹⁻¹¹

2 Results and discussion

2.1 Synthesis and characterization of *R*-Nb₂O₅

The *R* polymorph of Nb₂O₅ was prepared through soft chemical methods, specifically, through the drying and decomposition of

hydrated precursor $H_3ONb_3O_8$.³⁶ $H_3ONb_3O_8$ was produced from KNb_3O_8 , a layered niobate, through ion-exchange in nitric acid.^{36–38} KNb_3O_8 was prepared using molten salt synthesis in an excess of KCl, which facilitates the mass transport among precursors to form a phase pure product.^{18,39}

Bragg peak positions of the prepared *R*-Nb₂O₅ are consistent with the structure originally reported by Gruehn,²⁵ which is evident in neutron diffraction data (POWGEN at Oak Ridge National Laboratory, Fig. 3) as well as high-resolution synchrotron X-ray diffraction (XRD) data (11-BM at Argonne National Laboratory, Fig. S1[†]). In both types of diffraction data, peak broadening is evident. Scanning electron microscope (SEM) images show anisotropic particles with sub-micron features in two dimensions and one longer axis in the order of microns (Fig. 4c and d). This indicates that finite size effects give rise to the reflection-dependent broadening in diffraction data.

Gruehn's reported *R*-Nb₂O₅ structure was used as a starting model for refinements of the atomic structure model against neutron and X-ray diffraction data [International Crystal Structure Database (ICSD) collection code 25765].²⁵ Rietveld refinement against neutron diffraction data resulted in minor changes to lattice parameters and atom positions, resulting in a fit that captures the data well (Fig. 3). Similar parameters were found from the refinement of synchrotron XRD data (Fig. S1†). Details of refinement parameters and the resulting structures are provided in Tables S1 and S2.†

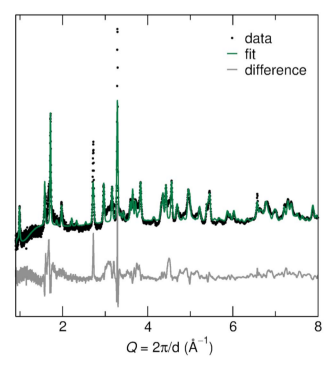


Fig. 3 Rietveld refinement against neutron diffraction data for the prepared R-Nb₂O₅ shows good agreement between the measured and calculated patterns. The largest differences occur in regions with diffuse scattering, as well as for two reflections that are underestimated in the calculated data. Details of the refined structure parameters are provided in the ESI.†

The differences between the measured data and calculated fit have two primary origins: diffuse scattering near Q of 1.5 Å⁻¹ and 3 Å⁻¹, and peak intensity mismatch near Q of 2.9 Å⁻¹ and 3.25 Å⁻¹ (Fig. 3). The diffuse scattering features suggest low crystallinity, disorder, or a combination. Even after accounting for anisotropic particle shape in refinements, there is a remaining intensity mismatch for the (113), (113), and (020) reflections, which are especially narrow and accordingly must be associated with the long axis of the particles. Similar reflection-dependent broadening has also been identified for KNb₃O₈,¹⁸ the precursor from which this *R*-Nb₂O₅ was prepared (Fig. 4a).

The elongated rectangular morphology of R-Nb₂O₅ comes from the morphology of the KNb₃O₈ and H₃ONb₃O₈ precursors (Fig. 4a and b). The layered nature of the KNb₃O₈ atomic structure drives preferential growth along specific crystallographic directions.¹⁸ The underlying particle shape is retained through ion-exchange and low temperature heating of the prepared product. This synthetic route did not allow for the preparation of larger or more isotropic R-Nb₂O₅ particles, which was beyond the scope of this work, but variations of particle size and shape may be accessible from more isotropic KNb₃O₈ precursors or through alternative synthetic methods.

A notable feature of the crystal structure of R-Nb₂O₅ is the off-centering of Nb within the octahedra, as well as the pattern of displacements from the center between neighboring octahedra. In V₂O₅, the nearest neighbor V–O distances range from 1.58 Å to 2.8 Å, and off-centering of V⁵⁺ away from the center of an octahedron occurs as pairs (two displaced lower, two displaced higher, Fig. 1d).³² In R-Nb₂O₅, Nb is also off-centered in the octahedra, but with a narrower distribution of Nb–O distances, ranging from 1.7 Å to 2.3 Å. In addition, off-centering occurs in a zig-zag pattern, alternating up and down for each octahedron, along the *c* direction (Fig. 2a).

2.2 Battery cycling behavior

R-Nb₂O₅ was cycled against Li metal in coin cells at various rates. At a relatively slow cycling rate of C/20 (based on the reaction of 2 mol Li per mol Nb₂O₅ in 20 h), galvanostatic cycling shows a plateau at \approx 1.8 V for $0 \leq x \leq$ 1, and a sloping region at low potentials (Fig. 5a), suggestive of two-phase and solid solution lithiation mechanisms at high and low potentials, respectively. The plateau for *R*-Nb₂O₅ is slightly higher than that for *H*-Nb₂O₅, which is at \approx 1.6 V.²⁰ Over the first \approx 55 cycles, unoptimized coin cells show gradual capacity fade (Fig. 5a).

Between the first and second discharge of *R*-Nb₂O₅, there are minor differences in the cycling profiles. Specifically, there is additional capacity before the plateau in the first discharge compared to later discharges. This difference is likely related to the irreversible capacity over the first cycle, with ≈ 0.3 mol Li per mol Nb₂O₅ remaining at the end of the first charge. This type of slight irreversibility is common in WR and related materials,¹⁸ and has been previously attributed to sluggish diffusion of Li⁺ from specific sites within WR-type structures, as well as an electronic transition from an initially electronic

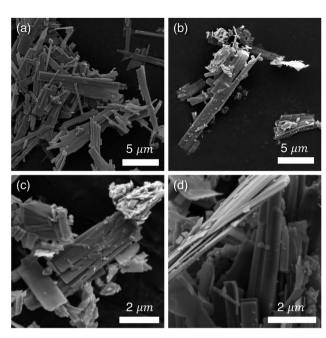


Fig. 4 SEM images of (a) precursor KNb_3O_8 , (b) intermediate product of ion-exchange, $H_3ONb_3O_8$, and (c, d) $R-Nb_2O_5$. The elongated, rectangular particles of $R-Nb_2O_5$ result from its precursors. The anisotropic particle size has crystallographic origins and results in the reflection-character-dependent broadening in the diffraction data (Fig. 3).

insulator to having delocalized electron states upon slight Nb reduction.^{10,11,17} Additionally, the differential capacity plot highlights minor differences in the shape of the first discharge curve relative to that of later cycles (Fig. 5). Even so, discharge and charge curves are generally symmetric, indicating similar reaction pathways on discharge and charge. This is evident in the positions of plateaus and sloped regions in galvanostatic cycling (Fig. 5a), and the corresponding peaks and diffuse features in the differential capacity plot (Fig. 5b). The symmetric cycling processes on discharge and charge are in contrast to those of the 'parent' compounds of R-Nb₂O₅, ReO₃ and V₂O₅, in which extensive and irreversible atomic structure changes during the first lithiation result in significant differences in the profile shape of the first discharge and following cycles.^{29,30,35}

At faster cycling rates, electrochemical curves show good reversibility and similar electrochemical features to those after cycling at slower rates, as well as lower capacities and higher overpotentials. Changes in cycling features with increasing rate are evident in Fig. 6a, which shows the second cycle at each rate from variable rate galvanostatic cycling. The increased overpotential and more sloped nature of the cycling curves with increasing rate are highlighted in the differential capacity plot (Fig. 6b). R-Nb₂O₅ has excellent reversibility and good capacity retention, visible in cycling at C/10 and 1C in the ESI† (Fig. S3 and S4).

The decrease in capacity with increasing rate is more significant for R-Nb₂O₅ than for typical block Wadsley–Roth materials. The reversible capacity of ≈ 150 mA h g⁻¹ at C/20 reduces to about 50 mA h g⁻¹ at 5C (Fig. 6c), whereas typical WR block materials, *e.g.*, TiNb₂O₇ and PNb₉O₂₅, maintain more capacity with proportional increases in cycling rates.^{11,40} While

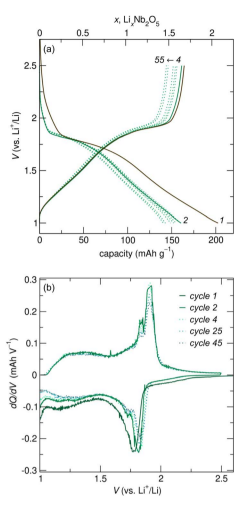


Fig. 5 (a) Galvanostatic cycling for coin cells cycled against Li metal at a rate of C/20. (a) A combination of plateaus and slopes in cycling profiles indicates two-phase and solid solution lithiation mechanisms, (b) exhibited as peaks and diffuse features in the differential capacity plot. Both plots indicate the symmetry of discharge and charge features, and their stability with cycling. In (a), the first and second cycles are represented with solid lines, and every tenth cycle following is represented by dashed lines, as numbered.

the capacities and rate capabilities of R-Nb₂O₅ do not meet those of typical WR block materials, they do exceed the reversibility and capacity retention of other related materials, such as ReO₃,^{4,35} and even 'slab' material Nb₃O₇F.¹⁵

Considering that R-Nb₂O₅ has an 'idealized' V₂O₅ structure, there is a significant difference between their electrochemical behaviors during lithiation. The asymmetry observed in the charge and discharge profiles for α -V₂O₅ during lithiation, due to phase transformations into ε -, δ -, and γ - polymorphs, is significantly different than that in the highly symmetric profiles we find for R-Nb₂O₅.^{29,41} The origins of these differences are discussed further in the following section.

2.3 Structural evolution with cycling

To understand the structural stability and cycling mechanism of R-Nb₂O₅, we analyzed *ex situ* synchrotron XRD data after the

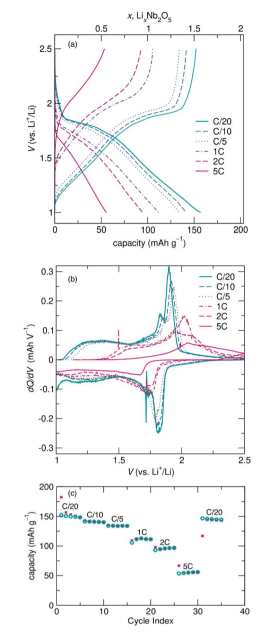


Fig. 6 (a) Galvanostatic cycling profiles and (b) differential capacity plots for coin cells cycled against Li metal at variable rates from C/20 to 5C. In (a), profiles are shown for the second cycle at each rate. (c) Capacity per cycle is shown for each rate, including the capacity recovered at C/20.

first discharge and the first charge (Fig. 7). For this, loose powder cathodes were cycled in Swagelok cells at C/20 and extracted after the first discharge (to 0.5 V, $\text{Li}_3\text{Nb}_2\text{O}_5$) and first charge (to 2.5 V, LiNb_2O_5). Cycling for loose powder Swagelok cells is qualitatively similar to that from coin cells (Fig. S2†). The deeper discharge relative to that of coin cells resulted in a higher first discharge capacity, but a similar reversible capacity, which suggests that the low potential capacity is associated with the formation of a solid-electrolyte interphase, storage by a conductive carbon additive, or a combination thereof.^{42,43}

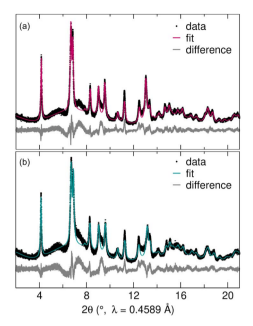


Fig. 7 Rietveld refinements of high-resolution synchrotron XRD data following (a) the first discharge and (b) the first charge of R-Nb₂O₅ indicate agreement between the measured and calculated patterns. Due to the poor sensitivity of X-rays to light elements, Li was excluded from the structural model and only the Nb–O framework was refined (Table 1).

Synchrotron XRD data of cycling products were used in Rietveld refinements of atomic structure models to determine the nature of these minor structural changes. For both states of charge, the R-Nb₂O₅ structure refined against neutron diffraction data of the prepared powder was used as the starting model. Due to the poor sensitivity of X-rays to light elements, Li was not included in the structural model, with an emphasis instead on the Nb–O framework. As in the pristine R-Nb₂O₅, reflection-dependant broadening was captured using isotropic crystallite size parameters, with a mix of Lorentzian and Gaussian shapes, and the preferred orientation (modeled with spherical harmonics) to account for the broadening caused by the anisotropic crystallite size. Additional information on Rietveld refinements and refined structures are provided in the ESI† (Tables S1, S2, S3, and S4).

There were no new reflections in the first discharge and charge products compared to pristine R-Nb₂O₅ (Fig. 3 and 7). Rather, changes in lattice parameters resulted in shifted peak positions, with the Nb–O framework retained through Li⁺ intercalation and deintercalation. As in the pristine R-Nb₂O₅, XRD data of cycled products have diffuse scattering and broad peaks. The qualitative similarity of diffraction data for the pristine and cycled products is reflected in the quantitative results of Rietveld refinements (Table 1 and Fig. 8). Lattice parameters undergo minor contraction (*a* and *c*) and expansion (*b*), resulting in 4.66% and 3.56% increases in the unit cell volume (V_{uc}) of the discharge and charge products, respectively, relative to those of pristine *R*-Nb₂O₅.

Table 1 Refined unit cell parameters for pristine, discharged, and charged R-Nb₂O₅ reflect the minimal changes that result from lithiation and delithiation. In addition, changes in the size and degree of distortion of octahedra over the three states show a decrease in distortion as Nb is reduced

Composition	Pristine Nb ₂ O ₅	Discharged Li ₃ Nb ₂ O ₅	Charged LiNb ₂ O ₂
a	3.9752(6) Å	3.94788(19) Å	3.9313(3) Å
b	3.8233(3) Å	4.04267(15) Å	4.0058(2) Å
с	12.7064(19) Å	12.6633(7) Å	12.6987(10) Å
β	90.603(13)°	90.0034(10)°	89.966(11)°
V _{uc}	193.11(4)Å ³	202.107(17)Å ³	199.98(2)Å ³
% Change $V_{\rm uc}^{\ a}$	_ ()	4.66%	3.56%
Octahedral volume	10.437 Å^3	10.81 \AA^3	10.6955 Å^3
Octahedral distortion index ^b	0.073	0.024	0.029

^a Percent change relative to that of the pristine one. ^b Baur's distortion index (based on bond length variation)⁴⁴ calculated with VESTA.⁴⁵

While lattice parameters are similar for pristine and cycled *R*-Nb₂O₅, there are notable differences in the [NbO₆] octahedra for the pristine and cycled products (Fig. 8). As mentioned, Nb in *R*-Nb₂O₅ is less off-centered than V in V₂O₅ (Fig. 1d and f), but still

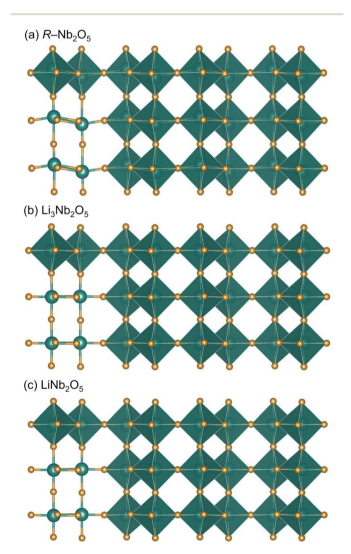


Fig. 8 Refined structures of (a) R-Nb₂O₅, and (b) lithiated product of the first discharge to 0.5 V, and (c) the following first charge to 2.5 V.

has a range of Nb-O bond lengths. Baur's distortion index is a single-value quantity that can be used to describe deviations from a perfect octahedron, with 6 M-O bonds of the same length, which would have a Baur's index of 0.44 Specifically, Baur's index represents octahedral (or tetrahedral) distortions as the deviation of bond lengths from an average bond length.44 In the cycled products, the reduction in Baur's index reflects the decrease in Nb off-centering with the reduction of Nb⁵⁺ during lithiation (Fig. 8). In pristine Nb₂O₅, Nb⁵⁺ has an electronic configuration $[Kr]5s^24d^0$; at the *ex situ* states of charge evaluated, there is at least some reduction of Nb away from this d⁰ electron configuration. The reduced oxidation state of Nb relieves the secondorder Jahn-Teller distortion,^{10,11,17} which is commonly active for fully oxidized early transition metals, such as Nb.46 The reduction of Nb, and thus Nb off-centering, following discharge and charge results in nearly identical crystal structures at these states of charge (Fig. 8, Tables 1, and S1, S2, S3, and S4).

In contrast to ReO₃ and V₂O₅, *R*-Nb₂O₅ has minimal structure evolution during the first discharge (Fig. 8).^{29,35} The difference in cycling behavior and structural stability between *R*-Nb₂O₅ and Nb₃O₇F is especially interesting as both have slab structures with ReO₃-like regions connected by parallel, non-intersecting crystallographic shear-planes.²⁷ The difference in their behavior is most likely a consequence of the difference in the width of their slabs. The extended-defect-based structures can be described as $(2 \times \infty)_{\infty}$ for *R*-Nb₂O₅, and $(3 \times \infty)_{\infty}$ for Nb₃O₇F.³

3 Conclusions

Based on our findings, *R*-Nb₂O₅ can serve as a 'structural bridge' between Wadsley–Roth block structures and V₂O₅ in terms of polyhedral distortions, polyhedral connectivity, and the consequences thereof for structural stability during cycling. Relative to V₂O₅, we show that bridging layers to form a three-dimensionally connected structure, as in its 'idealized' structure, stabilizes it against polyhedral rotations associated with irreversible structural changes. For WR structures, this work demonstrates for the first time that a $(2 \times \infty)_{\infty}$ slab structure, with only parallel (rather than intersecting) shear planes, is stable against polyhedral rotations at corner-sharing vertices during electrochemical cycling. This is in contrast to what has

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been found for Nb₃O₇F, in which non-intersecting crystallographic shear planes that link three-octahedron-wide slabs, denoted as $(3 \times \infty)_{\infty}$, do not provide structural stability during lithiation. Thus, the combination of three-dimensionally connected polyhedra and non-intersecting shear planes with a smaller slab width results in the stability of *R*-Nb₂O₅ against unfavorable distortions, making it a high voltage anode candidate, and moreso, a link between heavily studied material families.

In the context of published work on related structures, we illustrate how lithiation in isotypic materials induces varying degrees of structural changes and phase transformations, which have direct impacts on Li⁺ diffusion pathways. The highly symmetric charge discharge profile for R-Nb₂O₅ indicates that similar sequences of phase transformations occur during lithiation and delithiation. Furthermore, the minimal structural changes result in the retention of square channels that resemble the fast Li⁺ transport pathways in WR block structures. Future work to further characterize Li⁺ and electronic conduction, as well as long-term cycling stability, will provide additional insights into bridge classes of early transition metal oxide electrode materials, including Wadsley–Roth block structures and V₂O₅.

4 Experimental

4.1 Synthesis

Phase pure *R*-Nb₂O₅ was prepared from precursors synthesized using the molten salt reaction by ion-exchange methods adapted from reported methods.^{18,36,47} Precursor KNb₃O₈ was prepared by ball-milling K₂CO₃ (Sigma Aldrich, \geq 99%), Nb₂O₅ (Sigma Aldrich, 99.99%), and KCl (Sigma Aldrich, \geq 99%) in molar ratios of 1:3:12 in ethanol (Thermo Fisher, reagent grade) for 30 min. Ethanol was evaporated under ambient conditions and the resulting mixture was heated in a covered alumina crucible at 800 °C for 5 h in air, with a heating rate of 3° min⁻¹ and allowed to cool passively with a furnace. The reacted powder was washed with warm deionized (DI) water to remove KCl; the mixture was centrifuged and decanted to isolate KNb₃O₈ powder, which was dried in air.

The prepared KNb₃O₈, ≈ 1 g, was stirred in 100 mL of 7 M HNO₃ for 3 days to prepare H₃ONb₃O₈ *via* ion-exchange.³⁶ As before, the solid product was isolated and washed using DI water, centrifugation, and decanting. After drying at room temperature, H₃ONb₃O₈ powder was heated in air at 600 °C for 2 h, yielding *R*-Nb₂O₅.³⁶ Engineering controls (*i.e.*, ventilation hood) and personal protective equipment should always be used when handling hazardous materials, such as strong acids.

4.2 X-ray and neutron diffraction

The phase purity of the intermediates and final products was initially evaluated using Rigaku Miniflex and Panalytical X'Pert Powder diffractometers. High-resolution synchrotron XRD was carried out through mail-in from beamline 11-BM at the Advanced Photon Source at Argonne National Laboratory. Synchrotron X-rays had a calibrated wavelength of 0.4589 Å and powders were measured in transmission mode in 0.8 mm diameter Kapton capillaries sealed with epoxy at both ends.

For *ex situ* synchrotron XRD of cycled R-Nb₂O₅, powder cathodes (8:2 mass ratio of Nb₂O₅ and SuperP) were extracted from Swagelok half-cells cycled to a specified state of charge. The powders were washed with dimethyl carbonate (DMC), dried under vacuum, and ground with an agate mortar and pestle in an Ar-filled glove box. The samples were sealed under Ar in secondary containment for shipping and removed just prior to measurements to minimize exposure to moisture.

Neutron diffraction data of the as-prepared R-Nb₂O₅ were collected using a POWGEN diffractometer at the Spallation Neutron Source (SNS) at Oak Ridge National Laboratory (ORNL) through the mail-in program. The powders were loaded into a 6 mm diameter vanadium can. Data were collected in time-of-flight for 2 h with a center wavelength of 1.5 Å.

Structural data were refined using GSAS-II⁴⁸ and Topas Academic v7.⁴⁹ VESTA was used to visualize the crystal structures and for quantitative descriptions of [NbO₆] octahedra.⁴⁵

4.3 Electron microscopy

Scanning electron microscopy (SEM) was carried out using a TESCAN MIRA3 microscope with a beam voltage of 8 keV and a working distance of 15 mm. The samples were prepared on carbon tape and sputtered with a thin layer of gold to prevent charging.

4.4 Battery assembly and electrochemical studies

R-Nb₂O₅ was cycled in half-cell batteries with a Li metal anode (MTI, 99.9%). For preliminary cycling and *ex situ* synchrotron XRD, loose powder cathodes (Nb₂O₅ and SuperP in 8 : 2 mass ratio, mixed in an agate mortar and pestle) were assembled in Swagelok cells. For coin cells, cast cathode films were prepared by mixing *R*-Nb₂O₅ with Ketjen Black (MSE Supplies) and polyvinylidene floride (PVDF) in a 7.5 : 1.5 : 1 mass ratio, respectively; *N*-methyl-2-pyrollidone was used as a solvent for PVDF during casting on copper foil (MSE Supplies) using a 15 μ m doctor blade.

For both cell configurations, the electrolyte was 1 M LiPF₆ in a mixture (1:1 by volume) of ethylene carbonate and DMC. For Swagelok cells, the cathode and anode were separated by two Whatman glass filter dryer (GFD) disks; for coin cells, the separator was polypropylene (Celgard 2500, 25 μ m). The cells were assembled in an Ar-filled glovebox. Swagelok cells were cycled galvanostatically at various rates in terms of C, such that C is based on the reaction of 2 mol of Li. Coin cells were cycled galvanostatically at rates of C/20, C/10 and 1C with an upper potential limit of 2.5 V and a lower potential limit of 1 V. Additionally, cells were cycled with varying cycling rates of C/20, C/10, C/5, 1C, 2C, and 5C for 5 consecutive cycles, and a final set at C/20.

Author contributions

KP: conceptualization; data curation; formal analysis; investigation; visualization; writing – original draft, reviewing and editing. ADL: investigation; especially synthesis and initial characterization (laboratory XRD). SG: investigation; especially characterization (SEM imaging). MMB: conceptualization; formal analysis; project administration; resources; supervision; visualization; writing – original draft, reviewing and editing.

Conflicts of interest

There are no conflicts to declare.

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