

On the Active Components in Crystalline Li−**Nb**−**O and Li**−**Ta**−**O Coatings from First Principles**

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Cite This: *Chem. Mater.* 2023, 35, [5657−5670](https://pubs.acs.org/action/showCitFormats?doi=10.1021/acs.chemmater.3c01197&ref=pdf) **Read [Online](https://pubs.acs.org/doi/10.1021/acs.chemmater.3c01197?ref=pdf)**

ACCESS | **ILL** [Metrics](https://pubs.acs.org/doi/10.1021/acs.chemmater.3c01197?goto=articleMetrics&ref=pdf) & More | ILL Article [Recommendations](https://pubs.acs.org/doi/10.1021/acs.chemmater.3c01197?goto=recommendations&?ref=pdf) | G Supporting [Information](https://pubs.acs.org/doi/10.1021/acs.chemmater.3c01197?goto=supporting-info&ref=pdf) ABSTRACT: Layered-oxide LiNi*x*Mn*y*Co1[−]*x*−*y*O2 (NMC) positive Li_sNbO

electrodes with high nickel content deliver high voltages and energy densities. However, a high nickel content, e.g., $x = 0.8$ (NMC811), can lead to high surface reactivity, which can trigger thermal runaway and gas generation. While claimed safer, all-solidstate batteries still suffer from high interfacial resistance. Here, we investigate niobate and tantalate coating materials, which can mitigate the interfacial reactivities in Li-ion and all-solid-state batteries. First-principles calculations reveal the multiphasic nature

of Li–Nb–O and Li–Ta–O coatings, containing mixtures of LiNbO₃ and Li₃NbO₄ or of LiTaO₃ and Li₃TaO₄. The concurrence of several phases in Li−Nb−O or Li−Ta−O modulates the type of stable native defects in these coatings. Li−Nb−O and Li−Ta−O coating materials can favorably form lithium vacancies Vac_{Li}^\prime and antisite defects $\text{Nb}_{Li}^{\bullet\bullet\bullet\bullet}$ (Ta $_{Li}^{\bullet\bullet\bullet\bullet}$) combined into charge-neutral defect complexes. Even in defective crystalline LiNbO₃ (or LiTaO₃), we reveal poor Li-ion conduction properties. In contrast, Li₃NbO₄ and Li₃TaO₄ that are introduced by high-temperature calcinations can provide adequate Li-ion transport in these coatings. Our in-depth investigation of the structure−property relationships in the important Li−Nb−O and Li−Ta−O coating materials helps to develop more suitable calcination protocols to maximize the functional properties of these niobates and tantalates.

1. INTRODUCTION

Notwithstanding the maturity of the lithium (Li)-ion battery technology, stabilizing the electrode−electrolyte interfaces and interphases sets challenges in lithium-ion batteries (LIBs), as well as in novel all-solid-state batteries $(ASSBs).^{1,2}$ $(ASSBs).^{1,2}$ $(ASSBs).^{1,2}$ For example, in LIBs, hydrofluoric acid released upon $LipF_6$ decomposition can lead to the surface corrosion of cathode particles and subsequent leaching of transition-metal ions (TMs) into the electrolyte.^{[2](#page-11-0)} Furthermore, during reversible Li intercalation in cathode active materials (CAMs), significant volume variations of the active particles may induce cracks that lead to loss of contact among the active particles, and a subsequent reduction in performance.^{[3](#page-11-0)}

Thiophosphate solid electrolytes (SEs), e.g., $\rm Li_{10}GeP_2S_{12}$ (LGPS) and argyrodites Li_6PS_5X (with X = Cl, Br, I),^{[4](#page-11-0)} display high Li-ion conductivities and low-temperature processability and hence are promising candidates for ASSBs. However, the low oxidative (or anodic) stabilities of thiophosphate SEs can result in undesired side reactions, especially with high-voltage $($ >4.5 V vs. Li/Li⁺ $)$ CAMs.^{[5,6](#page-11-0)}

Introducing coating materials is one of the most effective solutions to mitigate possible interfacial instabilities in LIBs and ASSBs, improving their capacity retention, rate capabilities, and longevity of cells.[7](#page-11-0)−[9](#page-11-0) Coating materials can enhance the "adhesion" between cathode particles and act as physical protections from undesired chemical reactions. Coating materials also stabilize the cathode−electrolyte interfaces

chemically and electrochemically, thereby increasing or modulating the stability windows of electrolytes.

Simultaneously, coatings in LIBs are designed to have high ionic and electronic conductivities-both properties are essential for maintaining percolating ionic and electronic networks between cathode particles.² The requirements of coatings (for cathode electrodes) in ASSBs are different from LIBs: (i) deliver high Li-ion conductivity, and (ii) maintain sufficiently low electronic conductivity to prevent SEs with narrow stability windows from being easily oxidized by highvoltage CAMs.⁹

In particular, amorphous $LiNbO₃$ and $LiTaO₃$ materials with high ionic conductivities (∼10[−]⁵ −10[−]⁶ S cm[−]¹ at room temperature) and appreciable electronic resistance (∼10[−]¹¹− 10^{-12} Ω^{-1} cm⁻¹) have been effectively used as coatings in ASSBs.[10](#page-11-0)−[12](#page-11-0) Albeit with much lower Li-ion conductivities $(<10^{-12}$ S cm⁻¹ at 400 K) and exceedingly high migration barriers (1.16−1.33 eV),^{13,[14](#page-11-0)} crystalline LiNbO₃ and LiTaO₃ coatings show substantial enhancements in the electrochemical performance of LIBs (see [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S1).^{[15](#page-11-0)−[18](#page-11-0)}

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Figure 1. (a) Structures of LiNbO₃ (LiTaO₃) (R3*c*) with corner-sharing Nb(Ta)O₆ octahedra, transformed into pseudo-cubic structures using the [[1, 1,−1], [-1, 1, 1], [1,−1, 1]] transformation matrix. (b) Structure of Li₃NbO₄ (*I*43*m*) with edge-sharing Nb(Ta)O₆ octahedra. For the centered Li site in LiNbO₃ (LiTaO₃) in panel (a), there are two first nearest neighbors (NNs), four second NNs, four third NNs at the top and bottom Li layers, and four first NNs, two second NNs, two third NNs on the middle Li layer, as well as eight nearby Nb(Ta) atoms. The left panel in (b) describes the local environment of a Li site in Li₃NbO₄, and the right panel shows the arrangement of four Nb atoms within Nb_4O_{16} clusters at the center of orange and yellow planes. Additional views of these structures emphasizing the Li-ion environments are presented in [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S2.

As the information available in the literature on amorphous and crystalline phases of niobate and tantalate coatings appears scattered, it is important to set a firm baseline on the crystalline phases of these materials. While amorphous niobates and tantalates appear to provide the highest ionic conductivities, $10,13,19$ the crystalline analogues are reported to increase the hardness of cathode particles. $20,21$

Although higher levels of crystallinity in $LiNbO₃$ (or $LiTaO₃$) have been linked to a reduction of CAM particle cracking, $20,21$ there exists an apparent contradiction between the low Li-ion conductivity of crystalline $LiNbO₃$ (or $LiTaO₃$) and the good rate capabilities achieved in these cells.^{15,[19](#page-11-0)} Indeed, the texture and composition of these niobate and tantalate materials appear evidently more complicated than commonly perceived.

Previous reports 13,14 13,14 13,14 suggested that highly defective nanocrystalline LiNbO₃ and LiTaO₃ (from high-energy ball milling) containing amorphous-like components exhibit room-temperature Li-ion conductivities $(\sim\!10^{-6}$ S cm $^{-1})$, comparable with their amorphous analogues $(10^{-5} - 10^{-6} \text{ S cm}^{-1})$. These facts hint at the high sensitivity of Li-ion conductivity in $LiNbO₃$ and $LiTaO₃$ to the presence of defects in their crystalline phases.

Conventional synthesis protocols of crystalline niobate (or tantalate) coatings typically apply the wet-chemical method followed by prolonged (3−10 h long) calcinations at 773−1073 K ([Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S1). Furthermore, $LiMO₃$ (with M = Nb and Ta) tends to crystallize out from the congruent melts (Li-poor, Mrich) at off-stoichiometric compositions, facilitating the incorporation of intrinsic defects (see [Figure](#page-3-0) 3).[22](#page-11-0)−[25](#page-11-0) However, neutron reflectometry and secondary ion mass spectroscopy measurements did not suggest a marked increase in Li-ion transport in these materials.^{[26](#page-11-0),[27](#page-11-0)} Therefore, the role played by intrinsic defects in the functional properties of niobate and tantalate coatings remains ambiguous.

Recent investigations on the composition of Li−Nb−O coating highlight the coexistence of two primary phases: LiNbO₃ and Li₃NbO₄.^{[15](#page-11-0),[16,28](#page-11-0)} Zhang et al.¹⁵ suggested that the coexistence of $LiNbO₃$ and $Li₃NbO₄$ improves the electrochemical performances of coated cathodes. Several reports suggested that the Li⁺ migration barrier in crystalline $Li₃NbO₄$ (0.58−0.85 eV) is considerably lower than that of crystalline LiNbO₃ (1.16−1.33 eV).^{19,[29](#page-11-0)} Liao et al. and Yabuuchi et al. showed an increase in Li-ion conductivity by several orders of magnitude in Li_3NbO_4 upon doping with W^{6+} , or substituting of $Ni²⁺$ for Li⁺, which introduces Li vacancies.^{[29](#page-11-0),[30](#page-11-0)} This evidence suggests $Li₃NbO₄$ may be the active phase in transporting Li ions in niobate coatings.

Based on the facts exposed above, protocols employed for synthesizing crystalline niobate coatings can strongly affect: (i) the partition of the active phases of Li–Nb−O coatings,^{[16,28](#page-11-0)} (ii) the concentrations of intrinsic defects, and in turn affect (iii) the Li-ion diffusion in these materials. This knowledge gap requires immediate attention to optimize the functional properties of niobates and tantalates.

Leveraging a combination of first-principles calculations (density functional theory, DFT), bond valence mapping (through the $SoftBV)$, 31 nudged elastic band (NEB) calculations, and machine-learning (ML) molecular dynamics (MDs), we investigate how the Li migration properties of LiMO₃ and Li₃MO₄ (with M = Nb and Ta) are modulated by the occurrence of thermodynamically viable intrinsic defects. We demonstrate that what is commonly perceived as a $LiNbO₃$ (or $LiTaO₃$) coating material, in practice, is a multiphasic composite including variable fractions of $Li₃NbO₄$ (or $Li₃TaO₄$), altering the conduction property of the material. The occurrence of intrinsic defects in niobates (and tantalates) remains conditional to the coexistence of $LiMO₃$ and $Li₃MO₄$ at synthesis conditions, which regulates the availability of Li vacancies. A progressive transformation of $LiMO₃$ into $Li₃MO₄$ by prolonged high-temperature calcination results in a much less defective $Li₃MO₄$ coating material, which in turn hinders $Li⁺$ transport and deteriorates the functionality of the coating.

2. CRYSTAL STRUCTURES OF LIMO₃ AND LI₃MO₄

[Figure](#page-1-0) 1 shows $LiNbO₃$ and $LiTaO₃$ perovskite structures (space group *R*3*c*), which present octahedra tilting.[32](#page-12-0),[33](#page-12-0) In LiNbO₃ (LiTaO₃), surrounded by oxygen atoms, Li and Nb (Ta) cations occupying 2/3 octahedral sites are aligned along the *c*-axis (see [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S1) and follow a stacking sequence of Nb (Ta), Li (or vacancy), and Nb (Ta).^{[22](#page-11-0)} Both LiNbO₃ and LiTaO₃ show a single Li Wyckoff site.

We analyze the immediate environment of Li ions (gold atom in [Figure](#page-1-0) 1a), which is important to set up different models of intrinsic defect complexes, and investigate Li-ion migration pathways in LiNbO₃ and LiTaO₃. For a Li site in LiNbO₃ and $LiTaO₃$, there are six first nearest neighbors (NNs), six second NNs, and sixthird NNs Li sites in total from the top, middle, and bottom Li layers, respectively. The eight neighboring $Nb(Ta)O_6$ are shown in [Figure](#page-1-0) 1a as green octahedra.

 $Li₃NbO₄$ shows a disordered rock-salt structure with an average *Fm3_m* space group ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S3) when synthesized at a low temperature (at ∼220 °C).^{[34,35](#page-12-0)} At 700 °C, Li₃NbO₄ transforms into an *I*43*m* ordered phase [\(Figure](#page-1-0) 1b)[,36](#page-12-0)[−][38](#page-12-0) which highlights the edge-sharing $NbO₆$ octahedra. In the right panel of [Figure](#page-1-0) 1b, three Nb cations (at the center) form a tetramer (Nb_4O_{16}) with the other Nb cation in subsequent layers. Together with Li ions (and vacancies), these $Nb₄O₁₆$ clusters form a body-centered cubic lattice. Compared with the *R*3*c* closely packed LiNbO₃, this arrangement might allow larger voids that facilitate Li transport. For each Li site, there are two first NNs, four second NNs, and one third NN as shown in the left panel of [Figure](#page-1-0) 1b.

 $Li₃TaO₄$ exists in three main polymorphs ([Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S3): (i) a disordered low-temperature phase with an average space group *Fm*3*m* (similar to Li₃NbO₄, [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S3), (ii) an intermediate βphase (with a space group *C*2/*c*), and (iii) a high-temperature (>1450 °C) α -phase structure (with a space group $P2$).^{[34](#page-12-0),[38,39](#page-12-0)} Since the synthesis temperature (400−800 °C, [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S1) of crystalline Li−Ta−O coating materials is usually lower than 1450 °C, the formation of the high-temperature *α*-phase of Li₃TaO₄ can be excluded from the composition of typical Li− Ta−O coating materials. Therefore, we speculate that the ordered *β*-phase C_2/c Li₃TaO₄ phase can be the other constituent of the coating material together with the *R*3*c* LiTaO₃ phase. The ordered $C2/c$ Li₃TaO₄ shows zigzag chains formed by the edge-sharing TaO_6 octahedra surrounded by Li ions (lower panel of Figure 2). $Li₃TaO₄$ contains eight

Figure 2. Structure of β -Li₃TaO₄ in the *C*2/*c* space group. Unique Li sites are labeled and shown with different colors. The lower panel shows edge-sharing (green) TaO₆ octahedra zigzag along the *c*-direction. Li environments in β -Li₃TaO₄ are highlighted in [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S2c.

independent crystallographic sites (three Li, one Ta, and four O) in the unit cell, 40,41 40,41 40,41 which denotes the substantial difference in terms of local arrangements compared to *R*3*c* LiNbO₃, LiTaO₃, and $I43m$ Li₃NbO₄ displaying a unique Li site.

3. PHASE EQUILIBRIA AND ELECTROCHEMICAL STABILITIES OF LI−**NB**−**O AND LI**−**TA**−**O COATING MATERIALS**

[Figure](#page-3-0) 3a,b shows the experimental binary phase diagrams of Li₂O− $Nb₂O₅$ and $Li₂O-Ta₂O₅$ systems in the composition range spanning from 40 to 62 mol % $Li_2O^{24,33,42}$ $Li_2O^{24,33,42}$ $Li_2O^{24,33,42}$ $Li_2O^{24,33,42}$

In [Figure](#page-3-0) 3a, an exceeding (decreasing) $Li₂O$ content into $LiNbO₃$ leads to the formation of a secondary phase, $Li₃NbO₄$ (LiNb₃O₈), that coexists with $LiNbO₃$ over a large range of composition. In [Figure](#page-3-0) 3b, $Li₃TaO₄$ appears as a secondary phase with $Li_TaO₃$ in the $Li₂O$ -rich regions.

[Figure](#page-3-0) 3c,d shows the computed $(\varpi$ ²SCAN) phase diagrams at 0 K of Li₂O−Nb₂O₅ and Li₂O−Ta₂O₅ pseudo-binary lines including all experimentally known compositions. Green circles show the lowestenergy structures computed by DFT, while phases predicted as metastable by DFT but identified experimentally are shown by yellow circles. $Li_8Ta_2O_9$ has been excluded from [Figure](#page-3-0) 3d due to the unavailability of accurate structural information. As for the disordered $Li₃MO₄$ (M = Nb, Ta), 32 unique orderings of Li/M have been generated for each composition using a ∼2 × 2 × 1 supercell model (including 4 f.u.). Red squares show the formation energies of each Li/ M ordering.

Among all competing phases, $R3c$ LiNbO₃ and $\overline{I4}3m$ Li₃NbO₄ lie on the $Nb₂O₅-Li₂O$ convex hull—the convex envelope of structures with the lowest formation energies. Therefore, any Li-rich compositions of LiNbO₃ (addition of Li₂O to LiNbO₃) will be a mixture also containing Li3NbO4 and in excellent agreement with [Figure](#page-3-0) 3a. Similarly, *R*3*c* LiTaO₃ and *C*2/*c* Li₃TaO₄ are the most stable phases in the Ta₂O₅− $Li₂O$ convex hull. Note, the formation energy of $LiTa₃O₈$ falls slightly above (\sim 7 meV atom $^{-1}$) the convex hull set by Ta₂O₅ and LiTaO₃. This is in agreement with the absence of the $LiTa_{3}O_{8}$ phase as observed in the experiments, where only Ta_2O_5 and $LiTaO_3$ phases form under Li-

Li/M (M=Nb, Ta) orderings of disordered rock-salt Li₃MO₄ structure (average Fm3m) n.

Figure 3. Phase diagrams of (a) Li₂O-Nb₂O₅ and (b) Li₂O-Ta₂O₅ pseudo-binary systems redrawn from previous experimental data.^{[24](#page-11-0),[33,42](#page-12-0)} The melting points of LiNbO₃ and LiTaO₃ are ∼1240 and ∼1650 °C, respectively.³³ Phase diagrams of (c) Li₂O–Nb₂O₅ and (d) Li₂O–Ta₂O₅ computed from DFT (r²SCAN) at 0 K. Metastable or unstable Li/M orderings of Li₃MO₄ (M = Nb, Ta) are marked by red squares. The dashed line in panel (d) is a guide for the eye to show that $LiTa_3O_8$ sits slightly above the stability line formed by Ta_2O_5 and $LiTaO_3$.

Figure 4. DFT-computed (with r 2 SCAN and at 0 K) stability regions of niobate and tantalate phases in (a) Li−Nb−O and (b) Li−Ta−O domains. Thermodynamically stable regions are labeled, of which LiNbO₃, Li₃NbO₄, LiTaO₃, and Li₃TaO₄ are highlighted with different colors. Stable phases feature as a function of changes in the chemical potential ($\Delta \mu_{\rm L}$) $\Delta \mu_{\rm Nb}$, and $\Delta \mu_{\rm T}$) of Li, Nb (or Ta) referenced to their standard elemental states. Variations of $\Delta \mu_{Li}$ can be easily converted into voltages (see the main text).

poor conditions.[24](#page-11-0)[,42](#page-12-0) The computed phase diagrams of Figure 3 can

To evaluate the electrochemical stability windows of these coating materials, we construct ternary Li−M−O (M = Nb, Ta) phase diagrams based on the elemental chemical potential changes, $\Delta \mu_{\text{Li}}$ and $\Delta \mu_{\text{M}}$ as

qualitatively and quantitatively reproduce experimental observations.

Table 1. Computed Electrochemical Stability Windows (in V) of Coating Materials, and Their Reaction Energies **Δ***E*rxt (meV atom[−]¹) with Different Positive Electrodes and Solid Electrolyte Chemistries Obtained from Previous Computational Work[s43,44](#page-12-0)*^a*

	stability window		$\Delta E_{\rm{rst}}$ //cathode ⁴³				$\Delta E_{\rm{rxt}}$ //SE ⁴⁴	
coating	red.	OX.	NMC	LCO	LMO	LFP	Li ₃ PS ₄	LLZO
LiNbO ₃	1.89	3.86	$-4/0$	0/0	$0/-21$	$-23/-35$	-155	-76
Li ₃ NbO ₄	1.06	3.58					-132	-4
LiTaO ₃	1.25	3.94	0/0	0/0	$0/-7$	$-20/-24$	-49	-68
Li ₃ TaO ₄	0.44	3.55					-64	-3

a Red. and Ox. are for reductive (anodic) stability and oxidative (cathodic) stability, respectively. NMC is the abbreviation for LiNi*x*Mn*y*Co1[−]*x*−*y*O2, LCO for LiCoO₂, LMO for Li₂Mn₂O₄, LFP for LiFePO₄, and LLZO for Li₇La₃Zr₂O₁₂. Values of ΔE_{rxt}//cathode are reaction energies of full/halflithiated cathodes and coating interfaces.

Figure 5. Computed formation energies ΔE_D of intrinsic point defects in (a) LiNbO₃ and (b) Li₃NbO₄ vs. Fermi energy and under the most Li-poor conditions. The chemical potentials for panel (a) were defined by the equilibrium LiNbO₃–LiNb₃O₈–O₂ ($\Delta \mu_{Li}$ = −3.86 eV), and (b) by the Li3NbO4−LiNbO3−O2 equilibrium (Δ*μ*Li = −3.58 eV). Fermi energy is referenced to the valence-band maximum (VBM). The band gaps of ∼3.88 eV for LiNbO₃ and ~4.26 eV for Li₃NbO₄ set maximum values of the Fermi energy.

seen in [Figure](#page-3-0) 4. All competing phases available in the ICSD were included.

The feasible ranges of $\Delta \mu_{\text{Li}}$ and $\Delta \mu_{\text{M}}$ in which LiMO₃ and Li₃MO₄ are thermodynamically stable are shown by colored regions in [Figure](#page-3-0) [4a](#page-3-0),b. $\Delta \mu_{\text{Li}}$ is the Li chemical potential (μ_{Li}) referenced to the chemical potential of Li metal (μ^0_Li) . μ_Li is converted using the Nernst equation

into voltages (versus Li/Li⁺), $\phi = \frac{\mu_{\rm Li}^0 - \mu_{\rm Li}}{eF} = -\frac{\Delta \mu_{\rm Li}}{eF}.$

At more negative potential $(\Delta \mu_{\rm Li})$ of [Figure](#page-3-0) 4, which is more positive voltage vs. Li/Li^{+} , $LiNbO₃$ $(LiTaO₃)$ is in direct equilibrium with $LiNb₃O₈ (LiTa₃O₈)$ and in agreement with the phase diagram of [Figure](#page-3-0) [3.](#page-3-0) This condition is said to be Li-poor or oxidative and can be realized, for example, when the coating material interfaces with a high-voltage cathode material, such as >4.20 V vs. Li/Li⁺ in LiNi_xMn_yCo_{1−*x*−*y*}O₂, NMC. The Li-poor conditions can also be mimicked by a deficiency of Li precursors during synthesis. In contrast, at more positive potentials (lower voltages vs. Li/Li^{+}), $LiNbO₃$ ($LiTaO₃$) is in equilibrium with $Li₃NbO₄$ ($Li₃TaO₄$). This condition sets a chemically reducing environment (or Li-rich). From a synthesis point of view, this situation can be achieved by adding an excess of Li precursors or reacting with Li resources within cathodes.

The computed $(\varpi$ ²SCAN) electrochemical stability windows of $LiMO₃$ and $Li₃MO₄$ are summarized in Table 1, which are in good agreement with previous reports.⁴⁴ LiNbO₃, LiTaO₃, Li₃NbO₄, and Li₃TaO₄ show relatively poor reducing (cathodic) stabilities toward low-voltage electrode materials, with the lowest voltage (∼0.44 V vs. Li/Li^{+}) hit by $Li_{3}TaO_{4}$. Similarly, $LiNbO_{3}$ and $LiTaO_{3}$ show lowoxidative (anodic) limits below 4 V vs. Li/Li⁺ (see Table 1). Furthermore, the computed values of anodic limits of $Li₃NbO₄$ and $Li₃TaO₄$ appear slightly lower than their $LiMO₃$ analogues.

It is important to show whether the coating materials will mix spontaneously with the most common cathode electrodes, such as LiCoO2 (LCO), LiMn2O4 (LMO), LiNi*x*Mn*y*Co1[−]*x*−*y*O2 (NMC), and LiFePO₄ (LFP), or electrolytes, e.g., Li₃PS₄ and Li₇La₃Zr₂O₁₂ (LLZO). The ΔE_{rxt} of eq 1 is determined by the mixing ratio *x* providing the largest enthalpy of mixing 43

$$
\Delta E_{\rm rxt} = \min_{x \text{ in } [0,1]} \{ E_{\rm pd}[xc_a + (1-x)c_b] - xE(c_a) - (1-x)E(c_b) \}
$$
\n(1)

where E_{pd} , $E(c_{\text{a}})$, and $E(c_{\text{b}})$ are the DFT energies of the mixed phase, electrode (or electrolyte) material c_a , and coating material c_b , respectively. From Table 1, niobates and tantalates exhibit very low chemical reactivity $(\Delta E_{\rm{rst}})$ with most oxide-based positive electrodes.^{[43](#page-12-0)} A low chemical reactivity with high-voltage electrodes is further enhanced if these niobates and tantalates are crystalline instead of amorphous.

When paired with thiophosphate SEs, i.e., $Li₃PS₄$ in Table 1, the large ΔE_{rxt} suggests that niobates may not be as stable as tantalates with thiophosphate SEs. However, the limited reactivity of tantalates compared to niobates may be explained by the lack of reported compounds containing Ta and S in contrast to more studied phases incorporating Nb and S.

4. INTRINSIC DEFECTS IN LI−**NB**−**O AND LI**−**TA**−**O COATINGS**

As shown in the phase diagram [\(Figure](#page-3-0) 3a), the congruent composition of LiNbO₃ corresponds to ~48.55 mol % Li₂O and 51.45 mol % $Nb₂O₅$, with a [Li]/[Nb] ratio equal to 0.94, indicating a Li deficiency of ~6% in LiNbO₃.^{[45](#page-12-0)} Previously, the

nonstoichiometry in $LiNbO₃$ was explained by three possible defect models: (i) lithium and oxygen vacancy, [Li]_{1−2*x*}Nb-[O]3[−]*x*, (ii) lithium vacancy and niobium antisite, [Li]1[−]5*x*[NbLi]*x*NbO3, and (iii) niobium vacancy and niobium antisite, $Li[Nb_{Li}]_{5x}[Nb]_{1-4x}O_3$.^{[23](#page-11-0),[33](#page-12-0)} Model (i) can be immediately excluded as the density of the nonstoichiometric phase was found to increase with decreasing lithium content.^{22,[46](#page-12-0)}

To date, model (ii) is the most accepted to explain the nonstoichiometry of LiNbO₃. X-ray and neutron diffraction,^{[22](#page-11-0)[,47](#page-12-0)} and solid-state nuclear magnetic resonance^{[48](#page-12-0)} experiments of LiNbO₃ suggested that ∼1% of the Li sites are occupied by Nb, and ∼4% of the Li sites are vacant. This ratio of 1:4 is in line with the formation of a complex charge-neutral defect encompassing one Nb antisite defect (Nb^{oon}) accompanied by four Li vacancies (Vac'_{Li}) using the Kroger–Vink notation. Firstprinciples calculations confirmed that model (ii) is preferred compared to model (iii) under both Li-rich and Li-poor conditions.⁴⁹ Donnerberg et al.⁵⁰ investigated cation vacancies in $LiNbO₃$ and showed that the formation of Nb vacancies Vac $\frac{m}{N}$ is less favorable than the formation of Li vacancies Vac \dot{L} .

As both LiNbO₃ and Li₃NbO₄ are reported in the Li−Nb−O coating layer, $15,17,28$ $15,17,28$ $15,17,28$ here, we extend our defect analysis to $Li₃NbO₄$. Computational studies on the stability of chargeneutral defects in Li_3NbO_4 and $LiNbO_3$ suggested that a Li/Nb antisite neutral complex and Li-Frenkel defects could exist.^{[49,51](#page-12-0)} Therefore, we investigate the stability of Li vacancies (Vac $_{Li}$) and Nb antisites (Nb_{Li}) , as well as the charge-neutral complex defects $[4Vac'_{Li} + Nb_{Li}^{\bullet\bullet\bullet\bullet}]$ of model (ii).

[Figure](#page-4-0) 5 plots the computed $(\emptyset r^2\text{SCAN})$ formation energies $\Delta E_{\rm D}$ of isolated Li vacancies (Vac_{Li}) and Nb antisites (Nb_{Li}) in $LiNbO₃$ and $Li₃NbO₄$ following [eq](#page-10-0) 2. Results of isolated point defects in $LiTaO_3$ and Li_3TaO_4 are provided in [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S7 and [Tables](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S8, S9. $\Delta E_{\rm D}$ values are plotted as a function of the Fermi energy (E_{Fermi}). We referenced the absolute values of the Fermi energy to the energy of the valence-band maximum (VBM) of each bulk structure. The slopes of the defect lines represent different charge states of the defect (*q* in [eq](#page-10-0) 2).

High-temperature calcination facilitates the consumption of Li resources in positive electrode materials, hence setting a condition of Li-rich for LiNbO₃ and Li-poor for $Li₃NbO₄$ at the coexistence of these two phases ([Section](#page-8-0) 6). Upon charging, the availability of free Li in the cathode sets Li-rich conditions for the coating components (see [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S6). Therefore, it is crucial to investigate the formation of defects under different chemical/ electrochemical conditions of the coating.

Solid lines in [Figure](#page-4-0) 5 represent the ΔE _D values under the most Li-poor (oxidative) conditions for LiNbO₃ (LiNbO₃ $-$ LiNb₃O₈−O₂ equilibrium) and Li₃NbO₄ (Li₃NbO₄−LiNbO₃− $O₂$ equilibrium). The defect formation energies under the most Li-rich (reducing) conditions of LiNbO₃ and Li₃NbO₄ can be found in [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S6. By definition, values of $\Delta E_{\rm D}$ must be always positive quantities. In [Figure](#page-4-0) 5a,b at specific Fermi energies (<1 and >2.5 eV), the computed ΔE _D values are negative. This indicates that these defects are unstable in the phase space set by the chemical potentials of Li and Nb ([Figure](#page-3-0) 4).

Based on the temperature of calcination of coating materials $({\sim}1000 \text{ K})$,^{[15](#page-11-0),[16](#page-11-0),[28](#page-11-0)} both LiNbO₃ and Li₃NbO₄ phases coexist in the crystalline niobate coatings. Room-temperature (300 K) data are shown in [Tables](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S6 and S7. Vertical dotted (blue) lines are the equilibrium Fermi energy $(E_{F,eq})$ under the Li-poor condition. Note, the equilibrium Fermi energy informs about the availability of electrons (or holes) in the coating material. $52,53$ $52,53$ $52,53$

Under the most Li-poor condition for LiNbO₃ ([Figure](#page-4-0) 5a), the equilibrium Fermi Energy (*E*_{F,eq}) is ~1.271 eV at 1000 K. The defect formation energies for Vac_{Li} and $\text{Nb}_{\text{Li}}^{\bullet\bullet\bullet}$ are 0.62 and 0.74 eV, resulting in defect concentrations at 10^{19} and 10^{18} cm⁻³, respectively. The carrier (hole) concentration reaches the order of 10^{15} cm⁻³. Under the most Li-rich condition in LiNbO₃ ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S6a), $E_{\rm F,eq}$ is shifted to 2.89 eV at 1000 K, where $\Delta E_{\rm D}$ for $\text{Vac}_{\text{Li}}^{\prime}$ and $\text{Nb}_{\text{Li}}^{\bullet\bullet\bullet}$ are 0.96 and 1.09 eV, respectively. The corresponding defect concentrations decrease to 10^{17} and 10^{16} cm[−]³ . This trend shows that both Li vacancies and Nb antisites are less favored in the Li-rich condition of $LiNbO₃$ at the same temperature, resulting in a decrease in defect concentrations. $LiNbO₃$ in Li-rich conditions displays a higher free-electron concentration around 10^{16} cm⁻³.

Under the most Li-poor condition in $Li₃NbO₄$ at 1000 K ([Figure](#page-4-0) 5b), $E_{F,eq}$ is at 0.93 eV. The corresponding ΔE_{D} for Vac_{Li} and Nb^{***} are 1.04 and 1.28 eV, much higher than those of LiNbO₃. The defect concentrations of Vac_{Li} and Nb_{Li}^{•••} are at 10¹⁷ and 10¹⁶ cm[−]³ , respectively. The charge (hole) carrier concentration is 10^{17} cm⁻³. Under the most Li-rich condition for Li₃NbO₄ at 1000 K ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S6b solid lines), $E_{\text{F,eq}}$ moves to 2.85 eV, where ΔE _D for Vac_{Li} is 1.61 eV, and for less probable defects Nb_{Li}^{oo} and Nb_{Li}^{oo} , 1.71 and 1.75 eV, respectively. This indicates that Nb_{Li} with lower charge states are likely to be more stable than $Nb_{Li}^{••••}$ in Li_3NbO_4 under such conditions. But the defect concentrations of Li vacancies and Nb antisites are only at 10^{14} and 10^{13} cm⁻³ and hence several orders of magnitude lower than those in the Li-poor condition. The free-electron concentration of Li₃NbO₄ in the Li-rich condition is only around 10^{13} cm⁻³.

We now analyze the stability of neutral defect complex in LiMO₃ and Li₃MO₄ (M = Nb, Ta) comprising of four Li vacancies together with one Nb (Ta) antisite, thereafter indicated as $[4\text{Vac}_{Li} + M_{Li}^{\bullet\bullet\bullet\bullet}]$. The spatial arrangement of antisite and Li vacancies in LiNbO₃ and Li₃NbO₄ may vary and affect the relative stability of these defects. To this end, we investigated a number of possibilities (spatial orderings), whose details are in [Section](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S3. Defect complex models in $Li₃TaO₄$ with three unique crystallographic Li sites were constructed for each Li site. The configuration where the Li2 site is exchanged by a Ta atom shows the lowest formation energy. Table 2 summarizes the $\Delta E_{\rm D}$ of lowest-energy configurations among all defect models under Li-poor conditions.

As shown in Table 2, values of ΔE ²s of neutral defect complexes in $LiMO₃$ and $Li₃MO₄$ are comparatively lower than the sum of the individual point defects (last column). For example, the sum of defect formation energies for $4Vac'_{Li}$ and Nb<mark>°°°</mark>* in LiNbO₃ at *E*_{F,eq} under the Li-poor condition is ∼3.23

Table 2. Computed r 2 SCAN Defect Formation Energies $(\Delta E_{\text{D}}$ in eV) of Neutral Antisite Defect Complexes ([4Vac_{Li} + $\mathbf{M^{...}}$) Compared to the Sum of Isolated Defects (4Vac $'_{\text{Li}}$ + M_{Li}^{300} , $M = Nb$, $Ta)^a$

 a Calculations of antisite defect complexes $[4\text{Vac}_{\text{Li}}' + \text{M}_{\text{Li}}^{\bullet\bullet\bullet\bullet}]$ used large supercell models of $LimO_3$ $(Li_{64}M_{64}O_{192})$ and Li_3MO_4 $(Li_{96}M_{32}O_{128})$. The Li-poor reference states are provided in [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) [S10](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf).

eV, while that of the complex neutral defect is ∼2.26 eV. This indicates the favorable enthalpy gain of forming a stable complex as discussed by Li et al. 54 In the reducing (Li-poor) conditions, the minimum ΔE_D of the defect complex in Li₃NbO₄ is ~2.61 eV.

5. LI-ION MOBILITY IN LIMO3 AND LI3MO4 COATINGS

In this section, we address the mobility of Li ions in $LiMO₃$ and $Li₃MO₄$ by performing inexpensive but empirical SoftBV predictions, followed by accurate first-principles nudged elastic band (NEB) simulations.

Figure 6 shows the bond valence site energy (BVSE) landscape computed with SoftBV for each bulk structure

Figure 6. BVSE landscapes obtained from SoftBV of (a) *R*3*c* LiMO₃ (M $=$ Nb, Ta), (b) *I43m* Li₃NbO₄, and (c) *C*2/*c* Li₃TaO₄. Values of volumetric isosurfaces are set at 2000, 700, and 700 meV above the global minimum-energy set by Li sites, within LiNbO₃ (LiTaO₃), $\rm Li_3NbO_4$ and $\rm Li_3TaO_4$ respectively. Li sites setting global minima are colored in orange. Nb and Ta atoms are shown in green and serve as a guide for the eye.

whose volumetric isosurfaces are set to 2000 meV (in $LiMO₃$) and 700 meV (in $Li₃MO₄$), above the global minimum of Li-site energy computed by SoftBV. From the disconnected isosurfaces seen in Figure 6a, it is unlikely for Li ions to migrate in the defect-free crystalline phases of $LiNbO₃$ or $LiTaO₃$, as no Li "channels" are formed between adjacent sites. Figures 6b and [S8](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) clearly depict extended percolating Li-diffusion channels of Li₃NbO₄. For the ordered *β*-phase Li₃TaO₄, Figures 6c and [S9](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) show the percolating networks of Li ions. Insightful illustrations of the migration environments near the crystallographic Li1, Li2, and Li3 sites are provided in [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S13a−c.

From the SoftBV analysis, the percolating migration barriers-that give rise to a net transport of Li ions within the coating material-were further investigated using NEBs coupled with DFT@r²SCAN. Three main types of Li $^+$ migration pathway can be envisioned as below.

- 1. Pathway Vac_{Li}: One Li vacancy Vac_{Li} (low-vacancy limit) was introduced and compensated by a background charge (in the form of jellium).
- 2. Pathway $4Vac_{Li} + M_{Li}$ (M = Nb, Ta): One Li vacancy Vac_{Li} of the four vacancies of the charge-neutral defect complex $[4Vac'_{\lambda i} + M_{Li}^{\bullet \bullet \bullet \bullet}]$ migrates.
- 3. Pathway $4Vac\hat{L} + M_{Li}$: One Li vacancy Vac $'_{Li}$ of the four vacancies of the charge-neutral complex defect $[4Vac_{Li}^{\prime} +$

 $\mathbf{M}^{\bullet\bullet\bullet\bullet}_{\mathrm{Li}}$] migrates. All of the other three $\mathrm{Vac}_{\mathrm{Li}}$ were located near the migration event.

Based on these three distinct pathways, distinct migration models were identified by looking at the coordination environment of the migrating Li-ion [\(Figures](#page-1-0) 1 and [2](#page-2-0)). Values of computed migration barriers are summarized in [Figure](#page-7-0) 7 and [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S11.

From [Figure](#page-7-0) 7a,c, no matter which of the three pathways are chosen, the predicted migration barriers in $LiNbO₃$ and $LiTaO₃$ are consistently larger than 1100 meV. This data clearly demonstrates that Li-ion transport in LiNbO₃ and LiTaO₃ relying on intrinsic defects cannot guarantee sufficient Li-ion transport.

In particular, for $LiNbO₃$ and $LiTaO₃$, the Li-ion migration barriers in the low-vacancy limit model (Vac $_{Li}$) are 1273 and 1240 meV, respectively, in agreement with previous exper-imental results.^{13,[19](#page-11-0)} To investigate whether the presence of the neutral defect complex $([4\text{Vac}_{Li}^{\prime} + \text{M}_{Li}^{\bullet\bullet\bullet}])$ can lower the high migration barriers of crystalline $LiNbO₃$ (LiTaO₃), we studied the Li-ion migration in the vicinity of these defects (see [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) [S11b,c\)](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf). As shown in [Figure](#page-7-0) 7a,c, migration barriers are still as high as ~1296 meV for LiNbO₃ and ~1406 meV for LiTaO₃. The computed migration energies of Li-ion in congruent LiNbO₃ (\sim 1296 meV) appear in excellent agreement with the experimental values (\sim 1300 meV).^{26,27}

Starting from the hypothesis that the presence of Li vacancies near the migrating Li-ion may lower its migration barriers, in Li migration models labeled as $4Vac\Delta_{Li} + Ta_{Li}$, we forced all four Li vacancies to be close to the Li migration pathway ([Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S11d). Although the activation barriers (\sim 1126 meV for LiNbO₃ and $∼1146$ meV for LiTaO₃) are slightly lower than in the other cases, such barriers are still too high to achieve reasonable Li-ion transport across the coating layer.

 $Li₃NbO₄$ and $Li₃TaO₄$ exhibit much more promising transport properties with lower activation barriers, on average lower than 700 meV [\(Figure](#page-7-0) 7b,d). For example, in $Li₃NbO₄$, one Li vacancy can migrate to its two first NNs (∼467 meV), four second NNs (∼543 meV), and one third NN (∼701 meV, shown in [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S12b) sites, respectively. Among 11 distinct pathways considered between three types of Li sites in $Li₃TaO₄$, eight migration barriers are consistently lower than 700 meV (see [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S11). Therefore, in Li_3NbO_4 and Li_3TaO_4 , several migration pathways may enable macroscopic diffusion of Li ions ([Figures](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S8 and S9).

In analogy to $LiMO₃$, we also explore the Li migration in $Li₃MO₄$ (M = Nb, Ta) in the vicinity of the $M_{L,i}$ antisite and four Li vacancies. The computed paths of these Li migrations are shown in [Figures](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S12d and S13d−f, respectively. From [Figure](#page-7-0) $7b$ $7b$, in Li₃NbO₄, the Li-ion migration barriers generally increase in the proximity of Nb antisite defects when compared to the migration of a diluted Li vacancy (models Vac_{Li} in [Figure](#page-7-0) 7b). However, most pathways in [Figure](#page-7-0) 7b still show relatively low barriers (models 1a, 1b, and 1c with barriers of 535, 502, and 522 meV). A similar trend is found for $Li₃TaO₄$ as depicted in [Figure](#page-7-0) [7](#page-7-0)c. In both $Li₃NbO₄$ and $Li₃TaO₄$, specific arrangement of Li vacancies can lead to a sudden increase in the migration barriers, even beyond 1200 meV (see [Figure](#page-7-0) 7b,c).

This variation in the Li migration barriers is introduced by the immediate crystallographic environment of niobates (tantalates) and the proximity of the Li migration event to the complex antisite defect. In $LiMO₃$ (M = Nb, Ta), the NEB barriers for Li ions migrating in the presence of a defect complex vary slightly.

Figure 7. Li-ion migration barriers computed with NEB@r²SCAN in (a) LiNbO₃, (b) Li₃NbO₄, (c) LiTaO₃, and (d) Li₃TaO₄ supercell structures. In these structures, the environment of Li ions is set in terms of nearest neighbors, NNs [\(Figures](#page-1-0) 1 and [2](#page-2-0)). Filled bars show migration barriers computed in the low-vacancy limit (incorporating one Vac_{Li} per supercell). Hatched bars show migration barriers of models incorporating neutral complex antisitevacancy defects (4Vac_{Li} + M_{Li}). Bars with crossed patterns show migration barriers for LiMO₃ in the presence of the antisite defect complex, with all Li vacancies near the migration pathways and indicated as $4\text{Vac}_{\text{Li}}^{\triangle} + \text{M}_{\text{Li}}$. Configurations of all pathways are provided in [Figures](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S11−S13.

Figure 8. Arrhenius plots of Li-ion conductivities in (a) Li₃NbO₄ and (b) Li₃TaO₄ obtained from MTP-MDs trained on different defect types and concentrations. The temperatures modeled range from 500 to 850 K, except for model 0.5%. Dashed lines are the Arrhenius fits whose migration energies are in [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S13.

This is because the closely packed structures of $LiMO₃$ are very symmetric and structural changes introduced by point defects cause minimal variations in the migration environment and hence the Li-ion migrating barriers. In $Li₃MO₄$, the migration barriers of a Li vacancy (low-vacancy limit model) appear very sensitive to the migration environment as reflected in [Figure](#page-7-0) [7](#page-7-0)b,d. In the structures of $Li₃NbO₄$ (tetramer of Nb octahedra) and $Li₃TaO₄$ (zigzag chain of Ta octahedra) of [Figures](#page-1-0) 1b and [2](#page-2-0), the high-valence Nb^{5+} and Ta^{5+} ions arrange sparsely, in a way to provide additional voids enabling facile Li⁺ migration. In addition to the local environment, the introduction of the antisite defect complex will affect the Li migration pathways by either reducing the $Li⁺-Li⁺$ repulsion near the 4 Li vacancies or increasing the repulsion for pairs Li^+ – M^{5+}_{Li} .

So far, the computed migration barriers of [Figure](#page-7-0) 7, $Li₃MO₄$ (M = Nb, Ta), suggest facile Li-ion transport compared to $LiMO₃$. To gain more insight into the complex behavior of Liion transport in $Li₃MO₄$ (M = Nb, Ta), we perform machinelearning molecular dynamics (MDs) simulations based on the moment tensor potential (MTP) method. 55 In the MTP, we explore the effect of different concentrations of $\text{Vac}_{1,i}$ set to 0.5, 1, 2, and 4%. As calculated in [Section](#page-4-0) 4, the $\Delta E_{\rm D}$ values of neutral defect complexes in $Li₃MO₄$ are comparable with $LiMO₃$. Here, we only considered the case of 4% Li vacancies and 1% M antisites in $Li₃MO₄$, with the MTP-MD simulations performed in the most favorable defect models of $Li₃NbO₄$ and $Li₃TaO₄$ (see [Figures](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S12c and S13e).

[Figure](#page-7-0) 8 shows the Li-ion transport properties from MTP-MD simulations. Predicted migration barriers can be derived from the Arrhenius fits of [Figure](#page-7-0) 8, shown in [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S13. In [Figure](#page-7-0) 8, by including 0.5% Li vacancies in Li₃NbO₄, we achieved migration barriers ∼622 meV, in agreement with experimental values (∼580–850 meV) of crystalline Li₃NbO₄ and NEB barriers of [Figure](#page-7-0) 7.^{[19,29](#page-11-0),[56](#page-12-0)} Unsurprisingly, as the concentration of Li vacancies in $Li₃NbO₄$ increases, the inferred migration barriers of Li_3NbO_4 of 2 and 4% Li vacancies decrease significantly to ∼400 meV. Likewise, the Li-ion conductivities increase greatly by three to five orders of magnitude by increasing the Vac_{Li} from 0.5 to 4.0%. As shown in [Figure](#page-7-0) 8b, the ionic conductivities of $Li₃TaO₄$ are similar to those of $Li₃NbO₄$. The extrapolated Li migration barrier in $Li₃TaO₄$ with 0.5% Li vacancies is estimated to ∼527 meV, in agreement with the experimental measurements (~550 meV in Li_{2.98}Ta_{1.004}O₄).⁵⁷ Similar to $Li₃NbO₄$, as the concentration of Li vacancies increases, $Li₃TaO₄$ exhibits improved conduction properties.

In the presence of Nb or Ta antisite defect complexes, the Liion conduction of $Li₃MO₄$ of 4% Li vacancies and 1% M antisites is comparable with the ionic conductivity predicted in a model system with 4% Li vacancies. These results show that the M_{Li} antisite defects may not significantly affect Li-ion transport in $Li₃MO₄$, but M_{Li} defects ensure the availability of Li vacancies. The migration barriers of [Figure](#page-7-0) 7 support these observations.

6. DISCUSSION

To improve the cathode−electrolyte interfacial stability, extend the oxidation potential of electrolytes, and increase the longevity of lithium-ion batteries, coating materials are usually applied to positive electrode materials.² Commonly used coating materials are niobates and tantalates.^{[9](#page-11-0)}

Niobate coating materials are typically synthesized using wetchemical protocols. In these syntheses (see [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S1), the temperature of calcination mainly affects the crystallinity of $LiNbO₃$ and the distribution of $LiNbO₃$ with secondary phases,

as revealed by X-ray diffraction and transmission electron microscopy experiments.^{[15](#page-11-0)−[17](#page-11-0)}

As the temperature increases, up to four "stages" of calcination may be involved in the fabrication of these Li−Nb−O materials ([Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S1), 10,16,28,36 10,16,28,36 10,16,28,36 10,16,28,36 10,16,28,36 which are as follows.

- 1. At low calcination temperatures (<400 $^{\circ}$ C), there only exists an amorphous $LiNbO₃$ phase, which display high Li-ion conductivities $(10^{-5}-10^{-6} \text{ S cm}^{-1})$.
- 2. At intermediate temperatures (400−650 °C), the crystalline $LiNbO₃$ phase coexists with the disordered rock-salt $Li₃NbO₄$ phase.
- 3. At high temperatures (650−750 °C), a mixture of the crystalline $LiNbO₃$ phase with the ordered $I43mLi₃NbO₄$ phase is identified.
- 4. At temperature exceeding 750 $^{\circ}$ C, the niobate coating is composed of a single ordered $I\overline{4}3m$ Li₃NbO₄ phase. Therefore, at this stage, all of the $LiNbO₃$ phase is transformed into $Li₃NbO₄$.

The conversion temperatures between the Li−Nb−O phases mentioned above can have slight variations when the amount of precursors used in the synthesis and the calcination time vary. As an example, in the recent work of Xin et al., 28 28 28 the niobate coating material contains only Li₃NbO₄ when calcinated for 3 h at ~690 °C. In contrast, Zhang et al.^{[15](#page-11-0)} characterized that a Li–Nb−O coating still displays crystalline $LiNbO₃$ and $Li₃NbO₄$ phases after calcination at ∼700 °C for 10 h. This analysis suggests that niobate coating materials, commonly accepted as "LiNbO $_3$ ", may coexist with Li-rich materials, such as $Li₃NbO₄$ (see [Figures](#page-3-0) [3](#page-3-0)a and [4a](#page-3-0)). From our simulations, we extend these findings to tantalate coatings that may contain varying mixtures of $LiTaO₃$ and $Li₃TaO₄$.

From this investigation emerged that selected calcination temperatures control the coexistence of $LiMO₃$ and $Li₃MO₄$ (M = Nb, and Ta) phases in the coating materials. The concurrence of $LiMO₃$ and $Li₃MO₄$ phases can promote the formation of defects within $Li₃MO₄$ with promising ionic conductivities. Concomitantly, the presence of $LiMO₃$ extends the oxidative resistance to higher voltages, improving the electrochemical performance of batteries. Nevertheless, if calcination temperatures exceed 750 \degree C, LiMO₃ will be fully depleted, and the exceeding Li₃MO₄ will lower the onset voltage for oxidation to \sim 3.6 V vs Li/Li⁺.

The coexistence of $LiMO₃$ and $Li₃MO₄$ suggests that the functional properties of these coating materials, such as, (i) the Li-ion transport in these phases, (ii) the chemical and electrochemical stabilities (under an external bias), and (iii) the mechanical properties (not investigated here) are broadly linked to the multiphased nature of these materials. We built robust thermodynamic and kinetic models using state-of-the-art first-principles calculations to link the multiphased nature of niobates (tantalates) with their functional properties.

6.1. Intrinsic Defects in Coating Materials. Using DFT (at the r 2 SCAN level of theory), we verified the natural occurrence of intrinsic defects in these materials. The equilibrium of $LiNbO₃$ and $Li₃NbO₄$ in niobates will set a Lipoor (more oxidative) condition, highly favoring the formation of a neutral antisite defect complex in $Li₃NbO₄$, implying the simultaneous formation of four Li vacancies and a charged $\mathrm{Nb}_{\mathrm{Li}}^{\bullet\bullet\bullet\bullet}$ antisite defect, i.e., $[4\mathrm{Vac}_{\mathrm{Li}}^{\prime}+\mathrm{Nb}_{\mathrm{Li}}^{\bullet\bullet\bullet\bullet}]$ $(\mathrm{Tables}\ 2\ \text{and}\ 810).$ We arrived at similar findings in $Li₃TaO₄$, suggesting that the coexistence of $LiTaO_3$ and Li_3TaO_4 (Li-poor condition) will

favor the formation of a neutral defect complex in $Li₃TaO₄$, i.e., $[4\text{Vac}_{\text{Li}}' + \text{Ta}_{\text{Li}}^{\bullet\bullet\bullet\bullet}]$.

In general, in $LiNbO₃$, our DFT defect formation energies suggest that the neutral defect complex, i.e., $[4\text{Vac}_{\text{Li}}^{\prime} + \text{Nb}_{\text{Li}}^{\bullet\bullet\bullet\bullet}]$, is highly favored in the Li-poor condition. Isolated Li vacancies and $Nb_{Li}^{\bullet\bullet\bullet\bullet}$ antisites are also likely to form in $LiNbO_3$, with concentrations ~1.39 \times 10¹⁹ and ~3.46 \times 10¹⁸ cm⁻³, respectively, at 1000 K. Characterization with neutron and Xray diffraction, $22,47$ $22,47$ $22,47$ and later solid-state nuclear magnetic resonance^{[48](#page-12-0)} on LiNbO₃ suggested that ~1% of the Li sites are occupied by Nb, and ∼4% of the Li sites are vacant, hence supporting the existence of a neutral antisite defect complex and in excellent agreement with our computational models. While at the cathode interface, exchange of Nb (Ta) from the coating layer with transition metals of the positive electrode is possible;^{[58](#page-12-0)} here, we did not investigate the occurrence of these antisite defects.

6.2. Li-Ion Transport in Niobate and Tantalate Coatings. Subsequently, we inspected the properties of Li-ion transport (with SoftBV, NEB, and MTP-MD simulations) in the presence of defect types favored by the thermodynamic conditions in $LiNbO₃$ and $Li₃NbO₄$, and $LiTaO₃$ and $Li₃TaO₄$. In this work, we did not focus on simulating the behavior of amorphous niobates and tantalates. However, experimental investigations by Rahn et al. 26 26 26 suggested that amorphous $LiNbO₃$ synthesized in Li-poor (more oxidative) conditions exhibits better Li-ion transport $(E_a = 700 \text{ meV})$ than amorphous LiNbO₃ (E_a = 830 meV) synthesized in Li-rich conditions. This observation matches our computational results, with $LiNbO₃$ and $Li₃NbO₄$ likely to be highly defective in Lipoor conditions.

Clearly, coating materials are effective at migrating the Li-ion if the migration barriers are within some prescribed tolerance values. Previously, the maximum tolerable migration barriers (E_a^{max}) for Li ions in coating materials were estimated using the model proposed by Chen et al.,^{[59](#page-12-0)} suggesting that coating layers of different thicknesses are only viable when the Li migration barriers are smaller than a maximum value $E_{\rm a}^{\rm max}$. Based on manufacturing data of crystalline Li−Nb−O and Li−Ta−O coatings ([Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S1), we identified "reasonable" operating conditions to derive a value of $E_{\rm a}^{\rm max}$, which set a 10 nm coating thickness, operating at 25 °C, and at a discharge rate of 1C (1 h). This yields to $E_{\rm a}^{\rm max}$ of ~700 meV.

In crystalline $LiNbO₃$ and $LiTaO₃$, our data suggest that the migration of Li vacancies (in the dilute regimes) remains high (>1100 meV) with or without the presence of $[4\text{Vac}_{\text{Li}}' + \text{M}^{\bullet\bullet\bullet}_{\text{Li}}]$ in the materials. These values feature consistently above the maximum tolerable barriers, ∼700 meV for these coating materials. However, experimentally amorphous $LiNbO₃$ and $LiTaO₃$ both appear decent Li-ion conductors at room temperature ($\sim\!10^{-5}\!\!-\!10^{-6}\,\mathrm{S\ cm^{-1}}$). Therefore, future computational studies are required to elucidate the Li-ion transport properties of these amorphous phases.

The computed migration barriers in the crystalline $Li₃NbO₄$ (or $Li₃TaO₄$) phase are significantly lower (mostly <700 meV) than crystalline LiNbO₃ (or LiTaO₃). Li-ion conductivities obtained with machine-learned MDs also corroborate this finding. We claim that $Li₃NbO₄$ or $Li₃TaO₄$ are the active components in niobate or tantalate coating materials. Therefore, synthesis protocols and defect engineering remain crucial to the overall properties of these coating materials.

6.3. Low-Oxidative Stability Limits of LiMO₃ and Li₃MO₄. The high operating voltages \geq 4.2 V vs. Li/Li⁺ delivered

by high nickel content layered materials, such as Li- $\text{Ni}_{0.8}\text{Co}_{0.1}\text{Mn}_{0.1}\text{O}_2$ (NMC811), require coating materials with matched oxidative stability windows. In [Table](#page-4-0) 1, we estimated the oxidative (anodic) stabilities of all phases in niobate and tantalate coating materials. As widely applied coating materials, LiNbO₃ and LiTaO₃ show low-oxidative (anodic) limits below 4 V vs. Li/Li^{+} (see [Table](#page-4-0) 1), which are lower than the reported values of some phosphate coatings, such as $Li₃PO₄$ (~4.2 V vs. Li/Li⁺) and LiPO₃ (∼5.0 V).⁴⁴ The predicted anodic limits of $Li₃NbO₄$ and $Li₃TaO₄$ appear slightly lower than their $LiMO₃$ analogues. Indeed, the higher Li-ion transport in crystalline $Li₃NbO₄$ (and $Li₃TaO₄$) phases compared to $LiNbO₃$ (and LiTaO₃) can explain the limiting oxidative stabilities of $Li₃MO₄$ materials. Therefore, the selection of adequate calcination treatments for niobate and tantalate coatings should always ensure the concurrence of the highly conductive $Li₃MO₄$ component and $LiMO₃$ phases with better anodic stabilities.

Furthermore, from our calculations of intrinsic defects in the niobate and tantalate phases, the equilibrium Fermi energies, *E*_{F,eq} (see [Figure](#page-4-0) 5), can be linked to the electronic conductivity of these bulk materials.[53,60](#page-12-0) Across all of the chemical and electrochemical environments (reducing or oxidizing) explored here, the equilibrium Fermi energies of lithium niobates and tantalates are always at least ∼1 eV away from the valence or conduction bands. This indicates that intrinsic defects in lithium niobate and tantalate bulks alone may not impart significant electronic conductivities.

7. CONCLUSIONS

We investigated the structure−property relationship of lithium niobates and tantalates, which are an important class of coating materials for lithium-ion batteries. We underscored the importance of the multiphasic nature of niobate coating materials, $LiNbO₃$ and $Li₃NbO₄$. Similarly, tantalates appear in mixtures of LiTaO₃ and Li₃TaO₄.

We demonstrate that $LiNbO₃$ and $Li₃NbO₄$ may contain high concentrations of lithium vacancies Vac_{Li}' and antisites $\text{Nb}_{\text{Li}}^{\bullet\bullet\bullet}$ arranged into charge-neutral defect complexes, $[4Vac'_{Li} +$ $M_{Li}^{\bullet\bullet\bullet\bullet}$]. Our simulations reveal poor Li-ion transport of crystalline $LiNbO₃$ and $LiTaO₃$ even in the presence of intrinsic defects Vac_{Li} and a neutral antisite defect complex [4Vac_{Li} + $M_{Li}^{\bullet\bullet\bullet\bullet}$]. In contrast, our analysis demonstrates that the secondary phases $Li₃NbO₄$ and $Li₃TaO₄$ are much better ion conductors than $LiNbO₃$ and $LiTaO₃$.

Finally, the low-oxidative (anodic) voltages of $Li₃NbO₄$ and $Li₃TaO₄$ compared to $LiNbO₃$ and $LiTaO₃$ encourage the selection of adequate calcination treatments to ensure the concurrence of both phases.

Our findings stimulate more in-depth investigations on the structure−property relationships of these highly sought coating materials.

8. METHODS

8.1. First-Principles Calculations. Density functional theory (DFT) calculations were performed using the Vienna *ab initio* $Simulation$ Package. 61 The projector augmented wave potentials described the core electrons, and are as follows: Li 17Jan2003 $2s¹$, Nb_pv 08Apr2002 4p⁶ 5s¹ 4d⁴, Ta_pv 07Sep2000 5p⁶ 6s² 5d³, and O $08\overrightarrow{\text{A}p12002}$ $2s^2$ $2p^4$.^{[62](#page-12-0)} The kinetic energy cutoff describing the valence electrons was set to 520 eV. The meta-GGA functional r²SCAN approximates the exchange−correlation energy[.63](#page-12-0) The PBE functional was used for training the machine-learned potentials,^{[64](#page-12-0)} as it is ~10% faster than r^2 SCAN in the *ab initio* molecular dynamics (AIMDs). 5×5 × 5 Γ-centered *k*-point meshes were used to integrate the first Brillouin zone of $LiNbO₃$ and $LiTaO₃$ (with a *R3c* space group) and converged within 10^{-5} eV. k -point meshes of $3\times 3\times 3$ and $4\times 4\times 2$ were applied to Li_3NbO_4 ($\overline{I43}m$) and Li_3TaO_4 ($C2/c$). In larger supercells and metals (Li, Nb, and Ta), the *k*-point meshes were optimized. With the exception of defect calculations, volumes, shapes, and coordinates of each structure were optimized until the interatomic forces were $\leq 10^{-2}$ eV Å[−]¹ .

All crystal structures were taken from the Inorganic Chemical Structure Database. Disordered rock-salt structures of $Li₃NbO₄$ and Li3TaO4 were ordered by computing all possible Li/Nb (Li/Ta) arrangements at given supercell sizes.^{[65](#page-12-0)} O₂ gas was used as a reference for the oxygen $\mu_{\rm O}$ and was simulated by placing an oxygen dimer in a 10 \times 10 \times 10 Å³ box. We included O₂ vibrational contributions (~100 meV at 0 K).

As shown in [Tables](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S2 and S3, the DFT-calculated structures of LiNbO₃ (*R*3*c*), LiTaO₃ (*R*3*c*), Li₃NbO₄ (*I*43*m*), and *β*-Li₃TaO₄ (*C*2/ ϵ), using the meta-GGA r²SCAN exchange and correlation functional, are in much better agreement with existing experimental data on the lattice parameters and electronic band structures.^{40,6}

8.2. Defect Formation Energies. The following intrinsic point defects were studied: Li vacancy (Vac_{Li}) and M-Li antisite ($M_{L,i}$, M = Nb, Ta). We also investigated charge-neutral defect complexes $[4\text{Vac}_{\text{Li}}]$ + M_{Li}^{open}], $M = Nb$, Ta. Defects in LiNbO₃, LiTaO₃, Li₃NbO₄, and Li₃TaO₄ were modeled by large supercells containing 320, 320, 256, and 256 atoms, respectively. The defect formation energies $\Delta E_{\text{D}}(X^q)$ are calculated from eq 2

$$
\Delta E_{\rm D}(X^q) = E_{\rm defect}^{\rm tot} - E_{\rm bulk}^{\rm tot} - \sum_{i} n_i \mu_i + q E_{\rm Fermi} + E_{\rm corr}
$$
(2)

where $E_{\rm defect}^{\rm tot}$ and $E_{\rm bulk}^{\rm tot}$ are the r²SCAN energies of supercells with and without defect. μ_i is the chemical potential of species *i*, while n_i is the number of atoms of *i* added to $(n_i > 0)$ or removed from $(n_i < 0)$ the supercell model. μ _{Li} and μ _M were derived from the phase diagram of Li− Nb−O and Li−Ta−O [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S5). We explored defects with charges *q*: [−1, 0, and 1] for Vac_{Li} and [−1, 0, 1, 2, 3, and 4] for M_{Li}. The *E*_{Fermi} spans the r²SCAN band gaps of the respective bulk structures. $E_{\rm corr}$ corrects for the electrostatic energy of charged defects interacting with their periodic images and the potential alignment for the fictitious jellium background.^{[69](#page-12-0),[70](#page-12-0)} The correction of ref 70 was used for lowersymmetry $\rm LiNbO_{3}$, $\rm Li_{3}TaO_{3}$, and $\rm Li_{3}TaO_{4}$ ^{[70](#page-12-0)} whereas we implemented ref [69](#page-12-0) for Li₃NbO₄. The band structures computed with different GGA (PBE and PBEsol) and meta-GGA (SCAN and r 2 SCAN) exchange and correlation functionals are shown in [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S4. Details about the computation of defect concentrations and the identification of equilibrium Fermi energies are in refs [53](#page-12-0), [60](#page-12-0).

8.3. Li-Ion Migration in LiMO3 and Li3MO4 (M = Nb, Ta). The empirical softBV analysis was used to identify the topology of Li-ion diffusion in LiMO_3 and Li_3MO_4 .^{[71](#page-12-0)}

In the nudged elastic band (NEB) calculations, we used models as detailed in Section 8.2, which ensure a minimum distance ≥ 8 Å between image replicas. 59,72 59,72 59,72 59,72 59,72 In NEBs, spring forces were set to 5 eV Å $^{-1}\!$, and charge neutrality was imposed through a compensating background charge. Models of neutral defect complexes with the lowest formation energies were selected from Section 8.2 used in NEBs.

To fit the moment tensor potentials (MTPs), the training sets were generated with *ab initio* molecular dynamics (AIMD) simulations on 2 \times 2 \times 2 supercells (256 atoms) of Li₃MO₄, which are geometrically optimized by DFT using the r 2 SCAN functional. An NVT ensemble based on the Nosé−Hoover thermostat, and a time step of 0.5 fs were applied in the AIMDs.⁷³ A plane wave energy cutoff of the AIMDs was 400 eV. The total energy was integrated at the Γ point.⁷⁴

For $Li₃NbO₄$ AIMDs, we performed simulations at 600, 800, 1000, and 1200 K, each lasting 8 ps for the equilibration step, and after a temperature ramping of 0.5 ps. In total, 8000 snapshots of the last 1 ps were taken as the training sets.⁷⁴ While for Li_3TaO_4 , the temperature range was 600−1500 K with a step of 300 K. We investigate the role of defects on Li diffusion in ${\rm Li}_3{\rm MO}_4$ by considering two types of defects, i.e., the Li vacancy (Vac_{Li}) and the defect complex (4Vac_{Li} + Nb_{Li} (Ta_{Li})). In the model of Li vacancy, a Li vacancy was introduced into a supercell of 256 atoms (a concentration of ∼1%) and compensated

with a background charge. We replaced one Li atom with an Nb (Ta) atom for complex antisite-vacancy defects and created four Li vacancies based on the most stable models calculated from Section 8.2, with concentrations of antisites of ∼1% and Li vacancies of ∼4%.

In training the MTPs, we chose the radius cutoff of 5 Å, and a maximum level of 12 controlling the completeness of basis functions during the training of the bulk structures and the defective structures with 0.5, 1, and 2% Li vacancies.^{[74](#page-12-0),[75](#page-13-0)} For structures with 4% Li vacancies or with the antisite defect complex, we increased R_{cut} to 6 Å to better capture the structural complexity of this situation. Weights on energies, forces, and stresses were set to 100:10:1. The accuracy of the MTP fittings is reported in [Table](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S12.

MTP molecular dynamics (MTP-MD) simulations were conducted in LAMMPS^{[76](#page-13-0)} using the NVT ensemble (Nosé–Hoover).⁷³ We carried out long MDs of $Li₃MO₄$ for 10 ns with a time step of 1 fs, preceded by a temperature ramping of 10 ps, followed by 1 ns of equilibration. MDs used $4 \times 4 \times 4$ supercells with 2048 atoms. By varying the number of removed Li atoms, we computed the Li-ion conductivity of $Li₃MO₄$ structures including Vac_{Li} concentrations of 0.5, 1, 2, and 4%. By monitoring the mean-square displacements (see [Figure](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf) S14) of Li⁺ in MTP-MDs, we extracted the tracer diffusivity of eq 3

$$
D^*(T) = \lim_{t \to \infty} \frac{1}{2dt} \left(\frac{1}{N} \sum_{i=1}^N \left\langle [\vec{r}_i(t) - \vec{r}_i(0)]^2 \right\rangle \right) \tag{3}
$$

where $d = 3$, N is the number of lithium ions, $\vec{r}_i(t)$ is the displacement of *i*th Li at time *t*. The activation energy of Li-ion migration (E_a) was derived from the Arrhenius equation

$$
D^*(T) = D_0 \exp\left(-\frac{E_a}{k_B T}\right) \tag{4}
$$

where D_0 is the prefactor, *T* is the temperature, and k_B is the Boltzmann constant. Li-ion conductivities $\sigma(T)$ are extracted from the Nernst− Einstein equation

$$
\sigma(T) = \frac{nz^2e^2}{k_B T} D^*(T) \tag{5}
$$

where *n* is the volume density of the Li^+ , $z = +1$, and *e* is the electron charge.

■ **ASSOCIATED CONTENT**

\bullet Supporting Information

The Supporting Information is available free of charge at [https://pubs.acs.org/doi/10.1021/acs.chemmater.3c01197](https://pubs.acs.org/doi/10.1021/acs.chemmater.3c01197?goto=supporting-info).

(i) Synthesis and applications of lithium niobate and tantalate coating materials; (ii) structures of lithium niobate and tantalate compounds; (iii) analysis of intrinsic defects in lithium niobate and tantalate compounds; (iv) Li-ion migration barriers and migration paths vs. niobate or tantalate local environments; and (v) migration barriers extracted from moment tensor potential molecular dynamics [\(PDF](https://pubs.acs.org/doi/suppl/10.1021/acs.chemmater.3c01197/suppl_file/cm3c01197_si_001.pdf))

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Notes

The authors declare no competing financial interest.

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