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Probing Capacity Trends in MLi₂Ti₆O₁₄ Lithium-Ion Battery Anodes Using Calorimetric Studies

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ABSTRACT: Due to higher packing density, lower working potential, and area specific impedance, the $MLi_2Ti_6O_{14}$ (M = 2Na, Sr, Ba, and Pb) titanate family is a potential alternative to zero-strain $Li_4Ti_5O_{12}$ anodes used commercially in Li-ion batteries. However, the exact lithiation mechanism in these compounds remains unclear. Despite its structural similarity, $MLi_2Ti_6O_{14}$ behaves differently depending on charge and size of the metal ion, hosting 1.3, 2.7, 2.9, and 4.4 Li per formula unit, giving charge capacity values from 60 to 160 mAh/g in contrast to the theoretical capacity trend. However, high-temperature oxide melt solution calorimetry measurements confirm strong correlation between thermodynamic stability and the observed capacity. The main factors controlling energetics are



strong acid–base interactions between basic oxides MO, Li_2O and acidic TiO_2 , size of the cation, and compressive strain. Accordingly, the energetic stability diminishes in the order $Na_2Li_2Ti_6O_{14} > BaLi_2Ti_6O_{14} > SrLi_2Ti_6O_{14} > PbLi_2Ti_6O_{14}$. This sequence is similar to that in many other oxide systems. This work exhibits that thermodynamic systematics can serve as guidelines for the choice of composition for building better batteries.

INTRODUCTION

Lithium-ion batteries (LIBs) are the currently favored power sources for portable applications such as electric vehicles and electronic products.¹ Most commercial LIBs work via reversible intercalation² of lithium ions between a (mixed) transition metal oxide cathode and a layered graphite anode. For more than 30 years since the inception of LIBs by SONY (Tokyo, Japan) in the 1990s, graphite remains the most widely used and optimized anode material due to its high theoretical capacity (380 mAh/g) and low Li insertion voltage (ca. 0.2 V) while maintaining an intercalation mechanism, which guarantees structural integrity and long cycle life. Nevertheless, the graphite anode has several issues, such as solvent-dependent stability,^{4,5} large volume change, large irreversible capacity loss (ICL) from solid electrolyte interphase (SEI) formation (below 0.8 V),⁶ and fire risk due to lithium plating during overcharge or higher currents' some of which directly stem from its low voltage performance.

In this context, the zero-strain spinel $Li_4Ti_5O_{12}$ anode operating at flat and higher voltage (1.55 V vs Li, 175 mAh/g) was introduced^{8,9} and is commercially preferred¹⁰ over graphite in applications requiring fast charging and safety. Additionally, $Li_5Cr_7Ti_6O_{25}$ similar to $Li_4Ti_{15}O_{12}$ spinel synthesized at 800 °C exhibits large charge capacity, good cycle stability, superior rate performance, and outstanding electrochemical kinetics. When cycled at 200 mA/g, it displays a high initial reversible capacity of 146.6 mAh/g and retains a considerable capacity of 130.8 mAh/g after 300 cycles.¹¹ Among other Ti-based electrode materials,^{12–14} MLi₂Ti₆O₁₄ (M = 2Na, Sr, Ba, and Pb)^{15–17} shows lower voltage (1.28– 1.42 V vs Li), low area specific impedance,¹⁶ and higher packing density¹⁸ than Li₄Ti₅O₁₂ and has a high theoretical capacity (220–282 mAh/g) for redox activity of all six Ti (Figure 1i).¹⁹ The MLi₂Ti₆O₁₄ series exhibit open channels enabling reversible Li⁺ insertion in vacant sites like 8*c*, 4*a*, and 4*b* of the orthorhombic framework (*Cmca*). However, the observed capacity trend in this series does not follow the theoretical capacity trend. Previous studies identified crystallographic origins underlying the low capacity of the monovalent member (M = 2Na) vs the divalent members (M = Ba, Sr, and Pb).^{16,17} However, currently there is no explanation of the anomalous trend of the observed capacity in the isostructural divalent members—BaLi₂Ti₆O₁₄ (2.7 Li) < SrLi₂Ti₆O₁₄ (2.9 Li) < PbLi₂Ti₆O₁₄ (4.4 Li)—where PbLi₂Ti₆O₁₄ is opposite to the theoretical capacity trend.

We take a new approach to understand the electrochemical property trend in this isostructural $MLi_2Ti_6O_{14}$ series from their relative thermodynamic stabilities, attempting to elucidate the structure–property relationships in this family of materials. High-temperature oxide melt solution calorimetry is an established methodology to measure the thermodynamic stabilities of oxide and nonoxide systems.^{20–22} Recently, Abramchuk et al.²³ used this method to investigate the thermodynamic stability and its role in the mechanism of ionic

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Figure 1. (a–d) Rietveld refinement of XRD (Cu K α) patterns of members of the MLi₂Ti₆O₁₄ family (M = 2Na, Sr, Ba, and Pb) formed by solution combustion (900 °C, 2 h). Experimental data points (gray dots), calculated pattern (black), their difference (blue), and Bragg reflections (black bars) of the respective isostructural phases are shown. (e–h) SEM micrographs of the combustion synthesized MLi₂Ti₆O₁₄ anode indicating porous structure of secondary agglomerates. The inset shows a TEM image of primary nanometric particles (200 nm scale). (i) Structural illustration of the MLi₂Ti₆O₁₄ anodes. (j–m) Galvanostatic (dis)charge voltage profiles and (n–q) differential capacity plots depicting the electrochemical performance of the MLi₂Ti₆O₁₄ anodes.

transport in the NASICON-type solid-state electrolyte $Li_{1+x}Al_xTi_{2-x}(PO_4)_3$. They demonstrated a strong correlation between the energetic stability and ionic conductivity in LATPs with a pronounced change in the trends with composition. In the present work, we employ high-temperature oxide melt solution calorimetry to systematically assess the thermodynamic stability of $MLi_2Ti_6O_{14}$ (M = 2Na, Sr, Ba, and Pb), which provides insights into observed differences in the capacity. We found a strong correlation between the thermodynamic stability and reversible lithium insertion in $MLi_2Ti_6O_{14}$ anode materials and explain the trend in observed capacity. This work reflects the acid—base energetic contributions in this class of anode materials.

EXPERIMENTAL SECTION

Synthesis and Electrochemistry. $MLi_2Ti_6O_{14}$ was synthesized by a two-step solution combustion method. Metal nitrates as oxidizers (O) and glycine as fuel (F) were taken in a 1:1 ratio to ensure maximum exothermicity. Lithium nitrate (LiNO₃, Sigma-Aldrich, 99%), sodium nitrate (NaNO₃, SD Fine, 99.5%), barium nitrate (Ba(NO₃)₂, Sigma-Aldrich, >99%), lead nitrate (Pb(NO₃)₂, SD Fine, 99.5%), and strontium nitrate (Sr(NO₃)₂, Sigma-Aldrich, 99%) were used. Titanium (Ti⁴⁺) ions were sourced from freshly prepared titanyl nitrate (TiO(NO₃)₂ as it is unstable).²⁴ This was made by simultaneous hydrolysis—nitration of titanium isopropoxide (C₁₂H₂₈O₄Ti, Sigma-Aldrich, 97%) in isopropyl alcohol (Merck, analytical grade 99.7%) under ice-cold conditions. The mixture was heated to less than 150 °C to remove water and form a thick solution, which was reheated at 500 °C to trigger combustion. The resulting white intermediate powder was ground, pelletized, and annealed in air at 900 °C for 1 min to 2 h. The phase purity of combustion prepared MLi₂Ti₆O₁₄ samples was verified from powder X-ray diffraction patterns (2θ range = 10–50°, step size = 0.026° s⁻¹) obtained using a PANalytical X'Pert Pro X-ray diffractometer having a Cu-K α source ($\lambda_1 = 1.54$ Å). Rietveld refinement was done using the GSAS program.²⁵ The structures were illustrated using the VESTA software.²⁶

The surface morphology of the powders was characterized using a scanning electron microscope having a LaB₆ field emission source (Carl Zeiss ULTRA55 FESEM, 5 kV) and using a transmission electron microscope (FEI Tecnai F20 STwin, 200 kV). TG-DSC measurements were performed using a Setaram LABSYS EVO instrument. 25 mg of the powdered sample was pressed into a pellet, placed in a Pt crucible, and heated from 30 to 800 °C (10 °C/min) under a N_2 flow (20 mL/min). A buoyance correction was performed with an empty crucible prior to the experiment. The pristine powder was ground with carbon (Super P, Sigma-Aldrich, 99+ %) and a polyvinylidene fluoride (PVDF) binder in an 80:10:10 or 70:20:10 weight ratio. This electrode powder was mixed with N-methyl-2-pyrrolidone (Sigma-Aldrich, >99%) solvent and drop cast onto prewashed Cu coupons. The electrodes were dried at 80 °C overnight in a vacuum oven and were used for assembling CR2032 coin cells vs lithium (Sigma-Aldrich, >99%) in an Ar-filled glovebox (MBraun LABStar GmbH). Whatman GF/C glass fiber was used as the separator and 1 M LiPF₆ in 1:1:3 v/v % of ethylene carbonate/propylene carbonate/dimethyl carbonate (EC/PC/DMC) (chameleon reagent) was used as the electrolyte. The cells were cycled between 2 V to 1 or 0.5 V at C/20 and 50 mA/g current using a Neware BTS-4000 battery cycler to ensure we reach the maximum specific capacity values for each of the materials, which can then be rationalized with the thermodynamic trend.

High-Temperature Oxide Melt Solution Calorimetry. High-temperature oxide melt solution calorimetry was performed using a Setaram AlexSYS Tian-Calvet twin microcalorimeter using methods standard to our laboratory and described previously.²⁰⁻²² This calorimeter allows the direct determination of the enthalpy of formation $(\Delta H^{\circ}_{f,ox})$ of multicomponent compounds from their binary oxides. In a typical experiment, ~ 5 mg of the MLi₂Ti₆O₁₄ sample was pelletized and dropped from ambient temperature into a calorimeter at 800 °C containing the solvent molten sodium molybdate $(3Na_2O\cdot 4MoO_3)$ in a platinum crucible. The measured enthalpy of drop solution (ΔH_{ds}) is the sum of the sample heat content from ambient temperature to 800 $^\circ C$ and its heat of solution in the solvent at 800 °C. At least 8-10 experiments were done per sample, and the results are reported as average values with error being two standard deviations of the mean. The calorimetry glassware was flushed with oxygen gas at a flow rate of 65 mL/min to maintain a constant atmosphere, and the solvent was bubbled with the same gas at 5 mL/min to remove liberated evolved gases, aid dissolution, and prevent local saturation of the solvent. The calorimeter was calibrated using the heat content of 5 mg of pellets of α - Al_2O_3 (99.997%). The details of the calorimeter and procedures have been described previously.²⁰⁻²²

RESULTS AND DISCUSSION

Structure and Electrochemical Performance. The combustion synthesized MLi₂Ti₆O₁₄ powders were found to be phase pure from Rietveld refinement (Figure 1a-d and Tables S1-S4, Supporting Information). The powders were well crystallized into primary particles of about 1 micron, which agglomerated to a porous network (Figure 1e,f). The crystal structure is orthorhombic (symmetry: Cmca), containing a three-dimensional network of TiO₆ octahedral units. Two edge-shared units along the b-axis stitch the infinite c-axis directed edge-shared units in a ladderlike way. The ladders form a sheet in the *bc* plane. In the sheet, inside each *c*-directed ladder, there is an empty tunnel where structural lithium ions sit in tetrahedral sites with adjacent octahedral sites free. This sheet is three units thick along the *a*-axis and is glued to other sheets via larger M ions which populate the interlayer partially (M = Sr, Ba, and Pb) or completely (M = 2Na). One unit of MLi₂Ti₆O₁₄ can host up to six lithium atoms, for complete reduction of Ti4+ to Ti3+, offering a theoretical capacity of about 240 mAh/g depending on the atomic weight of M (Table 1). The mechanism of lithium insertion is unclear, and only about 75% of theoretical capacity has been attained.

Table 1. Space Group, Charge Capacity, and Li Uptake per Formula Unit of the $MLi_2Ti_6O_{14}$ Titanates

anode material	space group	charge capacity (mAh/g)	operating voltage (V) vs Li	Li uptake (per f.u.)
Na2Li2Ti6O14	Стса	62	1.28	1.32
SrLi ₂ Ti ₆ O ₁₄	Стса	111	1.42	2.74
BaLi ₂ Ti ₆ O ₁₄	Стса	125	1.38	2.85
$PbLi_{2}Ti_{6}O_{14} \\$	Стса	162	1.35	4.40

 $Na_2Li_2Ti_6O_{14}$ is the lightest among $MLi_2Ti_6O_{14}$ (M = 2Na, Sr, Ba, and Pb) and so it is expected to have the highest theoretical capacity of 282 mAh/g for six Ti redox (Figure 1i). However, due to the geometrical constraints, it cannot insert more than two Li (62 mAh/g, 1.3 Li) as the vacant 11 coordinated interlayer sites are fully filled with two monovalent Na atoms, compared to the half-filled interlayer by one divalent M atom (M = Ba, Sr, and Pb) in the remaining members (enabling them to incorporate up to 4.4 Li) (Figure 1j-m). Furthermore, even the divalent counterparts with half-filled interlayers are space limited as they reversibly allow only four lithium ions (instead of six) via reduction of four out of six Ti ions. Lithium ions (de)insert from 1.2 to 1.5 V in the system, with each member showing its signature (de)insertion voltage (Figure 1n-q). The observed charge capacity is in the order BaLi₂Ti₆O₁₄ (111 mAh/g, 2.7 Li) < SrLi₂Ti₆O₁₄ (125 mAh/g, 2.9 Li) < PbLi₂Ti₆O₁₄ (162 mAh/g, 4.4 Li). While a slightly higher observable capacity in the Sr analog compared to Ba is in line with the trend in their theoretical capacity, the reason why PbLi₂Ti₆O₁₄ shows the largest capacity and uptakes significantly more lithium despite being the heaviest is highly intriguing. To address this question, we embark on investigating the thermodynamic stability in this family of compounds.

Thermodynamic Studies. Thermochemical data are sensitive to the crystal structure, phase, and chemical purity of the sample. PXRD and TG-DSC measurements were performed before the calorimetric measurement to detect any possible contamination/decomposition and check for adsorbed water on the sample due to exposure in the ambient atmosphere prior to calorimetric experiments. Samples were phase pure as shown in Figure 1a-d, and TGA measurements detected no mass loss, confirming there was no adsorbed water on the samples.

The drop solution enthalpy (ΔH_{ds}) and formation enthalpy from oxides $(\Delta H^{\circ}_{f,ox})$ of MLi₂Ti₆O₁₄ (M = 2Na, Sr, Ba, and Pb) as a function of Li uptake per unit formula are shown in Table 2 and Figure 2a,b. The enthalpy of drop solution of

Table 2. Drop Solution Enthalpies (ΔH_{ds}) in $3Na_2O\cdot 4MoO_3$ at 800 °C and Formation Enthalpies ($\Delta H^{\circ}_{f,ox}$) at 25 °C of $MLi_2Ti_6O_{14}$ (M = 2Na, Sr, Ba, and Pb) Titanates

sample	$\Delta H_{\rm ds}$ (kJ/mol)	$\Delta H^{\circ}_{\rm f,ox}$ (kJ/mol)
$Na_2Li_2Ti_6O_{14}\\$	$502.41 \pm 0.35(8)$	-333.32 ± 4.91
BaLi ₂ Ti ₆ O ₁₄	$506.75 \pm 0.57(8)$	-323.54 ± 3.44
SrLi2Ti6O14	$498.16 \pm 0.68(8)$	-260.65 ± 3.15
PbLi ₂ Ti ₆ O ₁₄	$508.46 \pm 0.46(8)$	-157.81 ± 2.54
Li ₂ O	-77.21 ± 2.44^{23}	
Na ₂ O	-195.90 ± 4.23^{27}	
TiO ₂	$73.70 \pm 0.39(6)^{28}$	
SrO	-127.48 ± 1.84^{a}	
BaO	-181.22 ± 2.32^{a}	
РЬО	$-14.34 \pm 0.38 \ (6)^a$	
CO ₂	37.40	

^{*a*}This work. The value is the mean of the number of experiments indicated in parentheses; two standard deviations are given as errors.

PbO, SrO, and BaO is obtained from the drop solution enthalpy of lead oxide and strontium and barium carbonate (SrCO₃ and BaCO₃) at 800 °C using the thermochemical cycle in Tables S5 and S6 (Supporting Information). The enthalpies of drop solution of Li₂O, Na₂O, and TiO₂ are taken from our previous measurements.^{23,27,28} The formation enthalpy ($\Delta H^{\circ}_{f,ox}$) of MLi₂Ti₆O₁₄ (M = 2Na, Sr, Ba, and Pb) is calculated from the thermochemical cycles in Table 3. The $\Delta H_{\rm ds}$ for lithium titanates is endothermic ranging from 498.16 \pm 0.68 to 508.46 \pm 0.46 kJ/mol, and $\Delta H^{\circ}_{\rm f,ox}$ is exothermic ranging from -333.32 ± 4.91 to -157.81 ± 2.54 kJ/mol, confirming that the these materials are thermodynamically stable. $\Delta H_{\rm ds}$ does not follow any regular trend but $\Delta H^{\circ}_{\rm f,ox}$ is inversely proportional to lithium uptake in MLi₂Ti₆O₁₄ (Figure 2b). The least stable PbLi₂Ti₆O₁₄ ($\Delta H^{\circ}_{\rm f,ox} = -157.81 \pm 2.54$ kJ/mol) intercalates the most lithium (4.4 Li), followed by Sr₂Li₂Ti₆O₁₄ ($\Delta H^{\circ}_{\rm f,ox} = -260.65 \pm 3.15$ kJ/mol, 3.08 Li), then Ba₂Li₂Ti₆O₁₄ ($\Delta H^{\circ}_{\rm f,ox} = -323.54 \pm 3.44$ kJ/mol, 2.47 Li), while the most stable Na₂Li₂Ti₆O₁₄ ($\Delta H^{\circ}_{\rm f,ox} = -333.32 \pm 4.91$ kJ/mol) inserts the least lithium (1.32 Li). Factors such as size of the metal ion, geometrical and compressive strains, and "acid–base" interaction explains their stability trend.

The energetics of ternary oxide formation is dominated by the difference in the acid-base character of the cations. Binary oxides are classified as acidic, basic, or amphoteric. Smith's table³⁰ expresses all metal oxides in terms of acidity and the enthalpies of formation from binary oxides for lithium titanates change linearly with acidity. This validates the trends observed in the formation enthalpies, and in turn the lithium uptake (Figure 2c). The highly exothermic heats of formation reflect strong acid-base interactions among the MO, and Li₂O (basic oxides) with TiO_2 (a somewhat acidic oxide). The basic binary oxides MO and Li₂O interact with the acidic TiO₂ to form the MLi₂Ti₆O₁₄ structure. Alkali metal oxides exhibit greater basic character followed by alkaline earth metal oxides, among the divalent oxides those of larger cations are more basic compared to lighter cations, the order of basicity for MO is $Na_2O > BaO > SrO.^{31}$ Thus, the acid-base interaction between MO, Li₂O, and TiO₂ which stabilizes the structure most strongly in Na₂Li₂Ti₆O₁₄, followed by BaLi₂Ti₆O₁₄, then SrLi₂Ti₆O₁₄. This explains the trend observed in the ΔH°_{fox} $Na_2Li_2Ti_6O_{14} > BaLi_2Ti_6O_{14} > SrLi_2Ti_6O_{14}$. This correlation is consistent with the dominance of the acid-base chemistry in phase stability, as seen in trends of enthalpies of formation in



Figure 2. (a) Dissolution enthalpy (b) formation enthalpy vs reversible Li uptake (per formula unit) in the lithium titanates, and (c) enthalpies of formation from oxides for lithium titanates as a function of acidity based on Smith's scale of acidity for alkali, alkali-earth, and lead oxides.³⁰

reaction	ΔH (kJ/mol)
$MLi_{2}Ti_{6}O_{14(S,25^{\circ}C)} \rightarrow MO_{(soln,25^{\circ}C)} + Li_{2}O_{(soln,25^{\circ}C)} + 6TiO_{2(soln,25^{\circ}C)}$	$\Delta H_1 = \Delta H_{ds} (MLi_2Ti_6O_{14})$
$Li_2O_{(S,25^{\circ}C)} \rightarrow Li_2O_{(soln,25^{\circ}C)}$	$\Delta H_2 = -77.21 \pm 2.44^{23}$
$Na_2O_{(S,25^{\circ}C)} \rightarrow Na_2O_{(soln,25^{\circ}C)}$	$\Delta H_4 = -195.90 \pm 4.23^{27}$
$\text{TiO}_{2(S,25^{\circ}\text{C})} \rightarrow \text{TiO}_{2(\text{soln},25^{\circ}\text{C})}$	$\Delta H_3 = 73.70 \pm 0.39^{28}$
$\text{SrO}_{(s,25^{\circ}\text{C})} \rightarrow \text{SrO}_{(\text{soln},800^{\circ}\text{C})}$	$\Delta H_5 = -127.48 \pm 1.84^a$
$BaO_{(s,25^{\circ}C)} \rightarrow BaO_{(soln,800^{\circ}C)}$	$\Delta H_6 = -181.22 \pm 2.32^a$
$PbO_{(s,25^{\circ}C)} \rightarrow PbO_{(soln,25^{\circ}C)}$	$\Delta H_7 = -14.34 \pm 0.38^a$
$MO_{(s,25^{\circ}C)} + Li_2O_{(s,25^{\circ}C)} + 6TiO_{2(s,25^{\circ}C)} \rightarrow MLi_2Ti_6O_{14(s,25^{\circ}C)}$	$\Delta H_8 = \Delta H^{\circ}_{f,ox} (MLi_2Ti_6O_{14})$
$\Delta H_8 = \Delta H^{\circ}_{f} (\mathrm{Na}_2 \mathrm{Li}_2 \mathrm{Ti}_6 \mathrm{O}_{14}) = -\Delta H_1 + \Delta H_2 + 6\Delta H_3 + \Delta H_4$	
$\Delta H_8 = \Delta H^{\circ}_{f} (SrLi_2Ti_6O_{14}) = -\Delta H_1 + \Delta H_2 + 6\Delta H_3 + \Delta H_5$	
$\Delta H_8 = \Delta H^{\circ}_{f}(\text{BaLi}_2\text{Ti}_6\text{O}_{14}) = -\Delta H_1 + \Delta H_2 + 6\Delta H_3 + \Delta H_6$	
$\Delta H_8 = \Delta H^{\circ}_{f}(\text{PbLi}_2\text{Ti}_6\text{O}_{14}) = -\Delta H_1 + \Delta H_2 + 6\Delta H_3 + \Delta H_7$	

Table 3. Thermodynamic Cycles Used to Calculate Formation Enthalpies ($\Delta H^{\circ}_{f.ox}$) of MLi₂Ti₆O₁₄ at 25 °C

^aThis work.

many other compounds, including perovskites, olivines, tungstates, carbonates, nitrides and oxynitrides, and glasses.^{29,30,32-36}

The enthalpy of formation from oxides is additionally influenced by the space constraints. In Na₂Li₂Ti₆O₁₄, monovalent sodium occupies the 11-fold site completely as opposed to the divalent counterparts which occupy only half of it. Thus, for the sodium compound, there is no vacant site available for further lithium insertion in the structure. However, in $MLi_2Ti_6O_{14}$ (M = Sr, Ba, and Pb), despite having similar, additional vacant sites, the divalent members uptake lithium differently. This could be due to size of the metal ions, acid-base interactions as discussed earlier, and the compressive strain induced by the ion at the 11-fold coordinated site. Low bond valence sum (BVS) for Sr (1.75) and Pb (1.79) indicates no residual strain at these sites whereas in Ba (2.11)due to its high BVS has a shortened Ba-O length which results in a compressive strain, strangulating the lithium pathway to interstitial sites.

The difference in the Li uptake by MLi₂Ti₆O₁₄ (M = Sr and Pb), regardless of having no compressive strain and similar ionic radii of Sr and Pb, is puzzling. The trend of decreasing stability in the $\Delta H^{\circ}_{f,ox}$ is Na₂Li₂Ti₆O₁₄ > BaLi₂Ti₆O₁₄ > SrLi₂Ti₆O₁₄ > PbLi₂Ti₆O₁₄. The low stability of PbLi₂Ti₆O₁₄ is attributed to the amphoteric nature of PbO. Consequently, the acid—base interaction between PbO, Li₂O, and TiO₂ is weak compared to alkali and alkaline earth metal oxides; thus, there is decline in the stability of PbLi₂Ti₆O₁₄. Similar differences are seen in comparing lead and strontium silicate glasses.³⁷ This is in corroboration to the Smith's scale of acidity.

Depending on the stability of these lithium titanates, the amount of Li uptake varies during charge cycles. The least stable PbLi₂Ti₆O₁₄ gains stability by intercalating 4.4 Li, whereas the slightly more stable SrLi₂Ti₆O₁₄ intercalates only 2.9 Li. The most stable SrLi₂Ti₆O₁₄ and Na₂Li₂Ti₆O₁₄ intercalate 2.7 and 1.3 Li. Future calorimetric measurements on samples with intermediate charge cycles having a variable amount of Li in the 11-coordination site for the MLi₂Ti₆O₁₄ (M = Ba, Sr, and Pb) structures would provide more insight into the stability.

CONCLUSIONS

It can be concluded that there is strong correlation between the thermodynamic stability and the lithium uptake in this class of materials. Strong exothermic acid–base reactions between MO, Li_2O , and TiO_2 result in large exothermic enthalpies of formation, indicating that MLi₂Ti₆O₁₄ are thermodynamically stable. Depending on the nature of acidbase interactions, the stability varies, alkali metals exhibit greater basic character followed by alkaline earth metal oxides, and among the alkaline earth metal oxides, those of larger cations are more basic compared to lighter cations, and the order of basicity for MO is $Na_2O > BaO > SrO$. PbO is an amphoteric oxide for which the acid-base interaction is weak compared to alkali or alkaline earth metal oxides. Thus, the thermodynamic stability diminishes in the order Na₂Li₂Ti₆O₁₄ > BaLi₂Ti₆O₁₄ > SrLi₂Ti₆O₁₄ > PbLi₂Ti₆O₁₄ which is directly proportional to the acid-base interactions. The trends in stability also depend on the charge and size of the cation and the compressive strain which are typical for interoxidic compounds. The formation enthalpy in these materials is inversely proportional to the lithium uptake during charge cycles. Least stable PbLi₂Ti₆O₁₄ uptakes most Li (4.4 Li), and the most stable $Na_2Li_2Ti_6O_{14}$ uptakes least Li (1.3 Li).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.2c05701.

Rietveld refinement of $MLi_2Ti_6O_{14}$ (M = Sr, Ba, and 2Na) titanates and thermodynamic cycles used to calculate the drop solution enthalpy (ΔH_{ds}) of SrO and BaO in sodium molybdate at 800 °C (PDF)

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Notes

The authors declare no competing financial interest.

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