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About the Compatibility between High Voltage Spinel Cathode Materials and Solid Oxide Electrolytes as a Function of Temperature


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Supporting Information

ABSTRACT: The reactivity of mixtures of high voltage spinel cathode materials $\text{Li}_x\text{NiMn}_2\text{O}_4$, $\text{Li}_x\text{FeMn}_2\text{O}_4$, and $\text{LiCoMn}_2\text{O}_4$ cosintered with $\text{Li}_{1.5}\text{Al}_{0.5}\text{Ti}_{1.5}\text{(PO}_4\text{)}_3$ and $\text{La}_{6.6}\text{La}_1\text{Zr}_{1.6}\text{Ta}_{0.4}\text{O}_{12}$ electrolytes is studied by thermal analysis using X-ray-diffraction and differential thermoanalysis and thermogravimetry coupled with mass spectrometry. The results are compared with predicted decomposition reactions from first-principles calculations. Decomposition of the mixtures begins at 600 °C, significantly lower than the decomposition temperature of any component, especially the electrolytes. For the cathode + $\text{Li}_{6.6}\text{La}_1\text{Zr}_{1.6}\text{Ta}_{0.4}\text{O}_{12}$ mixtures, lithium and oxygen from the electrolyte react with the cathodes to form highly stable $\text{Li}_2\text{Mn}_2\text{O}_4$ and then decompose to form stable and often insulating phases such as $\text{La}_2\text{Zr}_2\text{O}_7$, $\text{La}_2\text{O}_3$, $\text{La}_3\text{TaO}_7$, $\text{TiO}_2$, and $\text{LaMnO}_3$ which are likely to increase the interfacial impedance of a cathode composite. The decomposition reactions are identified with high fidelity by first-principles calculations. For the cathode + $\text{Li}_{1.5}\text{Al}_{0.5}\text{Ti}_{1.5}\text{(PO}_4\text{)}_3$ mixtures, the Mn tends to oxidize to $\text{MnO}_2$ or $\text{Mn}_2\text{O}_3$, supplying lithium to the electrolyte for the formation of $\text{Li}_3\text{PO}_4$ and metal phosphates such as $\text{AlPO}_4$ and $\text{LiMPO}_4$ ($M = \text{Mn, Ni}$). The results indicate that high temperature cosintering to form dense cathode composites between spinel cathodes and oxide electrolytes will produce high impedance interfacial products, complicating solid state battery manufacturing.

KEYWORDS: solid ion conductor, garnet, interfacial reactivity, atomistic modeling, solid electrolyte

INTRODUCTION

The development of lithium ion batteries has made consistent progress during the previous decades and is reaching performance and cost targets that make them attractive for emerging markets such as electric vehicles, grid storage, and wearable electronics. However, despite the advances, the ceiling of conventional lithium ion batteries is limited by the capacity of the graphitic anode and the achievable voltage and capacity limits at the positive electrode. 1,2 In order to achieve higher energy densities, new architectures must allow for high energy densities anodes such as lithium metal or silicon and high voltage cathodes such as manganese spinel materials, all while improving upon the safety of lithium ion batteries.

A promising direction is to move toward the replacement of the liquid electrolyte and conventional separator with a lithium ion conducting solid forming a solid state battery. 3,4 Critical to the success of solid state batteries is the ability to form stable interfaces, impervious to reduction against lithium metal and oxidation at the cathode. When this is achieved, such as in thin film batteries with a lithium phosphorus oxynitride (LiPON) electrolyte, it is possible to demonstrate thousands of cycles using lithium metal and high voltage $\text{LiNi}_{0.5}\text{Mn}_{1.5}\text{O}_4$ cathode. 5 However, in practice, achieving this level of compatibility is difficult. Most materials have been shown to be unstable against lithium metal reduction, including the high conductivity sulfides and oxides. 6–9 In fact, even LiPON was recently shown to decompose into a stable passivation layer. 10 Similarly, prevention of the solid electrolyte toward oxidation at the positive electrode is difficult. For sulfide electrolytes, researchers tend to use an oxide coating deposited on the cathode to prevent reaction. 11–13 Although oxide electrolytes can be stable at high voltages, 14 their high conductivity is...
usually achieved when the number of grain boundaries is minimized, as they can dominate the resistivity of the electrolyte.\textsuperscript{15} Densifying oxide electrolytes often requires high temperature sintering and, in order to achieve intimate contact with the cathode, cosintering of the electrolyte and cathode into a dense pellet is necessary. This cosintering can lead to the formation of a resistive layer through interdiffusion or decomposition.\textsuperscript{16} Identifying the decomposition reactions that occur is important to designing a solid state battery. Further, the ability to predict \textit{a priori}, which interfacial products form, simplifies the design process. Toward this end, in this work, we show how a combination of ab initio calculations, experimental thermal analysis techniques, and X-ray-diffraction together elucidate the decomposition reactions between high voltage spinel $Li_xNiMnO_8$ (LNM0), $Li_xFeMnO_8$ (LFMO), and $LiCoMnO_4$ (LCMO) cathodes in contact with $Li_1.5Al_0.5Ti_1.5(PO_4)_3$ (LATP) and $Li_{6.6}La_3Zr_1.6Ta_0.4O_{12}$ (LLZ:Ta) oxide solid electrolytes during high temperature cosintering.

\section*{METHODS}

\textbf{Experimental Section.} Spinel cathode materials, LNM0, LFMO, or LCMO, were synthesized by the Pechini method.\textsuperscript{17} Prior to the Pechini synthesis, aqueous solutions of Li, Ni, Fe, Co, and Mn nitrates were prepared and their cation contents were quantified by inductively coupled plasma atomic emission spectroscopy (Thermo Elemental, IRIS Intrepid). Nitrate solutions were weighed and mixed together corresponding to the desired stoichiometry of the spinel samples. Citric acid was added into an individual solution in a molar ratio of $n_{\text{citric acid}}/n_{\text{cation}} = 1.8$. The suspensions were stirred until clear solutions were obtained. 120 mL of ethylene glycol was added to each solution (corresponding to a final sample weight of 30 g). The solutions were stirred continuously and kept at 90 °C for water evaporation. The residues were heat treated at 300 °C for decomposition for several hours on heating plates. The first calcination step was at 650 °C for 5 h in a furnace. The second calcination step was completed after another 5 h at 650 °C in a muffle furnace. Afterward, the sample was ball milled in ethanol using yttria-stabilized zirconia balls and heat treated again at 650 °C for 5 h. The LATP and LLZ:Ta powders were synthesized by sol–gel and solid state reactions, respectively, described elsewhere.\textsuperscript{18}

The spinels LNM0, LFMO, or LCMO were mixed with either LATP or LLZ:Ta in a volume ratio of 50:50 using a mortar and pestle to produce “spinel + electrolyte” powders. The mixed spinel + electrolyte powders were pressed into pellets with a pressure of 370 MPa using pressing dies with 13 mm diameter. Sintering of pellets was carried out on MgO plates in corundum crucibles at 600, 700, and 800 °C for the spinel + LATP pellets and at 400, 600, and 800 °C for the spinel + LLZ:Ta pellets. Dwell time was 1 h each and heating rate was 5 K/min.

X-ray-diffraction (XRD) measurements were performed on the heat treated spinel + electrolyte pellets using a D4 Endeavor (Bruker axs) with Cu Kα radiation from 10° to 80° with a 2θ step width of 0.02° and a 0.75 s accumulative time. Qualitative phase analysis was carried out on the basis of the powder diffraction file (PDF) database and the inorganic crystal structure database (ICSD, reference numbers listed in Table S1) using the software package EVA (Bruker axs). Due to the multiphase compositions of the samples and due to too many common peak positions of a series of low symmetric phases, reliable phase quantification was not attempted using Rietveld refinement.

Differential thermoanalysis and thermogravimetry coupled with mass spectroscopy (DTA/TG/MS) measurements were carried out using a NETZSCH STA 449F1 in $N_2/O_2$ atmosphere (technical air). Spinel powders LNM0, LFMO, and LCMO were analyzed by DTA/TG from room temperature to 1400 °C and back to room temperature at a heating and cooling rate of 10 K/min. DTA/TG was carried out from room temperature to 1200 °C and back to room temperature at a heating and cooling rate of 5 K/min on spinel + LATP mixtures. DTA/TG/MS was carried out on spinel + LLZ:Ta mixtures from room temperature to 1100 °C and back to room temperature at a heating and cooling rate of 5 K/min.

\textbf{Computation.} Multicomponent phase diagrams constructed with data from both density functional theory (DFT) calculations and experimental data has been shown to be useful for studying the mixing thermodynamics of solid–solid interfaces.\textsuperscript{13} In this work, we use this approach to predict the interfacial products that form between the solid electrolyte and cathode materials upon cosintering at elevated temperatures.

To evaluate the stability of compounds, we construct grand potential phase diagrams using the pymatgen software package.\textsuperscript{20} To obtain bulk energies, we employ DFT within the Projector Augmented Wave (PAW) formalism\textsuperscript{21} using the generalized gradient approximation\textsuperscript{22} to the exchange-correlation energy as implemented in the Vienna ab initio simulation package (VASP)\textsuperscript{23} to calculate the formation energy of each compound from the nearest phases present in the NIST-JANAF\textsuperscript{24} or Kubaschewski\textsuperscript{25} thermochemical tables or from the elements. A cutoff energy of 520 eV and a k-point grid of at least 500/n\textsubscript{atom} was used for all calculations. The nearest phases are uniquely defined as those that define the Gibbs triangle (the low energy facet) containing the desired composition in the phase diagram. This phase diagram is generated using the DFT computed energies to determine the convex hull. The stability of the electrolytes is calculated with reference to the phase diagram hull energy at the location of the electrolyte compositions.

On the basis of the methods of Wang et al.\textsuperscript{26} and Ong and coworkers,\textsuperscript{27,28} the effect of temperature upon phase diagrams can be approximated by generating the oxygen grand potential phase diagrams and then mapping the oxygen chemical potential to temperature. The normalized mixing energy and the decomposition products formed at the interfaces are determined by calculating the energy at the different facets along the tie-line between cathode and electrolyte.\textsuperscript{14,28} At a given oxygen chemical potential, the reaction energy is found by

$$\Delta E[c_i, c_j] = E_{\text{prod}}[x c_i + (1 - x) c_j] - x E[c_i] - (1 - x) E[c_j]$$

where $c_i$ and $c_j$ are the electrolyte and cathode compositions, respectively, and $x$ is a mixing parameter that can vary between 0 and 1. The energies considered are the grand potential free energies after applying the external oxygen chemical potential corresponding to eq 1. The first term on the right side ($E_{\text{prod}}$) represents the energy of the “products” along the tie-line. The other energy terms are the grand potential energies of the end members (i.e., electrolyte and cathode). The minimum along this tie-line is considered to be the most likely reaction at a given temperature. An example is seen in Figure 1 for the reactions between spinel + LATP and spinel + LLZ:Ta at 800 °C.
RESULTS

Spinel Decomposition. The thermal decomposition characteristics in air of all three investigated high voltage cathodes are comparable, as shown in Figure 2. After initial reactions for the desorption of surface-adsorbed H₂O below 300 °C, the TG signals remain constant and no DTA signals are observed. From T₂ = 650 °C and higher, mass loss occurs parallel to endothermic reactions at T₃ and T₄ in several steps. For both the LCMO and LNMO, a reaction takes place at T₃ = 750 °C but not for LFMO. The peak temperatures of the endothermic reactions at T₄ are slightly different for the different samples with T₄_LNMO < T₄_LCMO < T₄_LFMO. During cooling, a mass uptake of around 70–80% of the lost mass between T₃ and the maximum temperature (T_max) is measured for all three samples. Additionally, for LFMO and LCMO, exothermic reactions are measured at temperature T₅. These results are consistent with earlier reported properties for the thermal decomposition of lithium manganese spinels.23–32 It is known that LiMn₁₂O₄ spinels show three decomposition reactions upon heating. The first decomposition starts at around 750 °C by releasing oxygen from the lattice and formation of an oxygen defect spinel and Li₂MnO₃. Above 915 °C, as a second step, Li₂MnO₃ and spinel react to form LiMnO₂ while manganese is reduced and oxygen is released from the lattice.

Results from the ab initio grand potential phase diagram are in agreement with these findings. Oxygen loss and decomposition begins around 510 °C with the reactions:

\[
\text{Li}_2\text{Mn}_2\text{NiO}_8 \rightarrow 0.847\text{Li}_2\text{MnO}_3 + 0.667\text{Mn}_2\text{O}_4 + 0.50\text{Fe}_2\text{O}_3; \mu_0 = -1.39\text{eV}; T \approx 510°C
\]

\[
\mu_0 = -1.50\text{eV}
\]

\[
\text{Li}_2\text{Mn}_2\text{NiO}_8 \rightarrow 0.833\text{Li}_2\text{MnO}_3 + 1.0\text{NiO} + 0.333\text{Mn}_2\text{O}_4 + 2.0\text{LiMnO}_2; \mu_0 = -1.53\text{eV}; T \approx 560°C
\]

Decomposition of the electrolytes occurs at much higher temperatures, Figure S1. Oxygen loss from LATP is predicted to begin around 1270 °C (Table S2) with the reduction of Ti⁴⁺ to form Li₂Ti₂(PO₄)₃, Li₅P₂O₇, and AlPO₄. This is higher than the results that found LATP to be stable to 900 °C.33 Decomposition of LLZ:Ta is calculated to occur at about 1600 °C with the reduction of Ta and the formation of La₂O₃, Li₅ZrO₃, and Li₅TaO₅.

Spinel + LLZ:TA. The thermal analysis by DTA/TG/MS of the spinel + LLZ:Ta powders reveals a comparable behavior upon heating for all three mixtures (Figure 3). Several sequential mass loss steps are already observed below 500 °C, which can be assigned to the loss of CO₂ and H₂O by MS (not shown). The loss of CO₂ and H₂O at temperatures below 600 °C is characteristic for air sensitive LLZ:Ta.34 No significant endo- or exothermic reactions can be observed within the whole measured temperature range. A small endothermic signal is detected for LFMO + LLZ:Ta at 490 °C. Nevertheless, mass loss continues above 600 °C, but no elements were detected by MS. Between T₁ and T₅_max around 6.5 wt % mass is released from the LFMO + LLZ:Ta sample, 6.0 wt % from the LNMO + LLZ:Ta sample, and 4.5 wt % from the LMCO + LLZ:Ta sample, whereas around 13–17% of this mass is taken up again during cooling.

XRD performed on spinel + LLZ:Ta pellets that were heat treated at 400, 600, and 800 °C showed major phase changes starting at 600 °C. Figures 4–6. The evolution of phase content from 400 to 800 °C is comparable for all three mixtures. For the sample heat treated at 400 °C, the observed diffraction pattern is the same as the diffraction pattern of the as-mixed powders. Only LLZ:Ta and spinel (here described by a LiMn₁₂O₄ phase with adjusted lattice parameter a = 8.06
according to ref 35) can be detected. The XRD pattern for the spinel + LLZ:Ta pellet which was heat treated at 600 °C shows a contribution of spinel and LLZ:Ta but a significant decrease in diffraction peak intensities of the LLZ:Ta phase in comparison with the diffraction pattern of the as mixed sample. Additionally, the pattern includes a secondary garnet phase and monoclinic Li₂MnO₃. The decomposition of lithium manganese spinels to Li₂MnO₃ usually occurs at temperatures around 750 °C.29,31,32 The detection of larger quantities of Li₂MnO₃, as was shown here, therefore indicates an accelerated formation of Li₂MnO₃ at lower temperatures due to Li supply and oxygen uptake from LLZ:Ta. Accordingly, a low amount of LLZ:Ta was detected as well as a secondary garnet phase that very likely exhibits a Li-depleted stoichiometry.

For the samples heat treated at 800 °C, the peak intensities of the spinel phases are decreased and no more LLZ:Ta can be detected. Instead, a La₂Zr₂O₇ phase can be identified clearly. Additionally, for LCMO, Li₂MnO₃ and LiCoO₂ are found. For the LFMO + LLZ:Ta sample, LaMnO₃ and LiFe₂O₄ are also found, whereas LaMnO₃ and NiO is found in the LNMO + LLZ:Ta sample.

**Figure 7.** (a) DTA and (b) TG of spinel + LATP powders. Experiments were carried out from room temperature to 1200 °C at a heating and cooling rate of 5 K/min. DTA signal offset: 1 unit (for LFMO) and 2 units (for LCMO). TG signal offset: 2 units (for LFMO) and 4 units (for LCMO).
loss due to the release of surface-adsorbed H₂O and CO₂ occurs below 270 °C = T₁. A continuous mass loss starting at T₁ is very small for all three mixtures and occurs at around 860 °C, whereas T₁ lies at 980 °C for LCMO + LATP and slightly higher, at 990 °C, for the other two mixtures. A mass loss of around 5−6 wt % can be measured for all three samples from T₁ to Tₘₐₓ. Around 15% of the lost mass is taken up again during cooling. Additionally, an exothermic reaction was measured during cooling for LCMO + LATP and LFMO + LATP at around 935 and 930 °C, respectively, and for LNMO + LATP at around 880 °C. The products of the spinel + LATP reaction upon heating showed a significantly lower amount of mass uptake during cooling than the single spinel materials. A reversible mass uptake is very characteristic for lithium manganese spinels, thus proposing that the lithium manganese spinel phase almost completely is transformed into new phases by the reaction with LATP at maximal temperature of 1200 °C.

XRD performed on LCMO + LATP pellets which were heat treated at 600, 700, and 800 °C are shown in Figure 8. The XRD pattern for the LATP + LCMO pellet heat treated at 600 °C is described completely by a LiMn₂O₄ spinel phase with a cubic lattice parameter of a = 8.16 Å, representative for LNMO, and a Li₅Ti₃(PO₄)₄ phase. The intensities of the Li₅Ti₃(PO₄)₄ and LiMn₂O₄ phases decrease significantly for the sample heat treated at 700 °C in comparison with those at 600 °C. Additionally, a tetragonal phosphate phase and rutile can be identified. Therefore, the LNMO + LATP reaction at 700 °C majorly affects the chemistry of the LNMO phase, since LNMO components react with LATP to form new phosphate phases. For the sample heat treated at 800 °C, no more LATP can be detected. The peak positions of the LiMn₂O₄ phase are shifted significantly to lower angles and need to be described by another spinel phase with a = 8.27 Å. A series of tetragonal phosphates is also necessary to describe the observed diffraction pattern.

For the LFMO + LATP pellet heat treated at 600 °C two phases are detected by XRD (Figure 10). The first one is LiMn₂O₄ with a = 8.26 Å, representative for LFMO, and the second one is LATP. For the sample heat treated at 700 °C, LFMO can still be detected but intensities are decreased, whereas LATP cannot be identified anymore. The cathode and electrolyte have almost completely reacted to form a MnFeO₃-like phase and orthorhombic phosphates, which occurs at a lower temperature than for the other two cathode materials. Neither LFMO nor LATP can be identified for the sample heat treated at 800 °C. Instead, at both temperatures, MnFeO₃ emerges as the main phase and a series of secondary tetragonal phosphates are necessary to describe the pattern satisfactorily. The MnFeO₃-like phase is still the main phase indicating its dominant thermodynamic stability in the LATP + LFMO pseudobinary system at 700 °C as well as at 800 °C.

The composite mixtures of spinel + LATP react strongly with each other from at least 700 °C, and similar to the spinel + LLZO:Ta composites, the decomposition products are stabilized at room temperature. The large amount of AlPO₄

![Figure 8. XRD patterns of LCMO + LATP mixtures that were heat treated at 800, 700, and 600 °C and as mixed powder. Table S3 lists the consulted reference patterns.](image_url)

![Figure 9. XRD patterns of LNMO + LATP mixtures that were heat treated at 800, 700, and 600 °C and as mixed powder. Table S3 lists the consulted reference patterns.](image_url)
Predicted Decomposition Reactions. Mixing of the oxide electrolytes and spinel cathodes results in decompositions occurring at a temperature lower than what is observed for each separate phase. This can be rationalized by following the highly mobile lithium and oxygen species. The result of mixing the spinel cathodes with LLZ:Ta is that the decomposition of the cathodes acts as an oxidizer for the decomposition of LLZ:Ta. The predicted reactions are listed in Table 1 and appear to identify the correct reactions although some of the minority phases predicted could not be detected with XRD. The decomposition of LLZ:Ta is predicted to occur around 1600 °C when heated by itself; however, in the presence of the cathodes, decomposition reactions are predicted to occur at the onset of cathode decomposition. As was pointed out in recent theoretical work, the barrier for gaseous oxygen loss from solid ceramics is rather high, and thus, decomposition can be kinetically limited. When the two ceramics are in contact at elevated temperatures, however, oxygen and lithium exchange readily occur leading to decomposition. In these spinel based systems, the decomposition reactions are dominated by the formation of the highly stable Li2MnO3. From an analysis of the reaction products, it appears that oxygen lost from the cathode is absorbed by the LLZ:Ta electrolyte to form a lithium oxide compound which then drives the formation of Li2MnO3. A similar reaction likely forms LiCoO2 and LiFeO2 in LCMO and LFMO, respectively, while in the LNMO a reduced Ni is likely formed such as NiO.

The mixing reactions between spinel + LATP are considerably more complicated than with LLZ:Ta. There is significant mixing predicted to occur for temperatures above ~150 °C. The spinel + LATP decomposition products include Li3PO4 and TiO2 formed from the decomposed LATP, as well as mixed products such as LiMnPO4 in the LNMO and LFMO decomposition products. Unlike for the spinel + LLZ:Ta systems, the predicted reactions with the LATP electrolyte at experimental temperatures do not align with the experimental results. In all cases, the experimental results align best with predicted reactions at lower temperatures, Table 2. The two most likely explanations are that either the products are diffusion limited so that the formation of mixed metal compounds does not readily occur or our predictions of the most stable compounds are slightly off. For example, the

![Figure 10. XRD patterns of LFMO + LATP mixtures that were heat treated at 800, 700, and 600 °C and as mixed powder. Table S3 lists the consulted reference patterns.](image)

Table 1. LLZ/Cathode Reactions

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Reaction Equation</th>
</tr>
</thead>
<tbody>
<tr>
<td>LCMO: 600°C</td>
<td>11.6Li6.6La3Zr1.6Ta0.4O12 + 39.9LiMnCoO4 → 0.25O2 + 0.2La2MnCoO6 + 0.2La2O3 + 9.3La2Zr2O7 + 4.7La3TaO7 + 38.9Li2MnO4</td>
</tr>
<tr>
<td></td>
<td>+ 38.9Li2CoO2</td>
</tr>
<tr>
<td>LCMO: 800°C</td>
<td>11.6Li6.6La3Zr1.6Ta0.4O12 + 39.9LiMnCoO4 → 0.25O2 + 0.2La2MnCoO6 + 0.2La2O3 + 9.3La2Zr2O7 + 4.7La3TaO7 + 38.9Li2MnO4</td>
</tr>
<tr>
<td></td>
<td>+ 38.9Li2CoO2</td>
</tr>
<tr>
<td>LNMO: 600°C</td>
<td>36.9Li6.6La3Zr1.6Ta0.4O12 + 61.3Li2Mn3NiO8 → 0.25O2 + 3.2La2O3 + 29.5La2Zr2O7 + 14.7La3TaO7 + 2LaMnO3 + 182.9Li2MnO3</td>
</tr>
<tr>
<td></td>
<td>+ 61.3NiO</td>
</tr>
<tr>
<td>LNMO: 800°C</td>
<td>5.0Li6.6La3Zr1.6Ta0.4O12 + 8.8Li2Mn3NiO8 → 4.66O2 + 4.0La2Zr2O7 + 2.0La3TaO7 + 25.4Li2MnO3 + 2LaMnO3 + 8.8Ni</td>
</tr>
<tr>
<td>LFMO: 600°C</td>
<td>5.3Li6.6La3Zr1.6Ta0.4O12 + 7.25Li3Mn3FeO6 + 1.8O2 → 4.3La2Zr2O7 + 2.1La3TaO7 + LaFeO3 + 21.8Li2MnO3 + 6.25LiFeO2</td>
</tr>
<tr>
<td>LFMO: 800°C</td>
<td>5.0Li6.6La3Zr1.6Ta0.4O12 + 7.0Li3Mn3FeO6 + 1.5O2 → 4.0La2Zr2O7 + 2.0La3TaO7 + LaMnO3 + 20.0Li2MnO3 + 7.0LiFeO2</td>
</tr>
</tbody>
</table>

“Red phases were detected by XRD; blue phases were possibly detected, but their peaks overlap with other phases.”
transition between the compounds with different oxidation states, such as CoO/Co3O4 and MnO2/Mn3O4, and the stable phosphates, AlPO4, LiMnPO4, and LiCoPO4, have a large effect on which reactions are predicted to occur. Getting this redox equilibrium simultaneously correct in varying chemical environments is a known challenge in ab initio calculations. 27 Although beyond the scope of this work, repeating the tests with in situ XRD would further elucidate the reaction mechanisms. The reactions predicted at the experimental temperatures are shown in Table S3. An example of the oxygen transfer in the LCMO + LATP follows the reaction scheme:

\[
\begin{align*}
2Li_{1.5}Al_{0.5}Ti_{1.5}(PO_4)_3 + 6.05LiMnCoO_4 & \rightarrow 0.5O_2 + 2.0Li_3PO_4 + AlPO_4 + 3.0LiTiPO_5 + 6.02MnO_2 + 2.0Co_3O_4 + 0.02Li_2MnCoO_4 \\
2Li_{1.5}Al_{0.5}Ti_{1.5}(PO_4)_3 + 6.05LiMnCoO_4 & \rightarrow 0.5O_2 + 2.02Li_2PO_4 + AlPO_4 + 2.98LiTiPO_5 + 0.02tIO_2 + 6.05MnO_2 + 2.01Co_3O_4 \\
9.73Li_{1.5}Al_{0.5}Ti_{1.5}(PO_4)_3 + 5.86Li_2Mn3NiO_8 & \rightarrow 8.8O_2 + Li_3PO_4 + 4.86AlPO_4 + 17.46LiMnPO_4 + 14.59tIO_2 + 5.86LiNiPO_4 + 0.07Mn_3O_4 \\
71.72Li_{1.5}Al_{0.5}Ti_{1.5}(PO_4)_3 + 43.23Li_2Mn3NiO_8 & \rightarrow + 64.9O_2 + 7.37Li_3PO_4 + 35.86AlPO_4 + 129.7MnMnPO_4 + 106.7tIO_2 + 42.22LiNiPO_4 + 7TiNiO_4 \\
2.0Li_{1.5}Al_{0.5}Ti_{1.5}(PO_4)_3 + 5.93Li_3Mn_2FeO_8 & \rightarrow 3.0O_2 + AlPO_4 + 0.02Li_3Fe_2(PO_4)_3 + 4.93Li_3PO_4 + 5.96Mn_3O_3 + 5.89MnFeO_3 + 3.0tIO_2 \\
29.9Li_{1.5}Al_{0.5}Ti_{1.5}(PO_4)_3 + 88.6Li_2Mn_3FeO_8 & \rightarrow 44.5O_2 + 14.9AlPO_4 + 73.7Li_3PO_4 + LiMnPO_4 + 88.1Mn_3O_3 + 88.6MnFeO_3 + 44.8tIO_2 \\
\end{align*}
\]

(2)

(3)

From eqs 2 and 3, we see the decomposition of LCMO followed by the oxidation of LiMnO4, which then supplies lithium to react with the LATP system, the decomposition products of the spinel have a strong mixing energy with the LATP suggesting that the components will mix rather than simply decompose.

In a battery, a cathode composite layer is necessary which has interpenetrating electrically and ionically conductive networks that percolate through the composite layer. It is critical that the interfaces between the ionically conductive electrolyte and the cathode active material have low resistance. Although LLZ:Ta, in the spinel + LATP system, the decomposition products of the spinel have a strong mixing energy with the LATP suggesting that the components will mix rather than simply decompose.

A cosintered active material than the other spinels for use with LLZ:Ta. 36,39

The large reactivity between the spinels and LATP at even modest temperatures suggest that cosintering will be difficult, especially when there are very stable products such as Li2MnO3 or Li3PO4. This is unfortunate since it is predicted that LATP is stable up to high voltages, enabling the use of high voltage cathodes. 14 Encouragingly, the decomposition products include Li3PO4, which has modest ionic conductivity and may allow the battery to function. Therefore, a low temperature processing route to form well-bonded composites will likely be necessary to fabricate solid state cathodes in future solid state batteries.

**CONCLUSIONS**

Single spinel materials as well as spinel materials in combination with the LATP or LLZ:Ta electrolyte showed thermal decomposition at temperatures above 550–600 °C. Our results show the reaction reversibility of the single spinel materials is significantly altered if mixed with either LATP or LLZ:Ta electrolyte. XRD studies revealed that spinel + LATP mixtures are compatible with each other at 600 °C for the given sintering conditions. At higher temperatures, the materials react to form new phases. Spinel + LLZ:Ta were shown not to be compatible with each other even at 600 °C. The reaction products from these decomposition reactions were predicted from first-principles phase diagrams. The results suggest that insulating phases form above 600 °C which will likely add to the interfacial impedance of composite cathode in solid state batteries.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.6b09059.

DTA/TG data for LLZ and LATP powders, oxygen evolution profiles, and reaction energies for the mixed cathode-electrolyte systems (PDF)
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