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Exploration of NaSICON Frameworks as Calcium-Ion Battery Electrodes

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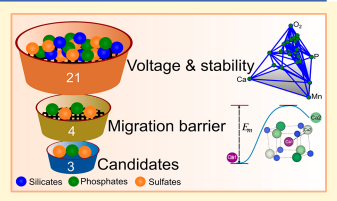
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ABSTRACT: The development of energy storage technologies that are alternative to state-of-the-art lithium-ion batteries but exhibit similar energy densities, lower cost, and better safety is an important step in ensuring a sustainable energy future. Electrochemical systems based on calcium (Ca)-intercalation or deintercalation form such an alternative energy storage technology but require the development of intercalation electrode materials that exhibit reversible Ca-exchange with reasonable energy density and power density performance. To address this issue, we use first-principles calculations to screen over the wide chemical space of sodium superionic conductor (NaSICON) frameworks, with a chemical formula of $Ca_xM_2(ZO_4)_3$ (where $M=Ti,\,V,\,Cr,\,Mn,\,Fe,\,Co,\,Or\,Ni\,Argeronal Ca,\,Po,\,Or\,Ni\,Argeronal Ca,\,Or\,Ni\,Argeronal Ca,\,Or\,Ni\,Arge$



calculate the average Ca^{2+} intercalation voltage, the thermodynamic stability (at 0 K) of charged and discharged Ca-NaSICON, and the migration barriers of (meta)stable Ca-NaSICON compositions. Importantly, our calculations indicate $Ca_xV_2(PO_4)_3$, $Ca_xMn_2(SO_4)_3$, and $Ca_xFe_2(SO_4)_3$ Ca-NaSICONs to be promising as Ca-cathodes. We find all silicate Ca-NaSICONs to be thermodynamically unstable and hence unsuitable as Ca-cathodes. We report the overall trends in the ground state Ca-vacancy configurations, besides voltages, stabilities, and migration barriers. Our work contributes to unearthing strategies for developing practical calcium-ion batteries, involving polyanionic intercalation frameworks.

1. INTRODUCTION

Li-ion batteries (LIBs) are well-known for high energy and power density, good cycle life, and reliable performance, making them state-of-the-art in the portable electronics and electric vehicle markets. However, looking into the future, there are various factors with respect to limited availability of lithium (also cobalt and nickel), ever-increasing energy demand, safety concerns, etc., which cast uncertainty over the ability of LIBs to power the world's requirements in the long run. 1-5 Therefore, the advancement and diversification of beyond-lithium (Li) electrochemical energy systems are imperative to fulfill both energy and power demands worldwide. Multivalent batteries, such as those based on magnesium, calcium (Ca), zinc, etc., are an emerging alternative to LIBs because of their high abundance in nature and their use of metal anodes leading to high (volumetric) energy density. 6-11 Among the multivalent systems, calcium is an attractive candidate because of its low standard reduction potential (-2.87 V versus standard hydrogen electrode, SHE) approaching that of Li⁺ (-3.04 V versus SHE), low cost, and abundance. 10,12

The development of Ca-ion batteries (CIBs) or Ca-based electrochemical systems is still in its infancy, especially due to

the challenge of finding "compact" cathode materials with high reversible capacity and reasonable rate performance, apart from the challenges associated with electrolyte development. Several cathode frameworks have been considered as Cacathodes, including 3-D "tunnel" structures, layered and spinel compounds, Chevrel phases, and Prussian blue analogues (PBA), with varying degrees of electrochemical performance. Theoretical screening approaches, such as using density functional theory (DFT^{17,18}) calculations by Arroyo-de Dompablo and collaborators, have investigated Ca intercalation in a wide range of chemistries, $^{19-21}$ such as CaFeSO, CaCoSO, CaNiN, Ca₃MnN₃, Ca(VO)₂(PO₄)₂, Ca₂Fe(Si₂O₇), CaM(P₂O₇) (M = V, Cr, Mn, Fe, and Co), CaV₂(P₂O₇)₂, and α -VOPO₄. The authors identified α -VOPO₄ as a promising cathode material due to its maximum theoretical specific capacity of 312 mAh g⁻¹, reasonable activation barrier for Ca

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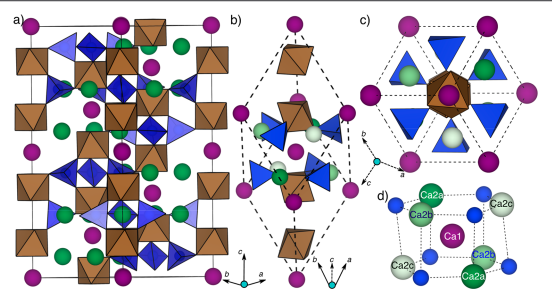


Figure 1. Crystal structures of the NaSICON conventional cell (panel a) and primitive cell (panel b). Brown and blue polyhedra indicate MO₆ and ZO₄ groups, respectively, where oxygen atoms occupy the polyhedral vertices. Purple and dark green spheres in panel (a) indicate Na1 and Na2 sites, respectively, which are equivalent to the Ca1 and Ca2 sites in Ca-NaSICONs. The different shades of green in panels (b), (c), and (d) signify the Ca2a, Ca2b, and Ca2c sites, which together form the Ca2 sites. Dashed black lines indicate the extent of the unit cell. (c) "Top" view of the NaSICON primitive cell, which clearly indicates the rhombohedral symmetry of the underlying structure. (d) A view of the Ca2 and Z sites that neighbor a Ca1 site, forming a hexagonal-prism-like block. Ca migration events in NaSICONs typically occur from a Ca1 site to a neighboring Ca2 site within the hexagonal prism.

diffusion (<0.65 eV), and a robust calculated average voltage of 2.8 V vs Ca. Although VOPO $_4$ ·2H $_2$ O has shown a reasonable electrochemical performance against Ca-containing electrolytes, ²² given the susceptibility of the framework to undergo proton intercalation, ²³ reversible Ca-intercalation in the VOPO $_4$ framework is yet to be rigorously demonstrated.

Several classes of oxide frameworks have been explored as Ca-cathode materials. Liu et al. 14 explored oxide spinels as Cacathodes using DFT calculations and found CaMn2O4 to be a potential cathode with an average intercalation voltage of ~3.1 V vs Ca and a theoretical gravimetric capacity of 250 mAh g⁻¹ with a theoretical energy density of ~800 Wh/kg. However, the calculated volume change with Ca²⁺ intercalation was found to be >20% which highlighted the instability of the lattice during (de)intercalation of Ca-ions, as later demonstrated by experiments.²⁴ In the case of (α) δ -V₂O₅, reversible (de)intercalation of Ca has been demonstrated at an average voltage of ~(3.13) 3.02 V versus Ca²⁺/Ca, with a theoretical capacity of 242 mAh g⁻¹ and activation barrier for Ca mobility of \sim (1700–1900 meV) 200 meV, respectively. ²⁵ However, the experimental investigation by Verrelli et al. ²⁶ elucidated the unsuitability of α - V_2O_5 as an intercalation host for CIB. Because the δ -V₂O₅ is not a stable polymorph, its synthesizability and electrochemical activity remain to be verified by experiment.

Other oxide chemistries that have been investigated as Cacathodes include $Ca_4Fe_9O_{17}$ as reported by Black et al., which exhibits a high calculated average voltage of 4.16 V vs Ca, a theoretical capacity of 230 mAh g⁻¹, as well as a reasonable activation barrier for Ca mobility of 0.72 eV.²⁷ Nonetheless, due to its high voltage, experimental validation of this compound is limited by the electrochemical stability windows of the currently available electrolytes. Recently, Lu et al. reported two promising Ca-cathode compositions, via a DFT-based screening procedure that included average voltages, charge neutrality, thermodynamic stability, and migration

barriers, namely, post-spinel- CaV_2O_4 and layered- $CaNb_2O_4$. While recent experiments indicate promise on reversible Ca intercalation in CaV_2O_4 , further optimization of the electrode framework is certainly required. ²⁹

In the chemical space of sulfide cathode materials, Palacin and co-workers have explored the electrochemical activity of Ca in TiS₂ for a variety of electrolytes. 30,31 The authors claimed reversible intercalation (with solvent cointercalation) of Ca in TiS2 in Ca(TFSI)2:PC electrolyte at 60 °C, 100 °C, and room temperature.³¹ Smeu et al. theoretically examined the Chevrel phases of $CaMo_6X_8$ (X = S, Se, Te) as viable Caelectrodes and found a high activation barrier for Ca mobility in this material (~780 meV) despite a reasonable intercalation voltage of ~1.4 V vs Ca²⁺/Ca.¹⁵ Kuperman et al. investigated the potassium iron hexacyanoferrate (a PBA framework) in a nonaqueous Ca electrolyte and reported reversible capacities of 150 mAh g⁻¹ at 23 mA g⁻¹ current density. 16 However, the experimental charge/discharge graphs exhibited capacitive-like behavior instead of the desired mechanism of Ca2+ intercalation. In the case of the Ca_{0.89}TaN₂ framework, Verrelli et al. reported irreversible electrochemical intercalation of Ca.³² Thus, the search for host materials enabling a reversible Ca-intercalation cathode with reasonable energy density and rate performance is still an active topic of research.

Given the similar size of Ca^{2+} (~ 1.00 Å in an octahedral coordination environment surrounded by O^{2-}) compared to Na^+ (~ 1.02 Å), 33,34 cathodes that exhibit reversible Naintercalation may be promising for Ca as well. Sodium superionic conductor (NaSICON) frameworks, which were first reported as Na solid electrolytes by Hong and Goodenough, 35,36 with the classical formula of $Na_{1+x}Zr_2Si_xP_{3-x}O_{12}$ ($0 \le x \le 3$), may be a potential class of tunable polyanionic Ca-cathodes. NaSICON materials comprise a chemical formula $A_xM_2(ZO_4)_3$, where A is the active ion (usually Na^+), M is a transition metal (or a combination of transition metals), and Z is usually Si, P, and/or S (and can also contain

elements, such as, Ge, As, etc.). ³⁶ With M being an open-shell transition metal, the NaSICON structure becomes redoxactive, and the framework can be a potential electrode material. NaSICON has been explored as a Na-electrode by multiple studies. ^{37–42} The NaSICON-NaV₂(PO₄)₃ framework has been recently shown experimentally to exhibit promising reversible Ca intercalation, with an energy density of ~220 Wh kg⁻¹, discharge capacity of ~70 mAh g⁻¹, and average voltage of 3.2 V vs Ca, signifying the promise of NaSICON frameworks as Ca-cathodes. ⁴³

Structurally, NaSICON contains octahedral MO₆ units (brown polyhedra in the conventional and primitive cells shown in panels (a) and (b) of Figure 1, respectively) connected via corner-sharing ZO₄ tetrahedra (blue polyhedra), forming a three-dimensional skeleton with two distinct intercalant sites, namely, 6-coordinated "Na1" (purple spheres in Figure 1a) and 8-coordinated "Na2" (dark green spheres) sites. Figure 1b shows the primitive cell of the NaSICON which consists of two M₂(ZO₄)₃ formula units, while the conventional cell of Figure 1a contains six formula units, with the rhombohedral symmetry of the structure highlighted in the "top" view of Figure 1c. In a Ca-NaSICON, both Na1 and Na2 sites can be occupied by Ca, resulting in "Ca1" and "Ca2" sites, respectively (Figure 1b,d). Note that there is one Ca1 site and three Ca2 sites (labeled as "Ca2a", "Ca2b", and "Ca2c" in Figure 1d) per M₂(ZO₄)₃ formula unit. Both Ca1 and Ca2 sites share faces with two MO₆ octahedra, while Ca2 sites also share edges with ZO₄ tetrahedra. Each Ca1 is surrounded by six Ca2 sites and six Z sites in its vicinity, forming a hexagonalprism-like block, as shown in Figure 1d. Also, the NaSICON framework can be considered to be made of "lantern" units comprising two MO₆ octahedra that are connected by three corner sharing ZO₄ tetrahedra (not shown in Figure 1).⁴

Here, we use DFT-based calculations to screen the chemical space of $Ca_xM_2(ZO_4)_3$, where Z = Si, P, or S and M is a 3dtransition metal among Ti, V, Cr, Mn, Fe, Co, and Ni, as possible Ca-cathodes. In principle, a $Ca_xM_2(ZO_4)_3$ framework can accommodate an exchange of a maximum of 2 Ca-ions, resulting in an oxidation state change of ± 2 per M atom (e.g., $M^{2+} \leftrightarrow M^{4+}$), making Ca-NaSICONs favorable in terms of energy density and motivating our choice of 3d transition metals. We calculate the average intercalation voltages, theoretical capacities, thermodynamic stabilities, and migration barriers upon Ca (de)intercalation in the Ca-NaSICON structures and benchmark our results with available experimental data. Specifically, we find $Ca_xV_2(PO_4)_3$, $Ca_xMn_2(SO_4)_3$, and $Ca_xFe_2(SO_4)_3$ to be promising cathode materials in terms of practicable intercalation voltages, thermodynamic (meta)stabilities, and reasonable migration barriers. The high thermodynamic instability eliminates silicate Ca-NaSICONs as Ca-cathode candidates. Ours is a screening study of the diverse chemical space of Ca-NaSICONs as potential calcium intercalation hosts, and we hope that it inspires further theoretical and experimental exploration of NaSICONs as Ca-cathodes.

2. METHODS

All spin-polarized total energy DFT calculations were performed using the Vienna *ab initio* simulation package (VASP), ^{45,46} by employing the projector-augmented-wave (PAW)^{47,48} formalism, where the specific PAW potentials used are identical to those of our previous work ^{49,50} and listed in the Supporting Information. Expanding the one-electron Kohn–Sham orbitals with a plane-wave kinetic energy

cutoff of 520 eV, we employed the Hubbard U corrected strongly constrained and appropriately normed (i.e., SCAN+U) functional to describe the electronic exchange and correlation. The U value applied for each transition metal was derived from previous work. We utilized Γ -point-centered, Monkhorst–Pack h-point meshes with a minimum of 32 subdivisions along each unit reciprocal lattice vector for all structures and a Gaussian smearing (of 0.05 eV) to integrate the Fermi surface. During the structure relaxation, the total energy of each structure was converged within 10^{-5} eV, and the atomic forces were converged to |0.03| eV/Å. Note, we have relaxed the cell volumes, shapes, and ionic positions of all structures without preserving any underlying symmetry in all our calculations.

Given the range of the transition metal oxidation states possible within the NaSICON framework (i.e., $M^{2+} \leftrightarrow M^{4+}$) and assigning oxidation states of +4, +5, and +6 for Si, P, and S, respectively, the Ca-NaSICON structure will remain electrostatically charge neutral for Ca contents in the range of $2 \le x \le 4$ in $Ca_xM_2(SiO_4)_3$, $0.5 \le x \le 2.5$ in $Ca_x M_2(PO_4)_3$, and $0 \le x \le 1$ in $Ca_x M_2(SO_4)_3$. Thus, $Ca_2 M_2(SiO_4)_3$, Ca_{0.5}M₂(PO₄)₃, and M₂(SO₄)₃ constitute "charged" Ca-NaSICON compositions, while $Ca_4M_2(SiO_4)_3$, $Ca_{2.5}M_2(PO_4)_3$, and Ca₁M₂(SO₄)₃ correspond to the "discharged" compositions. Among 3d transition metals, V can theoretically exhibit a +5 oxidation state. However, the extraction of the final Na in the V-phosphate NaSICON (i.e., Na removal from NaV2(PO4)3), which would allow V to exhibit a +5 oxidation state, has not reliably been achieved so far either chemically or electrochemically. 42 Hence, we preclude the possibility of V displaying a +5 oxidation state in Ca_xV₂(PO₄)₃. In summary, we assumed a possible exchange of 2 mol of Ca per NaSICON formula unit in silicates and phosphates, besides considering 1 mol of possible Ca exchange per formula unit in sulfates. Note, the theoretical capacities considered in this work should be considered as an upper limit to any experimental measurement.

To generate the initial Ca-NaSICON structures, we used the primitive cell of the Na₄Zr₂(SiO₄)₃ framework as the starting guess (2 formula units or 42 atoms), where each 3d transition metal substitutes Zr to generate the corresponding Ca-NaSICON framework.⁵⁵ The phosphate and sulfate-based Ca-NaSICONs are derived by substituting Si with P and S, respectively. After performing elemental substitutions, the lattice parameters of the theoretically generated Ca-NaSICON structures were scaled using the "RLSVolumePredictor" class of the pymatgen package.⁵⁶ All unique Ca-vacancy arrangements in the silicate (i.e., for Ca₂M₂(SiO₄)₃), phosphate (Ca_{0.5}M₂(PO₄)₃ and Ca_{2.5}M₂(PO₄)₃), and sulfate (Ca₁M₂(SO₄)₃) NaSICON primitive cells were enumerated using the "enumlib" library by Hart et al. 57-60 In total, we generated 70 $Ca_2M_2(SiO_4)_3$ (10 per transition metal), 14 $Ca_{0.5}M_2(PO_4)_3$, 42 $Ca_{2.5}M_2(PO_4)_3$, and 28 $Ca_1M_2(SO_4)_3$ configurations. The $Ca_4M_2(SiO_4)_3$ and $M_2(SO_4)_3$ compositions comprise one Ca-vacancy configuration per transition metal. Note that all the NaSICON structures considered in this work were generated theoretically, based on the Na₄Zr₂(SiO₄)₃ primitive cell.

A reversible intercalation battery based on Ca-NaSICON cathodes involves insertion/extraction of Ca^{2+} into/from the $Ca_xM_2(ZO_4)_3$ framework according to the redox reaction of eq 1.

$$Ca_yM_2(ZO_4)_3 + (x - y)Ca^{2+} + 2(x - y)e^- \leftrightarrow Ca_xM_2(ZO_4)_3$$

where y and x are the concentrations of Ca in the charged and discharged Ca-NaSICON structure, respectively. The average intercalation voltage due to (de)intercalation of (x-y) moles of Ca²⁺ can be calculated from the Gibbs energy change, which is approximated by the total energy change as calculated by DFT ($\Delta G \approx \Delta E$, eq 2), ignoring the p-V and entropic contributions. F in eq 2 is the Faraday constant, and $\mu_{\rm Ca}$ is the Ca chemical potential in pure Ca metal (i.e., in its ground state face-centered-cubic structure).

$$\begin{split} \langle V \rangle &= -\frac{\Delta G}{2(x-y)F} \\ &\approx -\frac{E(Ca_x M_2(ZO_4)_3) - [E(Ca_y M_2(ZO_4)_3) + (x-y)\mu_{Ca}]}{2(x-y)F} \end{split}$$
 (2)

To evaluate the thermodynamic stability, we compute the 0 K convex hull of quaternary Ca-M-Z-O systems by considering all possible elements, binaries, ternaries, and quaternaries whose structures are available in the inorganic crystal structure database (ICSD⁶²) and calculating their ground state energies with DFT. A list of the binary, ternary, and quaternary compounds calculated by us is provided in the Supporting Information, and the final geometries and energies of all computed structures (except binary transition metal oxides) are also available in the GitHub repository associated with this work.⁶³ We used the pymatgen⁵⁶ package to construct our convex hulls and quantify the extent of instability or stability by calculating the energy above/below the convex hull (E^{hull}) . In the case of unstable/metastable phases, (positive) E^{hull} indicates the largest decomposition energy that can be released by the unstable/metastable phase decomposing into its ground state(s). On the other hand, for a stable phase, (negative) E^{hull} is the lowest energy released by the formation of the stable phase from other stable phases within the chemical space.⁶⁵ Thus, compounds that are stable have $E^{hull} \leq 0$. Previous studies have used an approximate rule of thumb of $E^{hull} \sim 50$ meV/atom as a threshold of experimental synthesizability, 66 which we also consider here.

We calculated the Ca^{2+} migration barriers (E_m) to investigate Caion mobility within a subset of candidate Ca-NaSICON frameworks, using the DFT-based nudged elastic band (NEB⁶⁷) method. Notably, performing ab initio molecular dynamics simulations for Ca migration can be computationally prohibitive, especially when the migration barriers are high. 68,69 We calculated the E_m at high ("charged" compositions) and/or low ("discharged" compositions) Ca vacancy concentration limits. To set the Ca2+ migration path within the framework, we relaxed the initial and final end point configurations, assuming a vacancy-mediated migration mechanism, until the atomic forces were converged to |0.03| eV/Å. We considered all Ca migration events in all Ca-NaSICONs to occur between a Ca1 site and a Ca2 site within the hexagonal-prism displayed in Figure 1d, given previous studies that have established Na migration events to occur between Na1 and Na2 sites in Na-containing NaSICONs. 70,71 The minimum energy path (MEP) was initialized by linearly interpolating the lattice vectors and atomic positions to create seven intermediate images between the end points, with a spring constant of 5 eV per Å between images. The NEB calculation, done on Γ -centered Monkhorst-pack meshes with a density of at least 32 points per Å, was considered converged if the band forces became less than |0.05| eV/Å.

For the NEB calculations, we examined the effect of the choice of the exchange-correlation functional and cell size on E_m by using $Ca_{0.5}V_2(PO_4)_3$ and $Ca_{2.5}V_2(PO_4)_3$ as test cases, with the data presented in Figure S8 and Table S3 of the Supporting Information. Importantly, we found minimal difference in calculated E_m (~7–13% variation, or ~80-160 meV) between the Ca-NaSICON primitive cell (2 formula units of $M_2(ZO_4)_3$) and a supercell configuration (8 formula units),71 thus motivating the use of the primitive cell for further calculations. Note that the distance between migrating Ca²⁺ is >8 Å across its periodic image,⁷² which ensures minimal fictitious interactions between the migrating ions. Additionally, we found that the Perdew-Burke-Ernzerhof (PBE⁷³) parametrization of the generalized gradient approximation (GGA) consistently provided a lower estimate of E_m (by ~200 meV) compared to SCAN, consistent with our recent observations across multiple electrode and solid electrolyte materials.⁷⁴ Moreover, we found a negligible difference (~40 meV) in calculated E_m between SCAN and SCAN+U. Given that SCAN NEB calculations are computationally more expensive and are notoriously difficult to converge compared to GGA and that GGA provides reliable qualitative trends on calculated E_m ⁷⁴ the E_m data presented here, especially of the promising candidates, is calculated

using GGA. All calculated minimum energy pathways, including GGA and SCAN NEBs, are compiled in Figures S9, S10, and S11 of the Supporting Information.

3. RESULTS

3.1. Structural Features of $Ca_xM_2(SiO_4)_3$, $Ca_xM_2(PO_4)_3$, and $Ca_xM_2(SO_4)_3$. Table 1 lists the ground state Ca-vacancy

Table 1. Site Occupancy of the Ca1, Ca2a, Ca2b, and Ca2c Sites in the Calculated Ground State Configurations for Each Ca-NaSICON Composition Considered^a

	Ca site occupancy in ground state			
Composition	Ca1	Ca2a	Ca2b	Ca2c
Silicates				
$Ca_4M_2(SiO_4)_3$	1.0	1.0	1.0	1.0
$Ca_2M'_2(SiO_4)_3$, M' = Ti, V, Cr, Mn, Fe	1.0	1.0	0.0	0.0
$Ca_2Co_2(SiO_4)_3$	0.5	1.0	0.5	0.0
$Ca_2Ni_2(SiO_4)_3$	0.5	0.5	0.5	0.5
Phosphates				
$Ca_{2.5}M_2(PO_4)_3$	1.0	0.5	0.5	0.5
$Ca_{0.5}M_2(PO_4)_3$	0.5	0.0	0.0	0.0
Sulfates				
$Ca_1M_2(SO_4)_3$	1.0	0.0	0.0	0.0
$M_2(SO_4)_3$	0.0	0.0	0.0	0.0

^aIn a Ca-NaSICON primitive cell, there are two Ca sites of each type, indicating that the possible occupancy of each Ca site is 0.0, 0.5, or 1.0. M in the table with all transition metals considered (i.e., Ti, V, Cr, Mn, Fe, Co, and Ni). A schematic of all ground state configurations is provided in Figure S1.

configurations, in terms of occupancy of each Ca site type, for all charged and discharged Ca-NaSICON compositions considered here. Representations of these configurations and their simulated XRD pattern are shown in Figures S1 and S2—S4 of the Supporting Information, respectively.

For the discharged $(Ca_{2.5}M_2(PO_4)_3)$ and the charged $(Ca_{0.5}M_2(PO_4)_3)$ phosphates, the ground state Ca-vacancy configuration is identical for all transition metals within each compositional group. For example, in $Ca_{2.5}M_2(PO_4)_3$, the Ca1 site is fully occupied, and all Ca2 sites display a 0.5 occupancy. In the case of $Ca_{0.5}M_2(PO_4)_3$, the Ca1 site exhibits half occupancy for all transition metals, with all Ca2 sites being fully empty. Similarly, the discharged $Ca_1M_2(SO_4)_3$ shows an identical ground state Ca-vacancy configuration for all transition metals, with the Ca1 site being fully occupied and all Ca2 sites being fully empty. The presence of P^{S+} and P^{S+} in the P^{S+} detrahedra exert strong electrostatic repulsions on the P^{S+} forcing them to occupy fully the Ca1 sites (which only share corners with P^{S+} tetrahedra) before any of the Ca2 sites are occupied in both phosphates and sulfates.

Discharged silicate compounds, with composition $Ca_4M_2(SiO_4)_3$, involve Ca occupying all four Ca sites fully occupied (Figure S1d). Similar to the phosphates and sulfates, the ground state configuration of the charged compositions of $Ca_2M_2(SiO_4)_3$ for M=Ti, V, Cr, Mn, and Fe (Figure S1a) are identical, with Ca-ions fully occupying the Ca1 and the Ca2a sites. However, the ground state configuration of the charged $Ca_2Co_2(SiO_4)_3$ (Figure S1b) and $Ca_2Ni_2(SiO_4)_3$ (Figure S1c) are distinct from those of the other TM analogues. In the case of $Ca_2Co_2(SiO_4)_3$, the Ca2c site is fully occupied with the Ca1 and Ca2a sites being half occupied. In contrast, all Ca sites are

half-occupied in the case of $Ca_2Ni_2(SiO_4)_3$. Note that both Si and the transition metal have a nominal 4+ oxidation state at compositions $Ca_4M_2(SiO_4)_3$, indicating that the occupancy of Ca sites can be influenced by both electrostatic effects as well as the hybridization effects of the $M^{4+}-O^{2-}$ and $Si^{4+}-O^{2-}$ bonds. These effects explain the different occupancies observed in the charged Co- and Ni-silicates.

3.2. Intercalation Voltages and Theoretical Capacities of Ca-NaSICONs. Figure 2 displays a comparison of the

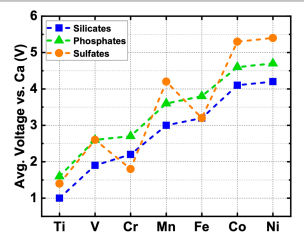


Figure 2. Computed average voltage as a function of transition metal, between the fully discharged and charged Ca-NaSICON compositions for silicate (blue squares), phosphate (green triangles), and sulfate (orange circles) chemistries.

computed average voltage (in Volts versus Ca metal), for silicate (blue symbols), phosphate (green), and sulfate (orange) Ca-NaSICONs across the entire range of chemically accessible compositions (i.e., $\text{Ca}_2\text{M}_2(\text{SiO}_4)_3 \leftrightarrow \text{Ca}_4\text{M}_2(\text{SiO}_4)_3$, $\text{Ca}_{0.5}\text{M}_2(\text{PO}_4)_3 \leftrightarrow \text{Ca}_{2.5}\text{M}_2(\text{PO}_4)_3$, and $\text{M}_2(\text{SO}_4)_3 \leftrightarrow \text{Ca}_1\text{M}_2(\text{SO}_4)_3$) for each 3d transition metal considered. We also compile all calculated voltages and the corresponding gravimetric capacities (in mAh g $^{-1}$) in Table 2. Expectedly, the theoretical capacities that are calculated with reference to the charged compositions decrease from Ti to Ni for silicates (from 237 to 226 mAh g $^{-1}$, see Table 2), phosphates (268 to 254 mAh g $^{-1}$), and sulfates (140 to 132 mAh g $^{-1}$), corresponding to an increase of atomic mass from Ti to Ni

Table 2. Average Intercalation Voltage (in Volts vs Ca) and the Theoretical Capacity (in mAh g^{-1}) of Silicates, Phosphates, and Sulfates-Based Ca-NaSICON Shown for the 3d Transition Metals^a

	Average Voltage (V)			Theoretical Capacity (mAh g ⁻¹)			
M	Silicate	Phosphate	Sulfate	Silicate	Phosphate	Sulfate	
Ti	1.0	1.6	1.4	237	268	140	
V	1.9	2.6	2.6	234	264	137	
Cr	2.2	2.7	1.8	233	262	137	
Mn	3.0	3.6	4.2	230	258	135	
Fe	3.2	3.8	3.2	229	257	134	
Co	4.1	4.6	5.3	226	254	132	
Ni	4.2	4.7	5.4	226	254	132	

^aThe theoretical capacities are computed with respect to the molar mass of $Ca_2M_2(SiO_4)_3$ for silicates, $Ca_{0.5}M_2(PO_4)_3$ for phosphates, and $M_2(SO_4)_3$ for sulfates.

Additionally, phosphates and silicates exhibit approximately twice the capacity of sulfates, indicating the possible exchange of two Ca^{2+} in phosphates and silicates compared to one Ca^{2+} in sulfates. Among phosphate and silicate compounds, phosphate NaSICONs exhibit consistently a higher theoretical capacity than silicates owing to the lower mass of the charged composition (i.e., $Ca_{0.5}M_2(PO_4)_3$ vs $Ca_2M_2(SiO_4)_3$).

The predicted average voltages for silicate and phosphate Ca-NaSICONs increase monotonically from Ti to Ni (Figure 2), consistent with trends in the crystal reduction potentials across the $+2 \leftrightarrow +4$ oxidation states of the transition metals. Note that trends in crystal reduction potentials are qualitatively consistent with the standard reduction potentials for several systems.⁷⁶ Indeed, the average Ca intercalation voltage raises from ~1.6 V (1.0 V) vs Ca in the Ti-phosphate (silicate) to ~4.7 V (4.2 V) in the Ni-phosphate (silicate). Additionally, the average voltages of the phosphate Ca-NaSICONs are persistently higher than the corresponding silicates for all transition metals, which is expected as a result of the stronger inductive effect exerted by P5+ compared to Si4+ on the transition metal octahedra. 77,78 For example, the estimated average Ca (de)intercalation voltage in Ca, Mn₂(PO₄)₃ is ~3.6 V vs Ca, which is ~0.6 V higher than the corresponding $Ca_xMn_2(SiO_4)_3$. Furthermore, our predicted Ca intercalation voltage in $Ca_xV_2(PO_4)_3$ (~2.6 V) is lower than the experimentally reported $\sim 3.2~V$ for Ca intercalation in NaV₂(PO₄)₃, 43,79 where the difference between theoretical and experimental voltages can be attributed to the presence of the residual Na in the Ca-NaSICON structure. Thus, the phosphate NaSICONs should exhibit higher energy densities compared to the silicate analogues owing to their higher intercalation voltages and marginally superior gravimetric capacities.

In the case of sulfates, the calculated average voltages exhibit a nonmonotonic trend, with Cr and Fe sulfates showing a markedly lower voltage than their adjacent transition metals. Importantly, Mn, Co, and Ni-based sulfates exhibit higher voltages than the corresponding silicates and phosphates. This observation can be partly attributed to the stronger inductive effect of S⁶⁺ compared to P⁵⁺ and Si^{4+,77,78} V-sulfate and Fesulfate Ca-NaSICONs exhibit identical voltages to Vphosphate (\sim 2.6 V) and Fe-silicate (\sim 3.2 V), respectively, highlighting the balancing effect of the inductive effect differences and the oxidation state of the transition metal undergoing redox. Note that higher oxidation states typically result in higher voltages and that all sulfate Ca-NaSICONs involve transition metals undergoing a $M^{3+} \leftrightarrow M^{2+}$ redox (as constrained by charge neutrality) compared to $M^{4+} \leftrightarrow M^{2+}$ in phosphate and silicate Ca-NaSICONs. Also, the high stability of the Cr3+ and Fe3+ oxidation states, attributed to their highspin t_{2g}^3 and $t_{2g}^3e_g^2$ electronic configurations, respectively, ^{80,81°} in their corresponding charged sulfate compositions is the primary reason for the local minimum in calculated voltages at Cr and Fe (Figure 2). Furthermore, the lower voltage of Cr sulfate (\sim 1.8 V), compared to the corresponding silicate (\sim 2.2 V) and phosphate (\sim 2.7 V), can also be largely attributed to the stability of Cr3+

Intuitively, we expect Ti-based Ca-NaSICONs to be candidate materials for Ca-anodes instead of cathodes, owing to their lower voltages across the polyanionic groups (1.0–1.6 V) investigated here. In terms of intercalation voltages alone, Co- and Ni-based Ca-NaSICONs are promising (>4 V across polyanionic systems), but their experimental accessibility given

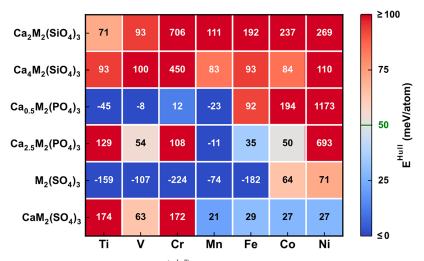


Figure 3. Energy above or below the ground-state convex hull (E^{hull}) extracted from the DFT calculations for charged and discharged silicate (top two rows), phosphate (middle two rows), and sulfate (bottom two rows) Ca-NaSICON. Each column represents a given 3d transition metal. Blue (red) squares indicate high degrees of stability (instability), with the specific E^{hull} value of each composition listed as a text annotation in the corresponding square. The green line on the legend bar indicates the rule-of-thumb $E^{hull} \sim 50$ meV/atom threshold for experimental synthesizability.

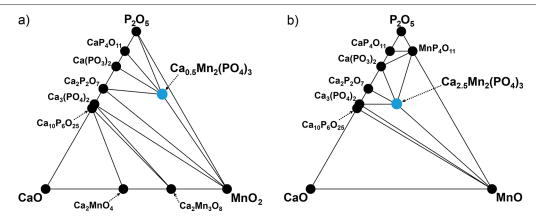


Figure 4. Calculated 0 K Ca–Mn–P–O quaternary phase diagrams showing the charged (panel a) and discharged (panel b) Ca-NaSICON (i.e., $Ca_xMn_2(PO_4)_3$) compositions. For ease of visualization, the pseudoternary projections of the Ca–Mn–P–O chemical space are plotted, namely, $CaO-P_2O_5$ –MnO $_2$ in panel (a) and $CaO-P_2O_5$ –MnO in panel (b). All stable compositions within the phase diagram are indicated by solid black circles, while solid black lines indicate tie-lines. The Ca-NaSICON compositions are indicated by solid light blue circles.

the stability limits^{11,82} of current Ca electrolytes is uncertain, while their potential synthesizability is discussed in the following section. With the exception of V- and Cr-silicates (showing average voltages <2.5 V), other V-, Cr-, Mn-, and Fesystems are candidates exhibiting moderate intercalation voltages (2.5–4.2 V). Combining the average voltages with the theoretical capacities, we observe the phosphate-based Ca-NaSICONs to be more promising than silicates and sulfates in terms of the gravimetric energy density. However, the practical realization of any Ca-NaSICON system as a Ca-cathode clearly depends on its thermodynamic stability and experimental accessibility.

3.3. Thermodynamic Stabilities of Ca-NaSICONs. Figure 3 depicts the thermodynamic (in)stability of the charged and discharged Ca-NaSICON compositions based on the silicate (top rows), phosphate (middle rows), and sulfate (bottom rows) host chemistry, with each column representing a transition metal. The green line on the legend bar indicates the $E^{hull} \sim 50 \text{ meV/atom rule-of-thumb}$ synthesizability threshold. Although we use an empirical synthesizability threshold here, the eventual formation, or not,

of a metastable compound depends critically upon the thermodynamic behavior of the competing phases. Text annotation in each square indicates the E^{hull} for the corresponding composition. The 0 K convex hulls for the charged and discharged $\mathrm{Ca_xMn_2(PO_4)_3}$ NaSICON are depicted in Figure 4, while the convex hulls for the other NaSICONs are compiled in Figures S5–S7 of the Supporting Information. For all unstable ($E^{hull} > 50$ meV/atom) or metastable (0 < $E^{hull} \leq 50$ meV/atom) Ca-NaSICONs, the predicted decomposition products are compiled in Table S1, while for the stable ($E^{hull} \leq 0$ meV/atom) Ca-NaSICONs, the "adjacent" phases 65 in their corresponding quaternary phase diagrams are listed in Table S2.

Importantly, for all transition metals considered (both charged and discharged), silicate Ca-NaSICON compositions exhibit a large positive E^{hull} (\sim 71–706 meV/atom), which is beyond the \sim 50 meV/atom threshold. This indicates their high thermodynamic instabilities. For instance, Ca₂Mn₂(SiO₄)₃ ($E^{hull} = \sim$ 111 meV/atom) is predicted to decompose to other stable phases in the Ca–Mn–Si–O quaternary phase diagram, including Ca₂Mn₃O₈, CaSiO₃, and SiO₂ (see Figure S5g and

Table S1). Thus, we expect all silicate-based Ca-NaSICONs to decompose into other stable phases and thereby are not promising for CIBs.

In the case of phosphates, both charged $(Ca_{0.5}Ni_2(PO_4)_3, E^{hull} \sim 1173 \text{ meV/atom})$ and discharged $(Ca_{2.5}Ni_2(PO_4)_3, E^{hull} \sim 693 \text{ meV/atom})$ Ni-NaSICONs show high E^{hull} , clearly indicating the lack of synthesizability of these potential high-voltage phases. Similarly, unstable phosphate Ca-NaSICONs include $Ca_{2.5}Ti_2(PO_4)_3$ ($E^{hull} \sim 129 \text{ meV/atom})$, $Ca_{2.5}Cr_2(PO_4)_3$ (108 meV/atom), $Ca_{0.5}Fe_2(PO_4)_3$ (92 meV/atom), and $Ca_{0.5}Co_2(PO_4)_3$ (194 meV/atom), all of which lie above the 50 meV/atom threshold. Notably, the E^{hull} for $Ca_{0.5}Cr_2(PO_4)_3$ (\sim 12 meV/atom), $Ca_{2.5}Fe_2(PO_4)_3$ (\sim 35 meV/atom), and $Ca_{2.5}Co_2(PO_4)_3$ (\sim 50 meV/atom) lie on/below the stabilization threshold, while $Ca_{2.5}V_2(PO_4)_3$ (\sim 54 meV/atom) lies only marginally above the threshold, signifying that these compounds may be obtained experimentally.

We find a set of phosphate and sulfate-based Ca-NaSICONs that are thermodynamically stable. Among the phosphate materials, the stable frameworks include $Ca_{0.5}V_2(PO_4)_3$ (~-8 meV/atom), $Ca_{0.5}Mn_2(PO_4)_3$ (~-23 meV/atom), $Ca_{2.5}Mn_2(PO_4)_3$ (~-11 meV/atom), and $Ca_{0.5}Ti_2(PO_4)_3$ $(E^{hull} \sim -45 \text{ meV/atom})$. Note that we depict the pseudoternary projections of the Ca-Mn-P-O chemical space corresponding to the charged Ca_{0.5}Mn₂(PO₄)₃ and the discharged Ca_{2.5}Mn₂(PO₄)₃ compositions in Figure 4. Ca has been experimentally shown to reversibly intercalate in NaV₂(PO₄)₃, 43,79 which is in good agreement with our predictions of Ca_{0.5}V₂(PO₄)₃ being thermodynamically stable. Furthermore, the presence of residual Na in $V_2(PO_4)_3$ may stabilize the Ca-NaSICON framework as the Ca concentration increases (i.e., toward the discharged composition), thus promoting electrochemical reversibility. Among sulfate Ca-NaSICONs, we predict all charged compositions, except $Co_2(SO_4)_3$ ($E^{hull} \sim 64$ meV/atom) and $Ni_2(SO_4)_3$ (~ 71 meV/atom), to be thermodynamically stable, in agreement with the fact that $Ti_2(SO_4)_3$, $Cr_2(SO_4)_3$, and $Fe_2(SO_4)_3$ are experimentally synthesizeable. 83–85 Additionally, our predictions of a high degree of thermodynamic stability of V₂(SO₄)₃ $(E^{hull} \sim -107 \text{ meV/atom})$ and $Mn_2(SO_4)_3$ $(E^{hull} \sim -74 \text{ meV/})$ atom) indicate a promise that these compounds may be synthesized in the future. Several discharged sulfate Ca-NaSICONs that are metastable include $CaMn_2(SO_4)_3$ (~21 meV/atom), CaFe₂(SO₄)₃ (~29 meV/atom), CaCo₂(SO₄)₃ (\sim 27 meV/atom), and CaNi₂(SO₄)₃ (\sim 27 meV/atom).

For practical realization of potential cathodes, both the charged and the discharged phases should be preferably thermodynamically stable to prevent any competing decomposition/conversion reactions⁸⁶ from occurring. Under such a constraint, we only find Ca₂Mn₂(PO₄)₃ (Figure 4) to be a candidate cathode. However, there are known examples of cathodes in other electrochemical systems, where either the charged or the discharged configurations are metastable but exhibit good, reversible performance. 87-90 Thus, if we allow for either the discharged or the charged phase to be stable with the corresponding charged or discharged phase to be metastable, we find Mn- and Fe-sulfate Ca-NaSICONs to be possible candidates, albeit with lower predicted energy densities compared to the Mn-phosphate (Table 2). Given that Ca has been experimentally intercalated into the NaV₂(PO₄)₃ structure⁴³ and given that Ca_{2.5}V₂(PO₄)₃ lies only marginally above the 50 meV/atom threshold, we observe the Vphosphate Ca-NaSICON to be a cathode candidate as well.

Other materials that may be promising from a purely synthesizability perspective (i.e., $E^{hull} \leq 50 \text{ meV/atom}$) and not necessarily with cathode applications are $Ca_{2.5}Fe_2(PO_4)_3$, $Ca_{2.5}Co_2(PO_4)_3$, $Ca_{0.5}Ti_2(PO_4)_3$, $Ca_{0.5}Cr_2(PO_4)_3$, $Ca_{0.5}Co_2(SO_4)_3$, $Ca_{0.5}Co_2(SO_4)_3$, and $Co_2(SO_4)_3$.

3.4. Migration Barriers of Ca^{2+} in Ca-NaSICONs. Apart from high intercalation voltage and thermodynamic stability, a Ca-cathode should also exhibit facile Ca^{2+} diffusion to facilitate charge/discharge processes at reasonable rates. The migration of divalent Ca^{2+} in solids is usually poorer than that of monovalent ions (e.g., Li^+ , Na^+), 68 typically attributed to Ca^{2+} 's stronger electrostatic interactions leading to local structural distortion during migration of ions. We consider a fairly liberal maximum allowed E_m of ~ 1000 meV for identifying potential candidates with reasonable rate performance (dashed green line in Figure 5), similar to our previous Ca-cathode screening study.

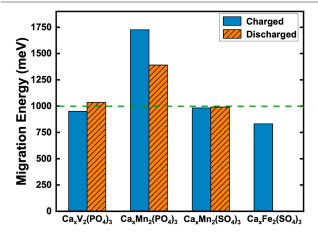


Figure 5. Calculated migration energy barriers of Ca^{2+} in the representative Ca-NaSICON cathode materials, such as $Ca_xV_2(PO_4)_3$, $Ca_xMn_2(PO_4)_3$, $Ca_xMn_2(SO_4)_3$, and $Ca_xFe_2(SO_4)_3$. The solid light blue and hatched orange bar indicate the Ca^{2+} migration energy barriers of the charged and discharged compositions. The horizontal dashed green line indicates the maximum tolerance of the migration energy barrier ($E_m = 1000 \text{ meV}$) for a facile migration of Ca-ion in a given anion framework. The minimum energy pathways of Ca^{2+} migration are shown in Figure S9.

The GGA-calculated E_m for the candidate materials, as identified by our voltage and stability calculations, namely, $Ca_xV_2(PO_4)_3$, $Ca_xMn_2(PO_4)_3$, $Ca_xMn_2(SO_4)_3$, and $Ca_xFe_2(SO_4)_3$, are displayed in Figure 5. The E_m values charged and discharged Ca-NaSICON compositions are represented by the light solid blue and hatched orange bars of Figure 5. Despite multiple restarts, the GGA-NEB calculation did not converge for the case of $CaFe_2(SO_4)_3$, explaining the lack of data for the discharged Fe-sulfate Ca-NaSICON. Apart from the candidate Ca-NaSICONs, we have also evaluated E_m for Ca^{2+} migration in a wider set of metastable Ca-NaSICONs using SCAN, including $Ca_{0.5}Cr_2(PO_4)_3$, $Ca_{0.5}Fe_2(PO_4)_3$, $Ca_{0.5}Co_2(PO_4)_3$, $Ca_{0.5}Co_2(PO_4)_3$, $Ca_{0.5}Co_2(SO_4)_3$, $CaV_2(SO_4)_3$, $CaCr_2(SO_4)_3$, $CaCr_2(SO_4)_3$, which is compiled in Figure S11.

Importantly, we find that the E_m of Ca^{2+} in charged V-phosphate Ca-NaSICON (~951 meV) is within the maximum tolerable E_m of ~1000 meV, while the discharged phase (E_m ~ 1035 meV) lies marginally above the tolerance limit. This is in

qualitative agreement with the experimental demonstration of Ca (de)intercalation in NaV₂(PO₄)₃.⁴³ Despite the similarity of the ionic size between Ca2+ and Na+ in an octahedral coordination environment and the migration pathways occurring across similar sites (Na1 to Na2 or Ca1 to Ca2), the $Ca^{2+}E_m$ is higher than that of Na^+ in the NaSICON frameworks. This can be attributed to the stability of Ca in the octahedral Ca1 (or Na1) site compared to Na. For example, Jeon et al. recently elucidated that, upon reversible Ca (de)intercalation in NaV₂(PO₄)₃, Ca preferably occupies the Na1 (or the Ca1) site by displacing the Na atoms during charge and discharge processes, signifying higher stability of Ca compared to Na at the Na1 (or the Ca1) site.⁷⁹ Thus, we believe that the higher migration barrier of Ca²⁺ in NaSICON frameworks is primarily due to the stability of Ca in Ca1 sites rather than any significant variations in the migration pathway itself.

In the case of $Ca_xMn_2(PO_4)_3$, the calculated E_m values in both the charged $(Ca_{0.5}Mn_2(PO_4)_3, \sim 1726 \text{ meV})$ and discharged $(Ca_{2.5}Mn_2(PO_4)_3, \sim 1391 \text{ meV})$ Mn-phosphate are well beyond the $\sim 1000 \text{ meV}$ feasibility limit. We do not expect $Ca_xMn_2(PO_4)_3$ to be a feasible Ca-cathode despite its attractive intercalation voltage and thermodynamic stability. The E_m values of both the charged and the discharged phases of $Ca_xMn_2(SO_4)_3$ ($\sim 983 \text{ meV}$ and $\sim 991 \text{ meV}$, respectively) and $Ca_xFe_2(SO_4)_3$ ($\sim 832 \text{ meV}$) lie below the $\sim 1000 \text{ meV}$ tolerance limit, signifying reasonable Ca-ion migration within these frameworks. In summary, we identify $Ca_xV_2(PO_4)_3$, $Ca_xMn_2(SO_4)_3$, and $Ca_xFe_2(SO_4)_3$ as potentially feasible Ca-cathode materials, given their intercalation voltage, thermodynamic (meta)stability, and accessible $Ca^{2+}E_m$.

4. DISCUSSION

Using DFT calculations, we have explored the Ca-NaSICON frameworks as potential cathode materials for CIBs, which are an important technological alternative to current state-of-theart LIBs. Specifically, we have evaluated the average Ca intercalation voltage, the theoretical capacity, and the 0 K thermodynamic stability for $\text{Ca}_x\text{M}_2(\text{ZO}_4)_3$ systems, where M is a 3d transition metal except Sc, Cu, and Zn, Z is either Si, P, or S, and x is determined by the overall charge neutrality of the Ca-NaSICON framework. Our calculated data indicates that V-phosphate, Mn- and Fe-sulfate Ca-NaSICONs are promising as Ca-cathodes, among the 21 NaSICON compositions considered, with the overall screening process shown in Figure 6.

In this work, we have used the SCAN+U calculations for both our voltage and stability predictions, while GGA was used to calculate the migration barriers. A recent assessment ⁹⁵ has shown that SCAN+U functional exhibits good qualitative trends but tends to overestimate voltages and instabilities in Liintercalation electrodes. We expect similar trends to be extended to Ca-NaSICONs as well, which is one of the reasons that we used a higher E^{hull} stabilization threshold (\sim 50 meV/atom) than in previous studies (\sim 25 meV/atom). ^{25,89} However, we expect SCAN+U's trends of high instabilities in silicate-NaSICON frameworks to be a fair evaluation of the Ca-M-Si-O chemical space.

Our calculated 0 K stability data indicates a high degree of instability ($E^{hull} \gg 50$ meV/atom), at both the charged and discharged states, for silicate Ca-NaSICONs. For example, Ca₂Mn₂(SiO₄)₃ and Ca₄Mn₂(SiO₄)₃ exhibit $E^{hull} \sim 111$ meV/atom and ~ 83 meV/atom and are expected to decompose into

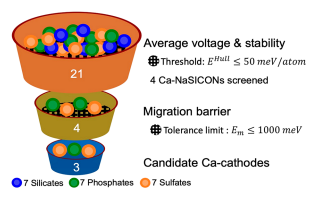


Figure 6. Summary plot depicting the screening process of NaSICON frameworks as Ca-cathodes based on calculated average voltage, thermodynamic stability, and migration barrier. The blue, green, and orange spheres represent silicate, phosphate, and sulfate-based Ca-NaSICONs, respectively. The mesh in the funnel plot represents the threshold for experimental synthesizability ($E^{Hull} \leq 50 \text{ meV/atom}$) and the maximum tolerance of the migration barrier ($E_m \leq 1000 \text{ meV}$), resulting in our three candidate compositions, $\text{Ca}_x \text{V}_2(\text{PO}_4)_3$, $\text{Ca}_x \text{Mn}_2(\text{SO}_4)_3$, and $\text{Ca}_x \text{Fe}_2(\text{SO}_4)_3$.

Ca₂Mn₃O₈, CaSiO₃, and SiO₂ and Mn₅Si₃O₁₂, Ca₂SiO₄, and Mn, respectively (Table S1). Such high instabilities among silicate-NaSICONs are driven largely by the high thermodynamic stability of competing ternary Ca-Si-O compounds, such as CaSiO₃, Ca₂SiO₄, Ca₃Si₂O₇, and Ca₃SiO, several of which are naturally occurring minerals.⁹⁶ Indeed, the E^{hull} of $Ca_2Mn_2(SiO_4)_3$ and $Ca_4Mn_2(SiO_4)_3$ reduce to ~98 meV/atom and ~24 meV/atom, respectively, if all calcium silicate phases are removed from the Ca-Mn-Si-O quaternary phase diagram. In the case of phosphates and sulfates, the number of thermodynamically stable Ca-P-O (Ca₃(PO₄)₂) and Ca-S-O (CaSO₄) are fewer, thereby driving the instability/ metastability of the corresponding Ca-NaSICON compositions to a lower extent compared to the silicates. On one hand, in the search for new Ca-intercalation frameworks, especially with polyanionic materials, chemical spaces containing several known Ca-containing ternary compounds may be avoided. On the other hand, given the diverse chemical space of NaSICON frameworks, tremendous efforts have been made by exploring several transition metal combinations for Na-ion battery cathodes.^{37,91–94} Apart from the identified candidate Ca-cathodes in this work, exploring combinations of transition metals within a single Ca-NaSICON framework thus represents a promising strategy for identifying potentially better Ca-cathodes.

5. CONCLUSIONS

Designing cathode intercalation hosts that can reversibly exchange Ca is an important requirement in developing CIBs, as an alternative for the state-of-the-art LIBs. Using first-principles calculations, we explored the Ca-NaSICON frameworks that classically act as Na intercalation frameworks, as potential cathode materials for CIBs. We evaluated the theoretical energy density (i.e., the average Ca intercalation voltage and the theoretical capacity), the 0 K thermodynamic stability for 21 different $Ca_xM_2(ZO_4)_3$ systems, where M=Ti, V, Cr, Mn, Fe, Co, or Ni and Z=Si, P, or S, and the Ca^{2+} migration barriers in select Ca-NaSICON frameworks. The calcium content (x in $Ca_xM_2(ZO_4)_3$) was determined by the overall charge neutrality of the Ca-NaSICON framework, for a possible range of $M^{4+} \leftrightarrow M^{2+}$ oxidation states, giving rise to

charged compositions of $Ca_2M_2(SiO_4)_3$, $Ca_{0.5}M_2(PO_4)_3$, and $M_2(SO_4)_3$ and discharged compositions of $Ca_4M_2(SiO_4)_3$, $Ca_{2.5}M_2(PO_4)_3$, and $CaM_2(SO_4)_3$ across all transition metals.

We find a monotonic increase in the calculated average voltages for silicates and phosphates through the 3d series (i.e., from Ti to Ni), while sulfate Ca-NaSICONs show local minima in calculated voltages at Cr and Fe, attributed to the stability of the Cr³⁺ and Fe³⁺ states. In terms of capacities, we observe phosphates (sulfates) consistently exhibiting the highest (lowest) capacity for all transition metals primarily due to the possibility of two (one) Ca exchange. In terms of thermodynamic stability, we find none of the silicate frameworks to be below our $E^{hull} = 50 \text{ meV/atom threshold}$, while several phosphate and sulfate Ca-NaSICONs are either stable or metastable. Combining our calculated voltage, capacity, thermodynamic stability, and migration barrier data, we find the $Ca_xV_2(PO_4)_3$, $Ca_xMn_2(SO_4)_3$, and $Ca_xFe_2(SO_4)_3$ to be promising candidate cathodes for CIBs. We hope that our work inspires further theoretical and experimental studies into Ca-intercalation frameworks, particularly within the chemically diverse Ca-NaSICON space, and drive the practical realization of CIBs.

ASSOCIATED CONTENT

Data Availability Statement

All the computational data presented in this study will be made freely available to all on our GitHub repository (https://github.com/sai-mat-group/ca-nasicon-electrodes).

5 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.chemmater.2c02841.

PAW potentials used, schematics of Ca-vacancy configurations of all Ca-NaSICONs considered, pictorial representations of all computed convex hulls at 0 K, list of decomposition products of unstable/metastable Ca-NaSICONs and adjacent phases of stable Ca-NaSI-CONs, and the migration barriers data (PDF)

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Notes

The authors declare no competing financial interest.

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