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First-principles evaluation of multi-valent cation insertion into orthorhombic $V_2O_5^{\dagger}$

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A systematic first-principles evaluation of the insertion behavior of multi-valent cations in orthorhombic V₂O₅ is performed. Layer spacing, voltage, phase stability, and ion mobility are computed for Li⁺, Mg²⁺, Zn²⁺, Ca²⁺, and Al³⁺ intercalation in the α and δ polymorphs.

A promising and realistic strategy to improve the energy density beyond the capability of current Li-ion battery technology is to transition to a battery architecture based on shuttling multivalent (MV) ions (*e.g.* Mg^{2+} or Ca^{2+}) between an intercalation cathode host and MV metal anode.^{1,2} Specifically, improvement in the volumetric energy density arises from the combination of using a multi-valent metal as the anode as opposed to an insertion structure (*e.g.* 3833 mAh cm⁻³ volumetric capacity for Mg metal compared to 800 mAh cm⁻³ for graphite), and storing more charge per ion in the cathode.^{3,4}

One of the major bottlenecks preventing the development of MV battery technology, however, is the poor electrochemical performance of potential MV cathode materials, thought to originate predominantly from poor MV ion mobility in the intercalation host structure.^{4–6} Moreover, the simultaneous challenge of developing functioning MV anodes and electrolytes compatible with candidate cathode materials has limited the ability to experimentally isolate and evaluate cathode electrochemical performance,⁷ and as such there is a general dearth of reliable data on MV ion intercalation in the literature to date to guide the ongoing search for new MV cathode materials with improved performance.

Nevertheless, reversible electrochemical Mg²⁺ intercalation has been successfully demonstrated in a handful of cathode hosts, namely Chevrel Mo₆S₈ (~135 mAh g⁻¹ capacity at ~1.0–1.3 V ν s. Mg metal),² as well as layered V₂O₅ (~150 mAh g⁻¹ at ~2.3–2.6 V)^{5,6} and MoO₃ (~220 mAh g⁻¹ at ~1.7–2.8 V).⁵ The orthorhombic V₂O₅ structure is especially interesting because it has also demonstrated the ability to reversibly intercalate Ca²⁺ and Y³⁺ in addition to Mg²⁺ ions.⁶ First-principles calculations (described in more detail in the ESI†) have proven to be an accurate and effective method to systematically assess the electrochemical properties of Li-ion batteries,^{8–10} and have also been used to study the process of ion intercalation in layered materials, such as graphite¹¹ and V₂O₅.^{12–15} In this work, we have performed a systematic first principles study of MV ion intercalation in the orthorhombic α - and δ -V₂O₅ polymorphs by evaluating the structural change, voltage, thermodynamic stability, and intercalant mobility for Li⁺, Mg²⁺, Zn²⁺, Ca²⁺, and Al³⁺ insertion and comparing to data in the literature when available.

The crystal structure and intercalation sites of the α - and δ -V₂O₅ polymorphs^{16–19} are shown in Fig. 1. Perpendicular to



Fig. 1 (a) The V₂O₅ structure of both the α and δ polymorphs on the *b*-*c* plane with the yellow spheres indicating the intercalant sites while (b) shows the α and δ polymorphs on the *a*-*b* plane. As indicated by the dashed blue regions, both the polymorphs differ by a change in the stacking of the V₂O₅ layers.

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the *b*-axis (*i.e.* in the a-c plane), the orthorhombic V₂O₅ structure consists of layers of alternating corner- and edgesharing VO₅ pyramids (shown in red), each consisting of 4 V-O bonds that form the base and one short V=O bond that forms the apex. The intercalation sites (yellow spheres) are situated in between the layers, and assuming no limitation in the number of redox centers, the theoretical gravimetric capacities for AV₂O₅ where A = Li, Mg, Zn, Ca and Al are 142, 260, 217, 242 and 385 mAh g^{-1} , respectively. Structurally, the main difference between the α and δ polymorphs is a shift in the layer stacking, indicated by the dashed blue lines in Fig. 1b, with alternate V_2O_5 layers displaced in the *a*-direction by half a lattice spacing, accompanied by a change in the interlayer distance and the anion coordination environment of the intercalation sites.¹⁶ While 8 oxygen atoms coordinate the intercalant ion in α (for Mg, there are two Mg–O bonds with length \sim 2.11 Å, two with \sim 2.39 Å, and four with ~ 2.46 Å, respectively), "4 + 2" oxygen atoms coordinate the intercalant in δ (for Mg, there are four Mg–O bonds with length ~2.05–2.07 Å, and two with ~2.33 Å).

In Fig. 2a, the interlayer spacings in the α and δ polymorphs (filled and hollow bars, respectively) are shown for empty V₂O₅ and intercalated AV₂O₅, where A = Li, Mg, Zn, Ca, and Al. To better capture the increased effect of van der Waals effects in the deintercalated limit, the interlayer spacings for empty V₂O₅ (4.46 Å for α ; 5.03 Å for δ) are calculated using the vdW-DF2 functional^{20,21} rather than standard DFT as the latter significantly overestimates this spacing (4.75 Å for α ; 5.27 Å for δ) compared to experiment (4.37 Å for α).^{12,15,18} As detailed in the ESI,† Al³⁺ intercalation in the α -V₂O₅ structure is found to be mechanically unstable and relaxes to the δ polymorph in our calculations, and we therefore remove it from further consideration in this study.

At the same intercalant composition, the δ structures consistently have larger layer spacings than α , ~ 3–5% larger for Li, Mg, and Zn and ~10-12% for Ca and empty V_2O_5 . With the exception of Ca intercalation, which increases the layer spacing by more than 10% in both polymorphs, the change in the layer spacing is much smaller in δ than α , less than 2% for Li⁺, Mg²⁺, Zn^{2+} , and Al^{3+} intercalation in δ -V₂O₅ compared to ~9–14% for Li⁺, Mg^{2+} , Zn^{2+} , and Ca^{2+} in α -V₂O₅. The behavior for Ca^{2+} is consistent with intercalation in the spinel system,²² where the volume change is also much larger than for Li⁺, Mg²⁺, Zn²⁺, and Al³⁺ intercalation, and in general may be attributed to the larger ionic radius of Ca²⁺ in comparison to the other ions.²³ Al³⁺ intercalation in δ -V₂O₅, in contrast to the other ions considered, is accompanied by a contraction of the layers, which is consistent with its small ionic radius and higher positive charge density that strengthens the attraction with nearby oxygen ions.

The average voltages of the compounds computed using the method of Aydinol *et al.*²⁴ are plotted in Fig. 2b and are referenced to the potential of the bulk metal of the corresponding intercalating ion (*i.e.*, Li metal for Li⁺ intercalation, *etc.*). The average voltages computed for Li, Mg, and Ca intercalation compare very well to available experimental data: \sim 3.2–3.4 V for Li measured by Delmas *et al.*,¹⁶ \sim 2.2–2.4 V for Mg measured by Gershinsky *et al.*,⁵ and \sim 2.4–3.1 V for Ca measured by



Fig. 2 (a) Plots the layer spacing values for the empty and intercalated versions of AV₂O₅ (A = Li, Mg, Ca, Zn and Al) for both the α and δ polymorphs. (b) Displays the calculated average voltage values for the intercalation of the different ions and (c) shows the energy above hull, which quantifies the stability of a structure, for the empty and intercalated versions of α and δ . The filled regions in all the graphs correspond to the α structure while the hollow regions correspond to the δ structure. Note that the energy above hull for α -CaV₂O₅ is 0 meV per atom, implying that it is a ground state configuration in the Ca–V–O system.

Amatucci *et al.*⁶ In general, the Li polymorphs have the highest voltage, followed by Ca, Mg, Al, and Zn, which reflects both the same order and approximately the same potential difference indicated by the electrochemical series ($-3.04 \text{ V} \nu s$. SHE for Li, -2.86 V for Ca, -2.37 V for Mg, -1.66 V for Al, and -0.76 V for Zn). In comparison, the voltage difference between the V₂O₅ polymorphs is much smaller for a given intercalation chemistry.

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For Li, Mg, and Zn the insertion voltage is higher in δ (3.36 V, 2.56 V, and 1.09 V, respectively) than in α (3.18 V, 2.21 V, and 0.68 V), unlike for Ca where α is higher (3.13 V for α ; 3.02 V for δ).

Fig. 2c displays the energy above the convex ground state energy hull (E^{hull}) of the deintercalated and intercalated V₂O₅ polymorphs with respect to the intercalant-V–O ternary phase diagram. The ternary ground state hulls were determined from the available calculated compounds in the Materials Project database.²⁵ A predicted thermodynamically stable structure will have a E^{hull} value of 0 meV per atom while higher (more positive) E^{hull} values indicate greater instability, which may be reflected in experimental difficulties in synthesis or decomposition during battery operation. Note that the E^{hull} values calculated here reflect the ground state (*i.e.* 0 K), and entropy contributions, which scale with $k_{\rm B}T$, can stabilize certain structures at higher temperatures.

In the deintercalated limit, V₂O₅ is thermodynamically stable in the α phase, but δ is only ~13 meV per atom higher in energy, indicating the possibility of metastability at room temperature. For Li intercalation, the α and δ structures are 82 meV per atom and 57 meV per atom more unstable than the ground state orthorhombic γ-LiV2O5 structure, which has a different orientation of the VO₅ pyramids¹⁶ along the *c*-direction shown in Fig. 1a, but the δ structure can remain metastable and has shown to be reversibly cycled electrochemically.¹⁶ δ-MgV₂O₅, which has been synthesized experimentally,¹⁷ is only ~ 27 meV per atom more unstable (compared to ~ 102 meV per atom for α) than the thermodynamic ground state, a two-phase equilibrium consisting of MgVO₃ and VO₂. Similarly δ -ZnV₂O₅ is only ~31 meV per atom more unstable than the ground state (ZnO and VO₂), indicating that a metastable synthesis comparable to the Mg system may be possible. As Al intercalated α -V₂O₅ displays mechanical instability in our calculations, when relaxed its energy is not defined, but the Al intercalated δ -phase is ~158 meV per atom unstable compared to the ground state ternary equilibrium of Al₂O₃, VO₂ and V_3O_5 . With the exception of α -CaV₂O₅, which is the ground state in the intercalated Ca– V_2O_5 system, the δ structures tend to be more stable than α in the discharged state (by 25 meV per atom for Li; 75 meV per atom for Mg; and 91 meV per atom for Zn), and accordingly the insertion voltages for δ are higher than α for Li, Mg, and Zn insertion but lower for Ca insertion, as observed in Fig. 2b. Given that the intercalant sites in α and δ are coordinated by 8 and "4 + 2" oxygen atoms respectively, the stability of the discharged δ -V₂O₅ structures for Li, Mg and Zn, and α -V₂O₅ for Ca align well with the preferred coordination environment of the respective ions, as tabulated by Brown.²⁶ Hence for intercalant ions that prefer a lower coordination number (*i.e.*, coordinated by a maximum of 6 neighboring atoms), an $\alpha \rightarrow \delta$ transition upon insertion in V₂O₅ is likely.

Fig. 3 displays the migration energies for intercalant diffusion along the *a*-direction in the α (Fig. 3a) and δ (Fig. 3b) polymorphs plotted against the normalized path distance calculated with the Nudged Elastic Band method.²⁷ The solid lines correspond to migration energies obtained in the empty lattice limit (charged state), and the dashed lines correspond to the fully intercalated limit (discharged state). As elaborated upon in the ESI,†



Fig. 3 The activation barriers for the diffusion of the different intercalating ions in the α and δ polymorphs are plotted in (a) and (b) respectively. The solid lines correspond to the empty lattice limit (charged state) while the hollow lines correspond to the full lattice limit (discharged state).

converging the migration energies in structures that exhibit a high degree of thermodynamic instability may not be possible, as was the case for Li, Mg, and Zn in the intercalated α -V₂O₅ structure, and for Ca in the intercalated δ -V₂O₅ structure. In lieu of determining the Mg migration barrier in the fully discharged α -V₂O₅ structure, we have computed the energy for Mg migration in a half intercalated structure with a specific ordering of Mg ions, referred to as the " ε " phase, which has also been observed in the Li-V₂O₅ system.¹⁶

In Fig. 3, the maximum energy difference encountered along the diffusion path defines the migration barrier (E_m) , which provides an approximate estimate of the ionic diffusivity. As a guide, at room temperature, $E_{\rm m} \sim 525$ meV corresponds to a diffusivity of $\sim 10^{-12}$ cm² s⁻¹, and a 60 meV increase (decrease) in the migration energy corresponds to an order of magnitude decrease (increase) in diffusivity. Due to stronger interactions between a multivalent intercalant and the surrounding anion environment, the migration barriers within the same host structure, for example Al³⁺, are generally higher than the divalent ion barriers (Mg²⁺, Zn²⁺, Ca²⁺), which are generally higher than the barriers for Li⁺. For the divalent intercalants, the trend in the migration barriers is Ca^{2+} (~1700–1900 meV) > Mg^{2+} $(\sim 975-1100 \text{ meV}) > Zn^{2+} (\sim 305 \text{ meV})$ in the α -phase, but Mg^{2+} $(\sim 600-800 \text{ meV}) > \text{Zn}^{2+} (\sim 375-425 \text{ meV}) > \text{Ca}^{2+} (\sim 200 \text{ meV})$ in the δ phase. The energy above the hull (Fig. 2c) ranked from the lowest to highest reflects this same trend, with $Ca^{2+} > Mg^{2+} > Zn^{2+}$ in α and Mg²⁺ > Zn²⁺ > Ca²⁺ for δ , and highlights the positive correlation between high intercalant mobility and low thermodynamic stability. For both V2O5 polymorphs considered, the change in the migration barrier from the deintercalated to intercalated limit for the same diffusing species is much smaller than the variation across intercalating ions.

Although the α and δ polymorphs of V₂O₅ are structurally very similar as earlier discussed, the anion coordination environment and therefore diffusion topology of the migrating intercalant

vary significantly, which accounts for the different shape of the migration energies seen in Fig. 3a and b. In the α phase, the stable insertion site is coordinated by 8 oxygen anions which is connected to the adjacent insertion site along the *a*-axis by a 3-coordinated shared face. The shape of the migration energies shown in Fig. 3a, therefore, reflect the change in coordination of $8 \rightarrow 3 \rightarrow 8$ encountered by the diffusing species with the migration barrier corresponding to passing through the shared face. For the δ phase, the stable insertion site adopts a "4 + 2" coordination and shares a corner with the adjacent insertion site along the *a*-axis. To migrate to this site, the intercalant passes through a 3-coordinated face shared with an intermediate 5-coordinated (pyramidal) site, and finally performs a symmetric hop to the next insertion site. The change in the anion coordination along the diffusion path is then "4 + 2" \rightarrow $3 \rightarrow 5 \rightarrow 3 \rightarrow$ "4 + 2", where occupation of the intermediate pyramidal site corresponds to a local minimum in the migration energy, as is reflected in Fig. 3b. Overall, the migration barriers are also lower in the δ phase compared to α (significantly lower for some cases), which we attribute in large part to the smaller coordination change during the migration process encountered in δ . Also, the change in the relative order of the migration barriers of divalent ions between α (Ca > Mg, Zn) and δ (Mg, Zn > Ca) can be explained by the correlation between the "preferred" coordination environments of the respective ions and the available anion coordination environments around the intercalation sites.²⁸ In a given structure, migration barriers are higher for an ion whose preferred coordination aligns with that of the coordination environment available for the intercalant site compared to an ion whose preferred coordination is different from that present in the structure. For example, Ca is in its preferred 8-coordinated site in α and hence has higher barriers than Mg and Zn, which are not in their respectively preferred 6 and 4 coordinated sites. Whereas in δ , Ca is present in an unfavored "4 + 2" coordinated site and hence has lower barriers than either of Mg or Zn, which are closer to their preferred coordination environments. Our results thus lend support to the hypothesis that coordination of the intercalation site is a good screening criterion for identifying fast multivalent cation diffusers.

An ideal MV cathode intercalation host must possess several properties – high capacity, high insertion voltage, and MV ion mobility, while simultaneously minimal structural change and thermodynamic instability. From the systematic first-principles study performed in this work, we are able to evaluate all of the candidate materials across each of these criteria. On the basis of ion mobility, Al^{3+} intercalation appears unfeasible at room temperature in V_2O_5 due to its prohibitively high migration barriers, and although Zn^{2+} intercalation is determined to be facile in both polymorphs and relatively stable in the δ phase, the insertion voltage is low. Mobility of Mg^{2+} and Ca^{2+} is determined to be poor in the α phase, but intercalation of these ions in the δ phase appear most promising, with sufficiently high voltage (3.02 V for Ca, and 2.56 V for Mg) and mobility ($E_m \sim 200$ meV for Ca and $\sim 600-800$ meV for Mg) albeit with moderate

thermodynamic instability (27 meV per atom for Mg and 40 meV per atom for Ca above the ground state hull in the discharged state).

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