

Charge Localization and Ordering in A$_2$Mn$_8$O$_{16}$ Hollandite Group Oxides: Impact of Density Functional Theory Approaches

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The phases of A$_2$Mn$_8$O$_{16}$ hollandite group oxides emerge from the competition between ionic interactions, Jahn-Teller effects, charge ordering, and magnetic interactions. Their balanced treatment with feasible computational approaches can be challenging for commonly used approximations in Density Functional Theory. Three examples (A = Ag, Li and K) are studied with a sequence of different approximate exchange-correlation functionals. Starting from a generalized gradient approximation (GGA), an extension to include van der Waals interactions and a recently proposed meta-GGA are considered. Then local Coulomb interactions for the Mn 3$d$ electrons are more explicitly considered with the DFT+$U$ approach. Finally selected results from a hybrid functional approach provide a reference. Results for the binding energy of the A species in the parent oxide highlight the role of van der Waals interactions. Relatively accurate results for insertion energies can be achieved with a low $U$ and a high $U$ approach. In the low $U$ case, the materials are described as band metals with a high symmetry, tetragonal crystal structure. In the high $U$ case, the electrons donated by A result in formation of local Mn$^{3+}$ centers and corresponding Jahn-Teller distortions characterized by a local order parameter. The resulting degree of monoclinic distortion depends on charge ordering and magnetic interactions in the phase formed. The reference hybrid functional results show charge localization and ordering. Comparison to low temperature experiments of related compounds suggests that charge localization is the physically correct result for the hollandite group oxides studied here. Finally, while competing effects in the local magnetic coupling are subtle, the fully anisotropic implementation of DFT+$U$ gives the best overall agreement with results from the hybrid functional.

Keywords: hollandite group oxides, manganese oxides, doped oxides, charge order, mixed valence, Jahn-Teller distortion, generalized gradient approximation, GGA, PBE, van der Waals interaction, Opt-B88, DFT+$U$, meta-GGA, SCAN, hybrid functional, HSE

I. INTRODUCTION

Manganese oxide minerals form in structures with an astonishing natural diversity and have a variety of practical applications. The hollandite group includes such specific minerals as hollandite, Ba$_2$Mn$_8$O$_{16}$, and cryptomelane, K$_x$Mn$_8$O$_{16}$. The backbone structure consists of edge-sharing MnO$_6$ octahedra that form double chains. These, in turn, connect through corner shared oxygens to form tunnels, as visualized in Fig. 1, resulting in an inherently one-dimensional structural feature. The additional ions such as Ba$^{2+}$, Ag$^+$, K$^+$, Na$^+$, Li$^+$, etc., occupy the larger cross-section tunnels. Water may also be incorporated in the tunnels. The ideal form of hollandite incorporates up to two ions per formula unit ($x = 2$) and the space group of the tetragonal cell is I4/m. The $\alpha$-MnO$_2$ structure corresponds to $x = 0$.

More broadly, the hollandite structure forms with tetravalent metal ions at the octahedral centers, e.g., with Mn$^{4+}$, Ti$^{4+}$, Cr$^{4+}$ and V$^{4+}$. With the incorporation of ions in the channels, there must be a corresponding reduction of ions in the backbone as well. The result can be a mix of Mn$^{4+}$ and Mn$^{3+}$ sites in the backbone or the inclusion of trivalent metal ions explicitly during synthesis, for instance Fe$^{3+}$. Naturally occurring minerals in the hollandite group often involve complex mixtures of cations both in the tunnels and within the octahedra. Depending on the ratio of the average ionic radii in the two positions, the crystal symmetry is observed to be lowered to monoclinic ($I2/m$), with the general trend indicating that relatively smaller ions in the tunnels correspond to monoclinic structures.

The hollandite group manganese oxides, and closely related structures with different sized tunnels, represent a tunable system with nanoscale, one-dimensional pores. The one-dimensional pores present a template for both fundamental studies of the impact of dimensionality on processes such as ionic diffusion and an internal surface support for chemical processes. In particular, hollandite group manganese oxides have been studied for use in catalysis and extensively for potential utility as cathodes in batteries based on Li$^+$, Na$^+$, and Mg$^{2+}$ as well. Facile synthesis of Ag-hollandite under mild conditions has made it more readily available for electrochemical studies. Furthermore, control of Ag insertion during synthesis results in clear changes in sample composition and structure with correlated changes in electrochemical characteristics. In particular, there is an interplay between Ag content, oxygen vacancy con-
structure oxides with the chemical formula $A_{\alpha}M_nO_{16}$ materials.

The present work is motivated by our need to assess the relationship of these structural degrees of freedom and the electrochemical response remains as an on-going challenge. The present work is motivated by our need to assess methods based on Density Functional Theory (DFT) for use in the broader exploration of phases formed in the Li$_x$Ag$_y$Mn$_8$O$_{16-z}$:(H$_2$O)$_w$ family of hollandite-derived materials.

The fundamental electronic structure of hollandite structure oxides with the chemical formula $A_2M_nO_{16}$ can be quite subtle, depending on the way the reduction implied by the tunnel cation $A$ is accommodated. Formally, the backbone transition metal $M$ is in a mixed valence state between $M^{4+}$ and $M^{3+}$. Furthermore, according to Hund’s rule, ions such as Mn$^{4+}$ and Mn$^{3+}$ with three and four $d$ electrons respectively should be in a high spin state locally. The octahedra in the parent $\alpha$-$\text{MnO}_2$ structure ($I4/m$) are close to ideal, with the six Mn-O bond lengths varying minimally (range of 0.02 Å near a value of 1.90 Å) implying a filled majority spin $t_{2g}$ manifold. The fourth $d$ electron for a local Mn$^{3+}$ will go to the $e_g$ manifold setting up the possibility for a local Jahn-Teller distortion to lower the electronic energy. However, the degree to which the excess electrons donated by the tunnel cations localize and local structural distortions reduce the symmetry, must emerge from competing factors: ionic radius ratio, local Coulomb interactions, the Jahn-Teller effect, and magnetic interactions. These factors will vary with the backbone metal cation $M$ and the tunnel cation $A$.

Not surprisingly, characterization of specific hollandite structure oxides has revealed phase diagrams with distinct magnetic and conducting phases. Two cases have been closely studied as a function of temperature. Near stoichiometric $K_2V_5O_{16}$ has a first order transition to a monoclinic phase at 170 K understood in terms of charge ordering of localized $V^{3+}$, dimerization along the tunnel axis and strong spin interactions. In $K_2Cr_3O_{16}$, an unusual sequence showed ferromagnetic ordering at 180 K followed by a metal-insulator transition at 95 K. The low temperature phase has a subtle monoclinic distortion, but there is no evidence for charge localization. For hollandite group manganese oxides, the magnetic measurements at low temperature have not been so cut, with the latest data interpreted to indicate a spin-glass phase. While naturally occurring cryptomelane exhibited a mild monoclinic distortion, room temperature characterization of synthesized hollandite group oxides showed a tetragonal phase ($I4/m$) for K, Li and Ag cases. In the Ba case, a monoclinic phase ($I2/m$) was found. However, no detailed studies at low temperature have been performed to our knowledge. Several studies based on DFT approaches have considered the ground state phase for $\alpha$-$\text{MnO}_2$ and ion insertion, also with some disagreement in the results regarding monoclinic distortion and magnetic order. Some of the differences can likely be traced to different choices in the approximate exchange-correlation functional.

In this paper, we revisit the application of the DFT-based tool-box to this challenging problem by critically comparing results obtained from several approximations to the treatment of exchange and correlation. We focus on $\alpha$-$\text{MnO}_2$ and the insertion of K, Ag and Li to form $A_2\text{Mn}_8\text{O}_{16}$. While in many cases synthesis of the ideal stoichiometry with two tunnel cations per unit cell has been difficult, we study this case to eliminate the additional complication of sampling configurations for occupancy of the A atom positions, as would be necessary for lower concentrations.

Starting with the version of the generalized gradient approximation (GGA) developed by Perdew, Burke and Ernzerhof (PBE), we consider the impact of van der Waals (vdW) interactions through the Opt-B88
functional.\textsuperscript{58} With these as a base, we then study the inclusion of the Hubbard correction term for the Mn 3d electrons, using the DFT+U approach,\textsuperscript{59,60} characterizing the physical and electronic structure as a function of the value of $U$ as well as the impact of the spherical approximation.\textsuperscript{62} This method has proven to be a cost effective way to substantially improve the treatment of systems with strong local Coulomb interactions and Jahn-Teller effects.\textsuperscript{60,62} For comparison, we also consider the Heyd, Scuseria and Ernzerhof (HSE) hybrid functional approach.\textsuperscript{63,64} Previous studies of perovskites suggest that it handles the subtelities of the open shell transition metals relatively accurately, albeit with increased computational cost.\textsuperscript{65} Finally, we consider the recently developed strongly constrained and appropriately normed (SCAN) meta-GGA for the exchange-correlation functional.\textsuperscript{66} Analysis of DFT+U and HSE for manganese oxide phases specifically suggest that both can account for structure and magnetic phases.\textsuperscript{67–69} Results for SCAN show that it may be an accurate and efficient alternative for manganese oxide energetics.\textsuperscript{70} Our study focuses on the treatment of physical properties resulting from mixed valence between Mn$^{3+}$ and Mn$^{4+}$.

Across the methodological choices, we examine energetics, charge localization, charge ordering, structural distortion and magnetic order. Using results based on HSE as a reference, we characterize the accuracy of the other functionals for the addition energy associated with Ag, Li and K, fundamental to predicting electrochemical properties. We then explore the implications for the crystal structure, finding a critical role for charge localization to form Mn$^{3+}$ centers. Charge localization is predicted by the HSE calculations and the GGA+U methods for moderate to large values of $U$. Correspondingly, electronically driven Jahn-Teller distortions also emerge, characterized by a local order parameter. They result in strong monoclinic distortions of the unit cell. We discuss the role of charge ordering in this case and the implications for the observed crystal structures. We also briefly discuss the magnetic phases formed. In agreement with prior work and experiment, we find that the competition between magnetic coupling mechanisms is delicate and presents a challenge. We do find that including the anisotropy of the Coulomb interaction in DFT+U with larger values of $U$ gives results closer to those from the HSE hybrid functional.

This paper is organized as follows. In the following section we summarize the expected Jahn-Teller effect in the hollandite group oxides, introduce a local order parameter to measure the corresponding distortion of the tetragonal unit cell and give a detailed description of all the DFT methods employed. In Sect. \textbf{III} we discuss our results comparing the various different DFT approaches for energetics, crystal structure, charge localization and charge ordering. We then discuss these results in Sect. \textbf{IV} in light of the literature for related compounds and compare two reasonable choices of method to study more general phase formation in Li$_x$Ag$_y$Mn$_8$O$_{16-z}$:$(H_2O)_w$. In the last section we summarize our work.

\section*{II. THEORY AND METHOD}

\subsection*{A. Structure and Jahn-Teller Effect}

The basic unit cell depicted in Fig. 1 which represents the formula unit A$_2$Mn$_8$O$_{16}$, will be the building block for our studies. The edge-sharing octahedra form the walls of the large 2 × 2 tunnels and point-sharing oxygen ions are the corners of the smaller 1 × 1 tunnels. Typically the 1 × 1 tunnels are too narrow to host large concentrations of guest ions,\textsuperscript{20} so that the inserted Li ions, or the structurally supporting ones like Ag or K, are located inside the larger 2 × 2 tunnels. Previous DFT studies on Li$_2$Mn$_8$O$_{16,2}$\textsuperscript{17,19} have shown that Li prefers to occupy the off-center positions at the concentration $x = 2$. Ag and K prefer a more symmetrical coordination, but generally with a different position along the tunnel direction. These are illustrated in Fig. 1.\textsuperscript{20,55,71}

In the absence of any cations in the tunnels, the structure of hollandite is tetragonal and has $I4/m$ symmetry (space group 87). In this case the cell contains exclusively Mn$^{4+}$ ions with an almost uniform octahedral crystal field. This field splits the degenerate atomic 3d states of Mn into the three-fold degenerate, occupied $t_2g$ states $d_{xy}, d_{xz}$ and $d_{yz}$ and the double degenerate, unoccupied $e_g$ states $d_{z^2}$ and $d_{x^2-y^2}$. Examples of the $e_g$ states in the local framework of the octahedra are illustrated in Fig. 2. In the ferromagnetic (FM) solution the $t_{2g}$ electrons of all Mn ions are aligned in parallel. In the case of anti-ferromagnetic (AF) order, there are a number of different ways for the spins to be organized, both within the basic unit cell shown in Fig. 1 as well as simple generalizations, such as a doubling along the indicated $c$ axis. Several alternatives have been enumerated in recent literature.\textsuperscript{22} In agreement with Ref. 54 we find that the ordering pattern designated C2-AFM, where the spin sign alternates between corner-sharing Mn ions, as illustrated in Fig. 1 has lower energy than the competing patterns. Results designated as AF in this paper refer to this order.

Independent of magnetic ordering, the atoms A in the tunnel typically ionize and transfer electrons to the Mn$^{4+}$ cations in the oxide backbone. One possibility is that the extra electrons partially occupy a sufficiently dispersed band, or multiple bands, such that they are delocalized. Another possibility is that the excess electrons are localized, in the extreme limit each one resulting in a local Mn$^{3+}$ cation. In the present case for monovalent atoms A in the tunnel, this can formally be written $(A^{+})_2(Mn^{3+})_2(Mn^{4+})_8(O^{2-})_{16}$. In a solution with local Mn$^{3+}$ cations, the ordering must also be determined. Several possible examples of structures within the basic hollandite unit cell and small supercells $(\sqrt{2} \times \sqrt{2} \times 1$ and $1 \times 1 \times 2)$ are discussed in detail in \textbf{III C}.

The 3$d^4$ Mn$^{3+}$ cation should be high spin according to...
Hund’s rule so the extra electron will occupy one of the nearly degenerate $e_g$ states. Following the usual Jahn-Teller argument,
the local octahedron can distort to a lower symmetry solution, split the degenerate $e_g$ states through a distorted crystal field and gain energy by singly occupying the lower energy state. The net energy gain is limited by the local elastic energy cost of the distortion. Two orthogonal distortions, $Q_2$ and $Q_3$, were introduced by Van Vleck to span distinct symmetry reduction pathways from the ideal octahedron that couple to the $e_g$ states.

Generalized to the slightly distorted initial octahedra found in $\alpha$-MnO$_2$ and written in terms of the Mn-O bond lengths defined in Fig. 2 these are

\begin{align}
Q_2 &= \frac{1}{\sqrt{2}} (l_1 + l_4 - l_2 - l_5) \\
Q_3 &= \frac{1}{\sqrt{6}} (2l_3 + 2l_6 - l_1 - l_4 - l_2 - l_5).
\end{align}

Here $l_{3,6}$ are the MnO bond lengths in the locally defined $z$ direction and $l_{1,4}, (l_{2,5})$ the corresponding bond lengths in the $x(y)$ direction.

The $Q_2$ order parameter measures the local orthorhombic distortion, and $Q_3$ the tetragonal distortion, of an octahedral crystal field.

Each solution with local Mn$^{3+}$ cations can be expected to have some degree of local Jahn-Teller distortion that can be quantified through $Q_2$ and $Q_3$. Furthermore, these distortions will couple to the crystal structure cooperatively in a particular way determined by how they are ordered, resulting in a final, possibly distorted unit cell. Finding the ground state at zero temperature requires sampling different patterns of charge order to determine the one with lowest energy. Target charge order is probed by locally imposing the distortion and relaxing the ionic positions to determine if the order is self consistently sustained.

All DFT calculations have been performed with the Vienna Ab initio Simulation Package (VASP). Specific exchange-correlation functionals utilized include PBE, Opt-B88 and SCAN. For the PBE and Opt-B88 calculations, the GGA$+U$ approach was used, specifically applied to the 3$d$ electrons of Mn. Survey calculations were done using the spherical approximation. In this case, one effective Coulomb interaction $U_{\text{eff}} = U - J$ is required, here treated as a parameter for analysis. For comparison, selected PBE$+U$ calculations were performed retaining the anisotropy of the Coulomb interaction, specifically with $U = 6$ eV and $J = 1$ eV and to be compared to $U_{\text{eff}} = 5.0$ eV. These values are similar to those chosen in a recent study of magnetic order in $\beta$-MnO$_2$ while the choice of $U$ is somewhat larger than those identified in a recent study of five manganese oxides. For the balance of this paper, we will use compact notation to distinguish the GGA$+U$ calculations according to the choice of GGA, the parameter values and the use of the spherical approximation, e.g., Opt-B88$+U_{\text{eff}}=1.6$ eV in the spherical approximation and PBE$+(U=6, J=1$ eV) considering full anisotropy.

For the projector augmented wave (PAW) basis set, a cutoff of 520 eV was employed in combination with the semi-core potentials Mn:3$s^23p^64s^23d^5$, O:2$s^22p^4$, Ag:4$s^24p^65s^14d^{10}$, K:3$s^23p^64s^4$ and Li:1s$^22s^1$ resulting in unit cells with roughly 250 electrons. The Brillouin zone of the unit cell was sampled with a Γ-centered, 2 $\times$ 2 $\times$ 6 grid of $k$ points. The Methfessel-Paxton smearing method of order 1 was used with a smearing parameter of $\sigma=0.1$ eV. Self-consistency was converged to a total energy convergence criterion of 10$^{-6}$ eV. Using these settings we have relaxed all structures with the conjugate gradient algorithm until the residual force (acting on each individual ion) was less than 20 meV/Å. As needed, cell shape was also relaxed, with accuracy in this procedure assured by the large basis set and by checking for changes upon restart following initial convergence of the cell parameters. After performing convergence tests with respect to $k$ point sampling and energy cutoff, we estimate that PBE, Opt-B88 and SCAN energy differences reported below are given with a precision of 5 meV/cell or smaller.

For the hybrid functional calculations, we have chosen HSE in the formulation of 2006 with 25% Hartree-Fock exchange and a range separation parameter of $\omega = 0.2$ Å$^{-1}$. Here the relaxations have been done employing a PAW basis set cutoff of 450 eV in combination with a 1 $\times$ 1 $\times$ 4 $k$ point sampling and a residual force threshold of 20 meV/Å. These cutoffs were slightly reduced, for computational efficiency, relative to the settings used otherwise. Note that semi-core electrons are retained, following experience that shows the importance of core-valence exchange, particularly for $d$ electron cases. For the relaxed unit cells, total energies were recalculated with the same settings as those used for the PBE, Opt-
B88 and SCAN calculations. Only small deviations in the forces and in the binding energies of A in $A_2Mn_8O_{16}$ were observed. We estimate that HSE-based relative energies reported below are given with a precision of 20 meV/atom or better.

The localized electronic structure was studied using a basis set of maximally localized Wannier functions. The projection of the PAW basis set on the valence states $Mn\,d$, $O\,p$, $K\,s$ and $Li\,s$ orbitals was done using the wannier90 tool with the VASP2WANNIER90 interface. For Ag the inclusion of the $d$ states in addition was necessary to reproduce the band structures with the Wannier interpolation method. The projected density of states (PDOS) has been interpolated using a $k$ point grid of $6 \times 6 \times 24$ points.

III. RESULTS

A. Binding energies

We first consider the binding energies of A in $A_2Mn_8O_{16}$. For this purpose we have calculated

$$E(A) = \frac{E_A - E_0 - 2\mu_A}{2},$$

(3)

where $E_A$ is the energy of the unit cell of $A_2Mn_8O_{16}$, $E_0$ the energy of pristine hollandite and $\mu_A$ the corresponding chemical potential of A. Here, both $Mn_8O_{16}$ and $A_2Mn_8O_{16}$ are computed for the AF ordering shown in Fig. 1 and correspondingly with the lowest energy charge ordering found. For the A chemical potential, we have chosen the energy per atom of bcc-Li, bcc-K and fcc-Ag metal, respectively. Thus Eq. (3) measures the difference in total electronic energy (i.e. $T = 0$ K and no correction for zero-point motion) of the reaction $Mn_8O_{16} + 2\times A \rightarrow A_2Mn_8O_{16}$. That is, negative (positive) energies $E(A)$ indicate a spontaneous formation (decomposition) of $A_2Mn_8O_{16}$ at zero temperature. A comparison of $E(A)$ for a set of different DFT functionals and values of $U_{\text{eff}}$ is given in Fig. 3.

It can be seen that for all investigated cations, the Opt-B88+$U_{\text{eff}}$ (points) and HSE functional (dashed lines) yield negative binding energies implying a spontaneous formation of $A_2Mn_8O_{16}$ at zero temperature. In contrast, SCAN (squares) as well as PBE+$U_{\text{eff}}$ (circles) for $U_{\text{eff}} < 2.5$ eV predict a positive formation energy for $A_2Mn_8O_{16}$ implying that additional energetic cost would be necessary to form silver hollandite at zero temperature. This suggests that there is insufficient description of vdW interactions in PBE and SCAN, a part of the correlation energy taken into account explicitly by Opt-B88 and empirically at short range with HSE. Evidently this is important for the chemical bonding of a noble metal like Ag in the tunnel. The vdW term in the exchange-correlation functional of Opt-B88+$U_{\text{eff}}$ shifts the binding energies of all three cations almost uniformly by $\approx 0.5$ eV downwards compared to PBE+$U_{\text{eff}}$, as seen by comparing the closed and open circles in Fig. 3. The same figure also reveals that in both PBE+$U_{\text{eff}}$ and Opt-B88+$U_{\text{eff}}$, the binding energy decreases monotonically for all investigated cations for $U_{\text{eff}} > 2$ eV and approaches the HSE result at specific values. Also, inclusion of anisotropy with PBE+$U_{\text{eff}}$ ($U=6, J=1$ eV) slightly improves the results compared to HSE.

In applications, such as comparison of energies for different compositions to build a phase diagram, one value of $U_{\text{eff}}$ must be selected for consistency. Here, Fig. 3 provides a calibration, using the HSE results as a reference. The best match with HSE binding energies for all three cations is obtained for the Opt-B88+$U_{\text{eff}}$ functional with moderate values of $U_{\text{eff}} \lesssim 2$ eV and for the anisotropic PBE+$U_{\text{eff}}$ ($U=6, J=1$ eV) method. A somewhat less accurate match is found with PBE+$U_{\text{eff}}$ for values of $U_{\text{eff}}$ in the range $3$ to $5$ eV.

Over all, K has the strongest binding energy in the hollandite tunnel followed by Li and the weakly bound Ag. This trend does track the ionization energy, a first measure for the energy gain possible upon formation of the ion in the tunnel. For instance, K has one of the lowest ionization energies in nature $82$ (4.34 eV). This is followed by Li with 5.39 eV $\approx 2$ which results overall in weaker ionic bonding for Li compared to K. In contrast, Ag has an ionization energy of 7.58 eV $\approx 2$ and, correspondingly shows relatively weak ionic binding. We have examined the binding of other ions in the hollandite tunnel.
The binding energy for Na (-3.10 eV) is found to be close to that for Li, in accordance with the trend above (the ionization energy of Na is 5.14 eV\(^{22}\)). However, this entire trend is violated by second row earth alkali elements like Mg, which has an ionization energy of 7.65 eV, but a binding energy (-2.90 eV) similar to Li. A simple model for the binding energy must include other factors beyond ionization energy.

**B. Structure and Monoclinic Distortion**

Next, we focus on the structure of pristine and doped hollandite compounds and compare how selected, different DFT methods affect the corresponding lattice constants. The results are reported in Tab. I. They all correspond to the AF order shown in Fig. I except for the case of PBE+(\(U=6, J=1\) eV) applied to Li\(_2\)Mn\(_8\)O\(_{16}\) where FM order was lowest in energy.

Starting with the results for the parent Mn\(_8\)O\(_{16}\) structure, Opt-B88, SCAN and Opt-B88+\(U_{\text{eff}}=1.6\) eV, yield a stable, slightly distorted structure as lowest in energy. A tetragonal structure is also stable (\(a = 9.68, 9.61\) and 9.70 Å respectively), but slightly higher in energy (1, 13 and 2 meV per cell respectively). On the other hand, using PBE+(\(U=6, J=1\) eV) and HSE a tetragonal structure has the lowest energy. Correspondingly, in those cases, a distorted structure can also be stabilized, but it has a slightly higher energy per cell (1 and 6 meV respectively). These are very small energy differences, comparable to our estimated precision, but the relative ordering is robust upon increasing the k-point sample. These results are indicative of quite soft degrees of freedom in the potential landscape that describes the hollandite materials, particularly deformations related to bond angles centered on the corner shared oxygen atoms.\(^{24}\)

Upon introducing cations into the tunnel, Opt-B88 and SCAN yield almost tetragonal structures for Ag and K, but predict a significant monoclinic distortion for Li. The latter distortion can be explained by the ionic radius of Li, located in the tunnel center. Introduction of Ag and K reduce the Mn\(^{4+}\) ions uniformly to Mn\(^{3.75+}\). In the absence of any Hubbard term, these compounds behave like simple band metals, with several empty bands of the parent Mn\(_8\)O\(_{16}\) partially occupied.

As one increases the interaction \(U_{\text{eff}}\) in the GGA+\(U_{\text{eff}}\) methods, the details of the distorted structure induced by Li insertion change, but no qualitative changes in the monoclinic cell emerge. For Ag and K insertion, there is a transition to a monoclinic cell at relatively small \(U_{\text{eff}}\). For Opt-B88+\(U_{\text{eff}}=1.6\) eV reported in Tab. I the K hollandite is distorted, whereas Ag hollandite remains tetragonal with slightly larger lattice constants. Increasing the interaction further triggers a monoclinic distortion for Ag\(_2\)Mn\(_8\)O\(_{16}\) as well. For the large \(U_{\text{eff}}\) limit, with PBE+\(U_{\text{eff}}\), the Li, K and Ag hollandites are predicted to have very similar monoclinic cells. For comparison, the hybrid HSE functional also predicts monoclinic cells for all three ions, with \(b - a\) values ranging from 0.62 Å for Li to 0.77 Å for K.

In contrast, the measured crystal structures indicate tetragonal cells in all cases. None of the measured crystals had ideal cation concentration, but variations in the lattice parameters among different cases is relatively small. For example, X-ray measurements for Ag\(_{1.22}\) and Ag\(_{1.66}\) yielded \(a = 9.770(2)\) and 9.738(2) Å respectively.\(^{24}\) There was variation in the stoichiometry of the compounds resulting from the ion exchange exper-

| TABLE I. For selected hollandite compounds, lattice constants and angle \(\gamma\) between primitive lattice vectors a and b, as calculated for several functionals in DFT. |
| --- | --- | --- | --- |
| | a [Å] | b [Å] | c [Å] | \(\gamma\) [deg] |
| Opt-B88 | | | | |
| Mn\(_8\)O\(_{16}\) | 9.58 | 9.78 | 2.85 | 89.43 |
| Ag\(_2\)Mn\(_8\)O\(_{16}\) | 9.66 | 9.70 | 2.84 | 90.00 |
| K\(_2\)Mn\(_8\)O\(_{16}\) | 9.70 | 9.70 | 2.86 | 90.00 |
| Li\(_2\)Mn\(_8\)O\(_{16}\) | 9.25 | 10.11 | 2.82 | 91.58 |
| Opt-B88+\(U_{\text{eff}}=1.6\) | | | | |
| Mn\(_8\)O\(_{16}\) | 9.57 | 9.79 | 2.85 | 90.47 |
| Ag\(_2\)Mn\(_8\)O\(_{16}\) | 9.70 | 9.70 | 2.85 | 90.00 |
| K\(_2\)Mn\(_8\)O\(_{16}\) | 9.49 | 10.18 | 2.88 | 91.57 |
| Li\(_2\)Mn\(_8\)O\(_{16}\) | 9.39 | 10.18 | 2.84 | 92.10 |
| PBE+(\(U=6, J=1\) eV) | | | | |
| Mn\(_8\)O\(_{16}\) | 9.80 | 9.80 | 2.90 | 90.00 |
| Ag\(_2\)Mn\(_8\)O\(_{16}\) | 9.51 | 10.35 | 2.90 | 91.77 |
| K\(_2\)Mn\(_8\)O\(_{16}\) | 9.64 | 10.34 | 2.91 | 91.03 |
| Li\(_2\)Mn\(_8\)O\(_{16}\) | 9.61 | 10.35 | 2.89 | 91.73 |
| SCAN | | | | |
| Mn\(_8\)O\(_{16}\) | 9.54 | 9.72 | 2.83 | 90.41 |
| Ag\(_2\)Mn\(_8\)O\(_{16}\) | 9.59 | 9.59 | 2.82 | 90.00 |
| K\(_2\)Mn\(_8\)O\(_{16}\) | 9.65 | 9.65 | 2.84 | 90.00 |
| Li\(_2\)Mn\(_8\)O\(_{16}\) | 9.30 | 10.11 | 2.81 | 91.88 |
| HSE | | | | |
| Mn\(_8\)O\(_{16}\) | 9.59 | 9.59 | 2.83 | 90.00 |
| Ag\(_2\)Mn\(_8\)O\(_{16}\) | 9.41 | 10.11 | 2.84 | 91.53 |
| K\(_2\)Mn\(_8\)O\(_{16}\) | 9.39 | 10.16 | 2.85 | 91.49 |
| Li\(_2\)Mn\(_8\)O\(_{16}\) | 9.40 | 10.02 | 2.81 | 92.19 |
| Experiment | | | | |
| Mn\(_8\)O\(_{16}\) | 9.777(2) | 2.8548(5) | 90.00 |
| Ag\(_{1.22}\)Mn\(_8\)O\(_{16}\) | 9.725(7) | 2.885(2) | 90.00 |
| K\(_{1.22}\)Mn\(_8\)O\(_{16}\) | 9.866(3) | 2.872(1) | 90.00 |
| Li\(_{1.22}\)K\(_{0.8}\)Mn\(_8\)O\(_{16}\) | 9.81-9.89 | 2.855(2) | 90.00 |

\(^{a}\text{Ref. 36,}^ {b}\text{Ref. 52,}^ {c}\text{Ref. 50,}^ {d}\text{Ref. 51}\)
ments reported in Ref. 51. The results were also affected by possible reactions with residual water in the tunnels. The range of lattice parameters reported for Li hollandite is indicated in Tab. I. Trends with cation are on a small scale, smaller than differences among calculations with different exchange-correlation functionals. However, the trend of reduced in-plane lattice parameter for the Ag case, and the increase for K, are reproduced by both Opt-B88 and SCAN functionals. These are room temperature experiments. Further discussion appears in Sect. IV below.

These results for K and Ag suggest that the structural deformation observed in the calculations is electronically driven. This may be significant for the Li case as well. The donated cation s electrons are becoming more localized with increasing $U_{\text{eff}}$ value, forming two distinct Mn$^{3+}$ ions per unit cell, so that at some point the system gains enough energy from orbital reordering to distort the tetragonal symmetry. This cooperative Jahn-Teller effect depends on the cation size as well as the coordination with the environment and therefore happens at different $U_{\text{eff}}$ values for the three cations.

Quantitatively, we find that the local Jahn-Teller effect in the $\mathrm{A}_2\mathrm{Mn}_8\mathrm{O}_{16}$ hollandite materials can be specifically tracked by means of the local order parameter $Q_3$, defined in Eq. (1). In those stable structures in which local Mn$^{3+}$ ions emerge, correspondingly the order parameter $Q_3$ becomes non-zero. For the case at hand, full localization of the two extra donated electrons per unit cell correspond to two Mn$^{3+}$ centered octahedra and six Mn$^{4+}$ centered octahedra. In each case, the octahedra naturally pair, with approximately similar distortions. To highlight the emergence of the Mn$^{3+}$ centers, we plot two averaged $Q_3$ parameters, one for the two Mn$^{3+}$ ions and one for the remaining six Mn$^{4+}$ ions. The results for all three cation doped hollandite unit cells as a function of $U_{\text{eff}}$ are given in Fig. 4. The insets illustrate the charge order that emerges in each case. The order parameter of the averaged six Mn$^{4+}$ ions can be further decomposed into three pairs, as shown in Fig. 4. While these three pairs of octahedra differ in minor details of structure, the magnitude of $Q_3$ remains near zero.

Looking at Fig. 4 one sees that, for all three compounds, $Q_3$ increases significantly between $U_{\text{eff}} = 1$ and 2 eV and indicates the formation of Mn$^{3+}$. The charge order has a different character in the case of Li, where as previously noted, the small ionic radius favors an off-center ion location. For large $U_{\text{eff}}$, this results in a local complex of two Li ions and two Mn$^{3+}$ centered octahedra in a single tunnel wall. This is distinct from the local structure found for $U_{\text{eff}} = 0$ and described above. In contrast, for the K and Ag cases, where the cation remains near the tunnel center, the two Mn$^{3+}$ centered octahedra are maximally separated. For reference, results based on HSE are also plotted. The values track the large $U_{\text{eff}}$ plateaux. The anisotropic PBE+$\left(U=6, J=1\right)$ eV) predicts a larger distortion, but also in overall good agreement with the HSE results. These results all correspond to AF order. For comparison, in the Ag case, with FM order and $U_{\text{eff}} = 5$ eV, $Q_3$ is a bit smaller, 0.38 Å, indicating some interplay with the magnetic order.

To understand this effect from the electronic structure perspective we have used Wannier orbitals and calculated the projected density of states (PDOS) for all Mn ions individually. To probe the local states in each octahedra.

FIG. 4. (color online) Averaged order parameter $Q_3$ as a function of $U_{\text{eff}}$ with OptB88+$U_{\text{eff}}$, calculated for Mn$^{3+}$ centered octahedra (points) and Mn$^{4+}$ centered octahedra (circles) for each of Ag, K and Li in $\mathrm{A}_2\mathrm{Mn}_8\mathrm{O}_{16}$. Results for HSE and PBE+$\left(U=6\right.)$ eV, $J=1$ eV) are also shown. Insets illustrate the charge order and provide a key to the octahedra sampled in each case.

FIG. 5. (color online) Same as in Fig. 4 but focusing on the order parameters $Q_3$ averaged in pairs for the six Mn$^{4+}$ octahedra, following the illustrative insets.
Drone in the most natural way, we have aligned locally the \((x,y,z)\) axes as depicted in Fig. (4). Physically, it is sufficient to restrict consideration to the local \(e_g\) derived empty states that receive the electrons from the A cations. In Fig. (6) we compare the \(e_g\) derived PDOS for the case of pure MnO\(_{16}\) with that of Ag\(_2\)MnO\(_{16}\) for PBE with \(U_{\text{eff}} = 0\) and with \(U_{\text{eff}} = 5\) eV. For each case, the results for an AF ordered state (Fig. (d)) are shown with the majority spin PDOS plotted upwards and the minority spin PDOS plotted downwards. The gray background shows the sum of all \(e_g\) states. The lines represent the \(d_{z^2}\) (blue) and \(d_{x^2-y^2}\) states (green) from specific octahedra with majority spin up (see insets).

In the case of pristine hollandite, the results shown are for a tetragonal cell and all eight octahedra are equivalent, except for the majority spin orientation. The increase in the interaction strength increases the band gap, as expected. Correspondingly, the hybridization that leads to a small amount of \(d_{z^2}\) and \(d_{x^2-y^2}\) character in the occupied portion of the PDOS, is reduced by the large \(U_{\text{eff}}\). But effectively, all Mn ions in MnO\(_{16}\) have a \(t_{2g}\) shell locally filled with majority spin electrons, but empty for minority spin electrons and no occupancy of the nominal \(e_g\) bands. The Jahn-Teller effect is absent.

For Ag\(_2\)MnO\(_{16}\), the additional \(s\) electrons from Ag partially occupy available \(e_g\) majority spin states. However, for \(U_{\text{eff}} = 0\), the system behaves like a band metal with partial occupation of several previously empty bands and the eight octahedra remain equivalent. The PDOS in Fig. (b) shows that the Fermi energy has moved into the bottom portion of the previously empty \(e_g\) derived bands and there is some detailed rearrangement of states. However, Fig. (d) clearly shows that, for a high value of \(U_{\text{eff}} = 5\) eV, the majority \(d_{z^2}\) state is fully occupied in two of the identifiable local Mn\(^{3+}\) octahedra. Correspondingly, the local \(d_{x^2-y^2}\) state is empty and a band gap is maintained. This large energy splitting goes with the large, local value of the \(Q_3\) order parameter illustrated in Fig. (4) and drives the energy gain from the Jahn-Teller effect. For the other six Mn\(^{3+}\) octahedra, the \(e_g\) states remain empty and the local value of the \(Q_3\) order parameter is close to zero. Overall, the consequence is that the unit cell becomes monoclinic for high \(U_{\text{eff}}\) due to the elongation of the Mn\(^{3+}\)O\(_6\) octahedra in the \(b\) direction.

In order to isolate the electronic effect from the ionic effect, we have also done self consistent calculations in which electrons have been added to pristine MnO\(_{16}\), compensated by a uniform positive background. This allows us to assess the electronic part of the Jahn-Teller effect in a continuous way as a function of added electrons to the unit cell, in essence studying (MnO\(_{16}\))\(^{-x}\). The results for three examples, using Opt-B88 and different \(U_{\text{eff}}\) values, are shown in Fig. (7).

For a low charge of \(x = 0.5\) and the restriction of a single unit cell a charge localized solution starts to emerge for \(U_{\text{eff}} \approx 3\) eV, with a local order parameter \(Q_3\) that grows to a modest value \((Q_3 = 0.17\) \(\AA)\) for larger \(U_{\text{eff}}\) (not shown). If a \(1\times1\times2\) supercell is used the order parameter \(Q_3\) increases roughly by a factor of two (see Fig. 7). In addition, the magnetic moment of the corresponding Mn ion increases from \(m = 3.4\) \(\mu_B\) to \(m = 3.7\) \(\mu_B\) indicating formation of a single Mn\(^{3+}\) ion for the supercell. For \(x = 1.0\) a single Mn\(^{3+}\) ion with a magnetic moment of \(m = 3.7\mu_B\) starts forming from a small value of \(U_{\text{eff}}\).

FIG. 6. Projected density of Mn \(e_g\) states \((d_{z^2}, d_{x^2-y^2})\) for MnO\(_{16}\) (Fig. (a),(c)) and Ag\(_2\)MnO\(_{16}\) (Fig. (b),(d)). Grey background represents the \(e_g\) states of all octahedra, while the colored lines in all figures represent the labeled projections of the majority spin up octahedra highlighted in the insets.

FIG. 7. Distortion index \(Q_3\) as function of effective parameter \(U_{\text{eff}}\) for charged unit cells (MnO\(_{16}\))\(^{-x}\), as defined in Eq. (4).
Finally, for the case of \( x = 2.0 \), which corresponds to the same number of electrons donated in \((\text{Ag,K,Li})_2\text{Mn}_8\text{O}_{16}\), the localized solutions already emerge with \( U_{\text{eff}} = 0 \). Furthermore, the saturated value of the distortion for larger \( U_{\text{eff}} \) exceeds that found for the real materials in Fig. 4. Together, these results suggest that the pure electronic driving force for charge localization and Jahn-Teller distortion is quite strong. However, along with these local effects, the overall lattice is expanded by roughly 4%. Evidently, the presence of the actual cations in the tunnels (instead of a simple neutralizing background) opposes this expansion and limits the Jahn-Teller distortion.

C. Charge Ordering

Having demonstrated the emergence of solutions with localized \( \text{Mn}^{3+} \) centered octahedra due to the Jahn-Teller effect, we further examine the impact of how those centers are ordered and the interplay with the magnetic order. We focus on \( \text{Ag}_2\text{Mn}_8\text{O}_{16} \). While a systematic study that extends beyond the basic unit cell is desirable, the computationally complexity grows rapidly. In particular, the number of distinct \( (\text{Mn}^{3+})_2(\text{Mn}^{4+})_6 \) configurations within even modest sized super cells, assessed using site occupancy disorder tool of Grau-Crespo,\(^{84} \) become quite large.\(^{85} \) Therefore, we have approached this problem empirically, identifying the smallest building blocks of \( \text{Mn}^{3+} \) patterns and building more complex structures based on them.

To this end, we have assumed that only \( d_{z^2} \) orbitals become occupied during charge localization, \( i.e. \), elongation of \( \text{Mn}^{4+}\text{O}_6 \) octahedra occurs only in the \((a,b)\) plane (see the Wannier projection in Fig. 2). This gives three different charge orderings for the unit cell: CO1, CO2, and CO3, visualized in Fig. 8.\(^{83} \) These orderings are the building blocks for three more extended charge orderings in supercells: CO4 \((\sqrt{2} \times \sqrt{2} \times 1)\) and CO5 and CO6 \((1 \times 1 \times 2)\). With the exception of CO3, this set coincides with the spin-charge ordering structures studied by Fukuzawa et al.\(^{82} \) by means of unrestricted Hartree-Fock calculations for \( \text{K}_2\text{Mn}_8\text{O}_{16} \). For our initial survey, we have used PBE+\( U_{\text{eff}} = 5 \text{ eV} \), which combines rough agreement with the HSE for binding energy of \( \text{Ag}_2\text{Mn}_8\text{O}_{16} \) and clear formation of local \( \text{Mn}^{3+} \) centers. The results are given in Tab. 11.

Several competing phases with different charge and magnetic order emerge. The lowest pair differ by only 0.04 eV per cell, but exhibit distinctive structures. CO1 and CO2 both have ferromagnetic (FM) order, but a different organization of the charge. CO1 has a strong monoclinic distortion, whereas CO2 is almost perfectly orthorhombic with an angle of \( \gamma = 93.58^\circ \) between \( a \) and \( b \). The next phase is the AF ordered CO1 structure, 0.06 eV above the FM CO1 structure. The charge order CO6, 0.13 eV above FM CO1, is essentially tetragonal in structure, achieved by organizing the Mn\(^{3+}\) evenly around the tunnel walls, but alternating along the \( c \) axis in a \( 1 \times 1 \times 2 \) unit cell.

The next distinct configuration, CO4 coincides formally with the charge ordering found in \( \text{K}_2\text{V}_8\text{O}_{16} \) at low temperatures.\(^{37} \) However, the Jahn-Teller effect in \( \text{Mn}^{3+} \) ions of CO4 acts in the \((a,b)\) plane (see Fig. 2 and Fig. 8) in contrast to the \( \text{V}^{3+} \) ions, where the same effect is active primarily along the \( c \) axis and causes a dimerization of \( \text{V}^{3+} \) (Peierls distortion) and a zig-zag pattern of \( \text{V}^{4+} \) along the tunnel direction.\(^{37} \) On the other hand, the distortion modes in \( \text{Ag}_2\text{Mn}_8\text{O}_{16} \) compensate each other and yield effectively a tetragonal structure for both, the ferromagnetic and antiferromagnetic ordering. Several other choices of charge and magnetic order also yield stable solutions, but with higher energy (Tab. 11).

With PBE+(\( U=6, J=1 \text{ eV} \), a similar picture emerges (Tab. 11). The lowest energy structure has a strong monoclinic distortion, but several structures close in energy exhibit smaller distortions. However, the interplay with
that found with PBE+

creased, but the nearly tetragonal structure is similar to

energy separation to the higher lying CO2 state is in-

CO1 show a similar degree of monoclinic distortion. The

HSE clearly con-

with charge order CO1 and CO2 both have AF order.

TABLE II. Energy (in eV per unit cell, relative to the lowest

energy case), magnetic ordering and lattice constants (in Å)
of Ag2Mn3+Mn4+O16 configurations of Fig. [5]obtained with

several different exchange-correlation approximations.

| conf. | \( E \) | mag. | \( a \) [Å] | \( b \) [Å] | \( c \) [Å] | \( \gamma \) [°] |
|-------|-------|------|---------|---------|---------|---------|
| PBE+ U_{eff} = 5 |
| CO1  | 0.00  | FM   | 9.55    | 10.32   | 2.93    | 91.14   |
| CO2  | 0.04  | FM   | 9.91    | 9.90    | 2.94    | 93.58   |
| CO1  | 0.06  | AF   | 9.54    | 10.35   | 2.92    | 91.49   |
| CO6  | 0.13  | FM   | 9.85    | 9.85    | 5.92    | 90.02   |
| CO4  | 0.16  | FM   | 13.93   | 13.97   | 2.96    | 90.21   |
| CO2  | 0.26  | AF   | 9.80    | 10.06   | 2.92    | 93.55   |
| CO4  | 0.45  | AF   | 14.06   | 13.99   | 2.92    | 90.25   |
| CO6  | 0.45  | AF   | 9.88    | 9.88    | 5.85    | 90.01   |
| CO3  | 0.55  | AF   | 9.91    | 9.94    | 2.92    | 89.55   |
| CO5  | 0.55  | AF   | 9.88    | 9.89    | 5.85    | 90.00   |
| PBE+ (U=6, J=1 eV) |
| CO1  | 0.00  | AF   | 9.53    | 10.35   | 2.90    | 91.38   |
| CO1  | 0.12  | FM   | 9.53    | 10.36   | 2.91    | 91.30   |
| CO2  | 0.17  | AF   | 9.78    | 10.06   | 2.91    | 93.38   |
| CO2  | 0.21  | FM   | 9.79    | 10.05   | 2.92    | 93.30   |
| CO6  | 0.38  | AF   | 9.87    | 9.88    | 5.83    | 90.02   |
| CO4  | 0.38  | AF   | 13.98   | 14.06   | 2.91    | 90.25   |
| CO4  | 0.39  | FM   | 13.98   | 14.05   | 2.92    | 90.25   |
| CO6  | 0.39  | FM   | 9.87    | 9.88    | 5.86    | 90.03   |
| CO3  | 0.48  | AF   | 9.90    | 9.94    | 2.91    | 90.46   |
| CO5  | 0.49  | AF   | 9.87    | 9.87    | 5.86    | 90.00   |
| CO5  | 0.50  | AF   | 9.87    | 9.88    | 5.83    | 90.00   |
| HSE  |
| CO1  | 0.00  | AF   | 9.41    | 10.10   | 2.84    | 91.64   |
| CO2  | 0.24  | AF   | 9.67    | 9.74    | 2.84    | 93.25   |
| CO1  | 0.26  | FM   | 9.39    | 10.10   | 2.84    | 91.63   |
| CO2  | 0.38  | FM   | 9.70    | 9.74    | 2.84    | 93.45   |
| Opt-B88 |
| Uniform | 0.00  | AF   | 9.66    | 9.66    | 2.84    | 90.00   |
| Uniform | 0.67  | AF   | 9.66    | 9.66    | 2.86    | 90.00   |
| Opt-B88+U_{eff}=1.6 |
| Uniform | 0.00  | AF   | 9.71    | 9.72    | 2.84    | 90.19   |
| CO1   | 0.02  | AF   | 9.40    | 10.18   | 2.87    | 91.27   |
| Uniform | 0.27  | FM   | 9.73    | 9.73    | 2.92    | 90.00   |

that from PBE+(\( U=6, J=1 \) eV) and the energy differ-
ence between the higher FM phase and the AF phase is
about the same.

For comparison, the Opt-B88 results show a uniform
phase, as expected for a band metal with no charge local-
ization, a tetragonal structure and a much larger stabi-
lization of the AF order over the FM order. Interestingly,
inclusion of a small \( U_{eff} \) does allow for a charge ordered
structure which is only slightly higher in energy than the
uniform structure and reduces the energy difference sep-
arrating the FM ordered phase.

For reference, the predicted magnetic order for the par-
ent \( \alpha-MnO_2 \) using Opt-B88 and PBE is AF, with an
energy difference per unit cell of 0.70 and 0.58 eV to
the FM phase. Explicit consideration of local Coulomb
interactions through GGA+\( U_{eff} \) systematically reduces
this energy splitting, to 0.01 and 0.06 eV respectively for
\( U_{eff}=5 \) eV. The same trend was found in Ref. 54, but we
do not find stabilization of the FM phase in the range
of \( U_{eff} \) we considered. With PBE+(\( U=6, J=1 \) eV), the
AF-FM energy splitting is increased (0.14 eV). Finally,
our results for HSE also indicate AF order to be favored,
with a similar splitting (0.24 eV). While internally con-
sistent, this final result for \( \alpha-MnO_2 \) is opposite to that
found in Ref. 54. We have included the semicon-
electrons and the structure for each phase was relaxed with
HSE.

Finally, we note that our HSE calculations also yield
an AF phase for the K and Li cases with an energy split-
ing of 0.20 and 0.14 eV respectively to the FM phase.
Interestingly, PBE+(\( U=6, J=1 \) eV) results show an AF
phase for K (0.12 eV), but a FM phase for Li (0.06 eV).

IV. DISCUSSION

Our results based on PBE+\( U_{eff} \) at high \( U_{eff} \) values show
a charge ordered ground state, but several close-
by competing charge and magnetic ordered phases. The
HSE results confirm the charge ordering in the ground
state. However, also considering the PBE+(\( U=6, J=1 \)
eV) results, the details of the interplay between charge
order and magnetic coupling are sensitive to the method
chosen. More generally, these results, particularly the
low energy scale, point to the idea that at higher tem-
perature, disordered phases may be more relevant to the
measured room temperature results. For context, we
briefly review what is known about exemplary, related
compounds.

Near stoichiometric \( K_2V_8O_{16} \) was obtained through
high pressure synthesis. While exhibiting the ideal,
tetragonal hollandite structure at room temperature, a
first order transition to a monoclinic phase was observed
at 170 K, with a corresponding change from conducting
to insulating behavior. Subsequent X-ray and neutron
diffraction studies clearly showed that the monoclinic low
temperature phase consisted of charge-ordered, localized
\( V^{3+} \) in a \( \sqrt{2} \times \sqrt{2} \times 2 \) supercell. Detailed calculations

the magnetic order is different. The lowest energy phase
with charge order CO1 and CO2 both have AF order.
Limited exploration with the hybrid HSE clearly con-
irms the charged ordered ground state. The results for
CO1 show a similar degree of monoclinic distortion. The
energy separation to the higher lying CO2 state is in-
creased, but the nearly tetragonal structure is similar to
that found with PBE+\( U_{eff}=5 \) eV and shows a similar
value of the \( Q_3 \) order parameter (see Fig. [4]). Further-
more, the magnetic order found with HSE agrees with

V. EXPERIMENTAL

TABLE III. Energy (in eV per unit cell, relative to the lowest
based on GGA+$U_{\text{eff}}=3$ eV provided a corresponding picture of the electronic structure. While local Jahn-Teller effects manifested for both $V^{3+}$ and $V^{4+}$ centered octahedra, strong, local antiferromagnetic coupling between dimers along the chains in the tunnel direction drove the opening of the energy gap and the metal to insulator transition.\textsuperscript{39} The reduced, monoclinic symmetry at low temperature was secondary. This agreed with the observation that $\text{Rb}_2\text{VsO}_{16}$ shows a metal-insulator transition, but remains tetragonal\textsuperscript{40}.

The strong coupling of $V^{3+}$ along the $c$ direction can be related to the partially filled $t_{2g}$ shell which is Jahn-Teller active. In contrast, it can be expected that the crystal distortions in hollandite chromates are weak due to the full subshell in $\text{Cr}^{3+}$. This was also observed for $\text{K}_2\text{Cr}_8\text{O}_{16}$, where ferromagnetic order emerged at 180 K, followed by a metal to insulator transition at 95 K (with ferromagnetic order maintained).\textsuperscript{41} While no structural transition was initially observed, a subsequent study detected a subtle, $\sqrt{2} \times \sqrt{2}$ monoclinic phase at low temperature ($P1\text{1}2_1/a$) with no measurable distortion of the lattice parameters, but small dimerization in the Cr-O bond lengths (scale 0.03 Å).\textsuperscript{42} However, bond valence sum analysis suggested no charge localization, that is, no evidence for local $\text{Cr}^{3+}$ ions. GGA+$U_{\text{eff}}=2.9$ eV calculations showed that the small dimerization was sufficient to open a small gap, while remaining consistent with the ferromagnetic order. A separate GGA+$U_{\text{eff}}=3$ eV based study identified a specific soft mode associated with this Peierls driven transition.\textsuperscript{43}.

The situation for hollandite group manganese oxides under study here has been much less clear. Early studies of the parent $\alpha-$MnO$_2$ showed antiferromagnetic order below 24.5 K.\textsuperscript{24} Studies of $\text{K}_x\text{Mn}_8\text{O}_{16}$ with $x$ near 1.5 showed a change in the activation energy associated with electrical conductivity around 200 K and changes in magnetic susceptibility at low temperature with some ambiguity as to the details of the ordering.\textsuperscript{25,26} A study examining $K$ concentration dependence of the magnetic ordering ($0.8 \leq x \leq 1.4$) suggested a spin glass for lower concentration and antiferromagnetic order for higher concentration.\textsuperscript{27} Recently, a study for samples with $x$ about 1.2 and 1.4 found that even for these concentrations, a spin glass was more likely at low temperature.\textsuperscript{49} At room temperature, structural characterization of these samples points to a tetragonal structure ($I4/m$). Turning to other examples, the original report of $\text{Ag}_{8.13}\text{Mn}_8\text{O}_{16}$ synthesis and characterization reported a tetragonal structure.\textsuperscript{28} On the other hand, $\text{Ba}_{1.2}\text{Mn}_8\text{O}_{16}$ was found to have a monoclinic structure ($I2\text{1}/m$) already at room temperature and weak ferromagnetic order (or more complex competing interactions) below 40 K.\textsuperscript{29,30} While some researchers have proposed localization of $\text{Mn}^{3+}$ ions\textsuperscript{31} there has been no compelling evidence. In particular, detailed structural studies at low temperatures have not been reported.

The chemically related system $\text{LiMn}_2\text{O}_4$, which would also be expected to show a mixed valence state between $\text{Mn}^{4+}$ and $\text{Mn}^{3+}$, exhibited a first order transition near 280 K to a structure with clear evidence of charge and orbital ordering driven by the Jahn-Teller effect.\textsuperscript{27,28} Calculations based on GGA+$U_{\text{eff}}=4.5$ eV showed localized $\text{Mn}^{3+}$ ions with corresponding Jahn-Teller distortions.\textsuperscript{29} The computed lattice parameters were also in good agreement with those measured for the low temperature, ordered phase.

The role of ionic radius has been analyzed early-on based on an extensive set of hollandite group samples, with the criterion for monoclinic distortion deduced to be $r_A < r_{\text{Mn}}/0.48$. Using Shannon ionic radii ($\text{Mn}^{4+}=0.53$ Å (6-fold), $\text{Li}^{1+}=0.59$ Å (4-fold) and $\text{Ba}^{2+}=1.42$ Å)\textsuperscript{30} this criterion predicts that $\text{Ba}_2\text{Mn}_8\text{O}_{16}$ should be tetragonal and that $\text{Li}_2\text{Mn}_8\text{O}_{16}$ should be monoclinic. The data for the ion exchanged samples that show a tetragonal structure for the Li case involve extra factors due to the role of water\textsuperscript{31} and may not be a definitive test. In the Ba case, room temperature measurements already show a mildly monoclinic structure\textsuperscript{32} suggesting that other factors are likely involved.

The detailed understanding of the V and Cr based hollandites and the mixed valence $\text{LiMn}_2\text{O}_4$ was fully supported by GGA+$U_{\text{eff}}$ calculations which explicitly took into account local Coulomb interactions among the transition metal 3$d$ electrons. The range of $U_{\text{eff}}$ values used were similar to those in which our study showed formation of local $\text{Mn}^{3+}$ centers in the $\text{A}_2\text{Mn}_8\text{O}_{16}$ systems and relatively accurate binding energy for the $\text{A}$ atoms in the tunnels. Our reference HSE calculations also clearly showed charge localization. Finally, the chemically closest analog for which clear temperature dependent structural measurements were available, $\text{LiMn}_2\text{O}_4$, showed a charge ordered phase at low temperature. Taken together, this evidence indicates that charge localization in the $\text{A}_2\text{Mn}_8\text{O}_{16}$ system, to form $\text{Mn}^{3+}$ centers is likely the correct physical picture.

The proper interpretation of the room temperature structural data remains an open question. One possibility would be that we simply have not yet identified the supercell with a robust charge order and a (near) tetragonal unit cell. We can not rule this out. The other possibility is that room temperature measurements are probing a disordered phase, relative to the localized $\text{Mn}^{3+}$ centers. This picture is supported by our PBE+$U_{\text{eff}} = 5$ eV and PBE+$\langle U = 6, J = 1 \text{ eV} \rangle$ results reported in Tab. II for the $\text{Ag}_2\text{Mn}_8\text{O}_{16}$ case. A series of charge ordered phases are separated by small energy differences and encompass phases with near tetragonal structure. While the Li case appears to be more complicated, with the fundamental unit emerging as two Li ions and two edge-sharing $\text{Mn}^{3+}$ octahedra, the same basic picture could hold.

The magnetic order and the details of the energy differences between the competing phases has emerged as a more subtle problem. As already discussed in Ref. 34, the competing effects that control the magnetic coupling between neighboring Mn ions are quite subtle for the hollandite structure, particularly the coupling across the
corner shared sites. The broadly applicable Goodenough-Kanamori-Anderson (GKA) rules clearly distinguish over two scenarios: Mn-O-Mn links with near 90° and near 180° bond angles on the connecting O center, favoring FM and AF coupling respectively.\(^1\) In hollandite, the Mn-O-Mn bond between two tunnel walls is roughly 130°, indicating that the balance of interactions should be expected to be quite delicate. The differences we find likely reflect tipping this balance one way or the other. The complexity of the observed magnetic signatures in experiment, described above, similarly indicate relatively weak magnetic coupling, possibly sensitive to other factors in the material composition or structure. The hollandites represent challenging systems for DFT-based methods. While our results based on HSE and \(\text{PBE}+\left(U = 6, J = 1\right)\) eV are rather consistent, the energy scale of the differences can be significant compared to the temperature scale, discussed above, at which phase transitions are observed in related compounds.

This study was motivated by the need to calibrate DFT-based methods, inherently applicable at \(T=0\), for use in studies of the broader phase diagram associated with \(\text{LiAg}_x\text{Mn}_8\text{O}_{16-\text{x}-(\text{H}_2\text{O})_x}\). Much of the associated practical, electrochemistry research is based on room temperature characterization. The emergence of strong monoclinic distortion, already for \(\text{Ag}_2\text{Mn}_8\text{O}_{16}\), signaled a disconnect that required further investigation. We have argued here that the formation of localized Mn\(^{3+}\) centers upon reduction by addition of Ag, Li or K, with the associated Jahn-Teller distortions, is likely correct. Parenthetically, we have found that the potential surfaces that describe the resulting cell distortions can be rather flat, making full convergence to the lowest energy structure challenging. In this picture, disorder at finite temperature explains the observation of higher symmetry structures. As a practical matter, calculations that fully account for the complexity of this picture can not be used for phase diagram exploration.

Our results suggest two alternatives. Referring back to Fig. 2 the GGA+\(U\) methods with moderate \(U\) in the range of 3-5 eV give relatively accurate addition energies for Ag, Li and K. Furthermore, inclusion of the full anisotropy in the Coulomb interaction improved the results. However, these calculations, at \(T=0\), will have charge ordered solutions with corresponding distortions of structure. This discrepancy from room temperature structure must be acknowledged, but may not be of practical significance for electrochemical properties, driven largely by energy. Alternatively, Opt-B88 with small \(U\), also gives relatively accurate addition energies. Below the thresholds identified in Fig. 1 electronically driven charge localization does not occur (as distinct from Li ion driven distortions). For mapping phase diagrams in this system, this alternative picture may also be adequate for electrochemical properties. It is also likely that a functional with explicit inclusion of van der Waals interactions will better describe \(\text{H}_2\text{O}\) in the tunnels. However, there is a strong caveat: in this approximation, the reduced system is modeled as a band metal and correspondingly, properties such as optical conductivity may be fundamentally wrong. Finally, we note that some of the features of this phase diagram that we have already described, Ag segregation upon addition of Li and the formation of sheet-like phases with Li, emerge with very similar energetics using either of these two approaches.\(^2\)

V. CONCLUSION

We have carefully mapped out the ground state phases of \(\text{A}_2\text{Mn}_8\text{O}_{16}\) in the hollandite structure for Ag, Li and K in the A position, as they are predicted by DFT-based methods with several different approximate exchange-correlation functionals. These methods included PBE, Opt-B88 (explicitly including van der Waals interactions), their generalization to approximately include local Coulomb interactions among the Mn 3d electrons (GGA+\(U\) approach), the meta-GGA SCAN and finally an exemplary hybrid functional, the HSE approach.

Using HSE as a reference for the A addition energy, we found that Opt-B88 with a small value of \(U\) was rather accurate across Ag, Li and K insertion. SCAN was also relatively accurate overall, but unfortunately predicted that Ag in \(\text{Ag}_2\text{Mn}_8\text{O}_{16}\) was not bound. Furthermore, these methods predicted that the reduced compounds were band metals, with a tetragonal unit cell for the Ag and K cases. This agrees with available, room temperature measurements of structure, but not with the HSE calculations. For Li, the small ionic radius generically led to different local structures such that the ionic interactions with the MnO\(_2\) walls drove local distortions and a monoclinic unit cell. Interestingly, this agrees with predictions of an empirical relationship derived from hollandite mineral data, based on the ratio of the Li ionic radius to that of Mn.

Alternatively, PBE+\(U\) with \(U_{\text{eff}}\) in the 3-5 eV range, showed similar accuracy for the A addition energy. For this range of \(U_{\text{eff}}\), localization of the electrons donated by Ag, Li and K to form specific Mn\(^{3+}\) centers was clearly observed. This agrees with the HSE calculations. The \(Q_3\) order parameter of Van Vleck was found to describe the associated local Jahn-Teller distortion driven by the introduction of the extra 3d electron on Mn\(^{3+}\) into the previously empty \(e_g\) derived states. The net result is strong monoclinic distortion of the unit cell. This does not agree with available room temperature structural data. However, further analysis of competing charge ordered phases shows a series of phases with relatively small energy difference from the predicted ground state phase. These include phases with nearly tetragonal unit cells. Based on comparison with studies of the temperature dependence of the properties of single crystal V and Cr based hollandites, as well as the mixed valence LiMn\(_2\text{O}_4\) spinel compound, we conclude that the charge localization to form Mn\(^{3+}\) centers is likely physically correct. Then the observed high symmetry structures at room temperature
would reflect disorder among the Mn$^{4+}$ centers.

At the same energy scale, the interplay with magnetic order is significant. The HSE calculations predict an AF phase for the parent α-MnO$_2$ and all three A$_2$Mn$_8$O$_{16}$ compounds studied. Here, the best agreement was obtained employing the anisotropic PBE+(U=6, J=1 eV) eV approach, although curiously, not for the Li case. This highlights the challenge in capturing all of the competing effects in these materials with sufficient accuracy for detailed simulation of the competing phases and transitions among them.

We have discussed the consequences of these results for use of these DFT-based methods to survey the phase diagram of Li$_2$Ag$_9$Mn$_8$O$_{16−x}$ where a large number of compounds and structures must be explored efficiently. We have outlined the trade-offs involved. Energetics may be captured with either low $U$ or high $U$ approaches. However, the convenience of avoiding the Jahn-Teller driven distortions will naturally mean sacrificing the ability to predict other properties that are sensitive to charge localization, such as optical conductivity.

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For example, we found 79 symmetrically distinct ways to distribute two Mn$^{3+}$ in the $1 \times 1 \times 2$ super cell and more than 2900 configurations for the $1 \times 1 \times 3$ super cell.

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