Comparative ab initio study of the structural, electronic, magnetic, and dynamical properties of LiOsO$_3$ and NaOsO$_3$

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Despite similar chemical compositions, LiOsO$_3$ and NaOsO$_3$ exhibit remarkably distinct structural, electronic, magnetic, and spectroscopic properties. At low temperature, LiOsO$_3$ is a polar bad metal with a rhombohedral $R3c$ structure without the presence of long-range magnetic order, whereas NaOsO$_3$ is a G-type antiferromagnetic insulator with an orthorhombic $Pnma$ structure. By means of comparative first-principles DFT+$U$ calculations with the inclusion of the spin-orbit coupling, we (i) identify the origin of the different structural ($R3c$ vs. $Pnma$) properties using a symmetry-adapted soft mode analysis, (ii) provide evidence that all considered exchange-correlation functionals (LDA, PBE, PBEsol, SCAN, and HSE06) and the spin disordered polymorphous descriptions are unsatisfactory to accurately describe the electronic and magnetic properties of both systems simultaneously, and (iii) clarify that the distinct electronic (metallic vs. insulating) properties originates mainly from a cooperative steric and magnetic effect. Finally, we find that although at ambient pressure LiOsO$_3$ with a $Pnma$ symmetry and NaOsO$_3$ with a $R3c$ symmetry are energetically unfavorable, they do not show soft phonons and therefore are dynamically stable. A pressure-induced structural phase transition from $R3c$ to $Pnma$ for LiOsO$_3$ is predicted, whereas for NaOsO$_3$ no symmetry change is discerned in the considered pressure range.

I. INTRODUCTION

Transition-metal oxide (TMO) perovskites represent a rich ground for the emergence of intriguing properties and novel phases originating from the complex interplay of different interactions with the cross coupling of spin, charge, orbital, and lattice degrees of freedom [1, 2]. When the transition-metal elements shift from $3d$ to $5d$, spin-orbit coupling (SOC) is enhanced owing to the increased atomic mass and correlation effects weaken due to the extended nature of $5d$ orbitals and associated widening of the band width [3]. Their comparable strength and cooperative interplay in $5d$ TMOs give rise to, e.g., a novel $J_{eff} = 1/2$ Mott-insulating state in an otherwise metallic Sr$_2$IrO$_4$ [4–6]. In addition to iridates, osmium TMOs have also stimulated a lot of interest [7], e.g., because of the observed unusual ferroelectric-like structural transition in metallic LiOsO$_3$ [8, 9] and continuous metal-insulator transition (MIT) [10, 11] and anomalously strong spin-phonon-electronic coupling [12] in NaOsO$_3$.

Despite similar chemical compositions, same electronic configurations ($5d^3$), and comparable electronic correlation and SOC strengths, LiOsO$_3$ and NaOsO$_3$ exhibit strikingly different structural, electronic, and magnetic properties. Experimentally, LiOsO$_3$ displays a bad metallic character over the whole temperature range [8]. It possesses a centrosymmetric $R3c$ rhombohedral structure at high temperature and undergoes a second-order ferroelectric-like structural transition to a noncentrosymmetric $R3c$ structure at $T_c = 140$ K [8]. The origin of this transition was understood by the instability of Li ions along the polar axis and the incomplete screening of the short-range dipole-dipole interactions [13–16]. Although a Curie-Weiss-like behaviour is observed below $T_c$, no evidence of long-range magnetic order is found even down to very low temperature [8, 17]. By contrast, NaOsO$_3$ displays an orthorhombic $Pnma$ structure and undergoes a continuous second-order MIT [10] with a small optical gap (∼ 0.1 eV) [18], which is accompanied by the onset of a long-range $G$-type antiferromagnetic (AFM) ordering at a Néel temperature $T_N = 410$ K with a magnetic moment of 1.0 $\mu_B$ [19]. The MIT in NaOsO$_3$ was initially explained by a Slater mechanism [10, 18–21] and later better interpreted in terms of a continuous Lifshitz-type transition driven by magnetic fluctuations [11, 22, 23]. A detailed comparison between characteristic ground-state (GS) properties and energy scales of LiOsO$_3$ and NaOsO$_3$ is summarized in Table 1.

In addition, the two compounds exhibit remarkably distinct spectroscopic properties [18, 26]. Upon raising the temperature, the AFM insulating state in NaOsO$_3$ develops into a bad metal (pseudogap) regime, which is transformed into a paramagnetic (PM) metallic phase with relatively good Fermi liquid properties at high temperature, as revealed by terahertz and infrared spectroscopy [18]. Conversely, the optical spectrum of LiOsO$_3$ rapidly loses the sign of metallic coherence as the temperature increases. At room temperature, the Drude peak is replaced by a slight low-frequency downturn [26], similar to the behavior observed in undoped V$_2$O$_3$, a prototypical material on the verge of a Mott MIT [27]. By conducting a first-principle many-body analysis we have demonstrated that the distinct high-temperature spectroscopic properties of these two compounds originate from their different degrees of proximity to an adjacent Hund’s-Mott insulating phase [25].

In this paper, by conducting a variety of comparative computational experiments rooted in density functional theory (DFT) plus an on-site Hubbard $U$ and SOC effects, we aim

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to cast some light on the the origin of the di
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openings (metallic vs. insulating state).

In order to seek a common and consistent XC functional that can describe both compounds reasonably well, we have assessed the local density approximation (LDA) [34] in the parametrization of Ceperly and Alder [35], the generalized gradient approximation (GGA) functional Perdew-Burke-Ernzerhof (PBE) [36] as well as its improved version for solid (PBEsol) [37], the strongly constrained appropriately normed (SCAN) meta-GGA functional [38] and the hybrid functional HSE06 [39]. All calculations where done within the noncollinear DFT+$U$ framework [40], based on experimental low-temperature structural parameters of LiOsO$_3$, and NaOsO$_3$ [10]. The conjugate gradient algorithm [41] was used for the electronic optimization with an accuracy such that the total energy difference was less than $10^{-6}$ eV between iterations. In order to quantify the strength of correlation effects, we computed $U$ from the cRPA within the “$t_{2g}/t_{2g}$” scheme [42] based on a nonmagnetic band structure. For more details and notations about cRPA, we refer to Refs. [43, 44]. The matrix elements of on-site Coulomb $U_{ij}$ and exchange $J_{ij}$ interactions calculated with LDA (without SOC) are given in Table II, which yields an orbital-averaged $U$ of 1.94 eV and 1.86 eV for LiOsO$_3$ and NaOsO$_3$, respectively. Note that the calculated $U$ values are insensitive to the specific functional used (e.g., the difference is less than 0.05 eV between LDA and PBE).

To clarify the geometric steric effect due to the different ionic radii of Li$^+$ and Na$^+$, two computational experiments were designed by considering Na-doped R3c-LiOsO$_3$ and Li-doped Pnma-NaOsO$_3$. The alloy structures were modeled using the special quasirandom structure (SQS) method [45] as implemented in the ATAT package [46, 47]. The SQS method mimics the disordered atomic configurations within a supercell of limited size in terms of the correlation functions in the cluster expansion method. For the Na-doped LiOsO$_3$, a supercell with 120 atoms was used, whereas for the Li-doped NaOsO$_3$, a supercell with 80 atoms was employed. A $4 \times 4 \times 4$ k-point grid was used to sampled the Brillouin zone (BZ) of all the supercells. The supercells were fully relaxed (including the cell shape and atomic positions) until the Hellmann-Feynman forces acting on each atom were less than $10^{-4}$ eV.

### Table I. Collection of the low-temperature GS properties of NaOsO$_3$ and LiOsO$_3$.

| Electronic configuration | NaOsO$_3$ | LiOsO$_3$ |
|--------------------------|-----------|-----------|
| Os$^{3+}$ ($t_{2g}$)    | Os$^{3+}$ ($t_{2g}$) |

### Table II. On-site Coulomb $U_{ij}$ and exchange $J_{ij}$ interactions (in eV) ($i$ and $j$ represent $t_{2g}$ orbitals) within the $t_{2g}/t_{2g}$ scheme for R3c-LiOsO$_3$ and NaOsO$_3$ using the LDA functional without SOC.

| $U_{ij}$ | $J_{ij}$ |
|----------|----------|
| LiOsO$_3$ | NaOsO$_3$ |
| $d_{x^2}$ | $d_{y^2}$ | $d_{z^2}$ | $d_{x^2}$ | $d_{y^2}$ | $d_{z^2}$ |

### II. COMPUTATIONAL DETAILS

All first-principles calculations were performed by employing the projector augmented wave method [29] as implemented in the Vienna *Ab initio* Simulation Package (VASP) [30, 31] with the inclusion of the SOC. A plane-wave cutoff of 600 eV was used for both LiOsO$_3$ and NaOsO$_3$. 10×10×10 and 8×8×8 Γ-centered k-point grids generated by the Monkhorst-Pack scheme were used for the rhombohedral LiOsO$_3$ unit cell and orthorhombic NaOsO$_3$ unit cell, respectivly. The ISOTROPY [32] and AMPLIMODES [33] programs were employed to determine the group-subgroup relationships and perform the symmetry-adapted soft mode analysis.
10 meV/Å.

The phonon dispersions and density of states (DOS) were calculated by finite displacements using the Phonopy code [48]. For the cubic perovskite phases, a supercell with 135 atoms was used, whereas a $2 \times 2 \times 2$ supercell was utilized for both $R3c$-LiOsO$_3$ (80 atoms) and $Pnma$-NaOsO$_3$ (160 atoms). For all phonon calculations, a $3 \times 3 \times 3$ $k$-point grid was used to sampled the BZ. Test results show that the phonon DOS are converged with respect to the chosen supercell size and $k$-point grid.

III. RESULTS

A. Crystal structure and symmetry mode analysis

From a theoretical perspective, the different structural symmetries of NaOsO$_3$ ($Pnma$) and LiOsO$_3$ ($R3c$) can be understood in terms of the different Goldschmidt tolerance factor $t = \frac{r_A}{\sqrt{2}(r_O + r_{O^\prime})}$ ($r$ denotes the ionic radius and $A=$Li/Na). In fact, the tolerance factor is often taken as an indicator for the degree of distortion of perovskites [49, 50]: $t=1$ represents the ideal conditions upon which the perovskite structure assumes its ideal cubic symmetry, which is generally stable in the range $0.9 < t < 1$; $t > 1$ favors a hexagonal structure, whereas $0.71 < t < 0.9$ yields rhombohedral or orthorhombic structures [51]. With a tolerance factor of 0.75 (LiOsO$_3$) and 0.84 (NaOsO$_3$), these two Os-based perovskites are predicted to assume rhombohedral or orthorhombic phase, respectively [51], since the slightly smaller $t$ of LiOsO$_3$ (originating from the smaller ionic radius of Li) should result in a more distorted rhombohedral structure. To confirm these expectation we have conducted a symmetry analysis of the phonon dispersions of LiOsO$_3$ and NaOsO$_3$.

Figs. 1(a) and (b) show the calculated phonon dispersions and partial phonon DOS for cubic NaOsO$_3$ and LiOsO$_3$ using the LDA functional, with optimized LDA lattice parameters. One can see that in general the two compounds show similar phonon dispersions except for the soft phonon modes. For cubic LiOsO$_3$ the soft phonons are dominated by Li and O atoms, whereas for cubic NaOsO$_3$ the negative frequencies originate only from O atoms. Both phonon dispersions share structural instability at the $R$ and $M$ points, but LiOsO$_3$ exhibits an additional soft mode at the zone center ($\Gamma$ point). The $R$ mode corresponds to the antiphase octahedral rotation mode $R_\Gamma$ along the [001] and [111] axis for NaOsO$_3$ [Fig. 1(d)] and LiOsO$_3$, respectively. By moving along this mode, the cubic phases of NaOsO$_3$ and LiOsO$_3$ reduce to the orthorhombic $Imma$ structure and centrosymmetric rhombohedral $R3c$ structure [Fig. 1(h)], respectively. Symmetry-adapted soft phonon analysis indicates that the instabilities at $M$ and $\Gamma$ are due to the in-phase octahedral rotations [$M_2^c$, Fig. 1(e)] and the ferroelectric distortions associated with the displacements of Li atoms along the polar [111] axis [$\Gamma_{5c}^c$, comparing Fig. 1(i) to Fig. 1(h)]. Further condensing these modes leads to the formation of $Pnma$-NaOsO$_3$ [Fig. 1(g)] and noncentrosymmetric $R3c$-LiOsO$_3$ [Fig. 1(i)]. A diagram showing the group-subgroup relationships is shown in Fig. 1(c). It is worthy noting that an additional octahedral tilt mode $X_5^-$ occurring in NaOsO$_3$ [Fig. 1(f)] is a secondary mode and it appears as a consequence of the combined effect of the primary modes $R_5$ and $M_2^c$. With respect to the parent cubic phase, the normalized amplitudes of the modes $R_5^c$, $M_2^c$, and $X_5^-$ for the experimental $Pnma$-NaOsO$_3$ structure are estimated to be about 0.72 Å, 0.50 Å, and 0.28 Å, respectively, while those associated with the modes $R_5^c$ and $\Gamma_{5c}^c$ for the rhombohedral $R3c$-LiOsO$_3$ structure are about 1.22 Å and 0.47 Å, respectively. Our mode analysis on LiOsO$_3$ is consistent with Ref. [14].

B. Assessing the XC functionals on electronic and magnetic properties

It is well known that the specific form of the XC functional plays an important role in first-principles DFT simulations and that finding an XC functional capable to account for the basic ground state properties is nontrivial, especially for complex materials. LiOsO$_3$ and NaOsO$_3$ represent typical examples that pose great challenges for the choice of an XC functional and no consensus has been achieved yet in literature due to a lack of proper scrutinization. For instance,
NaOsO$_3$ was studied using LDA in Refs. [10, 20, 21], and using PBE in Ref. [11]. Similarly, for LiOsO$_3$, LDA was used in Refs. [13, 15, 55], while PBE was employed in Refs. [14, 16] and PBEsol in Ref. [56]. Considering the quantitative and – to some extent– qualitative discrepancies between the results obtained by different XC functionals and in order to achieve a trustable and convincing comparative study between these two systems, we have performed a systematic assessment of the performance of LDA, PBE, PBEsol, and SCAN within a DFT+U+SOC framework (with $U$ ranging from 0 to 2.4 eV) as well as HSE06 for the prediction of band gaps and magnetic properties of NaOsO$_3$ and LiOsO$_3$. The GS magnetic states are determined by comparing the total energy difference between the two energetically favorable configurations, $G$-type AFM state and nonmagnetic state, for each $U$ value [55]. The results displayed in Fig. 2 show that none of the considered functionals is capable to simultaneously predict an insulating magnetic state for NaOsO$_3$ and a non-magnetic metallic state for LiOsO$_3$.

As a general and expected trend, we remark that the inclusion on the onsite $U$ tends to favor an insulating solution and to establish a magnetic ordering: Both band gap and local magnetic moment increase with increasing $U$, but a single value of $U$ cannot establish the desired ground states in both systems. The situation is particularly problematic for LiOsO$_3$, as discussed in more details in the following.

First, we note that SCAN, typically considered to be a rather accurate scheme, tends to overestimate the magnetic moments for both compounds [Figs. 2(b) and (d)], as it does for itinerant electron ferromagnets [57, 58]. Also, it overestimates the band gap for NaOsO$_3$ [Figs. 2(a) and (b)]. We note that similarly to SCAN, HSE06+SOC with default screening length ($\mu=0.2$) and exact exchange mixing parameters ($\alpha=0.25$) delivers an even larger band gap and magnetic moment: It gives an incorrect magnetic insulating state for LiOsO$_3$ with a gap of 1.25 eV and a moment of 1.70 $\mu_B$/Os, while for NaOsO$_3$ it predicts a gap of 1.43 eV and a moment of 1.67 $\mu_B$/Os. It should be noted, however, that hybrid functionals are sensitive to the choice of $\mu$ and $\alpha$ parameters and for moderately correlated itinerant systems the optimal value should deviate substantially from the default ones [59, 60]. In the following we will focus on a detailed discussion of the LDA, PBE, and PBEsol results.

In line with previous studies [10, 11, 20, 21], without $U$, LDA, PBE, or PBEsol fail to open the band gap in NaOsO$_3$ [Fig. 2(b)], and while PBE and PBEsol find a sizable local moment, LDA favors a nonmagnetic solution, in disagreement with experimental observations. The situation in LiOsO$_3$ is similar with the only exception that PBEsol does not stabilize any magnetic solution, in this case in line with the experimentally observed metallic nonmagnetic ground state.

With increasing $U$, the situation remains problematic. Above a certain critical $U_c$, both systems undergo a MIT and both the gap and magnetic moments grow almost linearly as a function of $U$. The values of $U_c$ are listed in Table III. A positive outcome of the calculations is that $U_c$ is systematically lower in NaOsO$_3$ than in LiOsO$_3$ implying that there exists a $U$ window for each functional where NaOsO$_3$ is insulating and LiOsO$_3$ metallic (LDA: 1.0-1.4 eV; PBEsol: 0.4-0.8 eV; PBE: 0.2-0.4 eV). The downside is that within these $U$ ranges, both systems are found to be magnetic, which is good for NaOsO$_3$ but in apparent disagreement with experiment for LiOsO$_3$. More precisely, PBE+U+SOC yields an ordered magnetic moment for LiOsO$_3$ for all $U$ values. On the other hand, LDA and PBEsol yield a nonmagnetic solution in the low-$U$ limit, but a magnetic moment develops for $U$ values larger than $U_c$, which is therefore outside the $U$ range indicated above. As soon as the $U$ reaches $U_c$, a well established magnetic moment $M_c$ of about 1 $\mu_B$ is found for all functionals as reported in Table III.

Summing up, LiOsO$_3$ is nonmagnetic and metallic for $U \leq 0.2$ eV (PBEsol) and $U \leq 0.8$ eV (LDA). In this $U$ range,
FIG. 3. LDA+U+SOC calculated magnetic moments \( M (\mu_B/\text{Os}) \) and band gaps \( E_g \) (eV) as a function of \( U \) for LiOsO\(_3\) and NaOsO\(_3\) with G-AFM and SQS-PM polymorphous descriptions.

However, NaOsO\(_3\) is magnetic but always metallic. The most likely cause of this apparent disagreement is the shortcoming of mean-field DFT in the LDA or GGA to account for magnetic fluctuations in itinerant magnets \([61, 62]\), and this leads to a systematic overestimation of the local ordered magnetic moment.

It needs to be noted that although at low temperature a long-range magnetic order is absent in LiOsO\(_3\) \([8]\), the Curie-Weiss-like behaviour observed below \( T_\chi \) \([8]\) and the \( \mu\)SR experiments \([17]\) suggest a disordered PM ground state. Within DFT it is challenging to model the PM state. Recently, G. Tirmarchi et al. \([28]\) proposed a polymorphous description for the spin disordered state, which is realized by a supercell calculation modeled in the SQS manner (termed SQS-PM) \([28]\).

Using this method, the gap opening and orbital ordering of the paramagnetic phases of the transition-metal monoxides \([28]\) as well as the 3\textit{d} perovskite oxides \([63]\) are reasonably well described.

Aiming to improve the description of the PM phase in LiOsO\(_3\), we have applied this SQS-PM approach using the LDA functional. The results are shown in Fig. 3 where we show the correlation between \( U \) and the magnetic moment \( M \) [Fig. 3(a)] and the correlation between \( M \) and the band gap \( E_g \) [Fig. 3(b)]. One can observe that for \( U \) smaller than 1.6 eV, the SQS-PM (stars) predicts a nonmagnetic state. As \( U \) increases, the magnitude of the disordered magnetic moment increases and a MIT appears for \( U \) larger than 2 eV. By contrast, the SQS-PM description of NaOsO\(_3\) always gives metallic solutions for the considered \( U \) values, in line with the observation that only G-AFM is capable to open the gap, whereas all other magnetic orderings yield a metallic solutions \([20]\).

Although the larger critical \( U_c \) required for the MIT in the SQS-PM phase seems to mitigate the above-mentioned issues of the XC functionals, the SQS-PM solutions of LiOsO\(_3\) are always higher in energy than the G-AFM ordered phases. Taking \( U=1.8 \) eV for instance, the energy difference is about 71 meV/f.u. Therefore, even the SQS-PM method is not a satisfactory solution and one might have to resort to either a new XC functional or a new method to treat itinerant magnetism in DFT \([61, 64, 65]\).

We conclude this section with a remark on the correlation strength of these compounds. From Fig. 2 one can see that the cRPA estimated \( U \) values (dashed lines) are very large and fall in a range in which both systems are magnetic insulator. We have previously reported that SOC effects could induce a considerable renormalization of the Coulomb interaction in NaOsO\(_3\) of about 1 eV, placing NaOsO\(_3\) in the moderately correlated regime. This spin-orbit renormalization was also found to be necessary to correctly describe the Lifshitz transition of NaOsO\(_3\) \([11]\). The comparison of the band structures with and without SOC for nonmagnetic LiOsO\(_3\) and NaOsO\(_3\) (Fig. 4) indeed shows that the inclusion of SOC increases the bandwidth of the \( t_{2g} \) states and thus enhances electron mobility. This reduces the correlation strength and leads to smaller \( U \) value as compared to the one obtained without SOC \([11, 66]\). However, a precise quantification of the SOC renormalization effect is a difficult task requiring the inclusion of SOC in the cRPA calculation. Unfortunately, to our knowledge, no cRPA implementation is available to compute \( U \) with SOC because of technical complexity in treating the complex-valued Wannier spinors within the cRPA scheme. cRPA calculations without SOC suggest that the two compounds have very similar Coulomb parameters (see Table II), and therefore, in the following calculations we will adopt the same \( U \) for both materials.

C. Cooperative steric-magnetic driven MIT

Since GGA overestimates the magnetic moment for itinerant magnets even more than LDA \([61]\) and LDA performs generally better than GGA in predicting ferroelectric properties \([67]\), in the following calculations the LDA with \( U = 1.2 \) eV is employed [results obtained for \( U=1.4 \) eV show very similar trends (not shown)].

As discussed previously, it is the difference in the tolerance factor that dictates the different crystal structures in LiOsO\(_3\) and NaOsO\(_3\). Then a natural question is whether this is also the origin of their distinct electronic properties. To verify this hypothesis, we designed two computational experiments to
track the transition between LiOsO$_3$ and NaOsO$_3$ via chemical doping using LDA+U+SOC in combination with SQS. We have inspected the following two scenarios. (i) Doping R3c-LiOsO$_3$ with Na. In this way, we study how Na doping affects the metallic ground state of LiOsO$_3$. (ii) Doping Pnma-NaOsO$_3$ with Li, where we control the influence of Li doping on the metallic state of NaOsO$_3$. For different Na (or Li) contents we have computed the optimized volume, the magnetic moment and the band gap in the R3c and Pnma phase. The results are shown in Fig. 5.

Let us first focus on Na-doped LiOsO$_3$ (squares in Fig. 5). As expected, the volume (tolerance factor) increases almost linearly as the Na doping concentration $x$ increases. At $x \approx 20\%$ a MIT appears and the band gap increases further as $x$ increases. The magnetic moment increases very slowly within the standard deviation of the fluctuating moments induced by disorder effects. Analogously, in Li-doped NaOsO$_3$ (circles in Fig. 5) as the Li concentration $(1-x)$ increases, the volume and the tolerance factor decrease and the insulator-metal transition occurs at a low Li concentration of 6.25%.

The overall similar trends in Na-doped LiOsO$_3$ and Li-doped NaOsO$_3$ convey a clear conclusion: The electronic ground state is mainly controlled by steric effects. The larger atomic radius of Na increases the volume and thus favors the onset of the insulating state. By replacing back Na with Li in the insulating phase of Na-doped LiOsO$_3$ and performing electronic self-consistent calculations while keeping the atom’s positions fixed at the corresponding Li sites, it is found that the band gap remains open. Expectedly, by fixing the volume, R3c-LiOsO$_3$ displays a larger magnetic moment and a larger tendency to become insulating than Pnma-NaOsO$_3$, as shown in Fig. 6(a). It is also obvious that it is the larger/smaller ground-state volume of NaOsO$_3$ that makes the system insulating/metallic. In addition, it is found that the band structure at a fixed crystal structure is essentially insensitive to the Na/Li cation (not shown). All these facts imply that the origin of the different electronic (metallic vs. insulating) properties of the two compounds is primarily driven by steric effects. The detailed changes on the effective band structure due to doping are displayed in Fig. 7, highlighting the emergence of the MIT in Na-doped LiOsO$_3$ and Li-doped NaOsO$_3$.

However, it is worth noting that the presence of magnetic order also plays an important role in the onset of the MIT. As shown in Figs. 6(b) and (c), the MIT appears only when the magnetic moment is larger than a critical value, which gets progressively reduced by increasing the volume. The fact that a larger volume favors a larger magnetic moment and the
FIG. 7. Evolution of the effective band structure (EBS) in Na-doped LiOsO$_3$ (upper panels) and Li-doped NaOsO$_3$ (bottom panels) (selected concentrations), unfolded in the corresponding primitive cell by means of the band unfolding technique [68, 69]. The lateral bar indicates the amount of the Bloch character. The sharpness of the EBS reflects the effect of the chemical disorder.

larger magnetic moment in turn assists the opening of the band gap suggests that the onset of MIT for the two compounds is driven by a cooperative steric and magnetic effect.

D. Structural stability and phase transition

Now we turn to discussing dynamical properties and possible structural phase transitions in LiOsO$_3$ and NaOsO$_3$. As expected, the GS phases of R$_3c$-LiOsO$_3$ and Pnma-NaOsO$_3$ are dynamically stable, as revealed by the phonon dispersions displayed in Figs. 8(a) and (d). Although the Pnma-LiOsO$_3$ and R$_3c$-NaOsO$_3$ phases at ambient pressure are energetically less favorable than the corresponding GS phases by <60 meV/f.u. (see Fig. 9), they turn out to be dynamically stable, since no soft mode appears in the vibrational spectra [see Figs. 8(b) and (c)]. We also mention that the NaOsO$_3$ with polar R$_3c$ symmetry is unstable and reduces to the nonpolar R$_3c$ symmetry after structural relaxations, suggesting that ferroelectric instabilities are not expected in NaOsO$_3$ under these conditions. This is consistent with the phonon calculations for cubic NaOsO$_3$, where no instability at the $\Gamma$ point is observed [Fig. 1(a)]. For LiOsO$_3$, on the other hand, we predict a structural phase transition from the R$_3c$ phase to the Pnma phase at a pressure of about 20 GPa [see the inset of Fig. 9(a)], consistent with the theoretical findings of E. Aulestia et al. [56]. Even above the transition pressure, we find that the Pnma-LiOsO$_3$ phase is dynamically stable (not shown), indicating that such a phase transition can be achievable for LiOsO$_3$ in high-pressure experiments. However, for NaOsO$_3$ our calculations do not discern any symmetry change within the considered pressure range: The Pnma phase is always stable and becomes progressively stabilized over the R$_3c$ phase by increasing pressure [Fig. 9(b)].

Interestingly, we also find that the Pnma-LiOsO$_3$ or R$_3c$-

NaOsO$_3$ phases seem to be dynamically stable only if the magnetic order is present. For instance, with a smaller $U=0.8$ eV, both Pnma-LiOsO$_3$ and R$_3c$-NaOsO$_3$ are found to be magnetically-ordered metals, without any soft phonons. If the magnetic moment is removed, e.g., by performing a non-spin-polarized LDA calculation, soft phonons appear and the system becomes dynamically unstable, a further indication of the
strong spin-lattice effects in this class of compounds [12].

IV. CONCLUSIONS

In conclusion, by comparative ab initio DFT+U+SOC calculations, we have systematically studied the strikingly distinct structural, electronic, magnetic, and dynamical properties of the two chemically similar osmates perovskites, LiOsO$_3$ and NaOsO$_3$.

First, we find that none of the considered XC functionals (LDA, PBE, PBEsol, SCAN and HSE06) is capable to accurately predict the correct electronic and magnetic ground state for both compounds simultaneously. This drawback is mostly due to the difficulties of DFT (within local, semilocal and nonlocal treatment of XC effects) in treating the magnetic fluctuations associated with the itinerant nature of LiOsO$_3$ and NaOsO$_3$. Neglecting fluctuations ultimately leads to an overestimation of magnetic moments. The SQS-PM approach allows for an improved description in stabilizing a PM state in LiOsO$_3$, but it incorrectly predicts a higher energy for the PM phase than for the G-AFM phase in LiOsO$_3$. In comparison to NaOsO$_3$, LiOsO$_3$ is less magnetic due to its smaller volume and to induce the MIT it requires a larger critical $U_c$. Though using a different $U$ in the two systems (for instance $U$=1.1 eV in NaOsO$_3$ and $U$=0.7 eV in LiOsO$_3$) would lead to a reasonable description of the two distinct ground states, the verification of this hypothesis would require the ab initio calculation of $U$ including SOC effects, which is, however, currently not possible.

Second, by following the transition from one compound to the other via chemical doping, we clarify that it is the cooperative steric and magnetic effects that controls the electronic properties and drives the formation of the distinct metallic/insulating state in the two systems: The larger/smaller volume of NaOsO$_3$/LiOsO$_3$ leads to a larger/smaller magnetic moment, which in turn assists the opening/closing of the band gap.

Finally, the different GS crystal structures ($R3c$ vs. $Pnma$) of LiOsO$_3$ and NaOsO$_3$ can be explained by purely steric effects and arise from the different Goldschmidt tolerance factors (0.75 vs. 0.84). Moreover, we show that the energetically unfavorable phases of $Pnma$-LiOsO$_3$ and $R3c$-NaOsO$_3$ at ambient pressure are dynamically stable. A pressure-induced structural phase transition from $R3c$ to $Pnma$ for LiOsO$_3$ is predicted, whereas for NaOsO$_3$ the $Pnma$ phase is stabilized over the $R3c$ phase by increasing pressure, suggesting that under these conditions NaOsO$_3$, unlike LiOsO$_3$, does not seem to be prone to ferroelectric instabilities.

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