I. INTRODUCTION

The physical origin of the insulating phases occurring in crystals with partially occupied $d$ shells exemplified by the transition metal (TM) monoxides MnO, FeO, CoO, and NiO has held the condensed matter physics community in constant fascination ever since Mott and Peierls proposed an explanation [1]. These oxides have a low-temperature spin-ordered antiferromagnetic (AFM) phase, in which they exhibit slightly distorted rock-salt structures (rhombohedral for NiO and MnO and monoclinic for CoO and FeO, the latter exhibiting Jahn-Teller atomic displacements), and a high-temperature spin disordered paramagnetic (PM) phase, having the macroscopically cubic rock-salt structure and globally null magnetization. In simplified band structure calculations [2–7], it has been customary to evaluate the electronic structure of the PM phase of these oxides using the macroscopically observed average rock-salt configuration $\Sigma_0$. Because there is but a single transition metal atom in such a unit cell, the condition of globally zero magnetization in the PM phase leads to vanishing local magnetic moments $\mu_i(\Sigma_0)$ at each metal site $i$, and, therefore, by symmetry, to zero band gaps $E_g(\Sigma_0)$ for such systems with partially filled $d$ orbitals. Here, $\Sigma_0$ was taken as the nonmagnetic, cubic rock-salt configuration in which all TM sites are equivalent (a monomorphous representation). As is generally taught [8,9], the ensuing electronic structure of compounds having partially occupied energy bands described in a structure where all the atoms are equivalent would be metallic [2,3] with the Fermi level intersecting a band. Yet, experimentally, MnO, FeO, CoO, and NiO are local-moment large band gap insulators, both in the AFM and PM phase [10–12] (see Table I).
This fundamental disagreement between such band structure theory and experiment set the historical stage for modeling the electronic structure of the PM phases of MnO, FeO, CoO, and NiO, and related quantum materials by many-body, correlated electron descriptions, such as the description based on the Hubbard Hamiltonian \(^{[13,14]}\), or, more recently, the dynamical mean-field theory (DMFT) \(^{[15,16]}\) rendering of the Hubbard Hamiltonian. Within such theories, the gap of the AFM and PM phases of these oxides emerges because the d electrons become localized due to the correlation-induced electron-electron repulsion, even in the absence of spatial symmetry breaking (symmetry can break afterwards, as a secondary fact). From the strongly correlated standpoint, the existence of local magnetic moments is a consequence of the electron localization and not an essential part of the gap opening mechanism itself. The Mott mechanism envisions that the electrons move across the lattice forming states on certain atomic sites with doubly occupied d orbitals and empty d orbitals on other sites. These types of excited configurations correspond respectively to the upper and lower Hubbard bands, which are envisioned to form the physical band edges, i.e., the valence-band maximum (VBM) and conduction-band minimum (CBM).

Indeed, one often finds in the literature comments that single determinant mean-field DFT band theory fails to reproduce the gap in the absence of long-range magnetic order \(^{[4,17–19]}\). But such conclusions may have been clouded by a few restrictions applied unwittingly to band theory itself. For example, the model that has been often used for the PM phase of MnO, FeO, CoO, and NiO restricts all metal sites to see identical local environments and potentials, i.e., such a model uses unit cells with a single TM per cell. Because such a monomorphous representation of a disordered PM phase forces upon us in band theory a zero magnetic moment on an atom-by-atom basis (instead of nonzero local moments with a global zero magnetization), the ensuing band gaps in such nonmagnetic models were always zero, irrespective of the quality of the description of the interelectronic interaction \(E_{\text{xc}}[n_i^+(r), n_i^-(r)]\) in DFT band theory. However, as shown here, upon examining the total DFT energies, one finds that the monomorphous nonmagnetic (NM) “phase” is unstable by more than 2 eV/formula unit than a proper DFT paramagnetic phase of CoO, so the NM phase is but a hypothetical state. In addition to confusing paramagnetic state with nonmagnetic state, other approximations may have perhaps prematurely disqualified DFT band theory from describing even simple Mott systems and related quantum materials. These approximations include (i) disallowing geometrical symmetry breaking (such as Jahn-Teller, atom pairing, or charge disproportionation) or (ii) using DFT functionals that do not distinguish occupied from unoccupied states (i.e., lacking exchange correlation discontinuity as in self interaction corrected DFT or its DFT + U approximant), thus forcing equal, and fractional, occupations of all components of open shell degenerate states at the Fermi energy. Since none of these simplifications are defining features of band theory itself, the failure of such approaches in explaining Mott insolation may have been prematurely viewed as a fundamental failure of the single determinant mean-field approach itself. The path then suggested in the literature to model the paramagnetic state of Mott insulators has been that of solving the Hubbard model by strongly correlated approaches. (Perhaps an early clue that d electron strong correlation is not the deciding factor for these binary oxide systems is the fact that the valence band and conduction band edges consist of oxygen p and transition metal s orbitals, shown below, not the proverbial d-like lower and upper Hubbard bands, envisioned by the founding thinkers in this field.) Here, we relax the restrictions often imposed previously but which are not an integral part of band theory per se, seeking to understand the minimum theoretical approach needed to describe the leading features of the Mott insulating behavior, namely, gap opening and amplitude and on-site magnetic moment formation in the AFM and PM phase of the classic, 3d monoxide Mott systems. To this end, we deliberately use a single-determinant, mean-field Bloch periodic band structure approach (based

<table>
<thead>
<tr>
<th>(3\overline{m})</th>
<th>MnO</th>
<th>NiO</th>
<th>(C2/m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exp.</td>
<td>DFT + U</td>
<td>Exp.</td>
<td>DFT + U</td>
</tr>
<tr>
<td>AFM</td>
<td>(\mu (\mu_r))</td>
<td>4.58</td>
<td>4.64</td>
</tr>
<tr>
<td></td>
<td>(E_{\text{gap}}) (eV)</td>
<td>3.5</td>
<td>1.88</td>
</tr>
<tr>
<td>FM</td>
<td>(\mu (\mu_r))</td>
<td>–</td>
<td>4.68</td>
</tr>
<tr>
<td></td>
<td>(E_{\text{gap}}) (eV)</td>
<td>–</td>
<td>0.82</td>
</tr>
<tr>
<td></td>
<td>(E_{\text{tot}}) – (E_{\text{tot}}) [AFM] (eV/fu)</td>
<td>–</td>
<td>0.059</td>
</tr>
<tr>
<td>PM</td>
<td>(\mu (\mu_r))</td>
<td>4.65</td>
<td>1.70</td>
</tr>
<tr>
<td></td>
<td>(E_{\text{gap}}) (eV)</td>
<td>3.7</td>
<td>2.22</td>
</tr>
<tr>
<td></td>
<td>(E_{\text{tot}}) – (E_{\text{tot}}) [AFM] (eV/fu)</td>
<td>0.014</td>
<td>0.061</td>
</tr>
</tbody>
</table>
on DFT + $U$), but allow for any energy-lowering symmetry breaking effects that can be captured by such a mean-field description.

We find here for the text-book MnO, FeO, CoO, and NiO Mott insulators that symmetry breaking afforded by using in band theory sufficiently flexible unit cells (the polymorphous representation of disorder) and allowing orbital occupation symmetry breaking and spin unrestricted symmetry breaking produce in DFT + $U$ [20–24] an insulating solution (band gaps of 1–3 eV) with strong local magnetic moments (2–4 Bohr magnetons), in general agreement with experiment. This symmetry breaking is not assumed, but obtained as an energy-lowering event within the mean-field-like variational theory, and differs from a correlation-induced gap opening mechanism that lacks spatial symmetry breaking. Previous authors of DMFT calculations [25–27] obtain large moments and large band gaps in the PM phases of these 3d monoxides, just as the present single determinant mean-field DFT band structure study does (see Table I). The DMFT authors state, however, that “This result is in remarkable contrast to that previously obtained by standard band-structure methods, which predict metallic behavior, e.g., for FeO and CoO. This implies the crucial importance of strong correlations of localized 3d electrons, to determine the electronic and magnetic properties of transition metal oxides.” We note that these authors refer to the nonmagnetic (monomorphous) metallic DFT description that is $\sim 2$ eV/formula unit higher in energy than the proper gapped PM phase which we discuss in the present work. Indeed, such a high energy NM metallic structure does not appear to be a reasonable representation of what band theory does. The present single-particle approach produces detailed geometrical information (such as atomic displacements and the equilibrium cell structure), magnetic moments, band gaps, bonding charge density, and the total energies of all phases, and permits the future use of precisely the same polymorphous description for studying defects and doping. The emerging physical picture is that at any given moment in time, (i.e., a snapshot) each TM atom will have a finite magnetic moment (but randomly pointing to different directions), and these moments are rotating with time. So, the average magnetic moment of each atom will not be equal to zero due to a quantum fluctuation. The conventional physics encoded in DFT with current nonlocal functionals—bonding, magnetism, spin-polarization, Jahn-Teller distortions, hybridization and selective occupation of the components of the $e_g$ and $t_2$ orbitals—suffices to produce large moments if not disallowed by imposing artificially high symmetry. As a result of these symmetry-breaking channels, electrons are exchanged between spin-up and spin-down bands to create an insulating closed shell configuration that lends itself to a single determinant description. Whereas the final expression would involve a (macroscopically symmetrized) combination of all individual symmetry broken configurations, the crucial fact is that such individual symmetry broken configurations have closed shells with large band gaps and magnetic moments, suggesting that such configurations would have but weak mutual interaction, so dynamically correlated approaches are not forced upon us by the physics at hand. We note that it is entirely possible that some other Mott compounds would remain metallic in a single-particle treatment even if the monomorphous description were replaced by a polymorphous representation. It is also possible that DFT with the current functional could miss properties other than the insulating character of the PM phases. We are not claiming otherwise. These are open, future research questions.

The fact that the main attribute of the classic Mott insulators (i.e., the existence of gaps) could be described by mean-field, single configuration band theory is consequential, as it redefines the minimal level of physics and computational effort required. Indeed, such single-determinant mean-field-like DFT approaches have recently successfully described in agreement with experiment the hole density in $p$-type doped NiO [28], the metal-insulator transition in rare-earth nickelates [29], and the $n$-type doping of SmNiO$_3$ [30], discussed in Ref. [31]. Such results may open the door for reexamination of the utility of non-naïve DFT based band theory methods for studying the basic attributes of more complex Mott systems, including ternary and quaternary oxides, as well as defects, doping, and interface characteristics.

II. DUAL INPUTS TO ELECTRONIC STRUCTURE THEORY

Any electronic structure method requires specifying (a) a representation for the crystal structure (and, for random systems such as the PM phases, the way the configurational average is performed), as well as (b) the type of electronic interactions allowed by the Hamiltonian and its solver (e.g., the forms of exchange and correlation in band theory or the dynamic correlation in explicitly correlated theories). In regard to (b), Table I shows the magnetic moments and band gaps calculated by ordinary DFT + $U$ for the AFM as well as the assumed ferromagnetic (FM) phases of MnO, FeO, CoO, and NiO. Recall that the DFT + $U$ method [24] as well as related methods such as the self-interaction corrected DFT [32] and the hybrid functionals [33] are all single-particle schemes in which the wave function of the $N$-electron system is a single determinant. We see from Table I that even in the single-determinant DFT + $U$ description with a reasonable value of $U$, i.e., 5 eV, which we used across the board for all compounds and spin configurations, these transition-metal monoxides, regardless of the type of magnetic ordering (AFM or FM), exhibit large local moments and band gaps. This opens the possibility that the actual magnetic order may not be the primary reason for these materials to be insulators, but the existence of on-site magnetic moments caused by the breaking of the symmetry between spin-up and spin-down electronic states may. We will follow this hint by removing in band theory the conceptual and computational barriers to the formation of energy-lowering symmetry breaking.

III. ALLOWING FOR A POLYMORPHOUS DESCRIPTION OF THE MAGNETIC STRUCTURE OF THE PARAMAGNETIC PHASES: BASIC CONCEPTS

Gapping in 3D metals can occur due to magnetic spin effects even without symmetry breaking (as is the case for the hypothetical FM phase in Table I). In this case, there is no atomic symmetry breaking and the presence of a band gap is a local electronic effect due to the possibility to make majority
spin and minority spin occupy different states with the result of forming spin up and spin down closed shells. (Note, however, that the FM phase is not a viable representation for the PM phase because the PM phase is disordered while the FM is ordered; what is more as Table I shows, the total energy $E(\text{FM})$ of the PM phase is higher than the total energy $E(\text{PM})$ of the PM phase even at low temperature.) However, gapping in a 3D metal can also occur due to nonmagnetic atom symmetry breaking effects because of their ability to create closed shell structures. This is illustrated by the emergence of a gap in a metal as a result of the geometric flexibility that enables electronic symmetry lowering. For example, the DFT total energy of cubic perovskite BaBiO$_3$ with a single Bi$^{4+}$ site per cell (a metal) is lowered by doubling the cubic perovskite primitive cell, allowing two Bi$^{4+}$ ions to express their multivalent nature by disproportionation into Bi$^{3+} + $ Bi$^{5+}$ (see Ref. [34]) with each site having its own local bond geometry, i.e., a symmetry lowering that causes gapping. Similarly (see Ref. [35]) for CsTlF$_3$ being allowed to express the multivalence of Tl in each site having its own local bond geometry, i.e., a symmetry lowering that causes gapping. In all such cases, a restricted structural description (one type of octahedron) incorrectly produced a metal, whereas a more flexible description of the cell lowered the total energy and produced the observed insulating gap in band theory. Such geometrical freedom needs to be exercised also for spin alloys to concomitantly examine its effect on the total energy and possible gapping.

Here, we wish to examine if significant on-site magnetic moments might produce a gap also in the PM phases within a single-determinant approach, had these moments not been eliminated at the outset by selecting for the representation for the crystal structure the monomorphous macroscopically averaged configuration $\Sigma_0$ where each TM site sees the same local environment. In the latter case, the global zero moment characteristic of the PM phase is interpreted on an atom by atom basis so $\mu_i(\Sigma_0) = 0$ at each metal site $i$, leading in band theory (where large moments mean large exchange splitting) to a band gap $E_g(\Sigma_0) = 0$. The polymorphous approach allows each TM atom to see a distinct local magnetic environment and a locally varying density-functional potential $V[\rho(r), m(r)]$ [where $\rho(r)$ and $m(r)$ are, respectively, the electron density and magnetization at position $r$] subject to the constraint that the total magnetic moment is zero as must be in a paramagnet. We will enquire if such a representation has sufficient geometrical freedom (e.g., unrestricted spatial symmetry) to allow in a self-consistent DFT($+U$) calculation the evolution of local magnetic moments on individual sites, if this would lower the total energy. In band theory language, large moments imply large exchange splitting which could enable large gaps.

**A. Using the average $\langle P \rangle$ of the properties $\{p_i\}$ of individual configurations $\{\sigma_i\}$ versus using the property $P(\Sigma_0)$ of the average configuration $\Sigma_0 = \langle \sigma_i \rangle$**

A common misconstrued naïve DFT approximation in calculating an observable macroscopic electronic or magnetic property $P$ of a disordered random alloy $A_iB_{1-x}$ of composition $x$ is to substitute the calculation of the ensemble average $\langle P \rangle$ for property $P$ by the calculation of the property $P(\Sigma_0)$ of the macroscopically averaged configuration $\Sigma_0$. This monomorphous approximation has been used in the single-site coherent potential approximation (s-CPA) for chemical alloys [36] where all $A$ ($B$) sites in the random alloy see the same potential $V_A$ ($V_B$), irrespective of the existence of different local environments for different $A$ ($B$) sites (characterized by different numbers of $A$ versus $B$ nearest neighbors to a given central atomic site). This approach in alloy theory forced vanishing charge fluctuations (hence zero Madelung contribution to the total energy [37,38]) and vanishing atomic displacements [39–42], both in conflict with more general theories (such as supercells [37,39,43,44] having a polymorphous distribution of different $A$ sites each with its local environment (and same for $B$ sites)).

The correct way to calculate the property $P$ of a phase that can have numerous individual configurations $\{\sigma_i^{(n)}\}$ each with property $P(\sigma_i^{(n)})$, where $n$ is the index of the configuration, is to calculate the polymorphous statistical average $\langle P \rangle = \sum_n c_n P(\sigma_i^{(n)})$ over the ensemble of microscopic spin configurations accessible to the system, instead of assuming $\langle P \rangle = P(\Sigma_0)$. The former approach has largely replaced the monomorphous approaches (s-CPA, virtual crystal approximation) in the theory of disordered substitutional alloys $A_iB_{1-x}$, producing qualitatively different results [37,39,43,44] in much better agreement with experiment. The same polymorphous approach can be applied to spin disordered phases, i.e., the PM phase of Mott insulators.

**B. Time average versus spatial average**

In a spin-disordered phase, the orientation of the on-site magnetic moments $\mu_i$ can change over time showing spin wave excitations representing the low-energy scale of the problem; such fluctuations in the orientation of the moments have zero overall average. In single-impurity DFMT, the Hubbard model is mapped onto an equivalent Anderson single impurity model in which the electrons at the impurity interact with a mean-field bath that models the rest of the system. There is but a single impurity, which fluctuates in time to give a time average of zero spin.

We examine instead how accurately can one predict the gaps of the PM phases of MnO, FeO, CoO, and NiO within a single-determinant description, if one correctly estimates the statistical average $\langle P \rangle = \sum_n c_n P(\sigma_i^{(n)})$ over the ensemble of microscopic spin configurations accessible to the system instead of forcing a zero moment on an atom-by-atom basis. Using a supercell consisting of $2 \times 2 \times 2$ or $3 \times 3 \times 3$ primitive cells, rather than a single NaCl primitive cell ($N = 1$) permits different local environments around each TM atom. Such a polymorphous representation allows for a number of degrees of freedom, consistent with an overall paramagnetic state, including (i) different numbers of TM neighbors to each central atom with spin up and spin down (geometric fluctuations), (ii) different occupations of the 3$d$ orbitals on each TM (occupation number fluctuation), as well as (iii) different local displacements (positional fluctuations), and consequently, (iv) different local magnetic moments. The magnitude (zero or otherwise) of these fluctuations is determined in a charge self-consistent DFT calculation by seeking lower total energies. This provides for an intrinsically multisite representation.
involving a distribution of transition metals, each with its own local environment. No mean-field-embedding bath is needed, and no arbitrarily selected symmetry breaking is involved.

Our underlying conjecture is that time fluctuations that could average the magnitude (not orientation) of the magnetic moment to zero would involve excitations across the band gap, so the time scale for such fluctuations is slow in relation to the relevant electron energy scale. Thus the magnitude of the on-site moments $|\mu_i|^2$ in gapped PM systems will not be zero, deciding the higher energy scale of the problem. As a result, the time average of the gaps of all configurations could be nonzero. Even though the average $\langle P \rangle = \sum_n c_n P(\sigma_i^{(n)})$ over the ensemble of microscopic spin configurations is involved, the band gaps and moments $P(\sigma_i^{(n)})$ of individual configurations are nonzero.

Indeed, the fact that the antiferromagnetic ordering disappears at $T_N$ does not mean that the magnetic moments at the transition metal sites go to zero too. On the contrary, the transition metal atoms retain robust local magnetic moments as evidenced by the Curie-Weiss behavior of the magnetic susceptibility as a function of the temperature. This means that the symmetry between spin-up and spin-down electronic states is instantaneously broken at each site thus allowing the localized magnetic moments to form. In contrast, in the naive DFT nonmagnetic approximation to the PM spin configuration [4,17–19], one imposes perfect symmetry between spin up and spin down, which is the symmetry that one obtains via time averages. However, properties such as band gaps calculated by using this zero average magnetic configuration does not correspond to the property calculated as an average of the properties of the spin configurations that the system traverses. A possible way to allow for a polymorphous description of the PM phase that permits the development of local moments in a variational calculation is the special quasirandom structure (SQS) construct.

C. The special quasirandom structure (SQS) as a finite supercell realization of a polymorphous paramagnet, not a snapshot configuration

Let us focus on the band gap as the property $P$ to be calculated. Instead of calculating the band gap for many snapshot configurations $[\sigma_i^{(n)}]$ and averaging the corresponding band gaps $P(\sigma_i^{(n)})$, we construct a single supercell of $N$ sites that approximates the polymorphous configurational average. This is done by requiring that the pair and multibody atom-atom correlation functions in this special $N$ site cell best match the analytically known correlation functions for the infinite, perfectly random configuration [45,46]. Convergence with respect to $N$ must be examined; we use $N \leq 216$ atoms/cell finding that the moments and the total energy have stabilized. The SQS fully complies with the polymorphous description of the PM phases that we want to apply here. An observable $P$ calculated for such a structure is not simply the property of a single snapshot configuration but approximates the ensemble average $\langle P \rangle$ for the random configuration (see Sec. A in the Supplemental Material, see Ref. [47], for the SQS construct [48–50] and the explanation of how an SQS approximates the ensemble average for a random system).

It is clear that describing random alloys by periodic structures will introduce spurious spatial correlations in the moment configuration beyond a certain distance ("periodicity errors"). However, many physical properties of solids are characterized by microscopic length scales that can be ordered according to their typical size so as to establish a hierarchy. For instance, interactions between distant neighbors generally contribute less to the total energy than do interactions between close neighbors. Therefore the guiding idea in the construction of special quasirandom structures is to obtain within such structures a close reproduction of the perfectly random atom arrangement for the first few shells around a given site, while the periodicity errors originate from the arrangement of the more distant neighbors. In this respect, the SQS construct is reminiscent of the “special k points” used for Brillouin zone integration [51,52] in the sense that the selected k points are not meant to reproduce properties that reflect mostly the long-range order. The accuracy of the SQS improves as one uses larger SQS cell representation (analogous to using more k points in BZ sampling methods) in which longer range correlation functions can be matched. We have used 64-atom $(2 \times 2 \times 2$ primitive cells) and 216-atom $(3 \times 3 \times 3$ primitive cells) SQSs.

The SQS, as we just pointed out, is a convenient computational tool to approximate ensemble averages. It has been shown that relatively small SQS produce numerically the same property values as well as larger (ergodic) randomly selected supercells do (see Ref. [53]). Note, however, that the SQS approach is not to be confused with the commonly practiced supercell approach. In the supercell approach, one occupies lattice sites by different spins using, say, a random statistics (i.e., via coin toss) or some choice of short-range order. However, each such occupation pattern corresponds to a single snapshot and in order to calculate the observable property $\langle P \rangle$, which is an ensemble average, one should average the properties $\{P_i\}$ of different supercell snapshots. In the SQS approach, the property $P_{\text{SQS}}$ calculated for one SQS provides an approximation of the average $\langle P \rangle$, which is progressively improved by increasing the size $N$ of the SQS and by extending the order and size of the figures that the SQS algorithm tries to hierarchically match. Because the SQS is a polymorphous approach, it allows chemically identical sites to develop their own, energy-lowering displacement patterns. In the transition metal monoxides investigated here, the minimization of the total energy for the PM phases shows negligible positional atomic displacements relative to the rock-salt positions (less than 0.07 Å in amplitude). Examples of previous works that use the SQS construct to model magnetic disorder are Refs. [54,55] in which magnetic SQSs were used to model UO$_2$ and CrN, respectively, finding encouraging results.

Figure 1 shows the SQS we use for the random PM phase. The histogram in Fig. 1 illustrates that, while in the AFM phase (formed by doubling of the primitive rock-salt cell) each metal atom has six spin-up and six spin-down metal neighbors [denoted by (6,6)], in the SQS representation of the high-temperature PM phase there is a distribution of local environments, e.g., (4,8), (6,6), (8,4), etc. The landscape of the self-consistent DFT potential $V[\rho(r),m(r)]$ corresponding to the SQS, in effect, allows each metal site to experience its own distinct “particle-in-a-box” type potential, simply because chemically identical metal sites that have different neighbors
number calculated averaging over a 216-atom SQS with distribution. The frequency of local environment types In the AFM phase, the metal sublattice of the underlying rock-salt structure is shown). by the 216-atom rock-salt SQS shown in the insert (in this model only cation site in (a) the AFM-II phase and (b) the PM phase modeled here.

varies between 0 and 12 and $\uparrow$$^3$NaCl with one TM/cell and nonmagnetic).

Within the SQS construct, we consider four types of fluctuations introduced by broken symmetry (with respect to NaCl with one TM/cell and nonmagnetic).

(a) Geometric fluctuation: different local geometrical environments for chemically identical TM atoms in the lattice. Specifically, a different number of spin-up vs spin-down sites can exist around each TM site, see Fig. 1. Here, the 12 (next-nearest) neighbor TM atoms to a central TM atom can have 6 up spin +6 down spin, or 4 up +8 down, etc.). This is afforded by the SQS construct, treating the PM phase as a spin alloy. The system is in PM phase (total magnetic moment equal to zero).

(b) Occupation number fluctuations whereby atomic sites with partial occupation of initially degenerate levels (e.g., two electrons in the threefold degenerate $t_2$ level) can have different assignments of the electrons to the degenerate partners. Occupation broken symmetry means that for every cobalt atom, the occupations on different $d$ states are always integer [such as (1,0,1) in two electron $t_2$, or (1,0) in single electron $e$ level, rather than using fractional and equal occupation such as (2/3,2/3,2/3) in the $t^2$ case, or (1/2,1/2) in the $e^1$ case. In a charge self-consistent DFT calculation, the fluctuations (a) and (b) could lead also to further energy lowering via the following mechanisms: (c) Site-to-site local magnetic moment fluctuations and (d) displacement fluctuations (i.e., atomic relaxation). The key point is that in the polymorphous approach used here all such symmetry breaking mechanisms—geometric [(a) and (d)] or non-geometric [(b) and (c)]—are considered simultaneously as long as they lower the energy. On the technical side, we note that to ensure that the occupation pattern does not correspond to a local minimum, one initially applies a “nudge” in the occupation matrix (see Refs. [54,56–58]) and then proceeds with the charge self-consistent DFT calculation. Section B in the Supplemental Material (Ref. [47]) describes the protocol used for nudging the site occupations and site relaxations.

E. The role of $U$ in DFT + $U$

To treat open shell systems with degenerate orbitals (such as one or two electrons in a triply degenerate $t_2$ level), one requires in DFT an exchange correlation (XC) functional that distinguishes occupied from unoccupied orbitals (so one or two of the degenerate $t_2$ partners will be occupied by integer electron(s) and the other will be kept empty, rather than occupying all partners by fractional electrons). This requirement means that the XC functional contains an exchange correlation derivative discontinuity, i.e., that the functional should belong to rung 4 or 5 of the DFT hierarchy: meta-GGAs (like SCAN) are “semilocal” functionals of the noninteracting density matrix, and belong to rung 3. In contrast, DFT + $U$, hybrid functionals and self-interaction corrected (SIC) functionals are all nonlocal functionals of the noninteracting density matrix and they are classified as rung 4. RPA, which is a nonlocal functional of all occupied and unoccupied orbitals and of their orbital energies, belongs to rung 5. The derivative discontinuity of the exchange correlation energy is missing in the first three rungs of DFT functionals, but present in the fourth and fifth rungs.

We use here the simplest nonlocal XC, i.e., DFT + $U$, a method that can be seen as an approximation to the more rigorous self-interaction correction. We used the PBE + $U$ nonlocal approximation to the exchange and correlation functional; for simplicity, we use a constant value of $U - J = 5.0 \text{eV}$ (where $U - J$ is the parameter in the DFT + $U$ formulation of Ref. [59]) for all materials in this study, although, most likely...
FIG. 2. Schematic of the sequence of level splittings and combinations for the \( d \) orbitals in MnO, NiO, FeO, and CoO as the exchange coupling and the crystal field of the symmetry appropriate to each phase are progressively imposed: (a) splitting of the \( d \) orbitals into the transition-metal atoms subjected to exchange coupling. (b) Splitting of the spin-up and spin-down \( d \) levels subjected to a cubic \( O_h \) crystal field: this is the case of a rock-salt structure with hypothetical \( \bar{Fm}3m \) magnetic ordering. (c) Splitting of the \( d \) level in the \( D_{3d} \) crystal field in the distorted rock-salt lattice with the rhombohedral \( \bar{R}3m \) AFM-II magnetic ordering. (d) Splitting of the \( d \) orbitals in a tetragonal crystal field as in the monoclinic \( C2/m \) phases of FeO and CoO. Note that in this schematic we emphasize the effect of the relevant interaction in progressively removing the degeneracy of the \( d \) orbitals, while we do not intend to reproduce to scale the position of the spin-up and spin-down energy levels. The reader should inspect the projected DOSs of Figs. 3 and 4 to obtain the actual calculated energy position of the \( d \) bands.

one can improve agreement with experiment by tweaking \( U \) separately for each compound. In the DFT + \( U \) method, the DFT total-energy functional is corrected by two terms (Refs. [20–24,59–63]). The first term is a mean-field approximation of the electron-electron interaction within a subset of localized orbitals (here, the \( d \) orbitals). The second term subtracts the contribution of the electron-electron interaction already accounted for in the approximate functional and largely consists of the self-Coulomb interaction (a manifestly one body effect).

Despite the impression suggested by the letter (Hubbard) “\( U \),” the DFT + \( U \) method (perhaps better renamed DFT + \( V \) to avoid such confusion) does not imply correlation in the Hubbard Hamiltonian sense. We note that numerous practitioners apply these methods with the belief that they model many-body correlations. DFT + \( U \), as well as the hybrid functionals and SIC DFT, are all methods in which the wave function of the \( N \)-electron system is a single determinant. In single-determinant, band structure approaches, each band structure calculation occupies its levels in a single specific manner by electrons (a single Slater determinant). Different possible patterns of occupation of levels by electrons (which can be built in separate band structure calculations) have no way of seeing each other.

F. The role of spin disorder versus spin order

We note from Table I that the FM spin arrangement is monomorphous and has a gap due to its long-range order, but it is obviously not a good model for the paramagnetic phase that is magnetically disordered. In a disordered phase, we allow for identical atoms to have the opportunity to experience different local structural environments. The exchange and crystal-field interactions modeled by DFT applied to SQSs are local effects that do not need a long-range magnetic ordering to mix and split the \( d \) levels, and as such these local effects drive the opening of a gap in the overall magnetically disordered phase. Furthermore, the SQS PM phase has lower energy than the FM phase by 59.5, 45.1, 67.6, and 79.0 meV/formula unit for NiO, MnO, FeO, and CoO, respectively (see Table I), so the FM description is not selected.

Table I shows that the internal energy at \( T = 0 \) of the spin disordered PM phase is higher than that of the AFM phase at \( T = 0 \), as expected. We clarify that for the spin disorder PM phase the free energy is \( F = E - TS \), where \( S \) is the entropy, so at \( T = 0 \), the AFM phase is lower in energy, and as \( T \) grows the free energy \( E - TS \) of the PM phase decreases until this becomes the lowest energy phase. However, we do not aim to calculate the Néel temperature. Such calculations within DFT are known in the literature (e.g., Franceschetti et al. [63] and Daene et al. [64]).

IV. SUMMARY OF THE MAIN RESULTS ON THE PM PHASES

Before we discuss the physical picture that emerges, we state the results obtained for the magnetically ordered AFM and FM phases and for the magnetically disordered PM phases, which we modeled by a polymorphous description. As we are not interested here in fitting the calculated gaps and
FIG. 3. Projected density of states (PDOS) on the transition metal $s$ and $d$ orbitals ($t_{2g}$ and $e_g$ components) calculated by DFT $+$ $U$ ($U = 5$ eV) for MnO and NiO in (a) and (b) the AFM phase with fully relaxed $R3m$ structures, and [(c) and (d)] the PM phase modeled by a cubic 64-atom $2 \times 2 \times 2$ SQS. The lattice parameters of the SQSs are set so that the volume per formula unit is equal to the calculated volume per formula unit of the DFT $+$ $U$ relaxed $R3m$ structures of MnO and NiO.

V. ANALYSIS OF THE RESULTS

A. Analysis of the occupations of the localized orbitals

For our analysis of the DFT $+$ $U$ results, we sought linear combinations of the $d$ orbitals that form a good representation of the point-group symmetry at the TM sites. The $d$-orbital occupation matrix that enters the “$+U$” term of the DFT $+$ $U$ energy functional is calculated using the $t_{2g}$ and $e_g$ orbitals as basis. However, the actual magnetocrystalline order in the AFM phases, or the lack of it in the PM phases, breaks the cubic point-group symmetry at the TM sites. In such a case, a good representation for the $d$ orbitals, which is often referred to as the “crystal-field representation,” is that defined by the eigenvectors of the occupations matrices. This representation is also meaningful in terms of the mechanism that drives the band gap opening. The sum of the probability distributions of the eigenvector functions with spin down each weighted by its occupation gives the distribution of the minority-spin electrons density around the transition metal sites that we inspect in the following. See Sec. B.1 in Ref. [47] for more details on this representation.

B. Making sure that the electronic structure DFT description does not get trapped in a high-symmetry basin

On the technical side, one needs to assure that the self-consistent procedure for the charge density optimization and relaxation of the atom positions does not get trapped in a high-symmetry solution but can explore a broad range of positional as well as wave function symmetries. In the case of systems in which the crystal-field produces degenerate states that are partially filled, one must explore lowered
FIG. 4. Projected density of states (PDOS) on the transition metal $s$ and $d$ orbitals ($t_{2g}$ and $e_g$ components) calculated by DFT + U ($U = 5\text{ eV}$) for FeO and CoO, in (a) and (b) the AFM phases with the fully relaxed monoclinic $C2/m$ structures, and in (c) and (d) the paramagnetic phases modeled by a cubic $64\times2\times2$ SQS. The lattice parameters of the SQSs are set so that the volume per formula unit is equal to the calculated volume per formula unit of the DFT + U relaxed $C2/m$ structures of FeO and CoO.

symmetries of the electronic state by allowing for distortions of the lattice [23,24,65]. Thus we permit an initial “nudging” of the atoms off the high-symmetry sites (and see if the quantum-mechanical forces tend to restore such high-symmetry positions or prefer Jahn-Teller-like displacements). At the same time, one needs to assure that the electronic self-consistency cycle could explore a broader range of wave function symmetries without getting trapped in high-symmetry solutions. To this end we avoid charge density symmetrization during the electronic self-consistent iterations. In the case of the PM phases, we “nudge” the systems initially with unequal $d$ orbital occupations. See Sec. B 2 in the Supplemental Material, Ref. [47] for the details of the nudging protocol. Starting from such an orbital configuration helps the self-consistent solver to converge towards a solution in which the $d$ orbitals mix to form linear combinations whose occupations ultimately are either close to one or zero.

VI. RESULTS: THE MAGNETICALLY ORDERED AFM PHASES

The AFM phases of MnO, FeO, CoO, and NiO have been studied by DFT [67–69] as well as its extensions and corrections, including DFT + U [23,66], hybrid functionals [70], and SIC [71]. Here we briefly describe our results of the evolution of the band gaps (Fig. 2) and provide the density of states [Figs. 3(a), 3(b), 4(a), and 4(b)] to establish a common basis for discussing later the generalized supercells needed to capture the physics of the PM phases. As shown in Fig. 2, in the ideal cubic rock-salt structure ($Fm\bar{3}m$ space group), the crystal-field splits the atomic $d$ levels into spin-up and spin-down $t_{2g}$ and $e_g$ levels. The AFM-II magnetic ordering already breaks the cubic space-group magnetocrystalline symmetry even without distortions to the ideal cubic lattice. The lattice relaxations that are experimentally observed in the low-temperature phases of these monoxides [72–74], lower the point-group symmetry of the crystal field at the TM sites with respect to that of the ideal cubic structure. See Sec. C in Ref. [47] for the details of the relaxed crystal structure of the AFM phases that we obtained by our DFT + U calculations. Qualitatively, a similar combined effect of the exchange and crystal-field interaction is the mechanism that drives the gap opening in the PM phases and provides a unifying, single-particle description of the insulating character of both the magnetically ordered and magnetically disordered phases. We, therefore, illustrate this mechanism starting with the AFM phases, as well as the hypothetical FM phases, through the same protocol used for the PM phases.

A. AFM MnO and NiO

AFM MnO and NiO exhibit a $R\bar{3}m$ magnetocrystalline structure in which the TM crystal field has the rhombohedral
$D_{3d}$ point-group symmetry. A crystal field of this symmetry mixes the $t_{2g}$ orbitals so as to give the $a_{1g}$ singlet and the $e'$ doublet [Fig. 2(c)]. The $e'$ orbitals are invariant to the $D_{3d}$ symmetry operations and are often indicated as $e''$. Figures 3(a) and 3(b) depict the DOS of MnO and NiO projected onto the cubic $t_{2g}$ and $e_g$ orbitals. In MnO, the Mn$^{2+}$ ions exhibit the $d^5$ electronic configuration which results into the five spin-up $d^5$ being orbitals fully occupied. A band gap opens in MnO between the fully occupied spin-up $d^5$ orbitals and the empty spin-down $d^5$ orbitals. In NiO, the Ni$^{2+}$ ions exhibit the $d^8$ configuration which corresponds to the five spin-up $d^5$ orbitals being fully occupied and the $t_{2g}^2$-derived orbitals being also fully occupied. A band gap opens in NiO between the occupied $t_{2g}^2$-derived levels and the empty $e_g^2$-derived levels.

**B. AFM FeO and CoO**

Fe$^{2+}$ and Co$^{2+}$ in FeO and CoO are, respectively, in the $d^8$ (meaning one electron in the spin-down $d$ states) and $d^7$ (meaning two electrons in the spin-down $d$ states) configurations. Our DFT $+ U$ calculations, in line with earlier studies [23,24,66], show that a gap opens in FeO and CoO already in the undistorted cubic lattice because of the symmetry lowering induced by the AFM-II ordering. FeO opens a gap by occupying the $a_{1g}$ singlet, while the $e_g'$ doublet is in the conduction. The opposite occurs in CoO with the $e_g'$ doublet occupied by two electrons and the $a_{1g}$ singlet in the conduction. FeO [75] and CoO [73,74] lower their total energies with respect to the ideal cubic lattice through tetragonal distortions of the metal-oxygen coordination octahedra that are accommodated within a monoclinic cell with $C2/m$ space-group symmetry (see Fig. 6 and Table A1 in the Supplemental Material, Ref. [47] for a description of the calculated equilibrium crystal structure). In FeO, there is a compression of the in-plane bonds and an expansion of the out-of-plane ones, while the opposite occurs in CoO. Figures 4(a) and 4(b) depict the DOS of monoclinic FeO and CoO projected onto the cubic $t_{2g}$ and $e_g$ orbitals. FeO and CoO continue to be insulating in the stable AFM monoclinic phase as in the higher-energy, AFM undistorted cubic phase. The orbital mixing that occurs due to the tetragonal distortion of the coordination octahedra is reflected by the shape of the spin-down electron density $\rho^\downarrow(r)$. In FeO, $\rho^\downarrow(r)$ has a square-planar shape rotated by 45° degrees around the $z$ axis [see Fig. 5(a)]. In CoO, $\rho^\downarrow(r)$ has an octahedral shape with the vertical axis lying along the diagonal of the $x-y$ plane [see Fig. 5(b)]. A detailed analysis of $\rho^\downarrow(r)$ in terms of the orbital mixing obtained in the DFT $+ U$ solution is performed in Sec. D1 in the Supplemental Material, Ref. [47].

**VII. RESULTS: THE MAGNETICALLY DISORDERED PARAMAGNETIC PHASES**

The DFT $+ U$ calculations of the PM phases of the four monoxides modeled with the magnetic SQS produce insulating solutions with strong magnetic moments at the TM sites (see Table I for the gaps and magnetic moments at the transition metal sites obtained in these SQS calculations). The minimization of the total energy for the PM phases of NiO, MnO, and FeO, modeled by the SQS shows negligible positional atomic displacements relative to the rock-salt positions, thus, no significant broadening of Bragg diffraction peaks is expected. The projected DOS (PDOS) on the metal $d$ orbitals are depicted in Figs. 3(c) and 3(d) for MnO and NiO and Figs. 4(c) and 4(d) for FeO and CoO. For the sake of comparing the PDOSs across the whole series of oxides and magnetic phases included in this study, we project the DFT $+ U$ wave functions on $t_{2g}$ and $e_g$ orbitals. However, in the polymorphous description, which is implemented through the magnetic SQSs, the crystal field at each TM site shows a low point-group symmetry, which in turn allows for the $t_{2g}$ orbitals to mix among themselves and possibly also with the $e_g$ orbitals. The magnetic disordered phases of MnO and NiO exhibit gaps that in both cases, as can be seen from the PDOSs in Figs. 3(c) and 3(d), open between subbands that derive predominantly from the $t_{2g}$ and $e_g$ orbitals and are both filled in a similar fashion as in the magnetically ordered AFM phases. The PDOS plots of FeO and CoO in Figs. 4(c) and 4(d), respectively, show that the gap originates in both systems mainly from a splitting in the $t_{2g}$-derived states. An inspection of the density $\rho^\downarrow(r)$ of spin-down electrons of PM FeO and CoO [see Figs. 6(a) and 6(b)] and of the eigenvectors of the occupation matrices calculated at the TM sites (see Sec. D2 in the Supplemental Material, Ref. [47]) shows that the gap is the result of the mixing of the $d$ orbitals in the low-symmetry crystal field that characterizes each site in the disordered
phases and of the splitting of these mixed orbitals that lifts the degeneracy of the unperturbed $d$ states. In PM FeO, $\rho^i(r)$ at the Fe sites [Fig. 6(a)] has a square-planar shape lying on one of the Cartesian axes and it is not tilted by 45° around a Cartesian axis as in AFM FeO [see Fig. 5(a) and 5(c)] for a comparison between $\rho^i(r)$ in AFM FeO and PM FeO, with the specific plane varying randomly from site to site. Sites 4 and 14 in Fig. 6(a) are examples of two distinct orientations of $\rho^i(r)$ around the Fe sites in PM FeO. The analysis of the eigenvectors of the occupation matrices in Sec. D2 in the Supplemental Material, Ref. [47] shows that this square planar shape originates from one $t_2g$ orbital almost completely filled.

In PM CoO, we observe [see Fig. 6(b)] that $\rho^i(r)$ shows two types of shapes at the Co sites: a cylinderlike shape [e.g., at site 14 in Fig. 6(b)] aligned to one of the Cartesian planes and points into a direction that varies randomly from site to site. Figures 5(b) and 5(d) display the Cartesian planes and points into a direction that varies from site to site. Figures 5(b) and 5(d) display $\rho^i(r)$ at the Co sites in AFM CoO and PM CoO, respectively, and show the different orientation of the octahedral shape in the two cases. From an inspection of the eigenvalues of the occupation matrices at the Co sites within the SQS (see Sec. D2 in the Supplemental Material, Ref. [47]), we observe that at each site two eigenvectors $\phi^{(1)}_{Co}$ and $\phi^{(2)}_{Co}$ have occupations of $\sim 0.9$, that is, are almost completely full and make for the dominant contribution to $\rho^i(r)$ at the Co sites. The two different types of the shape of $\rho^i(r)$, found, for example, at sites 4 and 14 in Fig. 6(b), originate from two distinct modes of mixing of the $d$ orbitals to which correspond two distinct pairs of nearly fully occupied eigenvectors $\{\phi^{(1)}_{Co}, \phi^{(2)}_{Co}\}$. At the sites where $\rho^i(r)$ shows a cylindrical shape [e.g., site 14 in Fig. 6(b)], the nearly fully occupied eigenvectors $\{\phi^{(1)}_{Co}, \phi^{(2)}_{Co}\}$ are two of the three $t_2g$ orbitals. At sites where $\rho^i(r)$ shows a tilted octahedral shape, the eigenvectors $\{\phi^{(1)}_{Co}, \phi^{(2)}_{Co}\}$ are normalized linear combinations of the three $t_2g$ orbitals with contributions also from the $e_g$ orbitals. Analytic models of the probability densities $|\phi^{(1)}_{Co}|^2$ and $|\phi^{(2)}_{Co}|^2$ and of their sum $|\phi^{(1)}_{Co}|^2 + |\phi^{(2)}_{Co}|^2$ are displayed in Fig. A3 in the Supplemental Material; these models are consistent with the tilted octahedral and the cylindrical shape of $\rho^i(r)$ at the Co sites in PM CoO obtained in the DFT $+ U$ solution.

The modality of orbital mixing and level splitting revealed by the present calculations is similar in the AFM and PM phases of the respective oxides. This helps explain the fact that the magnetic moments in the SQS PM configurations converge to values whose average is within less than 1% of the AFM values (Table I). At the same time, the magnetic disorder decreases the band gap of MnO, NiO, and CoO with respect to the value in the AFM phase.

**VIII. DISCUSSION**

**A. The physical picture of gapping of the PM phase:**

The role of different symmetry-breaking modes in the PM phase

We considered the following degrees of freedom that induce the symmetry breaking with respect to nonmagnetic NaCl-structure with one TM per primitive cell: (a) geometric symmetry breaking via the local environment distribution supplied by SQS; (b) occupation number broken symmetry (OBS); (c) displacement fluctuations (i.e., atomic relaxation); (d) site-to-site local magnetic moment fluctuations. In CoO, we illustrate the effects of these fundamental fluctuations in Fig. 7 by starting from the most complete calculation ("level IV" below) and "peeling of the onion" (i.e., removing the effects one by one), inspecting total energies, band gaps, local moments, and atomic displacements for the following 4 levels.

**Level IV theory: SQS + OBS, with relaxation; allows effects (a), (b), (c), and (d).** Figure 7(a) shows the projected density of states as well as the total energy, band gap, and average cobalt local magnetic moment for level IV of the 216-atom rock-salt CoO supercell. It represents a very large energy lowering of $-2219$ meV/formula units relative to the monomorphous nonmagnetic description (level I below) and yields a band gap of 2.39 eV and local moment of 2.75 BM. Some of its properties include the following.

(i) Although the $t_2g$ and $e_g$ representations are mixed by the various symmetry breaking channels, one can observe that the hole is localized predominantly on the $t_{2g}$ derived state rather than the $e_g$ state: the occupation broken symmetry is manifested by the fact that two electrons occupying the $T_1$, $T_2$, and $T_3$ components in the spin-down $t_g$ band cause the band to split into the fully occupied (by two electrons) VBM and an empty $t_g$ (CBM) derived spin-down conduction band. Together with the three electrons in the deeper spin-up $t_g$ band and two electrons in the spin-up $e_g$ band, this leads to the $t^{2}_{g}e^{2}_{g}t^{2}_{g}$ configuration akin to the $^{3}T_{1g}$ multiplet (spin 3/2) where the hole is distributed in the $t_{2g}$-like band. Placing instead the hole in the $e_g$-like band leads to a higher energy $^{3}E_{g}$ multiplet.

(ii) The atomic displacements are configuration-dependent and average to small amplitudes. Relaxation causes small atomic displacements, with displacement directions depending on the configuration selected, so the orientation average of the displacements is far smaller than in any particular configurations.

(iii) The band edges do not look like upper and lower Hubbard states but are oxygen derived. The orbital make up of the band edges [Table II and Fig. 7(a)] shows that the
FIG. 7. Analysis of the effects of different symmetry breaking modes on the electronic structure, total energy, band gap, and local moment of paramagnetic CoO. We show the projected density of states (PDOS) on the transition-metal $s$ and $d$ orbitals ($t_{2g}$ and $e_g$ components) and oxygen $p$ orbitals from levels (a) IV, (b) III, (c) II, and (d) I (see Sec. VII A for definition of these symmetry breaking levels) of rock-salt CoO. (a)–(c) are calculated in a 216-atom cubic rock-salt supercell with SQS, while (d) is calculated in a two-atom rock-salt primitive cell. All lattice parameters are the same as the one used in Fig. 4(d).

VBM is made predominantly of oxygen $p$ orbitals (down-spin VBM) and roughly equal amounts of oxygen and cobalt $d$ (up-spin VBM) whereas the lowest CBM is predominantly oxygen $p$ with some cobalt $d$ hybridization. Thus CoO (as well as the other oxides discussed here with late transition metal cations having rather deep 3$d$ orbitals) does not have the proverbial lower Hubbard $d$ VBM and upper Hubbard $d$ CBM, but are in fact charge transfer type band edges more similar to ordinary semiconductors such as ZnO rather than to early transition metal oxides (of V and Ti) that have more significant $d$ character in at least one of the two band edges.

Level III theory: SQS + OBS but no relaxation; allows effects (a), (b), and (d). This level of theory includes different local environments (via SQS) in the 216-atom rock-salt CoO supercell, but forces equal occupation of the members of a degenerate $t_{2g}$-like spin-down state with components $T_1$, $T_2$, and $T_3$, i.e., instead of placing two electrons in two of the partners and zero in the third, this central field like approximation places $(2/3, 2/3, 2/3)$ electrons in each partner. This places the Fermi level inside the down-spin $t_{2g}$ band leading to a metal. Figure 7(c) shows the density of states and total energy revealing a dramatic increase (494 meV/f.u.) in total energy is rather unchanged. By comparing the occupation matrices, we also find that in the unrelaxed solution the $e_g$ states do not mix with the $t_{2g}$ states, e.g., the spin-down valence states are a superposition of the type, e.g., $dxy$, $dyz$. After relaxation the valence band remained basically pure $t_{2g}$ with little contamination from the $e_g$ states.

Level II theory: SQS, no OBS (freezing equivalent occupation on each $t$-like state), no relaxation; allows effects (a) and (d). This level of theory includes different local environments (via SQS) in the 216-atom rock-salt CoO supercell, but forces equal occupation of the members of a degenerate $t_{2g}$-like spin-down state with components $T_1$, $T_2$, and $T_3$, i.e., instead of placing two electrons in two of the partners and zero in the third, this central field like approximation places $(2/3, 2/3, 2/3)$ electrons in each partner. This places the Fermi level inside the down-spin $t_{2g}$ band leading to a metal. Figure 7(d) shows the density of states and total energy revealing a dramatic increase (494 meV/f.u.) in total energy...
TABLE II. Energies (in eV) and orbital make ups of band edges from DFT $+ U \ (U = 5 \text{ eV})$ calculations for (a) level IV (SQS + OBS with atomic relaxations) and (b) level III (SQS + OBS but no atomic relaxations) using the 216-atom rock-salt CoO supercell. Level IV started initially from level III calculation. The occupation configuration given by level III in this table is also shown as pattern 1 in Table III.

<table>
<thead>
<tr>
<th>(a) CoO level IV</th>
<th>up-spin VBM</th>
<th>down-spin VBM</th>
<th>up-spin CBM</th>
<th>down-spin CBM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Energy (eV) ($E_{\text{Fermi}} = 0$)</td>
<td>-0.022</td>
<td>-0.022</td>
<td>2.367</td>
<td>2.453</td>
</tr>
<tr>
<td>Co $s$</td>
<td>0.64%</td>
<td>0.31%</td>
<td>9.00%</td>
<td>0.41%</td>
</tr>
<tr>
<td>Co $d_{eg}$</td>
<td>52.12%</td>
<td>4.47%</td>
<td>2.80%</td>
<td>2.41%</td>
</tr>
<tr>
<td>Co $d_{tg}$</td>
<td>1.96%</td>
<td>25.63%</td>
<td>20.78%</td>
<td>95.14%</td>
</tr>
<tr>
<td>O $p$</td>
<td>45.28%</td>
<td>69.59%</td>
<td>67.44%</td>
<td>2.04%</td>
</tr>
</tbody>
</table>

(b) CoO level III

| Energy (eV) ($E_{\text{Fermi}} = 0$) | -0.022 | -0.022 | 2.220 | 2.393 |
| Co $s$ | 0.74% | 0.32% | 6.66% | 0.31% |
| Co $d_{eg}$ | 51.68% | 4.96% | 1.94% | 1.50% |
| Co $d_{tg}$ | 1.93% | 24.92% | 23.86% | 96.15% |
| O $p$ | 45.65% | 69.80% | 67.55% | 2.04% |

relative to level III, just 1706 meV/formula unit below the level I. Despite the gap being zero, this approximation gives a nonzero local magnetic moment.

Level I theory: no SQS (nonmagnetic monomorphous description), no OBS, no relaxation; allows no symmetry breaking effects. This lowest level, often used in the past as the DFT rendering of paramagnets, involves a primitive NaCl-structure unit cell (1 Co + 1 O). There is no spin polarization, so the two spin channels (with plus and minus values) are exactly the same. The total energy is very high relative to all other theoretical levels; the band gap and moments vanish, and the Fermi energy lies in an $e_g$-like band instead of the correct $t_g$-like band as in the other levels of theory.

B. The existence of many broken symmetry configurations in the PM phase

Each independent broken symmetry configuration corresponding to a choice of occupation patterns can have, in principle, different charge density, total energy, local moments, and band gaps. The ultimate solution for the PM phase should correspond to a symmetrized combination of these configurations. Indeed, as explained in Sec. III A above, the correct way to calculate the property $P$ of a phase that can have numerous individual configurations $\{\sigma^{(n)}\}$ each with property $P(\sigma^{(n)})$, is to calculate the polymorphous statistical average $\langle P \rangle = \sum_{\sigma} c_{\sigma} P(\sigma^{(n)})$ over the ensemble of microscopic configurations. In general, the combination needs to be symmetrized, and if many-body configuration interaction is to be considered, the problem needs to be subjected to this additional diagonalization step. Table III illustrates different configurations for the 216 atom SQS of the PM phase of CoO (level III). We see that each of these configurations is closed shell and have rather similarly large band gaps and moments. According to our basic conjecture discussed in Sec. III B, we expect that the symmetrized superposition state will also have similarly large gaps and moments, i.e., that the interaction between these base configurations will be weak. The single determinant band approach used here neglects such dynamic interaction between configurations and would fail when the neglected interaction is strong. This is not the case in the currently considered compounds, which are closed shell (level IV or III) with large band gaps and are thus expected to be well represented by a single-determinant approach.

C. Mott localization and double occupation is not involved in the present physical picture

The bands obtained in the current single-determinant mean-field band structure theory are conventional single-particle states and should not be confused with the lower and upper Hubbard bands $[76,77]$ that characterize the solutions of the Hubbard model. The Mott mechanism requires that the electrons move across the lattice forming states on certain atomic sites with doubly occupied $d$ orbitals and empty $d$ orbitals on other sites (i.e., not a Bloch periodic band structure picture) while the overall charge number is conserved. These types of excited configurations correspond respectively to the upper and lower Hubbard bands, which are truly “dynamic” charge bands and correspond to many-body configurations, whereas the present approach produces gapping without such a mechanism, different than the Mott gap opening mechanism described in the textbooks. That it is not entirely surprising that $d$ electron strong correlation is not the deciding factor for these binary oxide systems as can be gleaned from Fig. 7 and Table II showing that the valence band and conduction band edges consist of oxygen $p$ and transition metal $s$ orbitals, not the proverbial $d$-like lower and upper Hubbard bands.

TABLE III. Comparison of the properties of four different configurations of SQS-PM CoO obtained by different initial occupation broken symmetries. All are closed shells. Shown are the total energies (in eV per formula unit), band gap (in eV), and average magnetic moment (in Bohr magnetons) at the transition-metal sites. Table A3 in the Supplemental Material also illustrates some typical output configurations.

<table>
<thead>
<tr>
<th>$d$ occupation pattern</th>
<th>$E_{\text{tot}}$ (eV/f.u.)</th>
<th>$E_{\text{gap}}$ (eV)</th>
<th>$\mu$ ($\mu_B$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-11.561</td>
<td>2.26</td>
<td>2.75</td>
</tr>
<tr>
<td>2</td>
<td>-11.560</td>
<td>2.37</td>
<td>2.75</td>
</tr>
<tr>
<td>3</td>
<td>-11.559</td>
<td>2.26</td>
<td>2.75</td>
</tr>
<tr>
<td>4</td>
<td>-11.560</td>
<td>2.35</td>
<td>2.75</td>
</tr>
</tbody>
</table>
D. Comparison with other approaches

Previously, the paramagnetic phases, including those of the present TM oxides, have been modeled by the disordered local moments (DLMs) \[78–80\] approach, which has been implemented within the single-site coherent-potential approximation (CPA) \[11,36,65,81\]. This approach assumes that the Schrödinger equation potential seen by chemically equivalent atoms in the disordered PM phase are all equal even though such atoms have distinctly different local environments (such as number of neighbors with spin up versus spin down, see Fig. 1). This picture automatically ignores the existence of inhomogeneous distribution of moments and charges. This is valid only when the local environment flips its spin so fast that a central atom does not distinguish if its environment is made of up spins or down spins but all can be described as some average. This is unlikely to be the case in insulators (such as Mott insulators) where the screening is ineffective. This DLM view leads to equal local moments on all TM atoms irrespective of their environments. The DLM approach is virtually equivalent to the so-called “Hubbard III” approximation \[14\] in regard to the treatment of the spin disorder. DLM and Hubbard III are in turn related to the DMFT approach, with the difference that DLM and Hubbard III are unable to describe quantum fluctuations, which are instead described by DMFT \[16\]. Similarly to the DLM description, DMFT is inherently a single-site theory in which all sites of a given species (e.g., all the Co sites in paramagnetic CoO) are geometrically equivalent.

The special quasirandom structure construct is an effective way to establish a physically grounded representation of the random magnetic configuration for three reasons. First, an SQS is constructed so that a property calculated using it is a close estimate of the ensemble statistical average that would be required to calculate that property for a fully disordered phase. Therefore, using one SQS, one can obtain reliable estimates of ensemble averages, i.e., the relevant quantities for the paramagnetic phases, by calculating one configuration instead of many randomly-generated configurations. Second, an SQS allows for a variety of local magnetic environments and for multiple patterns of uneven \(d\) orbital occupations that both concur to breaking the cubic symmetry and fully lifting the degeneracy of the \(d\) orbitals. Finally, it is straightforward to construct SQS that represent the property of imperfectly-random ensembles, i.e., those that have short-range order (SRO) and are thus better representative of PM phases closer to the Néel temperature. Instead of constructing the SQS by fitting to the analytically known random pair and many-body correlation functions (see Sec. A in the Supplemental Material), one can fit to independent measures or calculated correlation functions that incorporate SRO \[82\].

In conclusion, in the present study we find that the DFT + \(U\) method, which is a generalized Kohn-Sham approach, reproduces the insulating character and on-site magnetic moments of the prototypical Mott insulators MnO, NiO, CoO, and FeO when applied to SQSs, which approximate closely the ensemble average over the random magnetic configurations.

ACKNOWLEDGMENTS

The work of A. Z. was supported by Department of Energy, Office of Science, Basic Energy Science, MSE division under Grant No. DE-FG02-13ER46959 to CU Boulder. He would like to thank Chris Marianetti, David Singh, Sohrab Ismail Beigi, Gabi Kotliar, George Sawarzky, Igor Mazin, Lin Wang, and Olle Helleman for very interesting and extremely useful discussions. G. T. would like to thank also J. Ketterson of Northwestern University for useful discussions. The crystal structure figures displayed in this article were generated using the software VESTA \[83\]. In this research, G.T., Z.W., and A.Z. used resources of the National Energy Research Scientific Computing Center (NERSC), a US Department of Energy Office of Science User Facility, supported by the Office of Science of the US Department of Energy under Contract No. DE-AC02-05CH11231.

APPENDIX

For AFM phases, we use low-temperature lattice vector symmetry. Even in NiO and MnO, the spin symmetry alone changes the space group from \(\text{NaCl}\) to Rhombohedral \(R3\)-\(m\) [because we selected \((111)\) type AFM]. For FeO and CoO, the lowest energy AFM cell symmetry is monoclinic \(C2/m\). We then optimize the structure to achieve very small quantum mechanical forces. Note that the AFM structure can be Jahn Teller distorted (FeO, CoO, see Fig. 2).

For PM phases, we use the supercell shape as the macroscopically observed high-\(T\) cubic lattice vectors, keeping only the cell-internal atomic positions as variables during force calculations. In practice, this can be done in steps such as (i) freeze one choice at the time of OBS, and a frozen cell-internal atomic geometry, and run charge self-consistency. Do a few OBS choices independently, called “configuration i”; (ii) unfreeze the OBS choice starting from (i), while the geometry is still unrelaxed, and perform charge self-consistency; (iii) unfreeze the previously unrelaxed geometry, seeking a minimum. In practice, to avoid local minima, this requires an initial nudge in the form of small random atomic displacements, then following the calculated forces to geometries with vanishing forces.

Another protocol involves performing atomic relaxation concomitantly with steps (i) and (ii). Here in step (i’) one freezes a given choice of OBS, and performs a charge self-consistent calculation on that frozen OBS while at the same time relaxing the cell internal atomic positions. Do a few OBS choices independently called “configuration i’: (ii’) unfreeze OBS starting from (i’), while the geometry is being reoptimized and do charge self-consistency. In the present case both protocols (i) + (ii) or (i’) + (ii’) lead to virtually identical results.


