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# First-principles reinvestigation of bulk WO<sub>3</sub>

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Using first-principles calculations, we analyze the structural properties of tungsten trioxide WO<sub>3</sub>. Our calculations rely on density functional theory and the use of the B1-WC hybrid functional, which provides very good agreement with experimental data. We show that the hypothetical high-symmetry cubic reference structure combines several ferroelectric and antiferrodistortive (antipolar cation motions, rotations, and tilts of oxygen octahedra) structural instabilities. Although the ferroelectric instability is the largest, the instability related to antipolar W motions combines with those associated to oxygen rotations and tilts to produce the biggest energy reduction, yielding a  $P2_1/c$  ground state. This nonpolar  $P2_1/c$  phase is only different from the experimentally reported Pc ground state by the absence of a very tiny additional ferroelectric distortion. The calculations performed on a stoichiometric compound so suggest that the low-temperature phase of WO<sub>3</sub> is not intrinsically ferroelectric and that the experimentally observed ferroelectric character might arise from extrinsic defects such as oxygen vacancies. Independently, we also identify never observed R3m and R3c ferroelectric metastable phases with large polarizations and low energies close to the  $P2_1/c$  ground state, which makes WO<sub>3</sub> a potential antiferroelectric material. The relative stability of various phases is discussed in terms of the anharmonic couplings between different structural distortions, highlighting a very complex interplay.

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### I. INTRODUCTION

Tungsten trioxide, WO<sub>3</sub>, has been extensively studied due to its very attractive and rich properties for technological applications. WO<sub>3</sub> and its derivatives  $H_x$ WO<sub>3</sub> and WO<sub>3-x</sub> are electrochromic [1–6], thermochromic [7,8], and superconducting [3,8–14]. It has been envisaged that WO<sub>3</sub> may become one of the best materials for electrochromic applications such as in energy-efficient windows, smart glasses, antiglare automobile rear-view mirrors, sunroofs, displays, or even tunable photonic crystals [15] and to reduce photocorrosion [16]. The wide variety of the underlying electronic instabilities for these properties is mirrored by a multitude of related structural instabilities, which have been investigated ever since 1975 [17,18] and refined later [19–24].

All known WO<sub>3</sub> phases are characterized by very large distortions of the archetypal perovskite structure so that even the notion of octahedra tilts is to be taken with some caution. The WO<sub>6</sub> octahedra are so largely distorted that the variance of W–O distances in any observed structure is far greater than in most other known perovskite structures [25–29]. In this paper we make the fundamental connection between the electronic and structural structure properties of WO<sub>3</sub> and show that most, if not all, structural instabilities can be derived from a careful analysis of its intrinsic cubic phonon instabilities, despite these large deformation amplitudes.

The structural properties can be summarized as follows:  $WO_3$  shows no proper melt; surface melting of crystalline material occurs at 1746 K. Crystal growth proceeds typically by sublimation and gas transport at temperatures below 1400 K. At the highest temperatures the structure is tetragonal (space group P4/nmm) with strong antiferrodistortive cation movements so that the  $WO_6$  octahedra are strongly distorted [30] in an antipolar pattern. Additional rotational octahedral distortions condense in addition to the initial tetragonal displacements when lowering the temperature. They

further reduce the symmetry from tetragonal to orthorhombic, 57 monoclinic, triclinic, and finally to a second monoclinic phase. 58 A structural sequence, which contains phases stabilized by 59 temperature, is given in Fig. 1. A monoclinic phase  $(P2_1/n)$  60 [21] and a triclinic phase  $P\bar{1}$  exist at room temperature [25,26]. At higher temperatures, Vogt et~al. [22] and Locherer 62 et~al. [19] concluded a transition from Pbcn to the P4/ncc 63 phase and Howard et~al. [23] observed an intermediate  $P2_1/c$  64 phase. Locherer et~al. [19] and Woodward et~al. [26] found 65 an additional transition from P4/ncc to P4/nmm at 980 to 66 1200 K. Below room temperature, Salje et~al. [20] reported 67 a transition from the triclinic  $P\bar{1}$  phase to a polar phase (Pc) 68 with no further transitions down to 5 K.

WO<sub>3</sub> occurs (almost) always as oxygen deficient WO<sub>3-x</sub> 70 with a metal-insulator transition to a metallic phase for high 71 concentrations of oxygen vacancies or doping with alkali 72 metals. Superconductivity occurs in the metallic phase [9] 73 even if the reduced regions are restricted to nanoscale twin 74 boundaries. Bulk superconductivity in WO<sub>3-x</sub> was found 75 in a tetragonal phase with space group  $P\bar{4}2_1m$  [10]. (Bi-) 76 polaronic electron transport is a characteristic property of 77 WO<sub>3-x</sub> [31–34].

Numerous first-principles studies were performed on WO<sub>3</sub> 79 in order to characterize its electronic structure (bulk, thin films, 80 and cluster phases [35–41]), the role of oxygen vacancies 81 [42–45], and cation doping [46–53]. In this paper we do not 82 only focus on the electronic structure but also extensively 81 study the structural stabilities and metastabilities of the 84 various phases. We show that the hybrid functional B1-WC 85 is preferable for the study of the electronic and structural 86 properties of WO<sub>3</sub> over previous approaches. In the first 87 section we check the validity of the B1-WC functional against 88 six known crystallographic phases. The sole disagreement 89 exists for the crystallographic  $\varepsilon$  phase (Pc), which yields 80 a higher symmetric  $P2_1/c$  structure. In the second section 91 we analyze metastable phases starting from the phonon 92

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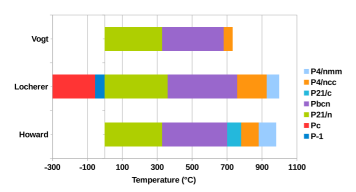


FIG. 1. Schematic summary of the temperature phase diagram WO<sub>3</sub> as reported by three main experimental sources (Vogt et al. from Ref. [22], Locherer et al. from Ref. [21], and Howard et al. from Ref. [23].

dispersions of the hypothetic cubic phase, identifying the main phonon instabilities. We then condense various possible combinations of these unstable modes in order to reproduce all experimentally observed structures. This procedure also reveals two new polar phases that are close in energy to the 97 ground-state.

### II. COMPUTATIONAL DETAILS

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Our first-principles calculations have been performed in the context of density functional theory, using the B1-WC hybrid functional [54] as implemented in the CRYSTAL code [55]. We have used the all-electron double- $\zeta$  basis sets for the oxygen atoms and small core Detlev Figgen pseudopotentials [56], associated with double- $\zeta$  valence basis sets for tungsten. We performed full structural relaxations with a convergence criteria on the root-mean square of the gradient and displacements smaller than  $5 \times 10^{-4}$  hartree/bohr and  $5 \times 10^{-4}$ bohrs, respectively. The electronic self-consistent calculations were converged until the difference of the total energy was smaller than  $10^{-9}$  hartree. The phonon frequencies and Born effective charges were computed using frozen phonon numerical differences [57,58] and the electric polarization through the Berry phase technique [59]. The integration in the Brillouin zone has been performed with a  $8 \times 8 \times 8$  grid of k points for the cubic unit cell and a  $4 \times 4 \times 4$  grid for cells doubled in the three directions with respect to the cubic one.

Our choice of a hybrid functional is in line with the results of Wang et al. [44], who have shown that hybrid functionals, and especially HSE06, provide good description of the structural and electronic properties of WO<sub>3</sub>. In our study, we selected the B1-WC functional that was specially designed for perovskite oxides [54] and was already successfully applied to a variety of other compounds [60–63].

#### III. ANALYSIS OF THE EXPERIMENTAL PHASES

Several density functional theory (DFT) studies of WO<sub>3</sub> have been performed previously [37,38,41-44,46-52,64] essentially focusing on the main and most common phases and on the electronic structure analysis with and without oxygen vacancies. A detailed analysis of the complex structural phase diagram of WO<sub>3</sub> is thus missing while a microscopic knowledge of the origin of these different phases would 132 be extremely valuable to understand the unique properties 133 of  $WO_3$ .

In this section we start by characterizing the different phases 135 of WO<sub>3</sub> observed experimentally to validate our approach and 136 we will discuss the possible origin of the Pc phase. We will 137 also analyze the electronic structure of these phases and we will 138 discuss how the B1-WC compares with the previous studies. 139 Further analysis of these phases and other never observed 140 metastable phases (comparison of relative internal energies, 141 symmetry mode analysis of the distortions, coupling of modes) will be reported in Sec. IV B).

### A. Structural and crystallographic analysis

In Table I we compare our calculated crystallographic data 145 of the P4/nmm, P4/ncc, Pbcn,  $P2_1/n$ ,  $P\bar{1}$ , and  $P2_1/c$  146 phases against the experimental measurements. Because the 147  $P2_1/c$  phase is not observed experimentally at low temperature, we compare it with the closely related experimental Pc 149 phase.

Our calculations of the P4/nmm phase are in very good 151 agreement with the observed cell parameters and the atomic 152 positions. The P4/nmm phase is antipolar and consists 153 of highly distorted WO<sub>3</sub> octahedra where the W–O bonds 154 dimerize in opposite direction along the [110] perovskite 155 direction. This W-O dimerization forms local dipole moments 156 that are aligned along the [001] direction and antialigned along 157 the [110] direction, so that the total dipole moment cancels. 158 The crystallographic unit cell is elongated along the [001] 159 direction and compressed along the [100] and [010] directions. 160 This antipolar distortion remains present in all the phases 161 discussed below in this section.

The P4/ncc phase shows additional octahedra rotations 163 around the z axis  $(a^0a^0c^-)$  in the Glazer notation [65]), 164 which induces a cell doubling along the [001] direction. The 165 calculated c cell parameter and the z component of the atomic 166 positions are in good agreement with experiments while the 167 in-plane displacements are less well reproduced (Table I). 168 The calculated a and b cell parameters are smaller than in  $_{169}$ experiments by 0.1 Å and the deviation from the tetragonal 170 O<sub>2</sub> position is about two times larger than observed. We 171 clearly overestimate the amplitude of the  $a^0a^0c^-$  distortion 172 (rotation angle of 13° instead of 7°; see also Fig. 7 further 173 discussed in Sec. IV B). Although this could be partly intrinsic 174 to the functional [66], it is worth noticing that our calculations 175 ignore thermal effects while experiments were performed at 176 high temperatures at which distortions might be reduced [67]. 177 Although such a reduction does not seem to appear for the 178 antipolar motions in the P4/nmm and P4/ncc phases, it 179 might be more substantial for the rotations and we observe 180 that the computed amplitudes of out-of-phase rotations are 181 in much better agreement with experimental data for the 182 low-temperature phases (see further discussed in Sec. IV B).

The orthorhombic *Pbcn* phase can be characterized by 184 an additional in-phase octahedra rotation about the crystal- 185 lographic y axis, yielding a rotation pattern  $a^0b^+c^-$ . We 186 find a similar overestimate of the octahedra distortions as 187 for the P4/ncc phase while the calculated cell parameters 188 are underestimated with respect to experiments. We note that 189

TABLE I. Calculated lattice parameters in Å and Wyckoff positions of distorted WO<sub>3</sub> phases fully relaxed with the B1-WC functional. For each phase, we specify the space group and the experimental parameters are reported for comparison.

P4/nmm		Present			Exp. [23]		
	а	b	c	а	b	c	
	5.299	5.299	3.930	5.297	5.297	3.929	
	X	У	z	X	У	Z	
$W_1$ (2c)	1/4	1/4	-0.0640	1/4	1/4	-0.0660	
O <sub>1</sub> (2c)	1/4	1/4	0.4900	1/4	1/4	0.4900	
$O_2$ (4d)	0	0	0	0	0	0	
P4/ncc		Present			Exp. [23]		
	<i>a</i> 5.168	<i>b</i> 5.168	<i>c</i> 7.870	а 5.278	<i>b</i> 5.278	<i>c</i> 7.849	
	3.108 X			3.278 X			
W <sub>1</sub> (4c)	1/4	у 1/4	z 0.2849	1/4	у 1/4	$\frac{z}{0.2832}$	
$O_1$ (4c)	1/4	1/4	0.2047	1/4	1/4	0.0030	
$O_1$ (8f)		-0.0570	1/4		-0.0250	1/4	
Pbcn	0.0270	Present	-7.	Exp. [22]			
	a	b	<i>c</i>	а	<i>b</i>	<i>c</i>	
	7.284	7.528	7.684	7.333	7.573	7.740	
	x	у	z	X	у	z	
$W_1$ (3d)	0.2510	0.0260	0.2800	0.2520	0.0290	0.2830	
$O_1$ (3d)	-0.0010	0.0430	0.2150	-0.0020	0.0320	0.2210	
$O_2$ (3d)	0.2930	0.2590	0.2590	0.2830	0.2690	0.2590	
$O_3$ (3d)	0.2870	0.0100	0.0060	0.2800	0.0130	0.0020	
$P2_1/n$		Present			Exp. [23]		
	а	b	с	а	b	с	
	7.359	7.486	7.544	7.303	7.538	7.692	
	α	β	γ	α	β	γ	
	90°	91.311°	90°	90°	90.855°	90°	
<b>33</b> 7 (4 )	x 0.2720	y 0.0074	Z 0.2700	x 0.2520	y 0.0260	Z 0.2055	
W <sub>1</sub> (4e)	0.2720	0.0074	0.2790	0.2528	0.0260	0.2855	
$W_2$ (4e)	0.2270	0.0133	0.7750	0.2497	0.0344	0.7805	
O <sub>1</sub> (4e)	0.0043	0.0410	0.2165	0.0003	0.0337	0.2122	
O <sub>2</sub> (4e)	-0.0056	0.4576		-0.0011	0.4632	0.2177	
$O_3$ (4e)	0.2883	0.2534	0.2924	0.2843	0.2598	0.2852	
$O_4$ (4e)	0.2029	0.2530	0.7198	0.2080	0.2588	0.7332	
$O_5$ (4e)	0.2795	0.0385	0.0059	0.2856	0.0410	0.0041	
$O_6$ (4e)	0.2790	0.4630	-0.0047	0.2841	0.4868	-0.0056	
$P\bar{1}$		Present			Exp. [25]		
	a	b	c	a	b	c	
	7.334	7.446	7.612	7.309	7.522	7.678	
	α	$oldsymbol{eta}$	γ	α	$oldsymbol{eta}$	γ	
	88.652°	91.022°	91.012°	88.810°	90.920°	90.930°	
W <sub>1</sub> (2i)	<i>x</i> 0.2603	y 0.0172	z 0.2826	<i>x</i> 0.2566	y 0.0259	$\frac{z}{0.2850}$	
$W_1$ (2i) $W_2$ (2i)	0.2540	0.5210	0.2820	0.2502	0.5280	0.2350	
	0.2340	0.0228	0.2183	0.2302	0.0313	0.7817	
$W_3$ (2i)			0.7793			0.7817	
$W_4$ (2i)	0.2456	0.5268	0.7216	0.2499	0.5338		
O <sub>1</sub> (2i)	0.0015	0.0395		0.0007	0.0386	0.2100	
$O_2$ (2i)	0.5022	0.5406	0.2115	0.5038	0.5361	0.2181	
O <sub>3</sub> (2i)	0.0026	0.4582	0.2897	0.0076	0.4660	0.2884	
O <sub>4</sub> (2i)	0.5012		0.2906	0.4972	-0.0362	0.2878	
	0.2892	0.2571	0.2836	0.2851	0.2574	0.2870	
$O_5$ (2i)							
O <sub>5</sub> (2i) O <sub>6</sub> (2i) O <sub>7</sub> (2i)	0.2081 0.2098	0.7575 0.2569	0.2174	0.2204 0.2186	0.7630 0.2627	0.2223 0.7258	

TABLE I. (Continued.)

$P\bar{1}$		Present		Exp. [25]				
O <sub>8</sub> (2i)	0.2927	0.7575	0.7772	0.2840	0.7583	0.7679		
O <sub>9</sub> (2i)	0.2911	0.0383	0.0060	0.2943	0.0422	-0.0002		
$O_10(2i)$	0.2889	0.5389	0.4941	0.2971	0.5446	0.4982		
O <sub>1</sub> 1 (2i)	0.2108	0.4767	-0.0061	0.2096	0.4820	-0.0072		
$O_12$ $(2i)$	0.2090	-0.0242	0.5063	0.2088	0.9830	0.5051		
$P2_1/c$	Present			Ex	p. (Pc) [20]			
	а	b	С	а	b	c		
	5.263	5.150	7.618	5.278	5.156	7.663		
	$\alpha$	$\beta$	γ	$\alpha$	$\beta$	γ		
	$90^{\circ}$	$91.787^{\circ}$	$90^{\circ}$	90°	91.759°	90°		
	X	y	z	X	у	z		
$W_1$ (2a)	-0.0093	-0.0173	0.6843	-0.0099	-0.0200	0.6743		
$W_{2}(2a)$	0.5011	0.4827	0.7530	0.5000	0.4710	0.7500		
O <sub>1</sub> (2a)	0.4975	0.5769	-0.0245	0.4920	0.5780	-0.0230		
O <sub>2</sub> (2a)	0.2087	0.2891	0.1794	0.2130	0.2890	0.1830		
O <sub>3</sub> (2a)	0.2830	0.7891	0.2580	0.2830	0.7860	0.2590		
O <sub>4</sub> (2a)	0.6999	0.2090	0.1795	0.7050	0.2070	0.1820		
O <sub>5</sub> (2a)	0.7918	0.7090	0.2579	0.7960	0.7110	0.2610		
O <sub>6</sub> (2a)	-0.0058	0.0769	0.4630	-0.0058	0.0730	0.4616		

the antipolar distortions along the z axis compare well with experiments for P4/nmm, P4/ncc, and Pbcn.

The  $P2_1/n$  structure still contains an additional octahedra rotation around the crystallographic x axis, yielding a rotation pattern  $a^-b^+c^-$ . The calculated cell volume is slightly too small (+0.7, -0.7, and -1.9% for a, b and c cell parameters respectively) and the oxygen motions related to the octahedra tilt are overestimated (Table I).

The  $P\bar{1}$  phase is similar to the  $P2_1/n$  phase if one replaces the in-phase rotation by an out-of-phase rotation, yielding a rotation pattern  $a^-b^-c^-$ . The distortions are anisotropic in all three directions, which causes the cell to be triclinic with the angles  $\alpha$ ,  $\beta$ , and  $\gamma$  close to 90°. The calculated a, b, and c coll parameters deviate from experiments by +0.4, -1.1, and -0.8%, respectively.

The  $P2_1/c$  differs from the  $P\bar{1}$  by the fact that two 205 out-of-phase rotations have the same amplitudes, yielding a 206 rotation pattern  $a^-a^-c^-$ . This phase was never reported at 207 low temperatures but is closely related to the experimental  $\varepsilon$  208 (Pc) phase, which only differs from the  $P2_1/c$  phase by an 209 additional polar distortion along the c axis. While relaxing 210 the low-temperature Pc phase, we observed that the system 211 always comes back in to the higher  $P2_1/c$  symmetry. Wijs et al. 212 using local-density approximation and generalized gradient 213 approximation exchange-correlation functionals [38] found a 214 similar effect. To further assess the dynamical stability of the 215  $P2_1/c$  phase with respect to a potential Pc ground state, we 216 computed the zone-center phonons and did not observe any 217 unstable mode: the lowest polar mode has a frequency of 218 158 cm<sup>-1</sup> and is far from being unstable. We also checked 219 whether a soft polar mode can be induced by increasing the 220 cell volume but did not observe any possibility to generate 221 a polar instability. From our calculations, the ground-state 222 structure of intrinsic WO<sub>3</sub> corresponds therefore to a  $P2_1/c$  223 phase. Following the argument by Wijs et al. [38], we suggest 224

TABLE II. Electronic band gap (in eV) of different phases of  $WO_3$  as calculated in the present work with the B1-WC hybrid functional. We compare our results with previous hybrid functional calculations (PBE0, HSE06, and B3LYP), GW calculations, and experimental measurements.

Ref.	B1-WC			HSE06 [42]	B3LYB [42]	Exp1 [35]	Exp2 [68]	Exp3 [43]
Cubic	1.50		2.25	1.67	1.89			
P4/mbm	1.27							
I4/mcm	1.48							
P4/nmm	2.12		2.28	1.71	1.85			1.75
P4/ncc	2.15							
Pcnb	2.65		3.35	2.57	2.89		3.21	2.35
$P2_1/n$	2.85	2.90	3.67	2.80	3.13	2.75	3.25	2.60
$P\bar{1}$	2.98	3.00	3.67	2.94	3.17			
P2 <sub>1</sub> /c	3.28	3.30						

that the polarity in the experimental  $\varepsilon$  (Pc) phase may be stabilized by the presence of oxygen vacancies or by another extrinsic parameter.

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In Table I we compare our calculated atomic positions and cell parameters of the  $P2_1/c$  phase with the experimentally determined Pc phase. The deviations are surprisingly small for lattice parameters (+0.2, -0.2, and -0.5%) for a, b, and c) and even smaller for the atomic positions. Comparing the structural parameters obtained with other hybrid functionals PBE0, B3LYP, and HSE06 reported by Wang et al. [42], we find close agreement with a typical smaller error margin for B1-WC. B1-WC gives a much better agreement for the  $P2_1/c$  phase with experimental data than using the three hybrid functionals tested by Wang et al.: HSE06, B3LYP, and PBE0 with errors of +0.6, +1.3, and +0.2% on a; +2.1, +2.5, and +0.6% on b; and +0.1, +3.0, and +1.7% on the c parameter. We notice, however, that the B1-WC often underestimates cell parameters while the three other hybrid functionals overestimate the cell parameters of WO<sub>3</sub>.

# B. Electronic structure

In Table II we compare the calculated electronic structures for the hypothetical cubic, P4/nmm, P4/ncc, Pbcn,  $P2_1/n$ ,  $P\bar{1}$ , and  $P2_1/c$  phases and compare them with the experiments and previous DFT calculations using PBE0, HSE06, and B3LYP hybrid functionals and GW. For the  $P2_1/n$  and  $P\bar{1}$  phases, experimental data coincide with the B1-WC band gaps. The B1-WC results are similar to those obtained with the HSE06 functional while the PBE0 gives a slightly smaller gap energy and B3LYP larger values. The B1-WC band gap is closest to the results of GW calculations, an agreement also observed for the  $P2_1/c$  phase.

Comparing the trend of band gaps between the different phases, we find that both the appearance of antipolar motions in the P4/nmm phase and out-of-phase rotations in the I4/mcm significantly open the band gap with respect to the cubic phase. Only the in-phase rotations in the P4/mbm phase seem to play a more minor role and slightly close the band gap. The calculated electronic gaps are in reasonable agreement with the experimental values for the three low-temperature structures:

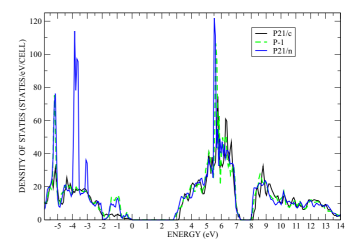


FIG. 2. Calculated density of states of the  $P\bar{1}$ ,  $P2_1/n$ , and  $P2_1/c$  phases with the B1-WC functional.

Eg = 2.85 eV for the room-temperature monoclinic phase  $P_{1}/n$ , Eg = 2.98 eV for the triclinic phase  $P_{1}/n$ , and Eg = 265 3.28 eV for the monoclinic phase  $P_{1}/c$ . Figure 2 shows the density of states of these three phases to demonstrate their similarity.

## IV. ORIGIN OF THE WO3 PHASES

The results presented so far give us confidence that  $^{270}$  the B1-WC functional reproduces well the experimental  $^{271}$  measurements so that we can now focus on the structural  $^{272}$  instabilities of the hypothetical  $Pm\bar{3}m$  cubic parent phase  $^{273}$  and explain how their condensation gives rise to the various  $^{274}$  known phases of WO<sub>3</sub>. This also allows us to identify novel  $^{275}$  ferroelectric metastable phases. In each case, we analyze  $^{276}$  the crystallographic structure through a decomposition of the  $^{277}$  distortions with respect to the cubic parent phase in terms of  $^{278}$  symmetry-adapted modes.

#### A. Unstable modes of the cubic reference

Figure 3 shows the calculated phonon dispersion curves 281 of hypothetical cubic WO<sub>3</sub>. Two branches of instabilities 282 (imaginary frequencies plotted as negative numbers in Fig. 3) 283 coexist in the Brillouin zone. 284

The first unstable branch has its largest imaginary value at  $\Gamma$ . The  $\Gamma$  unstable mode has the irreducible representation (irrep)  $\Gamma_4^-$  and corresponds to a polar mode. It suggests that the cubic phase of WO<sub>3</sub> is mostly unstable via this polar instability and might be ferroelectric, which we will see later is not exactly the case. The polar instability at  $\Gamma$  propagates toward the X and M points with weak dispersion while it strongly disperses towards the R point. Aside from  $\Gamma$ , the modes of this branch are antipolar. The dispersion of this unstable branch is very similar to the one reported in BaTiO<sub>3</sub> and corresponds to a ferroelectric instability requiring a chainlike correlation of displacements in real space [70].

The second branch of unstable modes appears between M  $_{297}$  and R points with smaller amplitudes and a nearly absent  $_{298}$  dispersion between these two points. The labels of the M  $_{299}$  and R point unstable phonon modes are  $M_3^+$  and  $R_4^+$  and  $_{300}$ 

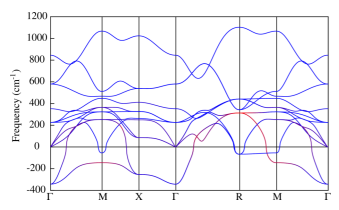


FIG. 3. Phonon dispersion curves of cubic WO<sub>3</sub> (negative frequencies refer to imaginary frequencies, i.e., to unstable modes). The coordinates of the high-symmetry points are as follows:  $\Gamma$  (0,0,0), X ( $\frac{1}{2}$ ,0,0), M ( $\frac{1}{2}$ , $\frac{1}{2}$ ,0), and R ( $\frac{1}{2}$ , $\frac{1}{2}$ , $\frac{1}{2}$ ). Thanks to the *band2eps* postprocessing script of ABINIT [69], the color of the bands is assigned to each point through the contribution of each atom type to the corresponding eigenvector: red for the tungsten atom and blue for the oxygens.

they correspond to rotations of the oxygen octahedra. The dispersion of this branch is comparable to what is observed for similar modes in SrTiO<sub>3</sub> or PbTiO<sub>3</sub> and linked to a planar character of the correlations of the atomic displacements in real space [70].

Figure 4 represents a schematic view of the eigenvectors related to the main instabilities of cubic WO<sub>3</sub>. The polar mode at  $\Gamma$  ( $\Gamma_4^-$ , 373i cm<sup>-1</sup>) shows motion of W against the O atoms, which is the source of a large electrical polarization. The antipolar modes at the X ( $X_5^-$ , 256i cm<sup>-1</sup>) and M points ( $M_3^-$ , 147i cm<sup>-1</sup>) are associated to opposite displacements from unit cell to unit cell along the [100] and [110] directions, respectively [71]. The  $M_3^+$  (62i cm<sup>-1</sup>) and  $R_4^+$  (69i cm<sup>-1</sup>) unstable modes correspond rotations of the oxygen octahedra about the central W atom with consecutive octahedra along the rotation direction moving, respectively, in the same or opposite directions. Using the Glazer notation [65], the  $M_3^+$  mode corresponds to  $a^0a^0a^-$ .

## B. Condensation and coupling of modes

Starting from the previous unstable modes, we now investigate how their individual and combined condensations in the hypothetical cubic structure give rise to various phases. We then compare their energies and analyze the amplitudes of distortions.

# 1. Condensation of modes of the unstable polar branch

We first consider the condensation of unstable  $\Gamma_4^-$ ,  $X_5^-$ , and  $M_3^-$  modes. Figure 5 shows the energy gain of the corresponding relaxed phase with respect to the cubic phase. We tested several condensation schemes: (i) condensation of the polar  $\Gamma_4^-$  mode along one (P4mm), two (Amm2), and three (R3m) directions; (ii) condensation of the  $X_5^-$  along one (Pmma) and two  $(P2_1/m)$  directions; and (iii) condensation of the  $M_3^-$  mode along one direction (P4/nmm).

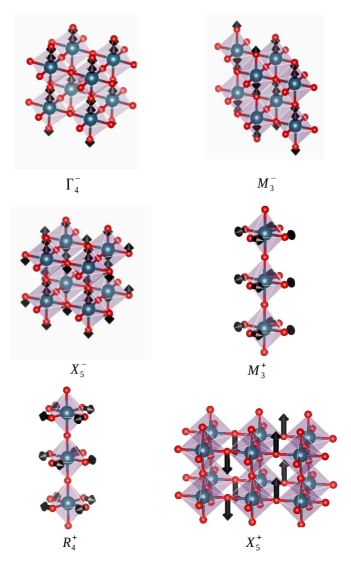


FIG. 4. Schematic view of the most important modes contributing to the distortions of WO<sub>3</sub>. Small red spheres represent the oxygens, and large blue spheres represent the tungsten atoms. All the modes are unstable but the  $X_5^+$  mode, which is discussed in Sec. IV B 4.

We observe that the energy gain of the polar instabilities  $_{355}$  is large and that the  $\Gamma_4^-$  polar mode drives a larger gain of energy (red columns in Fig. 5) than the antipolar  $X_5^-$  and  $M_3^ _{337}^-$  modes (green columns in Fig. 5). The space group related to the condensation of the  $M_3^-$  mode corresponds to the high-temperature phase observed experimentally (P4/nmm).

Condensation of the  $\Gamma_4^-$  mode along two and three directions produces energy gains larger than its condensation in a single direction so that the  $\Gamma_4^-$  mode alone will drive the system polar along the [111] direction with an energy difference between the Amm2 and R3m phases of 6 meV. We calculated the polarization amplitude in the three P4mm, Amm2, and R3m phases using the Berry phase technique and obtain 54, 69, and 69  $\mu$ C cm $^{-2}$ . These polarization values are comparable to those observed in robust ferroelectrics such as PbTiO3. They can be explained by the opposite motions of W and O atoms, associated with strongly anomalous Born effective charges (11.73 e for W and -8.78/-1.62 e for  $O_{\parallel}/O_{\perp}$  in 352 good agreement with previous calculations in Ref. [72]).

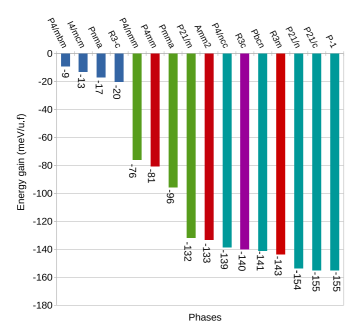


FIG. 5. Calculated energy gain (in meV/f.u.) with respect to the cubic phase of different phases of WO<sub>3</sub>. Red columns are the FE phases arising for the condensation of the polar unstable mode, blue columns are the phases arising from the condensation of the oxygen octahedral rotation unstable modes, green columns are the phases arising from the condensation of antipolar modes, the magenta column represents a phase combining FE and antipolar modes, and cyan columns represent phases combining oxygen octahedral rotations and antipolar modes. For clarity, the exact value of the energy gain is written in each case.

# 2. Condensation of modes of the oxygen rotation unstable branch

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Figure 5 (blue columns) shows the energy gain given by the condensation of the  $M_3^+$  and  $R_4^+$  modes along one direction (I4/mbm and I4/mcm), the  $R_4^+$  mode in three directions  $(R\bar{3}c)$ , and the orthorhombic Pnma phase where the  $R_4^+$  mode is condensed in two directions and the  $M_3^+$ mode in one direction  $(a^-a^-c^+)$ . These distortions lower the energy much less than the polar and antipolar motions. This observation is in line with the modest amplitude of the related phonon instabilities: the energy curvatures at the origin are less negative for the octahedral rotations than for the polar motion. Nevertheless the amplitudes of octahedral rotations are 10.7 and 11.7 deg in I4/mbm and I4/mcm, respectively. Such large distortions associated to a weak instability highlight relatively small anharmonicities, which might be explained by the absence of the A cation with respect to regular ABO<sub>3</sub> perovskites [73].

# 3. Combinations of modes

Beside the P4/nmm phase, none of the previous single irrep mode condensations correspond to observed phases. Thus, we now explore the condensation of combined octahedral rotations and polar/antipolar modes. We depict in Fig. 5 the energy gain given by the joint condensation of polar and oxygen rotation modes along three directions (R3c, in purple color) and of antipolar and oxygen rotation modes (P4/ncc, Pbcn,  $P2_1/n$ ,  $P2_1/c$ , and  $P\bar{1}$ , in cyan color).

Combining the polar distortion of the low-energy R3m 380 phase with additional oxygen rotation modes does not further reduce the energy. Instead, it yields a R3c phase slightly higher in energy but with a slightly amplified polarization 381 of  $71~\mu C cm^{-2}$  and a slightly reduced oxygen rotation (with respect to the  $R\bar{3}c$ ). This emphasizes an unusual competition 385 between these two types of distortions in WO<sub>3</sub> with respect to 386 regular perovskite compounds, where in WO<sub>3</sub> the R3c phase 387 forms a local minimum between the  $R\bar{3}c$  and R3m phases.

At the opposite, the mixing of the antipolar mode  $M_3^-$  with  $_{389}$ oxygen rotation modes can drive larger energy gains so that 390 the ferroelectric R3m phase is not the ground state. This is in 391 agreement with experimental observations where the observed 392 phases at all temperatures contain antipolar motions. Among 393 investigated phases, the  $P2_1/c$  phase appears as the most stable 394 but only marginally, as we observe that the  $P2_1/n$ ,  $P2_1/c$ , 395 and P1 phases are all extremely close in energy (energy 396 gains of 153, 155, and 155 meV, respectively, see Fig. 5). Consequently, within the precision of our calculations, we 398 cannot unambiguously assess which one is the ground state. Nevertheless, as discussed in Sec. III A and further exemplified 400 in the next section, the  $P2_1/c$  phase is in excellent agreement 401 with the experimental Pc ground state, except for a tiny 402 polar distortion. Our calculations highlight that, in fact, the 403  $P2_1/n$  and  $P\bar{1}$  phases observed at higher temperatures are 404 also extremely close in energy.

We further notice that the ferroelectric R3m phase, although 406 never observed experimentally, is also relatively close in 407 energy to the ground state (about 11 meV/f.u.). Following 408 Rabe [74], the nonpolar (or eventually weakly polar in the 409 experimental Pc phase) ground state of WO<sub>3</sub> combined 410 with an alternative low-energy ferroelectric phase obtained 411 by polar distortions of the same high-symmetry reference 412 structure makes it a potential antiferroelectric compound. 413 Indeed, applying an electric field, it might be possible to 414 open a typical double hysteresis loop from a field-induced 415 first-order transition from the  $P2_1/c$  ground state to the  $R3m_{416}$ polar phase. Estimating the critical electric field required 417 to stabilize the R3m phase from  $\mathcal{E}_c \sim \Delta E/\Omega_0 P_s$  [75], 418 where  $\Delta E$  is the energy difference between the two phases 419 (11.43 meV/f.u.), P<sub>s</sub> is the spontaneous polarization of the 420 polar phase (69  $\mu$ C cm<sup>-2</sup>), and  $\Omega_0$  is its unit-cell volume <sub>421</sub> (55 Å), we get the relatively modest value  $\mathcal{E}_c \sim 480 \text{ kV/cm}$ . 422 For the polar phase R3c we need to apply a greater electric 423 field  $\mathcal{E}_c \sim 638 \text{ kV/cm}$  to stabilize this phase. This allows us 424 to estimate that the critical field has similar value with respect 425 to other antiferroelectric material,  $\mathcal{E}_c \sim 470~\mathrm{kV/cm}$  for ZrO<sub>2</sub> 426 [75] and  $\mathcal{E}_c \sim 239 \text{ kV/cm}$  for PbZrO<sub>3</sub> [76]. Although this 427 might not be easy to check experimentally on real samples 428 that are typically oxygen deficient and highly conductive, the 429 calculations reveal that stoichiometric WO3 exhibits all the 430 features of an antiferroelectric compound.

# 4. Symmetry adapted mode analysis of the distorted phases

To quantify the distortions that appear in the various phases we project the structural distortions onto symmetry adapted modes of the cubic phase using AMPLIMODE software [77]. 435 The results in Fig. 6 show the amplitudes of the modes in the fully relaxed phases from the calculations but nonobserved 437

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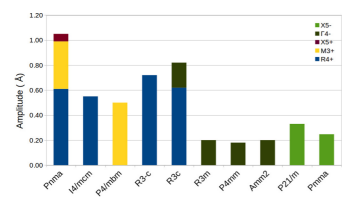


FIG. 6. Symmetry adapted mode decomposition of distorted phases of WO<sub>3</sub> explored in our study but not observed experimentally.

experimentally. In Fig. 7 we show the amplitudes of modes in both the fully relaxed and observed phases, which can be compared. 440

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In the following we discuss the competition or cooperation character of the mode distortions. In perovskite oxides, it is established that the oxygen rotations are in competition with the ferroelectric displacements but less attention has been given to the combinations of other types of mode. Often, this cooperation or competition comes from the biquadratic energy term in the free-energy expansion with respect to two order parameters. In WO<sub>3</sub>, we observe that the combination between the  $\Gamma_4^-$  mode and the  $R_4^+$  mode along the [111] direction in the R3c phase has the tendency to reduce the amplitude of the oxygen rotations with respect to the  $R\bar{3}c$  phase (the  $R_4^+$ mode is 13% smaller in the R3c phase than in the R3c phase, see Fig. 6) while the polar mode is unaffected. As discussed in the previous section, the combination of the  $\Gamma_{\perp}^{-}$  and  $R_{\perp}^{+}$ modes forms a local minimum (R3c phase) of higher energy than the R3m phase. This means that the polar distortions are in competition with the oxygen rotations as reported for perovskite oxides, with the difference that the polar mode amplitude is unaffected and that the R3c phase is locally

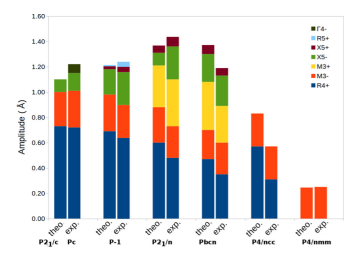


FIG. 7. Symmetry adapted mode decomposition of distorted phases WO<sub>3</sub>; comparison between experiments and our calculations with the B1-WC functional are shown.

stable (the system does not relax into the lowest-energy R3m 460 phase). The strain can also play an important role [78], but 461 when performing the same calculations at fixed cell parameters 462 (fixed to the cubic ones), we find that the R3c phase still forms 463 a local minimum of higher energy than the R3m phase. This 464 unusual energy landscape can come from the marginal gain 465 of energy of the oxygen rotations while large amplitudes of 466 rotations are present.

On the other hand, the association of the oxygen rotations 468 with the antipolar  $M_3^-$  mode is cooperative. When we compare 469 the amplitude of the  $R_4^+$  and  $M_3^-$  modes of the P4/nmm, 470 I4/mcm, and P4/ncc (Figs. 6 and 7) we find that when both 471 the  $R_4^+$  and  $M_3^-$  modes are present together in the P4/ncc 472 phase, their amplitude is slightly higher (4% larger) than 473 when condensed alone (P4/nmm and I4/mcm phases). Their 474 combination, however, drives a sizable gain of energy: the 475 P4/ncc phase is 63 and 126 meV lower in energy than the 476 P4/nmm and the I4/mcm phases, respectively. This means 477 that the combination of the oxygen rotations with the antipolar 478  $M_3^-$  mode is much more cooperative than the combination with 479 the polar mode  $\Gamma_4^-$ .

The *Pbcn* phase can be understood as a distorted P4/ncc 481 phase with additional M<sub>3</sub><sup>+</sup> oxygen rotations along [010]. The 482 resulting tilt pattern is  $a^0b^+c^-$  with a small energy gain of 2 483 meV with respect to the P4/ncc phase and a reduction of mode 484 amplitudes  $M_3^-$ ,  $R_4^+$ , and  $M_3^+$  (16, 7, and 22% reduction of the 485  $M_3^-$ ,  $R_4^+$ , and  $M_3^+$  modes with respect to the phases where they are condensed alone, i.e., P4/nmm, I4/mcm, and P4/mbm, 487 respectively). The  $M_3^+$  mode competes with the  $R_4^+$  and  $M_3^$ modes in the sense their combination reduces their amplitude, 489 but they cooperate to lower the energy of the system.

In the case of the monoclinic  $P2_1/n$ ,  $P2_1/c$ , and triclinic 491  $P\bar{1}$  phases, the combination of the  $M_3^-$  mode with several 492 oxygen rotations  $(a^-b^+c^-)$  for  $P2_1/n$ ,  $a^-a^-c^-$  for  $P2_1/c$ , 493 and  $a^-b^-c^-$  for  $P\bar{1}$ ) lowers the energy of the crystal and with 494 an increase of the mode amplitude with respect to the phases 495 where these modes are condensed independently. For example, 496 the antipolar  $M_3^-$  mode has his amplitude increased by 11, 10, 497 and 19% in the  $P2_1/n$ ,  $P2_1/c$ , and  $P\bar{1}$  phases, respectively. 498 This means that the dominant R<sub>4</sub><sup>+</sup> oxygen rotations cooperate 499 with the antipolar M<sub>3</sub><sup>-</sup> mode to promote the ground state 500 of WO<sub>3</sub>.

We note that in the P4/nmm, P4mm, Amm2, R3m, 502 I4/mcm, I4/mbm,  $R\bar{3}c$ , P4/ncc, and R3c phases the mode decomposition shows only the primary modes we have 504 condensed. This is different with the Pnma phase in which 505 an additional mode  $X_5^+$  appears with a small amplitude in the 506 mode projections while we have condensed only the primary 507  $R_4^+$  and  $M_3^+$  modes (see Fig. 6). This additional mode appears 508 by anharmonic coupling between the  $R_4^+$  and  $M_3^+$  modes such 509 that the symmetry of the Pnma structure allows the  $X_5^+$  mode 510 to develop even though the  $X_5^+$  mode is not unstable by itself 511 [79]. Similarly, we observe the apparition of several additional 512 modes in the Pbcn,  $P2_1/n$ ,  $P2_1/c$ , and  $P\bar{1}$  phases, which we 513 discuss in the next section.

## 5. Energy invariants

The different phases arise from the condensation of one or 516 several unstable modes (primary modes) of the cubic parent 517

TABLE III. Symmetry adapted modes decomposition and linear couplings of modes of the Pnma, Pbcn,  $P2_1/n$ ,  $P2_1/c$ , and  $P\bar{1}$  phases. From the left to right columns, we show the mode label (Irrep.) of the symmetry adapted mode, the direction of the mode condensation, the corresponding subgroup, the amplitude of the distortion in the calculated and in the experimental cases (the modes with an amplitude lower than 0.005 Å are not shown), and the linear coupling invariants of the most relevant modes where the letters correspond to the one given in the direction column (we highlight in bold the primary modes).

Space group	Irrep.	Direction	Subgroup	Amplitudes (Å)		
				Calc.	Exp.	Linear invariants
Pnma	$R_4^+$	( <b>a</b> , <b>a</b> ,0)	I4/mma	0.61		
	$\mathbf{M_3^+}$	$(0,0,\mathbf{c})$	P4/mbm	0.38		abc
	$X_5^+$	(0,a,0,0,0,0)	Cmcm	0.06		
Pbcn	$\mathbf{R_4^+}$	(0,0,a)	I4/mcm	0.47	0.35	
	$\mathbf{M_3^+}$	$(0, \mathbf{b}, 0)$	P4/mbm	0.39	0.30	abe, adce
	$\mathbf{M_3^-}$	$(c,0,\mathbf{d})$	Ibam	0.23	0.25	abf, adcf
	$X_5^-$	(0,0,e,-e,0,0)	Pmma	0.22	0.25	
	$X_5^+$	(0,0,0,0,f,f)	Pmma	0.07	0.06	
$P2_1/n$	$\mathbf{R_4^+}$	(0, a, b)	C2/m	0.60	0.48	aeg+bdf
	$\mathbf{M_3^+}$	$(\mathbf{c},0,0)$	P4/mbm	0.34	0.37	ach+aci-bch+bci
	$\mathbf{M_3^-}$	$(0, \mathbf{d}, \mathbf{e})$	Ibam	0.27	0.25	acdg-bcef
	$X_5^-$	(f,-f,0,0,g,g)	Pmmn	0.09	0.26	adeh+adei+bdeh-bdei
	$X_5^+$	(0,0,h,i,0,0)	$P2_1/m$	0.06	0.08	
$P2_1/c$	$\Gamma_4^+$	$(\mathbf{a},-\mathbf{a},-\mathbf{b})$	Cm	0	0.07	
	$\mathbf{R_4^+}$	(-b,a,-a)	C2/c	0.73	0.72	acd+bcd
	$\mathbf{M_3^-}$	( <b>c</b> ,0,0)	P4/nmm	0.27	0.30	
	$X_5^-$	(0,0,0,-d,0,0)	Cmcm	0.10	0.14	

structure but, in some cases, also include the further condensation of stable modes (secondary modes) with significant amplitudes. Often, the appearance of such secondary modes of large amplitude can be traced back in their linear coupling with the primary modes [80,81]. This can be rationalized from the existence in the energy expansion of the  $Pm\bar{3}m$  phase of invariant terms of the form  $\gamma Q_s \prod_{i=1}^N Q_p^i$  where  $Q_s$  is the amplitude of the secondary mode s and  $Q_p^i$  is the amplitude of the primary mode s. Depending of the number of primary mode involved, these coupling terms can be bilinear, trilinear, quadrilinear, etc.

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To clarify the origin of secondary modes in several low-symmetry phases of WO<sub>3</sub>, we restrict ourself here to the search of such multilinear invariant terms in the energy expansion around its cubic phase by using the INVARIANTS software [82]. In the last column of Table III we report these linear invariants up to the fourth order obtained for the *Pnma*, *Pbcn*,  $P2_1/n$ , and  $P2_1/c$  phases. The letters represent the mode amplitudes  $(Q^i)$  in the directions specified in the third column where the bold letters refer to the primary modes in the structure  $(Q^i_p)$  and the normal letters refer to secondary modes  $(Q^i_s)$ .

In the Pnma phase, we find that the  $X_5^+$  mode appears through a trilinear coupling with the oxygen rotations modes  $R_4^+$  and  $M_3^+$  (**ab**c invariant in Table III). This additional  $X_5^+$  mode is also found in Pnma of  $ABO_3$  perovskites [83,84] where the eigenvector corresponds to antipolar motions of the A cation. In  $WO_3$  the A cation is absent and  $X_5^+$  corresponds

to similar antipolar motions but of oxygen instead of the A site (see Fig. 4). 546

In *Pbcn* the primary  $M_3^-$  mode condenses along z, the  $R_4^+$  548 mode condenses along z, and the  $M_3^+$  mode condenses along 549 y and four additional secondary modes:  $X_5^-$  and  $X_5^+$  with a 550 large amplitude and  $M_5^+$  and  $M_4^+$  with a small amplitude as 551 well as an additional component of the M<sub>3</sub><sup>-</sup> mode about the 552 x direction. If we restrict ourself to the strongest  $X_5^-$  and  $X_5^+$ modes we find that both are coupled with the  $R_4^+$  and  $M_3^+$ modes through a trilinear coupling (abe and abf invariants in 555 Table III) but also through a quadrilinear coupling with the  $R_4^+$ mode and the two components of the  $M_3^-$  mode (adce and adcf 557 invariants in Table III). We can thus explain the appearance of 558 the  $X_5^+$  and  $X_5^-$  modes through a trilinear coupling with the 559 oxygen octahedral rotations and the appearance of the second 560 x component of the antipolar  $M_3$  mode through a coupling 561 with the secondary  $X_5^+$  and  $X_5^-$  modes and the primary  $R_4^+$ mode. The final structure can thus be seen as antipolar through 563 the  $M_3^-$  mode along z with a canting of its direction toward 564 the x axis and through the  $X_5^-$  mode along the y direction, the 565 whole distortions being associated with the  $a^-b^+c^-$  pattern of 566 oxygen rotation distortions.

The transition from Pbcn to  $P2_1/n$  can be seen as 568 being induced by the condensation of the  $R_4^+$  mode along 569 the remaining direction for the oxygen rotation octahedral 570 distortions to  $a^-b^+c^-$ . This means that we find the same mode 571 coupling as in the Pbcn phase plus some extra ones due to 572 the additional mode condensation. Because we do not induce 573

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a new irrep, the couplings are the same (i.e., trilinear and quadrilinear between the primary  $R_4^+$ ,  $M_3^+$ , and  $M_3^-$  modes and the secondary  $X_5^-$  and  $X_5^+$  modes) but in different directions from the *Pbcn* phase: we observe the  $X_5^-$  and  $X_5^+$  modes in two directions instead of one. Other modes also appear in the symmetry adapted mode analysis but with smaller amplitudes  $(M_4^+, M_5^-, M_5^+, \text{ and } M_2^+)$ , which we do not include in the invariant analysis.

P1 and  $P2_1/c$  are very similar in the sense that for both structures we can envisage the condensation of R<sub>4</sub><sup>+</sup> modes in three directions and the M<sub>3</sub>-mode in one direction. The difference is that in the  $P2_1/c$  phase the  $R_4^+$  mode is primary with the same amplitude in two directions and a different amplitude in the third direction  $(a^-a^-c^-$  where the  $M_3^-$  mode is primary in the z direction) while in the  $P\bar{1}$  phase the condensation of the  $R_4^+$  mode has different amplitudes in three directions  $(a^-b^-c^-)$ . In  $P2_1/c$  the presence of the  $X_5^-$  and  $M_5^$ secondary modes can be explained by trilinear coupling with the  $R_4^+$  and  $M_3^-$  primary modes (acd+bcd and abe invariants in Table III) in a similar way as in the  $P2_1/n$  phase.

This analysis shows that the low-symmetry phases of WO<sub>3</sub> are complex and involve numerous multilinear couplings of modes if one expands the energy with respect to the cubic phase. We note that, among possible couplings, the coupling with the secondary  $X_5^-$  mode is most important in all lowsymmetry phases.

Going further, in order to test whether symmetry arguments can lead to the polar Pc phase using anharmonic couplings, we have tested if there exists any bi-, tri-, or quadri-linear coupling involving a polar mode at the  $\Gamma$  point. We did not find any couplings with the  $R_4^+$ ,  $M_3^+$ , or  $M_3^-$  modes. We thus conclude that it is not possible to generate polarity in WO<sub>3</sub> in the limit to these primary modes, which are the ones appearing in other experimental phases.

## V. CONCLUSIONS

In this study, we have performed a first-principles study of WO<sub>3</sub> using the B1-WC hybrid exchange-correlation functional which appears to yield good overall agreement with experiments regarding electronic and structural properties together.

Starting from the inspection of the phonon dispersion curves of a hypothetical cubic structure taken as reference, we have identified two main branches of instabilities and characterized various phases arising from the condensation of one or more unstable modes. Although the dominant phonon instability is associated to a zone-center polar mode, we found 618 a nonpolar  $P2_1/c$  ground state arising from the combination 619 of cooperative antipolar distortions and oxygen octahedra 620 rotations. This phase is very similar to the experimentally reported polar Pc ground state, except for the absence of a 622 tiny polar distortion. Our calculations do not show, however, 623 any tendency of the  $P2_1/c$  phase to evolve to a Pc phase 624 suggesting that WO<sub>3</sub> is likely not intrinsically ferroelectric. 625 Instead the ferroelectric character might arise from extrinsic 626 defects such as oxygen vacancies. The  $P2_1/c$  phase is antipolar and defects could easily produce a slightly unbalanced antidipole structure, yielding a weak net polarization. In this 629 sense, off-stoichiometric WO<sub>3</sub> might be better described as a 630 ferrielectric compound [85]. The ground state is determined by 631 two antiparallel movements of W off-centerings which exactly 632 compensate each other in the  $P2_1/c$  phase. The displacements 633 are almost identical in the Pc phase, but the two displacements 634 do not fully compensate each other. We suspect that such 635 weak ferrielectricity can be induced by defects such as oxygen 636 vacancies.

At the level of our calculations, the  $P2_1/c$  ground state 638 is almost degenerated in energy with the  $P2_1/n$  and  $P\bar{1}$  639 phases observed at higher temperature. Also, we discovered 640 the existence of a never observed and low-energy ferroelectric 641 R3m phase with a large polarization. Although this might 642 not be of direct interest due to the conductive character 643 of usual off-stoichiometric samples, the proximity with the 644  $P2_1/c$  ground state of this structurally unrelated R3m polar 645 phase toward which the system could be switched through 646 a first-order transition under moderate electric fields makes 647 WO<sub>3</sub> a potential antiferroelectric material.

### **ACKNOWLEDGMENTS**

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