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1 Antiferromagnetic Order Breaks Inversion Symmetry in a Metallic 2 Double Perovskite, $\text{Pb}_2\text{NiOsO}_6$

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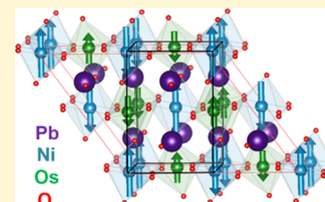


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Supporting Information

5 **ABSTRACT:** A polycrystalline sample of $\text{Pb}_2\text{NiOsO}_6$ was synthesized under high-pressure (6
6 GPa) and high-temperature (1575 K) conditions. $\text{Pb}_2\text{NiOsO}_6$ crystallizes in a monoclinic double
7 perovskite structure with a centrosymmetric space group $P2_1/n$ at room temperature. $\text{Pb}_2\text{NiOsO}_6$
8 is metallic down to 2 K and shows a single antiferromagnetic (AFM) transition at $T_N = 58$ K.
9 $\text{Pb}_2\text{NiOsO}_6$ is a new example of a metallic and AFM oxide with three-dimensional connectivity.
10 Neutron powder diffraction and first-principles calculation studies indicate that both Ni and Os
11 moments are ordered below T_N and the AFM magnetic order breaks inversion symmetry. This loss
12 of inversion symmetry driven by AFM order is unusual in metallic systems, and the 3d–5d double-
13 perovskite oxides represent a new class of noncentrosymmetric AFM metallic oxides.



1. INTRODUCTION

14 Transition-metal oxides (TMOs) exhibit unique correlations
15 between magnetism and electrical conductivity: ferromagnet-
16 ism (FM) in TMOs usually coexists with metallic conductivity,
17 whereas insulating TMOs usually exhibit antiferromagnetism
18 (AFM).¹ Exceptions from this behavior, such as FM insulating
19 oxides and AFM metallic oxides, are less common. CaCrO_3
20 and $\text{Nb}_{12}\text{O}_{29}$ are examples of AFM metallic oxides with three-
21 dimensional crystal structures.^{2–4} Other AFM metallic oxides
22 such as $\text{La}_{2-2x}\text{Sr}_{1+2x}\text{Mn}_2\text{O}_7$ and $\text{Ca}_3\text{Ru}_2\text{O}_7$ crystallize in layered
23 crystal structures, and FM couplings are dominant within the
24 layer.^{5,6} Recently, RuO_2 , and LaNiO_3 , which had been
25 described as paramagnetic metals, were found to be AFM-
26 ordered and are new examples of AFM metallic oxides with
27 three-dimensional crystal and electronic structures.^{7–9}

28 5d TMOs are unique correlated systems because of the
29 spatial extent of the 5d electrons, generally giving 5d TMOs
30 wider bandwidths (W), stronger spin–orbit coupling (SOC),
31 and smaller on-site Coulomb repulsion (U) compared with 3d
32 TMOs.¹⁰ For instance, metal–insulator transitions driven by
33 AFM orders were proposed in 5d oxides, $\text{Pb}_2\text{CaOsO}_6$,¹¹
34 $\text{Cd}_2\text{Os}_2\text{O}_7$,^{12,13} and NaOsO_3 ,¹⁴ and a ferroelectric-like
35 structural-transition-breaking inversion symmetry has been
36 observed in metallic LiOsO_3 .¹⁵ Recent studies on a 5d metallic
37 oxide $\text{Pb}_2\text{CoOsO}_6$ demonstrated that the AFM order breaks
38 inversion symmetry.^{16,17} In this work, we have built on this,
39 synthesizing a new 5d hybrid double perovskite oxide
40 $\text{Pb}_2\text{NiOsO}_6$, which is a new example of an AFM metallic
41 oxide. Characterization using neutron powder diffraction
42 (NPD) and property measurements suggest that the magnetic
43 order breaks inversion symmetry (similar to reports on
44 $\text{Pb}_2\text{CoOsO}_6$ ¹⁶). First-principles calculations confirm that
45 both Ni and Os moments are ordered, allowing us to confirm

the nature of the ground state (which has not been fully
explored previously). The 3d–5d double-perovskite oxides
establish a new class of noncentrosymmetric AFM metallic
oxides, and our symmetry analysis of $\text{Pb}_2\text{NiOsO}_6$ explores how
this understanding can be applied more widely to design new
magnetoelectrics.

2. EXPERIMENTAL SECTION

Polycrystalline $\text{Pb}_2\text{NiOsO}_6$ was synthesized via a solid-state reaction
from powders of PbO_2 (99%, Alfa), Os (99.95%, Heraeus Materials),
 NiO (99.997%, Alfa), and KClO_4 (99.99%, Alfa). The powders were
thoroughly mixed in a stoichiometric ratio in an Ar-filled glovebox,
followed by sealing in a Pt capsule. The Pt capsule was statically and
isotropically compressed in a belt-type high-pressure apparatus (Kobe
Steel, Ltd., Japan¹⁸), and a pressure of 6 GPa was applied while the
capsule was heated at 1400 °C for 1 h, followed by quenching to
room temperature in less than a minute. The pressure was then
gradually released over several hours.

A dense and black polycrystalline pellet was obtained, and several
pieces were cut out from it. A selected piece was finely ground for a
synchrotron X-ray diffraction (SXRD) study, which was conducted in
a large Debye–Scherrer camera in the BL15XU beamline, SPring-8,
Japan.^{19,20} The SXRD pattern was collected at room temperature, and
the wavelength was confirmed to be 0.65298 Å by measurement of a
standard material, CeO_2 . The absorption coefficient was measured in
the same line. The SXRD data were analyzed by the Rietveld method
with the RIETAN-VENUS software.²¹ The crystal structure was
depicted by VESTA.²²

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72 The electrical resistivity (ρ) of a polycrystalline pellet of
 73 $\text{Pb}_2\text{NiOsO}_6$ was measured by a four-point method at a gauge current
 74 of 0.1 mA in a physical property measurement system (PPMS,
 75 Quantum Design, Inc.). Electrical contacts were made with Pt wires
 76 and Ag paste in the longitudinal direction. The temperature
 77 dependence of the specific heat capacity (C_p) was measured in the
 78 same PPMS by a thermal relaxation method at temperatures between
 79 2 and 300 K with Apiezon N grease thermally connecting the material
 80 to the holder stage.

81 The magnetic susceptibility (χ) of $\text{Pb}_2\text{NiOsO}_6$ powder was
 82 measured in a magnetic property measurement system (Quantum
 83 Design, Inc.). The measurement was conducted in field-cooled (FC)
 84 and zero-FC conditions in the temperature range between 2 and 390
 85 K. The applied magnetic field was 10 kOe.

86 Time-of-flight NPD data were collected at the WISH diffrac-
 87 tometer (target station 2) at the ISIS Neutron and Muon Source.²³

88 For the NPD data collection, 3.26 g of $\text{Pb}_2\text{NiOsO}_6$ powder was placed
 89 in a 6 mm diameter cylindrical vanadium can under helium and sealed
 90 using indium wires. The sample was loaded into a helium cryostat and
 91 cooled to base temperature (1.5 K). A high-quality data set was
 92 collected at 1.5 K (~ 1 h, $\sim 40 \mu\text{A}$ h), and shorter scans (~ 15 min,
 93 $\sim 10 \mu\text{A}$ h) were collected every 2.5 K on warming to 100 K. A final
 94 higher-quality scan (~ 1 h, $\sim 40 \mu\text{A}$ h) was collected at 98 K in the
 95 paramagnetic phase. Data were analyzed and Rietveld refinements
 96 carried out using TopasAcademic,^{24,25} and the web-based ISO-
 97 DISTORT software²⁶ was used for symmetry analysis. Rietveld
 98 refinements for the antiferromagnetic system were carried out with a
 99 nuclear phase and a magnetic-only phase, with atomic displacement
 100 parameters (ADPs) for the magnetic sites constrained to be equal to
 101 those sites in the nuclear phase. A separate peak shape was refined for
 102 the magnetic-only phase.

103 The density functional theory (DFT) calculation was performed on
 104 $\text{Pb}_2\text{NiOsO}_6$ with the all-electron full-potential linearized augmented
 105 plane-wave method implemented in the WIEN2k code.²⁷ Generalized
 106 gradient approximation (GGA) of Perdew–Burke–Ernzerhof
 107 (PBE)²⁸ was used for the exchange–correlation functional. The
 108 SOC was taken into account in the second variation method. To
 109 consider the correlation effect, GGA + U was adopted within fully
 110 localized limits.^{29,30} The on-site Coulombic interaction parameters U
 111 = 4 and 2 eV for Ni and Os, respectively, and the Hund's coupling J_H
 112 = 0.8 eV, which was shown to describe a similar compound,
 113 $\text{Ca}_2\text{NiOsO}_6$ properly.³¹

3. RESULTS

114 **3.1. Crystal Structure.** Room-temperature SPXD data of
 115 $\text{Pb}_2\text{NiOsO}_6$ were successfully refined in a monoclinic double
 116 perovskite structure with space group $P2_1/n$ (see Supporting
 117 Information) similar to that reported for $\text{Pb}_2\text{CoOsO}_6$,¹⁶
 118 $\text{Pb}_2\text{MnReO}_6$,³² and $\text{Pb}_2\text{CoTeO}_6$.³³ Due to the weak X-ray
 119 scattering power of O (especially in the presence of strong
 120 scatterers Os and Pb), complementary NPD data were used to
 121 confirm this nuclear structure at 98 K. NPD data collected at
 122 98 K (above T_N) are consistent with the SPXD results and can
 123 be well fitted with a model of $P2_1/n$ symmetry (see the
 124 Supporting Information). The Ni and Os ions occupy 2a and
 125 2b sites, respectively. Allowing for antisite disorder in the
 126 model during the refinement (with constraints to maintain
 127 stoichiometry) revealed complete B-site ordering (100(6)%).
 128 Refinement of the occupancies of Pb and O sites (with a single
 129 global ADP) indicated that the material is very close to
 130 stoichiometric ($\text{Pb}_{1.940(1)}\text{NiOsO}_{5.90(1)}$). This stoichiometry was
 131 assumed for further analysis. Trace amounts of PbO_2 and NiO
 132 impurities were identified and included in the refinement (no
 133 Os impurity was detected). Final refined atomic parameters
 134 and selected bond lengths and angles are summarized in the
 135 Supporting Information. The bond valence sum calcula-
 136 tions^{34,35} support the nominal Ni^{2+} and Os^{6+} oxidation states

(see the Supporting Information). The refined crystal structure 137
 is shown in Figure 1a, where the corner-linked NiO_6 and OsO_6 138 f

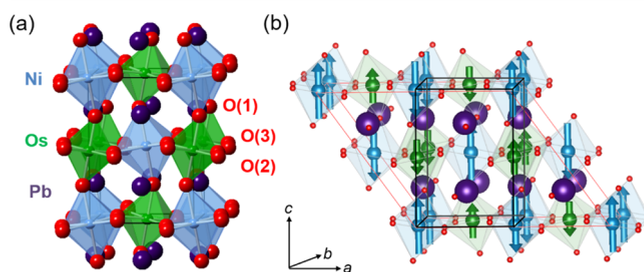


Figure 1. Illustration of the nuclear (a) and magnetic structures (b) of $\text{Pb}_2\text{NiOsO}_6$ at 1.5 K from Rietveld refinement using NPD data; Pb, Ni, Os, and O sites are shown in purple, blue, green, and red, respectively; Ni and Os moments are shown by arrows. The nuclear unit cell is shown by solid black lines and the larger, monoclinic $P2_1/n$ magnetic unit cell by solid red lines.

octahedra are ordered in the rock salt manner. The 139
 interoctahedral Ni–O–Os bond angles are 159.20(9), 140
 161.3(4), and 160.5(4) $^\circ$, which significantly deviate from 141
 180 $^\circ$ and imply substantial rotations of BO_6 octahedra. 142

3.2. Electrical and Magnetic Properties. The temper- 143
 ature dependence of resistivity (ρ) data decreases with cooling, 144
 as shown in Figure 2a, and shows the metallic nature of 145 f
 $\text{Pb}_2\text{NiOsO}_6$. The temperature dependence of magnetic 146
 susceptibility (χ) data shows a typical AFM transition with a 147
 peak at 58 K (see Figure 2b), which indicates the Néel 148

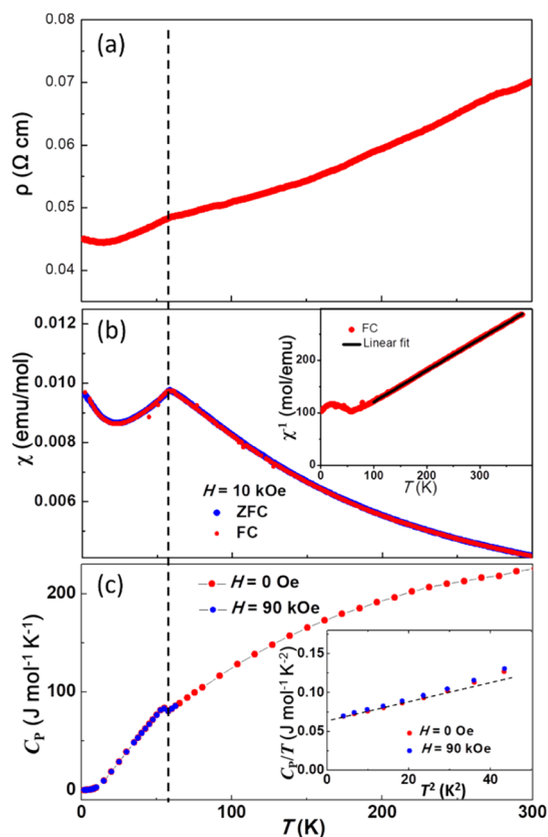


Figure 2. (a) Temperature-dependent resistivity, (b) temperature-dependent magnetic susceptibility, and (c) temperature-dependent specific heat of $\text{Pb}_2\text{NiOsO}_6$.

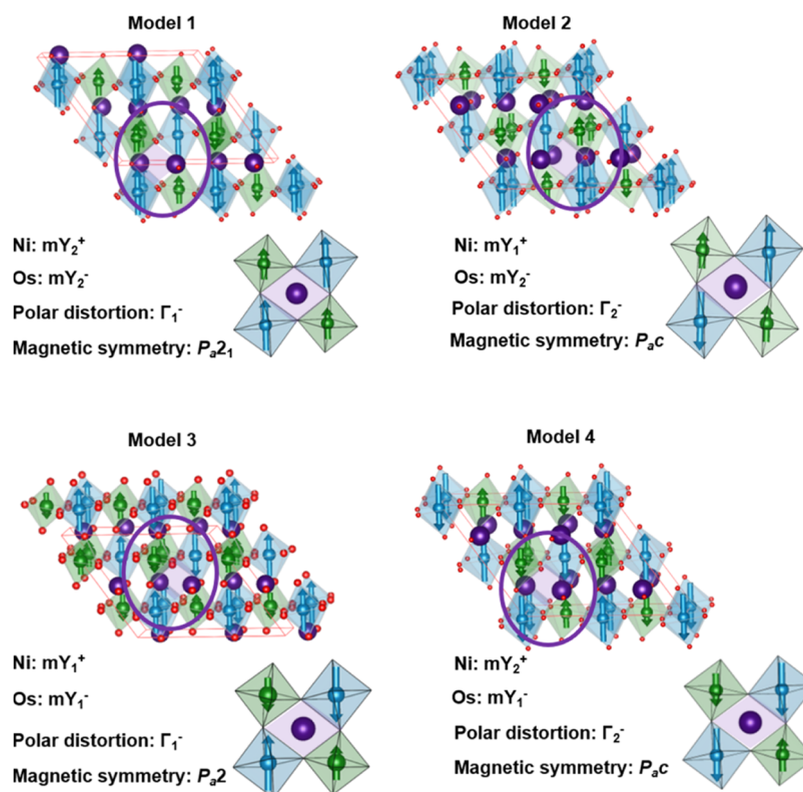


Figure 3. Four magnetic structures that result from the $k = (1/2 \ 0 \ 1/2)$ magnetic propagation vector for the $P2_1/n$ nuclear structure for $\text{Pb}_2\text{NiOsO}_6$ with a magnetic order on both Ni (blue) and Os (green) sites. To help visualize the difference between the four magnetic structures, the magnetic order around a PbO_{12} site (purple) viewed down the $[-1 \ 1 \ 0]$ direction of the nuclear unit cell is highlighted for each model. (Note that for model 2, the magnetic order around the PbO_{12} site is viewed along $[0 \ -1 \ 0]$ of the magnetic unit cell to show the magnetic moments about the same point in the nuclear structure).

149 temperature (T_N). The long-range AFM transition was further
 150 confirmed by specific heat data which display a λ -type anomaly
 151 at T_N (see Figure 2c). The χ^{-1} versus T data above the T_N
 152 show the Curie–Weiss behavior. Fitting the CW law to the
 153 data between 100 and 380 K results in an effective moment
 154 (μ_{eff}) of $3.66 \mu_B$ per formula unit (f.u.) and a Weiss
 155 temperature (θ_W) of -102 K. The obtained effective moment
 156 is comparable to that in other $\text{Ni}^{2+}\text{–Os}^{6+}$ double perovskites,
 157 $3.44 \mu_B/\text{f.u.}$ for $\text{Sr}_2\text{NiOsO}_6$ ³⁶ and $3.46 \mu_B/\text{f.u.}$ for
 158 $\text{Ba}_2\text{NiOsO}_6$.³⁷ These values are smaller than the spin-only
 159 moments of $4.0 \mu_B$ per formula unit for the Ni^{2+} ($3d^8$; $S = 1$)
 160 and Os^{6+} ($5d^2$; $S = 1$), which may be due to the SOC of Os^{6+} .
 161 The negative θ_W corroborates that AFM interactions are
 162 dominant in $\text{Pb}_2\text{NiOsO}_6$, which is consistent with the AFM
 163 order. The low-temperature part of specific heat data is plotted
 164 in the C_p/T versus T^2 , and the lowest temperature part can be
 165 characterized by an approximated Debye model ($C_p/T = \gamma +$
 166 $\beta_0 T^2$). The fitting gives a Sommerfeld coefficient (γ) of 63.5 mJ
 167 $\text{mol}^{-1} \text{K}^{-2}$. The large γ value is consistent with the metallic
 168 nature of $\text{Pb}_2\text{NiOsO}_6$. The deviation from linearity above 30
 169 K^2 could be due to the lattice contribution.

170 **3.3. Magnetic Structure.** To study the magnetic structure
 171 of $\text{Pb}_2\text{NiOsO}_6$, NPD data were collected from 1.5 to 98 K. On
 172 cooling below 57 K, additional reflections were observed in
 173 NPD patterns, which increased smoothly in intensity on
 174 cooling (see the Supporting Information). These were
 175 consistent with magnetic ordering described by magnetic
 176 propagation vector $k = (1/2 \ 0 \ 1/2)$. As described for the
 177 double perovskites $\text{Pb}_2\text{CoOsO}_6$ ¹⁶ and for the k_1 propagation
 178 vector for $\text{Sc}_2\text{NiMnO}_6$ ³⁸ there are four irreps associated with

the magnetic propagation vector $k = (1/2 \ 0 \ 1/2)$: mY_1^\pm and
 179 mY_2^\pm . The mY_n^+ (mY_n^-) irreps describe magnetic order on the
 180 Ni (Os) sites only. Magnetic susceptibility and heat capacity
 181 measurements for $\text{Pb}_2\text{NiOsO}_6$ (Figure 2b,c) and the evolution
 182 of magnetic Bragg intensity in NPD data collected on cooling
 183 (see the Supporting Information) suggest a single magnetic
 184 ordering transition which could result from one of three
 185 possible scenarios: (1) only Ni^{2+} moments order at T_N ; (2)
 186 only Os^{6+} moments order at T_N or (3), both Ni^{2+} and Os^{6+}
 187 moments order simultaneously at T_N . As described for related
 188 double perovskites,^{36,38–40} the magnetic moments on the two
 189 B sites are strongly correlated in refinements and NPD cannot
 190 unambiguously distinguish between these three scenarios.
 191 However, given the strong coupling between nearest Ni^{2+} and
 192 Os^{6+} ions in $\text{Ca}_2\text{NiOsO}_6$, it seems most likely that both Ni and
 193 Os sublattices order magnetically below T_N .³¹ Experiments on
 194 $\text{Pb}_2\text{CoOsO}_6$ including muon spin rotation experiments support
 195 magnetic ordering of both Co^{2+} and Os^{6+} moments,¹⁶
 196 consistent with our analysis for $\text{Pb}_2\text{NiOsO}_6$. Mode inclusion
 197 analysis^{41,42} using 1.5 K data suggested that the greatest
 198 improvement in fit was obtained with moments on the Os sites
 199 described by mY_2^- (R_{wp} decreased from 7.71% for a
 200 nonmagnetic model to 6.42% for the mY_2^- model) with Os
 201 moments close to the $[0 \ 0 \ 1]$ direction of the $P2_1/n$ nuclear
 202 unit cell. Magnetic ordering described by the mY_2^- irrep on the
 203 Os sites and the mY_1^+ (mY_2^+) irrep on the Ni sites breaks
 204 inversion symmetry, and the ferroelectric mode Γ_2^- (Γ_1^-) is
 205 coupled to both magnetic order parameters, allowing polar
 206 displacements in the ac plane ($[0 \ 1 \ 0]$ direction) of the $P2_1/n$
 207 nuclear unit cell. These two possible structures are very similar, 208

209 and our NPD data do not allow us to confirm which is more
 210 appropriate to describe the low-temperature nuclear and
 211 magnetic structure of $\text{Pb}_2\text{NiOsO}_6$. Attempts to investigate
 212 the polar distortions using both NPD analysis and electron
 213 diffraction were not successful, suggesting that these
 214 distortions are very subtle. Consistent with DFT calculations
 215 (see below), the $mY_2^- mY_1^+ \Gamma_2^-$ model was assumed for all
 216 further analysis. This magnetic structure is described by the
 217 monoclinic unit cell of symmetry P_2c which is related to the
 218 $P2_1/n$ nuclear unit cell by the basis vectors $(-2\ 0\ 0)$ $(0\ -1\ 0)$
 219 $(1\ 0\ 1)$ with an origin shift of $(0\ 1/4\ 0)$ (see Figure 1b). Given
 220 the complexity of the system, the moments on Ni and Os sites
 221 were constrained to be collinear (as observed in related
 222 systems^{16,31,34,38–40,43}) and the moments on Os sites were
 223 constrained to be eight times smaller than those on Ni sites, as
 224 might be expected for Ni^{2+} (d^8) and Os^{6+} (d^2) with significant
 225 covalent bonding.^{16,31,36,39} Allowing the moment direction to
 226 refine freely gave moments close to $[0\ 0\ 1]$ of the $P2_1/n$
 227 nuclear unit cell, and constraining the moments to lie exactly
 228 along this direction gave a similar fit (R_{wp} was the same to
 229 three decimal places) and was used in subsequent analysis.
 230 Allowing ADPs to refine anisotropically did not give a
 231 significant improvement in fit and ADPs were found to be
 232 fairly isotropic. The final refinement profiles and parameters
 233 are given in the Supporting Information.

234 Sequential Rietveld refinements were carried out using NPD
 235 data collected on warming to study the evolution of nuclear
 236 and magnetic structures. The 1.5 K model described above was
 237 used, and this sequential analysis suggested a fairly smooth
 238 expansion of the structure on warming (see the Supporting
 239 Information).

240 **3.4. First-Principles Calculations.** DFT calculations were
 241 carried out to explore whether both Ni and Os moments are
 242 ordered in the magnetic phase of $\text{Pb}_2\text{NiOsO}_6$ and to
 243 differentiate between the possible magnetically ordered
 244 structures. First, the total energies were calculated for magnetic
 245 models with either AFM order on both Ni and Os sublattices
 246 or AFM order only on the Ni sublattice (see the Supporting
 247 Information). These calculations indicate that the model with
 248 the AFM order on both Ni and Os sublattices is 0.102 eV per
 249 formula unit more stable than that with only Ni-ordered
 250 moments. These calculations support the noncentrosymmetric
 251 AFM models in which both sublattices are ordered.

252 As discussed above, group theory calculations assuming the
 253 propagation vector $(1/2\ 0\ 1/2)$ and magnetic order on both
 254 Ni and Os sublattices (from magnetic irreps mY_1^+ and mY_2^+ on
 255 Ni sites, mY_1^- and mY_2^- on Os sites) give four possible
 256 isotropy subgroups (Figure 3). These models give comparable
 257 fits to the NPD data, and we are not able to unambiguously
 258 determine the magnetic ground state from our experimental
 259 work. Although the relative orientation of magnetic spins is
 260 similar in these four structures (they all have the $\uparrow\uparrow\downarrow\downarrow\uparrow\uparrow$
 261 sequence of moments on the Ni–Os chains along the $[0\ 0\ 1]$
 262 direction of the nuclear unit cell), the superposition of the
 263 magnetic order on the nuclear structure (with monoclinic
 264 symmetry arising from rotations of NiO_6 and OsO_6 octahedra)
 265 results in different final symmetries and, as a consequence, in
 266 different distortions (e.g., polar degrees of freedom, bond
 267 distances, and angles) and hence different macroscopic
 268 properties.^{44–46} Since the four models derive from different
 269 combinations of irreducible representations, these are distinct
 270 structures and not translational domains.

DFT calculations were carried out to differentiate between 271
 these similar magnetic structures and to determine the ground 272
 state. Calculations were carried out using the GGA(PBE) + 273
 SOC + U scheme to determine the energy of the four magnetic 274
 structures shown in Figure 3, and the relative energies are 275
 given in Table 1. Model 2 (described above from analysis of 276 11

Table 1. Total Energy and Magnetic Moment Calculated for $\text{Pb}_2\text{NiOsO}_6$ for Models Shown in Figure 3^a

	energy (meV/atom)	total moment (μ_B)	spin moment (μ_B)	orbital moment (μ_B)
model 1	0	Ni: 1.85 Os: 0.78	Ni: 1.70 Os: 0.99	Ni: 0.15 Os: -0.21
model 2	-0.36	Ni: 1.86 Os: 0.78	Ni: 1.70 Os: 0.99	Ni: 0.16 Os: -0.21
model 3	+0.18	Ni: 1.85 Os: 0.77	Ni: 1.70 Os: 0.99	Ni: 0.15 Os: -0.22
model 4	+0.42	Ni: 1.85 Os: 0.76	Ni: 1.70 Os: 0.99	Ni: 0.15 Os: -0.23

^aUnit of the energy is meV per atom and is calculated by dividing the
 DFT total energy by the number of atoms in the unit cell (40 for the
 magnetic unit cells shown in Figure 3). Model 1 is chosen as the
 reference energy. Details about magnetic moments are discussed in
 the main text.

NPD data, Figure 1b) is found to be the lowest in energy for 277
 calculations including SOC (Table 1). These results suggest 278
 that the ground state of $\text{Pb}_2\text{NiOsO}_6$ is best described by P_2c 279
 magnetic symmetry, with Γ_2^- polar degrees of freedom, 280
 consistent with the ground state reported for $\text{Pb}_2\text{CoOsO}_6$.¹⁶ 281
 Notice that the same ground state was found for $\text{Pb}_2\text{NiOsO}_6$ 282
 from calculations without accounting for SOC. Mode 283
 decomposition of the relaxed structures from these DFT 284
 calculations was carried out using ISODISTORT,²⁶ but the 285
 amplitudes of polar displacements were very small 286
 (≤ 0.00035); this is consistent with NPD and electron 287
 diffraction analysis, both unable to confirm these displace- 288
 ments. The subtlety of these polar distortions (and that they 289
 are secondary rather than primary order parameters) is born out 290
 by the fact that the two lowest energy structures (models 1 and 291
 2) allow different polar distortions, suggesting that these 292
 distortions play a minor role in giving the noncentrosymmetric 293
 ground-state structure. 294

The electronic structure of $\text{Pb}_2\text{NiOsO}_6$ for model 2 (total 295
 and partial) is shown in Figure 4. Since Ni and Os atoms have 296 14
 local magnetic moments (Table 1), they show local spin 297
 polarization, as shown in Figure 4b,c. These local spin 298
 polarizations are summed to be zero, that is, the net total 299
 magnetic moment is zero, reflecting that $\text{Pb}_2\text{NiOsO}_6$ is 300
 antiferromagnetic, as demonstrated in Figure 4a. 301

The major contributions to the total DOS around the E_F are 302
 attributed to the Os 5d orbitals in both spin channels, which 303
 hybridize strongly with the O 2p orbitals. Occupation numbers 304
 for Ni 3d and Os 5d are 7.81 and 3.80, respectively. The huge 305
 hybridization indicated between Os 5d and O 2p orbitals 306
 suggests Ni^{2+} and Os^{6+} formal oxidation states in $\text{Pb}_2\text{NiOsO}_6$, 307
 consistent with the magnetic susceptibility experiment. Spin 308
 and orbital moments for Ni are 1.70 and 0.16 μ_B , respectively, 309
 thereby giving a total magnetic moment of 1.86 μ_B per Ni. For 310
 Os, spin and orbital moments are 0.99 and -0.21 μ_B , 311
 respectively, where the minus sign indicates that the orbital 312
 moment is opposite the spin direction; thus, the total moment 313
 is 0.78 μ_B per Os. These calculated moments are comparable 314

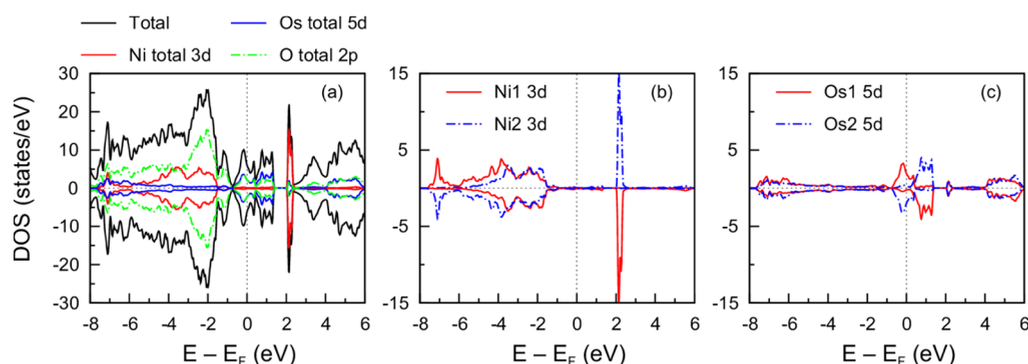


Figure 4. Total and partial density of states (PDOS) of $\text{Pb}_2\text{NiOsO}_6$ for model 2 from GGA + SOC + U calculation. (a) Black solid line corresponds to the total DOS. Red solid, blue solid, and green dashed dotted lines represent total Ni 3d, total Os 5d, and O 2p PDOS, respectively. (b) PDOS for each Ni 3d: Ni1 (Ni2) is presented for the spin majority as spin up (down). (c) PDOS for each Os 5d: Os1 (Os2) is presented for the spin majority as spin up (down). The positive and negative values in DOS correspond to spin up and down, respectively.

315 with those obtained from NPD analysis (see above). The
316 calculated γ is $7.1 \text{ mJ mol}^{-1} \text{ K}^{-2}$. This is much smaller than the
317 one obtained from the fitting of low-temperature specific heat
318 data ($63.5 \text{ mJ mol}^{-1} \text{ K}^{-2}$), which may be due to the fact that
319 DFT underestimates the electronic correlations in the
320 correlated systems, resulting in a relatively small γ value.

4. DISCUSSION

321 The A_2NiOsO_6 ($\text{A} = \text{Ca}^{2+}, \text{Sr}^{2+}, \text{Ba}^{2+}$, and now Pb^{2+}) oxides
322 adopt B-site ordered double perovskite structures and span a
323 range of properties, from insulating ($\text{A} = \text{Ca}^{31}$) to metallic (A
324 = Pb) and from ferromagnetic ($\text{A} = \text{Ba}$ at low temperatures³⁷)
325 to antiferromagnetic ($\text{A} = \text{Sr}^{36}$). While all these analogues
326 adopt the rocksalt ordering of NiO_6 and OsO_6 octahedra, the
327 degree of tilting of these octahedra increases with decreasing A
328 cation radius: $\text{Ba}_2\text{NiOsO}_6$ is cubic with 180° Ni–O–Os bond
329 angles; in tetragonal $\text{Sr}_2\text{NiOsO}_6$, octahedra are tilted around
330 the long axis, giving $180^\circ/166^\circ$ Ni–O–Os angles,³⁶ while
331 $\text{Ca}_2\text{NiOsO}_6$ adopts the monoclinic $P2_1/n$ structure (with
332 $a^-a^-c^+$ tilts) with Ni–O–Os angles of $\sim 151^\circ$.³⁶ $\text{Pb}_2\text{NiOsO}_6$
333 also adopts this $P2_1/n$ structure despite the ionic radius of Pb^{2+}
334 (1.49 \AA) being comparable to that of Sr^{2+} (1.44 \AA),⁴⁷ this
335 might in part be due to the inert pair Pb^{2+} ion favoring the
336 lower-symmetry coordination environment⁴⁸ possible in the
337 $P2_1/n$ structure: Pb occupies the 4e site of 1 symmetry in $P2_1/$
338 n compared with the higher-symmetry 4d site of -4 symmetry
339 in the $I4/m$ structure of $\text{Sr}_2\text{NiOsO}_6$.

340 The magnetic structure described here for $\text{Pb}_2\text{NiOsO}_6$ is of
341 the same symmetry as that reported for $\text{Pb}_2\text{CoOsO}_6$,¹⁶
342 although with a slightly different orientation of moments,
343 likely resulting from the different magnetic anisotropies of
344 Co^{2+} and Ni^{2+} ions in octahedral coordination environments.
345 In both $\text{Pb}_2\text{NiOsO}_6$ and $\text{Pb}_2\text{CoOsO}_6$, the magnetic order on
346 the Ni/Co and Os sublattices breaks inversion symmetry and
347 follows an $\uparrow\uparrow\downarrow\downarrow$ sequence along $[001]$ of the nuclear unit cell.
348 Magnetic ordering has been shown to break inversion
349 symmetry in other perovskites, including $\text{Sr}_2\text{NiMnO}_6$, but
350 with weak coupling between Ni and Mn sublattices,³⁸ in
351 contrast to Pb_2BOsO_6 ($\text{B} = \text{Co}, \text{Ni}$), which seem to have
352 collinear moments on both B and Os sublattices and a single
353 magnetic ordering transition. These observations are consistent
354 with strong couplings between Co/Ni and Os sublattices. This
355 $\uparrow\uparrow\downarrow\downarrow$ magnetic structure observed in $\text{Pb}_2\text{NiOsO}_6$ is signifi-
356 cantly different from those reported for other A_2NiOsO_6
357 double perovskites. Previous works have highlighted the

importance of both nearest-neighbor (likely FM) and next- 358
nearest-neighbor (likely AFM) interactions in these sys- 359
tems.^{31,49} The balance between these (competing) interactions 360
gives some magnetic frustration in $\text{Sr}_2\text{NiOsO}_6$ and makes the 361
magnetic structure of A_2NiOsO_6 phases very sensitive to bond 362
angles.³¹ 363

The symmetry requirements for magnetic order to break 364
inversion symmetry have been explored by Perez-Mato et al.⁵⁰ 365
and provide a recipe for designing new magnetoelectrics. If the 366
magnetic k vector is not compatible with the screw axes or 367
glide planes of the nuclear (paramagnetic) unit cell (when time 368
reversal symmetry is considered), then full magnetic order on a 369
lattice of magnetic atoms on special sites (of -1 symmetry) 370
will break the inversion symmetry, resulting in noncentrosym- 371
metric structures.⁵⁰ 372

The double perovskites considered here have rocksalt 373
ordering of B and B' cations on sites related by an origin 374
shift (and typically with symmetries including inversion 375
centers). If a single magnetic propagation vector k describes 376
the magnetically ordered phase, and full magnetic order is 377
expected on both B and B' sublattices, then, depending on k , 378
the irreps to describe the magnetic order on each sublattice 379
may be of opposite parity with respect to an inversion center at 380
the origin. These irreps couple to a noncentrosymmetric 381
distortion (Γ_x^-), breaking inversion symmetry. This is the case 382
described here for $\text{Pb}_2\text{NiOsO}_6$ for $k = (1/2 0 1/2)$ and also for 383
 $k = (-1 1/2 1/2)$. Likewise for a cubic double perovskite 384
 $\text{A}_2\text{BB}'\text{O}_6$ of $Fm\bar{3}m$ nuclear symmetry (with magnetic B and B' 385
ions on 4a and 4b sites, respectively), a magnetic $k = (1/2 1/2$ 386
 $1/2)$ would have a similar effect. This is also observed in the 387
hexagonal $\text{Ca}_3\text{CoMnO}_6$ ($R\bar{3}c$ nuclear symmetry, Mn^{3+} and 388
 Co^{3+} ions on 6a $(0 0 1/4)$ and 6b $(0 0 0)$ sites, respectively) 389
with magnetic $k = (0 0 0)$ giving the well-known $\uparrow\uparrow\downarrow\downarrow$ polar 390
magnetic structure.⁵¹ 391

If suitable cation-ordered structures with strong magnetic 392
coupling between the two sublattices (to favor them ordering 393
with the same magnetic k vector) can be identified, then new 394
magnetoelectrics might be designed if the magnetic exchange 395
interactions can be balanced to give the desired k vector. We 396
note that the improper ferroelectricity described here does not 397
require additional ordering of A-site cations (e.g., the 398
 $\text{AA}'\text{NiOsO}_6$ phases explored recently).⁵² 399

It is striking that both $\text{Pb}_2\text{NiOsO}_6$ and $\text{Pb}_2\text{CoOsO}_6$ ¹⁶ are 400
metallic, in contrast to the SOC Mott-insulating nature of 401
 $\text{Ca}_2\text{NiOsO}_6$ and $\text{Ca}_2\text{CoOsO}_6$,³¹ despite the structural 402

403 similarities between these Pb and Ca analogues. First, we note
404 that although $\text{Pb}_2\text{NiOsO}_6$ is metallic, its resistivity is several
405 orders of magnitude higher than that of $\text{Pb}_2\text{CoOsO}_6$ (300 K
406 resistivity is $\sim 0.07 \text{ } \Omega \text{ cm}$ (Figure 2a) and $\sim 3.5 \times 10^{-4} \text{ } \Omega \text{ cm}$
407 for Ni and Co¹⁶ analogues, respectively). This is similarly
408 observed for $\text{Ca}_2\text{NiOsO}_6$ and $\text{Ca}_2\text{CoOsO}_6$ and is ascribed to
409 the full occupancy of the $\text{Ni}^{2+} t_{2g}$ band reducing delocalization
410 of $\text{Os}^{6+} t_{2g}$ electrons.³¹

411 The half-metallic nature proposed for $\text{Sr}_2\text{NiOsO}_6$ results
412 from the partially-occupied Os t_{2g} states crossing the Fermi
413 level, with SOC broadening the Os 5d bands.⁵³ This scenario
414 can be applied to A_2NiOsO_6 (A = Ca, Pb), and our PDOS
415 calculations (Figure 4) are qualitatively similar to those
416 reported for $\text{Ca}_2\text{NiOsO}_6$ ³¹ (with Ni t_{2g} states below $\sim -2 \text{ eV}$
417 and a narrow band of Ni e_g states at $\sim 2 \text{ eV}$, with Os 5d and O
418 2p bands crossing E_F). However, the bandwidth in these
419 double perovskites is also influenced by Ni–O–Os bond
420 angles: in $\text{Ca}_2\text{NiOsO}_6$ with small Ca^{2+} ions, the Ni–O–Os
421 angles ($\sim 149.3\text{--}150.6^\circ$ at 4 K)³⁶ show much larger deviations
422 from the ideal 180° bond angles than in $\text{Pb}_2\text{NiOsO}_6$ (158.8--
423 161.0° at 1.5 K). The more distorted structure reported for the
424 Ca analogues is likely to decrease the orbital overlap and
425 bandwidth, giving wider band gaps than the less distorted Pb
426 analogues. This is consistent with the insulating and more
427 localized nature of $\text{Ca}_2\text{NiOsO}_6$ and its higher magnetic
428 ordering temperature (158 K, compared with $T_N = 58 \text{ K}$ for
429 $\text{Pb}_2\text{NiOsO}_6$).

5. CONCLUSIONS

430 A new 5d oxide $\text{Pb}_2\text{NiOsO}_6$ was synthesized under high
431 pressure. $\text{Pb}_2\text{NiOsO}_6$ crystallizes in a monoclinic double
432 perovskite structure with a centrosymmetric space group
433 $P2_1/n$ at room temperature. $\text{Pb}_2\text{NiOsO}_6$ is metallic down to 2
434 K and displays an AFM transition with $T_N = 58 \text{ K}$. $\text{Pb}_2\text{NiOsO}_6$
435 is a new example of AFM metallic oxides with three-
436 dimensional crystal and electronic structures. NPD and DFT
437 calculations indicate that both the Ni and Os moments are
438 ordered below T_N , breaking inversion symmetry, which is
439 similar to recently reported $\text{Pb}_2\text{CoOsO}_6$.^{16,17} The magnetically
440 driven loss of center of symmetry is similar to the type-II
441 multiferroics. The discovery of 5d oxides $\text{Pb}_2\text{NiOsO}_6$ together
442 with $\text{Pb}_2\text{CoOsO}_6$ establishes a new class of noncentrosym-
443 metric AFM metallic oxides.

ASSOCIATED CONTENT

Supporting Information

445 The Supporting Information is available free of charge at
446 <https://pubs.acs.org/doi/10.1021/acs.chemmater.1c01032>.

448 Refined room-temperature SXR D pattern and the
449 corresponding crystal structure information, refined
450 PND data at 98 and 1.5 K and the corresponding
451 crystal structure information, evolution of nuclear and
452 magnetic structures with temperature for $\text{Pb}_2\text{NiOsO}_6$
453 from sequential refinements using NPD data, and DFT
454 calculations of $\text{Pb}_2\text{NiOsO}_6$ with different magnetic states
455 (PDF)

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Notes

The authors declare no competing financial interest.

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