Antiferromagnetic $S = 1/2$ Spin Chain Driven by $p$- Orbital Ordering in CsO$_2$

Syarif Riyadi, Baomin Zhang, Robert A. de Groot, Antonio Caretta, Paul H. M. van Loosdrecht, Thomas T. M. Palstra, and Graeme R. Blake

1Zernike Institute for Advanced Materials, University of Groningen, Nijenborgh 4, 9747 AG Groningen, The Netherlands
2Radboud University, Institute for Molecules and Materials, Heyendaalseweg 135, 6525 AJ Nijmegen, The Netherlands

We demonstrate, using a combination of experiment and density functional theory, that orbital ordering drives the formation of a one-dimensional (1D) $S = 1/2$ antiferromagnetic spin chain in the 3D rocksalt structure of cesium superoxide (CsO$_2$). The magnetic superoxide anion (O$_2^-$) exhibits degeneracy of its 2$p$-derived molecular orbitals, which is lifted by a structural distortion on cooling. A spin chain is then formed by zigzag ordering of the half-filled superoxide orbitals, promoting a superexchange pathway mediated by the $p_z$ orbitals of Cs$^+$ along only one crystal direction. This scenario is analogous to the 3$d$-orbital-driven spin chain found in the perovskite KCuF$_3$ and is the first example of an inorganic quantum spin system with unpaired $p$ electrons.

DOI: 10.1103/PhysRevLett.108.217206
PACS numbers: 75.25.Dk, 71.20.−b, 75.10.Pq, 78.30.−j
served by Zumsteg et al. The original formulation for a expression reported by Feyerherm fitted the susceptibility maximum using an alternative ex-

The dominant feature in the magnetic susceptibility of polycrystalline CsO$_2$ is a broad maximum centered at $\sim$28 K [Fig. 1(a)]. A similar feature was previously observed by Zumsteg et al. [10], who remarked only that considerable magnetic order remained above $T_N = 9.6$ K.

Such a maximum is a typical indication of low-dimensional magnetic ordering [19]. The spins in AO$_2$ can rotate in the plane perpendicular to the molecular axis, but EPR shows only moderate anisotropy of the $g$ tensor [11,20]; thus, we can treat CsO$_2$ as an isotropic (Heisenberg) spin system. The susceptibility maximum was fitted using the $S = 1/2$ square-lattice Heisenberg model formulated by Keith et al. [21]. There are three variable parameters in this model: the exchange constant for a particular crystal direction $J/k_B$, the ratio $\alpha$ between two orthogonal exchange constants on a square lattice ($J' = \alpha J$), and the Curie constant. For 2D AFM ordering, $J = J'$ and $\alpha = 1$; for an AFM spin chain, $\alpha = 0$. This model can also be applied to 2D systems incorporating mixed AFM and FM interactions, for which $-1 \leq \alpha < 0$. The best fit to our data is shown in Fig. 1(a), which yielded $J/k_B = 20.27$ K, $\alpha = 0$, and $C = 0.325$. The zero value of $\alpha$ implies that CsO$_2$ is essentially an AFM spin chain. Fits performed with fixed nonzero positive or negative values of $\alpha$ were visibly worse. We also fitted the susceptibility maximum using an alternative expression reported by Feyertherm et al. [22], based on the original formulation for a $S = 1/2$ Heisenberg AFM chain by Bonner and Fisher [19]. We obtained $J/k_B = 20.0$ K, in good agreement with the model of Keith et al. [21]. Both models fitted the data equally well between 15 K and 70 K, at which a small anomaly associated with a structural transition (see below) is apparent. The spin chain appears to persist in applied fields up to at least 40 kOe [18]. A Curie-Weiss fit to the inverse susceptibility above 150 K [Fig. 1(b)] yielded an effective moment of 2.03 $\mu_B$. This is significantly higher than the expected spin-only value for $S = 1/2$ and is consistent with previous reports on CsO$_2$ and other AO$_2$ [10,20], suggesting unquenched orbital momentum. The fitted Weiss constant of $\theta = -40.3$ K, which corresponds to $\sim 47^\circ$, is more evidence for the quantum AFM nature of CsO$_2$.

Long-range 3D AFM ordering in CsO$_2$ was previously reported below $T_N = 9.6$ K on the basis of specific heat, magnetic susceptibility, and EPR measurements [11,10]. We observe a magnetic susceptibility anomaly at $\sim 9.5$ K [Fig. 1(b)] and the spin chain models no longer fit the data below $\sim 15$ K, supporting a transition to 3D ordering. The low-temperature susceptibility tail below 7 K is likely due to residual spins; a fit to the Curie law yielded a concentration of $\sim 8\%$ $S = 1/2$ disordered spins.

At room temperature, the XRPD pattern of CsO$_2$ is consistent with the previously reported tetragonal average structure (space group $I4/mmm$) [23]. However, Raman spectroscopy (see below) demonstrates that the true symmetry is lower. An impurity phase (CsOH,H$_2$O, $\sim 20\%$ by weight) was also present but does not contribute to the magnetic response. Splitting of the CsO$_2$ XRPD peaks with $h \neq k$ was observed below 70 K (see Fig. 2), consistent with a transition to orthorhombic symmetry. However, Rietveld refinements were rather insensitive to the oxygen positions and we were unable to determine details of the low-temperature structure. An earlier XRD study on CsO$_2$ single crystals reported a tetragonal-to-orthorhombic transition at the higher temperature of 190 K [23], below which incommensurate satellite reflections also appeared. No such peaks were apparent in our data. It was previously reported that different solution-grown CsO$_2$ single crystals from the same batch did not always exhibit a
tetragonal-orthorhombic transition [24], which suggests a sensitive dependence of the structure on oxygen stoichiometry. The evolution of the lattice parameters with temperature of our sample is shown in Fig. 2(c). Apart from the phase transition at ~70 K, the other noteworthy feature is a decrease of the c parameter above 200 K. This might be due to increasing librational freedom of the anions in the a and b directions on heating. The c axis also shortens on cooling below ~100 K, suggestive of coherent anion tilting away from the axis. The refinement of anisotropic microstrain peak-broadening parameters, using the Stephens formulation [25], was necessary at all temperatures to fit the peak profiles. The $S_{400}$ and $S_{040}$ parameters were much larger than $S_{004}$ [Fig. 2(d)], indicating either a degree of disorder or smaller coherently diffracting domain size in the pseudotetragonal basal plane.

Raman spectroscopy provided greater insight into the crystal structure. For the tetragonal average structure, three Raman active modes are apparent [Fig. 3(a)]. The stretching mode of $\text{O}_2^-$ is observed at 1134 cm$^{-1}$, which agrees well with the previously reported 1137 cm$^{-1}$ [26]. The broad peaks at 75 cm$^{-1}$ and 205 cm$^{-1}$ are probably a phonon mode involving the Cs cations and a $\text{O}_2^-$ librational mode, respectively (see below). The presence of the 75 cm$^{-1}$ mode is incompatible with $I4/mmm$ symmetry [18] and is evidence that the local symmetry is lower, as previously inferred from single-crystal XRD measurements [23].

Figure 3(b) shows the Raman spectrum at 4.5 K. Four peaks are apparent at low wave numbers, the origin of which was investigated by optimizing the orthorhombic structure of CsO$_2$ using DFT calculations without symmetry constraints. By using a supercell doubled along all three axes and the lattice parameters determined by XRPD at 20 K, we obtained two degenerate solutions. In the first structural solution, the anions are tilted by ~5° from c toward the basal plane diagonal, as shown in Fig. 4(b). The tilt directions are staggered along a and uniform along a, corresponding to a doubled b axis. The tilting of anions in the $z = 1/2$ layer of the supercell (which is the same as that in the $z = 1/4$ layer) is opposite to the tilting of anions in the $z = 0$ layer (the same as that in the $z = 3/4$ layer). Thus, a staggered pattern is formed when viewed along c and the c axis is also doubled. Within the ab plane there are staggered shifts of the Cs cations in the a direction, such that along b the cations form a zigzag arrangement. This is likely a consequence of the staggered anion tilts. In the second structural model the $z = 0$ and $z = 1/2$ layers are identical to those in the first structure, but the patterns of Cs shifts and anion tilts in these two layers are rotated by 90° in the $z = 1/4$ and $z = 3/4$ layers, respectively, such that the tilts become staggered along a rather than along b and the a axis is also doubled. This unit cell would be metrically tetragonal because overall there is no structural anisotropy in the ab plane. However, we observe a metrically orthorhombic cell (Fig. 2); thus, we favor the first model. Calculations of the Raman modes were carried out in VASP using the $a \times 2b \times 2c$ supercell [18]. Four Raman modes at low wave numbers were obtained: 66 cm$^{-1}$, 67 cm$^{-1}$, 75 cm$^{-1}$, and 200 cm$^{-1}$, which agree well with the experimental spectrum. The first three modes are associated with interlayer motions of the Cs cations along the c direction [18], and the broad feature observed at 209 cm$^{-1}$ is the librational mode ("swing-like" motion) of $\text{O}_2^-$. Finally, the calculated $\text{O}_2^-$ stretching mode at 1120 cm$^{-1}$ agrees well with the observed mode at 1135 cm$^{-1}$. Figure 3(c) shows the temperature dependence of the three low wave-number Cs modes, which merge to become a single broad peak.

![Figure 3](https://example.com/figure3.png)

**FIG. 3** (color online). Raman spectra of CsO$_2$ at (a) 110 K and (b) 4.5 K. (c) Raman modes associated with the interlayer motion of Cs and (d) librational mode of $\text{O}_2^-$ as a function of temperature.

![Figure 4](https://example.com/figure4.png)

**FIG. 4** (color online). (a) Partial oxygen DOS for anions A and B along the b axis in the $z = 0$ layer below 70 K. Because of the AFM configuration, the spin-up and spin-down DOS for anions in the $z = 1/4$ layer is reversed compared to the DOS shown here. (b) ab-plane view of the optimized structure of CsO$_2$ below 70 K showing anion tilting and ordering of the half-occupied $\pi_d^*$ and $\pi_u^*$ orbitals.
above the 70 K structural phase transition. The $\text{O}_2^-\text{librational mode broadens and weakens between 65 K and 110 K but does not disappear completely [Fig. 3(d)].}$

The phase transition at $\sim 70$ K results in OO, which is apparent from the site-projected partial oxygen DOS shown in Fig. 4(a) for two adjacent anions $A$ and $B$ along the $b$ axis of the optimized structure in Fig. 4(b). The four peaks in the DOS for each anion correspond to the spin-up and spin-down states of the two $\pi^*$ orbitals, three of which are fully occupied and one of which, above the Fermi level ($E_f$), is empty. The states in the energy range $-1$ to 0 eV correspond to a filled $\pi_c^*$ orbital for dimer $A$ and a filled $\pi_c^*$ orbital for dimer $B$. Here, $x$ and $y$ refer to the coordinate system indicated by axes $x'$ and $y'$ in Fig. 4(b), which are at $\sim 45^\circ$ to the unit cell axes $a$ and $b$. For any given dimer, the $\pi_c^*$ orbital is fully occupied on two nearest-neighbor dimers within the pseudotetragonal plane and the $\pi_c^*$ orbital is fully occupied on the other two. This corresponds to a zigzag configuration of alternating half-filled $\pi_c^*$ and $\pi_c^*$ orbitals along the $b$ axis, as shown in Fig. 4(b) (the anion axes tilt by $\sim 5^\circ$ from the $c$ axis and are almost perpendicular to the plane of the picture.)

The lobes of the magnetic orbitals do not point toward each other; thus, direct exchange will be weak and possible superexchange pathways via Cs must be examined. The occupied Cs $p_z$ orbitals have the correct orientation to mediate coupling along the pathways marked by red arrows in Fig. 4(b), whereas no obvious superexchange pathway is available in the perpendicular direction. The Cs $p_z$ orbitals lie at energies between $-8.5$ eV and $-5$ eV with respect to $E_f$ [18]. The sign of the superexchange along $b$ is AFM according to the Kugel-Khomskii picture [27]. The unit cell is doubled in the $c$ direction: when site $A$ at $z = 0$ in Fig. 4(b) has a half-occupied $\pi_c^*$ orbital, the corresponding site at $z = 1/2$ has a half-occupied $\pi_c^*$ orbital. Linear superexchange pathways along $c$ of the type $\text{O}_2^-\text{Cs} p_z^\text{O}_2^-\text{Cs} p_z^\text{O}_2^-\text{Cs} p_z$ are thus prevented. The result is an AFM spin chain in the $b$ direction.

The anion tilting and OO configuration in Fig. 4(b) is analogous to that calculated for $\text{KO}_2$ by Nandy et al. [15]. Although the interatomic distances along the $b$ direction in $\text{KO}_2$ are shorter, the magnetic susceptibility does not suggest low-dimensional ordering [10]. It is possible that the superexchange pathway along $b$ is disrupted by the much greater anion tilting in $\text{KO}_2$. Similarly, no spin chain seems to form in $\text{RbO}_2$, despite the crystal structure being very similar to $\text{CsO}_2$ [2]. This may be due to a different ground state OO configuration, calculated to be staggered along both basal plane axes with orbital lobes pointing between the Rb$^+$ cations [14]. We also note that the relevant $p$-levels of Rb and K lie significantly deeper than for Cs [28]. The marcasite phase of $\text{Na}_2$ below 196 K might exhibit a spin chain along the $c$ axis due to strong AFM direct exchange between anions [9]. A coupling constant of $J = -370$ K was calculated for the situation where the lobes of half-occupied orbitals on nearest neighbors point toward each other [29]. The magnetic susceptibility should thus rise with temperature without reaching the typical broad maximum associated with a spin chain; this result has been experimentally observed [10,29], but the data were not fitted to any model and OO has not been confirmed.

Although several examples of AFM $p$-electron spin chains are known in organic charge-transfer salts, these are a consequence of low structural dimensionality [30]. The closest analogy to $\text{CsO}_2$ is found in the $d$-electron spin chain $\text{KCuF}_3$ [31]. In this tetragonal perovskite, the distances between $\text{Cu}^{2+}$ cations are similar along the three principal crystallographic axes (4.14, 4.14, and 3.93 Å). A Jahn-Teller distortion removes the $\text{Cu}^{2+}$ orbital degeneracy, breaking the octahedral symmetry and favoring an alternating occupation of half-occupied $d_{x^2-y^2}$ and $d_{z^2}$ orbitals in the $ab$ plane. The lobes of the half-occupied orbitals on neighboring cations are orthogonal to each other in the $ab$ plane; thus, superexchange between nearest neighbors is extremely weak. In contrast, the lobes of both orbitals overlap with the fluorine $p_z$ orbitals in the $c$ direction, allowing superexchange along a linear pathway. An AFM spin chain is formed with $J_z/J_a = -100$ [32]. At $\sim 40$ K, the spin chain gives way to 3D $A$-type AFM order [33]. An explanation for this has recently been proposed by Lee et al. [34], who calculated that the half-occupied orbitals can more accurately be described as hybrids of $d_{x^2-y^2}$ and $d_{z^2}$, two distinct and almost degenerate states which exist in the spin-chain regime. One hybrid state favors AFM order in-plane and the other FM order in-plane, and rapid fluctuations between the two states occur. A structural distortion involving short-range-ordered rotations of the octahedra takes place at $\sim 40$ K, stabilizing the AFM hybrid state. In $\text{CsO}_2$, the 3D AFM ordering that sets in below 9.6 K requires the availability of superexchange pathways along both $a$ and $c$. Our Raman data (Fig. 3) reveal no obvious structural change down to 4.5 K; thus, the OO configuration and alternative superexchange pathways should be studied in more detail.

In conclusion, our observation of a $S = 1/2$ AFM spin chain in $\text{CsO}_2$ that is driven by OO is the clearest demonstration yet of correlated electron physics at play in an inorganic, open $p$-shell system. Our study shows that the concepts of orbital physics developed for transition metals are also valid for main-group systems with orbital degeneracy.

We thank G.A. de Wijs, S. Giriya, and P. Mahadevan for useful discussions. This Letter is part of the research program of the Foundation for Fundamental Research on Matter (FOM), which is financially supported by the Netherlands Organisation for Scientific Research (NWO).