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Iridates from the molecular side

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New exotic phenomena have recently been discovered in oxides of paramagnetic Ir^{4+} ions, widely known as 'iridates'. Their remarkable properties originate from concerted effects of the crystal field, magnetic interactions and strong spin-orbit coupling, characteristic of 5d metal ions. Despite numerous experimental reports, the electronic structure of these materials is still challenging to elucidate, and not attainable in the isolated, but chemically inaccessible, $[IrO_6]^{8-}$ species (the simplest molecular analogue of the elementary $\{IrO_6\}^{8-}$ fragment present in all iridates). Here, we introduce an alternative approach to circumvent this problem by substituting the oxide ions in $[IrO_6]^{8-}$ by isoelectronic fluorides to form the fluorido-iridate: $[IrF_6]^{2-}$. This molecular species has the same electronic ground state as the $\{IrO_6\}^{8-}$ fragment, and thus emerges as an ideal model for iridates. These results may open perspectives for using fluorido-iridates as building-blocks for electronic and magnetic quantum materials synthesized by soft chemistry routes.

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he 5d elements of the periodic table possess singular properties including a strong coupling of the electronic spin to its orbit as well as spatially extended valence orbitals leading to a reduced electronic repulsion and large effect on the ligand field. These intrinsic characteristics have been experimentally or theoretically demonstrated to be responsible for new exotic states of matter such as spin-orbit (SO) Mott insulators^{1,2}, topological insulators³⁻⁵, super-conductors⁶⁻⁸, spin-liquids and -ices⁹⁻¹¹ and for quantum metal-insulator transitions¹². The common denominator in these promising materials, collectively referred to as 'iridates' (herein named oxido-iridates), is the presence of octahedrally coordinated Ir^{IV} ions (Fig. 1a,b) featuring a t_{2g}^{5} electronic configuration. The half-filled $j_{\rm eff}=1/2$ level (Fig. 1c)¹³, resulting from the SO coupling, is at the origin of a narrow band gap responsible for the abovementioned phenomena. In all oxido-iridates including Sr₂IrO₄, the archetypal SO Mott insulator², corner- or edgesharing $\{IrO_6\}^{8-}$ distorted octahedra are present (Fig. 1a). While magnetic Ir^{IV} - Ir^{IV} interactions and their implications for the physical properties have been studied extensively in oxidoiridates 14,15, the intrinsic magnetic properties 16 of the elemental {IrO₆}^{8 -} moiety are consistently masked by the same long-range interactions. Birol and Haule 17 recently suggested that the design of materials incorporating isolated Ir^{IV} octahedra should facilitate smaller bandwidths and promote Mott ground states. If this strategy is developed to its logical end, a discouraging result is obtained as the isolation of the $[IrO_6]^{8-}$ ion (Fig. 1b) is chemically impossible. However for the realization of dimensionally reduced oxido-iridates, the fluoride ion appears as an ideal substitute to oxide being isoelectronic with comparable chemical and physical characteristics, but, importantly, with a reduced charge. Along this idea, we present the synthesis of molecular fluorido-iridates incorporating spatially isolated $[IrF_6]^{2-}$ units. Their local magnetic properties are probed by X-ray magnetic circular dichroism (XMCD) spectroscopy demonstrating that the $[IrF_6]^2$ and $[IrO_6]^8$ units possess virtually identical electronic ground states as suggested by theory¹⁷. Thus this fluorido-iridate moiety and its intrinsic properties, experimentally determined in this work, can be confidently used to model and emulate the basic {IrO₆}⁸ unit in oxido-iridates.

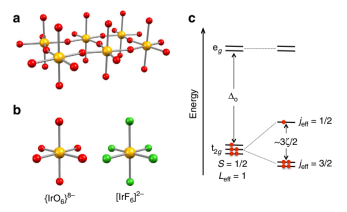


Figure 1 | Oxido-iridates versus fluorido-iridates. Ball and stick representations of (a) the oxido Ir^{IV} -based layer in Sr_2IrO_4 , (b) its smallest $\{IrO_6\}^8$ — unit and the isoelectronic fluoride counterpart, $[IrF_6]^2$ —. (c) Energy level diagram for Ir^{IV} (5d⁵; t_{2g}^5 electronic configuration). The octahedral ligand field splits the 5d orbitals into e_g and t_{2g} levels, and SO coupling further lifts the degeneracy of the t_{2g} levels into filled $j_{\rm eff}=3/2$ and half-filled $j_{\rm eff}=1/2$ levels. Note that this one-electron picture is equivalent to the SO splitting of the ${}^2T_{2g}$ (O_h) term into an $J_{\rm eff}=1/2$ ground state and excited $J_{\rm eff}=3/2$ state.

Results

Syntheses and structures. Highly water-soluble Na₂[IrF₆] was obtained by direct F₂ fluorination of Na₂[IrCl₆] · 6H₂O at elevated temperature (200 °C) and characterized by powder X-ray diffraction 18-20. Addition of a solution of PPh₄Cl (PPh₄⁺ = tetraphenylphosphonium(V) cation) to an aqueous $[IrF_6]^{2-}$ solution yielded $(PPh_4)_2[IrF_6] \cdot 2H_2O$ (1, Fig. 2a and Supplementary Fig. 1). Electrospray mass spectrometry on 1 indicated complete absence of any hydrolysis products such as [IrF₅(OH)]^{2 -} (Supplementary Fig. 2). 1 was subsequently used to synthesize Zn(viz)₄[IrF₆] (2, Fig. 2b and Supplementary Fig. 3; viz = 1-vinylimidazole). These compounds incorporate a low symmetry, but close-to-octahedral $[IrF_6]^2$ unit in 1 ($P\bar{1}$ space group) and a tetragonally distorted octahedral [IrF₆]² unit in 2 $(P4_2/n)$ space group). The axial elongation in both 1 and 2 is $\sim 1\%$, slightly smaller than, for example, the $\sim 3\%$ found in Sr₂IrO₄, and the bond angles are all within 1.5 and 0.1% of 90°, respectively. To elucidate any differences in the Ir^{IV} electronic structure when modifying the ligand field, we included in this study the related chloride complex, (PPh₄)₂[IrCl₆] (3, Fig. 2c and Supplementary Fig. 4), also featuring an approximately octahedral, but axially compressed ($\sim 0.7\%$) [IrCl₆]²⁻ ion. The nearest neighbour Ir-Ir distances (Å) are 10.0 (1), 8.1 (2) and 10.1 (3), which are much longer than in the oxido-iridates (for example, 3.9 Å in Sr_2IrO_4). Thus the Ir^{IV} - Ir^{IV} interactions are negligibly small and the $[IrX_6]^{2-}$ (X=F, Cl) unit can be considered as magnetically isolated in 1, 2 and 3.

Magnetic properties and X-ray spectroscopy. The susceptibility-temperature product, χT , of 1–3 (Supplementary Fig. 5) is practically temperature independent (Curie's law) confirming the lack of significant magnetic interactions between Ir^{IV} spins and thus the absence of any magnetic order at least down to 1.8 K. The complete set of magnetization data (M versus $\mu_0 H T^{-1}$; Supplementary Figs 6–8) could be fitted to the Brillouin function for an effective spin-1/2 (with $g\approx 2$), which corroborates

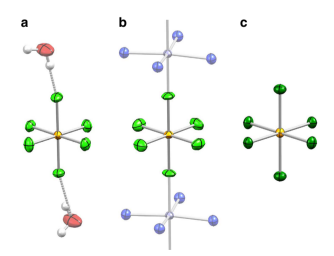


Figure 2 | X-ray structure view of the molecular iridates. Thermal ellipsoid plots of **1 (a)**, **2 (b)** and **3 (c)** are shown at 80% probability level. The counterions for **1** and **3** and the auxiliary parts of the 1-vinylimidazole ligands in **2** are omitted for clarity. Colour code: Ir, yellow; Zn, grey; Cl, dark green; F, pale green; O, red; N, blue; H, white. Selected bond lengths (Å) and angles (°) for **1**: Ir-F 1.9339(8)-1.9510(8), F-Ir-F $_{cis}$ 88.89(4)-91.35(4); for **2**: Ir-F 1.9281(1), 1.9449(1) (only two crystallographically different bond lengths), F-Ir-F $_{cis}$ 90, 90.0112(1); for **3**: Ir-Cl 2.3205(3)-2.3370(3), Cl-Ir-Cl $_{cis}$ 88.54(1)-91.47(1).

the Curie susceptibility and demonstrates the presence of an energetically isolated $J_{\rm eff}\!=\!1/2$ ground state.

Orbital $(M_{\rm orbital})$ and spin $(M_{\rm spin})$ contributions to the total magnetic moments $(M_{\rm total})$ have been determined experimentally using XMCD, which is defined as the difference between two X-ray absorption spectra (XAS) recorded with either opposite helicity or magnetization direction. XAS spectra were collected on 1–3 at the iridium $L_{2,3}$ edges under an external magnetic field of $\mu_0 H = \pm 17\,\mathrm{T}$ and at low temperatures $(T=2.6-2.9\,\mathrm{K})$. Since the measurements were performed on powdered samples, the isotropic $L_{2,3}$ XAS spectra could be approximated as $(\sigma^+ + \sigma^-)/2$, where $\sigma^+(\sigma^-)$ is the absorption cross-section obtained with helicity and magnetization aligned either parallel (+) or antiparallel (-) (Fig. 3). The XAS spectra are dominated by strong resonance peaks ('white lines') at both the L_3 $(2p_{3/2} \rightarrow 5d_{3/2,5/2})$ and L_2 $(2p_{1/2} \rightarrow 5d_{3/2})$ edges with the L_3 edge being significantly more intense.

The spectra of 1 and 2 are virtually identical, whereas for 3, a slight shift of the white line peak of *ca.* 2 eV towards lower energies is observed concomitantly with an additional component at *ca.* 12 eV higher energy (Supplementary Fig. 9). The latter feature is likely the signature of excitations into delocalized states

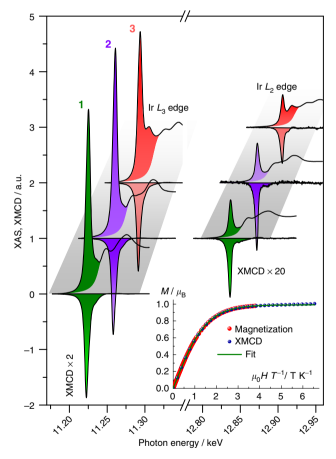


Figure 3 | **X-ray spectroscopy.** X-ray spectra of **1-3** showing the isotropic XAS (positive values) and XMCD (negative values) obtained in a magnetic field of +17 T (at 2.7, 2.6 and 2.5 K for **1**, **2** and **3** respectively). The spectra of **2** and **3** were shifted horizontally (35 and 70 eV, respectively) and vertically for clarity. The filled patterns are the integrals used for the sum rule analysis. Inset: Field dependence of the magnetization, M versus μ_0HT^{-1} , of a polycrystalline sample of **2** at T=2.0 K. 'XMCD' designates the field dependence of the XMCD maximum signal at the Ir L_3 edge at T=2.6 K. The green line is the best fit of the magnetization and scaled XMCD data to the Brillouin function.

originating from ligand to metal charge transfer that is not observed in 1 and 2 due to the more ionic Ir–F bond. The effect of SO coupling on the 5d states was quantified through the so-called SO sum rule²¹. It relates the branching ratio (BR) of the white line integrals at the SO split absorption edges to the expectation value of the angular part of the ground state SO operator per hole. For the $p \rightarrow d$ transitions (L_2 and L_3 edges), the branching ratio can be expressed as

$$BR = \frac{I_{L_3}^{XAS}}{I_{L_3}^{XAS} + I_{L_2}^{XAS}} = \frac{2}{3} - \frac{\langle \sum_{i} \mathbf{l}_i \cdot \mathbf{s}_i \rangle}{3 \langle n_h \rangle}$$
(1)

where, $\langle n_{\rm h} \rangle = \langle n_{\rm h}{}^{j=3/2} \rangle + \langle n_{\rm h}{}^{j=5/2} \rangle$ is the total number of holes in the 5d levels and $\langle \sum_i {\bf l}_i \cdot {\bf s}_i \rangle / \hbar^2 = -3/2 \times \langle n_e{}^{j=3/2} \rangle + \langle n_e{}^{j=5/2} \rangle$ is the expectation value of the one-electron $\mathbf{l}_i \cdot \mathbf{s}_i$ operator summed over all electrons; $\langle n_e \rangle$ being the occupation number of the corresponding levels. The white line integrals are larger in 3 than in 1 or 2, thus more holes in the 5d band should be present for 3 compared with 1 or 2. If one assumes that 1 and 2 are completely ionic, $\langle n_h \rangle = 5$, we obtain $\langle n_h \rangle = 5.26$ for 3. The corresponding results are shown in Table 1. The quantitative analysis of the XMCD spectra was performed by means of the magneto-optical sum rules providing direct access to $M_{\text{orbital}} = -\langle L_z \rangle$ μ_{B} and $M_{\rm spin,eff} = -2 \langle S_{\rm eff} \rangle \mu_{\rm B}$ (the details of this analysis are given in the 'Methods' section; Table 1)^{22,23}. The scaling of the field dependence of the XMCD to the bulk magnetization (inset Fig. 3 and Supplementary Fig. 10; see the 'Methods' sections) allows the determination of the absolute value of the magnetization at $\mu_0 H = +17 \text{ T}$: $M_{\text{total}} = 1.03$, 1.0 and 0.96 μ_{B} for 1, 2 and 3 respectively. From these values, the magnetic dipole contribution, $\langle T_z \rangle$, and $M_{\rm spin}$ (given by $M_{\rm spin,eff} + 7 \langle T_z \rangle$) can be estimated (Table 1) without the need of any sophisticated theoretical modelling from $M_{\text{total}} = M_{\text{spin}} + M_{\text{orbital}}$ considering $M_{\text{spin,eff}}$ and M_{orbital} deduced from the sum rules. The magnetic dipole contribution has never been experimentally determined for any iridate systems, but according to these results, $\langle T_z \rangle$ cannot be neglected without a significant underestimation of the Morbital/ $M_{\rm spin}$ ratio, as previously anticipated²⁴.

Theoretical considerations. The molecular nature of 1, 2 and 3 implies that intuitive localized bonding models should be able to disentangle the magnetic moment contributions. The angular overlap model (AOM) allows for a decomposition of the ligand field potential into σ - and π -bonding parameters²⁵. As a first approximation, a cubic [IrF₆]^{2−} model considering only σ -bonding (with Δ_O = 27,000 cm⁻¹, Racah interelectronic repulsion parameters B = 510 cm⁻¹ and C/B = 4.9, and the SO coupling constant ζ = 3,300 cm⁻¹; Fig. 1c)²⁶ affords $\langle \sum_i \mathbf{l}_i \cdot \mathbf{s}_i \rangle / \hbar^2$ = −2.65, M_{orbital} = 0.74 μ_{B} and M_{spin} = 0.36 μ_{B} , which are close to the ionic values (M_{orbital} = 0.67 μ_{B} and M_{spin} = 0.33 μ_{B}) for a pure J_{eff} = 1/2 system². Whereas the agreement with the experimental results of the fluorido-iridates (Table 1) for $\langle \sum_i \mathbf{l}_i \cdot \mathbf{s}_i \rangle / \hbar^2$ and M_{orbital} is

Table 1 Summary of the X-ray spectroscopy results.									
	1	2	3						
Branching ratio	0.85	0.85	0.82						
$\langle \sum_{i} \mathbf{I}_{i} \cdot \mathbf{s}_{i} \rangle (\hbar^{2})$	-2.8	- 2.7	– 2.5						
$\langle n_h^{j=5/2} \rangle / \langle n_h^{j=3/2} \rangle$	4.7	4.4	3.8						
$M_{\rm total}$ ($\mu_{\rm B}$)	1.03	1.00	0.96						
$M_{\rm spin}$ ($\mu_{\rm B}$)	0.24	0.23	0.31						
$M_{ m orbital}$ ($\mu_{ m B}$)	0.79	0.77	0.65						
$\langle T_z \rangle \; (\mu_{\rm B})$	– 0.11	- 0.10	- 0.081						
Obtained with a magnetic field of $\mu_0H=17$ T.									

surprisingly good, $M_{\rm spin}$ is overestimated in this simplistic approach which neglects the covalency. A more realistic model including π -bonding and scaled AOM parameters to account for the different Ir–F bond lengths (see the 'Methods' section) leads to the same conclusion: $\langle \sum_i \mathbf{l}_i \cdot \mathbf{s}_i \rangle / \hbar^2 = -2.65$, $M_{\rm orbital} = 0.72~\mu_{\rm B}$ and $M_{\rm spin} = 0.32~\mu_{\rm B}$. However, when covalency is explicitly accounted for by CASSCF calculations, the spin contribution $(M_{\rm spin} = 0.22~\mu_{\rm B})$ is in much better agreement with the experiment, but the orbital contribution becomes slightly underestimated $(M_{\rm orbital} = 0.53~\mu_{\rm B})$. Indeed, this departure from cubic symmetry has little influence on the energy of the first excited $J_{\rm eff} = 3/2$ state (see Fig. 1 caption) as evidenced by calculations (Supplementary Table 2) and near-IR absorption spectroscopy revealing absorption bands from $\sim 6,000~{\rm to}$ $\sim 8,500~{\rm cm}^{-1}$ (Supplementary Fig. 11).

Electron paramagnetic resonance spectroscopy. For an ideal $J_{\rm eff}=1/2$ state, magnetic anisotropy is absent, but minuscule deviations from cubic symmetry may result in strong g-factor anisotropy that can be probed experimentally by electron paramagnetic resonance (EPR) spectroscopy. The X-band ($\nu=9.634$ GHz) EPR spectrum of an $[{\rm IrF}_6]^2$ doped ${\rm Zn(viz)_4[ZrF_6]}$ single crystal ($\sim 1\%$ Ir) was measured at 5 K (Supplementary Figs 12–16; Table 2). The experimental eigenvalues of the g-tensor are indeed remarkably anisotropic with $g_z=1.37$ and $g_{xy}=2.11$ in good agreement with the CASSCF calculations leading to $g_z=1.30$ and $g_{xy}=2.24$ (Supplementary Table 2).

These combined experimental and theoretical results establish that the electronic ground state of the molecular $[IrF_6]^2$ species is $J_{\rm eff} = 1/2$ as suggested for the $\{IrO_6\}^8$ unit, present in all oxido-iridates.

Dynamic magnetic properties. The magnetization dynamics of the molecular fluorido-iridates was studied by a.c. susceptibility of **2** at different temperatures below 20 K (Fig. 4 and Supplementary Figs. 17–22 for **1**, **2** and **3**). The presence of peaks in the a.c. frequency (ν) dependence of the imaginary component, $\chi''T$ (Fig. 4b), which shift with temperature, clearly indicates the slow relaxation of the magnetization, while the vanishing of the real component, $\chi'T$, in the adiabatic limit ($\nu \rightarrow \infty$) reveals a blocking of the magnetization that concerns the whole volume of the material. Note that for the three compounds, the slow dynamics of the magnetization is observed only on the application of a small static magnetic field that likely serves to decouple the Ir^{IV} $J_{\rm eff} = 1/2$ from nuclear spins as justified by a comparable magnitude of the applied field (75 mT) to the width of the EPR spectra (Supplementary Fig. 15).

The real and imaginary components of the a.c. susceptibility could be well-fitted to a generalized Debye model²⁷ with small values of the distribution parameter (Supplementary Fig. 23) reflecting a single characteristic relaxation time. The temperature dependence of the extracted relaxation time is shown for 2 in Fig. 4 (and also Supplementary Fig. 24 for 1-3). In these molecular iridates, the spin-lattice relaxation rates τ^{-1} were modelled considering Raman and phonon-bottlenecked direct processes, described as a sum of power laws, $\tau^{-1} = CT^7 + DT^2$, and leading to $C = 46 \times 10^{-3} \, \text{s}^{-1} \, \text{K}^{-7}$, $D = 17 \, \text{s}^{-1} \, \text{K}^{-2}$ for 2 (ref. 28). The magnetization dynamics appears to be similar in 1, 2 and 3 (Supplementary Fig. 24), suggesting that this property is an intrinsic characteristic of the IrIV electronic structure. The slow relaxation of the magnetization in 2 was confirmed by muon spin relaxation (μ^+ SR) measurements at low temperatures (above 1.9 K; Supplementary Figs 25 and 26), where the implanted muons probe the local dynamics of magnetic fields, thereby ruling out any long-range order. This conclusion is further supported by

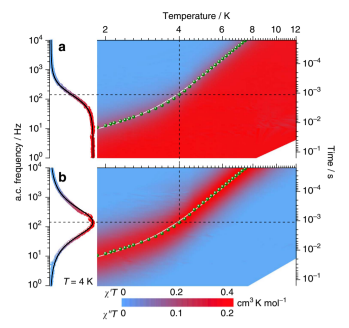


Figure 4 | Magnetization dynamics. Two-dimensional frequency/ temperature maps of the real ($\chi'T$, \mathbf{a}) and imaginary ($\chi''T$, \mathbf{b}) components of the a.c. susceptibility-temperature product for a polycrystalline sample of $\mathbf{2}$ obtained under $\mu_0H=75\,\mathrm{mT}$. Left part: representative 4-K data of the $\chi'T$ and $\chi''T$ frequency dependence with the solid black line being the best fit to the generalized Debye model²⁷, that was used for each temperature to determine the relaxation time (τ) shown as green dots on the contour plots. The best fit of τ^{-1} versus T is shown as solid white lines (see text). Dashed lines are guides for the eyes.

magnetization measurements on an oriented single crystal of 1 and 2 below 2 K (Supplementary Figs 27-30). The magnetic susceptibility ($\chi = dM/dH$) estimated from these experiments in the zero-field limit follows a Curie-Weiss law, $\chi = C/(T - \theta)$, down to lowest accessible temperature of 0.03 K, confirming the absence of magnetic order (Supplementary Fig. 31). For 2, the magnetization reaches saturation faster when the d.c. field is applied perpendicular to the C4 axis rather than along this axis (Supplementary Figs 29 and 30), confirming the easy-plane magnetic anisotropy in agreement with the extracted g-factors (vide supra). The temperature and magnetic field sweep rate dependence of the magnetization revealed the existence of a weak hysteretic behaviour. At the lowest temperatures (< 0.4 K) and in agreement with the a.c. susceptibility data (Fig. 4), butterfly-shaped M versus μ_0H hysteresis loops for both 1 and 2 (Supplementary Figs 27-30) confirm the presence of phononbottlenecked direct processes that dominate the magnetization relaxation²⁹.

Discussion

The principle of dimensional reduction in solid-state structures is based on the formation of a derived compound, $A_{na}MX_{x+m}$, from a parent MX_x precursor and an A_aX salt, thereby forcing termination of M–X–M polymerization³⁰. If A is voluminous, for instance an organic cation, child compounds with structurally and magnetically isolated molecular units can be formed. For many oxides, the ultimate dimensional reduction to molecular $\{MO_x\}^y$ is impeded by the progressive development of large localized negative charges (for example, for $\{IrO_6\}^8$). Thus, truly single-metal ion analogues of most oxides are difficult, if not impossible, to isolate. In the case of octahedral species, no example has been reported so far. As discussed herein for Ir^{IV} , the exchange of oxide with a less negatively charged fluoride results

in dimensional reduction of the iridate as compared with the ternary iridium oxide parents. Although their local structures are obviously comparable, the electronic resemblance between these compounds should be discussed here. The branching ratios extracted from XAS (0.85; Table 1) are identical for 1 and 2, and very close to the values found for oxido-iridates, for example, 0.87 for Sr₂IrO₄ and 0.85 for Y₂Ir₂O₇ (refs 31,32). In addition, the L_2/L_3 XMCD intensity ratio of 4.7% for 2 is almost identical to the value determined for Sr_2IrO_4 (~5%) (ref. 33). This striking agreement demonstrates the resemblance of the electronic structure of the {IrO₆}⁸ moieties in oxido-iridates and the molecular $[IrF_6]^2$ unit. A $M_{orbital}/M_{spin} = \langle L_z \rangle/2 \langle S_z \rangle$ ratio of ~ 2.5 in Sr₂IrO₄ was recently obtained by non-resonant magnetic X-ray scattering that circumvents the need for the estimation of $\langle T_z \rangle$ (ref. 34). For comparison, $M_{\text{orbital}}/M_{\text{spin}}$ ratios of 3.3, 3.3 and 2.1 are obtained for 1, 2 and 3, respectively. The slightly larger ratio for 1 and 2 as compared with Sr₂IrO₄, and 3 is attributed to the weaker covalency of the Ir-X bond for X=F than for X = O, Cl, as reflected in the nephelauxetic series³⁵. It is worth emphasizing that the fluoride ion is the least nephelauxetic of all known ligands, and thus the [IrF₆]² moiety has the closest proximity to a perfectly ionic iridate. As a less-pronounced covalency would induce a larger $M_{\rm orbital}/M_{\rm spin}$ ratio³⁶, a larger orbital magnetic moment is found in $[{\rm IrF_6}]^2$ over $\{{\rm IrO_6}\}^8$ and $[IrCl_6]^{2-}$, as expected.

In oxido-iridates, the purity of the $J_{\rm eff}=1/2$ state depends crucially on the structural deviation from the octahedral symmetry, rendering oxido-iridate systems with weakly distorted $\{{\rm IrO}_6\}^{8-}$ octahedra highly interesting³⁷. Remarkably, the $[{\rm IrF}_6]^{2-}$ moiety in **2** is structurally closer to cubic than any reported oxido-iridate system. Therefore fluorido-iridates are promising materials for stabilizing an ideal $J_{\rm eff}=1/2$ Mott ground state as was already concluded from the experimentally determined values of $M_{\rm orbital}$ and $M_{\rm spin}$. Paramagnetic relaxation in oxido-iridates has been reported for diamagnetically doped honeycomb systems where the imperfect stoichiometry resulted in the blocking of the magnetization at low temperature 38,39 . Our low-temperature experiments reveal that the slow dynamics of the magnetization is indeed intrinsic to the ${\rm Ir}^{\rm IV}$ centre, that might have implications for quantum magnetism.

Herein, we have established that a molecular $[IrF_6]^{2-}$ species and the {IrO₆}⁸ unit, present in all oxido-iridates, possess the same electronic $J_{\text{eff}} = 1/2$ ground state. The dominating role of the SO coupling on the peculiar magnetism intrinsic to IrIV was elucidated by studying spatially isolated, magnetically decoupled, [IrF₆]^{2 –} ions in solids. Whereas strongly perturbed by longrange magnetic interactions in iridates, the 5d spin and orbital moments, as well as their local anisotropy and dynamics have been quantified. These experimental results provide a solid basis for building a realistic model to describe the unusual physics discovered in oxido-iridates. Moreover, as illustrated by the one-dimensional $Zn(viz)_4[IrF_6]$ system (2), the $[IrF_6]^{2-}$ species appears to be a robust building-block for the construction of molecule-based architectures. It offers an unprecedented synthetic strategy using the versatile tools of soft chemistry to engineer new materials with exotic properties anticipated to be exclusively reserved for oxido-iridates.

Methods

Synthetic methods. Na₂[IrCl₆] · 6H₂O was heated to 150 °C *in vacuo* for 6 h, cooled to room temperature (exp. (calc. for Na₂[IrCl₆]) weight loss = 19% (19%)) and subsequently heated to 200 °C in a flow reactor under F₂ gas (10% diluted in Ar) and kept for 6 h. The last step was repeated to improve the crystallinity of the white product (exp. (calc. for Na₂[IrF₆]) weight loss = 19% (22%)) that was identified as Na₂[IrF₆] by powder X-ray diffraction (Supplementary Fig. 32). A solution of Na₂[IrF₆] (68 mg, 0.19 mmol) in water (1 ml) was added a solution of PPh₄Cl (200 mg, 0.53 mmol) in water (3 ml). The resulting solution was left

undisturbed for 1 h to produce crystalline 1 (114 mg, 59%). Anal. calcd. (found) for C₄₈H₄₄F₆IrP₂O₂: C: 56.46% (56.58%), H: 4.34% (4.19%). 1 can reversibly be oxidized to $[IrF_6]^-$ ($E_{1/2} = +0.8 \text{ V}$ versus Fc⁺/Fc; Supplementary Fig. 33). The addition of 1-vinylimidazole (0.80 g, 8.5 mmol) to a methanol solution (40 ml) of 1 (100 mg, 0.098 mmol) and Zn(NO₃)₂ · 6H₂O (100 mg, 0.34 mmol) afforded crystals of 2 after 1 day. Yield: 60 mg (82%). Anal. calcd. (found) for C₂₀H₂₄F₆IrN₈Zn: C: 32.11% (32.36%), H: 3.23% (3.07%), N: 14.98% (14.67%). Complex 3 was obtained from (NH₄)₂[IrCl₆] (203 mg, 0.460 mmol) that was first dissolved in aqueous HCl (0.1 M, 100 ml). The solution was heated to 50 °C and filtered hot. To this solution, a solution of PPh₄Cl (360 mg, 0.963 mmol) in aqueous HCl (0.1 M, 70 ml) was added slowly, inducing an immediate precipitation of orange-brown crystals. On complete addition, the reaction mixture was cooled to room temperature and stirred for 30 min leaving only a slightly coloured solution. The product was filtered off and washed three times with water and dried in a dynamic vacuum. Yield: 453 mg (91%). Anal. calcd. (found) for C₄₈H₄₀Cl₆IrP₂: C: 53.20% (53.24%), H: 3.72% (3.67%).

Crystallography. Powder X-ray diffraction patterns of Na₂[IrF₆] were obtained on samples mounted under nitrogen in an air-tight aluminium alloy cell and collected on a PANanalytical Bragg-Brentano θ-2θ geometry diffractometer (Cu Kα radiation) equipped with a secondary monochromator over an angular 2θ range of 5-80°. Rietveld refinement of the data was performed in the TOPAS (version 4) program⁴⁰. Atomic positions were allowed to refine freely without restraints, while thermal parameters for all atoms were fixed. A suitable fit ($R_{\rm wp} = 13.46\%$) was obtained using a structural model similar to that of Na₂[GeF₆] (ref. 41), with an expanded unit cell (space group P321, a = b = 9.32858(24) Å, c = 5.13417(19) Å, $\alpha = \beta = 90^{\circ}$, $\gamma = 120^{\circ}$; Supplementary Fig. 32). Single-crystal X-ray diffraction studies on 1-3 (Supplementary Table 1) were performed at 122(1) K on a Bruker D8 VENTURE diffractometer equipped with Mo Kα high-brilliance IμS radiation $(\lambda = 0.71073 \text{ Å})$, a multilayer X-ray mirror, a PHOTON 100 CMOS detector, and an Oxford Cryosystems low-temperature device. The instrument was controlled with the APEX2 software package. The structures were solved in Olex2 using the olex2.solve structure solution program (Charge Flipping) and refined using the olex2.refine program⁴². All non-hydrogen atoms were refined anisotropically and hydrogen atoms were placed at calculated positions and refined as riding atoms with isotropic displacement parameters.

Magnetometry. The magnetic measurements were carried out between 1.8 and 280 K with applied d.c. fields ranging from -9 to +9 T, with the use of a MPMS-XL Quantum Design SQUID magnetometer and a PPMS-9 Quantum Design susceptometer. Measurements were performed on polycrystalline samples sealed in polyethylene bags (typically $3 \times 0.5 \times 0.02$ cm; 15-30 mg) and immobilized in mineral oil. a.c. susceptibility measurements were performed with an oscillating field of 1-6 Oe with a frequency from 1 to 10,000 Hz. Before the experiments, the field-dependent magnetization was measured at 100 K to confirm the absence of any bulk ferromagnetic impurities. The magnetic data were corrected for the diamagnetic contributions from the sample, sample holder and mineral oil. Micro-SQUID magnetic measurements on single crystals were carried out in the field range from -1.4 to +1.4 T and in the 5.0-0.03 K temperature range. The field was applied in any direction of the micro-SQUID plane with precision $>0.1^{\circ}$ by separately driving three orthogonal coils⁴³.

X-ray spectroscopy. Polarization-dependent XAS spectra at the Ir L_2 and L_3 edges were acquired at the ID12 beam line at the European Synchrotron Radiation Facility (ESRF), Grenoble, France. A helical undulator of the APPLE-II type allowed the circular polarization of incoming X-rays to be changed for each spectrum. The spectra were obtained by using the total fluorescence yield detection mode in magnetic fields up to ± 17 T. Transmission detection was additionally used for 2 to check the validity of the self-absorption correction used for all samples, which took into account the chemical composition, the geometry of the experiment and the solid angle of the X-ray fluorescence detector. XMCD spectra were obtained as the difference of consecutive XAS spectra obtained with opposite photon helicities. In addition, the XMCD spectra were systematically recorded in both field directions to ensure the absence of experimental artefacts. The isotropic spectra were obtained as the sum of spectra with right and left circularly polarized X-rays and were normalized between zero before the absorption edge and one above the edge. The position of the step function describing transitions into the continuum was defined by the following procedure. The maximum of the L_3 white line was assigned to zero energy and the L_2 edge spectrum was shifted in energy to have the first EXAFS oscillations perfectly overlapping for the two spectra. The step function was placed by visual comparison of the shifted spectra. It was systematically checked that shifting the energy of the inflection point of the step function around the chosen position ($\pm 2 \,\mathrm{eV}$) was not significantly altering the white line integrals. A step function broadening of 6 eV was used, but this parameter does not affect the integrated values of the white line intensities. The white line integral at the L_2 edge was divided by 2.15 reflecting factors of (i) 2 from the occupation ratios of $2p_{3/2}$ and $2p_{1/2}$ core states and (ii) 0.15 from the fact that the sum rules are applied to the line intensities $I(\hbar\omega)$ (where $\hbar\omega$ is the photon energy) and not to the absorption cross sections, $\mu(\hbar\omega)$, which are related by $I(\hbar\omega) = 1/(4\pi^2\alpha \hbar\omega) \times$ $\mu(\hbar\omega)$ (binding energies for L_3 and L_2 edges are 11.225 and 12.835 keV,

Table 2 Hyperfine and superhyperfine coupling parameters for [IrF ₆] ^{2 -} doped in Zn(viz) ₄ [ZrF ₆] (\sim 1% Ir).										
$\mathbf{A_z}(^{191}\mathbf{Ir})$	$\mathbf{A_z}(^{193}\mathbf{Ir})$	$\mathbf{A}_{\mathbf{x}\mathbf{y}}(^{191}\mathbf{Ir})$	$\mathbf{A}_{\mathbf{x}\mathbf{y}}(^{193}\mathbf{Ir})$	A_z^Z	A_{xy}^{Z}	A_{xy}^{XY}	A_{yx}^{XY}	A_z^{XY}		
28.6 (12)	31.04 (6)	27.98 (11)	30.68 (7)	66.36 (6)	3.1 (2.5)	98.58 (5)	3.5 (2.7)	10.58 (6)		
Daramatar valuas	are in units of 10 - 4 cm	a = 1								

respectively). The same normalization procedure was applied to the corresponding XMCD integrals. The precise sample temperature of 2.6–2.9 K and the absolute magnitude of the magnetic moment at $\mu_0 H = \pm 17$ T was determined by scaling the field and temperature dependent XMCD signal intensity to coincide with the M versus $\mu_0 H T^{-1}$ master curve obtained from bulk magnetometry. The normalized spectra were analysed using the magneto-optical sum rules given by equation 2 for L_2 and L_3 absorption edges²²:

$$\begin{split} \langle L_z \rangle &= \frac{2 \langle n_{\rm h} \rangle}{3} \times \frac{I_{L_3}^{\rm XMCD} + I_{L_2}^{\rm XMCD}}{I_{L_3}^{\rm XAS} + I_{L_2}^{\rm XMS}} \\ \langle S_{\rm eff} \rangle &= \langle S_z \rangle + \frac{7}{2} \langle T_z \rangle = \frac{\langle n_{\rm h} \rangle}{2} \times \frac{I_{L_3}^{\rm XMCD} - 2 \times I_{L_2}^{\rm XMCD}}{I_{L_3}^{\rm XAS} + I_{L_2}^{\rm XAS}} \end{split} \tag{2}$$

Ligand field and *ab initio* **calculations**. The AOM calculations were performed using the Ligfield programme on the full $\rm d^5$ configuration 44,45 . All matrices were constructed in the weak-field basis and subsequently transformed to the eigen-basis corresponding to the parameter vectors given below. The octahedral component of the ligand field, $\Delta_{\rm O}=27,000~\rm cm^{-1}$, would agree with the AOM parameter $e_\sigma=9,000~\rm cm^{-1}$ in the cubic model. In the more realistic model, a e_σ/e_π ratio of 5, and a distance dependence of $e_\sigma \propto d_{\rm Ir-F}^{-5}$ and $e_\pi \propto d_{\rm Ir-F}^{-7}$ (where $d_{\rm Ir-F}$ is the Ir-F bond length) was assumed (yielding $e_\sigma^z=11,904~\rm cm^{-1}$, $e_\sigma^{3V}=12,457~\rm cm^{-1}$, $e_\pi^Z=2,356~\rm cm^{-1}$ and $e_\pi^{3V}=2,507~\rm cm^{-1}$). All ab initio calculations were performed with the ORCA 3.0 programme package 46 using ZORA 47,48 for relativistic corrections and the def2-TZVP basis set for the [IrX_6]^2- moieties and the def2-SVP basis set for the 1-vinylimidazole ligands in calculations when requested in the model. State-averaged CASSCF calculations were performed on experimental geometries using Kohn–Sham starting orbitals obtained with the TPSSh functional. Active spaces of CAS(5,3) (Ir t_{2g} orbitals—3 doublet CSFs) and CAS(5,5) (Ir d-orbitals—12 doublet configuration state functions (CSFs), 6 quartet CSFs, 1 sextet CSF) were used and the SO interaction taken into account through a mean field approximation (Supplementary Table 2) 49 .

EPR spectroscopy. The spectra were acquired at $T=5~\rm K$ on a Bruker Elexsys E500 spectrometer equipped with a Bruker ER 4116 DM dual mode cavity, an EIP 538B frequency counter and an ER035M NMR Gauss-meter, and with a microwave radiation frequency of 9.634 GHz. The spectra were simulated using home-written software considering an effective electronic $I_{\rm eff}$ of 1/2 and taking into account the natural isotopic abundance of $^{191}{\rm Ir}$ (37.3%) and $^{193}{\rm Ir}$ (62.3%), which have both a nuclear $I_{\rm Ir}=3/2$ spin, as well as the $^{19}{\rm F}$ nuclear spin ($I_{\rm F}=1/2$, 100%). The spectra were computed using the following spin Hamiltonian for each iridium, $^{191}{\rm Ir}$ and $^{193}{\rm Ir}$, isotope situated in a tetragonally distorted octahedral environment:

$$\begin{split} \hat{H} &= g_z \mu_{\rm B} \mu_0 \hat{S}_z H_z + g_{xy} \mu_{\rm B} \mu_0 \left(\hat{S}_x H_x + \hat{S}_y H_y \right) \\ &+ A_z^{\rm Ir} \hat{S}_z \hat{I}_z + A_{xy}^{\rm Ir} \left(\hat{S}_x \hat{I}_x + \hat{S}_y \hat{I}_y \right) \\ &+ A_z^{\rm Z} \hat{S}_z \left(\hat{I}_{5,z} + \hat{I}_{6,z} \right) + A_{xy}^{\rm Z} \left[\hat{S}_x \left(\hat{I}_{5,x} + \hat{I}_{6,x} \right) + \hat{S}_y \left(\hat{I}_{5,y} + \hat{I}_{6,y} \right) \right] \\ &+ A_{xy}^{\rm XY} \left[\hat{S}_x \left(\hat{I}_{1,x} + \hat{I}_{2,x} \right) + \hat{S}_y \left(\hat{I}_{3,y} + \hat{I}_{4,y} \right) \right] + A_{xy}^{\rm XY} \hat{S}_z \left(\hat{I}_{1,z} + \hat{I}_{2,z} + \hat{I}_{3,z} + \hat{I}_{4,z} \right) \\ &+ A_{yx}^{\rm XY} \left[\hat{S}_x \left(\hat{I}_{3,x} + \hat{I}_{4,x} \right) + \hat{S}_y \left(\hat{I}_{1,y} + \hat{I}_{2,y} \right) \right] \end{split}$$

Here, \hat{S} operators refer to the effective spin of $J_{\rm eff}=1/2$ pertinent to the electronic ground state. \hat{L}_i , \hat{J}_y and \hat{L}_z operators refer to the nuclear spin of the iridium nucleus and $\hat{L}_{n,x}$, $\hat{L}_{n,y}$, $\hat{L}_{n,z}$ (n=1, 2, 3, 4, 5 or 6) refer to the nuclear spin of the fluoride ligands. The fluoride ligands on the positive and negative X-axis are designated with subscripts '1' and '2', respectively. Similarly, subscripts '3' and '4', and subscripts '5' and '6', designate the fluoride ligands on the Y- and Z-axis, respectively. The extracted parameter values are given in Table 2. Remarkably, the estimated hyperfine interactions appear to be almost isotropic, in contrast to the strongly anisotropic superhyperfine interactions to the fluoride ions, which exhibit averaged values on the same order of magnitude as the hyperfine ones. Performing the same experiment on a concentrated crystal of 2 gave virtually identical g-factors, but the hyperfine interactions could not be resolved.

Muon-spin relaxation. The μ^+ SR measurements were performed on the Dolly spectrometer at the Paul Scherrer Institut (Villigen, Switzerland). Fully spin-polarized muons were implanted into the sample and acted as a local probe of the internal magnetic fields. The time evolution of the polarization, which was monitored via the anisotropic β-decay of the implanted muons (lifetime 2.2 μs), is determined by the temporal and spatial properties of the local magnetic field⁵⁰.

For example, fluctuating magnetic moments in the vicinity of a muon would produce a fluctuating dipolar magnetic field, and therefore, a polarization relaxation from its initial value to zero. The time scale of the relaxation can be extracted from the field dependence of the muon spin-lattice relaxation, λ , considering the following equation⁵¹,

$$\lambda = \frac{\Delta^2 \tau}{1 + \gamma^2 \tau^2 (\mu_0 H)^2} \tag{4}$$

where, Δ is the magnitude of the local magnetic field, γ is the gyromagnetic ratio of the muon, $\mu_0 H$ is the applied magnetic field and τ is the fluctuation time scale. The equation above assumes that Δ and τ are field-independent. From the fit of λ as a function of $\mu_0 H$ (Supplementary Fig. 26) $\tau \sim 0.016 \, \mu s$ is obtained. The small value of τ compared with that obtained from a.c. susceptibility measurements is due to the fact that μSR reflects the integrated contribution over q-space as a local, point-like probe, while a.c. susceptibility measures the response for q=0 (ref. 52).

Data availability. The X-ray crystallographic coordinates for structures reported in this Article have been deposited at the Cambridge Crystallographic Data Centre (CCDC), under deposition number CCDC 1431835–1431837. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif. All other relevant data are available from the authors on request.

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Author contributions

K.S.P., J.B. and R.C. designed the research project. K.S.P., A.T., E.D. and J.B. synthesized the compounds and developed the fluorination technologies. K.S.P. and D.N.W. performed the structural analysis. R.C., W.W., Y.L. and K.S.P. acquired and analysed the magnetic data. K.S.P., A.R., K.O., F.W., C.M., S.P. and R.C. acquired and analysed the X-ray spectroscopic data. H.W. and K.S.P. performed the EPR experiments and analysed the data. J.B., T.J.M., S.I.K. and S.O. performed the ligand field and quantum chemical calculations. Z.S. and K.S.P. obtained and analysed the μ^+ SR data.

Additional information

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

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Erratum: Iridates from the molecular side

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The original version of this Article contained an error in which the first affiliation was incorrectly given as 'CNRS, ICMCB, UPR 9048, Pessac 33600, France'. This has now been corrected in both the PDF and HTML versions of the Article.

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