Evidence of an odd-parity hidden order in a spin–orbit coupled correlated iridate

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A rare combination of strong spin–orbit coupling and electron–electron correlations makes the iridate Mott insulator Sr2IrO4 a promising host for novel electronic phases of matter1,2. The resemblance of its crystallographic, magnetic and electronic structures1–4 to La2CuO4, as well as the emergence on doping of a pseudogap region7–9 and a low-temperature d-wave gap10,11, has particularly strengthened analogies to cuprate high-Tc superconductors2. However, unlike the cuprate phase diagram, which features a plethora of broken symmetry phases9–12 in a pseudogap region that includes charge density wave, stripe, nematic and possibly intra-unit-cell loop-current orders, no broken symmetry phases proximate to the parent antiferromagnetic Mott insulating phase in Sr2IrO4 have been observed so far, making the comparison of iridate to cuprate phenomenology incomplete.

Using optical second-harmonic generation, we report evidence of a hidden non-dipolar magnetic order in Sr2IrO4 that breaks both the spatial inversion and rotational symmetries of the underlying tetragonal lattice. Four distinct domain types corresponding to discrete 90°-rotated orientations of a pseudovector order parameter are identified using nonlinear optical microscopy, which is expected from an electronic phase that possesses the symmetries of a magneto-electric loop-current order14–18. The onset temperature of this phase is monotonically suppressed with bulk hole doping, albeit much more weakly than the Néel temperature, revealing an extended region of the phase diagram with purely hidden order. Driving this hidden phase to its quantum critical point may be a path to realizing superconductivity in Sr2IrO4.

The crystal structure of the 5d transition metal oxide Sr2IrO4 is tetragonal and centrosymmetric with four-fold (C4) rotational symmetry about the c-axis19–21. It is composed of stacked IrO2 square lattices whose unit cell is doubled relative to the CuO2 square lattices found in high-Tc cuprates as a result of a staggered rotation of the octahedral oxygen cages (Fig. 1a). Despite the much larger spatial extent of 5d versus 3d orbitals, Sr2IrO4 is a Mott insulator by virtue of strong atomic spin–orbit coupling19–21. The electronic states near the Fermi level derive predominantly from the t2g d–orbitals of iridium ions, which are well approximated as two completely filled spin–orbital entangled $J_{\text{eff}} = 3/2$ bands and one half-filled $J_{\text{eff}} = 1/2$ band, which is split into an upper and lower Hubbard band by an on-site Coulomb interaction. Below a Néel temperature $T_N \sim 230$ K, the spin–orbital entangled $J_{\text{eff}} = 1/2$ magnetic dipole moments undergo three-dimensional long-range ordering into an orthorhombic antiferromagnetic structure2,3 that preserves global inversion symmetry but lowers the rotational symmetry of the system from C4 to C2. No additional symmetry breaking has otherwise been observed by neutron or X-ray diffraction.

Optical second-harmonic generation (SHG), a frequency doubling of light through its nonlinear interaction with a material, is strongly affected by point group symmetry changes19 and can be used to search for hidden phases with higher-rank tensor order parameters that are difficult to detect using conventional probes18,20. It is particularly sensitive to inversion symmetry breaking because the leading-order electric-dipole (ED) contribution to SHG, which is described by a third-rank susceptibility tensor $\chi_{\text{ED}}^{(3)}$ that relates the nonlinear polarization at the second-harmonic frequency $2\omega$ to the incident electric field via $P_2 (2\omega) \propto \chi_{\text{ED}}^{(3)} E_{\omega} E_{\omega} E_{\omega}$, is allowed only if the crystal lacks an inversion centre. Otherwise SHG can arise only from much weaker higher-rank multipole processes, as is the case for the Sr2IrO4 crystal structure described below. The rotational symmetry of a crystal can be directly determined by performing a rotational anisotropy (RA) SHG experiment where the intensity of obliquely reflected SHG is measured as a function of the angle $\phi$ through which the scattering plane is rotated about the surface normal of the crystal (Fig. 1b). For our experiment, the incident light is focused onto optically flat regions of the crystal surface with a spot size $d < 100 \mu m$. The polarizations of the incident (in) and reflected (out) beams can be independently selected to be either parallel (P) or perpendicular (S) to the scattering plane, thus allowing different nonlinear susceptibility tensor elements to be probed.

Figure 1c shows room-temperature RA-SHG patterns of Sr2IrO4 measured by rotating the scattering plane about its c-axis. The data measured under both $P_n - P_{\text{out}}$ and $P_n - S_{\text{out}}$ polarization geometries (see Supplementary Section 1 for other geometries) exhibit a clear $C_4$ symmetry and are completely accounted for by the leading-order non-local bulk contribution to SHG of electric-quadrupole (EQ) type4, which can be written as an effective nonlinear polarization $P_2 (2\omega) \propto \chi_{\text{EQ}}^{(3)} E_{\omega} \nabla \times (E_{\omega} E_{\omega})$. No surface ED contribution to SHG was detected within our instrument sensitivity6. As discussed in ref. 6, the rotation of the SHG intensity maxima away from the crystallographic a and b axes and their modulated amplitude in the $P_n - S_{\text{out}}$ pattern clearly signify the absence of mirror symmetry across the a–c and b–c planes. Expressions for the RA-SHG patterns (Supplementary Section 2) calculated using the set of independent non-vanishing tensor elements of the fourth-rank susceptibility tensor $\chi_{ijkl}^{(4)}$ derived from the experimentally

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established centrosymmetric 4/m crystallographic point group of Sr$_2$IrO$_4$, fit the data extremely well (Fig. 1c).

Remarkably, the rotational symmetry of the RA-SHG patterns measured at low temperature ($T = 170$ K) is reduced from $C_4v$ to $C_2$ (Fig. 1d). This is not caused by a structural distortion because extensive neutron and resonant X-ray diffraction studies show no change in crystallographic symmetry below room temperature. It cannot be accounted for by the antiferromagnetic structure that develops below $T_N \sim 230$ K because that has a centrosymmetric $4/m$ point group above $T = 170$ K (ref. 21). Instead, the ordering of a higher multipolar degree of freedom that coexists with the $J_{\text{eff}} = 1/2$ moment in each IrO$_2$ octahedron is left as the most plausible explanation. An ordering of higher-rank parity-even magnetic multipoles with the same propagation wavevector as the antiferromagnetic structure has, in fact, been proposed to occur in Sr$_2$IrO$_4$ below $T_N$ (refs 22,23). Such an order would naturally be difficult to detect using diffraction-based probes because it preserves the translational symmetry of the antiferromagnetic lattice and imparts no net magnetization to the crystal.

To investigate this possibility, we surveyed all magnetic subgroups of the crystallographic 4/m point group of Sr$_2$IrO$_4$ that do not include the two-fold rotation axis (2) as a group element. We find that our data are uniquely but equally well described by two subgroups ($2'/m$ and $m'1$) of the antiferromagnetic point group. Both $2'/m$ and $m'1$ break the global inversion symmetry of the crystal and thus allow a bulk ED contribution to SHG on top of the existing EQ contribution, which is consistent with the large changes observed in the SHG amplitude. Moreover, expressions for the RA-SHG pattern calculated using a coherent sum of the crystallographic EQ and the hidden-order-induced ED contributions (Supplementary Section 2), with the elements of $\chi^{\text{EQ}}$ derived from a 4/m point group and those of $\chi^{\text{ED}}$ derived from either a $2'/m$ or $m'1$ point group, fit the data extremely well (Fig. 1d). However this indicates that an ordering of higher-rank parity-even magnetic multipoles, which preserves global inversion symmetry, cannot be the origin of our RA-SHG results.

On the other hand, our results can be explained by an ordering of higher-rank parity-odd magnetic multipoles that preserves the translational symmetry of either the antiferromagnetic structure or the crystallographic structure. A microscopic model that satisfies this condition as well as the $2'/m$ or $m'1$ point group symmetries is the magneto-electric loop-current order (Supplementary Section 4), which is predicted to exist in the pseudogap region of the cuprates, but can in principle persist even at half-filling. This phase consists of a pair of counter-circulating current loops in each CuO$_2$ square plaquette (Fig. 1a inset) and can be described by a toroidal pseudovector order parameter defined as $Q = \Sigma_{\mathbf{r}} \times \mathbf{m}$, where $\mathbf{r}$ is the location of the orbital magnetic moment $\mathbf{m}$ inside the plaquette. Four degenerate configurations are possible because the two intra-unit-cell current loops can lie along either of two diagonals.

Figure 1 | Symmetry of the hidden order in Sr$_2$IrO$_4$. a, Crystal structure of a single perovskite layer in Sr$_2$IrO$_4$ (tetragonal point group 4/m). Inset shows the basic IrO$_2$ unit of the magneto-electric loop-current order, with the arrows pointing in the direction of current flow. b, Schematic of the RA-SHG experiment. The electric field polarizations of the obliquely incident fundamental beam (in) and outgoing SHG beam (out) can be independently selected to lie either parallel (P) or perpendicular (S) to the light scattering plane (shaded). RA-SHG data are acquired by measuring the SHG intensity $I(\omega)$ reflected from the (001) surface of Sr$_2$IrO$_4$ as a function of the angle $\psi$ between the light scattering plane and crystal a-c-plane. c, RA-SHG data collected under $P_{\text{in}}=P_{\text{out}}$ and $P_{\text{in}}=S_{\text{out}}$, polarization geometries using $\omega = 800$ nm incident light at $T = 295$ K. d, Analogous data to c collected at $T = 170$ K. All data sets are plotted on the same intensity scale, normalized to a value of 1, which corresponds to $\sim 20$ kW. The high-temperature data are fitted to time-reversal invariant bulk electric-quadrupole-induced SHG (orange lines). The low-temperature data can only be fitted to the coherent sum of time-reversal invariant bulk electric-quadrupole-induced and time-reversal broken bulk electric-dipole-induced SHG (purple lines), as described in the text.
in the square plaquette and can have either of two time-reversed configurations, which correspond to four 90°-rotated directions of the pseudovector. In a real material, one therefore expects domains of all four types to be populated.

To search for domains of the hidden order in Sr$_2$IrO$_4$, we performed wide-field reflection SHG microscopy measurements. A room-temperature P$_{in}$–S$_{out}$ SHG image collected on a clean ∼500 μm × 500 μm region parallel to the a–b plane is shown in Fig. 2a. This region produces a uniform SHG response consistent with the behaviour of a single crystallographic domain. On cooling to $T = 175$ K, brighter and darker patches separated by boundaries that are straight over a length scale of tens of micrometres become visible (Fig. 2b). The distribution and shapes of these patches can be rearranged on thermal cycling (Supplementary Section 5), which suggests that they are not pinned to structural defects in the crystal. To examine whether the different patches observed in Fig. 2b correspond to domains with different $\Omega$ orientation, we performed local RA-SHG measurements within each of the patches. An exhaustive study over the entire crystal area in Fig. 2b reveals a total of only four types of patches, characterized by the four distinct RA-SHG patterns (Supplementary Section 6) shown in Fig. 3, which are exactly 0°-, 90°-, 180°- and 270°-rotated copies of those shown in Fig. 1. These results are consistent with the hidden phase in Sr$_2$IrO$_4$ being a magneto-electric loop-current order—but other microscopic models that obey the same set of symmetries certainly cannot be ruled out.

The ordering temperature $T_\Omega$ of the hidden phase in Sr$_2$IrO$_4$ was determined by measuring the temperature dependence of the SHG intensity in the P$_{in}$–S$_{out}$ polarization geometry where the hidden-order-induced changes are largest. The onset of change is clearly observed at $T_\Omega \sim 232$ K, as shown in Fig. 4a, which is close to but slightly higher than the value of $T_N \sim 230$ K determined by d.c. magnetic susceptibility measurements, and evolves with an order-parameter-like behaviour on further cooling (Supplementary Section 7). In a Landau free energy expansion (Supplementary Section 8), a bilinear coupling between the antiferromagnetic Néel and hidden order parameters is forbidden by symmetry, so there is no a priori reason that $T_\Omega$ and $T_N$ should coincide. The close proximity of $T_\Omega$ and $T_N$ observed in Sr$_2$IrO$_4$ therefore suggests some microscopic mechanism by which one order can induce the other through a biquadratic coupling—an enhancement of the super-exchange coupling between $J_{ex} = 1/2$ moments due to the presence of hidden order being one possible scenario.

To experimentally examine whether or not the Néel and hidden orders are trivially tied, we performed analogous SHG experiments...
on hole-doped Sr$_{2}$Ir$_{1-x}$Rh$_x$O$_4$ crystals (Supplementary Section 9) to track the evolution of $T_N$ as a function of Rh concentration ($x$). Bulk magnetization$^{27}$ and resonant X-ray diffraction$^{28}$ studies have shown that Néel ordering persists for $x \leq 0.17$ and that $T_N$ is monotonically suppressed with $x$. But no evidence of any broken symmetry phases beyond the Néel phase has been reported in Sr$_{2}$Ir$_{1-x}$Rh$_x$O$_4$ so far. Remarkably, our SHG measurements show that the hidden phase transition observed in the parent compound also persists on hole doping (Fig. 4b,c) and that, although $T_D$ is suppressed with $x$, the splitting between $T_D$ and $T_N$ grows monotonically from approximately 2 K to 75 K between $x = 0$ and $x \approx 0.11$. This provides strong evidence that the Néel and hidden orders are not trivially tied, but are independent and distinct electronic phases.

Our finding of a hidden broken symmetry phase in proximity to an antiferromagnetic Mott insulator reveals a striking parallel between the cuprate and Sr$_{2}$IrO$_4$ phase diagrams (Fig. 4d), which is further strengthened by recent observations of a pseudogap region in Sr$_{2}$Ir$_{1-x}$Rh$_x$O$_4$ using angle-resolved photoemission spectroscopy$^9$. Driving the hidden phase to a quantum critical point$^{22}$ through higher doping may be a route to achieving high-$T_c$ (ref. 12) or parity-odd$^{29,30}$ superconductivity in the iridates. Although further theoretical studies are required to establish the microscopic origin of the hidden phase, the fact that it bears the symmetries of a magneto-electric loop-current order already suggests several interesting macroscopic responses, including a linear magneto-electric effect$^{18}$ and non-reciprocal optical rotation$^{17,18}$.

Methods

Methods and any associated references are available in the online version of the paper.

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Figure 4 | Temperature and hole-doping dependence of the hidden order. a–c, Change in SHG intensity from Sr$_{2}$Ir$_{1-x}$Rh$_x$O$_4$ at $x = 0$ (a), $x = 0.04$ (b) and $x = 0.11$ (c) measured relative to their room-temperature values as a function of temperature. Data were taken under $P_{\text{in}}$ polarization geometry at $\varphi = 78^\circ$. The error bars represent the standard deviation over 60 independent measurements. No difference was observed between curves measured on cooling and heating. Lines are guides to the eye. The dashed red lines mark the transition temperature $T_D$ of the hidden order phase deduced from our SHG data and the dashed black lines mark the Néel temperature $T_N$ determined from d.c. magnetic susceptibility measurements. d, Temperature versus doping phase diagram of Sr$_{2}$Ir$_{1-x}$Rh$_x$O$_4$ showing the boundaries of the hidden order and the long-range (LRO) and short-range (SRO) Néel ordered regions. Points where a pseudogap is present are also marked, although a pseudogap phase boundary is not yet experimentally known.
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Author contributions

L.Z. and D.H. planned the experiment. L.Z., D.H.T., H.C. and V.I. performed the measurements. L.Z. and D.H. performed the magnetic point group symmetry analysis. R.F. performed the Landau free energy calculation. T.Q. and G.C. prepared and characterized the samples. L.Z., R.F. and D.H. analysed the data and wrote the manuscript.

Additional information

Supplementary information is available in the online version of the paper. Reprints and permissions information is available online at www.nature.com/reprints. Correspondence and requests for materials should be addressed to D.H.

Competing financial interests

The authors declare no competing financial interests.
Methods

Material growth. Single crystals of Sr$_2$IrO$_4$ and Sr$_2$Ir$_{1-x}$Rh$_x$O$_4$ were grown using a self flux technique from off-stoichiometric quantities of IrO$_2$, SrCO$_3$, and SrCl$_2$, or RhO$_2$, IrO$_2$, SrCO$_3$, and SrCl$_2$, respectively. The ground mixtures of powders were melted at 1,470 °C in partially capped platinum crucibles. The soaking phase of the synthesis lasted for >20 h and was followed by a slow cooling at 2 °C h$^{-1}$ to reach 1,400 °C. From this point the crucible is brought to room temperature through a rapid cooling at a rate of 100 °C h$^{-1}$. The Rh concentration was determined by energy-dispersive X-ray spectroscopy.

RA-SHG measurements. Ultrashort optical pulses with 35 fs duration and 800 nm centre wavelength were produced at a 10 kHz repetition rate from a regeneratively amplified Ti:sapphire laser (KMLabs-Wyvern). The RA-SHG measurements were performed by means of a rotating scattering plane technique$^{35}$ using an SHG wavelength $\lambda = 400$ nm that is resonant with the O 2p to 4f band transition. Light was obliquely incident onto the sample at a 30° angle of incidence with a fluence that was maintained below 1 mJ cm$^{-2}$. Reflected SHG light was collected using a photomultiplier tube. Samples were cleaved either in air or in a nitrogen-purged environment and immediately pumped down to a pressure <5 x 10$^{-5}$ torr in an optical cryostat. Crystals were oriented before measurement using X-ray Laue diffraction.

SHG microscopy measurements. Wide-field SHG images with a spatial resolution of ≈1 μm were collected using the same light source as that used for the RA-SHG measurements. A 40° oblique angle of incidence was used and the incident light fluence was maintained below 1 mJ cm$^{-2}$. The reflected SHG light was collected by an objective lens and focused onto an electron-multiplying charge-coupled device (CCD) camera.

Fitting procedure. The low-($T < T_D$) and high-temperature ($T > T_D$) RA-SHG data were fitted to the expressions $I(2\omega_0, \varphi) = |A\tilde{e}(2\omega_0)\chi_{22}^{(2)}(\varphi)\tilde{e}(\omega_0)\tilde{e}(\omega_0) + A\tilde{e}(2\omega_0)\chi_{33}^{(2)}(\varphi)\tilde{e}(\omega_0)\tilde{e}(\omega_0)|^2I(\omega_0)^2$ and $I(2\omega_0, \varphi) = |A\tilde{e}(2\omega_0)\chi_{23}^{(2)}(\varphi)\tilde{e}(\omega_0)\tilde{e}(\omega_0) + A\tilde{e}(2\omega_0)\chi_{32}^{(2)}(\varphi)\tilde{e}(\omega_0)\tilde{e}(\omega_0)|^2I(\omega_0)^2$ respectively, where $A$ is a constant determined by the experimental geometry, $I(\omega)$ is the intensity of the incident beam, $\tilde{e}$ is the polarization of the incoming fundamental or outgoing SHG light and $\chi_{22}^{(2)}(\varphi)$ and $\chi_{23}^{(2)}(\varphi)$ are the bulk electric-quadrupole and electric-dipole susceptibility tensors, respectively, transformed into the rotated frame of the scattering plane. The non-zero independent elements of the tensors in the unrotated frame of the crystal are deduced by applying the appropriate point group (4/m for $\chi_{22}^{(2)}$ and 2/m or m1' for $\chi_{23}^{(2)}$) and degenerate SHG permutation symmetries. This reduces $\chi_{22}^{(2)}$ to 17 non-zero independent elements (xxxx = yyyz; zzzz = zyyz; xzxz = zyxx; yzyz = xyyz; xzzy = yyzz; zyzz = xyyz; yzxy = yxyz; xyyz = yxyz; yzxx = xyyz; xyyz = yxyz; xzxy = xxzy; zyxx = yxyz; yzxy = yxyz; xyyz = yxyz; zyxx = xyyz; xzyy = yxyz; yyxz = yxyz; xzyy = yxyz) and reduces $\chi_{23}^{(2)}$ to 10 non-zero independent elements (xxxx = yxyy; zzyz = zyxy; zyxx = xyyz; xyyz = yxyz; zyxz = yyzz; zyxz = yyzz; zyxz = yyzz; zyxz = yyzz).

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