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Ultrafast energy and momentum resolved dynamics of magnetic correlations in photo-doped Mott insulator Sr$_2$IrO$_4$

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Measuring how the magnetic correlations evolve in doped Mott insulators greatly improved our understanding of the pseudo-gap, non-Fermi liquids and high $T_C$ superconductivity [1–4]. Recently, photoexcitation has been used to induce similarly exotic states transiently [5–7]. However, the lack of available probes of magnetic correlations in the time domain hinders our understanding of these photo-induced states and how they could be controlled. Here, we implement magnetic resonant inelastic X-ray scattering at a free electron laser to directly determine the magnetic dynamics after photo-doping the Mott insulator Sr$_2$IrO$_4$. We find that the non-equilibrium state, 2 ps after the excitation, exhibits strongly suppressed long-range magnetic order, but hosts photo-carriers that induce strong, non-thermal magnetic correlations. These two-dimensional (2D) in-plane Néel correlations recover within a few picoseconds, while the three-dimensional (3D) long-range magnetic order restores on a fluence-dependent timescale of a few hundred picoseconds. The dramatic difference in these two timescales implies that the dimensionality of magnetic correlations is vital for our understanding of ultrafast magnetic dynamics.

In the layered perovskite Sr$_2$IrO$_4$, multiple interactions conspire to determine its electronic configuration. Strong spin-orbit coupling splits the Ir 5d states to form a narrow electronic band that can be further split by the modest on-site Coulomb repulsion to generate an antiferromagnetic Mott insulating state with close structural, and electronic analogies to the superconducting cuprates [2–4, 9, 10]. It has been well established that when a perturbation melts magnetic order in a Mott insulator, the resulting new state frequently exhibits unusual properties [11]. For example, surface-doping and Rh-Ir substitution in Sr$_2$IrO$_4$ have generated novel Fermi-arc and pseudogap behavior [2, 4] and some have argued that doped Sr$_2$IrO$_4$ might host high temperature superconductivity [12–13]. In both cases, magnetic correlations were argued to play a critical role in the formation of these states. Photo-doping a Mott insulator using ultrafast lasers provides an alternative route to create transient versions of these exotic states, with the advantage that the resulting states are tunable and reversible. To date, however, the appropriate tools have been lacking for probing the momentum and energy dependence of the electronic and magnetic correlations characterizing these ultrafast transient states.
**Figure 1. Experimental configuration.** a, The scattering setup. The vertically polarized pump pulse (shown in red) is incident on the ab-face of Sr$_2$IrO$_4$. X-ray pulses from a free electron laser (shown in purple) probe the resulting transient state. X-rays that are scattered close to 90° are either directly measured, to access the magnetic Bragg peak that probes the presence or absence of 3D magnetic order, or energy analyzed to access the inelastic spectrum that is particularly sensitive to the 2D magnetic correlations. The basic in-plane structural unit of Sr$_2$IrO$_4$ is outlined with a dotted black line. b, An illustration of the pump and probe processes. The 620 meV (2 µm) pump beam (in red) photo-dopes the sample exciting an electron from the lower Hubbard band (LHB) to the upper Hubbard band (UHB). Horizontally polarized 11.215 keV X-ray pulses from a free electron laser (shown in purple) probe the resulting transient state. The incident X-ray pulses excite an Ir 2p core electron into the 5d valence band, in order to couple to the spin degree of freedom. The resulting emitted photon encodes the magnetic and orbital configuration of the transient state [8]. c, Illustration of the detection of x-rays as a function of energy loss, momentum transfer and time delay, encoding the time-dependent magnetic correlations in the transient state. The RIXS planes plot simple spin wave calculations based on an increased thermal population of magnon after the pulse.

Figure 1 illustrates our experimental approach. Sr$_2$IrO$_4$ was cooled to 110 K, well below its Néel ordering temperature of 240 K [14]. Pump laser pulses with an energy of 620 meV (2 µm) drive carriers from the lower Hubbard band to the upper Hubbard band [15]. The transient magnetic response to this pump was characterized using a free electron laser. X-ray photons were tuned to the Ir L$_3$ resonance in order to couple to the spin degree of freedom via the resonant magnetic x-ray scattering mechanism and photons scattered around 90° were measured as a function of momentum transfer, $Q$, energy loss, $E$, and time delay, $t$. Further details are provided in the methods section.

Figure 2a,b plots the time and fluence dependence of the ($-3$, $-2$, 28) magnetic Bragg peak intensity in Sr$_2$IrO$_4$, which is sensitive to the presence of 3D magnetic order. This intensity is measured by an area detector without energy-analyzing the scattered photons. We find that fluences of $\gtrsim$ 5 mJ/cm$^2$ destroy the 3D magnetic order based on the criterion of having $\lesssim$ 10% remnant intensity in the magnetic Bragg peak. This fluence corresponds to exciting a substantial fraction of all the lattice sites within the illuminated volume. Indeed, comparable fluences were also required to destroy long-range magnetic order in other strongly correlated materials including manganites [10,17] and nickelates [18,20].

In order to characterize the charge response to the 620 meV (2 µm) pump excitation, we measured optical reflectivity at 1.55 eV (800 nm) in Fig. 2d. The photocarrier recombination is dominated by processes in the ps or sub-ps regime, far faster than the recovery of 3D magnetic order, suggesting that the charge and magnetic recovery processes are largely independent of one-another.

A detailed understanding of ultrafast magnetic dynamics, beyond the presence or absence of 3D magnetic order, is severely hampered by limited experimental information regarding the short range transient magnetic correlations. Other existing techniques such as X-ray magnetic dichroism [21], Faraday rotation [22] and the magneto-optical Kerr effect [23] capture only 3D magnetic order. This letter breaks new ground by energy analyzing the scattered X-rays i.e. by performing the first ever time resolved (tr) magnetic Resonant Inelastic X-ray Scattering (RIXS) experiment. RIXS probes the magnetic quasiparticle spectrum itself [8,24]. This is a fundamental expression of the nature of the correlated electron state – as it is the spatial and temporal Fourier transform of the spin-spin correlation function and it encodes the interactions present in the magnetic Hamiltonian. In the present case of the 5d valence electron compound Sr$_2$IrO$_4$, the rele-
vant X-ray $L$-edge is in the hard X-ray regime allowing full access to reciprocal space. Such $Q$-space resolution is not available in the complementary technique of time resolved two-magnon Raman scattering, due to the fact that visible photons carry negligible momentum [25].

Figure 3 plots the RIXS energy loss spectra measured in Sr$_2$IrO$_4$ after photo-excitation at 6 mJ/cm$^2$, as compared to the unperturbed state 50 ps before excitation. The chosen pump fluence corresponds to what was required to destroy 3D magnetic order, as seen in Fig. 2b. The RIXS spectra show two dominant features, identified as magnon and orbital excitations [10, 26, 27], which we address in turn.

Orbital excitations appear in the RIXS spectra around 600 meV, and correspond to exciting an electron from the $J_{\text{eff}} = \frac{1}{2}$ ground state orbital to the $J_{\text{eff}} = \frac{3}{2}$ state [26, 27]. The intensity of the orbital excitation is proportional to the $J_{\text{eff}} = \frac{1}{2} \rightarrow \frac{3}{2}$ transition cross-section, and thus directly reflects the electron population in these orbitals. This excitation is different from the pump excitation at 620 meV in that the RIXS process involves two dipole transitions so the process is allowed on a single site [8]. Given that a very similar amplitude of RIXS orbital excitations is seen before and after excitation, we infer that the majority, but not necessarily all, of the photo-excited carriers have decayed out of the $J_{\text{eff}} = \frac{3}{2}$ orbital even in the 2 ps time window. Charge dynamics can also be measured indirectly via optical reflectivity. We see in Fig. 3a that the optical reflectivity at 6 mJ/cm$^2$ is partially, but not fully, recovered at 2 ps delay. The residual reflectivity change likely results from a non-equilibrium charge distribution involving either spectral weight redistribution of the Hubbard bands or the population of in-gap charge states [28], but could also arise from transient structural modifications due to electron-phonon coupling.

The equilibrium excitation spectrum of undoped Sr$_2$IrO$_4$ can be accurately modeled in terms of magnons, or spin waves, that arise from Heisenberg spin-spin exchange interactions [10]. Figure 3a plots the magnetic dispersion from low energies at $(\pi, \pi)$ to $\sim 200$ meV at $(\pi, 0)$. Corresponding experimental RIXS spectra are plotted in Fig. 3b,c with data shown both in the equilibrium (50 ps before the pump) and in the transient state after the pump. Despite the almost-complete destruction of the magnetic Bragg peak in the transient state, magnons are still observed at both $Q$ points. Due to the relatively weak $c$-axis exchange interaction in Sr$_2$IrO$_4$, the intensity, energy-scale and dispersion of these magnons is most sensitive to the 2D Néel correlations between neighboring spins [10]. This indicates that 2D correlations largely retain their Néel-like nature in the transient state 2 ps after the pump. The fact that the magnetic Bragg peak, as shown in Fig. 2b, is very strongly attenuated in the transient state is likely to be due predominantly to the destruction of inter-plane correlations along the $c$-axis. Looking at the RIXS difference spectra in detail, we see that the magnon at $(\pi, 0)$ is identical before and after the pump. At $(\pi, \pi)$, however, there is an appreciable change. This indicates that the high energy $\sim 200$ meV correlations at $(\pi, 0)$ are more robust than the lower energy spin wave at $(\pi, \pi)$ that arises from a smaller disturbance of the Néel order. One interpretation of this observation is that the higher energy magnons recover to their equilibrium configuration in much less than 2 ps, which could be due to the fact that a higher energy excitation can decay into lower energy multi-particle excitations in a larger number of different ways than can the lower energy magnons.

We now consider the magnetic excitation spectrum around $(\pi, \pi)$ in more detail. Due to the finite ($\pm 0.5$ Å$^{-1}$) momentum resolution of our spectrometer, the observed spectrum is the sum of the very low energy magnons...
from precisely \( Q = (\pi, \pi) \), and of slightly higher energy magnons from closely neighboring \( Q \) values. This leads to the asymmetric peak in Fig. 3, which is further plotted as transient state difference spectra in Fig. 3f. We see that \( \sim 20\% \) of the spectral intensity around \( \sim 100 \) meV has been depleted and additional very low energy spectral intensity appears, which recovers on a ps timescale. Thermal heating effects on 2D quantum antiferromagnets have been studied extensively and result in a uniform relative broadening of the magnons across the Brillouin zone \([29, 30]\). Such broadening is not observed here, excluding a purely thermal explanation of our results. Given that we have established the presence of residual photoexcited carriers in the transient state, we suggest that these carriers are directly responsible for damping the magnetic correlations around \( \sim 100 \) meV and causing an apparent redistribution of the magnetic spectral weight to lower energy.

Having clarified the 2D correlations in the transient state, we reassess the behavior of the 3D magnetic order presented in Fig. 2. Even in the few ps regime (Fig. 2b), a small amount of magnetic recovery is evident. However, full recovery takes somewhere between 100 to over 1000 ps (Fig. 2c). We found that a minimal model for the magnetic correlations as a function of time, \( I(t) \), required one decay timescale \( \tau_{\text{decay}} \) and two recovery timescales, which for reasons that we will explain later, are labeled \( \tau_{\text{2D}} \) and \( \tau_{\text{3D}} \), where \( \tau_{\text{2D}} < \tau_{\text{3D}} \)

\[
I(t) = I_0 \left( \exp(-t/\tau_{\text{decay}}) + C \left[ 1 - \exp(-t/\tau_{\text{2D}}) \right] + (1 - C) \left[ 1 - \exp(-t/\tau_{\text{3D}}) \right] \right).
\]

(1)

This model was fit to the magnetic Bragg peak intensity data in Fig. 2b,c. In a similar way, we fitted the recovery of the optical reflectivity, which also required two charge timescales denoted \( T_{\text{fast}} \) and \( T_{\text{slow}} \).

Figure 4 summarizes the magnetic and charge dynamics in \( \text{Sr}_2\text{IrO}_4 \) after laser-excitation creates a large population of doublons. 3D magnetic order decays in \( 0.30 \pm 0.03 \) ps approximately independent of fluence (Fig. 4a), which is roughly equal to the jitter-limited time resolution of the experiment. This sets an upper limit on the timescale for the destruction of magnetic order in this system. The fast component of the charge recovery (Fig. 4a) is of a similar magnitude \( 0.27 \pm 0.04 \) ps. Panel c plots the faster magnetic recovery timescale which is \( \sim 1.7 \) ps and increases slowly with fluence. As discussed above, the 2D in-plane magnetic correlations recover on the ps timescale, so the lack of 3D magnetic order can be primarily attributed to the lack of coherence between the \( \text{IrO}_2 \) planes. Or, equivalently, after a few ps there is a large population of low-energy long-wavelength magnons that, on average, preserve the 2D Néel correlations. Consequently, we attribute the fast timescale to 2D in-plane correlations and label it \( \tau_{\text{2D}} \). On the hundreds of ps timescale the 2D correlations are largely restored and the recovery is dominated by the restoration of 3D inter-plane correlations, leading us to assign the slower timescale to \( \tau_{\text{3D}} \). The dramatic difference in these timescales reflects the strong anisotropy in the magnetic interactions. \( \text{Sr}_2\text{IrO}_4 \) has very strong in-plane magnetic exchange \( J_\parallel = 60 \) meV and very weak inter-plane magnetic exchange, which some researchers have estimated to be as low as \( J_\perp \approx 1 \) meV \([10, 27, 31, 32]\). \( \tau_{\text{3D}} \) is also seen to be linear with fluence (Fig. 4a), increasing up to \( 1130 \) ps at \( 13.8 \) mJ/cm\(^2\). This implies that the recovery of the long-range magnetic order relies crucially on the dissipation of energy in the material and is set by the timescale upon which the low-energy long-wavelength
magnons can dissipate their energy into the lattice degree of freedom, whereas \( \tau_{2D} \) has far weaker fluence dependence and seems to be related to a material property.

Time-resolved resonant X-ray scattering has provided a new window on the transient magnetic state in photo-doped \( \text{Sr}_2\text{IrO}_4 \). The 2D magnetic correlations we observe are non-thermal in nature and recover on a ps timescale denoted \( \tau_{2D} \). A striking similarity between \( \tau_{2D} \) and the slower charge recovery timescale \( T_{\text{slow}} \) is seen in Fig. 4. This may reflect the similar energy scale of the in-plane electronic hopping parameter, \( t_{||} \), and the magnetic exchange, \( J_{||} \), which are fundamentally linked in strongly correlated materials such as these via \( J_{||} \approx t_{||}^2/U \) where \( U \) is the Coulomb repulsion. The behavior of the long-range magnetic order, in contrast, depends on secondary processes, such as inter-plane magnetic coupling and the dissipation of the energy from the spins into other degrees of freedom.

This work shows that direct measurements of the 2D magnetic correlations are consequently crucial for a full understanding of magnetic dynamics in strongly correlated materials. With the continued improvement of free electron lasers, tr-RIXS is set to play a crucial role in understanding how magnetic correlations dictate the properties of doped Mott insulators and how they can be effectively manipulated by light.

**METHODS**

**Samples** The magnetic Bragg peak measurements were performed on 200 nm epitaxial films of \( \text{Sr}_2\text{IrO}_4 \), in order to match the volume of \( \text{Sr}_2\text{IrO}_4 \) to the penetration depth of the pump, as the X-ray penetration depth is longer than the pump. The disappearance of the magnetic Bragg peak in Fig. 2a,b confirms that the whole probed volume is excited. The film was deposited on \( \text{SrTiO}_3 \) using pulsed laser deposition as described in the supplementary information and Ref. [33]. Figures S1 and S2 in the supplementary information demonstrate good sample crystallinity and the lack of any detectable impurity phases. For RIXS, \( \sim 1^\circ \) grazing incidence X-rays were used to limit X-ray penetration depths to 80 nm on a bulk \( \text{Sr}_2\text{IrO}_4 \) crystal. Both samples have a c-axis surface normal. Reciprocal lattice notations are defined using the full unit cell with lattice constants \( a = b = 5.51 \, \text{Å} \) and \( c = 25.7 \, \text{Å} \). The high symmetry points in the in-plane Brillouin zone are defined in the reduced structural zone (which ignores the rotation of the IrO\(_6\) octahedra) as in Ref. [10]. The zone center, denoted \((\pi, \pi)\) and the zone boundary denoted \((\pi, 0)\) correspond to \((1, 0, L)\) and \((0.5, 0.5, L)\) respectively in the reciprocal lattice notation. In both experiments, the sample was cooled to about 110 K with a nitrogen cryostream, well below the Néel temperature of 240 K [13].

**Optical pump** For both tr-REXS and tr-RIXS experiments, 100 fs pump pulses were generated at 620 meV (2 μm) using an optical parametric amplifier. The pulses were polarized vertically in the \( ab \)-plane of the sample and were incident at 13° with respect to the sample surface. The choice of pump energy follows previous optical conductivity measurements [15] and resonates between the upper and lower Hubbard bands.

**The time-resolved resonant elastic X-ray scattering (tr-REXS) setup.** The tr-REXS experiment was performed at beamline 2 of the SP rings-8 Angstrom Compact free electron LAser (SACLA) with a 30 Hz pulse repetition rate. We adopted a horizontal scattering geometry as seen in Fig. 1a and tuned the X-ray energy to the peak in the Ir \( L_3 \)-edge resonance around 11.215 keV. A Multi-Port Charged Couples Device (MPCCD) area detector was placed at \( 2\theta = 88.7° \) to observe the magnetic Bragg peak \((-3, -2, 28)\). This geometry is chosen to optimize the X-ray resonant magnetic scattering cross sec-

**FIG. 4. Fluence dependence of the magnetic and charge dynamics timescales.**

(a) the decay time scale \( \tau_{\text{decay}} \), showing that the magnetic decay happens in \( \leq 0.3 \) ps, faster than the time resolution of the experiment. (b) the fast charge recovery time scale \( T_{\text{fast}} \). (c) the timescales for the slow component of the charge recovery \( T_{\text{slow}} \) and the faster component of the magnetic recovery. From our RIXS data we know that the 2D magnetic correlations recover on a ps timescale, so due to the similar timescales of this fitting parameter it is labelled \( T_{2D} \). These show strikingly similar magnitudes and fluence dependences. (d) \( \tau_{3D} \) the slow magnetic recovery, which is assigned to the restoration of 3D magnetic order. Grey lines are guides to the eye and error bars are the standard deviation from the least square fitting algorithm.
tion. We access the magnetic peak by rotating the sample around the vertical axis by $\phi = 12.8^\circ$ with the infrared and X-ray photons in an approximately co-linear geometry. The detector was read-out shot-by-shot and the signal was thresholded to suppress the background coming from X-ray fluorescence and electrical noise. The peak intensity was determined by binning the 2D MPCCD data into a 1D spectrum and fitting a Lorentzian lineshape with a uniform offset background. Each datapoint is the result of summing 1000-4000 shots. Previous characterization of the beamline found that the time resolution of this experiment was jitter-limited to approximately 300 fs.

The minimal model for the fitting is outlined in the main text (Equation 1). This formula was convolved with a 100 fs Gaussian to account for the pump pulse width. The other major contribution to the effective time resolution was the x-ray pulse jitter of approximately 300 fs, because this is only an approximate value this was not included in the fit, rather this is taken as an upper limit on the decay time. Apart from this quantity, all parameters were varied to fit the data in the long time delay scans at 1.0, 2.7 and 13.8 mJ/cm$^2$ fluence in Fig. 2b and these fits were used to constrain $\tau_{3D}$ in fits of the short time delay data in Fig. 2b, by interpolating the variation of $\tau_{3D}$ and $C$ as a function of fluence. In this way, equation 2 provides an accurate parametrization of the recovery dynamics at all fluences studied.

The time-resolved resonant inelastic X-ray scattering (tr-RIXS) setup. The tr-RIXS experiment was performed at the X-ray Pump Probe instrument [34] at the Linac Coherent Light Source (LCLS) with a 120 Hz repetition rate. We adopt a horizontal scattering plane, similar to the setup in the tr-REXS experiment. The $(\pi, 0)$ and $(\pi, \pi)$ were measured at $(-3.5, -3.5, 24.1)$ and $(-4, -4, 23.9)$. Non-integer values of $L$ were chosen to keep the X-ray incident angle around 1° as the RIXS spectrum is known to be essentially independent of $L$ [10]. A Si (333) monochromator produced a 50 meV incident energy bandpass. The RIXS spectrometer is conceptually similar to that used at Sector 27 at the Advanced Photon Source. Scattered photons from the sample are reflected from a segmented spherical Si(8,4,4) analyzer in a near-backscattering configuration and detected by a Princeton CCD. The sample, the analyzer crystal, and the photodetector are placed on a Rowland circle with a radius of 1 m in the vertical plane. The total energy resolution of the tr-RIXS setup was $\sim 70$ meV and the $Q$ resolution was defined by the $\sim 6^\circ$ angular acceptance of the analyzer. RIXS spectra were collected in a stationary mode without moving the spectrometer and the pixel-to-energy conversion was performed using well-established methods. The CCD was read out every 1800 shots. Jitter and timing drift were the main contributions to the time resolution, which was on the order of 500 fs.

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AUTHOR CONTRIBUTIONS

J.P.H., X.L., M.P.M.D. and M.F. initiated and planned the project. M.P.M.D., Y.C., X.L., S.W., D.Z., R.M., V.T. X.M.C., J.V., D.C., J.K., A.H.S., P.J., R.A.-M., M.G., A.R., J.R., M.S., S.S., H.L., L.P., S.O., T.K., M.Y., Y.T., T.T., L.H., C.-L.C., D.F.M., M.F. and J.P.H. prepared for and performed the experiments. M.P.M.D., Y.C., X.L., S.W., M.F., D.F.M. and J.P.H. analyzed and interpreted the data. J.L., C.R.S. and B.J.K. prepared the samples. M.P.M.D. and Y.C. wrote the paper with contributions from X.L., S.W., D.F.M., M.F. and J.P.H.

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