Pressure tuning of light-induced superconductivity in K₃C₆₀

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Optical excitation at terahertz frequencies has emerged as an effective means to dynamically manipulate complex materials. In the molecular solid K₃C₆₀, short mid-infrared pulses transform the high-temperature metal into a non-equilibrium state with the optical properties of a superconductor. Here we tune this effect with hydrostatic pressure and find that the superconducting-like features gradually disappear at around 0.3 GPa. Reduction with pressure underscores the similarity with the equilibrium superconducting phase of K₃C₆₀, in which a larger electronic bandwidth induced by pressure is also detrimental for pairing. Crucially, our observation excludes alternative interpretations based on a high-mobility metallic phase. The pressure dependence also suggests that transient, incipient superconductivity occurs far above the 150 K hypothesized previously, and rather extends all the way to room temperature.

Resonant optical excitation of infrared-active phonon modes can drive the crystal lattice of solids nonlinearly, excite other orders coherently, switch lattice polarization, drive insulator-to-metal or magnetic transitions, and even induce transient superconductivity above equilibrium Tc. In the potassium-doped fulleride K₃C₆₀, a superconductor with critical temperature Tc = 20 K, excitation of local molecular vibrations was shown to induce superconducting-like optical properties in the high-temperature metal (T > Tc). Key features of this state are a 12 meV wide gap in the frequency-dependent optical conductivity σ(ω), twice as large as the equilibrium 6 meV wide superconducting gap, and a divergent low-frequency imaginary conductivity σ₁(ω), indicative of high carrier mobility. This state was found to extend to at least 100 K, hence up to temperature scales far in excess of equilibrium Tc (20 K). For T > 100 K, a partially gapped state was reported. In this higher-temperature regime, the light-induced state can be interpreted either as a high-mobility metal en route to a transient superconducting state, or as an incipient superconductor, which is only partially coherent.

Many theoretical mechanisms have been invoked to explain these observations, ranging from a dynamical reduction of the electronic bandwidth to the parametric amplification of the pairing instability and to electron attraction in vibrationally excited molecular sites. Recent experiments in bilayer graphene are consistent with some of these suggestions, as the optical excitation of a vibrational mode similar to that driven in K₃C₆₀ appears to increase the electron-phonon interaction. Finally, recent theoretical work has raised the possibility that photostimulation may involve optical excitation and cooling of above-gap thermal quasiparticles into a superexcitonic state with high electronic heat capacity.

The face-centred cubic structure of doped fullerenes A₃C₆₀ is shown in Fig. 1a. Three electrons are donated by the alkali atoms to each C₆₀ molecule, which then form three narrow, half-filled bands near the Fermi level. Figure 1c shows how the equilibrium superconducting transition appears in the steady-state optical properties of K₃C₆₀, measured above and below Tc = 20 K. When cooling metallic K₃C₆₀ (red curves) below Tc, one observes large changes in the optical properties: a saturation of the low-frequency reflectivity to R = 1, a 6 meV gap in the real part of the optical conductivity σ₁(ω), and a divergence in the imaginary part σ₅(ω) (blue curves).

As superconductivity in K₃C₆₀ emerges from a combination of Jahn–Teller intramolecular distortions and electronic correlations, it is natural to explore the response of the material to direct excitation of optically accessible vibrational modes. In Fig. 1d we report the optical properties of polycrystalline powders of K₃C₆₀ 1 ps after the excitation tuned to 'on-ball' infrared-active modes of T₁u symmetry at 170 meV energy (7.3 μm wavelength), whose atomic distortion is displayed in Fig. 1b. At this frequency, strongly correlated metallic carriers are also excited, as the radiation is also resonant with a broad absorption peak extending from about 40 to 200 meV, whose precise origin is still unclear.

A broadband probe pulse was used to detect the light-induced changes in the optical reflectivity and complex optical conductivity between 1.6 and 7 THz (6.5–29 meV) using THz time-domain spectroscopy. Starting from the unperturbed metallic state at 100 K (red curves), we observed an increase in the reflectivity, which saturates to R = 1 for all probe photon energies below 12 meV, a gapped σ₁(ω) and a divergent σ₅(ω). These data confirm the results of ref. 9, but were recorded with an improved apparatus, involving higher pump fluence and a broader probe bandwidth (see Supplementary Information S3).

In this paper, we study how the features reported in Fig. 1d change with the application of hydrostatic pressure. At equilibrium, the application of pressure reduces the superconducting transition temperature Tc, because of the increase in the electronic bandwidth when the intermolecular spacing is reduced. As shown in Fig. 2, the size of the optical gap (2Δ₉) and the critical temperature Tc decay linearly even at relatively modest pressures. Due to the low bulk modulus (28 GPa; ref. 9), a pressure of 3 GPa reduces the superconducting gap to less than half of the ambient-pressure value, as the electronic bandwidth increases by about 25% (ref. 25).
rat ambient pressure, with a reflectivity approaching $\sigma_\omega$, C60 molecular distortion (blue) along the T1u(4) vibrational mode coordinates. The equilibrium lattice constant is 14.26 Å at room temperature. $b$ are represented by the green bonds connecting all the C atoms. The grey spheres are the K atoms, acting as spacers between neighbouring buckyballs. The structure, equilibrium phase transition, and transient light-induced phase of K3C60.

Figure 3 displays snapshots of the measured optical reflectivity $R(\omega)$ at the sample–diamond interface, along with complex conductivity spectra, $\sigma_\omega(\omega)$ and $\sigma_i(\omega)$, for different values of static pressure. The exact pressure was measured with calibrated ruby fluorescence (see Supplementary Information S3). In each panel, the red and blue curves trace the optical properties of the equilibrium metal and those of the non-equilibrium state induced by photoexcitation, respectively. For pressures up to 0.17 GPa (Fig. 3a–c) the transient optical response of K3C60 at equilibrium (red) and 1 ps after photoexcitation (blue) at $T = 100$ K. The light-blue curves show the data reported in ref. 8, measured with a fluence of 1 mJ cm$^{-2}$, while those in dark blue were measured with a broader probe spectrum and a higher pump fluence (3 mJ cm$^{-2}$).

As the applied pressure increases, a stronger suppression of the light-induced changes in both the reflectivity and complex optical conductivity is observed (Fig. 3d,e). Above 0.3 GPa the enhancement in the reflectivity is clearly less pronounced, and a progressively broader Drude peak appears at low frequency in the $\sigma_i(\omega)$ spectrum.

The reduction in light-induced coherence observed as a function of pressure is clearly not compatible with the behaviour expected for...
a light-induced metallic state, as a lattice compression in a metal is typically associated with larger electronic bandwidth, smaller effective mass and higher mobility. This is for example evident when analysing the equilibrium metallic properties in the red curves of Fig. 3 (see also Supplementary Information S2), where we observe higher plasma frequencies $\omega_p$ with increasing pressure.

In Fig. 4 we report the fractional spectral weight loss for frequencies inside the gapped region of the spectrum, obtained by integrating $\sigma_1(\omega)$ between 6.5 and 12.9 meV for different pressures and base temperatures of 100 K, 200 K and 300 K (see Supplementary Information S9 for full data sets at 200 and 300 K). Shaded blue areas indicate the pressure–temperature ranges where the light-
induced state is gapped. Overall, the light-induced gap fills even at moderate pressure values, becoming even smaller for increasing temperature. For $P \geq 0.3$ GPa, the pressure dependence of the light-induced effects is strongly reduced.

Fits to the optical properties of Fig. 3 make the qualitative analysis above quantitatively significant (see Supplementary Information S10). By fitting the transient optical response of $K_3C_60$, we extrapolated the value of the low-frequency optical conductivity $\sigma_\omega \approx \lim_{\omega \to 0} \sigma_i(\omega)$. To compare both superconducting-like and metallic-like states in a consistent fashion, we used a Drude–Lorentz fit for the entire pressure range, in which $\sigma_i$ was allowed to float from finite (metal) to infinite values (perfect conductor), and a single lorentzian was used to capture the mid-infrared absorption band.

The results of this analysis are summarized in Fig. 5, where we report the pressure dependence of $\sigma_i$ for three temperatures (100 K, 200 K, and 300 K). The red squares refer to the equilibrium metal, while the blue diamonds to the photoexcited state. As shown in these plots, the equilibrium metallic conductivity increases with applied pressure. In contrast, two pressure regimes are found for the light-induced state, one in which $\sigma_i$ decreases for small pressures ($d\sigma_i/dP < 0$, blue shaded area) and one where it eventually increases slightly for higher pressures ($d\sigma_i/dP > 0$, yellow shaded area). Several indications can be extracted from these data.

First, as mentioned above, from the optical properties alone reported in ref. 8, one could not uniquely differentiate a superconductor from a perfect conductor, as optics only identifies the density of charge carriers and the scattering rate. The hydrostatic pressure dependence reported here adds crucial information. At low pressures, the photoexcited state has clear superconducting-like pressure dependence ($d\sigma_i/dP < 0$), whereas for higher pressures the response is clearly metal-like ($d\sigma_i/dP > 0$). Furthermore, at high pressures the $\sigma_i$ of the photoexcited state follows the same slope as that of the equilibrium metal.

In this context the results reported for high temperatures ($T = 200$ K and $T = 300$ K) are surprising. In this temperature range, a high-mobility metallic state was proposed to interpret the data of ref. 8. However, this interpretation was also not unique, as a superconducting-like state with progressively lower coherence could also have explained the data. Figure 5 suggests that in the low-pressure regime the $d\sigma_i/dP < 0$ behaviour is retained all the way to 300 K, suggesting that some incipient features of transient superconductivity may already be present up to room temperature.

These observations also provide guidance for a microscopic explanation of our results. Indeed, as summarized in Fig. 6, we find a very strong dependence of the light-induced optical conductivity on pressure, and for the higher-pressure ranges (smaller lattice constants) the metallic phase (yellow) is stabilized. Our data set
an important benchmark for theories of photoinduced superconductivity\textsuperscript{13-15,26,27}, which should reproduce the observed pressure dependence. Figure 6 also indicates a clear path for future research in the broader context of $A_2C_{60}$ superconductivity, showing on the right-hand side the region of the phase diagram still to be accessed (b), with the interesting perspective of optimizing light-induced superconductivity further, for even larger lattice spacing.

**Data availability.** The data sets generated and analysed during the current study are available from the corresponding author on reasonable request.

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**Author contributions**

A. Cavalleri conceived the project together with M.M. and A. Cantaluppi. The time-resolved THz setup was built by A. Cantaluppi and M.B., who both made the pump–probe measurements and analysed the data with the support of G.J. and D.N. The equilibrium optical properties were measured by A. Cantaluppi and M.M., with the support of A.P. and F.D.P., and were then analysed by A. Cantaluppi and M.B. The samples were grown and characterized by D.P. and M.R. The manuscript was written by A. Cavalleri, with input from all co-authors.

**Competing financial interests**

The authors declare no competing financial interests.

**Additional information**

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