Beating the Stoner Criterion Using Molecular Interfaces

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It is accepted that only three elements are ferromagnetic at room temperature, the transition metals iron, cobalt and nickel. The Stoner criterion explains why, for example, iron is ferromagnetic but manganese is not, even though both elements have an unfilled 3d shell and are adjacent in the periodic table: the product of the density of states with the exchange integral must be greater than unity for spontaneous ordering to emerge. Here, we demonstrate that it is possible to alter the electronic states of non-ferromagnetic materials, such as diamagnetic copper and paramagnetic manganese, in

order to drive them ferromagnetic at room temperature. This remarkable effect is achieved via interfaces between metallic thin films and C₆₀ molecular layers. The emergent ferromagnetic state can exist over several layers of the metal before being quenched at large sample thicknesses by the material's bulk properties. While the induced magnetisation is easily measurable by magnetometry, low energy muon spin spectroscopy³ provides insight into its magnetic distribution by studying the depolarisation process of low energy muons implanted in the sample. This technique indicates localized spin-ordered states at and close to the metallo-molecular interface. Density functional theory simulations suggest a mechanism based on magnetic hardening of the metal atoms due to electron transfer.^{4,5} This opens a path for the exploitation of molecular coupling to design magnetic metamaterials using abundant, non-toxic elements such as organic semiconductors. Charge transfer at molecular interfaces can then be used to control spin polarisation or magnetisation, with far reaching consequences in the design of devices for electronic, power or computing applications.^{6,7}

Multifunctional materials with the spin degree of freedom such as multiferroics, magnetic semiconductors and molecular magnets have all aroused huge interest as potentially transformative components in quantum technologies. Strategies used to bring magnetic ordering to these materials typically rely on the inclusion of magnetic transition metals, heavy elements with a large atomic moment or rare earths. In thin film structures, proximity effects and coupling at interfaces play an essential role. This is especially the case for molecular spintronics, where organic thin films grown on copper have demonstrated spin filtering. The organic magnetic coupling can propagate for long distances in systems such as nanoscale vortex-like configurations or nanoskyrmion lattices.

We choose C_{60} as a model molecule due to its structural simplicity and robustness as well as its high electron affinity. C_{60} /transition metal complexes exhibit strong interfacial coupling between metal $3d_z$ electrons and molecular π -bonded p electrons. The potential created by the mismatch of molecular and metal work functions leads to a partial filling of the interface states. ¹⁹⁻²¹ Other molecules with close electron affinity and the potential for $3d_z/p$ coupling could be used to similar effect. In the case of C_{60} on metallic substrates such as Cu films, the charge transfer from the metal can be of up to $3e^-$ per molecule and leads to a metallisation of the interface. ²² Magnetic polarisation in fullerenes induced by spin injection or charge transfer may extend for long distances owing to low spin orbit coupling and the absence of a hyperfine interaction. ^{23,24}

In the metal, it is expected that the charge transfer will be quickly screened by free electrons. A priori, there would be no reason to consider that a spin-unpolarized molecule would change the magnetic state of a metallic film. However, we find that the charge transfer and surface reconstruction at the interface²⁵ can lead to an emergent magnetisation in both the metal and the molecule. Magnetometry measurements of C_{60} /Cu and C_{60} /Mn multilayers show hysteresis at room temperature. The magnetization disappears when all the transition metal-molecular interfaces are decoupled via an Al or Al₂O₃ spacer layer (Fig. 1).

Changes in the Density of States (DOS) of the metal may be larger close to the interface, but should be screened deeper within the material. If the film is thick enough, the bulk properties of the metal are expected to dominate and quench the magnetization. This effect is shown in figure 2: the magnetisation of C_{60} /Cu and C_{60} /Mn multilayers decays once the metallic film thickness exceeds 2-3 nm. Decreasing coupling between the top and bottom interfaces of a metal layer may also play a role.

The magnetisation of C_{60} /Cu samples is 3-4 times stronger than for C_{60} /Mn, which is probably due to the better lattice matching and larger charge transfer between copper and

 C_{60} . However, bulk manganese is paramagnetic and much closer to complying with the Stoner criterion than diamagnetic copper due to the larger exchange interactions and density of states at the Fermi level $(DOS(E_F))$. This may be correlated with the propagation length of the effect, which persists for sample thicknesses five times longer in Mn than in Cu. C_{60} /Mn multilayers also show a larger paramagnetic slope of the magnetisation than decoupled manganese –see supplementary information (SI), Figs. S.1-S.4. Both systems exhibit anisotropy with an easy axis that lies in the plane of the film, and out-of-plane saturation fields of ~10-15 kOe at room temperature (inset in Fig. 2b). The Cu and Mn samples degrade with time, and the magnetization drops over several days or weeks depending on the layer structure and protective cap used (SI, Fig. S.5).

With a view to explore the dependence of magnetisation on interfacial coupling, we have fabricated samples with different number of $C_{60}(15 \text{ nm})/Cu(2.5 \text{ nm})$ junctions. The magnetic moment of these multilayers is proportional to the number of Cu-C₆₀ interfaces, suggesting that the magnetism is indeed due to molecular coupling (Fig. 2a inset). However, the amount of Cu and C_{60} is also increased as we grow more layers. In order to ensure that the magnetisation is not simply proportional to the amount of material deposited, we perform a related set of measurements where the total sample thickness is kept constant: 9 nm of Cu and 81 nm of C_{60} , but split into different number of C_{60}/Cu repeats (SI, Fig. S.6). There, the magnetism also increases with the number of interfaces; e.g. magnetic moment of $[C_{60}(16.2 \text{ nm})/Cu(1.8 \text{ nm})]x5 > \text{moment of } [C_{60}(27 \text{ nm})/Cu(3 \text{ nm})]x3$. However, trying to split the sample into Cu films of ~1.5 nm or thinner results in discontinuous layers and a drop in the magnetisation. This thickness and interface dependence of the magnetisation could not arise from contaminants, and X-ray spectroscopy did not show the presence of impurities (SI, Figs. S.7-S.11).

Magnetometry measurements show that the magnetisation is dependent on the thickness of the metal but not on that of the molecular film as long as the latter is continuous and smooth (~10-20 nm). However, magnetometry by itself cannot determine where the magnetisation is located or how much of it corresponds to each material. Conversely, low energy muon spin rotation (μ SR) provides a magnetic profile of the sample²⁷ and has been applied successfully to other metallo-molecular systems.³ Here, a beam of almost fully polarised positive muons (μ ⁺) is moderated to keV energies so that their tunable stopping range is tens to hundreds of nm. The local polarisation at the μ ⁺ stopping depth is probed through the detection of decay positrons, preferentially emitted along the muons' spin direction.

We use this technique to study two samples: Cu-C_{60} is a magnetic multilayer with the structure (from the bottom): $\text{Ta}(5)/\text{C}_{60}(20)/\text{Cu}(2.5)/\text{C}_{60}(50)/\text{Au}(10)$ –thickness in brackets in nm. $\text{Cu-Al}_2\text{O}_3\text{-C}_{60}$ is a decoupled, diamagnetic reference multilayer with Al_2O_3 layers in between the Cu and C_{60} (Fig. 3a). The uppermost gold film slows injected μ^+ and protects the inner layers from oxidation. The total sample structure is designed to allow the active layers to be probed with a range of accessible μ^+ energies and to maximise the stopping profile at the regions of interest, i.e. close to the Cu-C_{60} interface. The Cu thickness is chosen to obtain the highest magnetisation (Fig. 2a). Muon stopping profiles and further experimental details can be found in the SI, section §S.3 and Figs. S.12-S.13.

Muons with 4 keV implantation energy probe the identical uppermost $C_{60}(50)/Au(10)$ layers of both samples. Nevertheless, the zero field μ SR measurements at 250 K demonstrate a significant difference: for the magnetic Cu-C₆₀ sample a fraction of \approx 10% of the muon spin polarization is rapidly lost -indicating that about 10% of the region sampled by the muons is affected by the magnetism (Fig. 3b). This points to additional sources of magnetic flux in the Cu-C₆₀ sample. In this region, the additional flux most likely arises from stray dipolar fields,

since at low temperatures (20 K) we find that 75% of the μ^+ implanted in the C_{60} layer form a bound electron-muon state called muonium which can only be observed in a non-magnetic environment. This strongly indicates that the C_{60} layer is for the most part free of magnetic moments, and suggests that the magnetism is localized at the Cu- C_{60} interface. Due to the presence of muonium in C_{60} the data analysis is difficult, but further support for this scenario comes from the energy/depth dependence of the μ SR data at low and high temperatures. At 20 K the observable muonium fraction decreases in the magnetic Cu- C_{60} sample as one approaches the Cu layer (SI, Fig. S.14). Analogously, at 250 K the difference between the spectra of the two samples increases for the 6 keV data and is even greater at 8 keV, the energy at which the muons most heavily sample the Cu layer (Fig. 3b). If the Cu were non-magnetic, one would expect an overall increase of the muon polarization, which is obviously not observed.

Another means of locating the magnetism in the Cu-C₆₀ sample is to study its μ SR response in the zero field remanent state. Both samples contain an oscillation at 0.4 MHz due to muonium formed in semiconducting C₆₀. After applying an external field of 300 Oe, this signal is shifted to 0.6 MHz, which we attribute to a small residual field of approximately 0.3 Oe in the apparatus. Nonetheless, the remanent 8 keV μ SR spectra shown in Fig. 3b are clearly different from the virgin spectra for the magnetic Cu-C₆₀ sample, while only subtle changes can be observed for the non-magnetic reference sample Cu-Al₂O₃-C₆₀. The striking new feature related to the remanent state of Cu-C₆₀ is an additional oscillation at \approx 0.75 MHz which is not observed for the non-magnetic Cu-Al₂O₃-C₆₀ (Fig. 3c). This new frequency can be naturally explained by an additional magnetic field of 0.1 Oe at the muonium site in close proximity to the Cu layer. The amplitude of this signal follows the fraction of μ ⁺ stopped in the Cu layer (Fig. 3d), while the non-magnetic signal at 0.6 MHz is anti-correlated to this

fraction. Altogether, the low energy μSR data fully support the notion of a magnetic moment being localized in the metallic layer and the immediate Cu-C₆₀ interface.

To search for the origins of the induced magnetisation, we have modelled the Cu- C_{60} interface via Density Functional Theory (DFT; SI Section §S.4). The molecular roughness of the C_{60} films has been accounted for via several interface models based on: *i.* the single crystal 7-vacancy Cu(111) reconstruction, 20 *ii.* C_{60} encapsulation into adsorbed Cu(111) films $(C_{60}@Cu)$, and *iii.* Cu(111) growth into the pits of the C_{60} -film $(Cu@C_{60})$, see Fig. 4a. Regardless of the adopted DFT flavour (SI, Figs. S.15-S.19), we find a non-magnetic ground state for all the considered interfaces. With the exception of the thicker C_{60} /slab interfaces and $C_{60}@Cu$ models prepared with short (1.5-2 Å) initial Cu-C distances, all the models exhibit a convex (positive) curvature of $DOS(E_F)$ –see figure 4b. Within the mean-field itinerant electron model (SI, Fig. S.20), convex $DOS(E_F)$ can lead, for sufficiently high external magnetic fields, to a spontaneous first-order para to ferromagnetic metamagnetic transition. For the computed $DOS(E_F)$ of the Cu- C_{60} interface models, the critical field (H_c) for the metamagnetic transition sharply decreases with increasing values of the Stoner exchange (I_s) as [1- $I_sDOS(E_F)]^{3/2}$.

Atom-resolved analysis of the I_s parameter reveals up to a factor 4 change of the exchange strength at the Cu-C₆₀ interfaces (Fig. 4c). Magnetic hardening up to a factor of three has been previously reported for magnetic cobalt atoms contacted to π -conjugated molecules.⁵ The computed values of I_s (from 0.86 eV for bulk Cu to up to >2.5 eV for interfacial Cu-atoms) suggest viable para to ferro metamagnetic transitions for fields lower than 1 kOe for thin Cu layers (Fig. 4d). On this basis, we may attribute the measured ferromagnetism to a transition of the Cu-C₆₀ system in magnetic fields of 0.3-5 kOe generated during sample deposition and preparation (see Methods). Our DFT calculations predict that 77 to 95% of the

magnetisation in the Cu- C_{60} system will then be distributed in the metal (SI, Table S.8 and Figs. S.21-S.26), in good agreement with the muon spectroscopy data.

Although the substantial electron transfer from the Cu-layers to C_{60} (≥ 1.6 e/ C_{60} depending on the model; SI, Table S.7) is effective in altering the curvature of DOS(E_F) and increasing I_s , the calculated DOS(E_F)× I_s product remains less than unity (SI, Table S.6), leaving the Stoner criterion unfulfilled. However, magnetometry and muon spectroscopy provide conclusive evidence for the emergence of magnetism at Cu/C_{60} interfaces. This is probably correlated with a sharp decrease of five orders of magnitude in the ferrometamagnetic critical field made possible by C_{60} -induced magnetic hardening of Cu. Similar effects due to charge transfer could also take place in other hybrid metallo-organic 17 and d^0 magnetic systems. 30 In order to maximise the potential of this effect, it should be possible to look for molecules with large electron affinity such as polyoxometalates and metals with a large exchange integral such as zinc. However, good band and structural matching will be needed to obtain significant results. Manipulating the charge transfer by applying electric potentials or through energy band matching could lead to applications in molecular memories or devices such as spin capacitors.

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Figure legends

Figure 1: Effect of molecular interfaces. Schematics and room temperature magnetisation for a $Ta(5)/[C_{60}(15)/Cu(2)]x5/Al(5)$ and a $Ta(5)/[C_{60}(15)/Al(3)/Cu(2)/Al(3)]x5$ samples -film thickness is in brackets in nm. The Cu-to- C_{60} charge transfer and interface reconstruction results in significant changes in the DOS of the metallic film and a band splitting that leads to magnetic ordering. On the other hand, an Al spacer between both materials screens the charge transfer from the 3d transition metal and stops the band splitting.

Figure 2: Room temperature magnetisation for Cu and Mn films. Dashed lines are exponentially modified Gaussian fits. Error bars in thickness constitute the film roughness and in magnetisation they are calculated as the error of the mean. **a.** Dependence of the magnetisation with the Cu film thickness for a total of 145 samples with the structure $Ta(5)/[C_{60}(10-20)/Cu(t)/C_{60}(10-20)]x1-5/Al(5)$. Films with $t \leq 1-1.5$ nm are discontinuous. Inset: Magnetic moment vs. number of $C_{60}(15)/Cu(2.5)$ interfaces —the moment is roughly proportional to the number of interfaces. **b.** As a. for Mn, with 96 samples measured. The magnetisation in Mn films is smaller than for Cu, but propagates for a longer distance. Inset: Out-of-plane and in-plane magnetisation measurements of a $[C_{60}(15)/Mn(2.5)]x4$ sample.

Figure 3: Muon spin spectroscopy (μ SR) at 250 K. Error bars are the standard error in \sim 10⁶ experiment events. Schematic of the and samples measured: Cu-C₆₀: $[Ta(5)/C_{60}(20)/Cu(2.5)/C_{60}(50)/Au(10)];$ and the control Cu-Al₂O₃-C₆₀: $[Ta(5)/C_{60}(20)/Al_2O_3(4)/Cu(2.5)/Al_2O_3(4)/C_{60}(50)/Au(10)]$. Far-right: raw hysteresis loops. b. Zero field µSR spectra for 4, 6, 8 and 16 keV muon implantation energies. Tables show the fraction of muons stopped in each layer/sample. At 8 keV, zero field spectra are shown before (closed symbols) and after an applied magnetic field of 300 Oe (remanent state – open symbols), evidencing clear differences in the Cu-C₆₀ sample. **c.** Oscillation frequencies from fits to the data plotted in **b**. Only in the Cu-C₆₀ sample, a new signal at \sim 0.75 MHz attributed to the emergent magnetisation is observed in remanence. **d**. The polarisation amplitude of the magnetic remanent signal at 0.75 MHz tracks the fraction of muons stopped in the Cu (maximum at 8 keV), whereas the signal associated to muonium at 0.6 MHz is anti-correlated to it.

Figure 4: DFT simulations and metamagnetic modellling. **a.** Schematic of the molecularly rough Cu-C₆₀ interface with: atomically flat C₆₀-Cu contacts (green square), C₆₀ inclusion in Cu-film (C₆₀@Cu; red square) and Cu-inclusion in pits of the C₆₀-film (Cu@C₆₀; black square). The C₆₀@Cu-1.5(55v) model includes 55 Cu vacancies. The optimised atomic structures are reported and labelled in SI, Figs. S.15-S.19. **b.** Total DOS around the Fermi energy (E_F) for bulk-Cu and the Cu-C₆₀ interface models. **c.** Atom-resolved analysis of the Stoner exchange integral for the Cu-atoms as a function of the shortest Cu-C₆₀ distance. **d.** The critical magnetic field (H_c) for spontanous ferromagnet metamagnetic transition as a function of Is for the models of computed positive DOS(E_F) curvature in (b). The horizontal black line marks a typical magnetic field during sample preparation. Shaded area are possible values for I_s in the different geometries.

Methods Magnetic measurements were taken using a superconducting quantum interference device operated as a vibrating sample magnetometer (SQUID-VSM or SVSM) model MPMS3 from Quantum Design with resolution better than 10⁻⁸ emu. The thin films were deposited on 0.5 mm thick Si/SiO₂ substrates. Metals were deposited by DC magnetron sputtering at a pressure of approx. 2.5 mbar (24 sccm of Ar; 10⁻⁸ mbar base pressure) with a deposition rate of 1-3 Å/s. C₆₀ films were deposited by thermal evaporation from a sublimed, 99.9% purity source in an alumina boat in the same chamber at $\sim 10^{-8}$ mbar and with deposition rates of 0.5-1 Å/s. Alumina films were grown via plasma oxidation of Al films: O₂ flow of 76 SCCM, 35 mA current. Oxygen is highly detrimental to the emergent magnetism, and samples grown in a poor vacuum ($P_{total} \ge 2 \times 10^{-8}$ or $P_{O2} \ge 5 \times 10^{-10}$ mbar) show no magnetisation. Ta seed layers are used to improve the sample roughness. Our thermally sublimated C₆₀ films are relatively rough when compared to sputtered metallic films (~1 nm RMS roughness for C_{60} vs. <0.5 nm in metals). The metallic films are continuous and there is no significant diffusion into the molecular film as seen in low angle X-rays (SI, Fig. S.3). Cross-sections of representative samples were analysed by transmission electron microscopy and showed the metallic layers to be continuous and the C₆₀ layers to be polycrystalline (Fig. S.27). The films experience a magnetic field of ~ 0.3 kOe during growth due to an in-plane magnet and the field from the magnetron gun. They are also subject to fields of ~1-5 kOe during loading and centring in the SVSM –needed to position the sample with respect to the SQUID sensor.

Low energy muon spin spectroscopy³¹ utilizes positive muons to provide a probe of local magnetisation. Positive muons (μ^+) are implanted into a sample and decay into a detectable positron and a neutrino/anti-neutrino pair. Due to charge parity violation, there is a preferred direction of emission of the positrons along the muons' spin vector. Determining the direction of the positron decay allows the precession of the muon spin to be determined and, therefore,

the local field at the muon implantation site. A polarized high intensity beam of energetic (MeV) positive muons³² is obtained from the decay of π^+ , generated by a proton beam impinging on a graphite target. After moderation in a cryogenic solid Ar moderator where the beam polarisation is conserved³³ the anti-muons are re-accelerated electrostatically to keV energies and transported by electrostatic elements to the sample. Positrons emitted from muon decay are detected by two plastic scintillator rings and the difference between the flux observed at these two detectors is used to determine the instantaneous spin direction of implanted positive muons as a function of time. The muon asymmetry data plot is then calculated as: A(t) = [NL(t)-NR(t)]/[NL(t)+NR(t)], where NL/R(t) are the background corrected decay histograms of the left and right positron detector, respectively. There, the error of each bin count *n* is given by the standard deviation of *n*. The errors of each bin in A(t) are then calculated by standard error propagation.

Standard, fixed spin-moment³⁴ and non-collinear van der Waals (vdW) corrected Density Functional Theory (DFT) simulations were executed via the Projected Augmented Wave (PAW) method as implemented in the VASP program.³⁵ We used the PBE exchange-correlation (XC) functional,³⁶ a 400 eV plane-wave energy cut-off, (0.2 eV, 1st order) Methfessel-Paxton electronic smearing,³⁷ and a 10 symmetry irreducible k-point grid for the C₆₀/7vac Cu(111)-4x4 models. For C₆₀@Cu(111)-8x8 and Cu(111)-8x8@C₆₀ models only one k-point was used. The adopted atomic-force threshold for geometry optimization was 0.02 eV Å-1. For the 7vac model we relaxed the topmost 5 Cu-layers together with the all the C₆₀ atoms. All the atoms of the C₆₀@Cu(111)-8x8 and Cu(111)-8x8@C₆₀ models were relaxed. In all cases, a vacuum separation of at least 12 Å was present between replicated images of the interface models.

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Supplementary Information is linked to the online version of the paper at www.nature.com/nature.

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