Single-molecule magnets (SMMs) are among the most promising building blocks for future magnetic data storage or quantum computing applications, owing to magnetic bistability and long magnetic relaxation times. The practical device integration requires realization of 2D surface assemblies of SMMs, where each magnetic unit shows magnetic relaxation being sufficiently slow at application-relevant temperatures. Using X-ray absorption spectroscopy and X-ray magnetic circular dichroism, it is shown that sub-monolayers of Dy$_2$@C$_{80}$(CH$_2$Ph) dimetallofullerenes prepared on graphene by electrospray deposition exhibit magnetic behavior fully comparable to that of the bulk.

Magnetic hysteresis and relaxation time measurements show that the magnetic moment remains stable for 100 s at 17 K, marking the blocking temperature $T_B$ being not only in excellent agreement with that of the bulk sample but also representing by far the highest one detected for a surface-supported single-molecule magnet. The reported findings give a boost to the efforts to stabilize and address the spin degree of freedom in molecular magnets aiming at the realization of SMM-based spintronic units.

1. Introduction

Single-molecule magnets (SMMs) belong to the smallest units wherein magnetic moments can remain intrinsically stable on time scales required for possible technological applications.[1–3] Intense research in the field of nanomagnetism within the last decade has further boosted technologically relevant performance indicators, resulting in SMMs magnetically stable above liquid-nitrogen temperature[4] or single atoms exhibiting extremely long magnetic relaxation times.[5–6] In particular, systems based on lanthanide family elements, like Dy and Tb, have been largely in focus, including single-molecule,[2,3] single-atom,[4,5] or single-chain magnets.[7,8] Adsorption of SMMs on surfaces allows to study individual molecular units, as well as to realize transport schemes essential for the implementation of SMMs in molecular-scale spintronics or quantum computing devices.[9–17] However the transition from bulk to surface-supported systems often goes along with a substantial change or even loss of SMM properties, that is, magnetic moment, magnetic anisotropy, or magnetization behavior.[18–21] On metallic surfaces, the interaction of the magnetic moments with the surface is rather strong, which is evidenced by the observation of the Kondo effect.[22,23] Thus, benchmark measurements during the last years demonstrating magnetic bistability of surface-adsorbed SMMs have been reported on substrates, where molecules are electronically weakly coupled to – TbPC$_2$ on HOPG,[24] on MgO/Ag(100)[25] and on graphene/SiC[26] pushing the blocking temperature ($T_B$) limit up to 9 K. On the other hand, DySc$_2$N@C$_{80}$ monolayers on Au(111)[27] recently showed a hysteresis opening at temperatures up to 10 K. In this sense, lanthanide ions encaged in C$_{80}$ molecules reportedly outperform most SMMs by their combination of chemical robustness with slow magnetic relaxation.[27–31] To further push the magnetic lifetime in the monolayer regime two important criteria have to be fulfilled: the first requirement is to synthesize SMM compounds showing intrinsically high $T_B$ in the bulk. The second requires implementation of the appropriate methods for molecular deposition on substrates, which provide sufficient decoupling of the SMM from the surface.

In this work we provide experimental evidence on outstanding slow magnetic relaxation in Dy$_2$@C$_{80}$(CH$_2$Ph) sub-monolayers on a graphene/Ir(111) surface. The Dy$_2$@C$_{80}$(CH$_2$Ph) molecules deposited by the electrospray deposition method are organized into islands as shown by low-temperature scanning tunneling microscopy (STM) imaging. We explore their magnetic properties by means of X-ray absorption spectroscopy (XAS) and X-ray magnetic circular dichroism (XMCD) measurements. The analysis of the magnetic relaxation behavior of Dy$_2$@C$_{80}$(CH$_2$Ph) adsorbed on graphene/Ir(111) yields a...
blocking temperature value not only similar to that of the bulk, but also representing the highest reported one for a molecular magnet on a surface.[25–27]

2. Results and Discussion

The outstanding bulk magnetic properties of Dy$_2$@C$_{80}$(CH$_2$Ph) dimetallocfullerene (see Figure 1a) stem from a single-electron bond connecting the two rare earth atoms within the C$_{80}$ cage. Dy$^{3+}$ ions with their spin states of $J_z = 15/2$ are ferromagnetically coupled in the ground state. Recent studies[28] showed that the easy magnetization axis is aligned along the [Dy$^{3+}$/e–Dy$^{3+}$] moiety, which is oriented roughly perpendicular to the C–C bond connecting the CH$_2$Ph side group to the C$_{80}$ cage. Necessary to stabilize the electronic structure of the molecular core, the benzyl addend represents one of the smallest linker group, which still allows to assemble and controllably address these SMMs in possible applications. On the other hand, it reduces the stability of the compound hampering thermal evaporation and thus requires an alternative deposition route to study their properties on a surface under clean environment.

Here, sub-monolayers of Dy$_2$@C$_{80}$(CH$_2$Ph) molecules were deposited on a graphene/Ir(111) substrate by employing the electrospray deposition (ESD) technique[32–34] (see Section 4 for further details). Our STM measurements show that the majority of molecules constitute isolated molecules lying on the surface. Figure 1b shows an STM topographic image of a molecular island with an apparent height of 0.93 ± 0.05 nm. Depending on the bias voltage, Dy$_2$@C$_{80}$(CH$_2$Ph) molecules show spherical or slightly elliptical shape with mean diameter of about 1 nm as can be seen in Figure 1c. We assign the asymmetry of appearances to be mainly reflecting the position of the CH$_2$Ph side group, and thus giving a reasonable guess for the respective molecule’s orientation. At a bias voltage of 1 V, both elongated shapes as well as double-lobe structures can be distinguished, indicating different spatial orientations of the molecules with respect to the side group, when embedded in close-packed islands.

Figure 1a shows the molecule–substrate interaction for Dy$_2$@C$_{80}$(CH$_2$Ph) on the graphene surface.[27] Furthermore, due to the presence of the side group, the molecules tend to order in a stripy pattern within the islands. Whereas these findings reflect the situation of the majority of molecules in the sample, the molecule–substrate interaction can be better evaluated by imaging a rarely found isolated SMM on the surface. Figure 1d shows the tip-induced displacement of a single molecule during scanning at moderate tunneling parameters. Albeit the weak pinning, we observe the favored adsorption site for Dy$_2$@C$_{80}$(CH$_2$Ph) to be the face-centered-cubic (fcc) region of the graphene moiré superlattice. Furthermore, a small protrusion to the side indicates the side group lying flat on the surface as preferred adsorption configuration. Upon going to dense-packed islands we however do not observe a clear correlation between the molecular assembly within the islands and the moiré superstructure of the substrate. Also the height variation of the molecules within the islands do not show a periodic modulation characteristic for the moiére structure, see Section S1, Supporting Information. We thus consider the substrate–molecule interaction to be substantially weaker than the molecule–molecule interaction for Dy$_2$@C$_{80}$(CH$_2$Ph) SMMs on graphene/Ir(111).

In order to explore the electronic and magnetic properties of the Dy$_2$@C$_{80}$(CH$_2$Ph) SMM monolayer we employ synchrotron-based XAS and XMCD techniques.[19] Figure 2a shows XAS spectra recorded at the Dy $M_5$-edge with both left ($\sigma_-$) and right ($\sigma_+$) circularly polarized light as well as the derived
dimetallofullerenes. The whole Dy coincides with measurements performed on related endohedral μz moment of μz as obtained from sum rule analysis. Values are expressed by the magnetic moments 1289 eV (≡ due to its reduced value compared to 10 B M 5-edge at normal incidence (θ = 0°) pointing at a weak molecule–substrate interaction. [Dy 3 the fullerene body with respect to the addend as well as of the preferred alignment of the easy axis is present. This behavior ordered assembly visible in STM only a minor effect toward a porting Information). These findings show that in spite of the σ evidence and surface normal from θ = 0° (out-of-plane) to θ = 80° (grazing). The red line denotes a fit with the function XMCD = cos(θ), revealing a preferred orientation of the magnetic easy axes of a fraction of all molecules in out-of-plane direction. Measurement parameters: B = 6.8 T, T = 2.5 K. c) Magnetic hysteresis loops (0° and 60°) obtained by recording the XMCD signal at 1289.0 eV upon sweeping the magnetic field with 2 T min⁻¹ at T = 2.5 K and a photon flux of 65.3Φ0. Φ0 = 1 × 10⁻³ photons nm⁻²s⁻¹. The saturation values are expressed by the magnetic moments μs as obtained from sum rule analysis.

XMCD signal calculated as the difference (σ+ - σ−). All spectra are normalized to the maximum in the (σ+ + σ−) curve at 1289 eV (≡XAS0) and the XMCD signal is expressed in percent. The spectral shape indicates a Dy³⁺ (4f⁶) oxidation state[16,37] and coincides with measurements performed on related endohedral dimetallofullerenes.[20,27,25,38] The whole Dy M4,5 range spectra together with the corresponding sum rule analysis is presented in Section S2, Supporting Information. We deduce a magnetic moment of μu = 4.8 ± 1.1 μB per Dy³⁺ ion at normal incidence (θ = 0°). This reflects the previously observed scatter in the adsorption geometry of Dy2@C80(CH2Ph) on graphene/Ir(111) due to its reduced value compared to 10 μB per Dy³⁺ ion in the single molecule.[28,38]

In order to obtain more detailed information about the orientation of the magnetic moments in the sample we track the XMCD signal upon changing the angle between beam incidence and surface normal from θ = 0° (out-of-plane) to θ = 80° (grazing), see Figure 2b and Figure S2b, Supporting Information. Whereas the spectral shape does not change we observe a variation of the relative XMCD intensity between 50–60%. The majority of Dy2@C80(CH2Ph) SMMs in the sub-monolayer thus show no preferred orientation of the magnetic easy axis[30] and only a fraction of molecules has their magnetic moments aligned in out-of-plane direction. The observed variation is also reflected as a slight reduction of the magnetic moment to μu = 3.5 ± 0.7 μB per Dy³⁺ ion at θ = 60° (see Figure S2b, Supporting Information). These findings show that in spite of the ordered assembly visible in STM only a minor effect toward a preferred alignment of the easy axis is present. This behavior is likely caused by the rotational degrees of freedom of both the fullerene body with respect to the addend as well as of the Dy³⁺–e–Dy³⁺ spin center with respect to the C80 cage, both pointing at a weak molecule–substrate interaction.

We further acquire magnetization curves of the sample at normal (θ = 0°) and grazing (θ = 60°) incidence, which are presented in Figure 2c. At a magnetic field sweep rate of 2 T min⁻¹ and T = 2.5 K we observe a hysteresis opening around ±4 T and a coercive field of about 1.2 T, comparable to values obtained by SQUID magnetometry at bulk samples.[28] On the other hand we obtain a reduced ratio of remanent to saturation magnetization Mrem/Msat = 30 % as compared to bulk, indicating a faster relaxation time for Dy2@C80(CH2Ph) in our experiment. The different saturation magnitude of both curves reflect the preferred orientation of Dy magnetic moments in out-of-plane direction.

One of the key parameters in the characterization of surface-supported molecular magnets is the closing temperature Tclose of the magnetic hysteresis curve at a given magnetic field sweep rate. We therefore acquire magnetization curves at sample temperatures of 2.5, 8, 15, and 22 K in normal incidence and plot the results in Figure 3a. The hysteresis loop gradually closes and suggests small but finite opening at 22 K. We calculate the loop area and obtain a linear trend with the temperature, see Figure 3b. Previous benchmark measurements on TbPc 2 on a MgO insulating layer rather suggest an exponential decay,[25] which is clearly not observed in our case. Even performing a defensive evaluation by omitting the point at 22 K due to a very small opening area, the linear fit in the range of 2.5–15 K (solid line in Figure 3b) suggests a closing temperature above 20 K, resembling the value obtained for the bulk.[28] We note that in this case the fit perfectly reproduces the point at 22 K (dashed line in Figure 3b), suggesting its overall validity albeit measurement uncertainty. Our findings demonstrate the robust molecular magnetism of Dy2@C80(CH2Ph) on graphene/Ir(111) that sustains up to record-high temperatures at about 20 K for surface-supported molecular magnets.

We now focus on the magnetization relaxation dynamics of the Dy2@C80(CH2Ph) sub-monolayer. In order to disentangle the contribution of X-ray photon-flux-induced and pure adsorption-related demagnetization we perform time-dependent XMCD relaxation measurements. The results are plotted in Figure 4a–e and Figure S3, Supporting Information. The XMCD signal was recorded as a function of time at Brest = 20 mT after saturating the magnetization at 6.8 T. In all measurements we observe a behavior that can be fitted with an exponential decay XMCD ∝ exp[−t/τ] + XMCD0. Whereas XMCD0 depends on
the thermodynamic equilibrium at $B_{\text{ext}}$, the relaxation time $\tau$ provides a measure for the internal demagnetization dynamics of Dy$_2$@C$_{80}$(CH$_2$Ph) on graphene/Ir(111).\cite{4,18,25} We note that small oscillations visible in the XMCD time traces are related to signal intensity fluctuations due to the top-up operation mode of the synchrotron.

Figure 4b shows the magnetic relaxation time $\tau$ as a function of X-ray photon flux $\Phi$ under normal incidence. We observe a clear flux-induced decrease of the relaxation time for values larger than $\Phi_c \approx 0.4 \Phi_0$, with $\Phi_0 = 1 \times 10^{13}$ photons nm$^{-2}$ s$^{-1}$. The dominant contribution of only one relaxation channel in this regime allows us to deduce the X-ray demagnetization cross-section (see Section S3, Supporting Information). The significant decrease explains the smooth shape of the hysteresis curves in Figures 2c and 3a, both recorded using a photon flux of $\Phi_{\text{large}} = 65.3 \Phi_0$ (see Section 2). The striking feature in the measurement however is a clear plateau at X-ray intensities below $\Phi_c$ where no change of the relaxation time is observed.

Figure 4. Magnetic relaxation of the Dy$_2$@C$_{80}$(CH$_2$Ph) sub-monolayer on graphene/Ir(111). a) Measurement scheme of the experiment as explained in the text. b) Magnetic relaxation time $\tau$ as function of X-ray photon flux in normal incidence and 2.5 K. A horizontal dashed line marks the average value in the regime $\Phi < \Phi_c$, a fit (red) denotes photon-induced demagnetization as dominant process for $\Phi > \Phi_c$. For the vertical axis we use a reciprocal scale. c) Time traces of the XMCD signal recorded at 2.5 K (blue), 8 K (green), and 15 K (orange). Exponential fits with a function $\text{XMCD} \propto \exp(-t/\tau) + \text{XMCD}_0$ are presented as solid lines. d) Temperature-dependency of the relaxation time $\tau$ at 20 mT. As lowest order approximation a linear fit is sketched as solid red line, the determination of $T_B(100)$ is marked. e) Magnetic-field dependency of the relaxation time $\tau$. Diamonds and solid curves denote data obtained by SQUID on a bulk sample (data marked by asterisk is for 0 T) and corresponding fits, respectively.\cite{28} Dashed lines serve as guide to the eye. Error bars for $\tau$ are smaller than the size of data points and thus not shown. f) Magnetic hysteresis recorded at a photon flux of 65$\Phi_0$ (black) and 13$\Phi_0$ (red). The curves are acquired at grazing incidence in order to maximize the signal-to-noise ratio. Measurement parameters: $\theta = 60^\circ, T = 2.5$ K.
marked as dashed line at $\tau_{\text{mol-sub}} = 463 \pm 11$ s. We explain this saturation by a crossover to a time scale that is dominated by intrinsic demagnetization processes. To minimize the influence of X-ray induced demagnetization we thus used a photon flux of 0.28 $\Phi_0 < \Phi_0$ for the following experiments in order to analyze the magnetization dynamics of the system.

As the closing temperature of the hysteresis loop is just a rough estimate of the blocking temperature of a molecular magnet and highly depends on the measurement parameters, $T_{B(100)}$ has been established as reliable characteristics in the SMM community. It denotes the temperature at which the relaxation time of a system amounts to 100 s, independently of the involved relaxation processes. In Figure 4c,d we plot XMCD time traces and the extracted values of $\tau$ as a function of sample temperature. In lowest order approximation the data can be fitted with a linear relation which yields $T_{B(100)} = 21.7 \pm 6.9$ K, being in good agreement with the value estimated from temperature dependence of the hysteresis loop area. This allows us to infer a magnetic blocking temperature of $T_{B(100)} = 17.1 \pm 2$ K, which matches the bulk value. Whereas the positive boundary denotes the standard error of the linear fit, the lower boundary is the difference to the 15 K measurement at which the relaxation time is still considerably larger.

In order to corroborate our experimental results we compare the surface-induced relaxation to bulk behavior studied by SQUID magnetometry,[24] see Figure 4e. Whereas $\tau$ resembles the bulk value at 15 K, it is strongly reduced compared to $\tau_{\text{bulk}}$ at lower temperatures. We assign this suppression to both X-ray induced demagnetization that limits the relaxation time as discussed before as well as to zero-field quantum tunneling of magnetization (QTM) observed in bulk,[28] which can be modified upon surface adsorption.[31] To further evaluate on this, we apply a magnetic field of 0.4 T, leading to an increase of $\tau$ to 941 ± 86 s at 2.5 K for the surface adsorbed SMM. We note that this value represents a lower margin for the magnetic relaxation time due to the present X-ray induced demagnetization, which remains unaffected by the magnetic field. The increase of $\tau$ in a magnetic field points to a suppression of QTM upon lifting the degeneracy of the spin ground states. At the same time it excludes an increased coupling to phonon modes that should become available with larger splitting.[39] Thus a take-over of Raman scattering as observed in bulk is likely to be the main driving mechanism of magnetic relaxation, additionally enhanced by the involvement of substrate phonons. In particular, phonon-induced relaxation pathways upon coupling to acoustic (ZA) phonons of graphene around 58 meV,[40,41] roughly corresponding to the effective barrier height of 615 K for Dy$_2$C$_8$(CH$_2$Ph)$_2$, should be considered. Moreover, the non-negligible coupling to the conduction electrons could open an additional relaxation channel involving spin-flip scattering.[21] However, a more detailed investigation is inevitable to address these points. Apart from these considerations for low temperatures, the exceptional concordance with bulk relaxation dynamics at 15 K as suggested from hysteresis and relaxation measurements supports the bulk-like value determined for $T_{B(100)}$. This furthermore points to a relaxation at 15 K that is likely dominated by an unusual thermally activated Orbach process with low barrier as observed in the bulk,[28] where the graphene surface plays no dominant role anymore. Conventional Orbach relaxation involving the first excited spin states is expected to prevail at even higher temperatures, which is however not assessed in these measurements.

We now record the magnetization curve using the lowest feasible photon flux of $\Phi_{\text{low}} = 0.2\Phi_{\text{large}} = 13.0\Phi_0$ (see Section 4 for further details). We note that the used X-ray flux still exceeds $\Phi_0$ by far, but turned out to be the best compromise between viable signal-to-noise ratio and total measurement time. The hysteresis loops of both photon fluxes for an angle of $\theta = 60^\circ$ are shown in Figure 4f. A strongly enhanced relaxation time leads to a further substantial opening of the hysteresis curve. First, this implies an increase of the coercive field by roughly +25% to 1.3 T. Second, despite a waiting time of 20 s at zero field it furthermore leads to an increase of the magnetic remanence ratio $M_{\text{rem}}/M_{\text{sat}}$ to roughly 50%, representing the largest value found for SMMs adsorbed on surfaces.[25] The substantial opening of the hysteresis curve at low X-ray flux as well as the remarkable agreement of the estimated blocking temperature $T_{B(100)}$ with the bulk value suggests only very weak impact of the graphene substrate on the [Dy$_3$] molecule core and its ligand field. Starting with the first observation of slow magnetic relaxation in surface-supported SMMs over 10 years ago in the sub-Kelvin regime[31] and subsequent improvements towards 10 K,[24–27] this work demonstrates a significant increase of the on-surface blocking temperature compared to all previous studies. Furthermore, our system is excellently suited to address the spin states and magnetization dynamics at a single-molecule level, for example using local probe methods.[42,43]

3. Conclusion

We have demonstrated that the Dy$_2@$C$_{80}$(CH$_2$Ph) SMM retains its magnetic properties upon surface deposition. This is achieved by a combination of the electrospray technique as gentle deposition method together with the use of graphene/Ir(111) as an appropriate substrate. The sub-monolayer shows a hysteresis curve that closes around 20 K, twice the value measured on surface-adsorbed molecular magnets so far. We disentangle the contribution of the X-ray photon flux on the demagnetization dynamics by employing time-dependent XMCD relaxation measurements and obtain a magnetic blocking temperature of $T_{B(100)} = 17$ K. This represents the highest value for surface-supported molecular magnets reported up to date and most notably matches at the same time the bulk blocking temperature. Our study thus provides a new milestone on the path toward on-surface systems with application-relevant working temperatures. Moreover, it demonstrates that the graphene–dimetallofullerene system represents a suitable platform for future studies of complex spin systems at the scale of single molecules, aiming at advanced information processing and memory units.

4. Experimental Section

All samples were prepared in situ. The Ir(111) single crystal (Surface Preparation Laboratory B. V.) was cleaned by repeated cycles of Ar$^+$...
sputtering at 2 kV, heating in an O₂ atmosphere of 5 × 10⁻² mbar at 900–1150 °C and flash annealing in ultra-high vacuum (UHV) up to 1500 °C. Graphene was prepared by exposing the clean Ir(111) surface to an ethylene atmosphere at a pressure of 1.1 × 10⁻² mbar for 20 min while keeping the sample at T = 1200°C. Dy₂@C₈₀(CH₂Ph) molecules were freshly solved in 1,2-dichlorobenzene and deposited in situ by ESD while the sample was kept at room temperature. The used ESD setup is described in detail elsewhere. In the presented XAS/XMCD study a total amount of 0.11 monolayer of Dy₂@C₈₀(CH₂Ph) has been deposited on graphene/Ir(111) to obtain a sub-monolayer coverage. The samples were prepared in situ using the very same procedure both at the low-temperature STM home lab and at the beamline in order to ensure comparable coverage and sample quality.

Scanning tunneling microscopy and spectroscopy experiments were performed in a two-chamber UHV system (base pressure 5 × 10⁻¹¹ mbar), equipped with an Omicron Cryogenic-STM. All STM measurements were carried out in the constant-current mode using grinded and polished PtIr tips (Nanoscore GmbH). The sign of the bias voltage (U) corresponds to the potential applied to the sample.

XAS and XMCD experiments were performed at the X-Treme beamline[44] of the Swiss Light Source at the Paul Scherrer Institute. The measurements were performed in the total electron yield mode, with the magnetic field being aligned antiparallel to the incoming X-ray beam and forming an angle θ with the surface normal of the sample. Special care had been taken in order to center the beam by rechecking the sample position and signal every multiple of 10°. At θ = 80°, the projected beam spot width on the surface is 2.8 mm, being still considerably smaller than the sample width of 4 mm. The measurements were performed using a defocused X-ray beam in order to minimize the photon flux. Accordingly, no significant beam damage could be observed in the spectra throughout the measurements. For magnetization curves no data points were collected within θ = ±0.2 T at a magnetic field sweep rate of 2 T min⁻¹ due to polarization reversal of the coil current, which also implied a waiting interval of around 20 s at zero field. XMCD relaxation measurements were performed by continuously alternating between both polarizations σ⁺ and σ⁻. The resulting curve was then calculated by interpolation and subtraction of both curves, XMCD(t) = σ⁺(t) − σ⁻(t). In order to minimize the photon flux the beam shutter was only opened for t shutter = 0.5 s with a waiting time of t reverse = 22.3 s in between each point due to polarization reversal. The flux for the relaxation measurements thus had been corrected using θ = Φ shutter / t shutter + t reverse [44] with Φ shutter being the respective continuous flux value depending on the beam spot size.[44,45]

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Author Contributions

F.L. and A.A.P. synthesized the Dy₂@C₈₀(CH₂Ph) molecules. F.P. and T.B. performed surface deposition and STM characterization. F.P., T.B., V.E., M.F., V.R., and J.D. performed the XAS/XMCD experiments. M.F. conceived and supervised the experiments. F.P. analyzed STM and XMCD data. F.P. and M.F. drafted the manuscript. All authors discussed the results and contributed to writing the paper.

Data Availability Statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Keywords

dimetallofullerenes, electrospray deposition, graphene, relaxation, single-molecule magnets

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