

Figure 1 | Supramolecular spin-valve device. **a**, Schematic and molecular representation of the $TbPc_2^*$ quantum nanomagnet. A Tb^{3+} ion (pink) is coordinated by two phthalocyanine groups; pyrenyl and hexyl substitutions maximize the supramolecular interaction with sp^2 -carbon materials. The quantum magnet consists of two spin systems: an organic $S=1/2$ radical delocalized over the two phthalocyanine rings and a highly anisotropic $J=6$ spin system localized on the Tb^{3+} metal ion. **b**, Magnetic characteristics of the $TbPc_2^*$ SMM showing the hysteresis loop at 0.04 K for a single crystal of diluted sample measured with a micro superconducting quantum interference device set-up at a field sweep rate of 1 mT s^{-1} . The marked steps are induced by quantum tunnelling at energy-level crossings presented in **c**. **c**, Zeeman diagram for the lowest states with $J_z = \pm 6$, which are split by the four nuclear spin states of $I = 3/2$. The calculation was done with the ligand field parameters of ref. 16. **d**, Top, atomic force micrograph of the supramolecular spin valve. The single-walled nanotube lies on a SiO_2 surface supported by a back gate and is connected to palladium source and drain electrodes. **d**, Bottom, scheme of the supramolecular spin-valve architecture (hexyl and pyrenyl groups are omitted for reasons of clarity).

been shown that the prominent magnetic properties of $TbPc_2$ SMMs remain robust when attached through supramolecular π - π interactions to sp^2 -carbon materials, such as carbon nanotubes²² and graphene²³. However, it is important to note that each molecule when deposited on the nanotube relaxes into a slightly different supramolecular position, leading to a local variation of the physical properties. As a matter of fact, some of them would have a huge QTM probability whereas others would mostly undergo a direct relaxation process.

To increase the non-invasive attachment to SWNTs at very low concentrations, the $TbPc_2$ SMMs used herein were modified by introducing one pyrene group and six hexyl groups into one of the two Pc rings (referred to hereafter as $TbPc_2^*$; see Fig. 1a). Both the pyrene group and the alkyl chains are known to exhibit attractive van der Waals interactions with sp^2 -carbon materials and are used as anchoring points on the nanotube. Moreover, the steric hindrance induced by this ligand prevents recrystallization of the SMM on the nanotube. The anchoring groups steer the supramolecular grafting of the quantum magnet and bring the substituted Pc ring into direct contact with the SWCNT wall. The latter and the Pc ring are conjugated systems, which strongly hybridize through π - π interactions²⁴. The supramolecular sample geometry is shown in Fig. 1d. The external magnetic field is

applied in the plane of the sample (0° corresponds to the normal to the nanotube axis) and a back gate fine-tunes the chemical potential of the SWCNT. Figure 2 shows the electronic-transport characteristics of the supramolecular $TbPc_2^*$ -SWCNT set-up. Differential conductance maps of dI/dV as a function of source-drain voltage V_{sd} and back-gate voltage V_g were taken at 0 T (Fig. 2a) and 1 T (Fig. 2b), corresponding to random and polarized orientations of the $TbPc_2^*$ magnetic moments, respectively. At both magnetic fields, the differential conductance maps exhibit the features of Coulomb diamonds²⁵, typically observed for SWCNT quantum dots in the weak-coupling regime²⁶. Intriguingly, the degeneracy points of most of the Coulomb diamonds are open in the absence of an external magnetic field (0 T, Fig. 2a) and closed in its presence (1 T, Fig. 2b). This is clear evidence of the modulation of an extra tunnelling barrier inside the quantum dot. Further insight can be gained when measuring the magnetic field hysteresis loops of the conductance at $V_{sd} = 0$. In Fig. 2c, the magnetic field is swept between 1 and -1 T resulting in an abrupt switch between a relatively high value ($\sim 1\ \mu\text{S}$) and a lower one ($\sim 100\ \text{nS}$). The characteristics of the hysteresis loops are consistent for each of the Coulomb peaks, as shown in Fig. 2d: the difference in conductance value is plotted between trace (from -1 to 1 T) and retrace (from 1 to -1 T) for each

Figure 2 | Molecular spin-valve electronic-transport characteristics. **a**, Coulomb map of the differential conductance as a function of source-drain voltage V_{sd} and back-gate voltage V_g without magnetic field. The differential conductance is given in the unit of the conductance quantum G_0 . A gap Δ is induced at the degeneracy point when all the magnetic moments are randomly oriented. Magnetic molecules induce effective tunnelling barriers in the SWCNT, hindering the electron flow, except in the case where the electrons have enough energy ($V_{sd} > \Delta/2$) to overcome these barriers. **b**, Coulomb map under a magnetic field of 1 T. The gap is closed, and the standard degeneracy points are recovered. This corresponds to the situation where all molecules are polarized in approximately the same direction. **c**, Zero-bias conductance measured as a function of the magnetic field. The red curve corresponds to the conductance under increasing field (-1 to $+1$ T) and the blue curve under decreasing field. The conductance jumps around zero field are attributed to molecules experiencing quantum tunnelling due to tunnel splittings. The last jumps around ± 500 mT are attributed to a direct relaxation process of a single molecule. When all the magnetic momenta are parallel only one spin carrier can flow easily, whereas when the magnetic moment are antiparallel the electron flow of both spin carriers is hindered. **d**, Intensity of the conductance hysteresis measured as a function of the gate voltage. The colour code corresponds to the difference between magnetic field trace and retrace, given in the unit of G_0 : blue corresponds to negative values, red to positive and white to zero hysteresis. The most intense regions are directly correlated to the Coulomb peaks. **e, f**, Schematic representation of the mechanism involving two TbPc_2^* SMM molecules (A and B) grafted on an SWCNT. With increasing magnetic field, molecule A switches first (**e**), thus leading to an antiparallel configuration of the spin valve with lowest conductance. Each molecule induces localized states in the nanotube through exchange interaction. The value of this interaction is estimated to be around $J = 0.5$ meV. The mismatch between spin levels induces effective tunnel barriers in the SWCNT for both spin polarizations. As a result, the electron flow through the SWCNT is hindered. When molecule B switches (**f**), a parallel configuration is recovered, and because of the level broadening (~ 0.2 meV) this configuration leads to high conductance. The hexyl and pyrenyl groups of the TbPc_2^* SMM are omitted for reasons of clarity.

gate step. The hysteretic behaviour disappears above a bias voltage of about 0.5 mV.

To relate the device characteristics to the magnetic signature of the TbPc_2^* quantum magnets, a detailed investigation of the magnetic field dependence of the largest conductance jumps was carried out. An analysis of the angular dependence of the switching field, which was made by plotting the difference between the trace and the retrace, reveals a very pronounced dependence. The smallest switching fields, which define the easy axis of magnetization, were found to be orthogonal to the SWCNT axis (x direction), whereas along the tube axis (y direction) a hard axis was found (Fig. 3a). Such a very strong angular dependence is in agreement with the Ising-like uniaxial anisotropy of the TbPc_2 -SMM family¹⁶. We compared the orientation of the easy axis with the SWCNT axis and found that the TbPc_2^* molecule is orientated with the Pc rings

parallel to the SWCNT wall, which is in accordance with the supramolecular design maximizing the π - π interactions between the Pc ring and the nanotube.

Quantum tunnelling of magnetization is an inherently stochastic process: the magnetization reversal at a tunnel splitting or through a direct relaxation process is expressed by transition probabilities. By cycling the hysteresis loop many times, the related stochasticity of the magnetic moment reversal can be elucidated (Fig. 3b). Increasing temperatures lead to a continuous reduction of the area of the hysteresis loop (Fig. 3c). At 40 mK the switching field is found to be around 0.5 T progressively declining and finally vanishing above 600 mK. The blocking temperature can be extrapolated to be around 1 K, which is in good agreement with recent reports on a TbPc_2 submonolayer²⁷. Other samples showed similar temperature dependencies (see Supplementary Fig. S5).

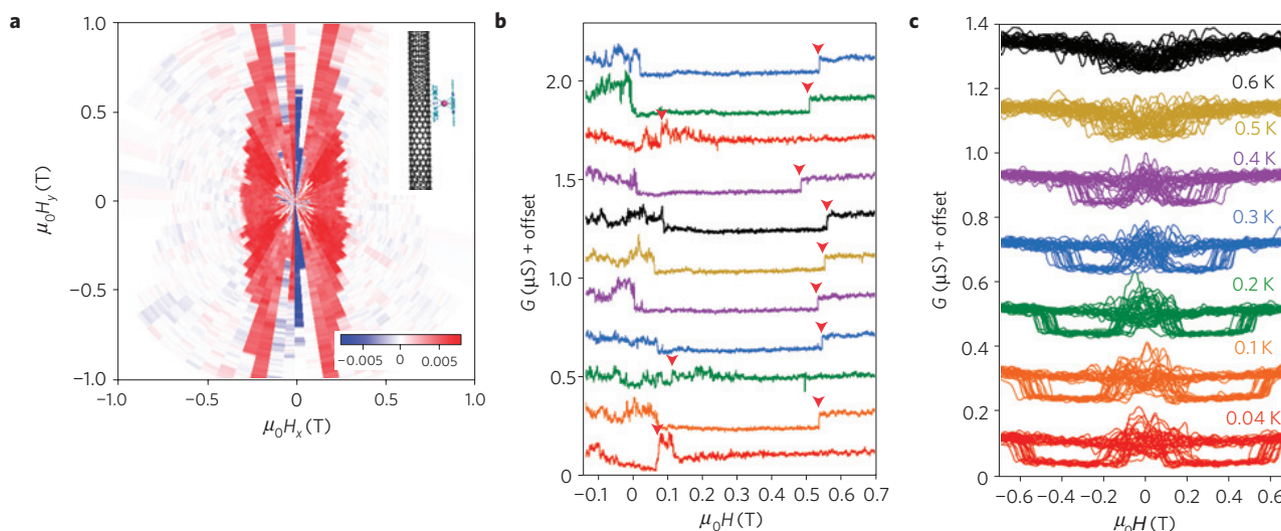


Figure 3 | Molecular spin-valve switching-field characteristics. **a**, Angular dependence of the switching field corresponding to the direct relaxation process. The difference between trace (from -1 to $+1$ T) and retrace (from $+1$ to -1 T) is plotted as a function of the angle of the applied field. The colour code is given in G_0 . The white code corresponds to no difference between trace and retrace for the corresponding field value whereas a red code corresponds to a bistable region. This means that in the white region both molecules are polarized in the same way and in the red region the antiparallel configuration is adopted. As a result, the border between the white and the red region corresponds to the switching field of the molecule experiencing a direct relaxation process. It is important to note that the switching field along the y axis cannot be resolved because our magnets are limited to 1 T. This ellipsoidal behaviour has been measured repeatedly on several samples (see Supplementary Fig. S3), and is in agreement with the Ising-like uniaxial anisotropy of the TbPc_2^* . The x axis can be attributed to the easy axis and the y axis to the hard axis. By comparing the orientation of the easy axis with the nanotube direction it can be deduced that the TbPc_2^* molecule is flat-landing on the nanotube, as shown in the inset. **b**, The conductance is measured 11 times as a function of the magnetic field applied at an angle of 30° with a sweep rate of 2 mT s^{-1} . The switching field of the studied molecule is marked by an arrow. Some of the curves present a transition close to zero instead of 0.5 T. This is attributed to a tunnel mechanism at an avoided level crossing. In this case all the TbPc_2^* molecules have experienced a tunnel transition and the spin-valve behaviour disappears. Most of the tunnel transitions happen at small positive field values, establishing that the Tb nuclear spins are cold enough to involve mainly the lowest magnetic spin states. **c**, 20 hysteresis loops at several temperatures between 0.04 and 0.6 K, using a field sweep rate of 70 mT s^{-1} . A few molecules switch around zero field through a tunnel process whereas one molecule switches at higher fields through a direct relaxation process. The blocking temperature is extrapolated to be around 1 K.

The observation of electronic coupling of the terbium-based $J = 6$ magnetic moment with the conduction electrons is explained by the intermediating presence of the $S = 1/2 \pi$ radical delocalized on the organic phthalocyanine ring systems. Owing to π - π interaction, this radical is supposed to be in close electronic contact with surfaces; for example, it was shown to lead to Kondo features in scanning probe spectroscopy studies²⁸. This means that the wavefunction of this unpaired spin can easily hybridize with the π -electron density of the nanotube. Moreover, transition-metal/Pc systems adsorbed on sp^2 -graphite surfaces show a pinning of the lowest unoccupied molecular orbital level close to the Fermi level²⁹ and recent investigations have shown a weak antiferromagnetic exchange coupling of the Tb magnetic moment of TbPc_2 to a ferromagnetic Ni substrate (P. Gambardella *et al.*, manuscript in preparation). Furthermore, the radical ligand state is close in energy to the Tb $4f$ states³⁰, enabling an efficient coupling of the Tb magnetic $J = 6$ state with the conduction electron of the SWCNT.

The observed magnetoresistance behaviour is explained by an effective tunnelling barrier induced by the magnetic configuration of very few SMMs coupled to the SWCNT quantum dot. The average number of molecules was previously determined to be about four molecules for a source-drain SWCNT segment of 300 nm (ref. 23). The conduction electrons of the SWCNTs are locally influenced through the π -radical-mediated exchange mechanism (*vide supra*) of the confined TbPc_2^* magnetic moment. For reasons of simplicity, the mechanism will be explained for the most ideal case involving two SMM molecules (A and B, Fig. 2e,f) and one spin-degenerate conduction channel: each molecule locally lifts the spin degeneracy, and as a result there is an energy mismatch in the antiparallel configuration between the orbitals

of the same spin, which creates an effective tunnelling barrier. In a spin-valve picture, the first of the molecules (A) plays the role of spin polarizer, whereas the second one (B) acts as spin analyser. The presence of two reversal mechanisms (tunnel splitting versus direct relaxation process) expressing significantly different tunnelling probabilities (see Supplementary Fig. S4a and b) leads to different switching fields for quantum magnets A and B. In the example presented herein molecule A has a close-to-unit QTM probability around zero field (but spread due to the hyperfine coupling) and consequently exhibits magnetic-moment flips close to zero field (Fig. 2e). With increasing magnetic field, the second molecule (B) remains still in the opposite magnetic state, rendering an antiparallel configuration of the spin valve, until this second molecule experiences a direct transition (Fig. 2f), reprogramming the valve into its parallel configuration. Moreover, the quantum behaviour of the second molecule (B) has been investigated through its magnetic field sweep-rate dependence (see Supplementary Fig. S4a). With decreasing fields, around a rate of 2 mT s^{-1} , the spin-valve characteristic disappears in one-third of the cases, which is in accordance with the Landau-Zener theory¹⁸. When molecules A and B tunnel close to zero field, the spin-valve behaviour vanishes (Fig. 3b). The supramolecular spin valve described here thus offers two distinct working regimes: (1) a bi-stable and (2) a sweeping-rate-independent regime. This allows for functional fine-tuning as well as electronic detection and manipulation of a single magnetic moment. It should be noted that other mechanisms invoking any shift of the Coulomb diamond to explain conductance changes (for example magneto-Coulomb effect, see Supplementary Fig. S6) are refuted by Fig. 2d, where no change of the magnetoconductance sign is observed.

In conclusion, a supramolecular spin valve consisting of quantum magnets (SMMs) and SWCNT components has been demonstrated. On reversing the magnetic field, the device exhibits magnetoresistance ratios up to 300%. Analysis of the differential conductance of the SWCNT with the switching-field angle and temperature reveals fingerprint-like characteristics of the magnetic molecules exhibiting Ising-like uniaxial anisotropy and quantum tunnelling phenomena. Our results open a pathway towards the design of operable molecular spintronic devices projecting the implementation of new electrical functionalities, high integration depth and an alternative fabrication scheme to cost-intensive lithographic technologies.

Methods

Sample preparation. SWCNTs of diameter about ~ 1.2 nm grown by the laser ablation method at the Rice University were dispersed in 1–2 ml dichloroethane and sonicated for 1 h. A droplet of this suspension was deposited onto a degenerately doped p-type silicon wafer with a ~ 450 nm surface oxide. SWCNTs were located by atomic force microscopy with respect to predefined markers and then contacted with 50-nm-thick Pd by standard electron-beam lithography with a contact spacing of ~ 300 nm. The TbPc_2^* SMMs were synthesized as previously reported²². The supramolecular grafting was carried out by drop casting a solution of TbPc_2^* diluted in dichloromethane with molarity $M = 10^{-8}$ mol l⁻¹ onto the sample. After 5 s, the sample was rinsed in dichloromethane and dried under nitrogen flow. Residual dichloromethane was removed by a second rinse with isopropanol.

Conductance experiments. Samples with high resistance (>100 k Ω) at room temperature were selected. The conductance measurements were carried out in a $^3\text{He}/^4\text{He}$ dilution refrigerator with a base temperature of 30 mK. Identical stable spin-valve characteristics have been seen on two different samples using this method and on five samples with the same method but with nanotubes grown by chemical vapour deposition (see Supplementary Section S2). The magnetic field in the sample plane was provided by two magnets, generating up to 1 T and up to 0.7 T, respectively. Electrical measurements of interest were made using a Stanford Research Systems SR-830-DSP lock-in amplifier or an ADwin real-time data acquisition system.

Received 16 February 2011; accepted 17 May 2011; published online 19 June 2011

References

- Rocha, A. R. *et al.* Towards molecular spintronics. *Nature Mater.* **4**, 335–339 (2005).
- Bogani, L. & Wernsdorfer, W. Molecular spintronics using single-molecule magnets. *Nature Mater.* **7**, 179–186 (2008).
- Lehn, J.-M. Supramolecular chemistry: From molecular information toward self-organization and complex matter. *Rep. Prog. Phys.* **67**, 249–265 (2004).
- Baibich, M. N. *et al.* Giant magnetoresistance of (001)Fe/(001)Cr magnetic superlattices. *Phys. Rev. Lett.* **61**, 2472–2475 (1988).
- Binasch, G., Grünberg, P., Saurenbach, F. & Zinn, W. Enhanced magnetoresistance in Fe–Cr layered structures with antiferromagnetic interlayer exchange. *Phys. Rev. B* **39**, 4828–4830 (1989).
- Dieny, B. Giant magnetoresistive in soft ferromagnetic multilayers. *Phys. Rev. B* **43**, 1297–1300 (1991).
- Awschalom, D. D. & Flatté, M. M. Challenges for semiconductor spintronics. *Nature Phys.* **3**, 153–159 (2007).
- Dediu, V., Hueso, L., Bergenti, I. & Taliani, C. Spin routes in organic semiconductors. *Nature Mater.* **8**, 707–716 (2009).
- Brede, J. *et al.* Spin- and energy-dependent tunnelling through a single molecule with intramolecular spatial resolution. *Phys. Rev. Lett.* **105**, 047204 (2010).
- Schmaus, S. *et al.* Giant magnetoresistance through a single molecule. *Nature Nanotech.* **6**, 185–189 (2011).
- Sahoo, S. *et al.* Electric field control of spin transport. *Nature Phys.* **1**, 99–102 (2005).
- Aurich, H. *et al.* Permalloy-based carbon nanotube spin-valve. *Appl. Phys. Lett.* **97**, 153116 (2010).
- Hueso, L. *et al.* Transformation of spin information into large electrical signals using carbon nanotubes. *Nature* **445**, 410–413 (2007).

- Ishikawa, N. *et al.* Upward temperature shift of the intrinsic phase lag of the magnetization of bis(phthalocyaninato)terbium by ligand oxidation creating an $S = 1/2$ spin. *Inorg. Chem.* **43**, 5498–5500 (2004).
- Stepanow, S. *et al.* Spin and orbital magnetic moment anisotropies of monodispersed bis(phthalocyaninato) terbium on a copper surface. *J. Am. Chem. Soc.* **132**, 11900–11901 (2010).
- Ishikawa, N., Sugita, M. & Wernsdorfer, W. Quantum tunnelling of magnetization in lanthanide single-molecule magnets: bis(phthalocyaninato)terbium and bis(phthalocyaninato)dysprosium anions. *Angew. Chem. Int. Edn.* **44**, 2931–2935 (2005).
- Christou, G., Gatteschi, D., Hendrickson, D. N. & Sessoli, R. Single-molecule magnets. *MRS Bull.* **25**, 66–71 (2000).
- Wernsdorfer, W. & Sessoli, R. Quantum phase interference and parity effects in magnetic molecular clusters. *Science* **284**, 133–135 (1999).
- Wernsdorfer, W. From micro- to nano-SQUIDS: Applications to nanomagnetism. *Supercond. Sci. Technol.* **22**, 064013 (2009).
- Ishikawa, N. *et al.* Determination of ligand-Field parameters and f electronic structures of double-decker bis(phthalocyaninato)lanthanide complexes. *Inorg. Chem.* **42**, 2440–2446 (2003).
- Zopellaro, G. *et al.* Spin dynamics in the negatively charged terbium (III) bis phthalocyaninato complex. *J. Am. Chem. Soc.* **131**, 4387–4396 (2009).
- Klyatskaya, S. *et al.* Anchoring of rare-earth-based single-molecule magnets on single-walled carbon nanotubes. *J. Am. Chem. Soc.* **131**, 15143–15151 (2009).
- Lopes, M. *et al.* Surface-enhanced Raman signal for terbium single-molecule magnets grafted on graphene. *ACS Nano* **4**, 7531–7537 (2010).
- Wang, Y. Y. *et al.* Single-walled carbon nanotube/cobalt phthalocyanine derivative hybrid material: Preparation, characterization and its gas sensing properties. *J. Mater. Chem.* **21**, 3779–3787 (2011).
- Hanson, R., Kouwenhoven, L. P., Petta, J. R., Tarucha, S. & Vandersypen, L. M. K. Spins in few-electron quantum dots. *Rev. Mod. Phys.* **79**, 1217–1265 (2007).
- Sapmaz, S., Jarillo-Herrero, P., Kouwenhoven, L. P. & van der Zant, H. S. J. Quantum dots in carbon nanotubes. *Semicond. Sci. Technol.* **21**, S52–S63 (2006).
- Otero, L. M., Caneschi, A. & Sessoli, R. X-Ray detected magnetic hysteresis of thermally evaporated terbium double-decker oriented films. *Adv. Mater.* **42**, 5488–5493 (2010).
- Katoh, K. *et al.* Direct observation of lanthanide(III)-phthalocyanine molecules on Au(111) by using scanning tunnelling microscopy and scanning tunnelling spectroscopy and thin-film field-effect transistor properties of Tb(III)- and Dy(III)-phthalocyanine molecules. *J. Am. Chem. Soc.* **131**, 9967–9971 (2009).
- Gopakumar, T. G., Müller, F. & Hietschold, M. STM and STS studies of planar and non-planar naphthalocyanines on graphite I & II. *J. Phys. Chem. B* **110**, 6051–6060 (2006).
- Vitali, L. *et al.* Electronic structure of surface-supported bis(phthalocyaninato) terbium(III) single molecular magnets. *Nano Lett.* **8**, 3364–3368 (2008).

Acknowledgements

This work is partially supported by the DFG programmes SPP 1459 and TRR 88, ANR-PNANO project MolNanoSpin No ANR-08-NANO-002, ERC Advanced Grant MolNanoSpin No 226558, STEP MolSpinQIP and the Nanosciences Foundation of Grenoble. Samples were fabricated in the NANOFAB facility of the Néel Institute. We thank M. Affronte, F. Balestro, N. Bendiab, L. Bogani, E. Bonet, V. Bouchiat, L. Calvet, A. Candini, D. Feinberg, J. Jarvinen, L. Marty, T. Novotny, R. Piquetel, C. Thirion and R. Vincent for discussion and software development. We thank D. Lepoittevin, E. Eyraud, R. Haettel, C. Hoarau and V. Reita for technical support. We thank N.-V. Nguyen and T. Crozes for help in device fabrication.

Author contributions

M.U., M.R. and W.W. designed, carried out and analysed the experiments; J.-P.C. helped to fabricate the devices; S.K. and M.R. designed, synthesized and characterized the molecule; M.U., M.R. and W.W. co-wrote the paper.

Additional information

The authors declare no competing financial interests. Supplementary information accompanies this paper on www.nature.com/naturematerials. Reprints and permissions information is available online at <http://www.nature.com/reprints>. Correspondence and requests for materials should be addressed to M.R. or W.W.