Engineering coherent interactions in molecular nanomagnet dimers

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Proposals for systems embodying condensed matter spin qubits cover a very wide range of length scales, from atomic defects in semiconductors all the way to micron-sized lithographically defined structures. Intermediate scale molecular components exhibit advantages of both limits: like atomic defects, large numbers of identical components can be fabricated; as for lithographically defined structures, each component can be tailored to optimise properties such as quantum coherence. Here we demonstrate what is perhaps the most potent advantage of molecular spin qubits, the scalability of quantum information processing structures using bottom-up chemical self-assembly. Using Cr₇Ni spin qubit building blocks, we have constructed several families of two-qubit molecular structures with a range of linking strategies. For each family, long coherence times are preserved, and we demonstrate control over the inter-qubit quantum interactions that can be used to mediate two-qubit quantum gates.

npj Quantum Information (2015) 1, 15012; doi:10.1038/npjqi.2015.12; published online 8 December 2015

INTRODUCTION

An information processing device whose elements are capable of storing and processing quantum superposition states (a quantum computer) would support algorithms for useful tasks such as searching1 and factoring2 that are much more efficient than the corresponding classical algorithms,3 and would allow efficient simulation of other quantum systems.4 One of the key challenges in realizing a quantum computer lies in identifying a physical system that hosts quantum states sufficiently coherently, and provides appropriate interactions for implementing logic operations.5 Among the molecular spin systems that have been proposed as qubit candidates are N@C₆₀, organic radicals10,11 and molecular magnets.12–14 We proposed exploiting molecular magnets based on heterometallic antiferromagnetic rings.15–16 These systems exhibit a number of favourable features supporting their application as components of a quantum computer: flexibility in their chemical composition allows control over both the total ground-state spin (by modifying the heteroatom) and the carboxylate ligands;17 their well-defined internal magnetic excitations may offer mechanisms for efficient single-qubit manipulations;15 and the ground-state spin is highly coherent,18 particularly when the chemical structure is optimised.19

In the context of molecular spin qubits, the simplest conceivable multi-qubit structure is a molecular dimer. This observation has motivated various efforts to synthesise dimers including, for example, N@C₆₀–N@C₆₀,10,11 radical–radical19–21 N@C₆₀–and molecular magnet (Kaminski D. et al. in preparation) and molecular magnet–molecular magnet.22–25 However, the design of a dimer specifically to host two-qubit experiments should take account of the importance of three key time scales relative to one another: T₂, the individual qubit phase relaxation time, must be longest; h/J, which is characteristic of the duration of two-qubit gates (where J is the inter-qubit interaction energy and h is Planck’s constant) should be intermediate; and the single-qubit manipulation time should be the shortest. In practice, phase relaxation times in heterometallic antiferromagnetic rings are in the 1–10 μs range at low temperatures,18,19 and they can be manipulated in a typical pulsed electron spin resonance (ESR) apparatus on the 10-ns timescale; thus an interaction offering h/J in the 100-ns range could be exploited, for example, in a multi-qubit experiment to generate controlled entanglement. In this Article we report the synthesis of two families of dimers of the antiferromagnetic ring Cr₇Ni and pulsed electron spin resonance (ESR) experiments probing the spin coherence times and the inter-dimer magnetic interactions. We demonstrate that the modular nature of our synthetic approach provides independent control of the molecular nanomagnet components and of the chemical coupling between them. This allows us the flexibility to optimise the physical properties of the dimers with respect to the three key time scales.

METHODS

Synthesis of multi-qubit molecular structures

The molecular structures of the dimers that we studied are shown in Figure 1. The monomer components are based on Cr₇Ni rings in which intra-ring nearest-neighbour antiferromagnetic coupling gives rise to a well-defined S=1/2 ground state. Each dimer family employs a different variety of Cr₇Ni ring, allowing very different approaches to chemical dimerisation.

The first family ([1]A, 1C and 1D in Figure 1) is a collection of hybrid [3]rotaxanes.26 In each of these compounds two Cr₇Ni rings are threaded by a rigid organic molecule. There is no covalent bonding between the two...
The structure of the two-ring compounds in the crystal.

Figure 1. The structure of the two-ring compounds in the crystal. B, yellow balls; C, grey balls; Cr, green octahedra; Cu, blue octahedra; Fe, blue trigonal prisms; N, blue balls; Ni, pale blue octahedra; O, red balls. (a) \([H_2\text{Th}1][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CtBu})_{16}]\) 1A (1B and 1Bd are very similar, differing only by the stopper on the thread); (b) \([H_2\text{Th}1][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CAd})_{16}]\) 1C (c) \([H\text{Th}3][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CtBu})_{16}]\) 2A; (d) \([\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CAd})_{16}]\) 1D; (e) \([\text{Cr}_7\text{NiF}_8(\text{Etglu})(\text{O}_2\text{CtBu})_{15}]\) 2B; (f) \([\text{Cr}_7\text{NiF}_8(\text{Etglu})(\text{O}_2\text{CAd})_{15}]\) 2C.

The [3]rotaxane structures are synthesized by growing the heterometallic rings around rigid thread molecules that contain amine-binding sites. Initially we studied two closely related threads: \([\text{R}-\text{CH}_2\text{CH}_2-\text{NH}-\text{CH}_2-\text{(C}_6\text{H}_4)_{3}-\text{NH}-\text{CH}_2-\text{CH}_2-\text{R}]\) \((\text{R} = \text{Ph}, \text{Th}1; \text{R} = \text{CH}_2\text{CHMe}_2, \text{Th}2)\). For both threads there are three aromatic rings between the two amines, and they differ only in the stopper on the thread. The reaction to make the [3]rotaxanes involves adding the organic thread to a solution of hydrated chromium trifluoride dissolved in pivalic acid (HO\text{2CtBu}); the heterometallic rings then grow around each ammonium cation and the ring is completed by addition of a nickel precursor. We made four related [3]rotaxanes (see Figure 1 and Supplementary Information): \([H_2\text{Th}1][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CtBu})_{16}]\) 1A; \([H_2\text{Th}2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CtBu})_{16}]\) 1B; \([H_2\text{Th}2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CAd})_{16}]\) 1D (with the pivalic acid perdeuterated); \([H\text{Th}1][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CAd})_{16}]\) 1C (HO\text{2CAd} = 1-adamantanecarboxylic acid) in yields between 17 and 35%.

In the [3]rotaxanes the two Cr-Ni rings each contain an octagonal array of seven chromium(III) sites and one nickel(II) site, with each edge of the octagon bridged by a single fluorine and two carboxylates; these molecules are green in colour. The dimer structures can essentially be regarded as an octagonal prism of metal sites. The fluorides lie inside the octagon and form H-bonds to the ammonium of the thread. The distance between the Cr-Ni ring centroids is 1.64 nm, whereas the longest metal-metal distance between rings is 1.94 nm. The ring cavity is sufficiently small that a phenyl- or iso-propyl group can act as a stopper for the rotaxane. In the crystal structure the nickel site is disordered about the eight metal sites of the octagon. We made compound 1Bd because we have found that deuterating ligands in Cr-Ni monomers can significantly prolong the spin coherence times. Similarly spin coherence times in isolated rings can be improved by reducing the mobility of nuclear spins in the ligands, leading us to prepare 1C, which contains carboxylate ligands with rigid adamantyl groups.

Attempts to increase the inter-ring separation by adding aromatic rings to the linker were limited by the low solubility of longer organic chains containing rigid aromatic groups. Instead, we made a new [2]rotaxane by growing the ring around Ph-CH=CH-Ni=CH=CH=CH=CH-Py \((\text{Py} = \text{pyridyl})\), giving \([\text{Th}3][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CtBu})_{16}]\) (Supplementary Figure S2); this [2]rotaxane contains a pyridyl group at the end of the thread and two of these are then coordinated to the axial position of a dimetal tetracarboxylate compound, in this case \([\text{Rh}_2(\text{O}_2\text{CMe})_4]\). This gives a new route to [3]rotaxanes, and results in \([H\text{Th}3][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CAd})_{16}]\) 1D (Figure 1c), where there is a distance of 2.5 nm between the Cr-Ni ring centroids of the [3]rotaxane.

The second family of dimers \((2A, 2B, 2C, and 2D in Figure 1) involves a chiral Cr-Ni ring, \([\text{Cr}_7\text{NiF}_8(\text{Etglu})(\text{O}_2\text{CtBu})_{15}]\) \((\text{H}_2\text{Etglu} = \text{N-ethyl-D-glucamine})\). This structure again contains an octagon of metal sites, but at the centre is a penta-deprotonated N-ethyl-D-glucamine, which bridges between Cr sites using the deprotonated alkoxide side-arms. Fluorides bridge three edges of the octagon, and there are again pivalate
inter-ring magnetic interaction is small compared with the ESR linewidth in energy. This is consistent with the requirement that two-qubit gates should take longer than single-qubit manipulations, a first indication that these dimers are more suitable for quantum information processing experiments than earlier heterometallic ring dimers. X-band pulsed ESR experiments yield spin coherence times, \( T_m \), (given in Table 1) that are broadly consistent with the times measured for monomers, \(^{15}\) reassuring reassurance that the formation of complexes and dimers is not intrinsically detrimental to the quantum phase coherence.

The form of the CW spectra imposes an upper bound on the intra-dimer magnetic interaction, but to measure this interaction precisely in this regime requires coherent methods; double electron–electron resonance (DEER, also known as pulsed electron double resonance, PELDOR) is the established method for so doing. Although originally applied to inorganic model systems, \(^{32}\) in recent years, DEER has been applied extensively to biological systems. \(^{34}\) By modifying pairs of sites in proteins with spin labels such as nitroxyl nitroxides, DEER can be used to measure the dipolar interaction, and therefore the distance, between the labels. \(^{35}\) DEER techniques exploit the fact that, if there is a magnetic interaction between two spin centres, the coherent precession rate of one of the centres is modified by an inversion of the spin state of the other centre. In practice this is achieved through pulsed ESR sequences using two frequencies, such as the standard four-pulse DEER sequence shown in Figure 3a. \(^{15}\) The first two pulses, \( \pi / 2 - \tau - \pi \) at frequency \( v_1 \), set up a Hahn echo \(^{31}\) at a time \( 2\tau \) on one of the spins. As this spin continues to precess, a ‘pump’ \( \pi \) pulse at a second frequency \( v_2 \) at time \( T \) (relative to the \( v_1 \) echo) inverts the second spin, \( v_2 \), inverting the first spin (and therefore the precession rate) experienced by the first spin. A subsequent refocusing \( \pi \) pulse at frequency \( v_1 \) at time \( t_2 \) after the first echo generates a second echo at a time \( 2\tau + t_2 \), with an amplitude that depends on the time at which the \( v_2 \) inversion pulse was applied and on the strength of the interaction between the two spins. Observing this echo as a function of the time, \( T \), of the pump pulse reveals oscillations at the frequency corresponding to the magnetic interaction energy between the two centres. A three-pulse variant, shown in Figure 3b, \(^{31}\) has the advantage of being shorter and having fewer pulses, thus enhancing the amplitude of the measured echo, particularly for systems in which the spin coherence times are short. The disadvantage is that for short times \( T \), the \( v_2 \) pump pulse overlaps the coherence-generating \( v_1 \) \( \pi / 2 \) pulse, distorting the spectra around the time \( T = 0 \) \(^{33}\).

This description of DEER depends on some heterogeneity between the two spin centres in the molecule under study, so that the frequencies \( v_1 \) and \( v_2 \) are resonant with different centres. We performed DEER experiments at low temperatures (2.5 K, to maximise spin coherence times) on dilute frozen solutions of dimers (0.1–0.2 mM, to minimise interdimer dipolar interactions). In this case differences between the two heterometallic rings may arise from conformational flexibility. In the first family, the [3]rotaxane dimers, there may be variation in the position of the Ni centre between the two rings, or some slight bending of the stiff threading molecule. In the second family, the covalently bound dimers, the rings could rotate about the axis defined by the linker. In either case, the distortions of individual rings (giving rise to \( g \)-strains) or variations in configurations of hyperfine-coupled nuclei (resulting in variations of the effective local magnetic field) offer asymmetry between the two rings within a dimer.

The CW spectra in Figure 1 extend over a magnetic field range of about 30 mT at X-band (about 9.5 GHz), corresponding to an absorption spectrum at a fixed magnetic field covering about 750 MHz, as plotted in Figure 3c. Intervals within this energy range can be identified with orientationally selected sub-populations of rings. Thus, microwave pulses at different frequencies within the spectrum excite orientations selectively, shown as coloured regions on spheres in Figure 3d. We anticipate that, notwithstanding a degree of flexibility, the orientations of two rings within a single dimer should be reasonably well correlated. With this in mind, we choose to separate \( v_1 \) and \( v_2 \) by a frequency that is small compared to the total width of the spectrum, so that they excite neighbouring orientational sub-populations of rings.

In practice, the response of the microwave resonator used in the experiment is not uniform over the frequency range of the absorption line. Instead we used fixed frequencies for \( v_1 \) and \( v_2 \), and we achieved orientation selection by adjusting the external magnetic field to bring the appropriate part of the absorption spectrum into resonance with the applied pulses. We used detection pulses (at frequency \( v_3 \)) of 40 ns and pump pulses (\( v_4 \)) of 24 ns, separated in frequency by 80 MHz.

### Table 1: Phase relaxation times and two-qubit gate times

| Compound (inter-ring distance) | \( T_m \) (ns) | \( x \) | Two-qubit gate time (ns) |
|-------------------------------|-------------|--------|------------------------|
| 1A (1.64 nm)                  | 801 ± 3     | 1.615 ± 0.007 | 77                     |
| 1Bd (1.64 nm)                 | 3239 ± 15   | 1.294 ± 0.008 | 84                     |
| 1C (1.64 nm)                  | 2442 ± 11   | 1.209 ± 0.007 | 82                     |
| 1D (2.50 nm)                  | 543 ± 8     | 1.295 ± 0.014 | 225                    |
| 2A (1.89 nm)                  | 608 ± 12    | 1.261 ± 0.002 | 157                    |
| 2B (2.71 nm)                  | 683 ± 4     | 1.446 ± 0.01 | 400                    |
| 2C (3.07 nm)                  | 543 ± 18    | 1.183 ± 0.03  | 550                    |

Hahn echo decays measured at 2.5 K were fitted to a phenomenological stretched exponential function of the form \( A \exp (-T_m/2\tau) \) where \( 2\tau \) was the total duration of the experiment, and \( T_m \) and \( x \) were fit parameters (\( x = 1 \) corresponds to a simple exponential, for which \( T_m = T_1 \)). The two- and \( \pi \)-pulses were, respectively, 140 and 280 ns long in order to suppress proton ESEEM. Solvents used were protonated toluene (samples 1A, 1B, 1C, 2B) or deuterated toluene (1D, 2A, 2C). Concentrations were 100 \( \mu \)M (samples 1A, 1B, 1C) or 200 \( \mu \)M (samples 1D, 2A, 2B, 2C). The two-qubit gate time corresponds to the first minimum in the oscillatory DEER signal. (Note that for compounds 2A, 2B, and 2C, the quoted inter-ring distance is a minimum; the rotational degree of freedom for rings in these structures may extend the ring separation.)
RESULTS

Figure 4 shows typical DEER spectra taken on compound 1A. The left panel shows the amplitude of the final echo as a function of the time, $T$, of the pump pulse for three different orientation selections. Open circles are raw data; the solid curves are filtered using standard tools (the Matlab package DeerAnalysis) to smooth the data and to remove oscillations (electron spin echo envelope modulation, or ESEEM) associated with couplings to proton nuclear moments. To the right of each trace, a spherical intensity map indicates the range of orientations probed at the given applied magnetic field and experimental detection frequency. The right panel shows the Fourier transforms of the time-domain data from the left panels (both raw data, open circles, and filtered data, solid lines).

The dependence of the DEER signal on magnetic field (or, equivalently, orientational sub-population) is strong, owing to several contributory factors. First, the inter-ring dipolar interaction depends on the orientation, $\theta$, of the dimer inter-ring axis with respect to the magnetic field as $3 \cos^2(\theta) - 1$. It therefore varies significantly as a function of the dimer orientation and passes through zero at the magic angle, $\theta \approx 54.7^\circ$. Second, the magnitude by which the detected echo is modulated in the DEER experiment depends on the proportion of dimers excited initially by $\nu_1$ that is also excited, at the other end, by $\nu_2$.

In a magnetic field of 377.84 mT (lowest trace, blue), dimers whose inter-spin axes are perpendicular to the field are predominantly excited. Within this sub-population the range of inter-ring magnetic dipolar interactions is strongly peaked around a single value, giving rise to clearly defined oscillations with a period of about 155 ns. The Fourier transform, which represents the distribution of coupling strengths within the selected orientational sub-population, is correspondingly sharply peaked at about 6.5 MHz. The first minimum in the time-domain data, occurring at about 77 ns, is the duration for inversion of the state of one of the qubits under the influence of the second. This evolution of the state of one qubit conditional on the state of the other is exactly the kind of physical interaction that allows for implementation of multi-qubit quantum logic; in our case, the time to the first minimum corresponds to the duration of a two-qubit conditional phase gate. We note that this time satisfies the criteria that we identified above for two-qubit devices: the two-qubit gate time (of order 100 ns) lies between the single-qubit
gate time (of order 10 ns) and the phase relaxation time (in the
microsecond range).

The sub-population excited at a magnetic field of 385.84 mT
(middle trace, green) is comprised of dimers whose inter-ring axes
lie at orientations peaked in between parallel and perpendicular
to the external magnetic field. The distribution of orientations
includes a significant proportion close to the magic angle at which
the dipolar interaction strength goes through zero, resulting in a
broad distribution of interaction strengths and an almost
monotonic dependence of the time-domain DEER signal on $T$

At 390.84 mT, $\nu_1$ excites principally rings whose planes are
perpendicular to the magnetic field (i.e. those whose effective $g$-factor is close to $g_z$). Deviations from this orientation lead to a
greater contribution to the effective $g$-factor from the larger $g_{\perp}$, and therefore to a higher resonant frequency. Thus, in dimers that
are orientationally selected by the excitation of one ring at
frequency $\nu_1$, $\nu_2$ (which is lower in frequency than $\nu_1$) is
comparatively unlikely to excite the other ring. This gives rise
directly to the comparatively weak modulation in the time-domain

Figure 4. DEER data for sample 1A. The detection and excitation frequencies were, respectively, $\nu_1 = 9.5632$ GHz, $\nu_2 = \nu_1 - 0.08$ GHz = 9.4832 GHz. The temperature was 2.5 K. The DEER signal was recorded as a function of the time of the $\nu_1$ pump pulse (left panels) at three magnetic fields, each corresponding to an orientational sub-population indicated by the intensity maps on spheres. The Fourier transforms of the time-domain data are shown on the right. Open circles show raw data; solid lines indicate filtered data. The principal difference between the unfiltered data and the filtered data is the absence in the latter of a component at about 18 MHz arising from proton ESEEM.

Figure 5. DEER data for sample 2A. The detection and excitation frequencies were, respectively, $\nu_1 = 9.5260$ GHz, $\nu_2 = \nu_1 - 0.08$ GHz = 9.4460 GHz.
axis) than in 1A (the centroids of whose rings are separated by 1.6 nm), and the fact that the dipolar interaction varies with separation, r, as 1/r³.

These trends are reflected in the DEER data for all of the compounds shown in Figure 1; the time-domain data and Fourier transforms for compounds other than 1A and 2A are presented in the Supplementary Figures S2 to S6, and the two-qubit gate times are listed in Table 1. There is a direct correlation between the interaction strength and the inter-ring separation, with larger separations giving rise to weaker interactions. The two-qubit gate times reach about 550 ns for the longest dimers. The interactions are weaker in the the second family than in the first, owing to the extra degree of freedom in the structure: rings that are attached to the linker through a single point on the rim of the ring are free to rotate about the linker axis, allowing a greater inter-ring separation.

We note also that our strategy allows for enhancement of the spin coherence times independently of the inter-ring interaction strength. In earlier studies of monomers,19 we found that deuteration and immobilisation of nuclear spins in the structure significantly improve the coherence. Thus, in compounds 1A, 1Bd and 1C, while the similar linkers ensure that h/|Ω| is similar for each, deuteration (1Bd) and confinement of hydrogen nuclei to rigid adamantyl groups (1C) result in extensions of the coherence times by factors of about four and three, respectively.

**DISCUSSION**

From the data presented here, we may draw several conclusions supporting the assertion that molecular spin systems may form useful components in future quantum technologies. Molecular magnets offer building blocks which, through supramolecular chemistry, can be assembled into multi-qubit structures. Identifying strict criteria for the relative timescales for single-qubit manipulations, two-qubit interactions and phase memory times, we have designed several families of structures to optimise the prospects of hosting two-qubit devices. For example, in compound 1A, the phase coherence time of about 800 ns significantly exceeds the two-qubit gate time of 77 ns, and both are much longer than the typical single-qubit manipulation time of about 10 ns. Thus, we build upon the wide range of proposals to exploit molecular nanomagnets in quantum information by demonstrating for the first time coherent manipulation of dimers of such species, including multi-frequency experiments probing the inter-qubit interaction, and by showing that we can chemically tune the interaction strength to meet architectural requirements. This prepares the path for experiments demonstrating controlled entanglement operations (as opposed to the observation of entanglement as an inherent property of a singlet ground state of two coupled spins27), and other two-qubit manipulations.

**ACKNOWLEDGEMENTS**

This work was supported by the EPSRC through the National EPR Service and grants EP/H012613/1, EP/H011714/1, and EP/L018470/1 and EP/K039547/1.

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Supplementary Information accompanies the paper on the *npj Quantum Information* website (http://www.nature.com/npjqi)