Reversible Mechanical Switching of Magnetic Interactions in a Molecular Shuttle

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An acid–base switchable molecular shuttle based on a [2]rotaxane, incorporating stable radical units in both the ring and dumbbell components, is reported. The [2]rotaxane comprises a dibenzo[24]crown-8 ring (DB24C8) interlocked with a dumbbell component that possesses a dialkylammonium (NH₂⁺) and a 4,4'-bipyridinium (BPY²⁺) recognition site. Deprotonation of the rotaxane NH₂⁺ centers affects a quantitative displacement of the DB24C8 macroring to the BPY²⁺ recognition site, a process that can be reversed by acid treatment. Interaction between stable 2,2,6,6-tetramethylpiperidine-1-oxyl (TEMPO) radicals connected to the ring and dumbbell components could be switched between noncoupled (three-line electron paramagnetic resonance (EPR) spectrum) and coupled (five-line EPR spectrum) upon displacement of the spin-labelled DB24C8 macroring. The complete base- and acid-induced switching cycle of the EPR pattern was repeated six times without an appreciable loss of signal, highlighting the reversibility of the process. Hence, this molecular machine is capable of switching on/off magnetic interactions by chemically driven reversible mechanical effects. A system of this kind represents an initial step towards a new generation of nanoscale magnetic switches that may be of interest for a variety of applications.

Living organisms make extensive use of biomolecular machines to perform functions crucial for life.[1] A large variety of synthetic molecular machines exhibiting controlled displacement of their components has been produced in the past three decades,[2,3] and laboratory demonstration of the potential utility of molecular machines in information and communication technology (ICT),[4] materials science,[5] catalysis,[6] and medicine[7] has been recently reported. Nevertheless, the exploitation of molecular movements to perform tasks is still a formidable research challenge, and more effort is necessary to extend the range of possible functionalities that may be accessed through the control of motion at the nanoscale.

An interesting possibility in this regard is the exploitation of molecular movements to enable or prevent distance-dependent processes. For example, mechanical switching of photoinduced intercomponent energy-, electron-, or charge-transfer processes has been observed in suitably designed systems.[8,9] On the other hand, the efficiency of such processes can be used to measure distances at the molecular scale[10] and therefore monitor the operation of the system, as is routinely done with Förster resonance energy transfer (FRET) processes in mechanical devices based on nucleic acids.[11]

Another intriguing option consists of incorporating radicals in the molecular components and using electron paramagnetic resonance (EPR) spectroscopy to investigate changes in the magnetic properties as the device operates.[12] So far, spin labelling for intramolecular magnetic switching purposes has been mainly limited to photochromic compounds,[12] and mechanically interlocked molecules containing persistent radicals have only been used to carry out EPR-based co-conformational investigations.[13]

Herein we present the first example of an acid–base-control-lable molecular shuttle in which through-space magnetic interactions between two mechanically interlocked nitrooxide units can be switched on and off reversibly by chemical stimulation. The mechanical switch (Figure 1) is based on a well-known [2]rotaxane architecture[14] comprising a dibenzo[24]crown-8 (DB24C8) ring interlocked with a dumbbell component that possesses two different recognition sites, namely, a dialkylammonium (NH₂⁺) and a 4,4'-bipyridinium (BPY²⁺) unit. A pair of stable 2,2,6,6-tetramethylpiperidine-1-oxyl (TEMPO) radicals are respectively attached to the ring and to one extremity of the dumbbell.

We envisaged that, in spite of the large (co-)conformational freedom of the rotaxane, the acid–base-driven displacement of the macrocycle between the NH₂⁺ and BPY²⁺ stations[14] could bring the two TEMPO radicals close together or far apart, thus affecting their through-space magnetic interaction (Figure 1). The synthesis of rotaxane 1-H3PF₆ and of its components 2 and 3-H3PF₆ is outlined in Scheme 1. The TEMPO-functionalized DB24C8 derivative 2 was obtained in 70% yield by condensation of the macrocyclic alcohol 4 with 4-carboxy-TEMPO 5 using N,N′-dicyclohexyl-carbodiimide (DCC) and 4-dimethylaminopyridine (DMAP) as the condensing reagents. The paramagnetic dumbbell 3-H3PF₆ was prepared in 50% yield by reacting the half-dumbbell 6-H2PF₆ and the commercially available 4-(2-iodoacetamido)-TEMPO (7) in acetonitrile under reflux for 18 h followed by anion exchange.

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To ensure that the TEMPO unit is large enough to behave as an efficient stopper for the DB24C8 ring, we initially studied the interlocking between the diamagnetic host and the half-dumbbell $6\cdot 2PF_6$. The reaction was carried out in chloroform to favor the formation of the hydrogen-bonded pseudorotaxane $4\cdot C27\cdot 6\cdot 2PF_6$, in which the ring encircles the NH$_2^+$ recognition site of $6\cdot 2H_2^+$, and the mixture was then held at reflux for four days in the presence of the paramagnetic stopper $7$. The reaction progress was followed by thin-layer chromatography, and the corresponding [2]rotaxane was isolated in 30% yield. The robustness of the mechanical link was checked by measuring a 1D Rotating-Frame Nuclear Overhauser Effect (ROESY) spectrum of the sample in CD$_3$CN (see Supporting Information) following selective irradiation of the peak of methylene protons close to the ammonium site. The Nuclear Overhauser Effect (NOE) spectrum reveals significant host–guest enhancement of signals corresponding to the OCH$_2$ protons of the ring, indicating that the NH$_2^+$ station is encircled by the crown ether and dethreading does not take place. Similarly, a threaded complex is not formed upon heating a mixture of $4$ and dumbbell $3\cdot 3PF_6$, confirming that the TEMPO moiety is a suitable stopper in the present system.

The target biradical rotaxane $1\cdot 3PF_6$ was made by threading $6\cdot 2PF_6$ with the radical-armed DB24C8 2 and using the TEMPO derivative 7 to introduce the second stopper (Scheme 1). Also, in this case, the reaction was monitored by thin-layer chromatography, and $1\cdot 3PF_6$ was isolated in 30% yield by means of a silica gel column and anion exchange. The interlocked nature of $1\cdot 3PF_6$ was confirmed by $^1H$ NMR spectroscopy (see Figure 2 a,b).

The shuttling of the macrocycle towards the secondary BPY$^{2+}$ station (Figure 1) was obtained by treating the $1\cdot 3PF_6$ rotaxane in CD$_3$CN with the non-nucleophilic Hüning’s base, diisopropylethylamine ($i$Pr$_2$EtN), which is strong enough to deprotonate the NH$_2^+$ center.$^{[14b]}$ Replacement of the ring onto the ammonium station was triggered by addition of trifluoro-
acetic acid. The addition of iPr$_2$EtN (2 equiv) caused the disappearance of CH$_3$NH$_2$H"CH$_2$ peaks in the $^1$H NMR spectrum of the rotaxane (Figure 2B) which were replaced by new signals of lower frequencies ascribed to CH$_3$NHCH$_2$ protons (chemical shift $\delta$ ~ 3.7 ppm, Figure 2C). Bipyridinium H$_x$ and H$_y$ undergo shifts of $\Delta\delta = -0.19$ and $+0.27$ ppm respectively, and H$_b$ and H$_c$ show downfield shifts of $\Delta\delta = +0.09$ and $+0.19$ ppm. A similar trend to H$_x$ and H$_y$ protons (shielding/deshielding, respectively, upon deprotonation) is also detected for the viologen-adjacent CH$_2$ hydrogens (protons f and e), which undergo shifts $\Delta\delta = -0.15$ and $+0.13$ ppm, respectively.

The $^1$H NMR spectrum recorded after the successive addition of CF$_3$COOH was superimposable to the original one (Figure 2B), indicating that the ring shuttles back to the NH$_2$H$^+$ station and highlighting the reversibility of the switching process.

An important result of these experiments is that the radical centers do not interfere with the movement of the ring. Finally, the changes in the magnetic interaction between the two TEMPO radical units of rotaxane 1H·3PF$_6$ (obtained from 1H·3PF$_6$ after addition of iPr$_2$EtN). Red dashed lines evidence the shift of guest protons caused by the movement of the paramagnetic crown ether, and blue dashed lines evidence the macrocycle signals. The signals marked with a star are relative to the protons of the amine.

The ratio of line intensities, however, differs substantially from the 1:2:3:2:1 pattern expected in case all the (co-)conformations of the [2]rotaxane were characterized by strong spin-exchange between the nitroxide units. The combination of three- and five-line EPR patterns observed after addition of iPr$_2$EtN is due to the overlap of the signals arising from 1) biradicals in which the two spin labels are too far apart to interact (three-line spectrum), and 2) biradicals in which the TEMPO moieties are similarly close to undergo exchange coupling (five-line spectrum). By measuring the relative EPR line intensities, we could estimate that, in acetonitrile at 298 K, about 15% of the deprotonated rotaxane molecules assume (co-)conformations such as that shown schematically in Figure 1, bottom, in which the two nitroxide units are sufficiently close to one another to allow a strong electron exchange ($J \gg q_a$).

Such behavior is a consequence of the relatively large flexibility of the connectors that link the TEMPO centers to the rotaxane components and of the (co-)conformational degrees of freedom of the ring and dumbbell relative to each another. More rigid systems are required to exploit the distance effect on through-space spin exchange interactions in a better way.

The addition of a stoichiometric amount of trifluoroacetic acid after the addition of the base caused the quantitative recovery of the initial three-line EPR spectrum (Figure 3C) as a consequence of the replacement of the macrocyclic ring to the ammonium station. The complete base- and acid-induced switching cycle of the EPR pattern was repeated for six times without an appreciable loss of signal, highlighting the reversibility of the process (see inset of Figure 3). Similar EPR results were also observed at a higher temperature (328 K).

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**Figure 2.** Partial $^1$H NMR spectra (600 MHz, CD$_3$CN, 298 K) of the dumbbell 3H·3PF$_6$ (A), the rotaxane 1H·3PF$_6$ (B), and the rotaxane 1·2PF$_6$ (C) (obtained from 1H·3PF$_6$ after addition of iPr$_2$EtN). Red dashed lines evidence the shift of guest protons caused by the movement of the paramagnetic crown ether, and blue dashed lines evidence the macrocycle signals. The signals marked with a star are relative to the protons of the amine.

**Figure 3.** Room temperature EPR spectra of rotaxane 1H·3PF$_6$ (0.15 mm) before (A) and after sequential addition of 2 equiv iPr$_2$EtN (B) and CF$_3$COOH (C). Right inset: enlargement of spectrum B showing exchange lines. Left inset: co-conformations (%) showing spin-exchange as function of sequential acid–base additions.
In summary, we have developed an acid–base switchable molecular shuttle based on a [2]rotaxane, incorporating stable radical units in both the ring and dumbbell components. The chemically driven shuttling motion was exploited to reversibly change the distance between the two spin centers, thereby enabling or preventing through-space electron exchange interactions that can be measured by EPR spectroscopy. Hence, this molecular machine is capable of switching on/off magnetic interactions by chemically driven reversible mechanical effects. A system of this kind can represent a first step towards a new generation of nanoscale magnetic switches that may be of interest for information and communication technology applications.[15] Spin labelling in molecular machines is also interesting because correlations between radical pairs, monitored by pulsed EPR methods, enable detailed conformational analyses, dynamics studies, and precise measurements of nanoscale distances.[18] Hence, they represent a powerful tool to increase our understanding of the intricate operation mechanisms of molecular machines. Experiments in this direction are underway in our laboratories.

Acknowledgements

This work was supported by the Italian Ministry of Education, University and Research (PRIN 2010CX2TLM InfoChem, PRIN MULTINANOITA) and the University of Bologna (Finanziamenti di Ateneo alla Ricerca di Base, SLaMM Project).

Keywords: EPR · molecular devices · rotaxane · spin labelling · supramolecular chemistry

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Received: September 26, 2014
Published online on October 22, 2014