Allosteric Initiation and Regulation of Catalysis with a Molecular Knot


School of Chemistry, University of Manchester, Oxford Road, Manchester, M13 9PL, UK

*Correspondence to: david.leigh@manchester.ac.uk

Abstract: Molecular knots occur in DNA, proteins and other macromolecules. However, the benefits that can potentially arise from tying molecules in knots are, for the most part, unclear. Here we report on a synthetic molecular pentafoil knot that allosterically initiates or regulates catalyzed chemical reactions by controlling the in situ generation of a carbocation formed through the knot-promoted cleavage of a carbon-halogen bond. The knot architecture is crucial to this function as it restricts the conformations that the molecular chain can adopt and prevents the formation of catalytically inactive species upon metal ion binding. Unknotted analogues are not catalytically active. Our results suggest that knotting molecules may be a useful strategy for reducing the degrees of freedom of flexible chains, enabling them to adopt what are otherwise thermodynamically inaccessible functional conformations.

One Sentence Summary: A synthetic molecular pentafoil knot is found to allosterically regulate or initiate catalyzed chemical reactions, requiring coordination to metal ions to generate a halide binding site that promotes formation of the key carbocation.

Molecular knots are found in circular DNA (1) and approximately 1% of proteins in the Protein Data Bank (PDB) (2), and form spontaneously in polymer chains of sufficient length and flexibility (3). Molecules with the topology of some of the simplest prime knots have been synthesized (4-13) but, although knots are fundamental elements of structure, the potential benefits that could arise from tying molecules in knots are mostly unclear (14-18). Recently, Fe(II) complexes of some synthetic molecular knots and links were found to strongly and selectively bind halide anions (X⁻) within their central cavities (19). This binding activity resembles a key feature of dehalogenase enzymes, which contain halide binding sites that facilitate the cleavage of carbon-halogen bonds (20, 21). Here we show that as little as 1 mol% of a synthetic molecular pentafoil knot can induce Lewis acid catalyzed reactions by the in situ generation of a carbocation by carbon-halogen bond scission promoted by CH···X⁻ hydrogen bonding and long range metal-cation···X⁻ electrostatic interactions.

We previously reported the synthesis of a molecular pentafoil knot (a 5₁ knot in Alexander-Briggs notation (22)), inspired by Lehn’s cyclic double helicates (23) but modified so as to be assembled, and the end groups connected, through imine chemistry (24, 25). The knot binds a chloride or bromide anion extremely strongly within its central cavity ($K_{\text{Cl}} = 3.6 \times 10^{10}$ M⁻¹, $K_{\text{Br}} = 1.7 \times 10^{10}$ M⁻¹ in MeCN) through a combination of CH···X⁻ hydrogen bonding and long range Fe(II)···X⁻ electrostatic interactions (19). However, attempts to remove the metal ions whilst maintaining the pentafoil knot topology proved unsuccessful due to the lability of uncoordinated imine bonds (25). We therefore investigated an alternative route to a metal-free pentafoil knot based on ring-closing olefin metathesis (RCM) of a tris(bipy) ligand strand (1) that had proved
successful in the synthesis of a related Star of David catenane (26) (Fig. 1, detailed experimental procedures and full characterization data are given in Sections S1, S4-S8 of the Supplementary Material).

Fig. 1. Assembly of wholly organic pentafoil knot 2 and its reversible metallation with zinc(II) ions to form [Zn₂2](BF₄)₁₀. Reagents and conditions: (i) FeCl₂, dimethylsulfoxide, 60 °C, 24 h; (ii) saturated aqueous KPF₆ solution (89 % [Fe₅₂•Cl](PF₆)₉ over two steps); (iii) Hoveyda-Grubbs 2nd generation catalyst, 1,2-dichloroethane/nitromethane (1/1), 60 °C, 24 h; (iv) saturated aqueous KPF₆ solution (98 % [Fe₂2•Cl](PF₆)₉ over two steps); (v) saturated aqueous NH₄BF₄
then saturated aqueous NH₄PF₆ (90 % [Fe₅2](PF₆)₁₀); (vi) Na₄EDTA, acetonitrile/water (1/1), 80 °C, 4 h (69 % 2); (vii) Zn(BF₄)₂, dichloromethane/methanol (3/1), 40 °C, 2 h (98 % [Zn₅2](BF₄)₁₀); (viii) Na₄EDTA, acetonitrile/water (1/1), RT, 3 h (98 % 2).

Heating ligand strand 1 with 1.25 equiv. of FeCl₂ in dimethylsulfoxide (DMSO) at 60 °C resulted in an intensely-colored red-purple solution indicative of the formation of low-spin Fe(II)–tris(bipy) complexes (Fig. 1, step i). After 24 h the pentameric circular helicate [Fe₅1•Cl](PF₆)₉ was isolated in 89 % yield by precipitation with aqueous KPF₆ (Fig. 1, step ii) and characterized by ¹H nuclear magnetic resonance (NMR) spectroscopy (Fig. 2B) and electrospray ionization mass spectrometry (ESI-MS) (Figs. S48 and S49). The end groups of each ligand strand were successfully connected to form the pentafoil knot [Fe₅2•Cl](PF₆)₀ by RCM using the Hoveyda-Grubbs second generation catalyst (Fig. 1, step iii) (27). The ¹H NMR spectrum of [Fe₅2•Cl](PF₆)₀ (Fig. 2C) confirms the absence of terminal alkene protons (cf. precursors 1 and [Fe₅1•Cl](PF₆)₀, Figs. 2A and 2B) and also two distinct environments for some sets of aromatic protons (e.g. H⁶, H⁷, H⁸) as a result of restricted rotamerization around the phenyl-bipyridine bond after RCM.

**Fig. 2.** Partial ¹H NMR spectra (600 MHz, 298 K, CD₃CN/CDCl₃ (3/1): A) ligand 1; B) cyclic pentameric helicate [Fe₅1•Cl](PF₆)₀; C) Fe(II)-pentafoil knot [Fe₅2•Cl](PF₆)₀; D) metal-free pentafoil knot 2; E) Zn(II)-pentafoil knot [Zn₅2](BF₄)₁₀. The signal assignments correspond to the labeling shown in Fig. 1. Signals from residual CHCl₃ are shown in grey.
Single crystals of [Fe₅₂•Cl](PF₆)₉ were obtained by slow evaporation of a solution of the knot in a mixture of acetonitrile and toluene containing excess NaBPh₄, and the solid-state structure determined by X-ray crystallography (Supplementary Section S8). The crystal structure (Fig. 3) confirmed the topology of the molecular pentafold knot as a single 190-atom loop woven around five Fe(II) cations to create the five crossing points. As with the related imine-pentafold knot, a chloride anion (present despite the large excesses of PF₆⁻ and BPh₄⁻ anions used in the purification and crystallization steps) is positioned within the central cavity of the pentafold knot. The chloride ion is displaced by ~1.47 Å from the plane of the Fe(II) cations (Fig. 3B) and is in close contact with the ten electron-poor aromatic CH⁺ protons oriented towards the middle of the cavity (CH···Cl distance 2.46–2.85 Å). The remaining anions are a ~2:1 mixture of PF₆⁻ and BPh₄⁻.

Fig. 3. X-ray crystal structure of pentafold knot [Fe₅₂•Cl](PF₆)₉(BPh₄)₃, shown as the Λ⁵- enantiomer (Λ⁵-[Fe₅₂•Cl](PF₆)₉(BPh₄)₃ is present in equal amounts in the unit cell). A) Viewed from above the central cavity of the knot. B) Viewed in the plane of the Fe(II) cations. Hydrogen atoms, solvent molecules and anions (except for the central Cl⁻) are omitted for clarity. The carbon atoms of one precursor ligand strand are colored light blue and the others light grey. N, dark blue; Fe, purple; Cl, green. Fe–Fe distances (Å): 8.305(7), 8.241(6), 8.313(7), 8.389(7), 8.431(7). Fe–Fe–Fe angles (°): 107.80(7), 106.86(7), 108.37(7), 108.14(7), 108.82(7). CH···Cl distances (Å): 2.461, 2.527, 2.539, 2.553, 2.629, 2.653, 2.669, 2.677, 2.714, 2.858. Hydrogen
bond distances were normalized using neutron diffraction data. Crystallographic data and experimental details of the structural refinement of $[\text{Fe}_2\cdot\text{Cl}]\text{(PF}_6)_9\text{(BPh}_4)_3$ are provided in Section S8 of the Supplementary Information.

Unlike the imine pentafoil knot (24), the metal ions (and simultaneously the chloride ion) could be removed from $[\text{Fe}_2\cdot\text{Cl}]\text{(PF}_6)_9$ by treatment with aqueous tetrasodium ethylenediaminetetraacetate (Na$_4$EDTA) solution at $80\,^\circ\text{C}$, forming the metal-free knotted ligand 2 in 69 % yield (Fig. 1, step vi; Supplementary Material, p. 9). The $^1\text{H}$ NMR spectrum of 2 in CD$_3$CN/CDCl$_3$ (3/1) (Fig. 2D) is relatively broad in comparison to the metallated knot. Molecular modeling suggests that 2 does not exist in a single well-defined conformation and the broadness of the $^1\text{H}$ NMR spectrum is apparently a consequence of reptation (the thermal motion of long entangled polymer chains, which influences properties such as the viscous flow of amorphous polymers (28)) as the 190-atom-long knot backbone slowly slithers through transiently tightened loops and across over and under crossings. Despite possessing no elements of Euclidean chirality, the metal-free knot 2 is intrinsically chiral by virtue of its topology. The enantiomers could be separated by chiral high performance liquid chromatography (HPLC) (Supplementary Section S4) and the pentafoil knots of opposite handedness shown to have circular dichroism spectra of equal and opposite shape and sign (Fig. S22).

The iron-coordinated pentafoil knot binds a chloride or bromide ion strongly within its central cavity ($K_{\text{CT}} \sim 2.6 \times 10^{10} \text{ M}^{-1}$ (MeCN), $K_{\text{Br}}^{-} 1.4 \pm 1.0 \times 10^{10} \text{ M}^{-1}$ (MeCN), Supplementary Section S9). Nevertheless, the halide anion could be removed while leaving the Fe(II) ions in place, forming $[\text{Fe}_2\cdot\text{Cl}]\text{(PF}_6)_{10}$, by treating $[\text{Fe}_2\cdot\text{Cl}]\text{(PF}_6)_9$ or $[\text{Fe}_2\cdot\text{Br}]\text{(PF}_6)_{10}$ first with saturated aqueous NH$_4$BF$_4$, which exchanges the majority of the anions—including the centrally bound halide—for BF$_4^-$, and then saturated aqueous NH$_4$PF$_6$, which replaces the BF$_4^-$ anions with PF$_6^-$ (19) (Fig. 1, step v; Supplementary Material, p. 7 and 8).

The reaction of tris(bipy) ligand strand 1 with divalent transition metal salts other than Fe(II) results in linear triple helicates, coordination polymers or complex product mixtures (29). Therefore it might seem that only iron(II)-complexed knots are accessible with this system. However, treatment of the metal-free pentafoil knot 2 with Zn(BF$_4$)$_2$ in a mixture of CD$_2$Cl$_2$/CD$_3$OD (3/1) (Fig. 1, step vii), gave an initially broad $^1\text{H}$ NMR spectrum (Fig. S4) that sharpened after two hours at 40 °C, indicating rearrangement to a single molecular species. After workup, the formation of the zinc(II) coordinated pentafoil knot $[\text{Zn}_2\cdot\text{Cl}]\text{(BF}_4)_{10}$ was confirmed by $^1\text{H}$ NMR spectroscopy (Fig. 2E) and ESI-MS (Figs. S57 and S58). The restrictions on the chain conformation imposed by the topology of 2 prevent formation of linear triple helicates (as predominantly occurs with 1 and Zn(II) salts) and enable the ligand to adopt the higher energy circular helicate structure with zinc(II) ions that is thermodynamically inaccessible with 1. Zinc(II) ions typically exhibit much faster coordination kinetics than low-spin iron(II) ions and the Zn(II)-pentafoil knot could be demetallated to reform 2 (Fig. 1, step viii) more readily and under milder conditions than the Fe(II)-pentafoil knot (Fig. 1, step vi). The Zn(II)-pentafoil knot binds a chloride or bromide anion within its central cavity approximately a tenth as strongly as its Fe(II) counterpart ($K_{\text{CT}} \sim 2.9 \times 10^{9} \text{ M}^{-1}$ (MeCN), $K_{\text{Br}}^{-} 7.7 \pm 2.0 \times 10^{8} \text{ M}^{-1}$ (MeCN); Supplementary Section S9).
Substantial recent catalysis research has leveraged anion-binding molecules to influence reactions by ion pairing (30) or, in some cases, promoting carbon-halogen bond cleavage (31). Even with the relatively modest anion binding affinities of the host molecules generally employed to date the approach has been shown to be effective across a range of reaction types (31). The empty cavity metallated pentafoil knots are amongst the strongest noncovalent binding synthetic hosts of Cl\(^-\) and Br\(^-\) known, readily sequestering traces of Cl\(^-\) from the environment, including solvents and glassware. We wondered whether such a strong halide binder might be able to carry out reactions that have traditionally been the preserve of halophilic silver salts. We envisaged that the knot might be able to promote the formation of carbocations by stabilizing the Br\(^-\) counter-anion formed by heterolytic carbon-bromine bond cleavage reactions.

We found that bromide ions bound within the cavities of either [Fe\(_5\)2•Br](PF\(_6\))\(_9\) or [Zn\(_5\)2•Br](BF\(_4\))\(_9\) react with methyl triflate to form bromomethane gas and the empty knots [Fe\(_5\)2](PF\(_6\))OTf and [Zn\(_5\)2](BF\(_4\))OTf, respectively (Supplementary Section S2.1). This reaction could be used to recycle the halide-binding cavity of the knot in situ, for example using the Zn(II)-pentafoil knot as a catalyst for the hydrolysis of bromodiphenylmethane (Ph\(_2\)CHBr) (Fig. 4A and Supplementary Section S2.2). Note that MeOTf does not promote the hydrolysis reaction in the absence of the knot. Other nucleophiles (e.g. MeCN) also react efficiently with the catalytically generated benzhydryl (Ph\(_2\)CH\(^+\)) carbocation (Supplementary Section S2.3).
In addition to acting as a catalyst in its own right, the metal-coordinated pentafoil knot could be used to generate other catalytic species, such as Lewis acidic trityl carbocations through anion abstraction (Fig. 4B). The Michael addition and Diels-Alder reactions of acrolein (3) with 2-acetylcylohexadiene (6), respectively (Fig. 4B, Tables S1 and S2), can each be promoted by Lewis acidic carbocations (32, 33). We confirmed that neither of these reactions occurs in the absence of trityl bromide (Tables S1 and S2, entry 1) and proceed only slowly in the presence of trityl bromide alone (Tables S1 and S2, entry 2). Addition of Fe(II)(bipy)$_3$, the basic metal coordination unit of the knot, does not increase the reaction rate above the background level (Tables S1 and S2, entries 3 and 4), nor does the addition of the tris(bipy) ligand strand 1, even in the presence of FeCl$_2$ (any pentameric circular helicate formed would have a Cl$^-$ anion already strongly bound to the halide binding site) (Tables S1 and S2, entry 5). Neither the Fe(II)- nor Zn(II)-pentafoil knots ([Fe$_2$]$_2$(PF$_6$)$_{10}$ and [Zn$_2$]$_2$(BF$_4$)$_{10}$) lead to catalysis of the reactions in the absence of trityl bromide (Tables S1 and S2, entries 11 and 14). However, in the presence of trityl bromide, both [Fe$_2$]$_2$(PF$_6$)$_{10}$ (Tables S1 and S2, entry 13) and [Zn$_2$]$_2$(BF$_4$)$_{10}$ (Tables S1 and S2, entry 15) initiate catalysis of the reactions effectively affording the endo-Diels-Alder (7) adduct in 62 % (Fe(II)-knot) and 82 % (Zn(II)-knot) and Michael adduct (5) in 83 % (Fe(II)-knot) and 98 % (Zn(II)-knot) yields, respectively (Figs. S6-S9).

Further evidence that the catalysis arises from in situ formation of the trityl cation by abstraction of a bromide anion into the knot cavity was provided by treating a solution of trityl bromide with [Fe$_2$]$_2$(PF$_6$)$_{10}$. UV-Vis spectroscopy confirmed the presence of the trityl carbocation (Fig. S14), showing its characteristic twin absorption features at $\lambda$ = 425 and 405 nm, whilst $^1$H NMR spectroscopy and mass spectrometry confirmed the contemporaneous formation of [Fe$_2$•Br](PF$_6$)$_9$. In contrast, treating trityl bromide with the demetallated form of pentafoil knot 2 did not generate the carbocation (Fig. S20) and no catalysis was observed upon addition of 3 and 4 or 6 (Tables S1 and S2, entry 12).

The binding of one type of molecular species (an effector) to induce structural change and alter binding or catalysis at a second site—allostery regulation—is a central part of the enzyme feedback loops biology uses to maintain homeostasis (34). Allostery control of Friedel-Crafts reactions has previously been demonstrated by a synthetic catalyst using coordination control of hydrogen bonding sites (35). However, treating the metal-free knot 2 with iron(II) salts led to
Our results suggest that tying molecules in knots may prove to be a useful strategy for reducing the degrees of freedom of flexible macrocycles and chains, enabling them to adopt otherwise thermodynamically inaccessible conformations that can perform useful tasks. By analogy it may be that one biological role of knotting is to prevent protein chains from adopting low energy, but inactive, folded states. The formation of a binding site by the organization of a long molecular strand by allosteric metal ion binding is also reminiscent of proteins. The use of a strongly halophilic, and chiral, metal-coordinated knot to promote the cleavage of carbon-halogen bonds may bring advantages of chemo- and stereoselectivity in chemical reactions traditionally promoted by silver salts.

References and Notes:


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46. OriginPro 8.51 from OriginLab®.

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**Supplementary Materials:**

Materials and Methods

Figures S1-S66

Tables S1-S3

References (36-46)