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Transmembrane Ion Channels Formed by a Star of David \([2]\)Catenane and a Molecular Pentafoil Knot

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Supporting Information Placeholder

ABSTRACT: A \((\text{Fe}^{II})_6\)-coordinated triply-interlocked ("Star of David") \([2]\)catenane \((6^7\text{ link})\) and a \((\text{Fe}^{II})_5\)-coordinated pentafoil \((5^1)\) knot are found to selectively transport anions across phospholipid bilayers. Allostery, topology and building block stoichiometry all play important roles in the efficacy of the ion transport properties. Multiple \(\text{Fe}^{II}\) cation coordination by the interlocked molecules is crucial: the demetallated catenane exhibits no anion binding in solution nor any transmembrane ion transport properties. However, the topologically-trivial, Lehn-type cyclic hexameric \(\text{Fe}^{II}\)-helicates—which have similar anion binding affinities to the metallated Star of David catenane in solution—also display no ion transport properties. The unanticipated difference in behavior between the open and closed-loop structures may arise from conformational restrictions in the linking groups that likely enhances the rigidity of the channel-forming topologically-complex molecules. The \((\text{Fe}^{II})_6\)-coordinated Star of David catenane, derived from a hexameric cyclic helicate, is two orders of magnitude more potent in terms of ion transport than the \((\text{Fe}^{II})_5\)-coordinated pentafoil knot, derived from a cyclic pentamer of the same building block. The reduced efficacy is reminiscent of multi-subunit protein ion channels assembled with incorrect monomer stoichiometries.

INTRODUCTION

The orderly entanglement of molecular strands within knots\(^1\) and links\(^2\) (catenanes) can induce properties and characteristics\(^1,12,22\) that are beginning to be explored in areas as diverse as anion binding,\(^3\) catalysis,\(^4\) materials,\(^5\) healthcare\(^6\) and the kinetic stabilization of supramolecular structures? One strategy to synthesize examples of complex molecular topologies is to join the ends of building blocks that assemble into cyclic metal double helicates.\(^1,8,9\) In a typical example, by tuning the assembly conditions the same set of components can be enticed to selectively form either a five-membered or six-membered Lehn-type cyclic helicate.\(^8\) Subsequent macrocyclization of the building block end-groups by ring-closing metathesis gives the corresponding pentafoil \((5^1)\) knot\(^4a\) or Star of David triply-interlocked \([2]\)catenane \((6^7\text{ link})\),\(^9c\) respectively (Scheme 1).

The metallated knot, link and the parent open cyclic helicates display good-to-very-strong halide binding affinities in their central cavities \((K_a \approx 10^{5–10^10} \text{ M}^{-1} \text{ in MeCN})\). The anion-binding properties of the interiors of these metal-coordinated molecular structures led us to investigate their potential as transmembrane ion channels or transporters.\(^10\) Protein ion channels typically have complex secondary, tertiary and quaternary structures that determine their transport properties.\(^11\) For example, membrane-spanning proteins often consist of \(\alpha\)-helix bundles or \(\beta\)-barrels, which confer rigidity and anchor the protein within the membrane. In contrast, small-molecule systems are generally much less complex\(^12\) and rely on intermolecular interactions, such as hydrogen bonding\(^13\) and aromatic stacking,\(^13c,14\) to promote their assembly into rigid membrane-spanning ion channels. Metal ion-ligand coordination can also be used to assemble ionophoric supramolecular structures, and a number of metalloorganic ion transporters have also been described.\(^15\) We reasoned that the high anion affinities\(^3a\) and the rigid shape with internal cavity imposed by the topology of the metallated pentafoil knot \(2^k\) and Star of David link \(3^c\) could make them suitable candidates for ion transportation (Scheme 1). Furthermore, the two molecular structures are assembled from the same repeat unit, \(1\), allowing the direct comparison of assemblies of the same building block in two different stoichiometries.

A series of assays in phospholipid vesicles were used to examine the ion transport behavior and selectivity of \(2\) and \(3\) (Figure 1). The metallated pentafoil knot was used as the mono-chloride salt (other anions \(\text{PF}_6^-\)), \(2\), because the fully hexafluorophosphate salt rapidly exchanges one \(\text{PF}_6^-\) for a Cl- ion by sequestering traces of chloride from glassware or solvent.\(^5\) The somewhat less halophilic metallated Star of David catenane was used as the fully hexafluorophosphate salt, \(3\). The ion transport abilities of the metallated knot and link were compared with that of the demetallated Star of David catenane, \(4\), which without transition metal coordination exhibits no anion binding in solution, and \(\text{Fe}^{II}\)-coordinated Lehn-type open cyclic hexameric helicates \((5\) and \(6\), to determine the influence of (i) metal-binding allostery and (ii) molecular topology (Figure 2). The results of the vesicle experiments
were corroborated by single-channel planar bilayer conductance measurements (Figure 3).

**EXPERIMENTAL DETAILS**

The Fe\(^{II}\)-coordinated pentafoil knot 2\(^a\) and the Star of David [2]catenane 3,\(^{9c}\) were prepared from ligand 1 in the presence of different Fe\(^{II}\) salts followed by ring closing olefin metathesis (Scheme 1). The ionophoric activity of the compounds in vesicles was initially assessed in 8-hydroxypyrene-1,3,6-trisulfonate (HPTS) assays using 4:1 v/v egg yolk phosphatidylcholine (EYPC)/cholesterol vesicles in 3-(N-morpholino)propanesulfonic acid (MOPS) buffer (15.2 mM lipid, [MOPS] = 20 mM, pH 7.4).\(^{16}\) In these experiments, vesicles containing HPTS and different salts were prepared ([salt] = 100 mM, see ESI, Section S3.1). Either knot 2 or link 3 (2.5 µM, 0.26 mol% relative to EYPC/cholesterol) was added to the solution containing the vesicles, followed by a pulse of NaOH (6.5 µM). Ionophoric activity was indicated by dissipation of the resultant pH gradient between the interior and exterior of the vesicles, which was monitored through the change in the relative fluorescence of HPTS (Figure 1A). Finally, the vesicles were lysed by addition of a detergent (Triton X-100) to allow normalization of the fluorescence response. The resulting kinetic data was fitted to pseudo first-order rate equations as an approximation\(^{13f,17}\) to compare the effectiveness of the analyzed compounds (see ESI, Table S1–S3). Background rates for ion leakage across the membranes for a range of salts were measured (see ESI, Figure S6, S7) and the values subtracted from the knot/link-mediated ion transport rates.

Scheme 1. Synthesis of (Fe\(^{II}\))\(^5\)-coordinated pentafoil knot complex 2 and (Fe\(^{II}\))\(^6\)-coordinated Star of David [2]catenane complex 3 from ligand 1. Pentafoil knot 2 is isolated and used with a chloride anion bound in the central cavity. The Star of David [2]catenane 3 was treated with Na\(_4\)EDTA to generate the demetallated link 4.

\(^a\)Reagents and conditions: (a) FeCl\(_2\), dimethylsulfoxide, 60 °C, 24 h followed by excess saturated aqueous KPF\(_6\) solution; (b) Hoveyda-Grubbs 2nd generation catalyst (H\(_2\)IMes)Cl\(_2\)RuCH(o-O\(\text{Pr}\)C\(_6\)H\(_4\)), 1,2-dichloroethane/MeNO\(_2\) (1:1), 60 °C, 24 h followed by excess saturated aqueous KPF\(_6\) solution; (c) FeSO\(_4\)·7H\(_2\)O, ethylene glycol, 170 °C, 24 h followed by excess saturated KPF\(_6\) in methanol; (d) Na\(_4\)EDTA (EDTA = ethylenediaminetetraacetic acid), H\(_2\)O/MeCN (1:1), 80 °C, 4 h.
Figure 1. (a) Schematic representation of the 8-hydroxypyrene-1,3,6-trisulfonate (HPTS) assay used to determine the ionophoric activity of 2 and 3 in vesicles; internal pH change can occur through any combination of antiport (M+/H+, A−/OH−) or symport (M+/OH−, A−/H+), as shown.13 (b) Normalized fluorescence data for HPTS assays with EYPC/cholesterol vesicles formed in the presence of KBr, following the addition of (FeII)5-coordinated pentafoil knot 2 (blue), (FeII)6-coordinated Star of David link 3 (red) or demetallated Star of David link 4 (grey). NaOH base pulse (13 µL, 1 M) at 1 min, Triton X-100 (20 µL, 10% v/v solution) at 6 mins (EYPC/cholesterol 4:1 v/v vesicles, [compound] = 2.5 µM, 100 mM KBr, 20 mM 3-(N-morpholino)propanesulfonic acid (MOPS), pH 7.4). (c) Observed pseudo-first order rate constants determined from HPTS assays for the transport of different salts ([3] = 2.5 µM, 100 mM MX (M = Li+, Na+, K+, Rb+, Cs+, X = F−, Cl−, Br−, I−; SCN−, NO3−), 20 mM MOPS, pH 7.4). The background rate was high for the small, basic F− anion, nonetheless it showed a low transport rate (< 0.005 s−1) when this background rate was accounted for.

RESULTS AND DISCUSSION

We first examined the ionophoric activity of the knot and link with respect to KBr (Figure 1B). The (FeII)5-coordinated pentafoil knot 2 displayed weak ion transport activity (kobs = 4.05 × 10−4 s−1, see ESI Figure S9), while (FeII)6-coordinated Star of David catenane 3 was >50-fold more active (kobs = 2.30 × 10−2 s−1, see ESI Figure S8). The pentafoil knot and Star of David link both showed significant ion transport, and differ only in the number of units of 1 included in their assembly (five and six, respectively). This difference alters both the size of the central hole and the number of cations associated with each structure. It is somewhat reminiscent of the outcome of subunit assembly with the wrong stoichiometry in some biological channels and pores; mis-assembly with respect to the number of building blocks still results in a functional ion channel but with greatly reduced efficacy of transport.18

The Fe6-coordinated knot and link are significantly different to the metal-free Star of David catenane ligand 4, in that the bound metal ions confer structural rigidity, add multiple positive charges, and inductively withdraw electron density from the pyridine rings, all of which create the strong anion binding site in the central cavity (ESI Figures S1, S3). As expected from

Figure 2. (a) Structure of Lehn-type hexameric cyclic helicates 5 and 6. (b) Representative HPTS assay with (FeIII)6-coordinated Star of David catenane 3 and hexameric cyclic helicate 6 ([compound] = 2.5 µM, 100 mM NaCl, 20 mM MOPS, pH 7.4). NaOH base pulse (13 µL, 1 M) at 1 min, Triton X-100 (20 µL, 10% v/v solution) at 6 mins.
its lack of anion binding affinity in solution, the demetallated Star of David link 4 showed no activity in the HPTS assay. An intact, in the form of the coordination of FeII cations, is an essential feature to enable both anion binding in solution and anion transmembrane transport by 2 and 3. Nonetheless, the charge provided by complexation to FeII is not in itself sufficient to cause ion transport; simple Fe(bipy)3 complexes, such as FeCl2(bipy)3, showed no ion transport in analogous experiments (ESI, Figure S12).

To determine the significance of the interlocked molecular topology in the ion transport experiments we also tested two Lenn-type circular metal helicates: methyl-derivative 5 and 6, the immediate precursor to 3 (Figure 2). These metal-coordinated complexes have anion affinities in solution that are very similar to the (FeII)6-coordinated Star of David link 3. Remarkably, given the activity of the topologically complex compounds, 2 and 3, neither 5 nor 6 showed any activity in HPTS assays with a range of salts (for HPTS assays with 5, see ESI Figure S10; for HPTS assays with 6, see Figure 3B and ESI Figure S2). The inactivity of 6 is particularly as it differs from the highly active (FeII)6-coordinated Star of David catenane 3 only in that it has not undergone ring-closing olefin metathesis. It appears that the conformational constraints imposed by joining of the strand ends is crucial for the metallated Star of David catenane 3 to be able to form transmembrane ion channels (although the knot is less effective in doing so). The less ordered ligand exterior is apparent in molecular modelling of helicate 6 (ESI, Figure S2).

Having established the strong ionophoric activity of the (FeII)6-coordinated Star of David link 3, we examined its anion selectivity (Figure 1C, ESI Figure S8). The metallated Star of David link 3 elicited ion transport decreasing in the order: KSCN ~ KI > KBr > KCl > KOAc ~ KN03 > KP. The halides and pseudo-halides follow the Hofmeister series, suggesting that desolvation may be the rate-limiting factor for ion transport. However, the transport rates of perchlorate and nitrate differ from that expected from their positions in the Hofmeister series (perchlorate sits between iodide and thiocyanate; nitrate sits between bromide and chloride). Haynes et al. observed a similar effect for Zn11V15 cages, which was attributed to size/shape-exclusion hindering passage of the larger perchlorate and nitrate anions. Changing the cation showed much less variation in transport rates (Figure 1C, ESI Figure S8d, Table S2), demonstrating that the ionophores are only selective between anions. Anion transport was confirmed by high levels of chloride transport in lucigenin assays (see ESI Section S3.2).

The mechanism of transport was probed to determine whether the (FeII)6-coordinated pentaofoil knot 2 and Star of David catenane 3 were merely disrupting the phospholipid bilayer, acting as an ion carrier or forming ion channels. 5(6)-Carboxyfluorescein assays ruled out vesicle lysis or the formation of large, nonspecific channels (see ESI, Section S3.3), while U-tube experiments demonstrated that both the metallated knot 2 and link 3 did not act as carriers (see ESI, Section S3.4). These observations suggest that the HPTS assay data arise from anion-selective channels formed in vesicle membranes by the metallated knot and link.

We further investigated the ion channel activity of the molecular knot and link by way of single-channel planar bilayer conductance (PBC) experiments (see ESI, Section S4 and Figure 3). A planar bilayer was formed between two wells containing buffer (20 mM MOPS, 1 M KCl, pH 7.4) and a potential applied across the membrane. The channel-forming compound was then added to one well (ESI, Figure S19); resulting step-changes in the ion current indicating the formation of ion channels. While we did not observe current steps with the metallated pentaofoil knot 2 within the 2 h duration of the measurement, addition of the metallated Star of David link 3 led to reproducible, quantized, square-topped ‘flicker’ events (Figures 3A, S20 and S21), indicative of short-lived ion channels. Increased channel formation was observed under negative applied potentials compared to when a positive potential was applied (Figure 3B). This is consistent with the positively charged link 3 being driven towards the membrane. The channels have a high conductance of around 0.22 nS (measured at +100 mV, Figure 3C), mirroring the high ionophoric activity observed in the HPTS assays (Figure 1). Hille analysis (ESI, Section S4.4) of the PBC data allowed the diameter of the channels formed by 3 to be estimated at 5.0–7.2 Å.

A Hill plot analysis of the HPTS assays for KCl transport by (FeII)6-coordinated Star of David catenane 3 (see ESI, Figure S11) gave an EC50 value of 1.3 μM (17.4 μg mL−1) and a channel stoichiometry of 1 (i.e. a channel is made by a single molecule of 3). While Hill plot analysis has many underlying assumptions, in combination with the observed ion selectivity it suggests that the formation of single catenane ion channels able to transport anions (e.g. perhaps through a ‘relay-race’ mechanism) is a reasonable explanation for the observed ionophoric activity.

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**Figure 3.** Single-channel planar bilayer measurements of ion channel formation by (FeII)6-coordinated Star of David cation 3 (EYPC lipid/cholesterol (4:1, w/w), [3] = 8 μM, 1 M KCl, 20 mM MOPS, pH 7.4, 293 K). (a) Single-channel recording at +100 mV showing the opening (red) and closing (grey) of a single ion channel. (b) All-points data analysis (see ESI, Section 4.3) of the ion currents recorded at +100 mV. The data were fitted to two Gaussian distributions and the difference in ion current measured. (c) Current-voltage curves showing the average ion current flow from +100 to −100 mV.

The overall picture that emerges from the experimental studies of the (FeII)6-coordinated Star of David 3 is that the high ionophoric activity arises from membrane insertion to form channels, which are likely to involve a single molecule of 3 (Hill plot analysis, n = 1, ESI, Figure S11). Given its size and relatively hydrophobic edges (ESI Figure S3), we speculate that the catenane could embed in the hydrophobic membrane with its faces parallel to either side of the otherwise intact membrane. Although the thickness of the catenane is signifi-
cantly less than the thickness of the bilayer, the bilayer may thin around the channel-former as observed for the shorter antibiotics in the peptaihol family.31

The markedly better ionophoric activity of the Star of David catenane 3 (constructed from six molecules of building block 1) over pentafoil knot 2 (made from five molecules of 1) may be a consequence of the weaker binding of anions (such as Cl\(^-\), Br\(^-\), I\(^-\)) in the larger diameter (\(\sim 0.46\) nm) central hole of 3 (Figure 4).34 The X-ray crystal structure of pentafoil knot 2 shows the tightly bound chloride anion is displaced above the central cavity (ESI Figure S1), with the hole diameter at its narrowest being smaller (\(\sim 0.24\) nm) than the ionic diameter of a chloride ion (\(\sim 0.36\) nm),22 suggesting that the anion might not easily pass through to the other side. This very tightly bound anion may also have a very slow rate of dissociation, effectively blocking the hole. The anion selectivity of the channel formed by 3 broadly follows the relative anion binding affinity of the Star of David catenane in MeCN solution.34 The calculated Hille diameter is also similar to the hole diameter in the solid state.9c The low rate of perchlorate and nitrate transport relative to their place in the Hofmeister series is also consistent with anion flow through rigid channels of well-defined size and shape, with a partial size-exclusion mechanism being the cause of the observed selectivity.19,21,22 Nonetheless, we do not rule out a random transport mode not involving the central cavity. If the compounds insert perpendicularly to the membrane the ions may flow around the molecule, as previously observed for certain DNA duplexes,32 although it is not clear how to rationalize such a mechanism with the observed size/shape departure from the Hofmeister series.

CONCLUSIONS

Our results show that a (Fe\(\text{II}\))\(_6\)-coordinated Star of David [2]catenane, 3, based on a double closed-loop hexameric assembly, exhibits high ionophoric activity in transmembrane ion transport experiments, with anion selectivity governed by anion desolvation energies and size-exclusion. The analogous (Fe\(\text{II}\))\(_3\)-coordinated pentafoil knot 2, based on a closed-loop cyclic pentamer of the same building block, also shows ion transport activity but is almost two orders of magnitude less active than the Star of David catenane. Allosteric binding of the Fe\(\text{II}\) ions creates a rigid central anion binding cavity and is crucial for forming ion channels. Remarkably, however, analogous Fe\(\text{II}\)-coordinated Lehn-type cyclic helicates—including the immediate precursor to the Star of David catenane prior to ring-closing metathesis—show no ionophoric activity despite having similar anion binding affinities to the Fe\(\text{II}\)-coordinated triply-interlocked [2]catenane in solution. This dramatic ‘topology effect’ is likely due to conformational restrictions imposed on the chains that link the building blocks in the catenate and knot. Control experiments rule out a carrier mechanism for transport membrane lysis, while single-channel planar bilayer experiments with the (Fe\(\text{II}\))\(_6\)-coordinated Star of David catenane, 3, confirm the formation of ion channels. The level of ion transport activity of the Star of David catenane (EC\(\text{50}\) for KCl transport = 1.3 \(\mu\)M, 17.4 \(\mu\)g mL\(^{-1}\)) is similar to that of some putative antibiotics.33 The discovery of a new class of synthetic ion channels for which both allosteric binding and ‘molecular nanotopology’34 play vital roles, opens up new research directions and the opportunity for creating ion channels that could potentially be tunable or switchable through the use of different metal ions, oxidation states or coordination geometries.35

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Experimental section, synthetic overview, fluorescence measurements, UV-vis measurements, planar bilayer measurements, extended data sets, analysis of previously reported crystal structures. (PDF)

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Notes
The authors declare no competing financial interest.

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