A Molecular Cage-Based [2]Rotaxane That Behaves as a Molecular Muscle

Chun-Ju Chuang, Wan-Sheung Li, Chien-Chen Lai, Yi-Hung Liu, Shie-Ming Peng, Ito Chao, and Sheng-Hsien Chiu

Department of Chemistry, National Taiwan University, No. 1, Sec. 4, Roosevelt Road, Taipei, Taiwan 10617, R.O.C., Institute of Chemistry, Academia Sinica, Nankang, Taiwan, R.O.C., and Institute of Molecular Biology, National Chung Hsing University and Department of Medical Genetics, China Medical University Hospital, Taichung, Taiwan, R.O.C.

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Abstract

We report a molecular cage-based [2]rotaxane that functions as an artificial molecular muscle through the control of the addition and removal of fluoride anions. The percentage change in molecular length of the [2]rotaxane is about 36% between the stretched and contracted states, which is larger than the percentage change (∼27%) in human muscle.

Skeletal muscles are responsible for the motion of many biological systems. On the molecular scale, these muscles are delicately controlled linear machines exhibiting reversible, programmed contraction and stretching movements. To mimic the unique function of biological muscles, several groups are developing artificial linear molecular assemblies that undergo controllable stretching and contraction. The design of these interlocked molecular muscles is elegant: two interlocked components of hermaphroditic rotaxane-like systems move with respect to one another upon the sequential application and removal of a stimulus, thereby changing the distance between the two termini in a controllable manner (Figure 1). Staying with the concept of rotaxane-based systems, we envisioned an alternative type of molecular muscle: one in which the thread-like component can exist in both extended and folded conformations within its encircling container-like macrocycle; muscle-like molecular motion would ensue if different recognition sites on the

1 National Taiwan University.
2 Academia Sinica.
3 Institute of Molecular Biology.


thread-like component interacted with the macrocyclic component under orthogonal conditions. Previously, we reported that the molecular cage 1 forms two types of [2]pseudorotaxane-like complexes, i.e., where complementary guest species are threaded through its two types of macrocyclic cavities.4 Herein, we report a [2]rotaxane, which incorporates this molecular cage, that functions as an artificial molecular muscle that can be controlled through the addition and removal of fluoride ions.

To mimic the function of biological muscle, [2]rotaxane 2-H2·4PF6, which contains two NH2+ and pyridinium units in the thread component linked by flexible n-alkyl chains, was designed. We anticipated that by controlling which of the recognition sites (NH2+ or pyridinium stations) of the dumbbell-structured component interacted with the DB24C8-like openings of the molecular cage component, we would obtain both stretched and contracted conformations of the [2]rotaxane, such that the distance between the termini of the thread-like component would vary substantially, thereby mimicking muscle-like behavior.

We synthesized the thread-like trication 3-H2·3PF6 from ditosylated 1,8-octanediamine5 in three steps (see Supporting Information). An equimolar (2 mM) mixture of the molecular cage 1 and 3-H2·3PF6 in CD3CN at room temperature provided a complicated 1H NMR spectrum that implied the presence of more than one type of complex, with corresponding rates of association and/or dissociation slower than the NMR spectroscopic time scale under these conditions (Figure 2a). Heating this solution at 323 K for 4 h gave a single set of signals for the complex in the 1H NMR spectrum (Figure 2c); the shifts in the positions of the signals of the pyridinium protons and their neighboring are minor compared with the dramatic shifts in the signals of the n-octyl unit’s protons, relative to those in the spectrum of the dumbbell 4-H2·4PF6 (Figure 2d), suggesting that the structure of this complex in solution was most likely the semirotaxane (1·3-H2)·3PF6. After adding 4-(3,5-di-tert-butylphenyl)pyridine 56 (200 mM) to a solution of the semirotaxane (1·3-H2)·3PF6, prepared by heating a CH3CN solution of the molecular cage 1 (50 mM) and the thread-like trication salt 3-H2·3PF6 (60 mM) for 4 h at 323 K, the solution mixture was stirred for 6 days, and the desired [2]rotaxane 2-H2·4PF6 was obtained in 30% yield after ion exchange and column chromatography (Scheme 1).

2D COSY and NOESY experiments helped us to identify most of the signals in the 1H NMR spectrum of the [2]rotaxane 2-H2·4PF6 in CD3CN at 298 K (see Supporting Information). The significant upfield shifts of the signals for the methylene units of the central n-octyl chain relative to those of the n-hexyl protons linking the ammonium and pyridinium stations—in addition to the negligible shifts in the signals of the protons of the pyridinium units compared with those of the dumbbell 4-H2·2PF6—suggested that, under these conditions, the molecular cage interacted noncovalently with the two NH2+ stations. Thus, the “stretched” form of the [2]rotaxane 2-H2·4PF6 predominated in solution under these conditions.

We grew single crystals suitable for X-ray crystallography through liquid diffusion of isopropyl ether into a solution mixture of CH3CN and MeOH (1:1) of 2-H2·4PF6. The solid-state structure7,8 (Figure 3) reveals the expected [2]rotaxane...
We suspect that responsibility for this behavior lies in (i) the two less-acidic complexed NH$_2^+$ centers in the [2]rotaxane and (ii) an insufficient gain in stability, through new ion–dipole and [C–H···O] hydrogen bonding interactions between the pyridinium and DB24C8-like units, to compensate for the loss of [N–H···O] hydrogen bonds.

Because the interactions between secondary dialkylammonium ions and crown ethers are affected by the nature of the counteranions, we turned our attention to controlling the muscle-like behavior of the [2]rotaxane 2-H$_2$4PF$_6$ through anion exchange (Figure 4). After adding 2 equiv of tetrabutylammonium fluoride (TBAF) to a CD$_3$CN solution of 2-H$_2$4PF$_6$, the $^1$H NMR spectrum revealed an equal amount of a new translational isomer in addition to the original one, i.e., translational isomerization of the [2]rotaxane 2-H$_2$4PF$_6$ (i) was slow under these conditions and (ii) required the exchange of all four of its counteranions. The addition of an additional 2 equiv of TBAF to this solution completely removed the signals of the original “stretched” isomer from the spectrum, resulting in the new translational isomer becoming the only species in solution (Figure 5). The significant downfield shift of the pyridinium protons suggests their possible [C–H···O] hydrogen bonding with the oxygen atoms of the DB24C8-like units; the slight broadening of the aromatic signal (H$_{Ar}$) and the complicated signals for the protons of the ethylene glycol units of the molecular cage component are consistent with extrusion of the n-octyl chain through one of the 34-membered-ring openings. Through 2D COSY and NOSY experiments, we identified most of the signals of this new translational isomer in the $^1$H NMR spectrum in CD$_3$CN at 298 K. In this translational isomer, the methylene protons of the n-hexyl chain of the dumbbell-shaped component are strongly shielded (see, for example, the signals of H$_3$ and H$_4$) by the aromatic units of molecular

### Scheme 1. Synthesis of a [2]Rotaxane That Functions As a Molecular Muscle Cage-Based System

![Scheme 1](image1.png)

**Figure 3.** Ball-and-stick representation of the solid state structure of the [2]rotaxane 2-H$_2$4PF$_6$. Atom labels: C, gray; H, pink; O, orange; N, blue.

molecular geometry: the dumbbell-shaped component is threaded through the two 24-membered-ring openings of the molecular cage unit, and the two NH$_2^+$ centers are located within the crown ether-like cavities.

Initially, we attempted to contract the [2]rotaxane 2-H$_2$4PF$_6$ through the addition of bases. We found, however, no conclusive $^1$H NMR spectroscopic evidence for contraction when we added 10 equiv of triethylamine (Et$_3$N), 1,8-diaminonaphthalene (proton sponge), or potassium tert-butoxide to a CD$_3$CN solution of the [2]rotaxane 2-H$_2$4PF$_6$ (see Supporting Information).

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cage 1, whereas those of the n-octyl group (see, for example, H2 and H4) are dramatically less shielded (Figure 5c).

In addition, the presence of a cross peak between the signals of the methylene protons adjacent to the pyridinium group and the aromatic protons of the molecular cage component in the 2D NOSY spectrum supports the “contracted” nature of this translational isomer.

We believe that the main driving force leading to this unique “contracted” form is the fluoride anions’ disruption of [N\(^+\)–H···O] hydrogen bonding between the NH\(^+\) centers and the DB24C8-like units of molecular cage 1 being relatively stronger than that of the [C–H···O] hydrogen bonding of the pyridinium ions. Subsequent addition of 2 equiv of CaBF\(_4\) to remove the fluoride anions from solution through precipitation of CaF\(_2\), provided a spectrum (Figure 5e) similar to that of the original [2]rotaxane 2-H\(_2\)4PF\(_6\) in the same solvent (Figure 5a), i.e., the stretched form of 2-H\(_2\)4\(^{4+}\) was regenerated. Thus, the contracted and elongated forms of the [2]rotaxane 2-H\(_2\)4\(^{4+}\) can be generated reversibly in CD\(_3\)CN through counteranion exchange (F\(^-\) vs PF\(_6\)\(^-\)/BF\(_4\)\(^-\)), providing a simple system for mimicking the function of skeletal muscles. The addition of Cl\(^-\), Br\(^-\), or I\(^-\) ions (as tetrabutylammonium salts) to a CD\(_3\)CN solution of the [2]rotaxane 2-H\(_2\)4PF\(_6\) failed to contract the molecular muscle; thus, the operation of the molecular muscle is selectively controlled by the formation of the tight binding NH\(^+\)/F\(^-\) ion pair.

To estimate the percentage change in the length of the thread-like component between the contracted and stretched states of the [2]rotaxane 2-H\(_2\)4\(^{4+}\), we performed molecular dynamics (MD) simulations in continuums of CHCl\(_3\) and water and in the gas phase (see Supporting Information). When starting the MD simulation from the solid state structure of the [2]rotaxane 2-H\(_2\)4\(^{4+}\), in which both pyridinium units are stacked with the aromatic component of the molecular cage, the molecular conformation of 2-H\(_2\)4\(^{4+}\) changed into the stretched form during the equilibration step under all solvent models. This result suggests that the stacking structure was not the predominant conformation of the [2]rotaxane 2-H\(_2\)4\(^{4+}\) in solution. Using the aromatic carbon atoms located between the two tert-butyl groups as reference points, our MD simulations revealed that the average lengths of the thread in the stretched and contracted states were 39.7 and 25.3 Å, respectively. Thus, the percentage change in molecular length is about 36% between the two states, which is larger than the percentage change (~27%) in human muscle.2b

Figure 5. 'H NMR spectra (400 MHz, CD\(_3\)CN, 298 K) of (a) the [2]rotaxane 2-H\(_2\)4PF\(_6\); (b) the mixture obtained after adding TBAF (2 equiv) to the solution in (a); (c) the mixture obtained after adding TBAF (2 equiv) to the solution in (b); (d) the mixture obtained after adding CaBF\(_4\) (1 equiv) to the solution in (c); and (e) the mixture obtained after adding CaBF\(_4\) (1 equiv) to the solution in (d).

Figure 6. Lowest-energy structures of 2-H\(_2\)4\(^{4+}\) obtained from MD simulations of the (a) stretched and (b) contracted forms. Atom labels: C, gray; O, red; N, blue.

energy structures of 2-H\(_2\)4\(^{4+}\) in the two different states obtained from simulations in the continuum water model.

We have prepared a molecular cage-based [2]rotaxane that functions as an artificial molecular muscle that is controlled through the addition and removal of fluoride anions. The unique container-like structure of molecular cage 1 compared to common macrocycles provides an alternative possibility in the design of molecular muscles. We are at present investigating the further use of container-like host molecules within other functional molecules.

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Supporting Information Available: Synthetic procedures and characterization data for the [2]rotaxane 2-H\(_2\)4PF\(_6\). This material is available free of charge via the Internet at http://pubs.acs.org.

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