Acid–Base Actuation of [c2]Daisy Chains
Lei Fang,‡ Mohamad Hmadeh,‡ Jishan Wu,§,∥ Mark A. Olson,‡ Jason M. Spruell,‡ Ali Trabolsi,† Ying-Wei Yang,§,∥ Mourad Elhabiri,‡ Anne-Marie Albrecht-Gary,*∥ and J. Fraser Stoddart*†

Department of Chemistry, Northwestern University, 2145 Sheridan Road, Evanston, Illinois 60208-3113, Laboratoire de Physico-Chimie Bioinorganique, UDS-CNRS (UMR 7177), Institut de Chimie de Strasbourg, Université de Strasbourg, ECPM, 25 Rue Becquerel, 67200 Strasbourg, France, and Department of Chemistry and Biochemistry, University of California, Los Angeles, 405 Hilgard Avenue, Los Angeles, California 90095-1569

Received February 3, 2009; E-mail: stoddart@northwestern.edu; amalibre@chimie.u-strasbg.fr

Abstract: A versatile synthetic strategy, which was conceived and employed to prepare doubly threaded, bistable [c2]daisy chain compounds, is described. Propargyl and 1-pentenyl groups have been grafted onto the stoppers of [c2]daisy chain molecules obtained using a template-directed synthetic protocol. Such [c2]daisy chain molecules undergo reversible extension and contraction upon treatment with acid and base, respectively. The dialkyne-functionalized [c2]daisy chain (AA) was subjected to an [AA+BB] type polymerization with an appropriate diazide (BB) to afford a linear, mechanically interlocked, main-chain polymer. The macromolecular properties of this polymer were characterized by chronocoulometry, size exclusion chromatography, and static light-scattering analysis. The acid–base switching properties of both the monomers and the polymer have been studied in solution, using 1H NMR spectroscopy, UV/vis absorption spectroscopy, and cyclic voltammetry. The experimental results demonstrate that the functionalized [c2]daisy chains, along with their polymeric derivatives, undergo quantitative, efficient, and fully reversible switching processes in solution. Kinetics measurements demonstrate that the acid/base-promoted extension/contraction movements of the polymeric [c2]daisy chain are actually faster than those of its monomeric counterpart. These observations open the door to correlated molecular motions and to changes in material properties.

Introduction
The mechanical bond1 is the distinguishing characteristic of mechanically interlocked molecules (MIMs) such as catenanes and rotaxanes.2 MIMs are true molecular entities as a result of their mechanically interlocked components being intrinsically linked to one another — resulting in a mechanical bond which prevents dissociation of the components unless one or more covalent bonds are cleaved. The dynamic nature of mechanical bonds allows for the components to undergo relative internal movements, i.e., translation and circumrotation, within the MIMs. Many examples of nanomachines/nanoswitches3 operating at the molecular level, such as molecular actuators,4 molecular electronic components,5 molecular nanovalves,6 etc., have been developed, based on either bistable catenanes7 or bistable rotaxanes8 as the MIMs.

Artificial molecular actuators, which can convert chemical,9 electrochemical,10 or photochemical11 energy into mechanical


motion, have the potential for spawning nanoelectromechanical systems (NEMS). Current investigations have focused on the development of the “bottom-up” approach, which is centered on the design and manipulation of macromolecules and molecular assemblies, both biological and artificial, with the aim of transferring molecular phenomena into nano/micromotions. To achieve macroscopic-matter-property changes from the molecular arena, much effort has been devoted to the design and synthesis of mechanically interlocked macromolecules, and such polymeric, oligomeric, or dendrimeric rotaxanes and catenanes. The incorporation of bistability into these macromolecules, resulting in mechanically interlocked switchable polymeric scaffolds (MISPS), is a valuable example of the “bottom-up” approach taken by chemists to transform motions at the molecular level into changes in properties in the nano-, micro-, and macroscale regimes. However, to date, formidable challenges still remain in the production of MISPS on account of their structural complexity and sensitivity to many reaction conditions, in addition to the insufficient, multistep synthetic sequences required for their preparation.

Acid–base switchable bistable systems have been studied extensively over the past decade. Typically a crown ether, e.g., dibenzoc[4]crown-8 ring (DB24C8), is employed as the encreding macrocycle, while a dialkylammonium center (−[CH2−N(CH3)2]−) and an N,N′-dialkylated-4,4′-bipyridinium unit (BYPm2+) are used as the recognition sites for the macrocycle. Since the association constants for the 1:1 complex formed between DB24C8 and (−[CH2−N(CH3)2]−)BYPm2+ in MeCN are 420/82 M−1 respectively, it is hardly surprising that the DB24C8 ring enircles the dialkylammonium center predominantly. Upon deprotonation, however, of the (−[CH2−N(CH3)2]−) center, the DB24C8 ring moves from this site to the BYPm2+ unit, on account of the drastically reduced binding affinity between the DB24C8 and the resulting neutral


-CH₂NH₂ function. Apparently, reprotonation of the neutral amino group back to a dialkylammonium center results in the return of the DB24C8 ring to this (-CH₂NH₂⁺CH₂-) center. Such acid–base switching has proven¹⁸ to be fast and reversible. Engineering the acid–base switchable² rotaxane into a [c2]daisy chain topology¹⁹ allows muscle-like contractions and expansions to occur as a result of the acid–base switching of the molecules. Moreover, the facile synthesis and symmetric nature of [c2]daisy chains allow further functionalization and elaboration of the molecule, especially into oligomeric or polymeric scaffolds, without confronting major synthetic challenges. Recently, a polymeric [c2]daisy chain was reported²⁰ and an acid–base switchable [c2]daisy chain was described.²¹

Herein, we report on (i) the preparation of a series of [c2]daisy chain compounds functionalized with various different end groups, (ii) the synthesis and characterization of a mechanically interlocked poly[c2]daisy chain, and (iii) the investigation of the acid–base switching behavior of both the monomers and the polymer.

### Results and Discussions

#### Design and Synthetic Strategy

A [c2]daisy chain topology has been identified¹⁹–²² wherein (Figure 1) two mechanically interlocked filaments glide along one another through terminal rings and in which the other end of each filament is attached to bulky stoppers to prevent dethreading of the rings. The C₂ axis and inversion center (Cₐ) along with the plane of symmetry (σ) present in this structure, make it highly symmetric. Thus, a synthetic approach which allows for the simultaneous elaboration of both filaments was adopted. In this case, to generate the stable [c2]daisy chain structure, a dimeric self-complex of the filament is attached with two end groups that contain bulky stoppers, e.g., 2,6-disopropyl- and 2,6-di-tert-butyl-phenyl groups.

To achieve molecular bistability, both recognition sites for DB24C8, namely, -CH₂NH₂⁺CH₂- and BPYM²², must be incorporated into the filament. By taking advantage of the strong binding affinity between DB24C8 and -CH₂NH₂⁺CH₂-, which is primarily on account of the convergent [N⁺−H⋯O] interaction, we are able to design a dimeric self-complex precursor in which the DB24C8 and -CH₂NH₂⁺CH₂- are covalently linked together. The -CH₂NH₂⁺CH₂- site on each filament can thread inside the DB24C8 ring on the other filament in aprotic solvents.

#### Figure 1

A graphic representation of a doubly switchable [c2]daisy chain with C₃ᵥ symmetry. The red rings will have the dibenzo[24]crown-8 constitution. The green stoppers are substituted disopropyl[phenyl] groups. The recognition sites on the rods attached to the red rings are the more preferred CH₂NH₂⁺CH₂- recognition sites (blue spheres) and the less preferred bipyrindinium (BPYM²²) recognition sites (elongated solid blue cylinders).

An Sₘ₂ alkylation of the brominated self-complex at both ends with bulky groups which contain the 4-pyridylpyridinium moiety completes the construction of the stoppered [c2]daisy chain molecules, while also generating the second BYPM²⁺ recognition site simultaneously. To simplify the NMR spectral interpretation of the resulting products, a DB24C8 ring containing a tetradeuterated catechol ring was incorporated into the model compound.

By functionalizing the stoppers, one can synthesize bistable [c2]daisy chain molecules with various terminal functional groups. Such an AA-type bifunctional compound can undergo polymerization with a compatible BB-type bifunctional compound to afford linear [c2]daisy chain polymers. The highly efficient Cu-catalyzed, Huisgen 1,3-dipolar cycloaddition¹⁴,²³ was chosen for the polymerization step on account of its mild nature and high conversion. The simple reaction conditions required for the polymerization, namely CuI addition with low heating, were mild enough to ensure that the [c2]daisy chain monomer units would not decompose or undergo side reactions. High conversions (typically >99%) are pivotal to stepwise growth polymerizations since the degree of polymerization is heavily dependent on the efficiency of the reaction.

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Based on the strategy which has been outlined, a series of monomeric and polymeric daisy chain derivatives (Figure 2) have been identified. The \([c2]\)daisy chain \(1a\cdot6PF_6\) (Figure 2a) is a model compound bearing a tetradeuterated catechol moiety on the DB24C8 rings. The derivatives \(1b\cdot6PF_6\) (Figure 2b) and \(1c\cdot6PF_6\) (Figure 2c) are terminally functionalized on their stoppers by 1-pentenyl and propargyl groups, respectively. \(2\cdot6nPF_6\) (Figure 2d) represents the linear polymeric \([c2]\)daisy chain derived from \(1c\cdot6PF_6\). With these compounds in hand, a muscle-like contraction-extension can be triggered by addition of bases and acids. In this report, the non-nucleophilic bases, 1,4-diaza-bicyclo[2,2,2]octane (DABCO) and \(N\)-tert-butyl-hexamethylphosphortriamide (phosphazene base referred to as P1-t-Bu), were chosen to initiate the base-triggered molecular contraction by fully deprotonating the \(-\text{CH}_2\text{NH}_2^+\text{CH}_2-\) centers, while leaving the BYPM\(^2+\) units intact. On the other hand, trifluoroacetic acid was used for reprotonation of \(-\text{CH}_2\text{NH}-\text{CH}_2-\) functions to promote the reverse extension movement.

**Synthesis.** The synthetic routes employed for the fabrication of both the tetraeterated and nondeuterated dimeric superstructures (\(5a\cdot-b\cdotPF_6\)), the \([c2]\)daisy chains \(1a\cdot-c\cdot6PF_6\), and the polymeric \([c2]\)daisy chain \(2\cdot6nPF_6\) are outlined in Schemes 1–3. The preparation of \((5a\cdot-b\cdotPF_6)_2\) began with reductive amination of the formylated DB24C8 derivatives \(3a\cdot-b\) with methyl 4-(aminomethyl)benzoate. Lithium aluminum hydride reduced the imine bonds and the ester groups to secondary amine and benzyl alcohol functions, respectively, in one pot. Bromination of the benzyl alcohol using phosphorus tribromide, followed by protonation of the amine using hydrobromic acid and counterion exchange using \(\text{NH}_4PF_6\), gave \(5a\cdot-b\cdotPF_6\), which forms a dimeric self-complex \((5a\cdot-b\cdotPF_6)_2\) in an aprotic solvent such as MeCN. Formation of \((5a\cdot-b\cdotPF_6)_2\) was confirmed (see Supporting Information) by the \([2M-2PF_6]^{2+}\) peak found in an electrospray ionization mass spectrum (ESI-MS) of \(5b\cdotPF_6\) using MeCN as the solvent.\(^{24}\)

To complete the construction (Scheme 2) of \([c2]\)daisy chain molecules, the 4-pyridylpyridinium-containing bulky stoppers \(7a\cdot-c\cdotPF_6\) were prepared by reacting 4,4’-bipyridine with the benzyl bromide derivatives \(6a\cdot-c\). A mixed solvent system (CHCl\(_3\)/MeCN, \(v/v\) = 1:1) was chosen to optimize the reaction yield by balancing both the solubility and polarity. By treating \(7a\cdotPF_6\) with \((5a\cdotPF_6)_2\) in this solvent mixture at 55 °C, the doubly threaded \([c2]\)daisy chain model compound \(1a\cdot6PF_6\) was formed and isolated in >30% yield. Using the same method,
1b−c6PF6 were synthesized successfully by reacting 7b−cPF6 with (5b−cPF6)2.

The synthesis of the linear main-chain poly[c2]daisy chain 2•6nPF6 is outlined in Scheme 3. As a terminally bifunctionalized compound, 1c•6PF6 was subjected to a step-growth AA+BB type polymerization using the Cu-catalyzed Huisgen 1,3-dipolar cycloaddition14i with 1 equiv of the diazide 8 in the presence of a stoichiometric amount of CuI in DMF-d7, to afford the linear main chain poly[c2]daisy chain 2•6nPF6. 1H NMR spectroscopy was employed to monitor the reaction process by tracking the disappearance of the alkyne signal and the appearance of the newly formed triazole proton resonance.

Structural Characterization. The 1H NMR spectrum of 1a•6PF6 in CD3CN reveals well-resolved peaks, which were assigned with the help of COSY-NMR 2D correlations. The spectra indicate that the DB24C8 rings exhibit an overwhelming selectivity to encircle the −CH2NH2+CH2− recognition sites at room temperature. Two sets of signals for the catechol rings were identified in the region 6.1−7.0 ppm, indicating the presence of two diastereoisomers. On account of the unsymmetrical substitution of the DB24C8 ring, dimerization gives (Figure 3) two C2-symmetrical (chiral) enantiomers and a C1-symmetrical (meso) diastereoisomer.

Compounds 1b−c•6PF6 exhibit 1H NMR resonances similar to 1a•6PF6, except for the addition of signals arising from the catechol protons and the terminal alkene and alkyne residues (see Supporting Information). High resolution ESI mass spectra of 1a−c•6PF6 all afforded multiply charged peaks, i.e., [M−2PF6]2+, as a result of the loss of multiple PF6− counterions during the ionization processes.

The 1H NMR spectrum of the polymer 2•6nPF6 in CD3CN shows well-resolved signals reminiscent of those present in the monomeric precursor 1c•6PF6. With each repeating unit connected to one another by mechanical bonds, 2•6nPF6 is a “true” macromolecule and is expected to have properties characteristic of a polymer. Chronocoulometry measurement (Figure 4a) carried out on 2•6nPF6 gave a diffusion coefficient of (2.52 ± 0.06) × 10−10 m2/s, a value which is much smaller than that [(7.95 ± 0.10) × 10−10 m2/s] of the monomer 1c•6PF6, suggesting that 2•6nPF6 has a much longer hydrodynamic radius than the monomer. Size exclusion chromatography/multiangle light scattering (SEC-MALS) analysis of 2•6nPF6, which was performed using a solution of NH4PF6 in DMF (0.2 mol/L) as the eluent,25 showed (Figure 4b) a major peak in the chromatograph, with a small amount of unreacted monomer. The calculated number average molecular weight (Mn) is 32.9 ± 2.5 kDa with a polydispersity index (PDI) of 1.85 from Zimm plot analysis. The Mn indicates that, on average, each polymer chain is composed of ~11 repeating units. As a control experiment, SEC-MALS analysis of the monomer 1c•6PF6 resulted in a sharp peak with molecular weight of 4 kDa, a value which is of the same order of magnitude (2.8 kDa) as the actual molecular weight.26 Moreover, differential scanning calorimetry (DSC) analysis of 2•6nPF6 showed (see Supporting Information)

Figure 3. Molecular formulas and graphical representation of isomeric superstructures of (5b•PF6)4: (a and c) a pair of enantiomers, (b) a meso molecule with a C1 symmetry; (b and a/c) diastereoisomers.

(25) SEC-MALS analysis of 2•6nPF6 using pure DMF as the eluent gave a molecular weight of 1466 kDa. Such a large molecular weight can be ascribed to aggregation in a non-electrolyte solution. It is well known that the aggregation of polyelectrolytes can be prevented by introducing ions in concentration into solutions.
protons ($H_d$ and $H_e$) on the phenylene spacers are shifted to higher field because of the mutually $\pi$ overlapping nature of the paraphenylen ring systems in the contracted geometry, (2) the separate peaks for $H_d$, $H_e$, and $H_c$ in the diastereoisomers are less anisochronous and give only one set of accidentally equivalent signals because of the lack of $\pi-\pi$ interactions of the monosubstituted catechol ring, and (3) $H_f$ is shifted to higher field while $H_f$ retains its original chemical shift, thus the encircling position of DB24C8 rings can be assigned primarily to that half-end of BYPM2$^+$ unit which is close to $H_f$. It is also interesting that the signal for the two protons of $H_f$ changed from an accidentally equivalent singlet to an AB-type system as a result of the increased anisochronous environment after the crown ether ring shuttles onto the BYPM$^+$ unit. This assignment is supported by the signal separation of the BYPM$^+$ proton ($\alpha$'s and $\beta$'s) as a result of amplification of the unsymmetrical BYPM$^+$ units when the DB24C8 rings encircle only half of them. Although more complicated spectra were recorded when conducting the acid–base switching cycle on 2$^6$PF$_6$, the observed $^1$H NMR signal movements, reminiscent (Figure 5d–f) of those of 1a$^6$PF$_6^-$, can be fully rationalized. This $^1$H NMR response demonstrates that the switchability of the polymer 2$^6$PF$_6$ under acid/base control is of the same magnitude as that of the monomer 1a$^6$PF$_6$.

Acid–Base Switching Monitored by UV/vis Absorption Spectroscopy. UV/vis absorption spectroscopic investigations of both the polymer 2$^6$PF$_6$ (Figure 6) and the monomer 1c$^6$PF$_6$ (see Supporting Information) in MeCN show similar reversible acid–base switching behavior. Upon the addition (Figure 6a) of DABCO, the absorption band of 2$^6$PF$_6$, centered at 260 nm, which originates from the BYPM$^+$ units, experiences a significant hypochromic shift, while a new absorption band appears at 390 nm as a result of the charge-transfer interactions between the $\pi$-electron deficient BYPM$^+$ units and the $\pi$-electron rich DB24C8 rings. Upon adding (Figure 6b) CF$_3$CO$_2$H to the basified solution, the same spectroscopic signatures undergo a reversal to lead to the initial state. Isosbestic points at 239, 328, and 343 nm in both titration experiments show unambiguously the contribution of only two species during these titrations. The acid–base titrations demonstrate (Figure 6c) that quantitative and reversible switching processes occur over 2 equiv of either acid or base. Switching cycles of the polymer were also investigated by absorption spectrophotometry at 390 nm (Figure 6d). The “on–off” behavior of 2$^6$PF$_6$, triggered by acid/base, remains clearly evident after 10 cycles.

Acid–Base Switching of the Polymer Monitored by Cyclic Voltammetry. It is well documented$^{18}$ that the switching of similar bistable systems can be monitored by cyclic voltammetry (CV) measurements which address the redox active BYPM$^+$ units. The bistability of polymer 2$^6$PF$_6$ was also investigated using this method. The CV scans of 2$^6$PF$_6$ display (Figure 7) two reduction peaks of the BYPM$^+$ units at $-370$ and $-797$ mV (vs Ag/AgCl), respectively. On addition of 2 equiv of P$_1$-$t$-Bu, deprotonation shifts the first reduction process of the BYPM$^+$ units strongly toward more negative potentials ($-477$ mV). Observation of this displacement, together with the $^1$H NMR and UV/vis spectroscopic results, indicate$^{27}$ that

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(26) This disparity in molecular weight from SEC-MALS is primarily on account of the small size of the monomer since low molecular weight species result in weaker scattered laser light and a low signal/noise ratio. Moreover, because the shape of the monomer is neither a random coil nor a perfect rigid rod, the Zimm plot analysis for static light scattering cannot fit the data perfectly to either of the polymer light-scattering function models.


(28) The separation of the second reduction peak after addition of CF$_3$CO$_2$H can be attributed to the presence of different counterions (PF$_6^-$ and CF$_3$CO$_2$-) associated with the BYPM$^+$ unit.

(29) For details of the kinetics mechanism, see Supporting Information.
the BYPM$^2{^+}$ units are engaged in donor–acceptor interactions within the polymer chain, a fact that is consistent with the shuttling of the DB24C8 rings from the $-$CH$_2$NH$_3^+$CH$_2$–centers to the BYPM$^2{^+}$ sites upon deprotonation of the former. Reprotonation of 2•6nPF$_6$ by CF$_3$CO$_2$H leads to the recovery of the first reduction peak$^{39}$ of the BYPM$^2{^+}$ unit at −350 mV.

Dynamics of the [c2]Daisy Chain Systems. The kinetics of the acid-induced extension and the base-promoted contraction processes of the [c2]daisy chain molecules in MeCN were studied using stopped-flow spectrophotometry techniques. Kinetic measurements were therefore carried out under pseudo-first-order conditions; the acid and base were added in large

![Figure 5](image5.png)

**Figure 5.** $^1$H NMR spectra of 1a•6PF$_6$ (400 MHz) and 2•6nPF$_6$ (500 MHz) in CD$_3$CN: (a) The original spectrum of 1a•6PF$_6$ with peak assignments, (b) 1a•6PF$_6$ after addition of 2 equiv of P$_t$-Bu, and (c) 1a•6PF$_6$ after further addition of 2.5 equiv of CF$_3$CO$_2$D; (d) the original spectrum of 2•6nPF$_6$ with peak assignments, (b) 2•6nPF$_6$ after addition of 2 equiv of P$_t$-Bu, and (c) 2•6nPF$_6$ after further addition of 2.5 equiv of CF$_3$CO$_2$D.

![Figure 6](image6.png)

**Figure 6.** UV–vis absorption spectrophotometric variations recorded in MeCN upon (a) addition of DABCO to a solution of 2•6nPF$_6$ and (b) further addition of CF$_3$CO$_2$H to a solution of deprotonated 2•6nPF$_6$. 2•6nPF$_6$ = 4.4 × 10$^{-5}$ mol/L based on the repeating units; $T$ = 25.0(2) °C; $l$ = 1 cm. (c) Variation in the absorbance of the charge transfer band ($\lambda$ = 390 nm) with the gradual addition of DABCO or CF$_3$CO$_2$H to 2•6nPF$_6$ in MeCN. (d) The absorbance at 390 nm on conducting 10 acid–base switching cycles (normalized by dilution factor).
excess, e.g., more than 10 equiv. A single-exponential signal versus time (see Figure 8a for 26nPF6; see Supporting Information for 1c6PF6) was recorded while monitoring the CT absorption band at 390 nm.29

For the two [2]daisy chain compounds (1c6PF6 and 26nPF6) considered in this work, the pseudo-first-order rate constants (kobs) were determined. For the polymer 26nPF6, the variations of the corresponding kobs with [DABCO]tot and with [CF3CO2H]tot are given in Figure 8b–c (for monomer 1c6PF6, see Supporting Information).29

We propose the acid/base switching mechanism in Figure 9 for the [2]daisy chain monomer 1c6+ and the polymer 26n+, respectively.29 Deprotonations of the (−CH2NH2CH2−) sites in 1c6+ and 26n+ as well as protonations of the (−CH2NHCH2−) centers within 1c6+ and 26n+, are acid–base equilibria which are reached within the mixing time (~3 ms) of the stopped-flow technique.18c For the deprotonation of the (−CH2NH2CH2−) units in 1c6+ and 26n+ systems, no spectrophotometric variation at 390 nm was observed during the mixing time.

Therefore, we suggest that the arrangement of the noncontracted 1c4n+ and 24n+ kinetic intermediates is comparable to that of 1c6+ and 26n+, most likely because of the π−π stacking interactions of the substituted catechol rings of the DB24C8 macrocycles.19c,21 Strikingly, protonation of the (−CH2NH−CH2−) units in 1c6+ and 24n+ leads to kinetic intermediates 1c6++ and 24n++, whose spectral properties are markedly different from those of the parent 1c4+ and 24n+ species. A significant absorption decrease at 390 nm (CT transitions) during the ~3 ms dead time suggests that significant electrostatic interactions between the two positively charged (−CH2NH4+CH2−) centers in 1c6++ and 24n++ intermediates destabilize the CT complexes between the dibenzo[24]crown-8 encircling macrocycles and the dialkylated viologen units. As a consequence, the shuttling process was found to be faster in this direction (Figure 8c).

The various acid–base reactions are therefore considered as fast pre-equilibria and afford kinetic intermediates with different protonation states. The rate-limiting steps therefore correspond to either the contraction of the kinetic intermediates 1c4n+ or 24n+ or the extension of 1c4++ or 24n++. Nonlinear least-squares fittings of the variations of kobs versus [DABCO]tot and [CF3CO2H]tot (Figure 8b–c and Supporting Information) allowed us to determine (Table 1) the monomolecular rate constants k1, k−1, k−2, and k−2 (s−1) relative to the contraction and extension processes of 1c6+ and of 26n+, respectively (with kobs = (k1/k2)[DABCO]tot/(1 + β3[DABCO]tot) + k−1 and kobs = (k3/k2)[CF3CO2H]tot/(1 + β3[CF3CO2H]tot) + k−2).30

Both 1c6+ and 26n+ exerted faster CF3CO2H-induced extension than DABCO-induced contraction. Indeed, k2 values [15.5(6) s−1 for ~1c6+ − 1c6+ and 24.7(8) s−1 for 24n+ − 26n+] are approximately three times higher than the corresponding k1 values [5.7(6) s−1 for ~1c6+ − 1c6+ and 7.2(5) s−1 for 24n+ − 26n+]. These data agree well with the kinetic parameters previously reported for a related acid–base-switchable [2]rotaxane.18c In both cases, backward rate constants k−1 or k−2 were also measured, suggesting possible redistribution processes. The association constants between DB24C8 and dibenzylammonium dications/4,4′-bipyridinium dications are 420 and 82 M−1 in MeCN, respectively.18a,d,18e The larger k/k−2 values (~4.0 for 1c6+ and ~15.4 for 26n+) compared to the k/...
k−1 values (∼2.0 for 1c+i+ and 24n+i+) are indicative of a better selectivity of DB24C8 rings for (−CH2NH2+CH2−) centers over BYPM2+ units than that on BYPM2+ units over neutral (−CH2NHCH2−) units. Interestingly, we also observed that the shuttling processes are faster for the polymeric derivative 26nPF6 than for the monomer 1c+6PF6. Speeding up of the reactions 24n+i+ f 24n+i+ or 26n+i+ f 26n+i+ is indeed measured, a trend which suggests that the shuttling of the other subunits becomes faster after the translocation of the first subunit in the polymer. A similar cooperative kinetics effect, which depends on π-electron donor−acceptor interactions for switching, was observed17 on a bistable side-chain polycatenane. This polymer-promoted switching rate enhancement is under further investigation.

Conclusions

The syntheses and characterization of a series of bistable [c2]daisy chain derivatives and a linear polymeric analogue have been reported. The versatile synthetic strategy enables amenable structural modifications of these acid−base switchable, doubly threaded architectures. The polymeric properties, namely, a low diffusion coefficient and a large SEC molecular weight, of the poly[c2]daisy chain demonstrate that the mechanical bonds are equivalent to covalent bonds in terms of their size effects. Extensive investigations of the pH-controlled switching mechanism revealed that both the monomer units and the polymeric ensemble display quantitative and fully reversible switching behavior in solution. Faster switching rates for both contraction and extension processes were observed on the polymer compared to its monomeric precursor. This robust polymeric switchable material sets the stage for the development of smart materials and functional nanomachinery.

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Supporting Information Available: Experimental procedures. Mass spectra and NMR spectra of key compounds. Additional UV/vis absorption spectra and kinetics measurement data. Complete ref 27. These materials are available free of charge via the Internet at http://pubs.acs.org.

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Table 1. Kinetic Parameters* for the Base-Induced Contraction (k1 and k−1) and Acid-Promoted Extension (k2 and k−2) Processes of the [c2]Daisy Chain Monomer 1c+6PF6 and Polymer 26nPF6

<table>
<thead>
<tr>
<th>Process</th>
<th>Reaction</th>
<th>k1 (s−1)</th>
<th>k−1 (s−1)</th>
<th>k2 (s−1)</th>
<th>k−2 (s−1)</th>
</tr>
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<tbody>
<tr>
<td>base-triggered contraction</td>
<td>1c+6− → 1c+i+</td>
<td>4.9 ± 0.2</td>
<td>5.7 ± 0.6</td>
<td>3.0 ± 0.5</td>
<td></td>
</tr>
<tr>
<td>acid-triggered extension</td>
<td>1c+i+ → 1c+6+</td>
<td>4.5 ± 0.3</td>
<td>7.2 ± 0.5</td>
<td>3.4 ± 0.3</td>
<td></td>
</tr>
</tbody>
</table>

*Solvent: MeCN, T: 25.0(2) °C. Errors = 3σ with σ = standard deviation.