Influence of *exo*-Adamantyl Groups and *endo*-OH Functions on the Threading of Calix[6]arene Macrocycle

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**ABSTRACT:** Calix[6]arenes bearing adamantyl groups at the *exo*-rim form pseudorotaxanes with dialkylammonium axles paired to the weakly coordinating \([B(Ar^1)_{4}]^-\) anion. The *exo*-adamantyl groups give rise to a more efficient threading with respect to the *exo*-tert-butyl ones, leading to apparent association constants more than one order of magnitude higher. This improved stability has been ascribed to the more favorable van der Waals interactions of the *exo*-adamantyls versus *exo*-tert-butyls with the cationic axle. Calix[6]arenes bearing *endo*-OH functions give rise to a less efficient threading with respect to the *endo*-OR ones, in line with what was known from the complexation of alkali metal cations.

**INTRODUCTION**

Mechanomolecules,\(^1^a\) such as rotaxanes and catenanes, have become increasingly popular thanks to their aesthetic appeal and to their applications as molecular machines\(^1^b\) or catalysts.\(^1^c\) They are most frequently obtained by threading a rod-like guest (axle) inside a macrocyclic host molecule (wheel) to give an interpenetrated pseudorotaxane precursor.\(^1^d\)

Beginning with crown ethers,\(^2\) a series of macrocyclic classes has been used over the years as the wheel component, which includes cyclodextrins,\(^3\) cucurbiturils,\(^4\) macrolactams,\(^5\) calixar-

ines,\(^6\) and pillararenes.\(^7\) As concerns the calixarene threading, it has been actively investigated by us and by Arduini and co-workers\(^9\) mainly using dialkylammonium and viologen axles, respectively. In particular, 10 years ago we found that scarcely preorganized calix[6]arene ethers (e.g., 1a,b) can be threaded by dialkylammonium axles only when they are coupled to the weakly coordinating tetraakis[3,5-bis(trifluoromethyl)phenylborate ([B(Ar^3)_{4}]) \(^-\) (Figure 1) “superweak” anion.\(^10\) During our studies, we have also found that the conformational mobility of the calix-wheel is strongly influencing the efficiency of the threading. In fact, the more preorganized hexahexyloxycalix[6]-wheel 1b is threaded more efficiently than the more mobile hexamethoxy-1a analogue, by dialkylammonium axles 2\(^-\)–4.\(^10^c\) In accordance, we have recently\(^1^1\) evidenced that the presence of alkyl substituents at the methylene bridges (e.g., 1c) also increases the threading efficiency as a result of the increased degree of preorganization. Another parameter strongly affecting the calix[6]arene threading is the nature of the substituents present at the *exo*-rim (commonly also called as the upper rim). Thus, the very common *p*-tert-butyl groups (like in 1a or 1b) give rise to more stable pseudorotaxane complexes than their simpler *p*-H-counterparts (1d,e), probably because of more extended favorable van der Waals interactions with the cationic axle.\(^10^a,12\) On the basis of this knowledge, we were intrigued to know the effect of bigger and more encumbering groups at the *exo*-rim, such as the *p*-adamantyl ones of 1f,g on the threading efficiency. In addition, we wonder whether the substitution of a few of the OR groups at the *endo*-rim with the OH ones could also lead to thermodynamically stable pseudorotaxane complexes. Prompted by these questions, we have investigated the threading ability by dialkylammonium axles of some *p*-adamantylcalix[6]arene ethers, including some examples of derivatives bearing free OH functions at the *endo*-rim, and we report here the result of this study.

**RESULT AND DISCUSSION**

**Synthesis and Conformational Properties of the Studied Hosts.** The studied hosts were easily obtained by exploiting the classical procedures reported in the literature for the synthesis of calix[6]arene derivatives.\(^1^3^–^1^5\) In particular, \(p\)-adamantylcalix[6]arene hexamethyl ether 1f was obtained in

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47% yield by methylation of the parent \( p \)-adamantylcalix[6]-arene-hexol with MeI, promoted by NaH. Its characterization was made easy by the sharp appearance of its \( ^1H \) and \( ^13C \) NMR signals because of the high conformational mobility of the macrocycle.

In particular, a single set of singlets was observed (Figure 5a) for the six equivalent \( ArCH_2 \) moieties (e.g., at 3.91 and 2.92 ppm for \( ArCH_2Ar \) and \( OCH_3 \) groups, respectively). The structure of \( 1f \) was also confirmed by X-ray crystallography which also provided useful information about its preferred conformation (vide infra). The 1,2,4,5-tetrahexyloxy-\( p \)-adamantyl-calix[6]arene-diol derivative \( 1g \) was obtained in 33% yield by an extension of the NaH-promoted 1,2,4,5-tetrasubstitution of calix[6]arenes originally reported by Gutsche. The characterization of \( 1g \) was made less easy by the broad appearance of its \( ^1H \) and \( ^13C \) NMR signals at room temperature (Figure 2a), due to a conformational mobility close to the NMR time scale. Therefore, all the relevant information was acquired at a high temperature (373 K) in CDCl\(_3\)CDCl\(_3\) (CDCl\(_3\)).

The 1,2,4,5-tetrahexyloxy-\( p \)-tert-butylcalix[6]arene-diol derivative \( 1h \) was obtained by a protection−deprotection procedure (Scheme 1) starting from the known 1,4-dibenzylxioxy-\( p \)-tert-butylcalix[6]arene-tetra-ol, which was first tetraalkylated with hexyl iodide (compound \( 6 \); 90% yield) and then debenzylated with \( H_2/\text{Pd/C} \) (95% yield).

The characterization of \( 1h \) was very similar to that of \( 1g \) for what concerns the broad appearance of its \( ^1H \) and \( ^13C \) NMR signals at room temperature, due to a conformational mobility close to the NMR time scale. The similarity was also extended to the unusual signals in the negative region of the spectrum because of the self-included hexyl chain. Also, in this case a characterization at low temperatures was performed that confirmed the peculiar pseudo[1]rotaxane partial-cone conformation (Figures S20−S22). This result leads to suppose that this conformational feature could be a characteristic of calix[6]arenes 1,2,4,5-tetrasubstituted with long alkyl chains.

**Figure 1.** Chemical drawing of calix[6]arene wheels 1a−h, ammonium cations \( 2^+−4^+ \) and \( [\text{B(ArF)}_4]^− \) anion.

**Figure 2.** \( ^1H \) NMR spectra (CDCl\(_3\), 600 MHz) of pseudo[1]rotaxane derivative \( 1g \) at 298 (a) and 243 K (b); (c,d) different portions of the HSQC spectrum of \( 1g \) (CDCl\(_3\), 600 MHz, 243 K); (e) portion of the COSY-45 spectrum of \( 1g \) (CDCl\(_3\), 600 MHz, 243 K).

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X-ray Analysis of 1f and 1g. Small colorless single crystals of 1f and 1g suitable for X-ray structure determination were analyzed using synchrotron radiation and cryo-cooling.
The asymmetric unit consists of a 1/2 molecule of 1f outside of the macrocycle. In the solid state, 1f exhibits a centrosymmetric 1,2,3-alternate conformation (Figure 3a).

The conformation of the adamantyl-substituted aryl rings is illustrated in Figure 3, where the molecules are viewed orthogonally (Figure 3c,d) with respect to the mean plane (in blue in Figure 3) of the calix[6]arene, as defined by the six methylene bridges. An absolute angle value greater/smaller than 90° indicates the outward/inward orientation of the adamantyl group, while a negative sign indicates an inverted orientation of the adamantyl group with respect to a given orientation of the macrocycle. In the case of 1f, the mean plane of one of the aryl rings (1) (Figure 3a), is almost perpendicular to the mean plane of the calix[6]arene, with a dihedral angle of 96° (Figure 3c). The adamantyl group thus leans slightly outward from the center of the macrocycle. The mean planes of the other two aryl rings make a large outward dihedral angle (2, 132°) and a slight inward angle (3, 72°), consequently, these adamantyl groups lean outward and inward, respectively. The symmetry of the molecule implies that the other three opposite phenyl groups of the macrocycle with an inverted orientation (1′, 2′, and 3′) have identical angles in the absolute value but with a negative sign. The methoxy groups are inward oriented for 2 (2′) and 3 (3′) and outward oriented for 1 (1′) (Figure 3c). The overall conformation, with two bulky adamantyl groups and four methoxy groups tilted toward the center of the macrocycle, results in a sealed molecular cavity (Figure 3a,c).

In the solid state, 1g exhibits an asymmetric partial-cone conformation, with just one of the phenyl rings with an inverted orientation with respect to the other five (Figure 3b). This phenyl (1) is opposite to a phenyl group (4) bearing a hexyloxy chain self-included into the macrocycle. The partial-cone conformation, combined with the mixed hydroxy/hexyloxy substitution pattern at the lower rim (1,2,4,5-hexyloxy) results in an asymmetric C3 molecular point symmetry. As it crystallized in the P1 space group, the structure is therefore composed of a racemic mixture of inherently chiral 1g molecules.

With regard to the overall conformation of 1g (Figure 3d), for the purposes of the following discussion, the side with the five adamantyl substituents is defined as the upper side. The mean plane of the inverted phenyl ring, with the adamantyl group on the lower side (1), is acutely tilted inward with a dihedral angle of −47° with respect to the mean plane defined by the six methylene bridges. The mean plane of the phenyl ring (4), located directly opposite the inverted ring, makes a very large outward dihedral angle (158°) on the upper side. Consequently, the bulk adamantyl group is tilted far from the center of the macrocycle, while the hexyl chain occupies the cavity of the macroring.

The internal hexyl chain is disordered over two positions with equal occupancy factors. In the first conformer, the two central bonds both assume gauche+ conformations (gauche− for the second conformer); while the other two relevant C−C bonds both assume an anti conformation (in both conformers). The mean planes of the other four phenyl groups 2 (hexyloxy-substituted), 3 (hydroxy-substituted), 5 (hexyloxy-substituted), and 6 (hydroxy-substituted) are all close to orthogonal with respect to the above-defined calix[6]arene mean plane; however, in all cases the adamantyl groups are tilted slightly outward, with angles of 94, 98, 96, and 98°, respectively. The conformation of these four phenyl groups is influenced by the formation of hydrogen bonds between the hydroxy group donors (3, 6) and the adjacent hexyloxy oxygen acceptor (2, 5) with O···O distances of 2.78 and 2.86 Å.

**Influence of exo-Adamantyl Groups on Calixarene Threading.** Initially, the influence of the exo-adamantyl groups was investigated by studying the threading abilities of p-adamantylcalix[6]arene hexamethyl ether 1f by dibenzylammonium axle 2′−[B(Ar)2]− (Figure 4). When this salt was added to a CDCl3 solution of 1f (1:1 ratio) then significant changes appeared in the 1H NMR spectrum (Figure S24) indicative of the formation of pseudorotaxane 2+⊂1f. The first piece of information was the appearance of a well-spaced AX system (at 3.46/4.39 ppm, Δδ = 0.93 ppm) for the ArCH2Ar groups of the calix-wheel 1f indicative of its cone conformation in pseudorotaxane 2+⊂1f.
The threading of the dibenzylammonium axle was clearly evident by the presence of upfield-shifted resonances for the benzylic unit hosted inside the cavity at 4.44, 5.26, and 6.00 ppm (ortho, meta-, and para-BnH, respectively) (Figure 5b); while the other outside the cavity was resonating at typical chemical shift values (7.90, 7.64, and 7.50 ppm, ortho-, meta-, and para-BnH, respectively). The threading equilibrium of $2^+ \subset 1f$ was reached immediately after mixing and slow on the NMR time scale. The determination of the corresponding apparent association constant was carried out by means of a competition experiment$^{10b,c}$ (Figure S48) with the native hexahexyloxy-tert-butylcalix[6]arene $1b$. In particular, a 1:1 mixture of $1f$ and $1b$ (in CDCl$_3$) was mixed with 1 equiv of $2^+ \cdot[B(\text{ArF})_4]^{-}$ and equilibrated for 10 min at 298 K. The $^1$H NMR spectrum of the mixture indicated that the pseudorotaxane $2^+ \subset 1f$ was favored over $2^+ \subset 1b$ in a 1:0.8 ratio (Figure S48). An apparent association constant value of $7.4 \pm 0.2 \times 10^4$ M$^{-1}$ was calculated from these data for the $2^+ \subset 1f$ complex, which is significantly higher than that previously observed for the corresponding tert-butylated pseudorotaxane $2^+ \subset 1a$ ($2.5 \pm 0.2 \times 10^3$ M$^{-1}$).$^{10d}$ To confirm this result, a competition experiment was performed in which $2^+ \subset 1a/1f$ were mixed in an equimolar ratio (5.2 mM) in CDCl$_3$, and the resulting $^1$H NMR spectrum of the mixture indicated that the $2^+ \subset 1f$ pseudorotaxane was favored over $2^+ \subset 1a$ (Figure S58). From this initial result, it is clear that the exo-adamantyl groups positively affect the efficiency of calix[6]arene threading probably due to more extensive favorable van der Waals interactions with the cationic guest. On the other hand, it is also evident that the bigger dimension of the para-adamantyl groups with respect to the p-tert-butyl ones does not hinder kinetically the equilibrium of the formation of the pseudorotaxane, being the equilibration time of $2^+ \subset 1f$ similar to that of $2^+ \subset 1a$.

The evaluation of the threading properties of $1f$ were then extended to the dipentylammonium axle $3^+$ by using similar experimental conditions. Thus, the addition of the $3^+ \cdot[B(\text{ArF})_4]^{-}$ salt to a CDCl$_3$ solution of $1f$ again caused the appearance of upfield-shifted resonances in the negative region of the $^1$H NMR spectrum (Figure 5c), demonstrating the formation of the pseudorotaxane $3^+ \subset 1f$, which was slowly exchanging in the NMR time scale. By means of a competition experiment$^{10a}$ (Figure S49) with $1b$, an apparent association constant of $5.0 \pm 0.2 \times 10^4$ M$^{-1}$ was found for $3^+ \subset 1f$ pseudorotaxane, which was again higher than that of $3^+ \subset 1a$ ($4.4 \pm 0.2 \times 10^3$ M$^{-1}$).$^{10b,c}$

Further insights on the higher stability of pseudorotaxanes $2^+ \subset 1f$ and $3^+ \subset 1f$ were obtained by density functional theory (DFT) calculations at the B3LYP/6-31G(d,p) level of theory using Grimme’s dispersion corrections (D3/124 = 3). As can be seen from the energy-minimized structures (Figures 6...
...and $\beta$ Additional C−Å and an average N−C, E, and F rings are more outward tilted (canting angles$^{18}$ of 51.6°, 42.1°, and 50.°(1), respectively) (Figure 6). As a result of these contributions, the calix[6]arene macrocycle is fixed in a cone conformation with D and E rings (Figure 6) almost orthogonal with respect to the mean plane defined by all bridging methylenes (canting angle 85.4°(1) and 88.8°(1), respectively), while the other A, C, E, and F rings are more outward tilted (canting angles$^{18}$ of 51.6°(1), 42.1°(1), 56.0°(1), and 51.0°(1), respectively) (Figure 6).

To further investigate the energy contribution of non-covalent interactions (NCI)$^{19}$ a second order perturbation theory (SOPT) analysis$^{20}$ of the Fock matrix in the natural bond orbital (NBO) basis was carried out. Interestingly, the SOPT analysis conducted on $2^+ \subset 1f$ pseudo$[2]rotaxane$ (Table S3) indicates that there is an articulate network of hydrogen bonding, C−H−$\pi$, and $\pi−\pi$ interactions. In particular, we evidenced the LP(2) → $\sigma^*$ interaction between O3 and the N248−H277 antibonding orbital (Figure S61) and the LP(2) → $\sigma^*$ interaction between O4 and the N248−H250 antibonding orbital. These interactions give an energetic contribution of 10.36 and 8.34 kcal/mol, respectively, for a total energy of 18.70 kcal/mol (Table S3). Moreover, the interactions of the O6 lone pairs with the C247−H276 antibonding orbital gives an interesting contribution of 2.62 kcal/mol. The overall energy analysis (Table S3) indicates that the total energetic contribution because of secondary NCI is 34.09 kcal/mol (see Supporting Information, page S68 for further detail). Very interestingly, a similar analysis for the corresponding tert-butylated $2^+ \subset 1a$ pseudo$[2]rotaxane$ gives a lower total energy contribution of 26.66 kcal/mol (Figure S65 and Table S5).

As concerns the DFT-optimized structure of $3^+ \subset 1f$ pseudorotaxane (Figure 6), two H-bonding interactions were identified between the NH$_2$ group of $3^+$ and the oxygen atoms of calix-wheel 1f, with a longer average distance of 1.98 Å and an average N−H···O angle of 161.45°. In this case, the SOPT analysis (Table S4) indicated an energy contribution of 30.18 kcal/mol for these interactions. An interesting contribution on the NCI energy is due to the LP(2) → $\sigma^*$ interaction between O6 and the N261−H281 antibonding orbital (11.61 kcal/mol). Also in this case, the calix[6]arene macrocycle is fixed in a cone conformation with D and E rings (Figure 6) almost orthogonal with respect to the mean plane defined by all bridging methylenes (canting angle 85.4°(1) and 88.8°(1), respectively), while the other A, B, C, and E rings are more opened with canting angle values of 47.2°(1), 62.8°(1), 53.4°(1), and 39.3°(1), respectively (Figure 6). From the above two examples, we can conclude that the presence of $p$-adamantyl groups gives rise to pseudorotaxane complexes with ammonium guests containing either linear alkyl chains or tert-butyl ones.

Figure 7. NCI plot by the sign of the second Hessian eigenvalue [gradient isosurfaces (s = 0.6 a.u.) for $2^+ \subset 1f$ (a), $2^+ \subset 1a$ (b) and $3^+ \subset 1f$ (c)]. In the coloring isosurface, blue and green colors represent strong and medium interactions (H-bonding and van der Waals).
Directional Threading with an Unsymmetrical Axle. During our studies,8,10,21 we have realized that the threading of directional calixarene-wheels with directional (or constitutionally asymmetric) alkybenzylammonium axles could give rise to two diastereoisomeric pseudo[2]rotaxanes differing by the moiety (alkyl or benzyl) included inside the cavity. Thus, we have found several examples of directional threading of calix[6]-wheels22 in which the endo-alkyl stereoisomer is preferentially formed over the endo-benzyl one. These general empirical observations have induced us to introduce the so-called “endo-alkyl rule”22 to shortly refer to this preferential formation of the endo-alkyl stereoisomer.

In order to verify if this “endo-alkyl rule” is also valid for p-adamantylcalix[6]arenes, we decided to study the threading of 1f with butylbenzylammonium axle 4+. Thus, the addition of the 4+[B(OrF)]4− salt to a CDCl3 solution of 1f again caused the appearance of upfield-shifted resonances in the negative region of the 1H NMR spectrum, demonstrating the preferential formation of the endo-alkyl-4+⊂ 1f pseudorotaxane (Figure S56). Interestingly, no hints of the endo-benzyl-4+⊂ 1f pseudorotaxane could be detected in the above experiment. In summary, we can conclude that the “endo-alkyl rule” is also valid for p-adamantylcalix[6]arenes.

Influence of endo-OH Functions on Calixarene Threading. As is known, the complexion of metal cations by calixarene hosts is less efficient for those bearing free OH functions with respect to the OR-analogues.64 Prompted by this consideration, we turned our attention to the endo-rim, and the question arises as to whether the presence of OH groups impairs the threading efficiency of the calix[6]-wheel. In fact, no information of this kind is currently available for the OH-bearing calix[6]-wheels.0.2 By moving to the 1,2,4,5-tetrahexyloxy-p-adamantyl-calix[6]arene-diol derivative 1g (Figure 9), we expect a slight improving in the threading efficiency with respect to the exotert-butyl analogue 1h, due to the above evidenced “adamantyl effect”. In fact, an apparent association constant of 8.5 ± 0.2 × 104 M−1 was found for 2+⊂ 1g (Figures 9b and S51), which is higher than that of 2+⊂ 1h (2.5 ± 0.2 × 105 M−1). Similar results were found for the dipentylammonium axle by comparing 3+⊂ 1g (3.4 ± 0.2 × 105 M−1) (Figures 9c and S52) with 3+⊂ 1h (7.1 ± 0.2 × 105 M−1). Interestingly, the threading with the unsymmetrical butylbenzylammonium axle 4+ confirmed the validity of the “endo-alkyl rule” also for endoOH-bearing calix[6]arene wheels 1g (Figure 9d) and 1h. Also in these instances, a higher thermodynamic stability was observed for the amantylated pseudorotaxane 4+⊂ 1g (9.3 ± 0.2 × 105 M−1) (Figures 9d and S53) with respect to the tert-butylated one 4+⊂ 1h (1.2 ± 0.2 × 105 M−1). DFT calculations at the B3LYP/6-31G(d,p) level of theory using Grimme’s dispersion corrections (10p(3/124 = 3)) were performed on pseudorotaxanes 2+⊂ 1g and 3+⊂ 1g and the corresponding energy-minimized structure are reported in Figure 10. For both the two supramolecular adducts, it is possible to observe the typical network of H-bonding interactions between the ammonium group and the oxygen atoms of the calixarene macrocycle (2+⊂ 1g: average N···O distance 1.73 Å, average N−H···O angle 166.45°; 3+⊂ 1g: average N−O distance 1.91 Å, average N−H···O angle 166.15°). Several NCI were identified between the endo-cavity benzyl unit of 2+ (or the endo-cavity alkyl chain of 3+) and the aromatic rings of 1g. All these interactions contribute to fix the calix[6]arene macrocycle in the cone conformation. An interesting observation was found for the two distal unsubstituted phenolic rings A and D of both pseudorotaxanes. These ArOH units are more tilted outward from cavity with canting angles 18° in the range 25.9−29.7° (Figure 10).

Figure 8. 1H NMR spectrum (CDCl3, 600 MHz) at 298 K of an equimolar mixture of 4+ and 1f. Inset: DFT-optimized structure of 4+⊂ 1f pseudorotaxane at the B3LYP/6-31G(d,p)-IOp(3/124 = 3) level of theory.
UV254 and visualized using UV light and nebulization with an μ pseudorotaxanes at the B3LYP/6-31G(d,p)-IOp(3/124) level of theory on a Bruker Avance-600 spectrometer [600 (1H) and 150 MHz (13C)], and Bruker Avance-300 spectrometer [300 (1H) and 75 MHz peak (CHCl3: Chemicals and were used without further puri- plates coated with silica gel (0.25 mm) with addition, we have also demonstrated that macrocycle threading and related interpenetrated architectures. OR ones, in line with what was known from the complexation of alkali metal cations. We do believe that these results can be considered useful reference points for future studies on exo-tert-butyl adamantyls versus exo-tert-butyls with the cationic axle. In addition, we have also demonstrated that endo-OH functions give rise to a less efficient threading with respect to the endo-OR ones, in line with what was known from the complexation of alkali metal cations. We do believe that these results can be considered useful reference points for future studies on macrocycle threading and related interpenetrated architectures.

CONCLUSIONS

In conclusion, we have here reported a study on the influence of the nature of the groups attached to the exo-rim and endo-rim of the calix[6]arene macrocycle on its threading properties with ammonium axles. We have here demonstrated that exo-adamantyl groups give rise to a more efficient threading with respect to exo-tert-butyl ones leading to apparent association constants more than one order of magnitude higher. This higher thermodynamic stability has been ascribed to the more favorable van der Waals interactions of higher. This higher thermodynamic stability has been ascribed.

EXPERIMENTAL SECTION

General Comments. Reactions under anhydrous conditions were conducted under an inert atmosphere (nitrogen) using dry solvents. The commercial reagents were purchased by Aldrich and TCI Chemicals and were used without further purification. The reactions were controlled by thin-layer chromatography with Macherey-Nagel plates coated with silica gel (0.25 mm) with fluorescence indicator UV365 and visualized using UV light and nebulization with an indicator solution of H2SO4-Ce(SO4)2. For reactions that require heating, the heat source used was an oil bath. The reaction temperatures were measured externally using electronic thermometers. The reaction products were purified by Macherey-Nagel silica gel chromatography (60, 70–230 mesh), NMR spectra were recorded on a Bruker Avance-600 spectrometer [600 (1H) and 150 MHz (13C)], Bruker Avance-400 spectrometer [400 (1H) and 100 MHz (13C)], and Bruker Avance-300 spectrometer [300 (1H) and 75 MHz (13C)]. Chemical shifts are reported relative to the residual solvent peak (CHCl3: δ 7.26, CDCl3: δ 77.16). Standard pulse programs, provided by the manufacturer, were used for 2D NMR experiments. Structural assignments were made with additional information from correlation spectroscopy (COSY) and heteronuclear single-quantum correlation spectroscopy (HSQC). Experiments. HR MALDI mass spectra were recorded on a Bruker Solaris FT-ICR mass spectrometer equipped with a 7 T magnet. The samples recorded in MALDI were prepared by mixing 10 μL of the analyte in chloroform (1 mg/mL) with 10 μL of the solution of 2,5-dihydroxybenzoic acid (10 mg/mL in methanol). The mass spectra were calibrated externally, and a linear calibration was applied.

Synthesis of Derivative 1f. To a stirred suspension of p-adamantylcalix[6]arene15 (0.43 g, 0.30 mmol) in dimethylformamide (DMF) (50 mL), sodium hydride (60% in mineral oil, 0.72 g, 18.00 mmol) was added. The mixture was stirred for 15 min at room temperature. Dimethylsulfate (2.72 mL, 28.80 mmol) was added and the reaction mixture was stirred at 90 °C for 2 h. After cooling, the reaction was quenched by the addition of methanol (10 mL). The solvents were evaporated under reduced pressure, and the residue was parted between dichloromethane and 2 M HCl. The organic phase was washed with water, dried with MgSO4 and concentrated to almost dryness. The residue was purified by column chromatography [SiO2, gradient from hexane to hexane/tetrahydrofuran (THF) (25:1)]. Derivative H was obtained as a white solid (0.21 g, 47%). 1H NMR (600 MHz, CDCl3, 298 K): δ 5.74 (s, 12H, ArH), 3.91 (s, 3H, ArCH3), 2.92 (s, 18H, OCH2), 2.02 (br s, 18H, Ad), 1.83–1.62 (m, 72H, Ad). 13C{1H} NMR (150 MHz, CDCl3, 298 K): δ 154.2, 145.9, 133.5, 125.7, 59.9, 43.4, 36.8, 35.7, 31.4, 29.0. HRMS (MALDI) m/z: [M + K]+ ca. 1564.9689; found, 1564.9690.

Synthesis of Derivative 1g. To a stirred suspension of p-adamantylcalix[6]arene15 (0.43 g, 0.30 mmol) in DMF (15 mL), sodium hydride (60% in mineral oil, 0.29 g, 7.2 mmol) was added. The mixture was stirred for 15 min at room temperature. 1-Bromohexane (1.01 mL, 7.20 mmol) was added and the reaction mixture was stirred at 90 °C for 20 h. After cooling, the reaction was quenched by addition of methanol (30 mL). The solid formed was collected, washed with methanol, dried, and dissolved in dichloromethane. The solution was washed with 2 M HCl, water, dried with MgSO4, and the solvent was evaporated under reduced pressure. The residue was purified by column chromatography [SiO2, gradient from hexane to hexane/chloroform (3:2)]. Derivative I9 was obtained as a white solid (0.18 g, 33%). 1H NMR (600 MHz, CDCl3, 298 K): δ 5.74 (br s, 2H, ArOH) 6.98–6.83 (overlapped, 4H, ArCH2Ar), 3.75–3.49 (overlapped, 2OH, ArCH2Ar, −CH2(CH2)2CH3), 1.91–0.69 (overlapped, 134H, −CH(CH2)2CH2CH2CH2CH2CH2Ad). 13C{1H} NMR (150 MHz, CDCl3, 298 K, TCDE, 373 K): δ 150.8, 148.3, 144.2, 140.1, 131.2, 130.8, 125.4, 124.2, 122.4, 41.8, 41.6, 35.3, 34.0, 33.8, 30.0, 29.4, 27.7, 27.6, 27.5, 23.4, 20.5, 12.1. HRMS (MALDI) m/z: [M + Na]+ ca. 1801.2725; found, 1801.2725.

Synthesis of Derivative 6. In a dry round flask, derivative 515 (1.05 g, 1.29 mmol) was dissolved in dry acetone (70 mL). Subsequently, Cs2CO3 (1.05 g, 1.29 mmol) was dissolved in dry acetone (70 mL). The raw was purified through precipitation by methanol. Derivative 6 was obtained as a white solid (1.81 g, 90%). 1H NMR (300 MHz, CDCl3, 373 K): δ 7.27 (bd, 4H, ArH), 7.03 (bd, 4H, ArH), 6.92 (br s, 4H, ArH), 6.89 (br s, 4H, ArH), 6.81 (br s, 4H, ArH), 4.65 (s, 4H, CH2PhCH2), 3.77 (overlapped, 12H, ArCH2Ar), 3.19 (br s, 8H, −CH2(CH2)2CH3), 2.23 (s, 6H, −OBeOH), 1.31–0.97 (overlapped, 86H, −CH2(CH2)2CH2−C(CH3)3), 0.74 (bt, 12H, −CH2(CH2)2CH3), 13C{1H} NMR (75 MHz, CDCl3, 373 K): δ 151.8, 151.1, 143.5, 143.1, 135.3, 133.6, 131.3, 131.2, 127.1, 126.4, 124.2, 72.9, 72.4, 32.1, 30.7, 29.7, 28.0, 24.2, 19.7, 12.1. HRMS (MALDI) m/z: [M + K]+ ca. 1557.0958; found, 1557.0941.

Synthesis of Derivative 1h. In a round flask, derivative 6 (0.70 g, 0.45 mmol) was dissolved in chloroform (50 mL). Subsequently, Pd/C was added. Stirring was continued for 18 h at room temperature under H2. After this time, the reaction was stopped by filtration on Celite. The solvent was evaporated under reduced pressure. Derivative 1h was obtained as a white solid (0.56 g, 95%). 1H NMR (600 MHz, CDCl3, 373 K): δ 6.97–6.62 (overlapped, 14H,

Figure 10. DFT-optimized structure of 2∗ ⊂ 1g and 3∗ ⊂ 1g pseudorotaxanes at the B3LYP/6-31G(d,p)-IOp(3/124) level of theory.
One fully occupied CHCl₃ molecule site shows a three-position disorder and partial occupancy are also observed for the solvent unit. These were refined anisotropically, while all other non-hydrogen atoms were refined anisotropically.

Complete contact information is available at: https://pubs.acs.org/doi/10.1021/acs.joc.0c01769.

Copies of ¹H and ¹³C NMR spectra for all products, HRMS, 2D NMR, ¹H VT NMR spectra, NMR titrations, and 2D NMR spectra of complexes, and computational studies (PDF)

Crystallographic data of 1f (CIF)

Crystallographic data of 1g (CIF)

Author Contributions

The manuscript was written through contributions of all the authors. All the authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.
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