Pathways for Conformational Interconversion of Calix[4]arenes

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Received June 27, 1994

Abstract: Conformational optimization and reaction path calculations are performed on [1]metacyclophane 3a and calix[4]arenes 1a and 2b using the CHARMM force field. For each of these compounds, a comprehensive search for all stable conformers was followed by an exhaustive exploration of the several hundred possible pathways between these conformers. The method employed for finding the reaction paths, Conjugate Peak Refinement, proved to be robust and reliable, allowing the connectivity of the complex potential energy surfaces to be charted. The relative stability of the four characteristic conformers agrees with experimental NMR data, except for the Cone form of 2b. The pathways for Cone inversion in [1]metacyclophane 3a show no preference for a pathway via the 1,2Alt or the 1,3Alt conformers. The conformational entropy corrected energy barriers $\Delta E_{\text{corr}}$ are 3.1 and 3.3 kcal/mol, respectively. For 1a, a stepwise pathway via the 1,2Alt conformer is found to be preferred for the Cone inverted-Cone conversion. The rate-limiting step is the transition from Cone to Paco, with a barrier of activation $\Delta E_{\text{act}} = 14.5$ kcal/mol, comparable to the experimental $\Delta H^\circ = 14.2$ kcal/mol. Conversion from the key Paco intermediate to the other characteristic conformers was investigated in detail in 2b. The $\Delta E_{\text{pot}}$ values for the conversion from the most stable Paco to 1,2Alt and 1,3Alt conformers are 19.6, 20.2, and 18.2 kcal/mol respectively, in qualitative agreement with the relative rates deduced from 2D EXSY NMR. Paths for the transition from inward to outward orientation of the methoxy moieties of 2b are calculated. The corresponding activation barriers for the rotation of a methoxy group in the 6–8 kcal/mol range, consistent with the upper bound obtained from the NMR time scale.

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Introduction

Calix[4]arenes are macrocyclic compounds which consist of four phenol rings connected via methylene bridges located ortho to the hydroxy groups. The rings cannot be coplanar and they adopt an orientation that makes the molecule appear like a cup or a chalice. They are being used increasingly in supramolecular chemistry as building blocks for larger molecules that are designed for the complexation of cations or neutral molecules.2 One interesting property of calix[4]arenes is that they have four characteristic nonplanar conformations, designated as the Cone, the partial Cone (Paco), the 1,2-alternate (1,2Alt), and the 1,3-alternate (1,3Alt) (see Figure 1). Several studies have been published in which the relative stability of the calix[4]arene conformations has been calculated with molecular mechanics,2–4 but the relative stability of the four conformations of calix[4]arenes 2 in CDC 13 found by NMR24a,b was not reproduced exactly. Grootenhuis et al.2 used MACROMODEL (AMBER and MM2 force fields), MM2P, AMBER,10 and QUANTA/CHARMM11 to calculate the relative stability of the four conformations of 2c. The force fields showed pronounced differences, which were mainly caused by differences in the electrostatic energy. Only MM2P and QUANTA/CHARMM calculated the partial cone to have the lowest energy, in accord with the NMR data. Using the MM32 force field, Shinkai et al.3 obtained good agreement for the relative stability of the four conformers of 2a and 2b between the NMR data and the calculated energies, including vibrational contributions estimated from normal modes.

Measurements with NMR spectroscopy have shown that calix[4]arenes with hydroxy moieties at the lower rim (1) adopt the Cone conformation in CDCl 3 solution,13 which is also the structure found in crystals.14 The stability of the Cone conformation is believed to be caused by a circular array of

(11) QUANTA 2.1A: Polygen Corp., Wallingford, MA.

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very strong hydrogen bonds, in which the OH groups at the lower rim act as both donors and acceptors. The calix[4]arenes I are still conformationally flexible, because the two equivalent Cone conformations (with the four oxygen atoms either all up or all down) can interconvert. Several mechanisms for interconversion have been proposed. One is a "broken-chain" mechanism, in which the array of H-bonds is broken and the 1,3Alt form is the intermediate; another is a "continuous-chain" mechanism, in which the hydrogen bonds remain mostly intact and a skewed 1,2Alt-like intermediate is involved. Some authors have suggested that the interconversion of tetrahydroxycalix[4]arenes is a concerted process in which more than one phenol ring rotate simultaneously. In all proposed mechanisms the hydroxyl moieties swing through the annulus, since the upper rim (bearing the R groups in Figure 1) is too large to pass through the annulus.

The calix[4]arene derivatives 2 exist in CDC$_3$ solution at room temperature as a mixture of four characteristic conformations, as determined by H and 2D NMR measurements. In decreasing order, the relative stability for 2b is Paco (most stable) > 1,2Alt > Cone > 1,3Alt. The conformers interconvert rapidly, with the Paco conformer as the key intermediate, as indicated by 2D EXSY NMR spectra showing cross-peaks for Paco/Cone and Paco/I,3Alt conformations. Each of the other three conformations can be formed from the Paco conformation by the rotation of only one anisole ring. The methoxy moiety passes through the annulus during the ring-flips. Conformational interconversion is no longer possible at room temperature when the methoxy moieties in 2b are replaced by ethoxy groups. Solvent effects in CDC$_3$ play a minor role, although compounds 2b and 3b are capable of complexing solvent molecules in the cavity formed by the four tert-butyl groups in the Cone form, as seen in the X-ray crystal structures of complexes with toluene, MeCN, or p-phenylcalix[4]arene complexed with CHCl$_3$. The [1]metacyclophanes 3 have recently been synthesized. These compounds are more flexible than either 1 or 2. No single characteristic conformation could be shown to be the most stable by NMR in CDC$_3$F solution. Because [1]metacyclophanes have the same skeleton as the calix[4]arenes, investigation of [1]metacyclophanes can yield insight into the influence of the lower rim substitute (X in Figure 1) and the skeleton on the properties of calix[4]arenes.

Recently, an algorithm has been developed for finding reaction pathways and transition states involving many degrees of freedom. This algorithm, called Conjugate Peak Refinement (CPR), yields a series of true saddle points which are connected to one another through minimum energy paths composed of a string of structures. The resulting path forms a continuous reaction coordinate between a given reactant and a given product. Unlike a trajectory obtained from molecular dynamics, which includes many fluctuations uncorrelated to the progress of the reaction, the reaction trajectory involves only the minimal amount of necessary motions. In the low-temperature limit, this sequence of conformations is the most probable path. At normal temperatures, it is likely to be a good approximation to the path obtained by averaging over the dynamic pathways, particularly if the conformational change is dominated by the energy rather than by entropy. CPR is currently the only available method for finding adiabatic reaction paths that is fully automated and reliable for systems that have complex energy surfaces involving a very large number of degrees of freedom. The CPR algorithm is integrated in the CHARMM program and has been successfully applied to the tyrosine ring-flip in BPTI, to the closure of the lid over the catalytic site of trypsin, and to the mechanism of rotamase catalysis by PKDP. Here, we use the CPR algorithm to exhaustively explore the conformational interconversions.

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version in [4]metacyclophanes 3a and in calix[4]arenes 1a and 2b. These compounds are representative of most compounds 1, 2, and 3, because the nature of the upper rim substituents has only a marginal effect on the total energy of the conformation.\textsuperscript{2} We propose detailed stepwise sequences for interconversion in [4]metacyclophanes 3 and in calix[4]arenes 1 and 2. The corresponding activation barriers of the conformational interconversion in calix[4]arenes 1a enable us to distinguish between the reaction mechanisms proposed for interconversions in 1.\textsuperscript{1,12} The energies of the transitions states in 2b are compared with kinetic data from 1H and 2D NMR. Results in agreement with experiment are obtained. They provide new insights into the nature of the transition states and indicate that version 22 of the all-hydrogen CHARM parameter set\textsuperscript{27} is appropriate for calix[4]arenes.

Methods

Energy Minimizations. For compounds 1a and 3a, the four characteristic conformations were considered and minimized. For compounds 2b, a systematic search with concomitant energy minimization of all possible in/out orientations of the methoxy groups in each of the four characteristic conformations was performed. The energies of the conformations were minimized by the Conjugate Gradient method and followed by the Newton–Raphson method. The root-mean-square (rms) value of the energy gradient was less than 0.0001 kcal mol\textsuperscript{-1} Å\textsuperscript{-1}. For compound 1a, the influence of cooperative polarization of the hydroxyl groups was investigated by varying the charge separation across the O–H bond. The bonded parameters, point charges, and van der Waals parameters were derived from a developmental version of the CHARM parameter set for the tyrosine and alliphatic amino acids side chains and for methyl acetate (see Appendix). A dielectric constant \(\varepsilon = 1\) was used with no cut-off for the nonbonded interactions. This seems appropriate, since most studies are carried out in weakly polar solvents. All calculations were done with the CHARM program.\textsuperscript{25}

Reaction Path Calculations. The Conjugate Peak Refinement (CPR) algorithm\textsuperscript{25} searches for maxima along the adiabatic energy valley connecting two local minima on an energy surface. These maxima are referred to saddle points and represent the transition states of the reaction pathway between the two minima. The algorithm has been integrated into the CHARM program\textsuperscript{27} in the TRAVEL module. The output of CPR consists of a series of \(N\) intermediate structures (each given by its coordinate vector \(x\)) along the reaction pathway from the reactant \(R\) to the product \(P\), including one or more saddle points. The one-dimensional reaction coordinate along this curvilinear path is given by

\[
l_i = \sum_{j=0}^{i-1} |r_j - r_i|
\]

The path-scanning step size \(\Delta l\) (the only parameter of the CPR method which has a system-dependent optimal value) was set initially to 1/6 of the rms difference between reactant and product coordinates. This step size was automatically reduced in TRAVEL until an optimum value is reached.\textsuperscript{24} The final value of \(\Delta l\) was 0.05 Å, except for the calculations on the polarized O–H bonds, where it got reduced to 0.01 Å. Presenting the optimum step size can save some computational overhead. The rms gradients of the saddle points were in the 10\textsuperscript{−6}–10\textsuperscript{−5} kcal mol\textsuperscript{-1} Å\textsuperscript{-1} range. In the absence of an initial guess for transition intermediates, the CPR method as implemented in TRAVEL generates a first guess for the pathway by linear interpolation between


(29) Mackinnon, A.; Karplus, M.; et al. To be submitted for publication.

the Cartesian coordinates of the reactant and the product. In a few cases, such an initial guess results in a path with some unlikely segments, for example in which a hydrogen passes through the aromatic ring. This problem is easily resolved by continuing the refinement after having excised the undesired segments from the path. After all the saddle points were fully refined by CPR, and for those paths shown here in the figures, the other path points were further relaxed into the adiabatic valleys of the energy by Synchronous Chain Minimization (SCM),\textsuperscript{29} a path optimization method related to the procedure by Chol and Eber.\textsuperscript{30} But better behaved in systems with many degrees of freedom.

The pathways for compounds 1a and 3a were generated by calculating the following transitions: Cone -- Pacc, Pacc -- 1,2,1Alt, Pacc -- 1,3Alt, Cone -- 1,2,1Alt, Cone -- 1,3Alt, and Cone -- Inverted Cone (Conc''), all without initial intermediates. Converged Cone -- Conc' pathways in 1a were obtained by increasing the charge separation on the OH moiety. The Pacc is the key intermediate for compound 2b, therefore the following transitions were calculated for this compound: all Pacc conformations to all Cone, 1,2Alt, and 1,3Alt conformations (conformations are described in Table 2). After the lowest-energy interconversion path was found from one characteristic conformation to the other, methyl rotation paths were calculated for each characteristic conformation, in order to find the complete lowest energy path from the ground state of each characteristic conformer to the ground state of every other characteristic conformer. Unless otherwise specified, the transition state energies given here are relative to the energy of the most stable conformer (global energy minimum) of 1a, 2b, or 3a, respectively.

Hammond Postulate. According to Hammond,\textsuperscript{31} "If two states, as for example, a transition state and an unstable intermediate, occur consecutively during a reaction process and have nearly the same energy content, their interconversion will involve only a small reorganization of the molecular structures". Although the Hammond postulate was formulated for chemical reactions, it is likely to apply also to conformational transitions. Thus, the change in geometry during a conformational interconversion should correlate with the change in energy. The applicability of the postulate was tested by comparing the progress along the reaction coordinate \(\lambda\) (taken relative to a given saddle point) to the change in the energy. For the structures \(k\) along the path on each side (\(k > 0\) or \(k < 0\)) of the saddle point (\(k = 0\)), we define:

\[
\Delta h_k = \lambda_k - \lambda_{\text{reactant}}
\]

\[
H_k = \text{MAX}(h_k) - h_k
\]

To allow a direct comparison of the energy and \(H_k\), the values \(H_k\) are scaled such that the amplitude of the change in \(H_k\) equals the energy barrier. \(H_k\) is plotted as a function of \(\lambda_k\) to show the correlation between changes in the energy and changes in the rms displacement of the atomic coordinates.

Results

(a) Geometry Optimizations. The minimized energies of the four characteristic conformers of compound 3a are listed in Table 1. These results are very similar to those previously obtained with the MM3 force field.\textsuperscript{40} Calculations with the MMX force field\textsuperscript{30} and MM2 force field\textsuperscript{30} also resulted in Cone as the lowest energy conformation. The energy differences between conformers is small, with a maximum value of 1.9 kcal/mol between the Cone and 1,3Alt conformers.

Also listed in Table 1 are the optimized energies of compound 1a, which are obtained when the four hydroxyls remain in the same orientation relative to their ring. It is the orientation that allows the formation of a circular array of four H-bonds in the Cone. The Cone is the lowest energy conformer, in accordance with 1H NMR data in various solvents.\textsuperscript{13} Earlier molecular mechanics calculations\textsuperscript{2,3} have yielded the same energy order: Cone < Pacc < 1,2Alt < 1,3Alt. Recent calculations


Table 1. Energy (kcal/mol) for the Characteristic Conformers of

\[ \text{[1]-Cyclophane 3a} \text{ and Calix[4]arene 1a} \]

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<tr>
<th>conf</th>
<th>MM3*</th>
<th>( E_{\text{vib}} )</th>
<th>( E_{\text{rot}} )</th>
<th>( E_{\text{elec}} )</th>
<th>( E_{\text{tot}} )</th>
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<td></td>
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<td>1.1</td>
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</table>

* Potential energy from ref 4h. \( E_{\text{vib}} = E_{\text{rot}} + E_{\text{elec}} + E_{\text{elec}} \).

Table 2. Energy (kcal/mol) of the Conformers of Calix[4]arene 2b

<table>
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<th>( E_{\text{rot}} )</th>
<th>( E_{\text{elec}} )</th>
<th>( E_{\text{tot}} )</th>
</tr>
</thead>
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<td>0.1</td>
<td>1.2</td>
<td>2.2</td>
<td>3.5</td>
</tr>
</tbody>
</table>

* The digits in the names of conformations indicate which anisole ring is inverted. The letters indicate whether the methoxy group attached to each aromatic ring points outwards (O) or inwards (I). For example, 0011~AAAA is a Paco in which the methoxy group on the invered aromatic ring points inwards. See also caption of Table 1.

(b) Interconversion pathways. (i) Ring-Flips in [1]-Metacyclophane 3a. The transition-state energies for the three saddle points connecting the Paco form to the Cone, 1,2Alt, and 1,3Alt conformers are listed in Table 3. The energy profiles of the corresponding reaction paths are shown in Figure 2. From the connectivity chart in Figure 3a, it is apparent that the Cone \( \rightarrow \) inverted Cone (Cone\') conversion with the lowest energy barrier is via the 1,2Alt intermediate, with the rate-limiting step (in terms of energy barriers) being the Cone \( \rightarrow \) Paco transformation, which has an activation barrier of \( \Delta E_{\text{act}} = 3.8 \text{kcal/mol} \). The energy barrier arises mainly from angle distortion

\[ \text{(2.4 kcal/mol) and unfavorable van der Waals interactions (} 2.8 \text{kcal/mol)} \], which are compensated somewhat by favorable electrostatic interactions \(-2.1 \text{kcal/mol}\). The Cone \( \rightarrow \) Cone' route via the 1,3Alt conformer is only slightly higher in energy; the rate-limiting steps are the Paco \( \rightarrow \) 1,3Alt \( \rightarrow \) 1,3Alt legs, whose transition-state energy relative to the Cone ground state is \( E_{\text{trans}} = 4.0 \text{kcal/mol} \). To search for a concerted reaction (i.e. without the Paco intermediate), a direct recombination of the Cone \( \rightarrow \) Cone' conversion was done with the Cone conformer taken as reactant and the Cone' conformer as product. The resulting pathway was found not to be concerted, but to proceed via the Paco and 1,2Alt intermediates.

Because there are four ways for a Cone conformer to convert into a Paco conformer (by flipping any one of the four benzyl rings), each with a potential energy barrier of \( \Delta E_{\text{act}} = 3.8 \text{kcal/mol} \).
Pathways for Conformational Interconversion of Calix[4]arenes


Figure 2. Energy profile (solid line, kcal/mol) and Hammond correlation (dashed line) for the ring-flipping isomerization reactions in compound 3a.

Figure 3. Connectivity chart and pathways for the Cone \( \rightarrow \) Cone' conversion: (A) cyclophane 3a, (B) calix[4]arene 1a. The numbers are the energies (\( \Delta H \)) of the stable conformers or \( \Delta H_{c} \) of the transition state of the reactions (in kcal/mol, corrected for conformational degeneracy, see text and Tables 1 and 3).

mol, the energy of activation of the Cone \( \rightarrow \) Paco transition corrected for this conformational entropy is \( \Delta H_{c} = \Delta H_{c} \rightarrow \Delta H_{c} \rightarrow kT \ln 4. \) At \(-30^\circ C\), \( \Delta H_{c} = 3.1 \) kcal/mol, which is below the upper limit of \( \Delta H \) \( < 5.9 \) kcal/mol deduced from 'H NMR measurements for the Cone \( \rightarrow \) Cone' conversion. \( ^{19} \) These results agree with the experimental observations that conformational interconversions in 1,4-metacyclophanes are very fast. \( ^{19} \) \( \Delta H_{c} \) is computed for all reactions listed in Table 3 by comparing the conformational degeneracy of the saddle points to the degeneracy of the ground state (see Discussion).

(ii) Stepwise Interconversion in 1a. For compound 1a, both stepwise (involving a single phenyl ring flip at a time) and direct (involving more than one ring-flip) interconversion pathways were calculated. The stepwise Cone \( \rightarrow \) Inverted Cone (Cone') reaction was investigated by individually refining the Cone \( \rightarrow \) Paco, Paco \( \rightarrow \) 2,2Alt, and Paco \( \rightarrow \) 1,3Alt paths. The direct Cone \( \rightarrow \) 2,2Alt, Cone \( \rightarrow \) 1,3Alt, and Cone \( \rightarrow \) Cone' paths were also refined. The direct Cone \( \rightarrow \) 2,2Alt path has the same saddle points as the stepwise Cone \( \rightarrow \) Paco, Paco \( \rightarrow \) 2,2Alt pathway. Similarly, the direct Cone \( \rightarrow \) 2,3Alt path proceeds via the stepwise Cone \( \rightarrow \) Paco \( \rightarrow \) 1,3Alt pathway. The direct Cone \( \rightarrow \) Cone' path also proceeds via a stepwise pathway, so that all interconversions in 1a can be broken down into three reaction steps, whose transition-state energies are listed in Table 3. The corresponding potential energy profiles along these reactions are shown in Figure 4. Converged paths, with two or more rings flipping simultaneously, were not found. Figure 4B shows the possible pathways for a Cone \( \rightarrow \) Cone' interconversion. The overall rate-limiting step (as defined by the highest energy barrier) of the Cone \( \rightarrow \) Cone' interconversion is the Cone \( \rightarrow \) Paco transition, with an activation barrier of \( \Delta H_{c} = 15.2 \) kcal/mol. The pathway then proceeds via the 1,2Alt conformer, which has a much lower barrier than the Paco \( \rightarrow \) 1,3Alt path. Accounting for the conformational entropy of the Cone \( \rightarrow \) Paco transition (see previous section and Discussion), \( \Delta H_{c} = 14.5 \) kcal/mol, in excellent agreement with the experimental value of 14.2 kcal/mol in CDCl\( _3 \) as determined by 'H NMR for the Cone \( \rightarrow \) Cone' interconversion. \( ^{19} \) Much of the barrier comes from breaking two hydrogen bonds, which amounts to \( \Delta H_{c} = 9.1 \) kcal/mol. Most of the remainder of the activation barrier comes from the deformation of some bond angles, such as the AR-C-H-AR angles (from 110° to 120°) of the two methylene groups to which the rotating ring is attached, with \( \Delta H_{c} = 5.9 \) kcal/mol. In the transition state, the rotating ring makes an angle of \( \sim 20^\circ \) with the mean plane of the four methylene carbon atoms, which means that the phenol group has partly passed through the plane of the methylene carbon atoms (Figure 5). During the transition, the opposite phenol ring rotates to be more in plane with the annulus, whereas the other two phenol rings move the other way and become closer to perpendicular to the plane of the methylene groups.

Earlier reaction path calculations have been performed on 1e by forcing internal rotations around an axis going through the two methylene groups attached to the phenolic ring. \( ^{3} \) This torsion angle was constrained and the rest of the calix[4]arene was relaxed by energy minimization. The resulting energy barriers for the interconversions Cone \( \rightarrow \) Paco, Paco \( \rightarrow \) 1,2Alt, and Paco \( \rightarrow \) 1,3Alt were 8.1, 8.9, and 9.4 kcal/mol, respectively. The force field used, MM2P, \( ^{9} \) treated the hydrogen bonds only in a qualitative way, so that lower barriers were obtained even though the optimal path was not followed. The predefined reaction coordinate did not ensure full relaxation during the interconversion and the maxima along this reduced adiabatic path were not resolved to true saddle points.

(iii) Converged Interconversion in 1a. Investigation of the strength of the hydrogen bonds in calix[4]arenes with IR spectroscopy in CCl\( _4 \) has suggested that the hydrogen bonding in calix[4]arenes may be cooperative, i.e. the breaking of the first H-bond in the circular array of four H-bonds is more difficult than the breaking of additional H-bonds. This is
believed to be caused by polarization of the O–H bonds in the circular array. The polarization effect in 1a was investigated here by adding a charge $\Delta q$ to the hydrogens and $-\Delta q$ to the oxygens of the hydroxyl groups. For every $\Delta q$, the conformers were reoptimized and the direct Cone $\rightarrow$ 1,2Alt and Cone $\rightarrow$ 3,1Alt conversion pathways were refined. In each case, the stepwise pathways through the Paco intermediate become a concerted movement of the phenol rings when $\Delta q \geq 0.35 \ e$, as denoted by the presence of a single saddle point instead of two successive saddle points along the path. With $\Delta q = 0.35$, resulting in a partial charge of $-0.89 \ e$ on O and $+0.78 \ e$ on H, the potential energy of the unique saddle point relative to the reoptimized Cone is 37.4 kcal/mol for the Cone $\rightarrow$ 1,2Alt path and 66.9 kcal/mol for the Cone $\rightarrow$ 1,3Alt path. The structure of the transition state is respectively 1,2Alt-like and 1,3Alt-like and it is characterized by the preservation of the four hydrogen bonds in a partially distorted circular array, at the cost of considerable distortion of the calix[4]arene framework (Figure 6). The 1,2Alt$^+$ conformer more readily accommodates the four hydrogen bonds than the 1,3Alt$^+$ conformer, because two H-bonds are still present in the stable 1,2Alt conformer, whereas the stable 1,3Alt conformer has all four H-bonds broken. This explains why the concerted pathway via the 1,2Alt$^+$ conformer is much more favorable than that via the 1,3Alt$^+$ conformer. The value of $\Delta q$ at which this pathway changes from a stepwise pathway to a concerted pathway coincides with the $\Delta q$ at which the Paco conformation becomes a hybrid Paco/1,2Alt conformation (Figure 7). The concerted pathway via the 1,2Alt$^+$ conformer resembles the proposed “continuous-chain” pathway. While these results indicate that a “continuous-chain” pathway is in principle possible, it is unlikely because of the unrealistically large partial charges on the hydroxyl groups necessary to obtain it and because the potential energy of the 1,2Alt and the 1,3Alt conformers relative to the Cone conformer increases significantly with $\Delta q$, reaching 35.7 and 66.4 kcal/mol, respectively, when $\Delta q = 0.35$, i.e. only 1.7 and 0.35 kcal/mol less than the respective transition-state energies. Most importantly, the corresponding activation barriers are irreconcilable with the kinetic data from NMR experiments (see previous section). Therefore, our calculations indicate that a concerted mechanism is unrealistic, i.e. the conversion from Cone conformer to inverted Cone conformer in 1 proceeds via a stepwise mechanism with the Paco and 1,2Alt conformers as intermediates (see Figure 3B).

(iv) Interconversions In Calix[4]arene 2b. For the trimethyl ether 2b the situation is more complicated because of the many minima that are introduced by the two possible orientations for each methoxy group (see Table 2). The orientation of the methoxy groups affects the energy barriers of the anisole ring rotations, which are large compared to the barriers of the methoxy rotations themselves. The transition-state potential energies of some methoxy rotations are listed in Table 4 and can be seen to be in the 8–10 kcal/mol range, whereas the lowest ring rotation transition states are in the 20–24 kcal/mol range. Taken relative to the ground states of each

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**Figure 5.** Saddle-point structure (stereo) of the ring-flipping isomerizations in compound 1a: (A) Cone $\rightarrow$ Paco, (B) Paco $\rightarrow$ 1,2Alt, (C) Paco $\rightarrow$ 1,3Alt. In each case, the converting ring is on the right and is rotating clockwise for the indicated reaction direction.

**Figure 6.** Saddle-point structure (stereo) of concerted ring-flipping in 1a with hydroxyl charge separation increased by $\Delta q = 0.35 \ e$: (A) Paco $\rightarrow$ 1,2Alt, (B) Paco $\rightarrow$ 1,3Alt.

**Figure 7.** Optimized structures (stereo) of the 1a Paco conformer obtained with increments in the hydroxyl charge separations $\Delta q$ of 0.0, 0.1, 0.2, 0.3 and 0.35 e.

**Table 4.** Potential Energy (kcal/mol) of the Transition State in Isomerization Reactions of 2b$^a$

<table>
<thead>
<tr>
<th>reaction</th>
<th>$E^\text{con}$</th>
<th>$E^\text{cone}$</th>
<th>$E^\text{cone}$</th>
<th>$E^\text{paco}$</th>
<th>$\Delta E^\text{pm}^b$</th>
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<tr>
<td>C$^\text{a}$, Cone $\rightarrow$ C$^\text{a}$, Con$^\text{a}$</td>
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<td>2.6</td>
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<td></td>
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<tr>
<td>0000$\rightarrow$AAAA$\rightarrow$AAAAA$\rightarrow$AAAAA</td>
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<td>5.9</td>
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<td>7.8</td>
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$^a$ The calculations listed here are the ones which have the lowest saddle-point energy for each given type of reaction (such as a methoxy rotation in a Cone, in a Paco, etc.). Energies are relative to the Paco ground state (0001$\rightarrow$AAAAA). See also caption of Tables 1 and 2. $^b$ $\Delta E^\text{pm}$ is the energy barrier relative to the ground state of the respective conformation (i.e. Cone, Paco, etc.). $^c$ Energies are relative to the 0000$\rightarrow$AAAAA Cone.
Figure 8. Energy profiles (kcal/mol) for the (A) Paco → Cone, (B) Paco → 1,2Alt, (C) Paco → 1,3Alt ring-flip conversions of 2b. See the caption of Table 2 for the nomenclature of the conformations.

of the four characteristic conformers of 2b, the activation barriers to methoxy rotation are only 6–8 kcal/mol (Table 4). For example, the 0000~AAAA → 0000~AABA reaction has a barrier of \( \Delta G^\star_{pm} = 5.9 \text{ kcal/mol} \) relative to the Cone ground state. A search for the preferred ring interconversion pathway cannot be limited to the conformational transitions between the reactant and product conformers with the most stable methoxy orientation, because paths for ring conversion with lower overall transition-state energy might connect characteristic conformers whose methoxy groups are not in the ground-state orientation. The conversion paths from the Paco conformer to the Cone, 1,2Alt, and 1,3Alt conformations were determined for all combinations of reactant and product methoxy orientations, for a total of 228 reactions.

The Paco → Cone conversion with the lowest saddle point potential energy of 19.6 kcal/mol is found with 0001~AABA as the Paco reactant (Table 4), which is 2.2 kcal/mol higher than the 0001~ABAA Paco ground state (Table 2). The Paco → Cone conversion directly from the Paco ground state yields a saddle point whose potential energy is slightly higher at 19.8 kcal/mol. The transition from the Paco ground state to 0001~AABA has a barrier of only 8.1 kcal/mol and therefore is not the rate-limiting step of the Paco → Cone ring-flip. The situation is similar for the Paco → 1,2Alt conversion, where the lowest transition state is also achieved when 0001~AABA is the Paco reactant, and for the Paco → 1,3Alt transition, which proceeds from the 0001~BAAA Paco state. The smallest activation barriers relative to the Paco ground state are then 19.6, 20.2, and 18.2 kcal/mol respectively for the Paco → Cone, Paco → 1,2Alt and Paco → 1,3Alt conversions. The corresponding potential energy profiles are shown in Figure 8, parts A, B, and C. The pathways connecting the four ground states of the characteristic conformers via methoxy rotation and anisole ring conversion are shown in Figure 9. In all cases, the activation energy for ring conversion is dominated by an increase of about 9 kcal/mol in van der Waals energy and an increase in bond stretch and bond angle energy of 3–6 kcal/mol. The electrostatic contribution to the barriers is smaller than in calix[4]arene 1a and comparable to that found in compound 3a (See Table 3), due to the lack of strong H-bonds. The conformations of the saddle points are shown in Figure 10.

The C_4v symmetric Cone of 2b has been proposed to be the saddle point of the interchange of two equivalent C_2 symmetric Cone conformations. In the C_4v form, the planes of two facing anisole rings are almost parallel to each other ("pinched") and
Figure 10. Saddle-point structures (stereo) of the ring-flipping isomutations in compound 2b: (A) 0001~AABA ~ 0000~AAAB, (B) 0001~AABA ~ 0011~AAAA, (C) 0001~BAAB ~ 0101~BAAA. In each case, the converting ring is on the right and is rotating clockwise for the indicated reaction direction. See footnote of Table 2 for an explanation of the conformation names.

Figure 11. Energy profile (solid line, kcal/mol) and Hammond correlation (dashed line) for the conversion from one C2v Cone to the other in 2b.

orthogonal to the annulus while the other two anisole rings are more coplanar with the annulus, unlike the C2v Cone which has the four rings at the same angle relative to the plane of the annulus. We have calculated the transition from one 0000~AAAA C2v Cone to the other (Figure 11). The saddle point is only 3.1 kcal/mol higher in potential energy than the C2v Cone and is indeed a C2v Cone. A comparable result was obtained by constrained energy minimization using the AMBER 3.0 force field.2 C2v ~ C2v' transitions have been observed during molecular dynamics simulations21 of lower ring substituted derivatives of 2. The type of lower ring substituent affects the propensity for the C2v form.2

Discussion

The distribution7,46 of the conformations of 2b as determined with 1H NMR in CDCl3 at −30 °C is 85% Paco, 8% 1,2Alt, 4% Cone, and 3% 1,3Alt. The accuracy in these particular NMR spectra is expected to be around 2% (absolute). The Boltzmann weighted probability of finding compound 2b in state x, corrected for the conformational degeneracy and neglecting vibrational effects on relative populations (vibrational and rotational contributions to the free energy will be presented elsewhere), is given by12

$$p(x) = \frac{\sum_n e^{-\beta E_n}}{\Omega}$$

where n is the degeneracy of conformer i and $E_i$ is the potential energy (both given in Table 2). $\Omega$ is the partition function

$$\Omega = \sum_n e^{-\beta E_n}$$

For example, there are two possible AAAA Cones for compound 2b, 0000~AAAA and 1111~AAAA, so the degeneracy of that conformer is n = 2. With the data in Table 2, p(Paco) = 53.9%, p(1,2Alt) = 18.4%, p(1,3Alt) = 18.3%, and p(Cone) = 0.006% at −30 °C. The Paco state is correctly predicted as the most probable conformation, but its calculated probability is somewhat lower than the experimental one. Also, the order of the abundance of the Cone and 1,3Alt forms is inverted.

If the relative energy differences due to methoxy rotation are assumed to be correct within each characteristic conformation of 2b and a constant correction term $E_r$ is added to the potential energy (as given in Table 2) of every conformer in the characteristic state x, then the exact experimental distribution is obtained for $\Delta E_{Paco} = 0.0, \Delta E_{1,2Alt} = +0.54$ kcal/mol, $\Delta E_{1,3Alt} = +1.0$ kcal/mol, and $\Delta E_{Cone} = -3.0$ kcal/mol. If only the lowest conformer of each type is considered, the result is $\Delta E_{Paco} = 0.0, \Delta E_{1,2Alt} = +0.44$ kcal/mol, $\Delta E_{1,3Alt} = +0.43$ kcal/mol, and $\Delta E_{Cone} = -3.0$ kcal/mol. With the MM3 force field, the following relative stabilities have been found:4 Paco > 1,3Alt ~ Cone > 1,2Alt; i.e., the ranking of the Cone was consistent with experimental data, but the 1,3Alt and 1,2Alt were inverted. A study with the AMBER 3.0 force field resulted in 1,3Alt > Paco > Cone > 1,2Alt. Quantitatively correct values are difficult to achieve, due to the complex balance of forces, particularly between covalent and electrostatic terms (see Table 2).

The Paco conformation has a high field position (δ = 1.99 ppm) for one of its three methoxy signals in the 1H NMR spectrum of 2b in CDCl3,7 indicating that one strongly shielded methoxy group is pointing inwards. This agrees with two of the three most stable Paco forms found here, 0001~AABA and 0001~AAAB (Table 2), which both have one methoxy pointing inward. The Cone has one single low field methoxy signal (δ = 3.83 ppm), indicating that the four unshielded methoxy groups are pointing outwards. This is consistent with the most stable Cone conformer, which has all methoxy groups pointing out (0000~AAAA in Table 2). In the 1,2Alt and 1,3Alt conformers, the four methoxy groups show one single intermediate field

signal ($\delta = 2.99$ ppm for the 1,2Alt conformer and $\delta = 2.87$ ppm for the 1,3Alt conformer). This means that either the four methoxy groups are pointing inward and have intermediate shielding or there is fast exchange between the strongly shielded inward and the unshielded outward orientations. From the calculations, having four methoxy groups pointing inward is very unfavorable. The 1,3Alt is only able to accommodate all four methoxy groups pointing inwards at a cost of 12.9 kcal/mol relative to the all-out 1,3Alt conformer. The all-inward conformation of 1,2Alt is unstable. This suggests that in both the 1,2Alt and the 1,3Alt conformers there is a fast exchange between conformations with methoxy groups pointing inward and outward. For example, the most populated 1,2Alt conformer (ABAB) has two methoxy groups in and two out. If the methoxy groups are rapidly interchanging (ABAB$\leftrightarrow$ABAB), a single intermediate field signal is expected. This is consistent with the calculations, which show that the methoxy rotations in the 1,2Alt and 1,3Alt conformers have low transition barriers relative to their respective ground states ($\Delta E_{rot}^{pe}$ in Table 4). The rate of interconversion between two species at a temperature where their signals have just coalesced in the spectrum (the coalescence temperature) can be estimated from the separation of the two signals (in Hz) at a low temperature where the two species are not exchanging. As an upper limit to the barrier for methoxy rotation can be estimated for the 1,2Alt and 1,3Alt conformers, whose inward and outward methoxy signals have coalesced. If one takes 1.84 ppm as an upper bound to the separation between the inward and outward methoxy signals (from the observed values of the methoxy signals, 1.99 ppm inward in Paco and 3.83 ppm outward in Cone), this corresponds to 1104 Hz at 600 MHz. Assuming Arrhenius kinetics, equal population of the two states, and $-50^\circ$C as the coalescence temperature, the energy barrier to methoxy rotation is $\Delta E_{rot}^{pe} \leq 10.7$ kcal/mol. Our calculated barriers for methoxy rotation ($\Delta E_{rot}^{pe}$ in Table 4) are in agreement with this limiting value.

The $^1$H NMR data for the Cone conformation of 2b are consistent with either a C$_{6v}$ symmetric Cone or the average structure resulting from the rapid interconversion between the two possible C$_{6v}$ Cone conformers, in which the C$_{6v}$ Cone is the transition state. The optimized geometry of the 2b Cone has C$_{6}$ symmetry (“pinched Cone”, see Results section). We found the C$_{6v}$ Cone to be the saddle point in the interconversion from one C$_{6v}$ Cone to the other, with an activation barrier of 3.1 kcal/mol, which suggests that fast interconversion is occurring.

The 2D EXSY NMR spectra of a calixcrowns derivative of 2a, which has two facing rings bridged at the upper rim so that the formation of the 1,2Alt conformer is prevented, have shown that Poco can convert to Cone and to 1,3Alt conformers. The
conversion to the 1,3-Alt conformer is faster than conversion to the Cone conformer. This is in accord with the respective activation barriers of 18.2 and 19.6 kcal/mol calculated here (see Table 4). Measurements on 2b with 1H and 2D EXSY NMR indicate that the conversions from Paco to Cone or to 1,3-Alt are faster than the conversion from Paco to the 1,2-Alt conformer.\(^7\) This is also consistent with the higher activation barrier of the Paco \(\rightarrow\) 1,2-Alt path (20.2 kcal/mol in Table 4).

The pertinence of the Hammond postulate to interconversion reactions in calix[4]arenes was tested by correlating the root-mean-square (rms) displacement of the atomic positions with the energy of the system (see Methods section). In most cases, the reaction energy profiles (see Figures 2, 4, 8, and 11) show an excellent correlation with the change in rms of the atomic positions along the path, indicating that the conformational changes in calix[4]arenes obey the Hammond postulate.

Although no solvent effects were included in the calculations, close agreement has been found between the calculated and experimental \(\Delta G^\circ\) for the Cone \(\rightarrow\) inverted-Cone conversion of 1a in CDCl\(_3\). This is not surprising, since the \(\Delta G^\circ\) as determined by \(^1\)H NMR is nearly solvent independent.\(^10\) In contrast, the conformational distribution of 2b found by \(^1\)H NMR is influenced by the solvent,\(^15\) mostly changes in the abundance of the Cone form at the cost of the other forms: 5% Cone in CDCl\(_3\), 14% Cone in CS\(_2\) and CCl\(_4\), 18% Cone in toluene. Explanations for this behavior have been sought either in terms of the polarity of the solvent\(^23\) or by the inclusion of a solvent molecule in the cavity of the Cone.\(^13\) Solvent polarity cannot explain all the solvent effect data: e.g. the Cone conformer of 2b is more abundant in CCl\(_4\) than in CDCl\(_3\), although the latter is the more polar solvent and the Cone is the conformation with the largest dipole moment, which would be expected to be more abundant in a more polar solvent.\(^15\) More likely, enlargement of the cavity in the tert-butyl-substituted calix[4]arenes allows for complex formation with the solvent.\(^14\,15\)

Conclusions

The energies and conformational transitions of calix[4]arenes 1a and 2b and [1]metacyclophane 3a have been studied using the CHARMM molecular mechanics potential with the new parameter set 22. Although these parameters were developed for amino acids, they could be used without modification. The relative stabilities for the four conformations of the three compounds are in good agreement with experimental NMR data. The same is true for the activation barriers of the ring interconversions between the characteristic conformers of 1a, 2b, and 3a and of the methoxy rotations in 2b. The present study provides detailed information concerning the calculated pathways connecting the various conformers. This was made possible by the CPR algorithm, which has been implemented in the CHARMM program. In the absence of an initial guess for transition intermediates, the method starts with a linear interpolation pathway from the reactant to the product conformation, which is then refined. In nearly every case, this was sufficient to yield the pathway with the lowest activation barrier for the Cone \(\rightarrow\) Cone' conversions. Only in 1a did a stepwise refinement of Cone \(\rightarrow\) Paco followed by Paco \(\rightarrow\) 1,2-Alt find a lower saddle point than the direct Cone \(\rightarrow\) Cone' calculation, which proceeded via the 1,3-Alt intermediate.

The pathways for [1]metacyclophane 3a Cone \(\rightarrow\) Cone' interconversion show no marked preference for the 1,2-Alt or 1,3-Alt intermediate. The structure of the 1a Cone form is close to the Cone form of the 3a [1]metacyclophane, indicating that the circular array of H-bonds in 1a puts little additional strain on the calix[4]arene frame. The most likely Cone \(\rightarrow\) Cone' conversion of 1a is along a stepwise Cone \(\rightarrow\) Paco \(\rightarrow\) 1,2-Alt \(\rightarrow\) Paco' \(\rightarrow\) Cone' pathway. The hypothesis of a concerted mechanism, in which the four phenol rings of 1a would co-rotate to conserve the four H-bonds, is not consistent with the present calculations, as demonstrated by the excessive hydroxyl O=H bond polarization necessary to achieve such a mechanism. The relative stability of the 2b characteristic conformers is affected by the conformational degeneracy of the energy states due to the symmetry of the molecule. Even though the 1,2-Alt form 0011/ABAB is ranked higher in terms of potential energy than the 1,3-Alt form 0101/AAAA, consideration of the conformational degeneracy reveals that the former is more stable than the latter, in accordance with experiment. Similarly, the stability ranking of the 1,3-Alt forms 0101/AAAA and 0101/ABAB, differing by their methoxy orientations, is inverted after inclusion of the conformational degeneracy of these structures.

Interconversions between the characteristic conformers of 2b occur with the Paco form as the key intermediate. The anisole ring interconversions with the lowest absolute transition-state energies are not between the ground states of the respective characteristic conformers but between conformations that are related to their ground states via one (in the case of 1,3Alt), two (in the case of Cone and 1,2Alt), or up to three (in the case of Paco) rotations of the methoxy groups. Rotations of the methoxy moieties have energy barriers well below the energy barriers for anisole ring rotation, explaining why the methoxy hydrogens give a single peak in \(^1\)H NMR. The NMR data in favor of a \(C_\alpha\) symmetric Cone, i.e., the aromatic protons appear as one singlet and the protons on the methylene bridges appear as an ABq system, can be explained by the low barrier between the two stable \(C_\beta\) forms, which allows fast exchange on the NMR time scale.

The present analysis demonstrates the power of molecular mechanics combined with an efficient and general technique for finding reaction paths to explore in detail the connectivity on potential energy surfaces of complex molecules with multiple conformers.

Note Added in Proof: M. Hosseini pointed out the paper by K. B. Lipkowitz and G. Pearl [J. Org. Chem. 1993, 58, 5729–5736] in which molecular mechanics calculations with different programs for a number of calix[4]arenes were reported. Species 2b is the only one that was calculated by Lipkowitz and Pearl, as well as in the present paper. In contrast to their results, we find good agreement with experiment. G. Wipff and M. Lauterbach (private communication) have obtained energy differences similar to those reported in the present paper; they have also studied the effect of solvent on the energy differences.

Acknowledgment. The work reported here was supported in part by a grant from the National Science Foundation and a gift from Molecular Simulations Inc. to Harvard University.

Appendix

New molecular mechanics parameters were used for the calix[4]arenes in this study. Figure 12 shows the atom types and their partial charges. The nonbonded van der Waals parameters are given in Table 5 and the bonded parameters in Table 6.