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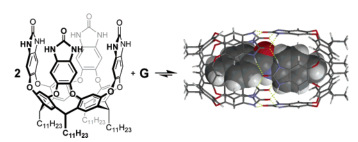
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ABSTRACT



An unusually stable molecular capsule was formed by heating phenyleneurea-spanned resorcinarene cavitand with 4-methyl-N-p-tolylbenzamide. The molecular capsule behaved as a discrete molecular entity showing a cylindrical D_{4d} structure and showed no guest exchange in toluene- d_8 even at 100 °C.

Self-assembled molecular capsules held together by hydrogen bonds or metal—ligand interaction have attracted much interest.^{1–3} The successful application of the complementary hydrogen-bonding theme was first introduced by the groups

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of Rebek and de Mendoza for self-assembling glycoluril-derived capsules such as "tennis balls" and "softballs".^{4–10}

Cavitand 1 reported by Rebek and co-workers self-assembled to form a cylinder-shape capsule 1·1 by eight bifurcated hydrogen bonds between an imide hydrogen atom in a cavitand with two neighboring carbonyl oxygen atoms as shown in Figure 1.^{11–23} Capsule 1·1 can accommodate various guests such as 4-methyl-*N*-*p*-tolylbenzamide 4 in a

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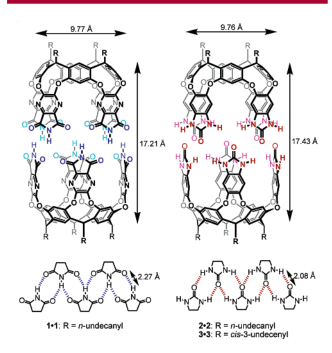


Figure 1. Hydrogen bonding modes of self-assembled pyrazinimide capsule 1·1 and phenyleneurea capsule 2·2. The bond lengths were calculated from the energy-minimized structures (Spartan 04, V1.0.1, AM1 semiempirical).

nonpolar mesitylene- d_{12} . The inclusion complexes **G@1·1** show interesting guest-exchange phenomena and unprecedented isomerism. $^{11-23}$

Phenyleneurea cavitands 2, 3, and other analogues were first reported by de Mendoza et al.24 These cavitands aggregated, forming different self-organized structures such as vesicles or filaments, depending on the nature and length of the four alkyl feet. Cavitand 2 formed large reverse vesicles through side to side extensive stacking. In contrast, cavitand 3 formed dimeric capsules with carboxylic acids.

We report the observation of the unusually stable capsule formation of cavitand 2. At elevated temperature guest molecules could template cavitand 2 to form a dimeric capsule 2.2 by reorganizing the intermolecular hydrogen bonds of the aggregate of cavitand 2.

Cavitand 2 was directly prepared as reported by de Mendoza,²⁴ and the intermediates, i.e., the corresponding octanitro cavitand and octaamino cavitand reported by Rebek^{23,25-27} were similarly prepared by the method reported by Cram et al.28

At room temperature, cavitand 2 was barely soluble in CH₂Cl₂, CHCl₃, toluene, or mesitylene but reasonably soluble in DMF or DMSO, as reported by de Mendoza.²⁴ However, at elevated temperature it became soluble in these nonpolar solvents and then stayed in homogeneous solution at room

The encapsulation studies of cavitand 2 and 4-methyl-Np-tolylbenzamide 4 were performed in mesitylene as devised by Rebek. 12,13 A mixture of cavitand 2 and guest 4 in mesitylene remained as a heterogeneous mixture at room temperature even after 5 days, but at above 100 °C the mixture was slowly homogenized. The encapsulation complex 4@2.2 was prepared by heating the mixture under reflux until it became homogeneous and then removing the solvent by vacuum distillation at 70 °C. The solid residue was dried at 100 °C under vacuum. The solid complex 4@2.2 is then soluble in CDCl₃, DMSO- d_6 , or toluene- d_8 at room temper-

An equimolar mixture of cavitand 2 and guest 4 (1.67 mM: 1.67 mM) in mesitylene gave a soluble complex in a relatively short period (30 min at 170 °C), but incomplete encapsulation was observed by ¹H NMR spectroscopy, which showed three different chemical shifts of the N-H of urea moieties for cavitand 2 (10.35, 10.36, and 10.40 ppm). The chemical shift of the N-H of cavitand 2 was a good indicator of whether the encapsulation complex 4@2.2 was formed completely or partially: the chemical shift of 2.2 in toluene d_8 was observed at 10.35 ppm.

A complete encapsulated complex 4@2.2 was prepared from a mixture of cavitand 2 and guest 4 (1.67 mM:6 mM, respectively) in mesitylene by heating under reflux for 30 min. The ¹H NMR spectrum of $4@2\cdot2$ in toluene- d_8 at 100 °C showed the chemical shifts for guest 4 at 5.41 (d, H_c), 5.20 (d, H_e), 3.29 (d, H_b), 3.14 (d, H_f), -2.33 (s, H_g), and -2.41 (s, H_a) ppm (Figure 2 and Table 1). Like the isomorphous cavitand 1·1,11 the large upfield chemical shifts of encapsulated guest 4 in capsule 2.2 are observed up to 4.50 ppm for the methyl group (H_a, 2.09 ppm) of free 4 in the ¹H NMR spectrum as a result of the shielding by aromatic ring current of capsule 2.2. Compared to the chemical shifts $(\Delta \delta_1)$ for **4** in **1·1**, those $(\Delta \delta_2)$ in **2·2** are slightly smaller $(\Delta\Delta\delta = 0.40, 0.37 \text{ for } H_a \text{ and } H_g, \text{ respectively}), \text{ which is}$ consistent with the molecular dimensions (17.21 vs 17.43 Å through the long C_4 axis of $1\cdot 1$ and $2\cdot 2$, respectively) calculated using semiempirical AM1.

The two different chemical shifts of the N-H of urea moieties for cavitand 2 (10.40 and 10.36 ppm) confirm that

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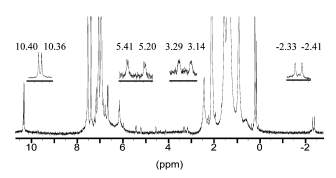


Figure 2. ¹H NMR spectrum of encapsulation complex 4-methyl-N-p-tolylbenzamide@**2**•**2** in toluene-d₈ at 100 °C.

two chemically nonequivalent methyl groups of guest 4 occupy each cavitand of the dimeric capsule and the guest's head-to-tail rotation is tightly restricted on the ¹H NMR time scale even at 100 °C.

The capsular complex $4@2\cdot2$ was unusually inert in nonpolar solvents such as toluene- d_8 . No guest exchange with the solvent molecule was observed in toluene- d_8 at 100 °C for 24 h. However, in polar solvents such as DMSO- d_6 the guest 4 in capsular complex $4@2\cdot2$ escaped rapidly at room temperature.

Table 1. Comparison of ¹H NMR Chemical Shifts (ppm) of 4-Methyl-N-p-tolylbenzamide **4** and Encapsulation Complexes **4@1·1** and **4@2·2** in Toluene- d_8^a

	protons						
	Ha	H_{b}	H_{c}	H_{d}	$\mathrm{H_{e}}$	${ m H_f}$	${ m H_g}$
free 4	2.09	6.96(d)	7.54(d)	d	7.40(d)	6.92(d)	2.12
$4@1 \cdot 1^{b}$	-2.81	c	c	c	c	c	-2.70
$\Delta\delta_1$	4.90						4.82
$4@2 \cdot 2$	-2.41	3.29(d)	5.41(d)	4.54	5.20(d)	3.14(d)	-2.33
$\Delta\delta_2$	4.50	3.67	2.13		2.20	3.78	4.45
$\Delta\Delta\delta$	0.40						0.37

 $^a\Delta\delta_1=\delta$ of free $\mathbf{4}-\delta$ of $\mathbf{4@1\cdot 1}$, $\Delta\delta_2=\delta$ of free $\mathbf{4}-\delta$ of $\mathbf{4@2\cdot 2}$. $\Delta\Delta\delta=\Delta\delta_1$ - $\Delta\delta_2$. b In mesitylene- d_{12} at 295 K. 11 c Not reported. d Obscured with solvent peak.

Capsular complexes with smaller guests such as p-xylene were prepared by heating cavitand 2 in p-xylene. In the toluene- d_8 , p-xylene escaped rapidly as a result of the dynamic equilibrium with toluene- d_8 by mass law ($t_{1/2} = 37$ min at 50 °C).

It is obvious that the hydrogen bonding mode in the self-assembly 2·2 is superior to that in 1·1, which has been already observed by de Mendoza.²⁴ For molecular capsule 2·2, two sets of lone pair electrons in the sp² hybridized carbonyl oxygen complementarily hydrogen bond with two adjacent H-N protons, which consumes all of the potential hydrogen bond donors and acceptors. However, in the self-assembly 1·1 only one set of lone pair electrons of the carbonyl oxygen hydrogen bonds to the adjacent H-N proton, as shown in Figure 1.

The unusual stability of capsule $4@2\cdot2$ in toluene- d_8 without exchanging guest 4 with solvent even at 100 °C suggests that, in addition to the exceptional tight hydrogen bond between two cavitands 2, van der Waals interaction between guest 4 and capsule $2\cdot2$ and the dipolar attraction or hydrogen bond between the amide group of guest 4 and the urea groups of the capsule $2\cdot2$ are efficiently operating.

However, in polar solvents such as DMSO- d_6 the guest 4 in capsular complex $4@2\cdot2$ escaped rapidly at room temperature. The polar DMSO- d_6 molecules compete for the hydrogen bond acceptors in the dimeric capsule $2\cdot2$, which weakens the stability of capsule $2\cdot2$ and allows the fast escape of guest 4 and/or the fast exchange of guest 4 with solvent. The observed vase conformation of 2 in DMSO- d_6 confirms that cavitand 2 still exists as a dimeric capsule DMSO- $d_6@2\cdot2$ because monomeric cavitand 4 could only exist as rectangular kite conformation as a result of the repulsion between adjacent urea groups. 28

In summary, the solvent- and/or guest-assisted molecular capsule formations $G@2\cdot2$ were observed and characterized by heating a mixture of cavitand 2 and guests in a nonpolar solvent at the elevated temperature.

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Supporting Information Available: Synthetic procedures, encapsulation experiments, molecular modeling, and selected NMR spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

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