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Cavitand-Based Pd-Pyridyl Coordination Capsules: Guest-Induced Homo- or Heterocapsule Selection and Applications of Homocapsule to Protection of Photosensitive Guest and Chiral Capsule Formation

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Abstract: A 2:4 mixture of tetrakis[4-(4-pyridyl)phenyl]cavitand (1) or tetrakis[4-(4-pyridyl)phenylethynyl]cavitand (2) and Pd(dppp)(OTf)₂ self-assembles into a homocapsule $\{\mathbf{1}_2 \cdot [\mathsf{Pd}(\mathsf{dppp})]_4\}^{8+} \cdot (\mathsf{TfO}^-)_8$ (C1) or $\{\mathbf{2}_2 \cdot [Pd(dppp)]_4\}^{8+} \cdot (TfO^-)_8$ (**C2**), respectively, through Pd–Npy coordination bonds. A 1:1:4 mixture of 1, 2, and Pd(dppp)(OTf)₂ produced a mixture of homocapsules C1, C2, and a heterocapsule $\{1\cdot 2\cdot [Pd(dppp)]_4\}^{8+} \cdot (TfO^{-})_8$ (C3) in a 1:1:0.98 mole ratio. Selective formation (self-sorting) of homocapsules C1 and C2 or heterocapsule C3 was controlled by guest-induced encapsulation under thermodynamic control. Applications of Pd-Npy coordination capsules with the use of 1 were demonstrated. Capsule C1 serves as a guard nanocontainer for trans-4,4'-diacetoxyazobenzene to protect against the trans-to-cis photoisomerization by encapsulation. A chiral capsule $\{\mathbf{1}_2 \cdot [Pd((R)-BINAP)]_4\}^{8+} \cdot (TfO^-)_8$ (C5) was also constructed. Capsule C5 induces supramolecular chirality with respect to prochiral 2,2'-bis(alkoxycarbonyl)-4,4'-bis(1-propynyl)biphenyls by diastereomeric encapsulation through the asymmetric suppression of rotation around the axis of the prochiral biphenyl moiety.

Introduction

Supramolecular capsules constructed by self-assembly of preorganized modular subunits provide an isolated nanospace.^[1] Guest molecules confined in an isolated nanospace often show unique properties that are not observed in their free forms. Encapsulation of guest molecules in the capsules can be used for various applications such as separation techniques, capture and containment of hazardous chemicals, stabilization of reactive intermediates, catalysts, and sensing techniques.^[1,2] Self-

assembled capsules based on metal-coordination bonds have been extensively studied. $^{\left[^{3-6\right] }}$

Self-sorting has been defined as high-fidelity recognition of self from nonself within complex mixtures.^[7] Self-sorting during self-assembled capsule formation through selection between homomeric assembly (narcissistic self-sorting or self-recognition) and heteromeric assembly (social self-sorting or selfdiscrimination) is an interesting topic in supramolecular chemistry, with a view to mimicking biological processes.^[7-12] In almost all cases, selective formation of homomeric- or heteromericassembled capsules has depended on the structural demands of the modular subunits. However, control of selective formation between homomeric- and heteromeric-assembled capsules depending on guest encapsulation is almost unprecedented.[8a,b,9a]

As another topic, encapsulation strategies that use covalently bound and self-assembled capsules enable effective protection of reactive compounds against chemical or photochemical reactions.[13-16] trans-Azobenzene is а well-known photoresponsive molecule that undergoes trans-to-cis and cis-totrans photoisomerizations upon UV and visible light irradiation, respectively. Upon UV light irradiation, the encapsulated transazobenzene is usually released from self-assembled capsules because of the trans-to-cis photoisomerization;[17] indeed, the preservation of trans-azobenzene encapsulated in selfassembled capsules upon UV light irradiation is without precedent.[16e]

Chiral capsules are another interesting topic in supramolecular nanospace chemistry. Chiral molecular recognition of a racemic guest upon encapsulation in chiral capsules has been studied extensively.^[11a,18,19] In a type of twisted

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capsule composed of north and south hemispheres, it is known that the equilibrium between (*P*)- and (*M*)-twistomers can be controlled either by the introduction of a chiral group onto a racemic self-assembled capsule^[19b] or by the encapsulation of a chiral guest into a racemic self-assembled capsule.^[6f,g,20] Chiral induction of a prochiral guest upon encapsulation in a chiral capsule would also be important for supramolecular nanospace chemistry^[21] because such an encapsulation design is related to asymmetric capsular catalysts^[22] and to the emergence of novel stereoisomerisms.^[23]

Calix[4]resorcinarene-based tetrafunctionalized cavitands possess a bowl-shaped electron-rich aromatic cavity. They have been widely used as scaffolds for covalently bound and selfassembled capsules.^[1c] Previously, we reported that a 2:4 mixture of tetra(4-pyridyl)cavitand (3) and Pd(dppp)(OTf)₂ in CDCl₃ gives a complex mixture, whereas the components of a 2:4 mixture of tetrakis(4-pyridylethynyl)cavitand (4) and Pd(dppp)(OTf)₂ selfassemble into Pd-pyridyl (Pd-Npy) coordination homocapsule $\{\mathbf{4}_2 \ [Pd(dppp)]_4\}^{8+} \ (TfO^-)_8 \ (\mathbf{C4}) \ (Chart 1)^{[6c,d]} \ However, we have$ not observed quest encapsulation of C4 to date. Therefore, the characteristics of cavitand-based Pd-Npv coordination capsules upon guest encapsulation have remained unclear. The present work is concerned with a combination of expanded tetra(4derivatives. pyridyl)cavitand i.e., tetrakis[4-(4-**(1)**^[24] pyridyl)phenyl]cavitand tetrakis[4-(4or pyridyl)phenylethynyl]cavitand (2), and Pd(dppp)(OTf)₂ (Chart 1). Our attention has focused on what functionalities (features) are exhibited by Pd-Npy coordination capsules that are composed of cavitand 1 or 2. Here, we conducted a comprehensive study of cavitand-based Pd-Npy coordination capsules. We report (1) the formation of homocapsules $\{\mathbf{1}_2 \cdot [Pd(dppp)]_4\}^{8+} \cdot (TfO^-)_8$ (C1) and $\{\mathbf{2}_2 \cdot [\mathsf{Pd}(\mathsf{dppp})]_4\}^{8+} \cdot (\mathsf{TfO}^-)_8$ (C2) and heterocapsule а $\{1\cdot 2\cdot [Pd(dppp)]_4\}^{8+} \cdot (TfO^{-})_8$ (C3), (2) their guest-encapsulation ability, and (3) the control of self-sorting on homo- or heterocapsule selection by guest-induced encapsulation. We also describe applications of Pd-Npy coordination capsules with the use of 1 such as (4) usability of capsule C1 as a guard nanocontainer for a photoresponsive guest and (5) the formation of a chiral capsule $\{\mathbf{1}_2 \cdot [Pd((R)-BINAP)]_4\}^{8+} \cdot (TfO^-)_8$ (C5) with the use of Pd((R)-BINAP)(OTf)₂ in place of Pd(dppp)(OTf)₂, wherein C5 induces supramolecular chirality with respect to prochiral biphenyl guests by diastereomeric encapsulation.



Chart 1. Structures of cavitands 1-4.

Results and Discussion

Formation of homocapsules C1 and C2

Pd(dppp)(OTf)₂ has limited solubility in CDCl₃; however, a 2:4 mixture of tetrakis[4-(4-pyridyl)phenyl]cavitand (1, 2 mM) and

Pd(dppp)(OTf)₂ in CDCl₃ becomes soluble at 25 °C after heating the mixture at 50 °C for 1 h. The ¹H NMR spectrum of the mixture showed the formation of a highly symmetrical single species, indicative of a homocapsule $\{1_2 \cdot [Pd(dppp)]_4\}^{8+} \cdot (TfO^-)_8$ (C1) through Pd-Npy coordination bonds (Scheme 1a), and the complete disappearance of the signals of free 1 and free Pd(dppp)(OTf)₂ (Figure 1c vs. Figures 1a, b and Figure S3). The signal assignments of C1 were supported by the ¹H-¹H COSY spectrum (Figure S4). The signals of the pyridyl α -proton and the outer proton of the methylene-bridge rim (O-CH_{in}H_{out}-O) of C1 were shifted downfield by 0.40 and 0.10 ppm, respectively, and the signals of the pyridyl β -proton and the inner proton of the methylene-bridge rim of C1 were shifted upfield by 0.30 and 0.12 ppm, respectively, relative to those of free 1. The large downfield shift of the pyridyl α -proton of the subunit **1** ($\Delta \delta$ = 0.40 ppm) indicates the formation of the Pd-Npy coordination bond.[6c,d] Further evidence for the formation of C1 was provided by the ESI-TOF-MS spectrum and guest encapsulation experiments (vide infra).

A 2:4 mixture of tetrakis[4-(4-pyridyl)phenylethynyl]cavitand (2; 2 mM) and Pd(dppp)(OTf)₂ in CDCl₃ also becomes soluble at 25 °C after heating the mixture at 50 °C for 1 h. The ¹H NMR spectrum of the mixture showed the formation of a highly symmetrical single species, indicative of a homocapsule {2₂·[Pd(dppp)]₄}⁸⁺.(TfO⁻)₈ (C2) through Pd–Npy coordination bonds (Figure 1e vs. Figure 1d, Scheme 1b, and Figures S5 and S6). The signal of the pyridyl α -proton of C2 was shifted downfield by 0.39 ppm, and the signals of the pyridyl β -proton and the inner and outer protons of the methylene-bridge rim of C2 were shifted upfield by 0.25, 0.11, and 0.06 ppm, respectively, relative to those of free 2.^[6c,d] Further evidence for the formation of C2 was provided by the ESI-TOF-MS spectrum (vide infra).



Figure 1. Association behavior of cavitands 1 or 2 with $Pd(dppp)(OTf)_2$ monitored by ¹H NMR (400 MHz, CDCl₃, 298 K): (a) $Pd(dppp)(OTf)_2$ alone, (b) 1 alone (2 mM), (c) capsule C1 (after heating a mixture of 1 (2 mM) and $Pd(dppp)(OTf)_2$ (4 mM) at 50 °C for 1 h), (d) 2 alone (2 mM), and (e) capsule C2 (after heating a mixture of 2 (2 mM) and $Pd(dppp)(OTf)_2$ (4 mM) at 50 °C for 1 h). The signals marked "a–e" are assigned in Scheme 1. The representative signals of free species and complexes are shown in black and red, respectively.

Guest encapsulation in homocapsules C1 and C2

Capsule **C1** encapsulates one guest molecule such as bis(4acetoxyphenyl)acetylene (**G1**) or 4,4'-bis(1-propynyl)biphenyl (**G2**) to form guest-encapsulating capsule **G@C1** (Scheme 1a).

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Scheme 1. (a) Self-assembly of a 2:4 mixture of cavitand 1 and Pd(dppp)(OTf)₂ into homocapsule C1 and its guest encapsulation; (b) self-assembly of a 2:4 mixture of cavitand 2 and Pd(dppp)(OTf)₂ into homocapsule C2.



Figure 2. Association behavior of capsule C1 or C2 with guests G1~G3 monitored by ¹H NMR (400 MHz, CDCI₃, 298 K): (a) C1 alone (1 mM), (b) G1@C1 ([C1] = 1 mM + [G1] = 10 mM), (c) G1 alone, (d) G2@C1 ([C1] = 1 mM + [G2] = 2 mM), (e) G2 alone, (f) [C1] = 1 mM + [G3] = 6 mM, and (g) G3 alone. The signals marked "a–e" and "x–z" are assigned in Scheme 1. The

representative signals of free species and complexes are shown in black and red, respectively.

The ¹H NMR spectrum of a 1:10 mixture of C1 (1 mM; [1] = 2 mM and [Pd(dppp)(OTf)₂] = 4 mM) and G1 (10 mM) in CDCl₃ at 298 K showed formation of G1@C1 (Figures 2b and S7). The ¹H NMR signals of G1@C1, free C1, and free G1 were independently observed. This result indicates that one molecule of G1 is encapsulated in C1 and that the exchange of G1 in and out of C1 is slow on the NMR time-scale. The ¹H NMR chemical shift changes of the signals of the encapsulated G1 relative to those of free **G1** ($\Delta \delta = \delta_{\text{encapsulated-guest}} - \delta_{\text{free-guest}}$) were -3.88 ppm for the protons of the acetoxy groups (signal-x, OAc) at the parapositions and -0.87 ppm for the aromatic *meta*-proton (signal-y). The very large upfield shift of the acetoxy groups of the encapsulated G1 is due to the shielding effect of the aromatic cavity ends of C1. This result indicates that the acetoxy groups of G1 are oriented toward both aromatic cavity ends of C1 (Scheme 1a) to maximize CH $\cdots\pi$ interaction between the methyl moiety of the acetoxy group of G1 and the electron-rich aromatic cavity of the subunit 1. Based on the integration ratios of the three species as a function of concentration, the association constant (K_a) of C1 with **G1** was estimated to be 2.27×10^3 M⁻¹ in CDCl₃ at 298 K.

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Scheme 2. Control of self-sorting by guest encapsulation: (a) self-assembly of a 1:1:4 mixture of cavitand 1, cavitand 2, and Pd(dppp)(OTf)₂ into a mixture of homocapsules C1, C2, and heterocapsule C3; (b) upon addition of G1, thermodynamic equilibration shift of a mixture of C1, C2, and C3 into a G1@C1-enriched and C2-enriched mixture; (c) upon addition of G3, thermodynamic equilibration shift of a mixture of C1, C2, and C3 into a G3@C3-enriched mixture.



Figure 3. ¹H NMR spectra (400 MHz, CDCl₃, 298 K) of (a) homocapsule C1 alone (1 mM), (b) a mixture of C1, C2, and heterocapsule C3 (after heating a mixture of 1 (1 mM), 2 (1 mM), and Pd(dppp)(OTf)₂ (4 mM) at 50 °C for 1 h), and (c) homocapsule C2 alone (1 mM). The representative signals of C1, C2, and C3 are marked with blue, red, and purple open circles, respectively. See also Scheme 2a.

The ¹H NMR spectrum of a 1:2 mixture of **C1** (1 mM) and **G2** (2 mM) in CDCl₃ at 298 K showed formation of **G2@C1** (Figures 2d and S8). The ¹H NMR signals of **G2@C1**, free **C1**, and free **G2** were independently observed. The $\Delta\delta$ value for the protons of the 1-propynyl groups (signal-x, CCMe) at the *para*-positions was – 3.52 ppm. This result indicates that the 1-propynyl groups of **G2** are oriented toward both aromatic cavity ends of **C1**. The K_a value of **C1** with **G2** was estimated to be 9.53×10^3 M⁻¹ in CDCl₃ at 298 K.

Capsule **C1** is sensitive to the guest molecular length for guest encapsulation, and does not encapsulate molecules such as 4,4''diacetoxy-*p*-terphenyl (**G3**), which do not fit the cavity size of **C1** (Figures 2f and S9). The cavity size of **C2** is larger than that of **C1**. Capsule **C2** does not encapsulate **G1**, **G2**, or **G3** because the molecular sizes of these guests are small relative to the cavity size of **C2** (Figure S9). At this stage, we have not been able to observe guest encapsulation of **C2**.^[25]

Self-sorting in capsule formation: homomeric vs. heteromeric capsule

Self-sorting through selection between homomeric assembly and heteromeric assembly is an interesting topic. However, control of selective formation between homomeric and heteromeric assembled capsules depending on guest encapsulation is almost unprecedented.^[8a,b,9a]

A 1:1:4 mixture of **1** (1 mM), **2**, and Pd(dppp)(OTf)₂ in CDCl₃ becomes soluble at 25 °C after heating the mixture at 50 °C for 1 h. The ¹H NMR spectrum of the mixture showed the expected signals for homocapsules **C1** and **C2**, and signals for the formation of a new species also appeared at 4.33 ppm (independent), 4.58 ppm partially overlapped with the signal of the inner proton (H_{in}) of the methylene-bridge rim (O-CH_{in}H_{out}-O) of **C2**, and 5.46 ppm partially overlapped with the signal of the outer proton (H_{out}) of the methylene-bridge rim of **C1** (Figures 3b and S10), although other signals of **c1** or **C2**. Based on comparisons of chemical shifts and integration ratios of the ¹H NMR signals of the new species with those of **C1** and **C2**, it is suggested that the

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Figure 5. Association behavior of a mixture of C1, C2, and C3 with G1 monitored by ¹H NMR (400 MHz, $CDCl_3$, 298 K): (a) G1 alone; (b) a mixture of C1, C2, and C3 after heating a mixture of 1 (1 mM), 2 (1 mM), and $Pd(dppp)(OTf)_2$ (4 mM) at 50 °C for 1 h; after heating a mixture of C1, C2, C3, and G1 (6–18 mM) at 50 °C for 1 h: [G1] = (c) 6 mM, (d) 12 mM, and (e) 18 mM. The representative signals of G1@C1, C1, C2, and C3 are marked with blue solid circle, and blue, red, and purple open circles, respectively. The signals "x and y" of encapsulated G1 and those of free G1 are shown in red and black, respectively. See also Scheme 2b.



Figure 4. ESI-TOF-MS spectrum of a 1:1:4 mixture of **1** (0.01 mM), **2**, and Pd(dppp)(OTf)₂ (a mixture of capsules **C1**, **C2**, and **C3**) in CHCl₃ (capillary = -3500 V and ion source temperature = 34 °C): (a) top: observed spectrum of [M $- 4(\text{TfO}^-)$]⁴⁺ (M = **C1**, **C2**, and **C3**), bottom: calculated spectrum of [**C3** $- 4(\text{TfO}^-)$]⁴⁺; (b) top: observed spectrum of [**M** $- 3(\text{TfO}^-)$]³⁺ (M = **C1**, **C2**, and **C3**), bottom: calculated spectrum of **C3**, bottom: calculated spectrum of [**C3** $- 3(\text{TfO}^-)$]³⁺. See also Figure S12.

new species is a hetero-capsule $\{1 \cdot 2 \cdot [Pd(dppp)]_4\}^{8*} \cdot (TfO^-)_8$ (C3) composed of 1, 2, and $Pd(dppp)(OTf)_2$ in a 1:1:4 mole ratio (Scheme 2a). The three aforementioned signals of C3 were assigned to the H_{in} of the subunit-1, the H_{in} of the subunit-2, and the H_{out} of subunit-1, the detail of which is shown in Figure 7a (vide infra). Based on the signal integration ratios of the H_{in} of the methylene-bridge rims of C1, C2, and C3, the product ratio of C1, C2, and C3 was estimated to be 1:1:0.98 (Figure 3b). This product ratio remained unchanged after 24 h at room temperature and also after additional heating at 50 °C for 12 h (Figure S11), indicating that the thermodynamic equilibration between C1, C2, and C3 was reached after initial heating the mixture at 50 °C for

1 h (Figure 3b). The product ratio of homocapsules C1 and C2 is inevitably 1:1, when 1, 2, and Pd(dppp)(OTf)₂ are mixed in a 1:1:4 mole ratio. The formation of C1 and C2 was preferred, because C1 is thermodynamically most stable among C1, C2, and C3. Concrete evidence for the formation of C3 was provided by the ESI-TOF-MS spectrum of the mixture (Figures 4 and S12), wherein the molecular ion peaks of C3 were observed at *m*/z 2149.7278 [C3 – 3(TfO⁻)]³⁺ (calcd 2149.7085) and 1575.0162 [C3 – 4(TfO⁻)]⁴⁺ (calcd 1575.0433), in addition to those of C1 at *m*/z 2117.7201 [C1 – 3(TfO⁻)]³⁺ (calcd 2117.7084) and 1551.0265 [C1 – 4(TfO⁻)]⁴⁺ (calcd 1591.0432) and those of C2 at *m*/z 2181.4004 [C2 – 3(TfO⁻)]³⁺ (calcd 2181.7086) and 1599.0076 [C2 – 4(TfO⁻)]⁴⁺ (calcd 1599.0433). Further evidence for the formation of C3 was supported by guest-encapsulation experiments (vide infra).

As mentioned above, guest G1 is encapsulated in homocapsule C1, but not C2. Figure 5 shows the ¹H NMR spectra of the mixture upon addition of G1 to the equilibrated mixture of C1, C2, and C3 (1:1:0.98) derived from a 1:1:4 mixture of 1 (1 mM), 2, and Pd(dppp)(OTf)₂ in CDCl₃ as described above. Upon addition of G1 (6 mM), the signals of guest-encapsulating G1@C1 and free C2 increased with a decrease of those of free C1 and with complete disappearance of those of C3 (Figure 5c). The product ratio of G1@C1, C1, C2, and C3 was shifted to 41:9:50:0, wherein a thermodynamic equilibration was shifted most favorably toward formation of G1@C1 through G1-induced stabilization of C1 by encapsulation (Scheme 2b). Upon further addition of G1 (18 mM in total), the product ratio of G1@C1, C1, C2, and C3 changed to 45:5:50:0 (Figures 5e and S13). Thus, selective formation of homocapsules C1 and C2 was achieved upon addition of G1. Guest G2 also induced selective formation of the homocapsules. Upon addition of G2 (18 mM) to the mixture of C1, C2, and C3 (1:1:0.98) described above, the product ratio of G2@C1, C1, C2, and C3 was shifted to 50:0:50:0 (Figure S14).

As noted above, homocapsules C1 and C2 do not encapsulate G3 because the cavity sizes of C1 and C2 are too small or large, respectively, relative to the molecular size of G3. The cavity size of heterocapsule C3 is between those of

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Figure 6. Association behavior of a mixture of C1, C2, and C3 with G3 monitored by ¹H NMR (400 MHz, CDCl₃, 298 K): (a) a mixture of C1, C2, and C3 after heating a mixture of 1 (1 mM), 2 (1 mM), and Pd(dppp)(OTf)₂ (4 mM) at 50 °C for 1 h; after heating a mixture of C1, C2, C3, and G3 (5–50 mM) at 50 °C for 1 h: [G3] = (b) 6 mM, (c) 15 mM, (d) 20 mM, (e) 30 mM, (f) 40 mM, and (g) 50 mM; and (h) G3 alone. The representative signals of G3@C3, C1, C2, and C3 are marked with purple solid circle, and blue, red, and purple open circles, respectively. The signals "x–z" and "x'–z" shown in red indicate the encapsulated G3 oriented to the subunits-1 and -2 of C3, respectively. See also Scheme 2c.



Figure 7. Expanded spectra of (a) Figure 6a, (b) Figure 6c, (c) Figure 6e, and (d) Figure 6g in the region of the inner (H_{in}) and outer (H_{out}) protons of the methylenebridge rims (O-CH_{in}H_{out}-O) and the basal CH of side chain R of capsules. The signals marked "a" and "b" indicate **C1** and **C2**, respectively. The signals marked "a" and "b" indicate the subunits-**1** and -**2** of **C3**, respectively. The signals marked "a" and "b" indicate the subunits-**1** and -**2** of **C3**, respectively.



Figure 8. Plots of the product ratio of **G3@C3**, **C1**, **C2**, and **C3** as a function of **[G3]**, based on the ¹H NMR data of Figure 6.

homocapsules C1 and C2. As a result, it was found that heterocapsule C3 selectively encapsulates G3 to form G3@C3 (Scheme 2c). Figure 6 shows the ¹H NMR spectra of the mixture upon addition of G3 to the equilibrium mixture of C1, C2, and C3 (1:1:0.98) derived from a 1:1:4 mixture of 1 (1 mM), 2, and Pd(dppp)(OTf)₂ in CDCl₃, as described above. Figure 7 shows the expanded spectra of Figure 6 in the region of the inner (Hin) and outer (Hout) protons of the methylene-bridge rims (O-CHinHout-O) of capsules. The signals of C1, C2, and C3 were assigned as shown in Figures 7a and 6a. The signals-a indicate the Hin at 4.25 ppm and the Hout at 5.45 ppm (partially overlap) of homocapsule C1, and the signals-b indicate the H_{in} at 4.56 ppm (partially overlap) and the H_{out} at 6.02 ppm (overlap) of homocapsule C2. For heterocapsule C3, the signals-a' indicate the H_{in} at 4.33 ppm and the Hout at 5.46 ppm (partial overlap) of the subunit-1, and the signals-b' indicate the Hin at 4.58 ppm (partial overlap) and the

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Figure 9. ¹H NMR spectra (400 MHz, CDCl₃, 298 K) of (a) C1 alone (1 mM), (b) *trans*-G4@C1 ([C1] = 1 mM + [*trans*-G4] = 3 mM), (c) *trans*-G4 alone (3 mM), (d) after 350 nm irradiation of a mixture of C1 (1 mM) and *trans*-G4 (3 mM) for 60 min, and (e) after 350 nm irradiation of *trans*-G4 (3 mM) for 60 min. The signals "x-z" shown in red, purple, and blue indicate *trans*-G4 encapsulated in C1, free *trans*-G4, and free *cis*-G4, respectively, which are assigned in Scheme 3.



Scheme 3. Almost no photoisomerization of trans-G4 encapsulated in C1.

Hout at 6.02 ppm (overlap) of the subunit-2. For guestencapsulating heterocapsule G3@C3, the signals-a" indicate the H_{in} at 4.35 ppm (overlap) and the H_{out} at 5.41 ppm of the subunit-1, and the signals-b" indicate the H_{in} at 4.62 ppm (partial overlap) and the Hout at 6.04 ppm (overlap) of subunit-2, which were assigned based on 2D NOESY experiments (Figure S15) on the sample (Figures 7d and 6g). Upon increasing the amount of G3, new signals indicating G3@C3 increased with decreases of signals for free C1, free C2, and free C3 (Figures 6 and 7), wherein a thermodynamic equilibration was shifted most favorably toward formation of G3@C3 through G3-induced stabilization of C3 by encapsulation. Upon further addition of G3 (50 mM in total), the product ratio of C1, C2, C3, and G3@C3 reached 12:12:8:68 (Figures 6g and 7d). Figure 8 shows the plots of the product ratio of G3@C3, C1, C2, and C3 as a function of [G3].

As the electronic environment of the subunit-1 differs from that of the subunit-2 in heterocapsule C3, the G3 encapsulated in C3 is desymmetrized. Thus, the ¹H NMR signals of G3 encapsulated in C3 appeared as two sets of signals (Figure 6). Representative ¹H NMR signals of G3 encapsulated in C3 appeared at –1.48 ppm ($\Delta \delta = -3.83$ ppm) and –1.92 ppm ($\Delta \delta = -4.27$ ppm) for the acetoxy protons (signal-OAc, x and x') at the *para*-positions and at 6.58

ppm ($\Delta\delta$ = -0.61 ppm) and 6.28 ppm ($\Delta\delta$ = -0.91 ppm) for the aromatic meta-protons (signals-y' and y). In other words, this G3encapsulation behavior in a capsule provides definitive evidence for the formation of heterocapsule C3. The very large upfield shift of the acetoxy protons of the encapsulated G3 indicates that the acetoxy groups of G3 are oriented toward both aromatic cavity ends of C3. The 2D NOESY spectrum (Figure S15b) of the sample (Figures 6g and 7d) disclosed the orientation of G3 encapsulated in C3. The acetoxy proton signal-x of -1.48 ppm ($\Delta\delta$ = -3.83 ppm) showed the NOE correlations to the signals-a" of H_{in} and H_{out} of the subunit-1; i.e., the acetoxy group-x is oriented to the subunit-1. On the other hand, the more upfield-shifted acetoxy proton signal-x' of -1.92 ppm ($\Delta \delta = -4.27$ ppm) showed the NOE correlations to the signals-b" of H_{in} and H_{out} of the subunit-2; i.e., the acetoxy group-x' is oriented to the subunit-2. The NOE correlations were also observed between the aromatic proton signal-y' of G3 and the signals-b" of H_{in} of the subunit-2, and between the aromatic proton signal-y' and the aromatic proton signal-z' of G3.

The G3-encapsulation in the equilibrium mixture of C1, C2, and C3 was investigated in more detail. The ¹H NMR spectrum just after addition of G3 (40 mM) at room temperature to the equilibrium mixture of C1, C2, and C3 (33.6:33.6:32.8) derived from a 1:1:4 mixture of 1 (1 mM), 2, and Pd(dppp)(OTf)₂, showed the product ratio of C1/C2/C3/G3@C3 = 26:26:4:44 (Figure S16b). The ¹H NMR spectrum of this mixture after 24 h at 25 °C indicated a change in the product ratio of C1/C2/C3/G3@C3 to 13:13:11:63 (Figure S16c). However, the product ratio remained unchanged after additional heating of this mixture at 50 °C for 12 h (Figure S16d). These results suggest that the encapsulation of G3 in C3 is relatively fast, and the equilibrium shift of C1 and C2 toward C3 is relatively slow. This product ratio was the same as that of the sample after heating the initial mixture at 50 °C for 1 h (Figure 6f).

Capsule C1 as a guard nanocontainer for *trans*-4,4'diacetoxyazobenzene

In the ¹H NMR study (Figures 9b and S17), capsule **C1** also encapsulates *trans*-4,4'-diacetoxyazobenzene (*trans*-**G4**) to form

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Scheme 4. Self-assembly of a 2:4 mixture of cavitand 1 and $Pd((R)-BINAP)(OTf)_2$ into chiral capsule C5.



Figure 10. Association behavior of cavitand 1 with $Pd((R)-BINAP)(OTf)_2$ monitored by ¹H NMR (400 MHz, CDCI₃, 298 K): (a) $Pd((R)-BINAP)(OTf)_2$ alone, (b) chiral capsule **C5** (after heating a mixture of **1** (2 mM) and $Pd((R)-BINAP)(OTf)_2$ (4 mM) at 50 °C for 1 h), and (c) **1** alone (2 mM). The signals marked "a–g" are assigned in Scheme 4. The representative signals of free species and **C5** are shown in black and red, respectively.

trans-G4@C1 with $\Delta \delta$ = -3.80 ppm for the acetoxy proton signal (δ = -1.44 ppm) and K_a = 14.0 × 10³ M⁻¹ in CDCl₃ at 298 K.

Irradiation of trans-G4 (3 mM) in CDCI3 with 350 nm light[26] resulted in the photoisomerization to cis-G4, wherein the ¹H NMR signals of trans-G4 (δ = 7.96, 7.26, and 2.36 ppm) decreased with increasing signal intensity for cis-G4 (δ = 7.04, 6.91, and 2.29 ppm) (Figure 9c vs. Figures 9e and S18). The photostationary state was reached within 60 min (trans-G4/cis-G4 = 2:98). In marked contrast, the trans-G4 encapsulated in C1, derived from a 1:3 mixture of C1 (1 mM) and trans-G4 (3 mM) in CDCl₃, underwent almost no photoisomerization to cis-G4 (Figures 9d and S19). Upon 350 nm irradiation for 60 min, the encapsulation ratio of trans-G4 encapsulated in C1 remained almost unchanged (only 6% reduction of encapsulation), although excess amount of trans-G4, which was not encapsulated in C1, photoisomerized to cis-G4 (Scheme 3). The acetoxy groups of trans-G4 encapsulated in C1 are oriented toward both aromatic cavity ends of C1 ($\Delta \delta$ = -3.80 ppm), leading to tight encapsulation of trans-G4. As a result, confined space of trans-G4@C1 would interfere with geometrical change in the transition state of trans-to-cis photoisomerization of trans-G4. Thus, capsule C1 serves as a guard nanocontainer for trans-G4 to protect against the trans-to-cis photoisomerization of *trans*-**G4** by encapsulation in **C1**. Furthermore, this protection may be explained by the very slow exchange of *trans*-**G4** in and out of **C1**, in which the exchange cross-peak between the encapsulated and free *trans*-**G4** in the 2D NOESY spectrum was not observed even at 323 K (Figure S20), and/or overlap of UV– vis absorption bands between **C1** and *trans*-**G4** at 350 nm (Figure S21).



Figure 11. Schematic illustration of structural interconversion between (a) capsule **C5'** ($R = CH_2CH(CH_3)_2$) in solution and (b) an infinite 2D coordination-bonded network sheet structure **S5'** in the crystalline state.

Formation of chiral capsule C5

We found the formation of a chiral capsule $\{1_2 \cdot [Pd((R)-BINAP)]_4\}^{8+} \cdot (TfO^-)_8$ (C5) with the use of Pd((R)-BINAP)(OTf)_2 in place of Pd(dppp)(OTf)_2.^[27]

A 2:4 mixture of 1 (R = $(CH_2)_{10}CH_3$) (2 mM) and Pd((R)-BINAP)(OTf)₂ in CDCl₃ became soluble at 25 °C after heating the mixture at 50 °C for 1 h. The ¹H NMR spectrum of the mixture showed the formation of a highly symmetrical single species, indicative of a chiral capsule $\{\mathbf{1}_2 \cdot [Pd((R)-BINAP)]_4\}^{8+} \cdot (TfO^-)_8$ (C5) through Pd-Npy coordination bonds (Scheme 4), and the complete disappearance of the signals of free 1 and free Pd((R)-BINAP)(OTf)₂ (Figure 10b vs. Figure 10a, c). The signal assignments of **C5** were supported by the ¹H–¹H COSY spectrum (Figure S22). The signals of the pyridyl α -proton and the outer proton of the methylene-bridge rim (O-CH_{in}H_{out}-O) of C5 were shifted downfield by 0.04 and 0.11 ppm, respectively, and the signals of the pyridyl β -proton and the inner proton of the methylene-bridge rim of C5 were shifted upfield by 0.47 and 0.15 ppm, respectively, relative to those of free 1. The very small downfield shift of the pyridyl a-proton and the relatively large upfield shift of the pyridyl β -proton of the subunit 1 of C5, compared with those of C1, suggest the shielding effect of the subunit (R)-BINAP. Further evidence for the formation of C5 was supported by guest encapsulation experiments (vide infra).

Single crystals of the components of a 2:4 mixture of **1'** (R = CH₂CH(CH₃)₂) and Pd((*R*)-BINAP)(OTf)₂, suitable for X-ray diffraction analysis, were obtained by slow diffusion of EtOH into a CHCl₃ solution of {**1'**₂·[Pd((*R*)-BINAP)]₄}⁸⁺·(TfO⁻)₈ (**C5'**). However, surprisingly, the crystal structure was not the discrete capsule **C5'**, but was found to be an infinite two-dimensional Pd–Npy coordination-bonded porous network sheet structure

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Figure 12. X-ray crystal structure of S5': (a) one unit structure, (b) 2D coordination-bonded network sheet structure, (c) 3D packing structure (assembly of four sheets), and (d) its space filling representation. In Figure 12a, triflate ions are omitted for clarity. In Figures 12b–d, triflate ions and hydrogen atoms are omitted for clarity.



Scheme 5. Schematic illustration of diastereomeric encapsulation of prochiral biphenyl guests G5~G9 in chiral capsule C5.

 $[{\mathbf{1'} [Pd((R)-BINAP)]_2}^{4+} (TfO^{-})_4]_n$ (**S5'**) (Figure 11). Figure 12 shows the one-unit structure, 2D network sheet structure, and 3D packing structure of S5' in the crystal structure.^[28] Crystal data and structural refinement of S5' are listed in Table S1, and an ORTEP view is shown in Figure S23A. Each 1' is placed in the ab plane in a parallel fashion along the c axis, and is connected with Pd((R)-BINAP) by the Pd–Npy coordination bonds to form 2D coordination-bonded porous network sheet S5', with square cavities of dimension ca. 8.4 Å × 8.4 Å including van der Waals radii (Figure 12a, b). The bond distances of Pd-N and Pd-P are 2.111 Å and 2.280 Å, respectively, and the bond angles of N-Pd-N, P-Pd-P, and N-Pd-P are 84.601°, 90.814°, and 92.907°, respectively. The adjacent 2D coordination-bonded porous network sheets of S5', which are separated by 16.83 Å (1/2c of the unit cell) and are translated by 9.89 Å along the a axis (1/2a of the unit cell) and 9.89 Å along the b axis (1/2b of the unit cell), are layered in an offset manner (ABAB pattern) to give a 3D packing structure with chambers (but no channels) (Figure 12c, d). The ¹H NMR spectrum of a solution upon dissolving the single crystals of S5' in CD₂Cl₂ showed that single crystals of S5' include CHCl₃ and EtOH as cocrystal solvents in a ratio of $\{1' \cdot [Pd((R) - R)]\}$ $BINAP_{2}^{4+}(TfO^{-})_{4}/CHCl_{3}/EtOH = 1:1:2$ (Figure S23B). Interestingly, the ¹H NMR spectrum of this solution indicated reproduction of the capsule structure C5'. These results suggest that the capsule C5' is kinetically as well as thermodynamically stable in solution, but the 2D coordination-bonded porous network sheet S5' is thermodynamically stable in the crystalline state (Figure 11). This structural interconversion between C5' in solution and S5' in the crystalline state occurs through reversible Pd-Npy coordination bonds and probably because the crystal packing force of S5' is thermodynamically more favorable than that of C5'.

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Figure 13. Association behavior of chiral capsule C5 with guest G2 or G5~G9 monitored by ¹H NMR (400 MHz, CDCl₃, 298 K): (a) C5 alone (1 mM), (b) G2@C5 ([C5] = 1 mM + [G2] = 10 mM), (c) G5@C5 ([C5] = 1 mM + [G5] = 10 mM), (d) G6@C5 ([C5] = 1 mM + [G6] = 10 mM), (e) G7@C5 ([C5] = 1 mM + [G7] = 18 mM), (f) G8@C5 ([C5] = 1 mM + [G8] = 10 mM), and (g) G9@C5 ([C5] = 1 mM + [G9] = 18 mM). The representative signals of encapsulated and free guests are marked with solid and open circles, respectively. The signals of the propynyl-CH₃ groups, the terminal CH₃ groups of the CO₂(CH₂)_nCH₃ groups (G6~G9; n = 1~4) are marked with red, blue, and green circles, respectively. See also Scheme 5.

Chiral induction of prochiral biphenyl guest upon encapsulation in chiral capsule C5

Chiral induction of a prochiral guest upon encapsulation in a chiral capsule would be important for supramolecular nanospace chemistry^[21] because such an encapsulation design is related to asymmetric capsular catalysts^[22] and to the emergence of novel stereoisomerisms.^[23] Chiral capsule {**1**₂·[Pd((*R*)-BINAP)]₄}⁸⁺ (TfO⁻)₈ (C5) encapsulates one guest molecule such as 4,4'-bis(1-propynyl)biphenyl (G2) and prochiral 2,2'bis(alkoxycarbonyl)-4,4'-bis(1-propynyl)biphenyls (G5~G9) (Scheme 5). We found that chiral capsule C5 induces supramolecular chirality with respect to prochiral G5~G9 upon encapsulation. Guest-encapsulation behaviors of C5 were investigated by ¹H NMR spectroscopy in CDCI₃ at 298 K (Figure 13).

The K_a value of **C5** with **G2** was estimated to be $1.64 \times 10^3 \text{ M}^{-1}$, and the $\Delta\delta$ value for the protons of the 1-propynyl groups of **G2** was -3.73 ppm (Figures 13b and S24). Compared with $K_a = 9.53 \times 10^3 \text{ M}^{-1}$ and $\Delta\delta = -3.52$ ppm for **G2@C1** (Figures 2d and S8), this result indicates that the cavity length of **C5** composed of **1** with Pd((*R*)-BINAP) is somewhat shorter than that of **C1** composed of **1** with Pd(dppp).

The K_a value of **C5** with 2,2'-bis(methoxycarbonyl)-4,4'-bis(1propynyl)biphenyl (**G5**) (R' = CH₃; n = 0) was estimated to be 1.32 × 10³ M⁻¹, and the $\Delta\delta$ value for the 1-propynyl group of **G5** was – 3.65 ppm (Figures 13c and S25). Compared with $K_a = 3.32 \times 10^3$ M⁻¹ and $\Delta\delta = -3.53$ ppm for **G5@C1** (Figure S26), this result also supports the conclusion that the cavity length of **C5** is somewhat shorter than that of **C1**. The $\Delta\delta$ values for the CO₂CH₃ groups of **G5** were –1.08 and –1.11 ppm for **G5@C5** and –1.10 ppm for **G5@C1**, the values of which were much smaller than the $\Delta\delta$ values for the 1-propynyl groups of **G5@C5** and **G5@C1**. These results indicate that the 1-propynyl groups at the 4,4'-positions of **G5** are oriented toward both aromatic cavity ends of **C5** and **C1**, and the CO_2CH_3 groups at the 2,2'-positions of **G5** are directed to two of the four equatorial windows of **C5** and **C1** (Scheme 5).

The ¹H NMR signal of the CO₂CH₃ group of **G5** encapsulated in chiral capsule C5 notably appeared as two sets of singlets at 2.54 and 2.51 ppm ($\Delta \delta$ = -1.08 and -1.11 ppm) (Figures 13c and S25f), whereas the signal of the CO₂CH₃ group of G5 encapsulated in achiral capsule C1 appeared as one singlet at 2.52 ppm ($\Delta \delta$ = -1.10 ppm) (Figure S26f). This result indicates that chiral induction of prochiral G5 occurred upon encapsulation in the chiral capsule C5 through asymmetric suppression of rotation around the axis of the prochiral biphenyl moiety in G5; that is, diastereomeric complexes (R)-G5@C5 and (S)-G5@C5 were produced on the NMR time-scale (Scheme 5).^[21] The diastereomeric excess (d.e.) resulting from diastereomeric encapsulation selectivity of G5@C5 based on the ¹H NMR signal integration ratios of the two CO₂CH₃ signals was estimated to be 20%. At this stage, it is not easy to establish which enantiomer of (R)- or (S)-G5 is more favorably encapsulated in chiral capsule C5, and further studies are required in this regard.

The K_a values, the *d.e.* values, and the $\Delta\delta$ values of the propynyl CH₃ groups and the terminal CH₃ groups of the CO₂(CH₂)_nCH₃ groups of guests for the association of **C5** with **G5~G9** (alkyl chains R' = (CH₂)_nCH₃; n = 0~4) in CDCl₃ at 298 K are summarized in Table 1.

The K_a value of **C5** with 2,2'-bis(ethoxycarbonyl)-4,4'-bis(1propynyl)biphenyl (**G6**) (n = 1) was estimated to be 257 M⁻¹ (Figures 13d and S27). For **G6@C5**, the ¹H NMR signals of the 1-propynyl group and the CO₂CH₂CH₃ group as well as the CO₂CH₂CH₃ group split into two sets of signals at –1.57 and –1.59 ppm ($\Delta \delta = -3.66$ and –3.68 ppm), 0.031 and –0.008 ppm ($\Delta \delta = -$ 0.993 and –1.032 ppm), and 3.22 and 3.10 ppm ($\Delta \delta = -0.83$ and -0.95 ppm), respectively (Figures 13d and S27e). The diastereomeric encapsulation selectivity of **G6@C5** was estimated to be 22%.

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Table 1. The K_a values, the *d.e.* values, and the $\Delta \delta$ values of the propynyl CH₃ groups (A) and the terminal CH₃ groups (B) of the CO₂(CH₂)_nCH₃ groups of guests for the association of **C5** with **G5~G9** (alkyl chains R' = (CH₂)_nCH₃; *n* = 0~4) in CDCl₃ at 298 K.

G@C5	<i>K</i> _a (M ⁻¹)	d.e. (%)	A, $\Delta\delta$ (ppm)	B, $\Delta\delta$ (ppm)
G5 (<i>n</i> = 0)	1,320	20	-3.65	-1.08, -1.11
G6 (<i>n</i> = 1)	257	22	-3.66, -3.68	-0.99, -1.03
G7 (<i>n</i> = 2)	143	40	-3.66, -3.69	-0.92, -0.97
G8 (<i>n</i> = 3)	58	13	-3.62, -3.65	-0.92, -0.94
G9 (<i>n</i> = 4)	37	<3	-3.65, -3.66	-0.68, -0.70

The K_a value of **C5** with 2,2'-bis(*n*-propoxycarbonyl)-4,4'bis(1-propynyl)biphenyl (**G7**) (n = 2) was estimated to be 143 m⁻¹ (Figures 13e and S28). The ¹H NMR signals of the 1-propynyl group and the CO₂(CH₂)₂CH₃ group split into two sets of signals at -1.58 and -1.60 ppm ($\Delta \delta = -3.66$ and -3.69 ppm) and -0.147 and -0.200 ppm ($\Delta \delta = -0.918$ and -0.970 ppm), respectively (Figures 13e and S28h). The diastereomeric encapsulation selectivity of **G7@C5** was estimated to be 40%. The ¹H NMR spectra for the association of **C5** with **G8** (n = 3) or **G9** (n = 4) are shown in Figures 13f and S29e for **G8@C5** and Figures 13g and S30h for **G9@C5**.

As shown in Table 1, the K_a values and the $\Delta\delta$ values of the terminal CH₃ groups of the CO₂(CH₂)_nCH₃ groups of guests for the association of C5 with G5~G9 (n = 0~4) sequentially decreased with the elongation of the alkyl chains of the CO₂(CH₂)_nCH₃ groups of G5~G9. These results suggest that the alkyl chains of the CO₂(CH₂)_nCH₃ groups of G5~G9 encapsulated in C5 cannot protrude from the equatorial windows of C5, and buckling (bending) of longer alkyl chains of the CO₂(CH₂)_nCH₃ groups occurs inside the cavity of C5,[29] probably because the equatorial windows of C5 are too small or narrow to allow protrusion of longer alkyl chains of the CO₂(CH₂)_nCH₃ groups of guests. If the equatorial windows of C5 are large enough to allow protrusion of longer alkyl chains of the CO₂(CH₂)_nCH₃ groups of guests, the decrease of the K_a values would be prevented^[16d,30,31] and increase of the d.e. values might be expected with the elongation of the alkyl chains of the CO₂(CH₂)_nCH₃ groups of G5~G9.^[21] This is not the case for the association of C5 with G5~G9. The diastereomeric encapsulation selectivity for the association of C5 with G5~G9 showed maximum value (d.e. = 40%) for G7 (n = 2) among them.

Conclusion

We have demonstrated that the components of a 2:4 mixture of tetrakis[4-(4-pyridyl)phenyl]cavitand (1) or tetrakis[4-(4-pyridyl)phenylethynyl]cavitand (2) and Pd(dppp)(OTf)₂ self-assemble into homocapsules { $1_2 \cdot [Pd(dppp)]_4$ }⁸⁺ (TfO⁻)₈ (C1) and { $2_2 \cdot [Pd(dppp)]_4$ }⁸⁺ (TfO⁻)₈ (C2), respectively, through Pd–Npy coordination bonds. We comprehensively investigated cavitand-based Pd–Npy coordination capsules to disclose their properties upon guest encapsulation.

Self-sorting in the self-assembled capsule formation through selection between homomeric assembly and heteromeric

assembly is an interesting topic in supramolecular chemistry. We have demonstrated that selective formation of homo- or heterocapsule is controllable by guest-induced encapsulation. A 1:1:4 mixture of 1, 2, and Pd(dppp)(OTf)₂ in the absence of guest produced a mixture of homocapsules C1, C2, and a heterocapsule $\{1 \cdot 2 \cdot [Pd(dppp)]_4\}^{8+} \cdot (TfO^-)_8$ (C3) in a 1:1:0.98 mole ratio under thermodynamic control. Upon addition of excess bis(4acetoxyphenyl)acetylene (G1), the mixture of C1, C2, and C3 described above was thermodynamically shifted to G1@C1/C1/C2/C3 = 45:5:50:0. In contrast, upon addition of excess 4,4"-diacetoxy-p-terphenyl (G3), the above-mentioned mixture of C1, C2, and C3 was thermodynamically shifted to C1/C2/C3/G3@C3 = 13:13:11:63, wherein a thermodynamic equilibration was shifted most favorably toward formation of G3@C3 through G3-induced stabilization of C3 by encapsulation.

Applications of Pd-Npy coordination capsules with the use of 1 have been investigated. We have demonstrated that capsule C1 serves as a guard nanocontainer for trans-4,4'diacetoxyazobenzene (trans-G4) to protect against the trans-tocis photoisomerization of trans-G4 by encapsulation in C1, wherein confined space of trans-G4@C1 interferes with geometrical change in the transition state of trans-to-cis photoisomerization of trans-G4. A chiral capsule {12 · [Pd((R)-BINAP)]₄)⁸⁺·(TfO⁻)₈ (C5: R = (CH₂)₁₀CH₃) made up of the components of a 2:4 mixture of 1 and Pd((R)-BINAP)(OTf)2 was also constructed. The unique structural interconversion between capsule C5' (R = $CH_2CH(CH_3)_2$) in solution and an infinite twodimensional Pd-Npy coordination-bonded porous network sheet structure $[{1' [Pd((R)-BINAP)]_2}^{4+} (TfO^-)_4]_n$ (**S5**') in the crystalline state was observed. We found that chiral capsule C5 induces supramolecular chirality with respect to prochiral biphenyl guests 2,2'-bis(alkoxycarbonyl)-4,4'-bis(1-propynyl)biphenyls (G5~G9: alkyl chains R' = $(CH_2)_n CH_3$; n = 0~4) by diastereomeric encapsulation through the asymmetric suppression of rotation around the axis of the prochiral biphenyl moiety upon encapsulation in C5. The diastereomeric encapsulation selectivity for the association of C5 with G5~G9 showed maximum value (*d.e.* = 40%) for **G7** (*n* = 2) among them.

Further studies of chiral capsules composed of cavitand **1** and various Pd-chiral ligands are expected to endow this type of capsule with characteristics that should help in the development of functional materials, which would constitute an important advance in supramolecular nanospace chemistry.

Experimental Section

The detailed synthesis and characterization of all the compounds are described in the Supporting Information. CCDC 1998707 contains the supplementary crystallographic data for this paper. This data is provided free of charge by The Cambridge Crystallographic Data Centre and Fachinformationszentrum Karlsruhe Access Structures service.

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Keywords: cage compounds • cavitands • host-guest systems • self-assembly • supramolecular chemistry

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Expanded pyridyl-cavitands form Pd–Npy coordination capsules. These capsules have multiple functions: (1) control of self-sorting on homo- or heterocapsule selection by guest-induced encapsulation, (2) a guard nanocontainer for a photoresponsive guest, and (3) a chiral capsule that induces supramolecular chirality with respect to a prochiral guest.