## **Synthesis and Binding Properties of a Family of Potentially Chiral Spherands**

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A new type of strongly binding and chiral spherand is reported, which is composed of a phenanthroline and four alkoxy benzene units incorporated in 18-membered rings whose binding free energies and **1H** NMR spectra suggest the systems to be preorganized for complexation.

This paper describes the first incorporation of a phenanthroline unit into a number of spherands, and the resulting effects on their structures and binding properties. With the help of Corey-Pauling-Koltun (CPK) molecular models, we designed the compounds to be highly preorganized, strong binders of alkali metal, ammonium and alkylammonium ions, to be chiral but possessing  $C_2$  axes, and to be potentially useful as chiral catalysts for reactions involving carbanide ion pairs.1

Earlier papers reported that while prototypical spherand **1**  was the strongest binder of  $Li<sup>+</sup>$  and  $Na<sup>+</sup>$  known at the time,<sup>2</sup> chiral spherand  $2$  of  $D_3$  symmetry was a relatively weak binder of these ions.<sup>3</sup> The EtO groups of chiral  $C_2$  system 3 can pass through the centre of its 18-membered ring at working temperatures, and therefore its enantiomers are subject to racemization **.4** Here we report the results of an investigation of hosts **5** and **6** to establish synthetic feasibility, to determine their configurations, and to assess their configurational stabilities. Model examination of **4** and *5* provided the conclusion that configurationally pure material would probably racemize at working temperatures by sequential passage of all of the RO groups through the centre of the macroring, but that **6** should be enantiomerically stable. We chose *5* over **4** to study since in CPK models, one Me group of the four present in **4** can turn inward to fill its cavity, whereas an Et group of *5* turned inward appears to be a higher energy conformation than one with all Et groups turned outward. Systems enantiomerically stable in the latter conformation, such as  $6$ , possess  $C_2$  symmetry, and appear in models to be good candidates for use as asymmetric catalysts.

Spherands *5* and **6** were synthesized by the reaction sequences,  $7^5 \rightarrow 8^+ \rightarrow 9^+ \rightarrow 10^+ \rightarrow 11,11+ 12^6 \rightarrow 13^+ \rightarrow 14 \rightarrow$  $15^{\frac{1}{2}} \rightarrow 16^{\frac{1}{2}} \rightarrow 5^{\frac{1}{2}}$  and  $16 \rightarrow 17^{\frac{1}{2}} \rightarrow 6^{\frac{1}{2}} + 18^{\frac{1}{2}}$  Compound 18 is the meso-form of **6,** characterized most convincingly by its crystal structure (see below). The 1H NMR spectra of **5** and 5-NaBr in  $(CD_3)_2$ SO, 360 Hz, clearly show the CH<sub>2</sub>Me

t These new compounds gave elemental analyses within 0.30% of theory, and the expected <sup>1</sup>H NMR and mass  $(m/z, M + H<sup>+</sup>)$  spectra.

 $\ddagger$  Diol 7<sup>5</sup> was monoethylated with 1.1 equiv. of K<sub>2</sub>CO<sub>3</sub> and excess of Et<sub>2</sub>SO<sub>4</sub> in acetone at  $25^{\circ}$ C  $(22 h)$  to give  $8\dagger$   $(95\%)$  as a colourless oil, which was brominated with  $Br_2$  in CHCl<sub>3</sub> at 25 °C to provide 9<sup>†</sup> (78%) as prisms, m.p. *97-98°C.* This phenol was ethylated with excess of  $K_2CO_3$  and  $Et_2SO_4$  in acetone (25 °C, 26 h) to give 10<sup>†</sup> (96%) as a colourless oil. This bromide with BuLi in tetrahydrofuran at -78 °C (10 min) gave the corresponding lithiate. This material was treated with  $(MeO)_{3}B$  at  $-78^{\circ}C$ ; the boronic ester was allowed to warm to  $25^{\circ}$ C and was hydrolysed with  $H_3O^{\dagger}$ -Et<sub>2</sub>O to the boronic acid 11 (75%) and *25%* of reduced material. Without separation, the mixture was submitted to a Suzuki<sup>7</sup> coupling with 12<sup>6</sup> [Ar, Na<sub>2</sub>CO<sub>3</sub>,  $Pd(PPh<sub>3</sub>)<sub>4</sub>$ , MeC<sub>6</sub>H<sub>5</sub>, H<sub>2</sub>O at reflux] to give after alumina gel chromatography *(80%* cyclohexane *20%* CH2C12) **13t** *(90%)* as a white foam. This product was deethylated by refluxing **it** in 48% aqueous HBr and AcOH for 48 h to give tetraphenol **14** *(ca.* loo%),  $m.p. > 300 °C$ , whose elemental analysis was unsatisfactory but whose lH **NMR** and mass spectra were as expected. Bromination of **14** with **2,4,4,6-tetrabromocyclohexa-2,5-dienone~** gave **1st** (77%) as an orange powder,  $m.p. > 300 °C$  (decomposition). This material when ethylated with NaH, Et<sub>2</sub>SO<sub>4</sub>, tetrahydrofuran under Ar at 25 °C for 24 h gave 16<sup>†</sup> (83% after filter chromatography on alumina gel-Et<sub>2</sub>O)



protons to be diastereotopic, being split into four distinct multiplets lying between 6 **3** and 4, (the same protons of noncyclic 13 and 16 were enantiotopic). This spectrum shows that  $5$  possesses  $C_2$  symmetry and that its OEt groups possess the up-down-up-down arrangement formulated in *5.* The 1H NMR spectra of 6 and  $6$ -NaBr in  $(CD_3)_2CO$  also indicated these compounds to possess *C,* symmetry. Further splitting of its protons was observed in the  $1H NMR$  spectrum of  $6.Na$ 

The crystal structure of  $18.2Me<sub>2</sub>CO$  (crystallized from acetonewater, determined at 25 °C) belongs to the monoclinic space group  $C_2/c$ :  $a = 24.507(3)$ ,  $b = 25.582(4)$ ,  $c = 18.356(3)$  Å,  $\beta = 116.683(4)$ °,  $Z = 8$ ; 3313 reflections with  $I > 3\sigma(I)$ ;  $2\theta_{\text{max}} = 100^{\circ}$ ; Cu-K $\alpha$  radiation;  $R = 0.08$ ,  $R_w = 0.12$ , GOF = 3.48. Atomic coordinates, bond lengths and angles, and thermal parameters have been deposited at the Cambridge Crystallographic Data Centre. See Notice to Authors, Issue No. 1.

(+ )-camphorsulfonate, whose chiral counterion produces diastereomeric complexes, which results in the doubling of at least four **of** its aromatic proton signals.

A stereoview of the crystal structure of 18  $(R = 0.08)$  is formulated in 21. The conformational structures **19** and 20 are consistent with very preliminary crystal structures, respectively, of *5* and 5.NaBr (not given here), as well as with their 1H NMR spectra and conclusions drawn from inspection of their CPK molecular models.

These results taken as a whole provide the following structural conclusions. *(i)* The host of 5.NaBr and free *5* are sufficiently similar to indicate *5* to be fully preorganized for binding, and are members of the spherand-spheraplex family of compounds. *(ii)* Both 19 and 20 possess a down-up-downup arrangement of their OEt groups, and belong to the chiral  $C_2$  point group. *(iii)* The arrangement of ligands about the  $Na<sup>+</sup>$  in 20 cannot assume an ideal octahedral geometry, as evident from model examination. *(iv)* Host 18 possesses an up-down-down-up arrangement of Me<sub>2</sub>CHCH<sub>2</sub>O groups as shown in **21.** Host **18** is preorganized to not bind cations, and is a diastereoisomer of *6.* The latter undoubtedly possesses the down-up-down-up arrangement, as indicated by its 1H NMR spectrum.

The free energies for binding  $(-\Delta G^{\circ})$ , kcal mol<sup>-1</sup>; 1 cal = 4.184 J) by  $5$  of the alkali metals and  $NH<sub>4</sub>$ , MeNH<sub>3</sub> and  $Me<sub>3</sub>CNH<sub>3</sub>$  picrates (Pic) were determined at 25 °C in CDCl<sub>3</sub> saturated with  $D_2O$  by the extraction of the salts from  $D_2O$  by the host at  $10^{-3}$  l mol<sup>-1</sup> in CDCl<sub>3</sub>.<sup>11</sup> The respective values of  $-\Delta G^{\circ}$  were as follows: Li<sup>+</sup>, 15.8; Na<sup>+</sup>, 15.7; K<sup>+</sup>, 15.0; Rb<sup>+</sup>, 13.6; Cs<sup>+</sup>, 13.7; NH<sub>4</sub><sup>+</sup>, 14.3; MeNH<sub>3</sub><sup>+</sup>, 13.4; Me<sub>3</sub>CNH<sub>3</sub><sup>+</sup>, 11.5 kcal mol-1. Thus **5** is a remarkably strong general binder of all of these ions, the range spanning from 15.8 for Li+ to 11.5 kcal mol<sup>-1</sup> for  $Me<sub>3</sub>CNH<sub>3</sub><sup>+</sup>$ , the smaller Li<sup>+</sup>, Na<sup>+</sup>, and K<sup>+</sup> being the more strongly complexed, but by small amounts.

These results show reduced binding and loss of specificity when the phenanthroline group is substituted for two alkoxyaryl units in prototypical spherand, **1,** which binds only LiPic

as a white foam. Similarly, tetraphenol 15 was isobutylated with BuiI-NaH-Me2NCH0 to give **17t** (62%) as a white foam. Ring closure of dibromide 16 was accomplished under dry conditions under Ar by lithiation at **-78** "C in tetrahydrofuran, and by cannulating the lithiate (20 min) into a dry, refluxing  $MeC<sub>6</sub>H<sub>5</sub>$  solution of tris(pentanedionato)iron(m) (high dilution). The mixture was refluxed for 1 h, cooled, stirred with aqueous HC1, evaporated under vacuum, and the residual solids were extracted with CH<sub>2</sub>Cl<sub>2</sub>. The resulting extracts were evaporated, and the residue was submitted to reverse phase chromatography on coated silica gel<sup>9</sup> with 6% (v/v) NaBr in  $60:40$ (v/v) acetone- $H_2O$  as the mobile phase. During this chromatographic step,  $5 \cdot Li^{+}$  underwent cation exchange to give  $5 \cdot Na^{+}$ . The fractions containing product were evaporated and the dried material was extracted into  $CH_2Cl_2$ . The combined extracts were filtered and the filtrate was evaporated and dried to give 5-NaBr (14%) as a foam, whose <sup>1</sup>H NMR and mass spectra were as expected. This complex suspended in MeOH and deionized water was heated to reflux and the MeOH allowed to evaporate. The water was decanted, and the procedure repeated three times. The residual solid was dried at 110 *"C*  for **6** h to give **57** (86%) as a powder. Application of the same ring-closing and decomplexing procedurelo to dibromide 17 gave **6t**  (0.4%) as a powder, and a trace of isomeric host **18** as a crystalline solid.













and NaPic, but with  $-\Delta G^{\circ}$  values of  $>23$  and 19.3, kcal mol<sup>-1</sup>, respectively.<sup>12</sup> Host 1 is not only preorganized for binding, but also its ligating sites are shielded from solvation, which is not true in **5**. The planar phenanthroline ring spreads the ends of the quateraryl moiety, which provides the system with great flexibility regarding the placement of solvent molecules and of its ligating oxygens, and determination of its cavity size. What is surprising is that **5** binds Li+ so well, since in CPK models, only two nitrogens and two of the alkoxy oxygens can simultaneously ligate the metal. The most similarly shaped analogue of *5* is 3, which binds Li+ with only 7 kcal mol<sup>-1</sup> ( $-\Delta G^{\circ}$ ).<sup>4</sup> The  $-\Delta G^{\circ}$  values for 5 binding Li<sup>+</sup>, Na<sup>+</sup> and **K+** resemble those observed for **2213** which contains two phenanthroline units, and provides the respective values of 15.4, 15.5 and 14.4 kcal mol<sup> $-1$ </sup>. In overall binding ability, 5 resembles hemispherand  $23^{14}$  whose  $-\Delta G^{\circ}$  values (kcal mol-1) are: Li+, 12.1; Na+, 15.4; **K+,** 15.6; **Rb+,** 14.2; **Cs+,**  13.1; NH4+, 14.4; MeNH3+, 14.4; and Me3CNH3+, 13.2. The low yield in the synthesis of 6, whose enantiomers are undoubtedly stable to racemization, unfortunately makes it a

poor candidate for asymmetric induction studies. The authors warmly thank the US Public Health Service for

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