LARGE MOLECULAR CAVITIES BEARING SIDEROPHORE TYPE FUNCTIONS

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Abstract. Novel host skeletons (I), (2) and (3) of type I spacered by triphenylbenzene or trityl- or benzene units and bearing three catechol functions, have been synthesized. They constitute the largest molecular host cavities known so far. The very strong Fe^{3@} binding of the new cyclic and acyclic ligands and their capability to enclose organic guest molecules was investigated by spectroscopic methods and by competition experiments.

1. INTRODUCTION

The macrobicyclic ligand 4b forms a tailor-shaped molecular cavity with six catechol oxygen atoms allowing the insertion of small matching cations like Fe^{33} .¹ The complex constant of the $4b$ Fe^{3@} complex is 10⁵⁹, the highest ever reported for a complex of this type. 2 In contrast to **4b,** the vastly enhanced cavity insids hitherto unknown skeletons like lb is spacered by six additional p-phenylene units and should rather be suited to recognize and enclose large organic molecules of complementary size and shape as guest particles than to bind $Fe^{3\theta}$ intramolecularly.

In this contribution we describe the first syntheses of the largest molecular cavities of type I known so far.

2. SYNTHESES

The synthetic steps yielding the novel skeletons 1, 2, 3 and in particular the siderophore type ligands 1b, 2b, 3b (cyclic) and 7AbB, 7BbB (acyclic) are outlined below.

The critical cyclization steps 7Cay + 5B + 3a, 7Bay + 5B + 2a and 7Aay + 5A + 1a were carried out under dilution conditions in chlorobenzene as solvent at 132° C. A step-by-step synthesis of the macrobicyclic ligands using the triamines $5(A, B,$ C) and 2,3-dimethoxy-4-methoxycarbonylbenzoylchloride (6) 3 leads to yields of 5%, 12% and 3%, respectively. Splitting the OCH₃ bonds was achieved by use of BBr₃ and yielded 3b, 2b and lb. The general procedures are described in the experimental section.

3. UV/VIS SPECTROSCOPIC INVESTIGATION OF Fe³⁰ COMPLEXATION

CPK models (Figure 1) show that 2b should be conformationally flexible enough to form octahedral Fe^{30} complexes inside the cavity. On the other hand, the more spacered cavity of 1b is definitely too large for one Fe^{30} ion being enclosed in its centre.

Figure 1. A comparison of the space filling and cavity sizes of lb, 2b and 4b (from left to right)

According to molecular models the six OH groups of lb can be easily directed towards the cavities' centre all at once without significant steric hindrance. The space filling of lb, shown in Figure 1, gives an idea of the size and shape of the cavities' interior. The formal transition from a small cavity like 4 to a cavity like 1 using analogous structural principles and synthetic strategies, only differing by the application of larger spacer units, can be used as a concept to construct or even to tailor-shape functionalized molecular cavities of type I and similar ones bearing modified donor systems.

As UV spectra show, 3b, 2b and 1b as well as 4b form complexes with $Fe^{3\theta}$ ions λ_{max} = 600, 598 and 581 nm, resp.; in DMSO (4b: λ_{max} = 538 nm)]. Competition experimax
ments between the ligands and Fe^{3@} (equal amounts of 4b and 3b, 4b and 2b, 4b and 1b and the open chained ligands 7Ab β , 7Cb β are added to solutions of FeCl₃ in DMSO) show the following results: At room temperature there is no significant equilibration between the complexes, which are all formed to a certain extent. Even the open chained ligand 7Ab8 under these conditions competes successfully with the macrobicyclic ligand 4b in the complexation of $\text{Fe}^{3\theta}$. The situation is changed drastically by heating the solutions. The UV spectra of all mixtures mentioned now in all cases are identical with that of the $4b\cdot Fe^{3\theta}$ complex, showing that this specific complex as the strongest one is formed exclusively.² All other ligands mentioned cannot compete under these conditions.⁴

Figure 2. UV spectra of the **lb,** 2b and 4b.Fe3@ complexes (in DMSO)

4. ¹H-NMR STUDIES OF HOST/GUEST INCLUSION

In contrast to $4b$, 3b and 2b which may be used for photometric Fe³⁰ detection and determination, lb should also form complexes with larger guest particles, e.g. organic molecules of complementary size and shape, as has been revealed for hosts bearing cavities fitted with similar ionizable functions.⁵ Neither 1,3,5-tri-tertbutylbenzene (8) as an uncharged guest leads to highfield shifted 1 H-n.m.r. signals $(in$ DMSO- $d_c)$ compared to the isolated components, nor do molecules of complementary functionality like 1,3,5-benzenetriyl-tris (methaneamine) (9) (after protonation by the hosts' OH groups) effect highfield chemical shifts. Only the proton transfer from the OH groups to the NH₂ groups can be detected: The failure of an encapsulation of a sterically fitting tris-cation by an anionic macrobicyclic large cavity is in accord with previous findings.⁵ We conclude that the binding of a morefold charged guest inside a morefold opposite charged host with matching cavity size rises the problem of a sincere electrostatic barrier.⁶ Similar electrostatic barriers may be relevant to some extent to transport processes at biological membranes.

Whereas space filling models indicate toluene, benzene and 1,2-dichloroethane to be matching guests for the cavity of 3b, experiments using this potential host/ guest combination in D_2O as the solvent did not result in high field shifts for host or guest protons. We suppose that the three catechol units do not form favourable walls for the inclusion of a guest molecule, because their aromatic hydrogen atoms project into the interior **Of** the cavity. The OH groups must likely being turned outward in an appropriate hydrophilic solvent.

Therefrom we conclude that in the future design of host cavities it might be reasonable to build up walls facing the cavity sphere rather than partly projecting into it perpendicular to the potential guest.

As can be seen from the $1H-n,m,r$. spectrum, 3a forms a clathrate with 1 mol crystal-chloroform. The latter can be removed by heating for 5 d at $80^{\circ}C/0.05$

Figure 3. ¹H-n.m.r. spectra of 1a (400 MHz), 2a (90 MHz) (in CDCl₃) and of the 3a clathrate with chloroform (90 MHz, in CD_2Cl_2)

Ilost systems **of** the types lb and Zb 7 are of additional interest beyond their receptor model properties, not only because these spacer units can be intramolecularly combined (cf. 3a, 3b; see experimental section), but also as they allow further functionalization utilizing their OH groups.

The hexaoxa acetic acid 10b, e.g. (from 1b and ethyl- α -bromoacetate) should be capable of forming a smaller cavity compared to lb and therefore again be able to coordinate metal cations. A novel family of acidic, EDTA type "concave complexes" should evoke through alternation of the $\texttt{(CH}_2)_{n}$ chain.

EXPERIMENTAL SECTION

Physical Measurements. Melting points are taken on a Reichert microscope melting point apparatus and are uncorrected. IH-n.m.r. spectra were recorded on a Bruker WH-90 and WM-400 instrument. Ultraviolet-visible spectra were recorded on a Cary model 219 spectrophotometer with quartz cuvettes. Mass spectra were recorded on MS-30 and MS-50 A.E.I., Manchester, U.K. FAB mass spectra were performed at the Institut für Physikalische Chemie, Universität Bonn, by Prof. Dr. F. Röllgen and Dipl.-Chem. K.P.Wirth and at the Institut fUr Physiologische Chemie, Bonn, by Mrs. Dr.J.Peter-Katalinic. TLC and column chromatography were performed on silica gel coated AL sheets and silica gel (0.063-0.1 mm) from E.Merck, Darmstadt.

Trimethyl-2,2',2", 3,3',3"-hexamethoxy-4,4'.4 "-[1,3,5-benzenetriyItris-f4,2_phenyienmethyieniminocarbonylljtris benzoate **(7aaa)j** *Trimethyl-2,2',2".3,3',3"-hexamethoxy-#,#I, 4"-[l,i,f-ethantriyitris(4,l-phenylenmethyleniminocarbonyl)~tris benzoate (7Bao); Trimethyl-2,2',2",3,3',3~-hexemethoxy-4,4',4~-[f,3,5-bensenetriyltris(methyleniminocarbonyl)]tris benzoate* **(7Caa):** Triethylamine (8.3 ml., 60.0 mmol, dried with KOH and distilled after addition of 2% phenylisocyanate) and triamine SA (SB, SC) (13.05 mmol) are dissolved in 100 mL of N,N-dimethylacetamide (Janssen Nr. 18588-4), methyl 2,3-dimethoxyterephthaloyl chloride (6) (IO.13 g, 39.16 mmol) is added dropwise to the stirred solution. Colourless ammonium salts precipitate. After 4 hrs stirring in the closed flask the precipitated salts are filtered by suction. The solvent is evaporated. The **residue is** dissolved in **chlo**roform, rated. washed with 2% NaOH, 2% HCl and water, dried with ${\rm Na}_2$ SO_A and again evapo To remove the solvent completely, the crude product is melted at 15O*C and evacuated (0.05 torr, 6.6 Pa) for 1 hr. After cooling to room temperature the material is purified by column chromatography on silica gel with CHCl₃/EtOH:

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7Aa¤ (chloroform/ethanol 93:7, v/v): R<sub>r</sub>= 0.50
7Ba\alpha (chloroform/ethanol 97:3, v/v): R_{\overline{v}} = 0.507Caa (chloroform/ethanol 25:2, v/v): R<sub>F</sub>= 0.57.
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The solvent is evaporated and the residue dried at 100° C/0.05 torr.- The m.p. are
The solvent is evaporated and the residue dried at 100° C/0.05 torr.- The m.p. are 138-140°C, 146-148°C and 44-48°C, respectively. ¹H-n.m.r. and MS data see Table 1.
Anal.calcd for **7Aaα**, C₆₀H₅₇N₃O₁₅ (1060.12): C 67,98, H 5.42, N 3.96; found: C
67.90, H 5.70, N 3.89.- Anal. calcd for **7Baα**, C 5.68, N 4.15, found: + $\frac{1}{2}$ $\frac{1$ C 66.45, H 5.87, N 4.39. 7Ca α : $C_{A}^{}$ α _{A 5} α ₃ C_{5} (831.83):

2,2',2",3,3', 3 *"-Hexamethory-d,d',d "-[1,3,5-benzenetriyltris-(4,1_phenylenmethylen*iminocarbonyl)] *tris benzoic acid (7AaB)*; 2,2',2",3,3',3"-Hexamethoxy-4,4',4"-[1,1, *l-ethantriyltris(4,1-phenylenmethy~eniminocarbo~yl)~tris benzoic acid (7BaEf:* To a solution of 7Aaa (7Baa) (6.06 mmol) dissolved in 180 mL of diglycol dimethylether and 70 mL of water sodium hydroxide (0.84 g, 20.98 mmol, p.a.) is added. The mixture is stirred until the sodium hydroxide is dissolved completely and refluxed for 5 hrs. After cooling the solvents are distilled off. The residue is dissolve in water. The solution is acidified using diluted HCl. The flocky precipitate is filtered by suction, washed with water and finally dried at 100°C/0.05 torr (6.6 Pa); the yield is nearly quantitative. IB-n.m.r. data see Table I. Anal. calcd for **7AaB**, C₅₇H₅₁N₃O₁₅ (1018.05): C 67.25, R 5.05, N 4.13; found C 67.10, H 5.17, N 4.12.-"Anāl."cālcd for 7Baß, C₅₃H₅₁N₃O₁₅ (970.00): C 65.63, H 5.30, N 4.33; found C 65.87, H 5.44, N 4.51.

l'risodium-2,2',2',3,3',3n-hexamethoxy-4,4',4 "-[1,3,5-benzenetrlyItrisfmethy.Zeniminocarbonyllltris benzoate (sodium salt* **of** 7Ced): To a solution of 7Caa (7.64 g, 9.18 mmol) dissolved in 150 mL of methanol (distilled) and 3 mL of water sodium hydroxide (I.15 g, 27.7 mmol, p.a.1 is added. The mixture is stirred until the sodium hydroxide is dissolved completely and refluxed over night. After cooling the solvents are distilled off. The foam produced is dried in vacua, pulverized and dried at 100°C/O.O5 torr (6.6 Pa). The substance decomposes at 245°C, the yield
is nearly quantitative. ¹H-n.m.r. data see Table 1. C₃₀H₃₆N₃Na₃O₁₅ (855.69).

2,2',2",3,3 ', *3"-Hexamethoxy-#,#I,4 "-[1,3,5-benzenetriyltris(4,1-phenylenmethylen*iminocarbonyl)]tris *benzoyl* chloride (7AaY), 2,2',2",3,3',3"-Hexamethoxy-4,4',4"-[1,1, *f-athantriyltris(4,1-phenylenmethyleniminocarbo~yl~~tr~s benzoyz chloride (7BaYl: A* 1-ethantriyitris(4,1-phenylehmethylehimihocarbong1); is benzoyi chloride (7ba1): A
100 mL flask fitted with 50 mL of thionyl chloride (colourless, otherwise distilled) is cooled with liquid nitrogen until the thionyl chloride begins to solidify. After that, 3.90 mmol of the tris benzoic acid 7Aa8 (7Ba8) are added. The mixture is stirred, while the flask is warmed to room temperature by the surrounding air. Stirring is continued at 20°C for additional 0.5 hr. The thionyl chloride is distilled off in vacuo from the yellow solution at a temperature <25°C. A pale yellow foam is formed, from which the thionyl chloride is completely removed in vacuo (0.05 torr, 6.6 Pa). After some hrs the material changes to a pale solid. The yield is quantitative. 'H-n.m.r. data see Table 1. 7AaY: C₅₇H₄₈C1₃N₃O₁₂ (1073.38), 7Bay: $c_{53}H_{48}c_{13}N_3o_{12}$ (1025.33).

2,2',2",3,3',3 *"-Hexamethoxy-d,d',d "-[i,3,5-benzenetriyltris(methyleniminocarbonyl)~* tris *benzoyl chloride (7Cayl: A* 100 mL flask fitted with 50 mL of thionyl chloride is cooled with liquid nitrogen until the thionyl chloride begins to solidify. After that, 5.00 g (ca. 5.8 mmol) of the trisodium salt of 7CaB are added. The mixture

is stirred, while the flask is warmed to room temperature by the surrounding air. Stirring is continued at 20°C for an additional 0.5 hr; after this, no educt should swim on the thionyl chloride surface. The mixture is filtered by suction through a glass frit. Usually, NaCl is easily filtered off in this way. The thionyl chloride is distilled off in vacua from the yellow filtrate at a temperature <2S°C. A pale yellow foam is formed, from which the thionyl chloride is completely removed in vacua (0.05 torr, 6.6 Pa). After some hrs the material changes to a pale solid with m.p. 130-132°C. The yield is quantitative. 1_H -n.m.r. data see Table 1. $C_{39}H_{36}Cl_{3}N_{3}O_{12}$ (845.08).

2,2',2",3,3',3"-Hexahydroxy-4,4',4"-[1,3,5-benzenetriyltris(4,1-phenylenmethylen*iminocarbonyl>]tris* benzoic acid *17AbBl; 2,2',2",3,3',3"-Hexahydroxg-4.4',4"-f1.t. f-ethantriyltris~4,J-phenylenmethyleniminocarbo~y~~]tris benzoic acid (ISbE);* 2,2',2",3,3',3"-Hexahydroxy-4,4',4"-[1,3,5-benzenetriyltris(methyleniminocarbonyl)] tr*is benzoic acid (7CbB):* To 1 1M solution of BBr₃
mmol) under cooling with ice a solution of **7Aac (7**B tris benzoic acid (**7Cb8**): To 1 1M solution of BBr₃ in dichloromethane (7.5 mL, 7.5 m
mmol) under cooling with ice a solution of **7Aaα (7Baα, 7Caα**) (0.22 mmol) in 20 mL of dichloromethane (distilled from $P_A O_{10}$) is added dropwise. After completed addition the mixture is stirred at room temperature for **24 hrs. Under cooling with ice** 20 mL of water are added dropwise. The mixture is stirred intensively at room temperature for 4 hrs, afterwards filtered by suction (frit) and washed several times with water and dichloromethane. The residue is dissolved in some mL of water and the pH set to 11 using NaOE. The nearly clear solution is filtered, the filtrate is acidified using diluted HCl. The **fine** flocky precipitate is filtered by suction, washed five times with water and finally dried at 100°C/0.05 torr (6.6 Pa). The yields are 7AbS: 81%, **7BbE:** 78% and in the case of 7CbE: 73% colourless powder, insensitive to air, thermal decomposition at 330, 250 and 205°C, respectively
¹H-n.m.r. and MS data see Table 1. 7AbB: C_{si}H_{oo}N₂O_{ts} (933.88), 7BbB: C₄₇H_{oo}N. and MS data see Table 1. 7AbB: C₅, H₃₀N₃O₁₅ (933.88), 7BbB: C₄₇H₃₀N₃O₁ (885.85), $7CbB: C_{33}H_{27}N_{3}O_{15}$ (705.57).

12,13,35,36,55,56-Hexamethoxy-9,16,32,39,52,59-hexaazatridecacyclo[22.22.18.
2^{11,14},2^{18,21},2^{27,30},2³⁴,³⁷,2^{41,44},2^{47,50},2^{54,57},2^{61,64},1^{3,45},1^{22,26}]tetrao **conta-1,3(71/,4,6,11.13,18,20,22,24,26(78~,27,29,34,36,4~,43,45,47,4~,54,56,6!,** *63,65,67,69,72,74,76,79,81,83-tritriacontaene-l0,15,33,38,53,58-hexone (la); 1,20- Dimethyl-10,11,29,30,47,48-hexamethoxy-7,14,26,33,44,51-hexaazaundecacyclo*
16.2^{2,5}.2^{9,12}.2^{16,19}.2^{21,24},2^{28,31}.2^{35,32}.2^{39,42}.2^{46,49}.2^{53,56}]tetraheptac *4,9,11,16,18,21,23,28,30,35,37,39,41,46,48,53,55,57,59,61,63,65,67,69,71,73-heptacosaene-8,13,27,32,45,50-hexone (2a); 1-Methyl-10,11,25,26,43,44-hexamethoxy-7,14,*
22*,29,40,47-hexaazanonacyclo*[16.16.12.2^{2,5}.2^{9,12}.2^{24,27}.2^{31,34}.2^{35,38}.2^{42,45}.1^{16,} *20]henhexa-2,4,9,11,16,18,20(57~,24,26,3l,33,35,37,42,44,49,5f,53,55,58,6O-heneicosaene-8,13,23,28,41,46-hexone r'3a): 8,9,23,24,35,36-Hexamethoxy-5,12,20,27,32,39 hexaazaheptacyclo[l4.l4.lO.27~1o.222~2s.23483'.l3~29.~14f~~* joctatetraconta-1,3(431, *7,9,J4,16,J8(46~,22,24,29,34,36,4J,44,47-pentadecaene-6,IJ,2f,26,33,38-hexone (4&I:*

The following components are used for the cyclization reactions:

In a high dilution apparatus (according to V6gtle, NORMAG) a solution of the trichloride and a solution of the trinmine are added dropwise and synchronously within 8 hrs in a argon atmosphere to 2.2 L **of** vigorously stirred solvent. During this addition colourless fluffy precipitates are formed. After cooling to room temperature the suspension is filtered and the residue is washed well with the solvent. The solvents of the filtrate are distilled off, the residue stirred with *little* methanol, the resulting solid collected on a glass frit. The residue is dissolved in 100 mt of chloroform and the solution is washed with 2% HCl, 2% NaOH and water, dried with Na₂SO₄ and evaporated. The residue is chromatographed on a silica gel
column (35 x 500⁴mm): ia (chloroform/ethanol 93:7, v/v): <u>R.</u>w 0.41; 2a (chloroform ethanol 93:7, v/v): $R_{\rm p} = 0.41$; 3a (dichloromethane/acetone/ethanol 80:15:5, v/v/v):
 $R_{\rm p} = 0.45$; 4a (chloroform/ethanol 9:1, v/v): $R_{\rm p} = 0.48$.- The solvent is evaporated
and the residue dried at 100°C/0.05 tor **la:** C₈₄H₇₂N₆O₁₂ (1357.53); 2a: C₇₆H₇₂N₆O₁₂ (1261.45); 3a: C₆₂H₆₀N₆O₁₂ (1081.20);

4a: $C_{48}H_{48}N_6O_{12}$ (900.93).

12, 13, 35, 36, 55, 56-Hexahydroxy-9, 16, 32, 39, 52, 59-hexaazatridecacyclo (22.22.18.2^{4,7}.

2^{11, 14}. 2^{18, 21}. 2^{27, 30}. 2³⁴, ³⁷. 2⁴¹, ⁴⁴. 2⁴⁷, ⁵⁰. 2⁵⁴, ⁵⁷. 2^{61, 64}. 1³, ⁴⁵. 1^{22, 26} ltetrao conta-1,3(71),4,6,11,13,18,20,22,24,26(78),27,29,34,36,41,43,45,47,49,54,56,61, 63,65,67,69,72,74,76,79,81,83-tritriacontaene-10,15,33,38,53,58-hexone (1b); 1,20-Dimethyl-10, 11, 29, 30, 47, 48-hexahydroxy-7, 14, 26, 33, 44, 51-hexaazaundecacyclo [20.20.
16.2², 3.2³, 12, 11, 29, 30, 47, 48-hexahydroxy-7, 14, 26, 33, 44, 51-hexaazaundecacyclo [20.20. 22, 29, 40, 47-hexaazanonacyclo [16.16.12.2^{2, 5}.2^{9,11}.2², 2³, 2 20]henhexa-2, 4, 9, 11, 16, 18, 20(57), 24, 26, 31, 33, 35, 37, 42, 44, 49, 51, 53, 55, 58, 60-heneicosaene - 8, 13, 23, 28, 41, 46-hexone (3b); 8, 9, 23, 24, 35, 36-Hexahydroxy - 5, 12, 20, 27, 32,
39-hexaazaheptacyclo [14.14.10.2^{7, 10}, 2², 2², 2⁵, 2³, 1³, 2⁹, 1¹, ¹⁹ octatetraconta-1,
3(43), 7, 9, 14, one (4b): To a 1 M solution of BBr, in dichloromethane (7.5 mL, 7.5 mmol) under
cooling with ice a solution of BBr, in dichloromethane (7.5 mL, 7.5 mmol) under
cooling with ice a solution of 1a (2a, 3a, 4a) (0.22 mmol) in of water are added dropwise. The mixture is stirred intensively at room temperature for 4 hrs, afterwards filtered by suction (frit) and washed several times with water and dichloromethane. The residue is dried at 100°C/0.05 torr (6.6 Pa). The yields are 1b: 79%, 2b: 82%, 3b: 72% and 4b: 78% colourless powder, insensitive
to air, thermal decomposition at 248, 280, 285 and 250°C, respectively.- $B-n.m.r.$
and MS data see Table 1.- 1b: $C_{78}B_{60}N_6O_{12}$ (1273

 $\overline{\texttt{Table 1.}}$ $\overline{}$ H-n.m.r. and MS data

2072

Table 1, Contd.

Compound	¹ H-n.m.r. data δ [ppm]	MS $[*]$
7 Ca γ	laj l 3.95 (s, 18H, OCH ₂), 4.70 (d, 6H, Ar-H, $J_{\text{ab}} = 8$ Hz), 7.30 (s, 3H, Ar-H), 7.74, 7.90 (XB, 6H, Ar-H, $J_{AB} = 8$ Hz), 8.30 (t, broad, 3H, NH, $J = 6$ Hz)	
7AbB	ĮЫ 4.57 (s, broad, 6H, CH ₂), 7.20-7.82 (m, 21H, Ar-H), 9.69 (s, broad, 3H, NH)	
7вь8	[b] 2.07 (s, broad, 3H, CH ₃), 4.49 (s, broad, 6H, CH ₂), 6.98-7.47 (m, 18H, Ar-H), 9.42 (s, broad, $3H$, NH)	
7съ8	(b) 4,59 (d, 6H, CH ₂ , J= 6 Hz), 7.20-7.45 (m, 9H, Ar-H), 9.57 (m, 3H, NH)	FAB-MS in glycerol matrix: 706 [M+H] ^e
1a	$\lceil a \rceil$ 3.86 (s, 18H, OCH ₂), 4.76 (d, 12H, CH ₂ , J= 5.5 Hz), 7.49, 7.69 (AA'BB', 24H, Ar-H, $\frac{3}{4} = 8$ Hz), 7.78 (s, 6H, Ar-H), 8.03 (s, 6H, Ar-H), 8.18 $(t, 6H, NH, J = 5.5 Hz)$	FAB-MS in m-nitrobenzyl alcohol matrix: 1358 [M+H] Θ
2а	[a] 2.18 (s, 6H, CH ₃), 3.84, 3.88 (2s, 18H, OCH ₃), PAB-MS in m-nitrobenzyl alcohol 4.69 (d, 12H, CH ₂ , $\frac{3}{2}$ = 6 Hz), 7.09, 7.29 (AA'BB', 24H, Ar-H, $J_{AB} = 6.5$ Hz), 7.97, 8.02 (2s, 6H, Ar-H), 8.19 (t, 6H, NH, $J = 6$ Hz)	matrix: 1261 $[M+H]^{\#}$
3a	(c) 2.16 (s, $3H$, CH_2), 3.51 (s, $9H$, OCH_3), 3.61 (s, 9H, OCH_2), 4.52 ² 4.61 (m, 12H, CH ₂ , $\underline{J} = 6$ Hz), 6.73, 7.01 (AA'BB', 12H, Ar-H, $J_{AB} = 8 \text{ Hz}$), 7.50 $(s, 3H, Ar-H)$, 7.78 $(s, 6H, Ar-H)$, 8.01 $(t, 6H,$ $NH, J = 6 Hz$	FAB-MS in thioglycerol matrix: 1358 $[M+H]$ ^w
4a	ia) 2.82 (s, 18H, OCH ₃), 4.55 (d, 12H, CH ₂ , I= 6 Hz), 7.41 (s, 6H, Ař-H), 7.90 (s, 6H, Ař-H), 7.93 (t, 6H, NH, J= 6 Hz)	$EI: 900 [M^T]$
1b	$^{[b]}$ 4.66 (s, broad, 12H, CH ₂), 7.1-8.0 (m, 36H, Ar-H), 9.58 (s, broad, 6H, NH)	FAB-MS in m-nitrobenzyl alcohol matrix: 1271 [(M-2H)+H] ^e
2ь	$[15]$ 2.18 (s, broad, 6H, CH ₃), 4.57 (s, broad, 12H, CH ₂), 7.07, 7.24 (AA'BB', 24H, Ar-H, J_{AR} ⁼ 8.5 Hz), 7.42 (s, 6H, Ar-H), 9.46 (s, broad, 6H, NH)	FAB-MS in m-nitrobenzyl alcohol matrix: $1177 [M+H]$ ⁰
ЗЬ	$^{[b]}$ 2.09 (s, broad, 3H, CH ₃), 4.47 (s, broad, 12H, CH ₂), 6.82, 6.98 (AA'BB', 12H, Ar-H, J_{nn} = 8 Hz), 7.20 (s, 6H, Ar-H), 7.42 (s, 3H, Ar-H), 9.16 (s, broad, 6H, NH)	FAB-MS in thioglycerol matrix: 997 $[M+H]^\Theta$
4Ь	4.35 (s, broad, 12H, CH ₂), 6.92 (s, 6H, Ar-H), 7.31 (s, 6H, Ar-H), 8.63 (s, broad, 6H, NH)	FAB-MS in glycerol matrix: $817 [M+H]$ ^t

[a] CDCl_3 . [b] DMSO-d_{6} . [c] $\text{CD}_{2}\text{Cl}_{2}$.

^[*] For the new macrobicyclic pyrocatechols ib-3b and their methylethers no electron induced mass spectra could be obtained. Fast atom bombardment mass spectrometry supplied intensive molecular peaks using thioglycerol- and m-nitrobenzyl alcohol matrices.

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- 7. In concord with the systematization of the names of ligands (ending 'and') and complexes (ending 'plex') by *Cram* $^{\circ}$ we propose the family name 'siderands' for the iron binding ligands lb-lb and 7AbB, 'IBbB, 7Cb6 and 'sideroplexes' to their complexes. The natural siderophores then are a subclass of the siderands.
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