\AA). Insertion of a water molecule at this site in Co(d, l - (N) -Me₆[14]4,11-dieneN₄)(OH₂)CH₃²⁺ forces these geminal methyl groups back toward the mean N_4 plane (see Figure *5;* the water-geminal methyl distance is 3.26 **A;** the geminal methyl-geminal methyl distance is 6.14 **A).** In this regard it is interesting that we have found no evidence for the Co- $(N_4)(OH_2)CH_3^{2+}$ isomer with the methyl group in this coordination site. Apparently the title compound is at least 2 kcal mol⁻¹ more stable than the isomer with $Co-CH_3$ on the same side of the molecule as the geminal methyl groups.

While it is not possible to determine the magnitude of the strain energies involved, relaxation of the strained equatorial ligand in the five-coordinate intermediates would contribute to a diminished $Co-CH_3$ bond energy and could also effect a diminished activation barrier for water loss in the Co- $(Me_6[14]4, 11$ -diene $N_4)(OH_2)CH_3^{2+}$ complexes.

Registry No. $[Co(d, l(N)-Me_6[14]4, 11\text{-dieneN₄)(OH₂)CH₃] (CIO₄)₂$ 3H₂O, 76581-80-1.

Supplementary Material Available: Tables of anisotropic thermal parameters, hydrogen atom positional parameters, bond lengths to hydrogen, and structural factors and a figure showing the crystal structure of the title compound (18 pages). Ordering information is given on any current masthead page.

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Tetracarbon Metallacarboranes. 10.' On the Thermodynamically Favored Geometry of Large Nido Cages. Structure of the Thermal Rearrangement Product (Isomer 111) of $(\eta^5$ -C₅H₅)Co(CH₃)₄C₄B₇H₇

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The preferred structure of 12-vertex, 28-electron (nido) polyhedral cages, which contain two skeletal electrons beyond the 26 required for a regular closo icosahedral system and which have thus far been found in at least seven different structural classes, was the focus of this investigation. The crystal and molecular structure of the title compound, which was formed by thermal rearrangement of isomer I at 140 °C, were determined by single-crystal X-ray diffraction. Isomer III has an open-cage geometry in which all four carbon atoms and two borons reside on a 6-membered open face, with one of the carbons isolated from the other three. The cage is isostructural with the previously characterized metallacarboranes $(\eta^5$ -C₃H₅)Fe(CH₃)₄C₄B₇H₈ and $(\eta^5$ -C₃H₅)₂Co₂C₄B₆H₁₀ (isomer VII); however, the structures of the three isomers of $(\eta^5-C_5H_5)Co(CH_3)_4C_4B_7H_7$ are grossly different from each other. The observed geometry of isomer III implies a thermodynamic preference of skeletal carbon atoms for low-coordinate vertices on the open rim, even though this requires three of the four carbons to remain adjacent. The adoption of a high-coordinate vertex by cobalt is also significant. The seven established classes of 12-vertex, 28-electron cage systems are discussed in light of the present study. Crystal data are as follows: mol wt 314.9, space group $C2/c$, $Z = 8$; $a = 25.944$ (8), $b = 8.686$ (3), $c = 15.410$ (5 carbons isolated from the other three. The cage is isostructural with the previously characterized metallacarboranes $(\eta^5$ -C₃H₃)Ee(CH₃)4C₄B₇H₈ and $(\eta^5$ -C₃H₃)₂C₂_{C4}B₈H₁₀ (isomer VII); however, t = 3294 Å³; $R = 0.052$ for 1542 independent reflections having $F_0^2 > 3\sigma(F_0^2)$.

Introduction

The four-carbon carboranes $(CH_3)_4C_4B_8H_8$, $(CH₃)₄C₄B₈H₈²⁻, and (CH₃)₄C₄B₇H₉ and the metallic$ carboranes derived from them form a structurally diverse family of cages having $11-14$ vertices. As revealed by X-ray investigations,² the framework geometries of many of these systems are highly irregular, in that they deviate markedly from ideal closo polyhedra or fragments thereof. In most of the metal-containing species it is apparent that the cage structures are dictated primarily by the mode of metal attack on the carborane framework³ and are kinetically rather than thermodynamically stabilized. This has been conclusively established⁴ for the 14-vertex $(\eta^5$ -C₅H₅)₂Fe₂(CH₃)₄C₄B₈H₈ isomers I-IV, which form by metal insertion into $(CH_3)_4C_4B_8H_8^2$ under mild conditions and feature asymmetric, open-cage structures; at 300 °C these isomers ulti-

(1) Part **9:** Grimes, R. N.; Sinn, E.; Pipal, J. R. *Inorg. Chem.* **1980,** *19,* **2087.**

(3) Grimes, R. N.; Pipal, J. R.; Sinn, E. *J. Am. Chem.* **Soc. 1979,101,4172.**

mately rearrange to the thermodynamically favored *Dzd* closo polyhedron. For most other C_4 metallacarborane systems, however, the most stable geometry has not been determined.

An intrinsically important class of cage structures consists of species having 12 framework atoms and 28 skeletal electrons (Figure 1).⁵ These molecules are $(2n + 4)$ -electron (nido) systems⁶ whose special significance lies in the presence of two "extra" electrons beyond the normal allotment of 26 for a regular 12-vertex icosahedron. Known 26-electron species such as $B_{12}H_{12}^2$ and $C_2B_{10}H_{12}$ are closed-shell electronic systems in which the 13 available bonding molecular orbitals in the cage skeleton are precisely filled;⁷ introduction of more electrons must therefore induce distortion from icosahedral symmetry. However, the nature of such distortions depends cru-

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⁽⁴⁾ Maxwell, W. **M.;** Web, R.; **Sinn,** E.; Grimes, R. N. *J. Am. Chem* **Soc. 1977,** *99,* **4016.**

⁽⁵⁾ Pipal, J. R.; Grimes, R. N. *J. Am. Chem.* **Soc. 1978,** *100,* **3083.**

⁽⁶⁾ For descriptions of skeletal electron-counting procedures and structural classifications see: (a) Wade, K. Adv. Inorg. Chem. Radiochem. 1976, 18, 1. (b) Mingos, D. M. P. Nature (London), Phys. Sci. 1972, 236, 99. (c) Rudolph, R. W. Acc. Chem. Res. 1976, 9, 446. (d) Williams, R. E. *Inorg. Chem.* **1971,** *10,* **210.**

^{(7) (}a) Lipscomb, W. N. "Boron Hydrides"; Benjamin: New York, 1963.

(b) Longuet-Higgins, H. C.; Roberts, M. de V. Proc. R. Soc. London,

Ser. A 1954, 224, 336; 1955, 230, 110. (c) Moore, E. B., Jr; Lohr, L.

L., Jr.; Lips

Figure **1.** Structural types of 12-vertex, 28-electron (nido) cages. All cage structures depicted have been crystallographically characterized except that shown in type 2: type 1, $R_2C_2B_{10}H_{11}$ (R = CH₃, C₆H₅);²⁴ type 2, proposed structure of (CH₃)₄C₄B₈H₈ (isomer B)³ and of $(C_5H_5)_2C_02C_4B_6H_{10}$ (isomer VI);¹⁹ type 3, $(CH_3)_4C_4B_6H_8$ (isomer A);²³ type 4, $(C_5H_5)_2C_02(CH_3)_4C_4B_6H_6$ (left, isomer V)¹⁹ and (C_5H_5) - $Co(CH_3)_4C_4B_7H_7$ (right, isomer I);^{5,18} type 5, $(C_5H_5)Fe(CH_3)_4C_4B_7H_8$ (left),¹⁴ $(C_5H_5)_2Co_2C_4B_6H_{10}$ (right, isomer VII),¹⁵ and $(C_5H_5)Co(C-$ H₃)₄C₄B₇H₇ (bottom, isomer III, this work); type 6, (C₅H₅)Co(CH₃)₄C₄B₇H₇, (isomer II);¹ type 7, (C₅H₅)CoSe₂B₉H₉.²⁵

cially on the type, number, and location of skeletal heteroatoms⁵ and leads to a fascinating variety of cage structures, as can be seen in Figure 1.

Earlier papers¹⁻⁵ have dealt with a number of 28-electron and 30-electron (nido and arachno) 12-vertex metallacarboranes, most of which are obtained under mild conditions and hence could not be presumed to reflect thermodynamically preferred geometries. In the present article we report the structure of a new isomer (111) of a 28-electron, 12-vertex species, $(\eta^5$ -C₅H₅)Co(CH₃)₄C₄B₇H₇, whose isomers I and II have previously been characterized.^{1,8} Since isomer III was formed by thermal rearrangement of isomer I at $140 \degree C$,⁸ its geometry is obviously relevant to the question of thermodynamic stability in MC_4B_7 nido systems and in 12-vertex nido cages in general.

Experimental Section

Orange plates of $(\eta^5$ -C₅H₅)Co(CH₃)₄C₄B₇H₇ (isomer III), obtained by pyrolysis of isomer I as described elsewhere,⁸ were grown over 14 days by vapor diffusion of hexane into a tetrahydrofuran (THF) solution at 0 °C. A crystal was selected for data collection and mounted on a glass fiber in an arbitrary orientation. Crystal data: $CoC_{13}B_7H_{24}$, mol wt 314.9, space group $C2/c$, $Z = 8$; $a = 25.944$ (8), $b = 8.686$ (3), $c = 15.410$ (5) $\mathbf{\hat{A}}$; $\beta = 108.47$ (3)^o; $V = 3294$ $\mathbf{\hat{A}}^3$; μ (Mo $K\alpha$) = 10.7 cm⁻¹; $\rho_{\text{calcd}} = 1.27$ g cm⁻³; crystal dimensions (mm from centroid) (100) 0.125, (100) 0.125, (010) 0.21, (010) 0.21, (001) 0.02, **(001)** 0.02. The Enraf-Nonius program **SEARCH** was used to obtain 25 accurately centered reflections which were then employed in the program **INDEX** to obtain an orientation matrix for data collection and also to provide approximate cell constants. Refined **cell** dimensions and their estimated standard deviations were obtained from least**squares** refinement of 25 accurately centered reflections. The masaicity

of the crystal was examined by the ω -scan technique and found to be satisfactory. Systematic absences of $h + k = 2n + 1$ for *hkl*, *l* $= 2n + 1$ for *hOl*, and $k = 2n + 1$ for *OkO* indicated possible space groups *C2lc* or *Cc;* since the centric space group *C2/c* was found to provide a satisfactory model, the noncentric group *Cc* was not considered further. For $Z = 8$ this is consistent with the molecular formula on the assumption of 19.6 **A'** per nonhydrogen atom.

Collection and Reduction of the hta. Diffraction data were collected at 295 K on an Enraf-Nonius four-circle CAD-4 diffractometer controlled by a PDP8/M computer, using **Mo** *Ka* radiation from a highly oriented graphite-crystal monochromater. The θ -2 θ scan technique was used to record the intensitis for the reflections from a highly oriented graphite-crystal monochromater. The θ -2 θ
scan technique was used to record the intensities for the reflections
for which $1^{\circ} \le 2\theta \le 46^{\circ}$. Scan widths were calculated from the
formula SW for which $1^{\circ} \le 2\theta \le 46^{\circ}$. Scan widths were calculated from the formula SW = $A + B \tan \theta$, where *A* is estimated from the mosaicity of the crystal and *B* compensates for the increase in the width of the peak due to $K\alpha_1-K\alpha_2$ splitting. The values of *A* and *B* respectively were 0.60 and 0.35° for both compounds. This calculated scan angle was extended at each side by 25% for background determination (BG1 and BG2). The net count (NC) was then calculated as NC = TOT - 2(BG1 + BG2), where TOT is the estimated peak intensity. The intensities of three standard reflections were monitored at intervals of 100 reflections and showed no systematic trends. Raw intensity data were corrected for Lorentz-polarization effects, and their standard deviations were calculated in the **usual** manner from **counting** statistics $(p = 0.03)$.⁹ This resulted in 2437 reflections of which 1542 had $\tilde{F}_o^2 > 3\sigma(F_o^2)$ after averaging of equivalent reflections. Only those data for which $F_0^2 > 3\sigma(F_0^2)$ were used in the refinement of structural parameters.

Solution and Refinement of the Structure. A three-dimensional Patterson synthesis was **used** to locate the metal atom, whose position permitted the location of all other nonhydrogen atoms and B-H hydrogen atoms from difference Fourier maps. Some methyl **hy-**

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Table I. Positional and Thermal Parameters and Their Estimated Standard Deviations^{a, b}

atom	x	у	z	B_{11}	B_{22}	B_{33}	B_{12}	B_{13}	B_{23}
Co	0.16769(3)	$-0.0231(1)$	0.06401(6)	2.53(3)	2.84(3)	4.93(3)	0.04(4)	1.28(2)	$-0.32(4)$
C(7)	0.0849(2)	0.0050(7)	0.0340(4)	2.0(2)	3.1(3)	4.2 (2)	0.4(2)	1.2(2)	$-0.0(2)$
C(9)	0.0855(2)	$-0.2813(7)$	$-0.0399(4)$	3.5(3)	2.9(3)	4.1(3)	$-0.2(2)$	0.9(2)	$-0.8(3)$
C(11)	0.0950(2)	$-0.2293(8)$	0.1434(4)	3.8(3)	5.5(4)	3.5(3)	$-0.3(3)$	1.4(2)	0.9(3)
C(12)	0.1029(2)	$-0.0492(8)$	0.1265(4)	3.2(2)	5.2(4)	4.5(3)	0.3(3)	2.0(2)	$-0.9(3)$
C(7M)	0.0585(3)	0.1646(9)	0.0144(5)	3.8(3)	5.0(4)	7.0(4)	1.3(3)	1.3(3)	$-0.1(3)$
C(9M)	0.0512(3)	$-0.3528(10)$	$-0.1298(5)$	5.5(4)	5.8(4)	5.7(4)	$-1.1(3)$	1,3(3)	$-1.1(3)$
C(11M)	0.0699(3)	$-0.2615(11)$	0.2185(5)	7.2(4)	9.5(6)	5.0(3)	$-1.2(4)$	2.9(3)	1.2(4)
C(12M)	0.0981(3)	0.0572(10)	0.2017(5)	7.1(4)	7.6(5)	6.2(3)	0.5(4)	3.5(3)	$-1.6(4)$
C(1P)	0.1914(3)	0.2043(8)	0.0922(5)	5.2(3)	3.1(3)	8.8(4)	$-1.3(3)$	2.6(3)	$-1.6(3)$
C(2P)	0.1839(3)	0.1708(9)	0.0017(5)	5.1(3)	5.3(4)	7.9(4)	$-1.9(3)$	1.6(3)	1.0(4)
C(3P)	0.2189(3)	0.0583(9)	$-0.0022(6)$	8.9(3)	5.4(4)	14.0(4)	$-4.1(3)$	8.9(2)	$-3.7(4)$
C(4P)	0.2487(3)	0.0141(9)	0.0854(8)	3.1(3)	3.5(4)	19.9(8)	$-0.4(3)$	3.5(4)	0.6(5)
C(5P)	0.2316(3)	0.1089(10)	0.1450(6)	5.7(4)	7.8(5)	7.3(5)	$-4.3(3)$	$-1.4(4)$	1.0(4)
B(1)	0.1860(3)	$-0.2612(9)$	0.0884(6)	2.9(3)	3.3(4)	6.1(4)	0.5(3)	1.4(3)	0.2(3)
B(3)	0.1491(3)	$-0.2238(9)$	$-0.0251(5)$	3.6(3)	3.0(4)	4.9(4)	0.2(3)	1.6(3)	$-0.6(3)$
B(4)	0.1355(3)	$-0.3891(9)$	0.0248(6)	4.2(3)	2.4(4)	7.1(4)	0.4(3)	2.1(3)	$-0.0(4)$
B(5)	0.1424(3)	$-0.3577(10)$	0.1382(6)	4.1(4)	3.6(4)	5.9(4)	0.1(3)	0.3(3)	1.7(4)
B(6)	0.1625(3)	$-0.1717(11)$	0.1692(6)	4.1 (4)	4.9 (5)	4.7(4)	0.2(4)	0.5(3)	0.4(4)
B(8)	0.0877(3)	$-0.0892(9)$	$-0.0428(5)$	2.4(3)	3.6(3)	3.8(3)	0.5(3)	0.9(2)	0.9(3)
B(10)	0.0752(3)	$-0.3331(9)$	0.0529(5)	3.0(3)	3.5(4)	5.1(4)	$-0.2(3)$	1.2(3)	0.6(3)
atom	\mathbf{x}	\mathcal{Y}	\mathbf{z}	B, A ²	atom	$\pmb{\chi}$	\mathcal{Y}	z	B, A ²
H(1)	0.233(2)	$-0.294(7)$	0.114(4)	4(1)	H(71M)	0.022(2)	0.168(7)	$-0.000(3)$	4(1)
H(3)	0.160(2)	$-0.215(6)$	$-0.094(3)$	4(1)	H(72M)	0.062(2)	0.194(7)	$-0.040(4)$	5(2)
H(4)	0.144(2)	$-0.499(6)$	0.002(3)	4(1)	H(73M)	0.077(2)	0.213(7)	0.071(4)	6(2)
H(5)	0.156(2)	$-0.460(7)$	0.192(3)	5(1)	H(91M)	0.011(2)	$-0.340(7)$	$-0.153(4)$	6(2)
H(6)	0.181(2)	$-0.129(7)$	0.246(4)	4(1)	H(92M)	0.059(3)	$-0.457(8)$	$-0.127(4)$	8(2)
H(8)	0.070(2)	$-0.040(6)$	$-0.116(3)$	3(1)	H(93M)	0.065(2)	$-0.316(8)$	$-0.174(4)$	7(2)
H(10)	0.043(2)	$-0.421(7)$	0.043(3)	4(1)	H(111M)	0.089(2)	$-0.195(7)$	0.274(4)	6(2)
H(1P)	0.172(2)	0.277(8)	0.118(4)	6(2)	H(112M)	0.075(2)	$-0.365(8)$	0.237(4)	7(2)
H(2P)	0.158(2)	0.221(7)	$-0.047(4)$	5(1)	H(113M)	0.034(2)	$-0.239(7)$	0.201(4)	5(2)
H(3P)	0.220(2)	0.016(8)	0.062(4)	7(2)	H(121M)	0.125(2)	0.007(7)	0.268(4)	6(2)
H(4P)	0.274(2)	$-0.065(8)$	0.102(4)	7(2)	H(122M)	0.062(2)	0.061(8)	0.202(4)	6(2)
H(5P)	0.244(2)	0.100(8)	0.208(4)	6(2)	H(123M)	0.110(3)	0.160(8)	0.194(4)	7(2)

The form of the anisotropic thermal parameter is $\exp[-(B_{11}h^2(a^*)^2 + B_{22}k^2(b^*)^2 + B_{33}l^2(c^*)^2)]/4 + (B_{13}hk^2b^* + B_{13}h^2b^* + b^2b^2b^2)$ $B_{23}klb*c*$)/2]. b For all hydrogen atoms, standard isotropic B values are reported.

Table **I1**

drogens were located from the Fourier maps; the remaining methyl hydrogens, and all cyclopentadienyl hydrogens, were introduced into calculated positions. All hydrogens were included in the refinement for several cycles and thereafter held fixed. Following absorption

Table **111.** Selected Bond Angles (Deg)

corrections (maximum and **minimum** absorption coefficients were 0.95 and 0.91), refinement was continued to convergence with final values of $R = 0.052$ and $R_w = 0.055$, where $R = \sum ||F_0|| - |F_c|| / \sum |F_0|$ and $R_w = (\sum w(|F_0| - |F_c|)^2 / \sum w|F_0|^2)^{1/2}$.
Full-matrix least-squares refinement was based on *F*, and the

Fun-matrix least-squares refinement was based on r , and the function minimized was $\sum w([F_0] - [F_c])^2$. The weights *w* were taken as $[2F_o/\sigma(F_o^2)]^2$ where $|F_o|$ and $|F_c|$ are the observed and calculated structure factor amplitudes. The atomic scattering factors for non-

Figure 2. Stereoview of molecular structure of $(\eta^5$ -C₃H₃)Co(CH₃)₄C₄B₇H₇ (isomer III) (hydrogen atoms omitted).

Figure 3. Stereoview of unit cell packing.

 a M = Co or Fe. b Framework carbon atoms C(11) and C(12) were numbered C(8) and C(9), respectively, in ref 14.

hydrogen atoms were taken from Cromer and Waber¹⁰ and those for hydrogen from Stewart.¹¹ The effects of anomalous dispersion were included in F_c by using Cromer and Ibers'¹² values of $\Delta f'$ and $\Delta f''$. The computing system and programs are described elsewhere.¹³ Tables of observed and calculated structure factors are available **(see** paragraph at end of paper regarding supplementary material).

The error in an observation of unit weight was 2.0, and the largest parameter shift in the last cycle of refinement was 0.05 times the estimated standard deviation. There are no close intermolecular contacts, the shortest (nonhydrogen) distance being 3.448 **(7) A** between C(2P) and C(3P).

Results and Discussion

Final positional and thermal parameters are given in Table I, while Tables I1 and I11 list interatomic distances and angles;

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- (11) Stewart, R. F.; Davidson, E. R.; Simpson, W. T. *J.* Chem. *Phys.* **1965,** *42,* 3175.
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Figures 2 and 3 present stereoviews of the molecular structure and unit cell packing, respectively. The cage framework is an open basket with a well-defined open face on which all four framework carbons are located. This structure is entirely consistent with previously reported^{8 1}H and ¹¹B spectra, which indicated a $CoC₄B₇$ cage of low symmetry in which all boron and framework carbons are nonequivalent.

Comparison with Isostructural Complexes. The molecule clearly belongs in the structural class designated as type *5* in Figure 1; it is isostructural (and also cage-isoelectronic) with the previously characterized type 5 species $(\eta^5$ -C₅H₅)Fe- $(CH_3)_4C_4B_7H_8^{14}$ and $(\eta^5-C_5H_5)_2C_0C_4B_6H_{10}$ (isomer VII),¹⁵ both of which are also depicted in Figure 1. The type *5* geometry can be described as a 13-vertex closo polyhedron¹⁶ from which a high-coordinate vertex has been removed. **As**

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Figure 4. Structures of the structurally characterized isomers of $(\eta^5$ -C₅H₅)Co(CH₃)₄C₄B₇H₇.

such, it is the only one of the seven structural classes in Figure 1 that readily fits the standard definition^{6d} of a nido cage as a closo polyhedron minus one vertex.

Although the three type *5* cages are of similar gross shape, the arrangement of skeletal carbon atoms is different. In the dicobalt species, all four carbons occupy contiguous vertices on the open face; in the iron complex, only three carbons are on the open face, the fourth being as far from it as possible;¹⁷ and in the present (monocobalt) species, all four carbons are on the open face, but one is separated from the others. Moreover, a comparison of the analogous framework C-C and Co-C bond distances in the three species (Table IV) reveals a striking difference between the dicobalt compound and the other two: the central carbon-carbon interaction $[C(11)-C (12)$] in $(C_5H_5)_2Co_2C_4B_6H_{10}$ is much shorter (by 0.14 Å) than the corresponding C-C distances in the FeC_4B_7 and CoC_4B_7 complexes. On the other hand, the $C(7)-C(12)$ bond lengths in the three molecules are similar and relatively short (1.42-1.45 **A),** suggesting that there is significant localized bonding between the low-coordinate carbon $C(7)$ and its neighboring carbon atom in each system; in the $CoC₄B₇$ and $FeC₄B₇$ cages, this localized C(7)–C(12) bonding is evidently at the expense of the $C(11)-C(12)$ interaction which is depleted in electron density and hence relatively weak. In the dicobalt complex, it would appear that the presence of a second metal atom at the other end of the cage induces a more even distribution of bonding electron density through the threecarbon chain on the open rim, with more nearly equal C-C distances [the $C(11)$ – $C(10)$ length, which has no counterpart in the CoC₄B₇ and FeC₄B₇ cages, is 1.530 (8) \AA ¹⁵ If this interpretation is valid, one should of course expect to see corresponding effects in other type *5* metallacarboranes **as** their X-ray structures become available.

Comparison with $(\eta^5\text{-}C_5H_5Co(CH_3)_4C_4B_7H_7$ (Isomers I and **11).** The structures of isomers I and I1 (Figure 4) have previously been established via X-ray studies of the parent compounds^{1,18} and a B-ethoxy derivative⁵ of isomer I. Although the cage geometries of isomers I, 11, and I11 are of types 4, 6, and *5,* respectively (Figure l), this is not the first example of metallacarborane isomers involving three different structural classes; it was shown in earlier work^{15,19} that the dicobalt system $(\eta^5-C_5H_5)_2Co_2C_4B_6H_{10}$ forms isomers of types 2, 4, and *5* (only the latter two have been crystallographically confirmed). Similarly, the 14-vertex $(\eta^5 \text{-} C_5H_5)_2\text{Fe}_2(\text{CH}_3)_4C_4B_8H_8$ (20) Dunks, G. B.; Hawthorne, M. F. In "Boron Hydride Chemistry"; isomers also exhibit at least three different types of cage

structure.⁴ In contrast, gross structural variation among isomers is extremely rare in *dicarbon* metallacarboranes,²⁰ where isomerism usually is restricted to interchange of heteroatoms among different vertices **of** a common polyhedral framework.

The synthetic origins of isomers I and II of $(\eta^5$ -C₅H₅)Co- $(CH_3)_4C_4B_7H_7$ have been described earlier.⁸ Isomer I was produced, together with two isomers of $(\eta^5$ -C₅H₅)Co- $\overline{(CH_3)_4C_4B_6H_6}$, by reaction of $(\eta^5-C_5H_5)Co(CO)_2$ with (C- H_3)₄C₄B₈H₈ in THF under UV light; it has also been obtained by oxidation of the sandwich compound *closo,nido-* $[(CH₃)₂C₂B₄H₄]COH[(CH₃)₂C₂B₃H₅]$ in the presence of $CoCl₂$ and $C₅H₅$. Isomer II was a minor product of the treatment of $(\overline{CH}_3)_4C_4B_8H_8^2$ ion with CoCl₂, C₅H₅⁻, and HCl in THF. *All* of these syntheses were conducted at room temperature.

The thermolysis of isomer I at 140° C for 18 h in nonane solution under a nitrogen atmosphere formed isomer I11 in *5096* yield, with no other products detected.⁸ Thus, the cage structure of isomer I11 is thermodynamically favored relative to that of I; moreover, we believe it to be the most stable arrangement for this system (see below) although this has not been conclusively proved. We will not discuss here the cage rearrangement mechanism but will note that a possible pathway for interconversion of $(C_5H_5)_2Co_2C_4B_6H_{10}$ isomers of type 4 and type *5* geometry has been presented elsewhere (see ref 19, Figure 3); an analogous scheme would apply equally to the present isomerization of $CoC₄B₇$ species.

Several trends are evident in the $I \rightarrow III$ rearrangement: (1) adoption of a true nido (type *5)* geometry; (2) retention of low-coordinate vertices on the open face by the carbon atoms; (3) movement of one carbon to an isolated position with respect to the other carbons; (4) adoption of a 6-coordinate vertex by the cobalt atom. The first trend is not surprising, since the type *5* arrangement is more compact and probably permits more efficient framework electron delocalization, and hence a lower-energy system, than does type 4 (isomer I). Trend 2 is probably not a major driving force in the isomerization, since all cage carbons are low coordinate in both isomers. Trend 3 reflects the well-known tendency **of** framework carbon atoms to separate from each other; it is noteworthy, however, that *none of the carbons chooses to leave the open face in order to achieve carbon-carbon separation.* Thus, clearly the preference of carbon atoms to achieve (or retain) low coordination^{6d} outweighs their mutual replusion, and the observed structure represents the maximum possible C-C separation for a type *5* cage in which all carbons are on the open rim.

Finally, the increase in coordination of cobalt from five to six with respect to the cage may be a factor in the rearrangement since there is a general (but by no means invariable) tendency for iron and cobalt atoms to adopt high-coordinate vertices when given a choice; one can argue that the higher coordination sites permit slightly better overlap between metal and cage orbitals (with electron-rich metals such as nickel, however, the situation is often reversed so that low-coordinate vertices are favored $2¹$).

All of the evidence to date leads us to believe that type *5* geometry is the preferred arrangement for 12-vertex, 28 electron metallacarboranes. This does not, however, hold true for 12-vertex, 28-electron carboranes lacking metal atoms, in which the tendency to retain a nearly icosahedral arrangement

⁽¹⁷⁾ A mechanism has ken **proposed to account for the formation of this** complex and related ferracarboranes from $(CH_3)_4C_4B_8H_8^2$: see ref 3. **(18) Welch, A.** J., **private communication.**

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is clearly evident (Figure 1). **As** earlier work has shown, $(CH₃)₄C₄B₈H₈$ exists in solution²² as an equilibrium mixture of two forms (evidently types 2 and **3,** Figure 1) and in the solid²³ as type 3 only; the compound does not rearrange up to its decomposition temperature. Thus, the presence of one

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or more metal atoms in the cage can have a profound influence on the cage geometry even when the formal skeletal electron count is held constant.

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Supplementary Material Available: Listing of observed and calculated structure factor amplitudes **(7** pages). Ordering information is given on any current masthead page.

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Properties and Structures of Five-Coordinated Cobalt (11) and Copper(I1) Complexes of N,N'-Bis[(5-chloro-2- hydroxyphenyl)phenylmethylene]-4-oxaheptane- 1,7-diamine

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The title complexes, [Co(cbpO)] and [Cu(cbpO)], have been synthesized and studied by single-crystal X-ray diffraction. The copper complex contains the metal atom in a distorted square pyramid whose apex is formed by the Cu-0 bond to the ether oxygen atom. The 02N2 base of the square pyramid is distorted *so* as to give some trigonal-bipyramidal character to the geometry, the N atoms being raised above **(0.35 A)** the base and the Cu atom and the 0 atoms below it **(0.06** and **0.35 A,** respectively). The **Cu-0** bond to the ether oxygen is markedly elongated **(2.575 A),** indicating a rather weak linkage. This compares with apical bond elongations of **2.374 (10)** and **2.686 (1) A,** respectively, in the related five-coordinated [Cu(mbpN)] and [Cu(cbpS)]. If the long Cu-0 bond were absent, the copper environment would approximate a flattened tetrahedron. The metal environment in [Co(cbpO)] is distorted trigonal bipyramidal with the two Cc-N bonds axial and the three Co-O bonds equatorial. The Co-O bond to the ether oxygen is relatively strong $(2.124 \, (3) \, \text{\AA})$ in contrast with the [Cu(cbpO)] analogue. The observation of metal to ether oxygen bonding at all in these complexes contrasts with the apparent absence of such bonds in a series of related salicylaldimine complexes. [Co(cbpO)] readily forms a six-coordinated pyridine adduct, while [Cu(cbpO)] appears not to do so, even in pyridine solution. Crystal data for [Cu(cbpO)]: space
group $C2/c$, $Z = 8$, $a = 13.993$ (5) A, $b = 13.652$ (8) A, $c = 29.40$ (1) A, $\beta = 91.46$ (3)°, $V = 5615$ **1840** reflections. Crystal data for [Co(cbpO)]: space group *C2/c, Z* = **8,** *a* = **13.914 (7) A,** *b* = **13.477 (4) A,** *c* = **29.86** (1) \hat{A} , $\beta = 90.57$ (3)^o, $V = 5599$ \hat{A}^3 , $R = 6.1\%$ for 3397 reflections.

Introduction

Bivalent metal complexes of potentially pentadentate ligands of the type H_2 cbp**X** (1) and H_2 sal**X** (2) may form either

2, H,salX

four-coordinated or five-coordinated structures, depending on whether the central donor atom X is bonded to the metal. For H_2 salX complexes, where $X = S$, O, Taylor et al. have suggested, on the basis of spectral data, that the central X group does not usually coordinate to the metal in salX complexes, producing distorted square-planar copper(II) complexes,² tetrahedral cobalt(II) complexes,³ and possibly weakly coordinated low-spin five-coordinated nickel(II) complexes.⁴ These suggested structures have not yet been confirmed by X-ray crystal structural data. Subsequent experiments have provided evidence of M-X bonding in complexes of H_2 salX ($\dot{X} = NR$, PCH₃),^{3,5,6} H₂cbpX (X = NH, S)⁷⁻¹⁰ and H₂mbpX (X = NH).^{7,9-12} In most cases the M-X bonding has been confirmed by X-ray crystal structure data.^{5-8,10,11} Due to the weak donor properties of ether oxygen atoms, the $X = O$ ligand gives

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