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# Further Studies of the Tetraborane Carbonyl B<sub>4</sub>H<sub>8</sub>CO

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Tetraborane-8 carbonyl ( $B_4H_8CO$ ) is made by the action of CO on either  $B_4H_{10}$  or  $B_5H_{11}$ , suggesting the processes  $B_4H_{10} \rightarrow B_4H_8 + H_2$  and  $B_5H_{11} \rightarrow B_4H_8 + BH_3$ . The reversal of these (significant for polyborane interconversion mechanisms) is shown by the high yields of  $B_4H_{10}$  and  $B_5H_{11}$  obtained when  $H_2$  or  $B_2H_6$  reacts with  $B_4H_8CO$ . Basic reagents attack  $B_4H_8CO$ to form nonvolatiles without liberation of any CO; dimethyl ether and water behave so, and trimethylamine forms some ( $CH_3)_3NBH_3$ . Ethylene also fixes the CO, in a slightly volatile unstable compound, ( $C_2H_4)_4B_4H_8CO$  (structure uncertain), whereas  $BH_3CO + 3C_2H_4 \rightarrow (C_2H_5)_3B + CO$  is quantitative. With excess  $PF_3$ ,  $B_5H_{11}$  forms  $BH_3PF_3$  and  $B_4H_8PF_3$ , from the latter of which CO reversibly displaces  $PF_3$ ; in these reactions CO is not irreversibly fixed. In support of structural elucidation of  $B_4H_8CO$  by infrared and nuclear magnetic resonance spectra,  $B_4D_{10}$ ,  $B_5D_{11}$ , and  $B_4D_8CO$  were made in nearly pure form. The structure of  $B_4H_3CO$  remains uncertain but the possibilities are limited. There may be a tautomeric equilibrium such that at least one H atom alternates between two B-H-B bridgings and a terminal B-H situation—much like the tautomerism of  $B_8H_{11}$ .

Tetraborane-8 carbonyl (B<sub>4</sub>H<sub>8</sub>CO)<sup>1</sup> undergoes chemical reactions suggesting two different initial dissociations:  $B_4H_8CO \rightarrow B_4H_8 + CO$  and possibly  $B_4H_8CO \rightarrow$  $BH_3 + B_3H_5CO$ . Thus the room-temperature decomposition gives yields of B<sub>5</sub>H<sub>11</sub> near 30%, as expected from the secondary process  $B_4H_8 + BH_3 \rightarrow B_5H_{11}$ . The yield of B5H11 is still higher if much B2H6 is present during the decomposition, and the availability of  $B_4H_8$  is still further indicated when B4H8CO decomposes in the presence of much  $H_2$  to give high yields of  $B_4H_{10}$ , suggesting the reaction  $B_4H_8 + H_2 \rightarrow B_4H_{10}$ . The direct reverse of this reaction is indicated by the formation of B<sub>4</sub>H<sub>8</sub>CO from B<sub>4</sub>H<sub>10</sub> and CO,<sup>2</sup> at CO-independent rates.<sup>3</sup> Thus the easily reversible reactions  $\rm B_4H_{10} \rightarrow B_4H_8$  + $H_2$  and  $B_4H_8 + 1/_2B_2H_6 \rightarrow B_5H_{11}$  (already suggested by arguments from rate data)<sup>4</sup> gain more direct support, contributing toward a full understanding of the longknown reversible interconversions involving H<sub>2</sub>, B<sub>2</sub>H<sub>6</sub>,  $B_4H_{10}$ , and  $B_5H_{11}$ .<sup>5</sup> The  $B_4H_8$  group again plays an obvious role in the nearly quantitative formation of the new compound B4H3PF3 from B5H11 and PF3 and in the reversible displacement of PF<sub>3</sub> to make B<sub>4</sub>H<sub>8</sub>CO.

The initial formation of the  $B_3H_5CO$  unit (whether by the action of  $B_4H_8$  to remove  $BH_3$  from  $B_4H_3CO$  or by a direct dissociation) would account for much of the 34%yield of CO-containing brown solids (formed during the decomposition of  $B_4H_8CO$ ), for one would expect  $B_3H_5CO$  to condense very quickly to polymeric boron hydride material with permanent incorporation of the CO; in fact a strong heating of the brown solids *in vacuo* yields only very little CO.

The CO also is permanently incorporated in the products of the reactions of  $B_4H_8CO$  with substances having a surplus of electrons available for bonding, such as amine or oxygen bases or ethylene. The extreme contrast to the corresponding reactions of  $BH_8CO$  may be

(5) A. B. Burg and H. I. Schlesinger, J. Am. Chem. Soc., 55, 4009 (1933).

illustrated by the following examples, of which (1) and (2) are quoted from an earlier paper.<sup>6</sup>

 $BH_{3}CO + (CH_{3})_{3}N \longrightarrow (CH_{3})_{3}NBH_{3} + CO \qquad (1)$ 

 $BH_{3}CO + 3H_{2}O \longrightarrow B(OH)_{3} + 3H_{2} + CO \qquad (2)$ 

$$BH_{3}CO + 3C_{2}H_{4} \longrightarrow (C_{2}H_{5})_{3}B + CO$$
(3)

 $\begin{array}{c} B_{4}H_{8}CO + 1.03(CH_{3})_{3}N \longrightarrow [(CH_{3})_{8}N]_{1.03}B_{3.87}H_{7.61}(CO)_{0.96} + \\ 0.04BH_{3}CO + 0.045B_{2}H_{6} \quad (4a) \end{array}$ 

+ 1.47(CH<sub>3</sub>)<sub>8</sub>N 
$$\longrightarrow$$
 [(CH<sub>3</sub>)<sub>2</sub>N]<sub>1.78</sub>B<sub>3.18</sub>H<sub>5.4</sub>(CO)<sub>0.94</sub> + 0.74(CH<sub>3</sub>)<sub>8</sub>NBH<sub>5</sub> (4b)

 $B_4H_8CO + xH_2O \longrightarrow 8.8H_2 + yB(OH)_8 + residue + 0.035CO$ (5)

$$B_4H_8CO + (CH_3)_2O \longrightarrow "(CH_3)_2OB_4H_8CO"$$
(6)

$$B_4H_8CO + 4C_2H_4 \longrightarrow (C_2H_5)_4B_4H_4CO$$
(7)

Reaction 4a occurred during 6 days at  $-78^{\circ}$  with no excess amine; 4b during a further 3 months at  $-78^{\circ}$  with excess amine acting upon the residue of 4a. It is not known whether some (CH<sub>3</sub>)<sub>3</sub>NBH<sub>3</sub> formed early in the process or exclusively at later stages. The hydrolysis (5) also lacks quantitative neatness, since 10H<sub>2</sub> would be expected from each B<sub>4</sub>H<sub>8</sub> group. The H<sub>2</sub> deficiency could be explained partly by the transfer of one H from B to C and partly by the retention of some B-B bonds, but there is no direct evidence whereby to judge the relative importance of these effects. The dimethyl ether reaction (6), occurring during 40 hr. at  $-78^{\circ}$ , seemed neater; however the main product was nonvolatile and uncertain as to content of (CH<sub>3</sub>)<sub>2</sub>O, nearly all of which could be removed in no specific steps as the mixture warmed slowly to room temperature. Traces of unidentified polyboranes also were evolved (leaving a resinous product very different from the original B<sub>4</sub>H<sub>8</sub>CO), but under no conditions was there any such main formation of polyboranes as in the  $B_5H_{11}$ -(CH<sub>3</sub>)<sub>2</sub>O reaction.<sup>7</sup>

The ethylene reaction (7) seemed much like the corresponding  $BH_3CO-C_2H_4$  reaction (3) except for the retention of CO in stable combination. The slightly volatile liquid product, of molecular formula as written,

<sup>(1)</sup> A. B. Burg and J. R. Spielman, J. Am. Chem. Soc., 81, 3479 (1959).

<sup>(2)</sup> A. B. Burg, Eighth Annual Technical Report to the Office of Naval Research, Nov., 1954, p. 12.

<sup>(3)</sup> R. Schaeffer, ARL Technical Report 60-334, Contract No. 33(616)-5827 (1960), p. 15.

<sup>(4)</sup> J. A. Dupont and R. Schaeffer, J. Inorg. Nucl. Chem., 15, 310 (1960).

<sup>(6)</sup> A. B. Burg and H. I. Schlesinger, *ibid.*, **59**, 780 (1937).

<sup>(7)</sup> J. L. Boone and A. B. Burg, ibid., 81, 1766 (1959).



Fig. 1.—Column for removal of  $B_{\delta}H_{\vartheta}$  from  $B_{\delta}H_{11}$ . A nearly total reflux under high vacuum is maintained by Dry Ice (in isohexanes) in the large outer chamber, from which the inner chamber is air-gap insulated to aid development of a thermal gradient favoring separation. External insulation is by glass wool. For temperatures down to  $-100^{\circ}$  a pump connects to the upper right opening.

proved to be unstable in the sense of forming hydrogen and a nonvolatile oil, but no carbon monoxide.

The permanent capture of the CO unit in such reactions prevented the use of  $B_4H_8CO$  for further attempts to make polyboranes such as  $B_5H_9$ ,  $B_6H_{10}$ , or B<sub>10</sub>H<sub>14</sub> by the action of basic reagents.<sup>8</sup> Indeed, no simply connected adduct of  $B_4H_8$  with a base lacking the electron acceptor character of PF<sub>3</sub> or CO seems yet to have been clearly demonstrated. On the other hand, base adducts of B<sub>3</sub>H<sub>7</sub> seem stable enough,<sup>9</sup> whereas our best attempts failed to demonstrate any B<sub>3</sub>H<sub>7</sub>CO as a byproduct of the synthesis of B<sub>4</sub>H<sub>8</sub>CO from either B<sub>4</sub>H<sub>10</sub> or  $B_5H_{11}$ . Two distinctly different approaches by Schaeffer also failed to produce any B<sub>3</sub>H<sub>7</sub>CO.<sup>3</sup> It is suggested that the dissociation  $B_4H_{10} \rightarrow B_3H_7 + BH_3$  makes good progress only in the presence of a base which can combine firmly with the very strong Lewis acid  $B_3H_7$ ; otherwise the reaction  $B_4H_{10} \rightarrow B_4H_8 + H_2$  is more probable.<sup>4</sup> Indeed, we find that B<sub>4</sub>H<sub>10</sub> has no appreciable influence on the decomposition of B<sub>4</sub>H<sub>8</sub>CO, showing that BH<sub>3</sub> is not nearly so readily available from  $B_4H_{10}$  as from  $B_2H_6$ .

Speculations concerning the molecular structure of  $B_4H_8CO$  are aided by its chemical behavior. For example, the easy loss of CO in some reactions, such as the cleanly reversible  $B_4H_8CO + PF_3 \rightleftharpoons B_4H_8PF_3 + CO$  (for the forward reaction, see reference 3, pp. 21–22), argues strongly against any B–C–O–B pattern like the B–C–C–B bonding in  $B_4H_8C_2H_4$ .<sup>10</sup> It is reasonable to suggest that both CO and PF<sub>3</sub> act as  $\sigma$ -bond donors to boron and as  $\pi$ -acceptors for electrons in the  $B_4H_8$  unit. Such a system of weak  $\pi$ -bonds, wherein  $B_4H_8$  electrons interact with  $C_{2p}$  and  $P_{3d}$  orbitals, would be like the  $\pi$ -bonding which we assume to occur in BH<sub>3</sub>CO and BH<sub>3</sub>PF<sub>3</sub>, accounting for their low polarity and high volatility.

The infrared spectra of  $B_4H_8CO$  and  $B_4D_8CO$  include a C==O stretching band closely similar to that in  $BH_3$ -CO, suggesting a very similar kind of B-C-O bonding. The B<sup>11</sup> nuclear magnetic resonance spectrum is even more indicative of structure, suggesting a three-way proton-shift tautomerism like that found in  $B_5H_{11}$ .<sup>11</sup> However, a reliable determination of the dominant structure remains a problem for the future.

## Syntheses

The Reaction of Pentaborane-11 with CO.—The quantitative reaction  $B_{5}H_{11} + 2CO \rightarrow B_{4}H_{8}CO + BH_{8}CO$  has been mentioned briefly<sup>1</sup> but requires fuller description. To make pure  $B_{4}H_{8}CO$ (free of  $B_{5}H_{9}$ ) by this method, one must begin with pure  $B_{5}H_{11}$ . This was obtained from a flow-process product<sup>12</sup> by the use of the small and relatively simple fractionating column shown in Fig. 1. Designed for work at the lowest possible pressures, this column used a wire-spiral packing and operated under a high reflux ratio, with Dry Ice in hexane ( $-78^{\circ}$ ) in the reflux head. The  $B_{5}H_{9}$  impurity slowly escaped to a high-vacuum trap at  $-196^{\circ}$ , behaving as the more volatile component of the liquid even though solid  $B_{5}H_{9}$  probably is less volatile than liquid  $B_{8}H_{11}$ at  $-78^{\circ}$ . Our purest  $B_{6}H_{11}$  melted at  $-122.0 \pm 0.1^{\circ}$  (ear ier value,  $-123.3^{\circ})^{5}$ ; vapor tensions 52.8 mm. at  $0^{\circ}$ , 103.1 mm. at

<sup>(8)</sup> J. L. Boone and A. B. Burg, J. Am. Chem. Soc., 80, 1519 (1958).

 <sup>(9) (</sup>a) W. V. Hough, M. D. Ford, G. T. Hefferan, and L. J. Edwards,
 U. S. Atomic Energy Comm. Rept. CCC-1024-TR-274 (1957); (b) G.

Kodama, R. W. Parry, and J. D. Carter, J. Am; Chem; Soc., 81, 3534 (1959).
 (10) B. C. Harrison, I. J. Solomon, R. D. Hites, and M. J. Klein, J. Inorg.

<sup>(10)</sup> D. C. Anthenson (1, 9, 600 minute), 1.22 minute, 1.22 mi

<sup>(11)</sup> R. E. Williams, ibid., 20, 198 (1961).

<sup>(12)</sup> A. B. Burg and F. G. A. Stone, J. Am. Chem. Soc., 75, 228 (1953).

13.5°, and 116.1 mm. at 16.1°. Its  $B^{11}$  n.m.r. spectrum showed no trace of the spurious peak attributed to  $B_{\delta}H_{\vartheta}$  impurity.<sup>13</sup>

The conversion of B<sub>5</sub>H<sub>11</sub> to B<sub>4</sub>H<sub>8</sub>CO required a great excess of dry, oxygen-free carbon monoxide. This was made by warming a mixture of formic and sulfuric acids in an evacuated flask attached directly to the high-vacuum system, wherein it could be condensed and redistilled. For such distillations the evaporation was from traps immersed in liquid nitrogen  $(-196^{\circ})$  and the receiving traps were cooled by pumped liquid nitrogen which had attained temperatures as low as  $-210^{\circ}$ . After being measured as a gas, the CO could be condensed almost completely in the reaction tube at  $-210^{\circ}$ , and that which remained in the apparatus after the seal-off could be collected by an automatic Sprengel pump for measurement over mercury. Then, after the bombtube reaction, the CO could be distilled out of the reaction tube (opened in vacuo) and measured back by the same methods, for a rough estimate of the consumed CO. In one experiment a top fraction of the CO was collected by means of the Sprengel pump, and CuO combustion showed less than 0.01 H<sub>2</sub> per B<sub>5</sub>H<sub>11</sub> originally present.

The stoichiometry of two such quantitative experiments has been presented.<sup>1</sup> Times no longer than 2.5 hr. at room temperature were sufficient for complete reaction. Longer times led to lower yields because the decomposition of  $B_4H_8CO$  to nonvolatile solids is not reversible, nor entirely inhibited by carbon monoxide. This method of making  $B_4H_8CO$  is fast and it virtually proved the formula  $B_4H_8CO$  (confirmed by analysis),<sup>1</sup> but the tedium of making pure  $B_6H_{11}$  for such use might lead to a preference for the  $B_4H_{10}$  method.

The Reaction of Tetraborane-10 with CO.—For the reaction  $B_4H_{10} + CO \rightarrow H_2 + B_4H_8CO$ , pure  $B_4H_{10}$  was made from  $B_5H_{11}^{5,12}$  and measured as a gas. The sample was condensed from the high-vacuum system into a 230-ml. stainless-steel pressure cylinder (having a Teflon-seated safety valve); then a much larger measured sample of CO was condensed into the cylinder by means of pumped-down liquid nitrogen. In most experiments relatively high CO pressures were used in order to minimize the decomposition of  $B_4H_8CO$  during the relatively long times which were required for extensive reaction.

The conditions and results of our most informative experiments are presented in Table I. The reaction ran at mean tempera-

TABLE I THE REACTION OF  $B_4H_{10}$  with CO

	~ Volatiles recovered,									
Expt.	$B_4H_{10}$	$B_4H_{10}$ , Pco,			mmoles			104k,		
no.	mmoles	atm.	hr.	$B_4H_{10}$	BH3CO	B <sub>4</sub> H <sub>8</sub> CO	yield	min1		
1	2.68	10	55	2.01	0.02	0.27	40	0.87		
<b>2</b>	3.62	9	23	2.68	.09	.36	38	2.17		
3	5.75	89	23	5.04	.45	.36	51	0.96		
4	5.45	59	62	3 79	.07	.71	43	.97		
5	3.62	56	75	2.72	.09	.67	<b>74</b>	.63		
6	2.59	53	222	1.21	.24	1.12	81	.57		
7	1.21	50	261	0.49	.17	0.49	68	.57		
8	4.53	67	235	0.56	1.06	1,44	<b>27</b>	1.49		
9	4.97	71	66	2.35	2.00	1.40	54	1.91		

tures in the range 20-30°, with rates varying accordingly. In expt. 9 the formation of  $H_2$  was roughly 2 mmoles, exceeding that required by the formation of  $B_4H_8CO$  according to the ideal equation  $B_4H_{10} + CO \rightarrow H_2 + B_4H_8CO$ ; the difference is attributable to a parallel decomposition of  $B_4H_{10}$ . Experiments 8 and 9 began with 1.17 and 2.70 mmoles of  $BH_3CO$ , respectively, resulting in an indication that this component is moderately consumed if present at the start, while other experiments show that some  $BH_3CO$  is formed if not initially present. In each case the per cent yield is based upon the consumed  $B_4H_{10}$ .

Although these experiments (of which 1–7 were performed in the years 1951–1954) were not planned as accurate rate and mechanism studies, the values here found for  $10^4k$ , based upon the equation  $-d(B_4H_{10})/dt = k(B_4H_{10})$  (independent of CO

pressure), are quite reasonably consistent with Schaeffer's flowprocess rate studies at 80–110°, with CO pressures around 0.5 atm.<sup>3</sup> From his results 10<sup>4</sup>k can be estimated as 0.55 at 20° and 2.21 at 30°, in close agreement with our results in this temperature range. Thus it is confirmed that the rate is independent of extreme variation of pressure of carbon monoxide, and it is evident that Schaeffer's results are dependable for long-range extrapolation.

Synthesis of Deuterioboron Compounds.—For infrared spectroscopic comparisons it was important to have samples of  $B_2D_6$ ,  $B_4D_{10}$ ,  $B_5D_{11}$ , and  $B_4D_5CO$ . Accordingly, a 15.3-mmole sample of  $B_2D_6$  was made from LiD and  $BF_3$ .<sup>14</sup> Its initial D-content was estimated as 98% by the intensity of the infrared band at 2585 cm.<sup>-1</sup> in relation to that of the 2705 cm.<sup>-1</sup> band, using published data<sup>15</sup> for calibration. Then the D content was raised above 99% by two equilibrations (sealed bulbs, 20 hr. at 65°) with 99.5+% D<sub>2</sub>. The vacuum-fractionated sample showed a 238.0-mm. vapor tension at  $-111.9^{\circ}$ , in agreement with an earlier estimate.<sup>16</sup>

This  $B_2D_6$  sample was employed in a miniature vapor-flow apparatus to produce 0.42 mmole of  $B_4D_{10}$  and 1.50 mmoles of  $B_5D_{11}$ . Treatment of 0.964 mmole of this  $B_5D_{11}$  with high-pressure CO gave 0.965 mmole of  $BD_3CO$  (57 mm. at  $-102^\circ$ ; calcd., 55 mm.)<sup>16</sup> and  $B_4D_8CO$  presumably in similar yield.

**Physical Properties.**—The volatility of  $B_4H_8CO$  is well described by the original vapor-tension equation.<sup>1</sup> Each of the deuterioboron compounds is slightly more volatile than its protium counterpart, as shown by the comparisons in Table II.

 TABLE II

 INCREASED VOLATILITY OF DEUTERIUM COMPOUNDS

	,	Vapo	r tension va	alues, P <sub>mm</sub>	, corr	
t, °C.	$B_4H_{10}$	$B_4D_{10}$	$B_{\delta}H_{11}$	B <sub>5</sub> D <sub>11</sub>	B <sub>4</sub> H <sub>8</sub> CO	B4D8CO
0.0	387	423	52.8	56.7	71.3	74.2
-40.1	49.0	59.3	•••			• • • •
-78.5	3.2	3.8				• • •

Five new observations of the melting point of  $B_4H_8CO$  gave readings ranging from -110.3 to  $-113.2^\circ$ . The highest result was for  $B_4H_8CO$  which had been freshly made from  $B_6H_{11}$  + 2CO; older samples probably had formed some  $B_6H_{11}$ , which could be removed only by reaction with CO at high pressures. The melting points of two of the deuterium compounds were observed as slightly lower than for the protium compounds:  $B_4D_8$ -CO,  $-112.2^\circ$  (vs.  $-110.3^\circ$ );  $B_8D_{11}$ ,  $-122.9^\circ$  (vs.  $-122.0^\circ$ ). However, a trace of  $B_5D_9$  impurity might have affected both results.

#### Chemistry Involving the Tetraborane-8 Group

The New Trifluorophosphine Adduct.—The reaction of  $B_2H_6$ with PF<sub>8</sub> to give  $BH_8PF_8^{17}$  is so like the synthesis of borine carbonyl as to suggest the analogous reaction of  $B_6H_{11}$  with PF<sub>8</sub> to make  $BH_8PF_8$  and  $B_4H_8PF_8$ . In fact, this reaction proved to be quite easy and straightforward.

Phosphorus trifluoride was purified through the formation and dissociation of the adduct  $(CH_3)_8NPF_{3}$ ,<sup>18</sup> and after high-vacuum distillation from a trap at  $-160^{\circ}$  its vapor tension at  $-112^{\circ}$  was 372 mm. Two experiments on its reaction with pure  $B_6H_{11}$  are summarized in Table III (quantities in mmoles). In each experiment the reaction occurred in a sealed 25-ml. tube, so that the partial pressure of PF<sub>3</sub> was near 10 atm. The BH<sub>3</sub>PF<sub>3</sub> and unused PF<sub>8</sub> were isolated by high-vacuum fractional condensation, with identification of the former by its 77.9-mm. vapor ten-

<sup>(13)</sup> R. E. Williams, S. G. Gibbins, and I. Shapiro, J. Chem. Phys., **30**, 320 (1959). We are grateful to Dr. Williams for obtaining this spectrum for our sample.

<sup>(14)</sup> H. I. Schlesinger, H. C. Brown, J. R. Gilbreath, and J. J. Katz, J. Am. Chem. Soc., 75, 195 (1953).

<sup>(15)</sup> W. J. Lehmann, J. F. Ditter, and I. Shapiro, J. Chem. Phys., 29, 1248 (1958).

<sup>(16)</sup> A. B. Burg, J. Am. Chem. Soc., 74, 1340 (1952).

<sup>(17)</sup> R. W. Parry and T. C. Bissot, ibid., 78, 1524 (1956).

<sup>(18)</sup> J. E. Griffiths and A. B. Burg, ibid., 82, 1507 (1960).

TABLE III Synthesis of B4H8PF3

-Reactants		Temp.,	Time,	Products		PF:
$B_{\delta}H_{11}$	$PF_3$	°C.	hr.	BH3PF3	$B_4H_8PF_3$	used
0.465	9.85	32	4.0	0.437	0.424	0.87
0.651	9.95	29	2.0	0.642	0.623	1.20

sion at  $-97.6^{\circ}$  (known, 77.6 mm.)<sup>17</sup> and its molecular weight (100.6; calcd., 101.8).

The fraction which passed a high-vacuum trap at  $-78^{\circ}$  and condensed at  $-97^{\circ}$  seemed uniform, for three successive fractions showed respective vapor tensions of 6.0, 6.2, and 6.1 mm. at  $-47.7^{\circ}$ . Two vapor-phase molecular weight results (corrected for dissociation) were 136 and 141; calcd. for B<sub>4</sub>H<sub>8</sub>PF<sub>3</sub>, 139.3.

The molecular formula  $B_4H_8PF_3$ , strongly indicated by the nearly quantitative syntheses and the molecular weight results, was confirmed by a quantitative conversion to  $B_4H_8CO$ . A  $B_4H_8PF_3$  sample measured as 0.215 mmole, standing in a mixture with CO at a partial pressure of 18 atm. during 1 hr. at 28°, yielded 0.216 mmole of PF<sub>3</sub> (377 mm. at  $-112^\circ$ ) and 0.220 mmole of  $B_4H_8CO$ , identified by its 70.5-mm. vapor tension at 0° and by its infrared spectrum. The reverse reaction,  $B_4H_8CO$ + PF<sub>3</sub>  $\rightarrow$  CO +  $B_4H_8PF_3$ , also can be made nearly quantitative (ref. 3, p. 22).

The decomposition of  $B_4H_8PF_3$  is somewhat faster than that of  $B_4H_8CO$  and is not quite analogous, for less  $PF_3$  than CO is permanently trapped in the condensing solids. Thus 0.42 mmole of  $B_4H_8PF_3$ , left for 3 days at room temperature, produced a slightly volatile white solid, 0.126 mmole of  $H_2$ , 0.044 mmole of  $B_5H_9$ , and a mixture equivalent to 0.060 mmole of  $B_2H_6$  and 0.387 mmole of  $PF_3$ . This 92% recovery of the  $PF_3$  indicated that  $PF_3$  is not much concerned with the further conversion of the  $B_4H_8$  moiety, and presumably the dissociation produces no polymerizable fragment containing  $PF_3$ . Also of interest is the complete displacement of  $PF_3$  from  $B_4H_8PF_3$  by trimethylamine,<sup>3</sup> in contrast to the total retention of CO when  $B_4H_8CO$  reacts with this amine. Thus it appears that  $B_4H_8PF_3$  is the better source of  $B_4H_8$  for most chemical purposes.

Reactions Related to Polyborane Conversions .- The thermal decomposition of B<sub>4</sub>H<sub>8</sub>CO yields H<sub>2</sub>, B<sub>2</sub>H<sub>6</sub>, BH<sub>8</sub>CO, B<sub>4</sub>H<sub>10</sub>, B<sub>5</sub>H<sub>11</sub>, and nonvolatile solid material containing CO. The yield of  $B_4H_{10}$  is strongly increased by a large excess of  $H_2$ , while an abundance of B<sub>2</sub>H<sub>6</sub> leads to far higher yields of B<sub>5</sub>H<sub>11</sub>, but B<sub>4</sub>H<sub>10</sub> seems not to affect the course of the reaction very much. Since CO inhibits the dissociation  $B_4H_8CO \rightarrow B_4H_8 + CO$ , its frequent removal is necessary if the process is to go forward faster than the decomposition of such unstable products as  $B_4H_{10}$  or  $B_5H_{11}$ . Accordingly, experiments 1, 3, and 4 of Table IV were interrupted at 5-min. intervals for removal of CO and H<sub>2</sub> through a  $-196^{\circ}$  trap by means of an automatic Sprengel pump. At some of these intervals and at the end there was a complete resolution of the products: the CO and H<sub>2</sub> were collected for measurement over mercury and analyzed by CuO combustion, and the other volatile components were separated into three main fractions by high-vacuum fractional condensation. The  $B_2H_6$ -BH<sub>3</sub>CO fraction was analyzed by allowing the  ${\rm BH}_{\delta}{\rm CO}$  to dissociate to  ${\rm B}_{2}{\rm H}_{\delta}$ and CO. Tetraborane was isolated directly. The B5H11-B<sub>4</sub>H<sub>8</sub>CO fraction was analyzed by exposure to water during 90 sec. at 0°, whereby each  $B_5H_{11}$  formed one  $B_4H_{10}$ <sup>8</sup>; in the meantime the hydrolysis of  $B_4H_8CO$  yielded no volatiles except  $H_2$  and a trace of CO, and any  $B_{\delta}H_{9}$  would have been observed as such, but could not be found.

The experiments described by Table IV employed a 130-ml. Pyrex tube with a stopcock protected by mercury to avoid the otherwise quite noticeable reaction between Apiezon L grease and  $B_4H_8CO$ . After each interval the mixture was warmed rapidly (within 30 sec.) from  $-196^\circ$  to room temperature and rapidly quenched at  $-196^\circ$  at the end of the interval.

In expt. 2 it is evident that the large proportion of  $H_2$  caused an almost complete conversion of  $B_4H_8$  to  $B_4H_{10}$ , if we allow for some decomposition of this product during the process. In con-

TABLE IV B<sub>4</sub>H<sub>6</sub>CO Dissociation Reactions  $\mathbf{2}$ Experiment no. 1 3 4 B<sub>4</sub>H<sub>8</sub>CO, mmoles 0.3540.2360.4070.673Added reagent, per B<sub>4</sub>H<sub>8</sub>CO  $14B_2H_6$  $13B_4H_{10}$ None  $41H_2$ Time, total min. 2316 30 27Av. temp., °C. 26252626Per cent B<sub>4</sub>H<sub>8</sub>CO 78consumed 39 94 86 Yields of prod. per 100B<sub>4</sub>H<sub>8</sub>CO consumed: 63 CO 85 5759 $H_2$ 39 -89<sup>a</sup> 7 1517 $-34^{a}$  $B_2H_6$ 1819BH<sub>3</sub>CO 4 Nil 10 $\mathbf{2}$ 2167 4  $-1^{a}$  $B_4H_{10}$  $B_5H_{11}$ 28Nil 7528

<sup>a</sup> Material consumed.

trast, expt. 3 afforded little  $H_2$  for the conversion to  $B_4H_{10}$ ; instead, the large proportion of  $B_2H_6$  caused almost a complete conversion to  $B_5H_{11}$ , some of which must have decomposed. In expt. 4 the slight loss of  $B_4H_{10}$  would be ascribed to the decomposition of slightly more of it than was formed.

### Additions of $B_4H_8CO$ to Electron-Rich Reactants

In its reactions with basic reagents (or even ethylene)  $B_4H_8CO$ never liberates more than a trace of CO. Apparently there is a direct attack at a rate far faster than the dissociation  $B_4H_8CO \rightarrow$  $B_4H_8 + CO$ . Hence we suppose that the first step is the formation of an adduct of  $B_4H_8CO$ , and then it can be argued that the resulting increase of electron density in the  $B_4H_8$  part would increase the hydridic activity of some B-H bonds enough for a rapid attack upon the CO part. Such an enhancement of hydridic activity through complex formation is well known for BH<sub>3</sub> complexes,<sup>19</sup> and it would seem that just such a more effective hydridic tendency would explain why the methyldiboranes react with CO not to form a methylated borine carbonyl, but to destroy all B-H bonds and capture the CO,<sup>20</sup> probably forming  $(-B-C-O-)_2$  rings.

The Trimethylamine Reaction.—A 0.790-mmole sample of  $B_4H_8CO$  absorbed all of a 0.817-mmole sample of  $(CH_8)_8N$  during 6 days at  $-78^\circ$  (in a 30-ml. tube with a stopcock attached to the vacuum system), with an apparently accidental liberation of 0.036 mmole of  $B_2H_6$  and 0.031 mmole of  $BH_2CO$  and formation of a slightly yellow solid. A further 1.750-mmole sample of  $(CH_3)_8N$  was partially absorbed as shown in Table V. The only

	TAB	LE V	
Absor	ption of $(CH_3)_3N$	ву 0.790 ммоl	e of B <sub>4</sub> H <sub>8</sub> CO
Time interval, h <b>r</b> .	Interim temp., °C.	Total amine absorbed, mmoles	Ratio (CH3)3N/B4H8CO
144	-78	0.817	1.034
511	-78	1.553	1.96
479	-78	1.703	2.16
25	Warming to 0°	1.903	2.41
15	25	1.930	2.44
930	25	1.969	2.49

volatile product now was 0.581 mmole of  $(CH_3)_8 NBH_3$ , which was isolated by sublimation from the nonvolatile glassy residue. Its formation from this CO-containing polyborane is reminiscent of

<sup>(19)</sup> A. B. Burg, Angew. Chem., **72**, 185 (1960); XVIth International Congress of Pure and Applied Chemistry, Butterworths, London, 1960, p. 44.

<sup>(20)</sup> F. L. McKennon, Ph.D. Dissertation, University of Chicago Libraries, 1937.

the  $(CH_3)_8N-B_8H_9$  reaction, which forms  $2(CH_3)_8NBH_8$  per  $B_5H_9$ ; but this does not necessarily mean that a BH<sub>8</sub> pattern is to be found in the structure of  $B_4H_8CO$ .

The Hydrolysis.—Measured samples of  $B_4H_8CO$  were allowed to warm to room temperature with vacuum-degassed pure water, in stopcock tubes attached to the high-vacuum system. The evolved  $H_2$  and CO were collected for measurement and analyzed by CuO combustion, with the results shown in Table VI. The

TABLE VI THE HYDROLYSIS OF B4H8CO

Expt. no.⇒	B <sub>4</sub> H <sub>8</sub> CO, mmoles	CO, mmoles	H2, mmoles	Ratio H2/B4H8CO
1	0.13	0.008	1.14	8.8
<b>2</b>	0.318	0.012	2.80	8.7
3	0.426	0.015	3.73	8.8

only other volatile component was water. The white solid residue was dried *in vacuo* at  $110^{\circ}$  and mulled with hexachlorobutadiene, for an infrared spectrum which showed only the bands of the mull liquid and of boric acid.<sup>21</sup> One of the three residues was heated for 1 hr. with aqueous HCl at 90°, giving only negligible further evolution of hydrogen.

The Dimethyl Ether Reaction.—An initial experiment used 0.266 mmole of  $B_4H_8CO$  with 1.33 mmoles of  $(CH_3)_2O$ , reacting during 4 days in a 15-ml. tube at  $-78^\circ$ . Slow warming to  $-26^\circ$  permitted recovery of 1.15 mmoles of the ether and a trace of an unidentified boron hydride, leaving a yellow nonvolatile solid. A fresh measured sample of  $(CH_3)_2O$  was added and withdrawn in stages at  $-78^\circ$ , in an attempt to recognize a definite etherate, but as the ether content diminished the pressure at  $-78^\circ$  only decreased smoothly, except for a fairly sharp drop from 12.9 to 5.8 mm. as the ratio of the ether to  $B_4H_8CO$  dropped from 0.86 to 0.78. It appeared that the  $B_4H_8CO$  had been converted to a nonvolatile solid which combined only very weakly and indefinitely with the dimethyl ether.

In a second experiment,  $B_4H_8CO$  was treated with  $(CH_3)_2O$ in an apparatus designed to measure the molecular weight of the solid product by the vapor tension lowering method.<sup>22</sup> With dimethyl ether as the solvent at  $-26^\circ$  the results at mole fractions 0.064 and 0.032 were 192 and 189, respectively, indicating an average molecular weight about 2.4 times that of  $B_4H_8CO$ .

The Ethylene Reaction.-In a typical experiment, 0.528 mmole of B<sub>4</sub>H<sub>8</sub>CO, stabilized by 2.08 mmoles of CO, was allowed to react at room temperature with 9.47 mmoles of C2H4 in a 24ml. sealed tube ( $C_2H_4$  pressure nearly 10 atm.) for 2 hr. Then the mixture was resolved into 0.014 mmole of H<sub>2</sub>, 2.13 mmoles of CO (1% liberation from B<sub>4</sub>H<sub>8</sub>CO), 7.62 mmoles of C<sub>2</sub>H<sub>4</sub> (consumed, 1.85 mmoles), 0.142 mmole of (C<sub>2</sub>H<sub>5</sub>)<sub>3</sub>B, and approximately 0.25 mmole of a new volatile compound, the molecular formula of which proved to be  $(C_{2}H_{4})_{4}B_{4}H_{8}CO$ . The low yield of this (not quite 50%) could be explained by its instability, for on standing at room temperature it forms a nonvolatile oil, such as was observed in major proportion. A purified sample of the new compound had 0.5 mm. vapor tension at 25°; values at higher temperatures were not sought because the decomposition was too rapid. Indeed, its tendency to form more volatile material was such that it could be distilled quantitatively only by pumping continually beyond the receiving trap. On exposure to the open air, it demonstrated an aromatic or fruity odor, very different from the typical boron hydride odor.

The formula  $(C_2H_4)_4B_4H_8CO$  was confirmed by analysis, using the "wet combustion" method: oxidation by nitric acid in a sealed soda-lime glass tube at 270°. The resulting CO<sub>2</sub> was passed over hot copper to reduce the oxides of nitrogen and through a trap at  $-78^{\circ}$  to remove water, but the infrared spectrum showed the presence of some N<sub>2</sub>O. For separation, the CO<sub>2</sub> was absorbed in a slight excess of 0.2 N Ba(OH)<sub>2</sub>, so that the N<sub>2</sub>O could be isolated and measured, and then the CO<sub>2</sub> was liberTHE TETRABORANE CARBONYL B<sub>4</sub>H<sub>8</sub>CO 1143



Fig. 2.—The  $B^{11}$  magnetic resonance spectrum of  $(C_2H_4)_4B_4H_8CO$ .

ated by aqueous HCl and measured, for a recheck of the result obtained by subtracting the  $N_2O$  from the measured volume of the mixture. The boric acid remaining in the original reaction tube was titrated, using the usual mannitol method. Of the results shown in Table VII, those from the largest sample (15.4 mg.) are regarded as the most dependable.

TABLE VII Analysis of  $(C_2H_4)_4B_4H_8CO$ 

Sample weight, mg.		7.1	15.4
mmoles of CO <sub>2</sub> found	0.234	0.334	0.724
calcd.		.334	. 724
mmoles of B(OH) <sub>3</sub> found	1 0.102	.176	.330
	•••	.148	.322
Ratio C:B found	2.29	1.90	2.19
calcd.	2.25	2.25	2.25

The molecular weight determination was attempted by the vapor-tension lowering method, using as the solvent very pure *n*-butane which had been made by the Grignard method (vapor tension 781 mm. at 0°); however, the result was low (159 vs. calcd. 191) because of traces (estimated less than 11 mole %) of more volatile material which developed during the measurements at 0°.

The infrared spectrum of  $(C_2H_4)_4B_4H_8CO$  in solution in cyclohexane showed a strong band at 2121 cm.<sup>-1</sup>. This might represent the C=O stretching frequency (2150 cm.<sup>-1</sup> in B<sub>4</sub>H<sub>8</sub>CO), but it cannot be distinguished from a possible symmetric in-phase B-H-B bridge frequency; a more complete spectrum is needed, but cannot be obtained dependably except by special low-temperature methods. B-H stretching was observed at 2480 cm.<sup>-1</sup> and C-H bending at 1370 cm.<sup>-1</sup>.

The B<sup>11</sup> n.m.r. spectrum of pure liquid  $(C_2H_4)_4B_4H_8CO$  (run at 12.83 Mc.) is shown in Fig. 2, wherein the chemical shifts relate to boron trifluoride diethyl etherate as a reference zero. The doublet centered at  $+53\delta$  represents apical B-H. The peaks at  $-8.8\delta$  and  $+2.3\delta$  might represent a doublet (J = 140 c.p.s.) with one peak raised by overlap with the  $-16.7\delta$  peak, or two singlets. Any conclusion concerning structure would be premature. For example, the ethylene might go in to become a  $C_2H_5$  group (in good analogy to the BH<sub>3</sub>CO reaction with ethylene) or at least some of it might form B-C<sub>2</sub>H<sub>4</sub>-B connections.

Borine Carbonyl with Ethylene.—The reaction of  $BH_3CO$  with  $C_2H_4$  is a typical hydroboration,<sup>28</sup> almost quantitatively forming  $(C_2H_5)_3B$  and CO. The numerical data of three such experiments are given by Table VIII. All of these occurred near room temperature. Direct observation in the case of expt. 3 showed a

<sup>(21)</sup> R. R. Servos and H. M. Clark, J. Chem. Phys., 26, 1175 (1857).

<sup>(22)</sup> A. B. Burg, J. Am. Chem. Soc., 65, 1630 (1943).

<sup>(23)</sup> H. C. Brown and B. C. Subba Rao, ibid., 81, 6423 (1959).

The Reaction of Borine	CARBONYL	<b>WITH ETH</b>	YLENE
Expt. no.	1	$^{2}$	3
Container volume, ml.	24	115	50
BH₃CO, mmoles	$0.407^{a}$	0.750	0.544
C <sub>2</sub> H <sub>4</sub> , mmoles	9.07	3.79	1.71
Time, min.	120	30	5
C <sub>2</sub> H <sub>4</sub> consumed, mmoles	1.23	1.99	1.47
Products, mmoles			
$(C_2H_5)_3B$	0.399	0.390	0.359
CO	0.383	0.526	0.535
$H_2$	Nil	0.310	0.190

TABLE VIII

<sup>*a*</sup> Corrected for 0.039 mmole of  $B_2H_6$  impurity, as demonstrated by subsequent treatment of the ethylene fraction with  $(CH_3)_8N$ ; by this method no  $B_2H_6$  was found after expt. 2 and 3.

fairly sudden precipitation of white droplets 2.5 min. after the tube was warmed to  $25^{\circ}$ . Table VIII does not include unidentified volatile products resulting from expt. 2 and 3, wherein the relatively low pressure of  $C_2H_4$  (near 1 atm., vs. 9 atm. in expt. 1) permitted extensive side reactions correlated with the formation of H<sub>2</sub>. One volatile product showed 50 mm. vapor tension at 0° (mol. wt. 89); another, 0.9 mm. at  $25^{\circ}$ . Nonvolatile products also were observed in these experiments.

The product triethylboron was recognized by its vapor tension (12.5 mm. at  $0^{\circ}$ )<sup>24</sup> and mol. wt. (98.4; calcd., 98.0); also its nearly quantitative formation in expt. 1 left no doubt.

#### Instrumental Approaches to Structure

Infrared Spectra.—The infrared spectra of ordinary and completely deuterated  $B_4H_{10}$ ,  $B_4H_{11}$ , and  $B_4H_8CO$ were recorded in the sodium chloride range by a Perkin Elmer Infracord spectrophotometer, whereby complete scans occurred fast enough to avoid appreciable effects of decomposition. The path length was 10 cm. and the pressures approximately 20 mm. Calibration by polystyrene film reduced the probable error to 1%.

The spectra of the six samples are compared in Table IX, with suggested assignments based upon correla-

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ratios R show which bands represent BH and BD modes. Although some assignments remain uncertain, the spectra do give structural information.

The two B-H stretching bands and the scissors-bending frequency (1160 cm.<sup>-1</sup>) show that  $B_4H_8CO$  has at least one BH<sub>2</sub> group. The very strong band at 2150 cm.<sup>-1</sup>, unchanged by deuteration, compares directly with the known C-O stretching frequencies of CO (2143 cm.<sup>-1</sup>) and BH<sub>3</sub>CO (2165 cm.<sup>-1</sup>).<sup>27</sup> It is reasonable that the B-C-O bonding in BH<sub>3</sub>CO and B<sub>4</sub>H<sub>8</sub>CO is nearly the same, with carbon bonded only to one boron atom. Another mode not much affected by deuteration is the B-B stretching, indicated by the shoulder at 1080 cm.<sup>-1</sup> for  $B_4H_8CO$  and 975 cm.<sup>-1</sup> for  $B_4D_8CO$ . The comparison seems analogous to the B<sub>5</sub>H<sub>9</sub>-B<sub>5</sub>D<sub>9</sub> case, for which B-B stretching has been recognized at 1126 and 960 cm.<sup>-1, 28</sup> Less certain are the assignments for the 899 and 804 cm.<sup>-1</sup> bands for B<sub>4</sub>H<sub>8</sub>CO. The absence of similar bands for  $B_4D_8CO$  probably means that these would have been found if the scan could have gone below 670 cm.<sup>-1</sup> and then would have been assigned to a BH<sub>2</sub> in-plane rocking mode. On the whole, it appears that the B<sub>4</sub>H<sub>8</sub> unit suffered no extreme structural change on conversion from its environment in  $B_5H_{11}$  to that in  $B_4H_8CO$ .

Nuclear Magnetic Resonance.—The B<sup>11</sup> n.m.r. spectra of "neat" liquid  $B_4H_8CO$  and  $B_4D_8CO$  were obtained by means of a Varian V-4300 high-resolution spectrometer operating at 12.83 Mc., through the courtesy of National Engineering Science Co., with the operation directed by Dr. R. E. Williams. The chemical shift and spin coupling values were measured by reference to triethylboron and boron trichloride as secondary standards, in turn referred to boron trifluoride etherate as zero. The first traces were made as

				TABLE	IX			
		Comparison	s of Infrare	d Spectr.	A OF POLYBORA	NE COMPOU	$NDS^{a,b}$	
$B_4D_{10}$	R	BBH11	$B_{\delta}D_{11}$	R	B <sub>4</sub> H <sub>8</sub> CO	B4D8CO	R	Suggested assignments
1950 vs	1.33	2600 vs	1960 vs	1.33	2570  vs	1940  vs	1.32	B—H asymm. stretch
1830 vs	1.37	2500  vs	1840 vs	1.36	2490  vs	1830 vs	1.36	B—H symm. stretch
1730 w	1.32	2330 w			2320 w	1750 w	1.33	$2 \times BH_2$ scissors
					2150  vvs	2150  vvs	1.00	C==O stretching
1580 s	1.37	2050 s	1515  m	1.35	1940 s	1440 s	1.35	
		1910 w	1400 w,sh	1.36	1810 w,sh			
		•••						Hydrogen bridge modes
1360 w	1.36		• • • •		1510 s	1110 s,sh	1.36	
1040  m	1.35	1440 s,b	10 <b>9</b> 0 s,b	1.32	1440 m	1080 s	1.34	
960 w	1.31							3
854 s	1.35	1170 vs	850 s	1.38	1160 s	880 m	1.32	BH <sub>2</sub> scissors bend
844 s	1.35	•••	• • •		1056 vs	815 vs	1.30	B—-H bending
805 s	1.34	· · · · ·						
		1050 s	1030 m,sh	1.02	1080 sh	975 m	1.10	B-B stretching
$725 \mathrm{m}$	1.34	960 s	760 m,b	1.26	944 w	700  mw	1.34	BH <sub>2</sub> wagging
(Off-sc	ale)	905 s,vb	690 w,vb	1.31	.899 ms	(Off-sca	le)	BH <sub>2</sub> in-plane rocking?
(Off-sc	ale)				<b>8</b> 04 m	(Off-sca	1e)	Uncertain
	$\begin{array}{c} B_4 D_{10} \\ 1950 \ vs \\ 1830 \ vs \\ 1730 \ w \\ \dots \\ 1580 \ s \\ \dots \\ 1360 \ w \\ 1040 \ m \\ 960 \ w \\ 854 \ s \\ 844 \ s \\ 805 \ s \\ \dots \\ 725 \ m \\ (Off-sc \\ (Off-sc \\ (Off-sc \\ )) \end{array}$	$\begin{array}{cccccccc} B_4 D_{10} & R \\ 1950 \ vs & 1.33 \\ 1830 \ vs & 1.37 \\ 1730 \ w & 1.32 \\ \dots & \dots \\ 1580 \ s & 1.37 \\ \dots & \dots \\ 1360 \ w & 1.36 \\ 1040 \ m & 1.35 \\ 960 \ w & 1.31 \\ 854 \ s & 1.35 \\ 844 \ s & 1.35 \\ 844 \ s & 1.35 \\ 805 \ s & 1.34 \\ \dots & \dots \\ 725 \ m & 1.34 \\ (Off-scale) \\ (Off-scale) \\ (Off-scale) \end{array}$	$\begin{array}{c ccccc} COMPARISON\\ B_4 D_{10} & R & B_5 H_{11}\\ 1950 \ vs & 1.33 & 2600 \ vs \\ 1830 \ vs & 1.37 & 2500 \ vs \\ 1730 \ w & 1.32 & 2330 \ w \\ \dots & \dots & \dots \\ 1580 \ s & 1.37 & 2050 \ s \\ \dots & 1910 \ w \\ \dots & \dots & 1910 \ w \\ \dots & \dots & 1910 \ w \\ \dots & \dots & 1360 \ w & 1.36 & \dots \\ 1040 \ m & 1.35 & 1440 \ s,b \\ 960 \ w & 1.31 & \dots \\ 1364 \ s & 1.35 & 1170 \ vs \\ 844 \ s & 1.35 & \dots \\ 805 \ s & 1.34 & \dots \\ \dots & 1050 \ s \\ 725 \ m & 1.34 & 960 \ s \\ (Off-scale) & 905 \ s,vb \\ (Off-scale) & \dots \end{array}$	COMPARISONS OF INFRARE $B_4D_{16}$ $R$ $B_8H_{11}$ $B_8D_{11}$ 1950 vs         1.33         2600 vs         1960 vs           1830 vs         1.37         2500 vs         1840 vs           1730 w         1.32         2330 w            1730 s         1.37         2050 s         1515 m           1580 s         1.37         2050 s         1515 m                 1580 s         1.37         2050 s         1515 m                 1360 w         1.36             1360 w         1.35         1440 s,b         1090 s,b           960 w         1.31               1050 s         1030 m,sh           925 s         1.34               1050 s         690 w, vb	TABLE           COMPARISONS OF INFRARED SPECTR. $B_4D_{10}$ $R$ $B_8H_{11}$ $B_8D_{11}$ $R$ 1950 vs         1.33         2600 vs         1960 vs         1.33           1830 vs         1.37         2500 vs         1840 vs         1.36           1730 w         1.32         2330 w             1.580 s         1.37         2050 s         1515 m         1.35                  1580 s         1.37         2050 s         1515 m         1.35                   1580 s         1.37         2050 s         1515 m         1.35                   1360 w         1.36               1040 m         1.35         1440 s,b         1090 s,b         1.32           960 w         1.31               135         1170 vs         850 s         1.38 <td>TABLE IX           COMPARISONS OF INFRARED SPECTRA OF POLYBORA           <math>B_4D_{16}</math> <math>R</math> <math>B_8H_{11}</math> <math>B_6D_{11}</math> <math>R</math> <math>B_4H_8CO</math>           1950 vs         1.33         2600 vs         1960 vs         1.33         2570 vs           1830 vs         1.37         2500 vs         1840 vs         1.36         2490 vs           1730 w         1.32         2330 w          2320 w              2150 vvs         1840 vs         1.36         2490 vs           1730 w         1.32         2330 w          2320 w          2320 w               2150 vvs         150 vvs           1580 s         1.37         2050 s         1515 m         1.35         1940 s                    1360 w         1.36                1360 w         1.35         1440 s,b         1090 s,b         1.32         1440 m           960 w         1.31        </td> <td>TABLE IX           COMPARISONS OF INFRARED SPECTRA OF POLYBORANE COMPOUND           <math>B_4D_{16}</math>         R         <math>B_6H_{11}</math> <math>B_8D_{11}</math>         R         <math>B_4H_6CO</math> <math>B_4D_9CO</math>           1950 vs         1.33         2600 vs         1960 vs         1.33         2570 vs         1940 vs           1830 vs         1.37         2500 vs         1840 vs         1.36         2490 vs         1830 vs           1730 w         1.32         2330 w          2320 w         1750 w              2150 vvs         2150 vvs         2150 vvs           1580 s         1.37         2050 s         1515 m         1.35         1940 s         1440 s                    1360 s         1.37         2050 s         1515 m         1.36         1810 w,sh                     1360 w         1.36                1360 w         1.35         1440 s,b         1090 s,b         1.</td> <td>TABLE IX           COMPARISONS OF INFRARED SPECTRA OF POLYBORANE COMPOUNDS<sup>a,b</sup> <math>B_4D_{16}</math> <math>R</math> <math>B_8H_n</math> <math>B_6D_n</math> <math>R</math> <math>B_8H_9CO</math> <math>B_4D_9CO</math> <math>R</math>           1950 vs         1.33         2600 vs         1960 vs         1.33         2570 vs         1940 vs         1.32           1830 vs         1.37         2500 vs         1840 vs         1.36         2490 vs         1830 vs         1.36           1730 w         1.32         2330 w          2320 w         1750 w         1.33              2150 vvs         1210 vvs         1.00           1580 s         1.37         2050 s         1515 m         1.35         1940 s         1440 s         1.35   </td>	TABLE IX           COMPARISONS OF INFRARED SPECTRA OF POLYBORA $B_4D_{16}$ $R$ $B_8H_{11}$ $B_6D_{11}$ $R$ $B_4H_8CO$ 1950 vs         1.33         2600 vs         1960 vs         1.33         2570 vs           1830 vs         1.37         2500 vs         1840 vs         1.36         2490 vs           1730 w         1.32         2330 w          2320 w              2150 vvs         1840 vs         1.36         2490 vs           1730 w         1.32         2330 w          2320 w          2320 w               2150 vvs         150 vvs           1580 s         1.37         2050 s         1515 m         1.35         1940 s                    1360 w         1.36                1360 w         1.35         1440 s,b         1090 s,b         1.32         1440 m           960 w         1.31	TABLE IX           COMPARISONS OF INFRARED SPECTRA OF POLYBORANE COMPOUND $B_4D_{16}$ R $B_6H_{11}$ $B_8D_{11}$ R $B_4H_6CO$ $B_4D_9CO$ 1950 vs         1.33         2600 vs         1960 vs         1.33         2570 vs         1940 vs           1830 vs         1.37         2500 vs         1840 vs         1.36         2490 vs         1830 vs           1730 w         1.32         2330 w          2320 w         1750 w              2150 vvs         2150 vvs         2150 vvs           1580 s         1.37         2050 s         1515 m         1.35         1940 s         1440 s                    1360 s         1.37         2050 s         1515 m         1.36         1810 w,sh                     1360 w         1.36                1360 w         1.35         1440 s,b         1090 s,b         1.	TABLE IX           COMPARISONS OF INFRARED SPECTRA OF POLYBORANE COMPOUNDS <sup>a,b</sup> $B_4D_{16}$ $R$ $B_8H_n$ $B_6D_n$ $R$ $B_8H_9CO$ $B_4D_9CO$ $R$ 1950 vs         1.33         2600 vs         1960 vs         1.33         2570 vs         1940 vs         1.32           1830 vs         1.37         2500 vs         1840 vs         1.36         2490 vs         1830 vs         1.36           1730 w         1.32         2330 w          2320 w         1750 w         1.33              2150 vvs         1210 vvs         1.00           1580 s         1.37         2050 s         1515 m         1.35         1940 s         1440 s         1.35

<sup>a</sup> Frequencies in cm. <sup>b</sup>v, very; s, strong; m, medium; w, weak; b, broad; sh, shoulder.

tions reported in the literature.<sup>25,26</sup> The frequency

quickly as possible, with the samples at temperatures just high enough for a stable spectrum. However,

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partial decomposition at higher temperatures did not appreciably change the spectra.

The n.m.r. record for  $B_4H_8CO$  (Fig. 3) consists of a low-field triplet, the slight asymmetry of which was repeatedly confirmed, and a high-field doublet (of area one-third as great) typically representing the apical B-H group found in most polyboranes. The pattern for  $B_4D_8CO$  is simpler because the spin coupling constant for B-D is only about 17% as large as for B-H, so that multiplets are not resolved. Thus the  $B_4H_8CO$ low-field triplet becomes for  $B_4D_8CO$  only a single peak centered at 2.3 $\delta$ , and the high-field doublet is similarly collapsed to a singlet.

The asymmetry of the  $B_4H_8CO$  low-field triplet must be due to some kind of composite character. Its components cannot be either a pair of doublets, two closely adjacent peaks forming the central peak, or a doublet nearly coincident with two peaks of a triplet, for in such cases the low-field pattern of  $B_4D_8CO$  would have been a doublet instead of a singlet. However the components could be either (1) a triplet and a doublet having nearly the same chemical shift, with the doublet so broadened as to show only as a distortion of the triplet, or (2) a triplet and a singlet having nearly the same chemical shift. Present evidence does not support any decision between these two cases.

If case 1 is correct, the most probable structure is one derived from that proposed for the  $B_4H_8$  intermediate in the decomposition of  $B_4H_{10}$ ,<sup>4</sup> namely



However, if case 2 is correct, we may consider a threeway tautomerism in which an apical B–H forms a bridge to either of two other boron atoms. This situation would be like the apical vs. bridging effect of the unique hydrogen atom in  $B_5H_{11}$ ,<sup>11</sup> which is shown for comparison with the three  $B_4H_8CO$  patterns in Fig. 4.

The three views of B<sub>5</sub>H<sub>11</sub> with apex-to-base bridge bonding are shown with dotted lines indicating prospective shifts of bonds, such as to form the suggested tautomers of B4H8CO with minimal reorganization. It can be argued that the easiest point of attack by CO upon B<sub>5</sub>H<sub>11</sub> would be at one of the upper-corner boron atoms, either of which must be especially electrondeficient because of participation in three three-center bonds. As BH<sub>3</sub> is removed (as shown by the dotted line slicing if off), the central (apical) boron atom would become a good place to attach the second CO. Such a mechanism would be consistent with the hydrolysis of  $B_5H_{11}$  to give mole-for-mole yields of  $B_4H_{10}^8$ ; in that case, however, a BH unit would be hydrolyzed off at the sensitive upper corner, leaving the B<sub>4</sub>H<sub>10</sub> pattern with scarcely the motion of even one atom.







Although it seems more difficult to find any way whereby  $B_{\delta}H_{11}$  could go so directly to the case 1 structure for  $B_4H_8CO$ , it still can be argued that the conversion from any of the tautomeric patterns to the case 1 structure is not very difficult, involving only minor redirection of B-H-B bridge vs. B-B and B-H terminal bonding. Hence no speculative discussion of possible easy mechanisms for the formation of  $B_4H_8CO$  from either  $B_{\delta}H_{11}$  or  $B_4H_{10}$  can aid in the selection of the most appropriate structural pattern.

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