

CONTRIBUTION FROM THE DEPARTMENT OF CHEMISTRY, UNIVERSITY OF SOUTHERN CALIFORNIA, LOS ANGELES, CALIFORNIA 90007

Phosphinopentaboranes<sup>1</sup>

BY ANTON B. BURG AND HERBERT HEINEN

Received December 8, 1967

Phosphine derivatives of the type  $R_2PCl$  react with  $LiB_5H_8$  in ether to form 2,3- $\mu$ -( $CH_3$ )<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> (mp 17°; bp estd 191°), 2,3- $\mu$ -CH<sub>3</sub>CF<sub>3</sub>PB<sub>5</sub>H<sub>8</sub>-A (mp 14°; 1 mm at 25°) which isomerizes irreversibly by an apparently first-order rate process to 2,3- $\mu$ -CH<sub>3</sub>CF<sub>3</sub>PB<sub>5</sub>H<sub>8</sub>-B (mp -28°; 9 mm at 25°; bp estd 137°), and 1-(CF<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> (mp -42°; bp estd 184°). The bridging *vs.* peak positions of phosphorus are proved by the B, F, and H nmr spectra and understood in terms of different phosphine-base strengths. The phosphorus nmr spectra show an interesting decet for 1-(CF<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> but no resolution for the B-P-B bridged compounds. Both infrared and nmr results indicate weaker-than-normal terminal B-H bonds (with more than normal B<sub>2p</sub> character) adjacent to bridging P atoms. The CH<sub>3</sub>CF<sub>3</sub>PB<sub>5</sub>H<sub>8</sub> isomers show no discernible B-F coupling, whereas 1-(CF<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> shows  $J_{B-F} = 6.0$  cps.

The recent discovery of the compound  $\mu$ -(CH<sub>3</sub>)<sub>3</sub>-SiB<sub>5</sub>H<sub>8</sub>, in which a three-center B-Si-B bond slowly changes to the more stable situation in 2-(CH<sub>3</sub>)<sub>3</sub>Si-B<sub>5</sub>H<sub>8</sub>,<sup>2</sup> suggested the similar synthesis of  $\mu$ -R<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> compounds whose properties might contribute to the development of chemical principles. Indeed, LiB<sub>5</sub>H<sub>8</sub><sup>3</sup> (in ether at -120 to -78°) reacted easily with (CH<sub>3</sub>)<sub>2</sub>-PCl to produce the fairly stable  $\mu$ -(CH<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub>, the B-P-B bridged structure of which was confirmed by a proton nmr spectrum showing different environments for the two methyl groups. The similar CH<sub>3</sub>CF<sub>3</sub>PCl reaction also led to B-P-B bridging, here proved by the existence of two isomers whose <sup>11</sup>B nmr spectra included a characteristic triplet quite similar to that found for  $\mu$ -(CH<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub>. These isomers were easily distinguishable by their different melting points, volatility, infrared spectra, and nmr chemical shifts. The initial product was almost entirely the higher melting isomer A, which converts completely to isomer B, about 9 times as volatile as A.

The reaction product of LiB<sub>5</sub>H<sub>8</sub> with (CF<sub>3</sub>)<sub>2</sub>PCl or (CF<sub>3</sub>)<sub>2</sub>PI was still different, consisting solely of 1-(CF<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub>, with no evidence found for any B-P-B bridged intermediate. It seems that phosphorus here cannot use its lone-pair electrons for base action toward boron, nor can it develop an electron-deficient bridge in the presence of the lone-pair electrons. This result is curious, for we might have expected the environment of phosphorus in a  $\mu$ -(CF<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> to be similar to that in the stable trimer [(CF<sub>3</sub>)<sub>2</sub>PBH<sub>2</sub>]<sub>3</sub>,<sup>4</sup> except for half as many B-H bonds and electron-deficient boron. The absence of  $\mu$ -(CF<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> would support the idea that the stability of [(CF<sub>3</sub>)<sub>2</sub>PBH<sub>2</sub>]<sub>3</sub> depends upon interaction between P and four adjacent B-H bonds.<sup>5</sup> However, this argument would lose

force if some special stabilization effect favors 1-(CF<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub>.

## Syntheses and Characterizations

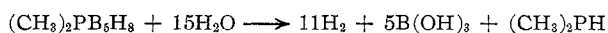
The synthesis of each R<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> compound began with the reaction of B<sub>5</sub>H<sub>9</sub> with a 0.1 M solution of LiC<sub>4</sub>H<sub>9</sub> in diethyl ether at -78°, in a vertical tube leading through a stopcock to the high-vacuum manifold. The resulting solution of LiB<sub>5</sub>H<sub>8</sub> (2-3 mmoles) was treated with an amount of the R<sub>2</sub>PX compound roughly equimolar to the B<sub>5</sub>H<sub>9</sub> (both slightly in excess over the original LiB<sub>5</sub>H<sub>8</sub>). As the mixture warmed slowly from -120 to -78° (or sometimes to -50°), the occurrence of the desired reaction was indicated by a milky precipitate which redissolved during further standing at a fixed temperature (*e.g.*, 2 hr at -50°). No H<sub>2</sub> was formed. The ether and excess reactants (B<sub>5</sub>H<sub>9</sub> and sometimes R<sub>2</sub>PCl) were distilled from the tube at -50 to -40° into the vacuum system; then the product R<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> could be delivered similarly as the tube warmed to 25°. Each product was purified by repeated high-vacuum fractional condensation through a series of low-temperature U traps.

**Dimethylphosphinopentaborane(9).**—The product (CH<sub>3</sub>)<sub>2</sub>-PB<sub>5</sub>H<sub>8</sub> was made from (CH<sub>3</sub>)<sub>2</sub>PCl,<sup>6</sup> in a yield representing 90% of the original LiC<sub>4</sub>H<sub>9</sub>. One fractional condensation at -15° was sufficient for its purification. The pure compound melted sharply at 16.8° and showed the normal set of vapor tensions given by Table I. Its vapor-phase decomposition became quite appreciable at 85°; or the liquid might suffer as much as 20% conversion to an amber-brown resin during 1 week at 25°. This resin, like others derived from B<sub>5</sub>H<sub>9</sub> with bases, showed a very complex <sup>11</sup>B nmr spectrum (Figure 1): boron in many environments.

TABLE I

| VOLATILITY OF (CH <sub>3</sub> ) <sub>2</sub> PB <sub>5</sub> H <sub>8</sub> |      |      |      |      |      |      |
|------------------------------------------------------------------------------|------|------|------|------|------|------|
| Log $P = 7.0157 + 1.75 \log T - 0.005T - (3010/T)$                           |      |      |      |      |      |      |
| $t_{760} = 191.4^\circ$ ; Trouton constant = 22.5 eu                         |      |      |      |      |      |      |
| Temp, °C                                                                     | 52.0 | 58.5 | 61.6 | 66.0 | 70.4 | 80.1 |
| $P_{\text{obsd}}$ , mm                                                       | 3.36 | 4.93 | 5.90 | 7.47 | 9.43 | 15.5 |
| $P_{\text{calcd}}$ , mm                                                      | 3.38 | 4.93 | 5.89 | 7.47 | 9.43 | 15.4 |

The vapor-phase molecular weight of (CH<sub>3</sub>)<sub>2</sub>PB<sub>5</sub>H<sub>8</sub> was determined in a 627-ml immiscible tensimeter. At 84° and 14.25 mm pressure, the result was 122.8; at 78° and 10.7 mm, 123.8; calcd, 123.1. For analysis, a 15.4-mg sample (0.125 mmole) was hydrolyzed in the presence of HCl, requiring 24 hr at 100° for completion. The yield of H<sub>2</sub> was 1.37 mmoles, or 99.6% of the value calculated from the equation



**Methyltrifluoromethylphosphinopentaboranes.**—The slightly volatile isomer CH<sub>3</sub>CF<sub>3</sub>PB<sub>5</sub>H<sub>8</sub>-A was the main product of the reac-

(6) A. B. Burg and P. J. Slota, Jr., *ibid.*, **80**, 1107 (1958).

(1) It is a pleasure to acknowledge the generous support of this research by the Office of Naval Research. Reproduction in whole or in part is permitted for any purpose of the United States Government. We are grateful also to our colleague Dr. K. L. Servis for making possible our use of the HA-100 and A-60 nmr instruments, to Dr. R. E. Williams of Space-General Corp. for the loan of a 32.1-Mc radiofrequency unit and probe, and to both for useful discussions.

(2) D. F. Gaines and T. V. Iorns, *J. Am. Chem. Soc.*, **89**, 4249 (1967).

(3) D. F. Gaines and T. V. Iorns, *ibid.*, **89**, 3375 (1967).

(4) A. B. Burg and G. Brendel, *ibid.*, **80**, 3198 (1958).

(5) A. P. Lane and A. B. Burg, *ibid.*, **89**, 1040 (1967).

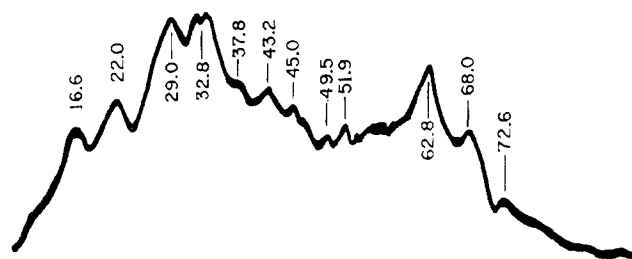


Figure 1.—Boron nmr spectrum of the nonvolatile decomposition product of  $(\text{CH}_3)_2\text{PB}_5\text{H}_8$ . The line width in each region measures the noise level. The chemical shifts are measured upfield from trimethyl borate.

tion of  $\text{LiB}_5\text{H}_8$  with  $\text{CH}_3\text{CF}_3\text{PCl}$ ;<sup>7</sup> the yield represented 91% of the  $\text{LiC}_4\text{H}_9$ . Also found was a 2.8% yield of the more volatile  $\text{CH}_3\text{CF}_3\text{PB}_5\text{H}_8\text{-B}$ , probably formed by conversion of isomer A during the processes of isolation and purification. Repeated runs confirmed that the synthesis is quite specific for isomer A. The conversion to B was complete in the liquid phase during 2 hr at 50° or during 48 hr in the vapor phase at 25° (1 mm pressure in a 1-l. bulb). The conversion could be observed by the change in the infrared spectrum in the gas phase at 35° or by the growth of the proton and  $^{19}\text{F}$  nmr peaks for isomer B during repeated scans of the "neat" liquid.

Isomer A melted sharply at 13.8° and showed 0.7 mm vapor tension at 20°. It was not feasible to determine the molecular weight in the vapor phase; and in ether solution at 0° the differential pressure method gave 270 at 0.6 *M* or 260 at 0.5 *M* (calcd, 177.1). Under similar conditions the B isomer gave the value 179.9. The mass spectra of both isomers (recorded by West Coast Technical Service, through the kind assistance of Dr. J. F. Ditter, of Space-General Corp.) showed a cutoff peak at 178, corresponding to  $\text{CH}_3\text{CF}_3\text{P}(^{11}\text{B})_5\text{H}_8$ . It was quite clear that isomer A was associated in solution for some structural reason not applicable to isomer B. The results could not be explained on the basis of dimer A in equilibrium with monomer B, for the conversion proved to be irreversible.

The  $\text{CH}_3\text{CF}_3\text{PB}_5\text{H}_8$  isomer B melted in the range  $-28.5$  to  $-28.3^\circ$ . This indication of purity was confirmed by the normal vapor tension data shown in Table II. The vapor-phase molecular weight results at 58–64° were 177.6 and 176.4 (calcd, 177.1).

TABLE II

VOLATILITY OF  $\text{CH}_3\text{CF}_3\text{PB}_5\text{H}_8\text{-B}$ 

$$\log P = 6.3335 + 1.75 \log T - 0.005T - (2450/T)$$

$$t_{760} = 136.9^\circ; \text{Trouton constant} = 21.45 \text{ eu}$$

| Temp, °C                | 24.1 | 30.0  | 40.3  | 45.4  | 49.7  | 53.7  |
|-------------------------|------|-------|-------|-------|-------|-------|
| $P_{\text{obsd}}$ , mm  | 8.55 | 11.96 | 20.82 | 26.89 | 33.23 | 40.12 |
| $P_{\text{calcd}}$ , mm | 8.57 | 11.99 | 20.79 | 26.93 | 33.22 | 40.14 |

For analysis, 11.0 mg (0.062 mmole) of  $\text{CH}_3\text{CF}_3\text{PB}_5\text{H}_8\text{-B}$ , hydrolyzed in 10 ml of water (sealed tube, 48 hr at 100°), gave 0.67 mmole of  $\text{H}_2$  (98% of the expected 11  $\text{H}_2$  per molecule). The yield of  $\text{CH}_3\text{CF}_3\text{PH}$  was 0.0625 mmole (100.8%); it was identified by its molecular weight (116 as calcd) and infrared spectrum.<sup>7</sup> The boric acid was titrated as 0.312 mmole (100.5% of calcd). Finally, a direct basic hydrolysis of the compound gave 0.97  $\text{HCF}_3$  per mole, after 48 hr at 100°. This full confirmation of the formula  $\text{CH}_3\text{CF}_3\text{PB}_5\text{H}_8$  for isomer B serves equally well for its precursor, isomer A, since the conversion was quantitative.

**Bis(trifluoromethyl)phosphinopentaborane(9).**—The compound  $(\text{CF}_3)_2\text{PB}_5\text{H}_8$  was made first from 2 mmoles of  $\text{LiB}_5\text{H}_8$  with 3.21 mmoles of  $(\text{CF}_3)_2\text{PCl}$  in 20 ml of diethyl ether at  $-78^\circ$  (15 hr). The ether, removed *in vacuo* at  $-50^\circ$ , was not accompanied by any of the expected excess of  $(\text{CF}_3)_2\text{PCl}$ , which could have been identified quite easily by its strong infrared spectrum. The slightly volatile product was distilled out *in vacuo*

during 5 hr at 24°, leaving a milky oil which apparently included the elements of the missing  $(\text{CF}_3)_2\text{PCl}$ . Complete removal of the ether from the product  $(\text{CF}_3)_2\text{PB}_5\text{H}_8$  required five-times repeated high-vacuum fractional condensation through a trap at  $-30^\circ$ . The yield of the pure product was 266 mg (1.15 mmoles; 57%).

The use of  $(\text{CF}_3)_2\text{PI}$  for this synthesis was essentially the same in all respects.

The sharp melting point of  $(\text{CF}_3)_2\text{PB}_5\text{H}_8$  ( $-42.0^\circ$ ; no range) and the normal character of its vapor tensions (Table III) left no doubt of its purity; no isomerism was observed.

The vapor-phase molecular weight determinations gave 236 and 231; calcd, 231.1. The hydrolytic analysis is described with millimole quantities, as follows.

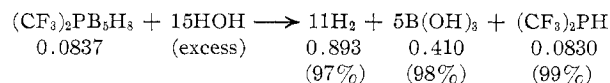


TABLE III

VOLATILITY OF  $(\text{CF}_3)_2\text{PB}_5\text{H}_8$ 

$$\log P = 6.6114 + 1.75 \log T - 0.005T - (2790/T)$$

$$t_{760} = 184.3^\circ; \text{Trouton constant} = 21.0 \text{ eu}$$

| Temp, °C                | 25.0 | 35.5 | 45.3 | 57.0 | 68.6 | 74.0 | 82.0 |
|-------------------------|------|------|------|------|------|------|------|
| $P_{\text{obsd}}$ , mm  | 1.20 | 2.41 | 4.39 | 8.29 | 14.9 | 19.3 | 27.7 |
| $P_{\text{calcd}}$ , mm | 1.24 | 2.43 | 4.35 | 8.28 | 14.9 | 19.3 | 27.7 |

**Nuclear Magnetic Resonance.**—The chemical shifts ( $\delta$ , ppm) and observable coupling constants ( $J$ , cps), determined by means of the Varian HA-100 instrument ( $^1\text{H}$  at 100 Mc,  $^{19}\text{F}$  at 94.1 Mc,  $^{31}\text{P}$  at 40.5 Mc, and  $^{11}\text{B}$  at 32.1 Mc) for the four phosphinopentaboranes, are shown in Table IV. Here the  $\delta$  values for protons relate to tetramethylsilane as a 5% internal standard, and were rechecked by the Varian A-60 instrument. For the other nuclei, the  $\delta$  values were measured by substitution of tubes containing neat-liquid samples of reference compounds, the side bands of which determined the scale in cps. For boron, trimethyl borate proved convenient for avoiding superposition and was more sharply measurable than the polyborane peaks; for phosphorus, 85%  $\text{H}_3\text{PO}_4$  was used; and for fluorine,  $\text{Cl}_3\text{CF}$  was used in one case, and the others were corrected from  $\text{C}_6\text{F}_6$ . Since the new compounds and the standards were in tubes of precisely similar bore and wall thickness, the systematic errors due to differences of diamagnetic susceptibility probably did not exceed 0.5 ppm. For consistency, all  $\delta$  values here are measured upfield from the standards; a minus sign means a downfield shift.

TABLE IV

## NMR SPECTRA OF PHOSPHINOPENTABORANES

|               | $\text{CH}_3\text{CF}_3\text{PB}_5\text{H}_8$ |     |                   |          |                 |      | $(\text{CF}_3)_2\text{PB}_5\text{H}_8$ |
|---------------|-----------------------------------------------|-----|-------------------|----------|-----------------|------|----------------------------------------|
|               | Isomer A                                      |     |                   | Isomer B |                 |      |                                        |
|               | $\delta$                                      | $J$ | $\delta$          | $J$      | $\delta$        | $J$  | $\delta$                               |
| 1-B           | 64.8                                          | 154 | 63.7              | 163      | 64.2            | 163  | 68.8                                   |
| 2,3-B         | 41.3                                          | 114 | 42.1              | 105      | 41.1            | 112  | ...                                    |
| 4,5-B         | 19.1                                          | 156 | 19.8              | 160      | 18.6            | 164  | 30.4                                   |
| $\text{CH}_3$ | -1.72                                         | 12  | -1.37             | 10.0     | -1.97           | 12.6 | ...                                    |
|               | -1.09                                         | 10  |                   |          |                 |      | ...                                    |
| P             | 85 <sup>a</sup>                               | ... | 33.5 <sup>a</sup> | ...      | 47 <sup>a</sup> | ...  | 25.1 <sup>b</sup> "73"                 |
| F             | ...                                           | ... | 59 <sup>c</sup>   | 68       | 71 <sup>c</sup> | 61   | 49.9 P 71.3<br>B 6.0                   |

<sup>a</sup> Unresolvable peaks, 750–800 cps wide at the base. <sup>b</sup> Decet; cf. Figure 5. <sup>c</sup> Corrected to  $\text{Cl}_3\text{CF}$  reference by adding 165 ppm to the  $\delta$  measured from  $\text{C}_6\text{F}_6$ , in accord with the measured difference between the two reference tubes actually used.

The  $^{11}\text{B}$  spectra of  $(\text{CH}_3)_2\text{PB}_5\text{H}_8$  and the  $\text{CH}_3\text{CF}_3\text{PB}_5\text{H}_8$  isomers all were so very similar that Figure 2 almost could represent any of them. The integration of each showed accurately a 2:2:1 intensity pattern, confirming the B–P–B bridge interpretation. The most interesting feature of this spectral pattern is the central triplet. It cannot be interpreted as a superposed pair of doublets, for a 12.8-Mc spectrum of  $\text{CH}_3\text{CF}_3\text{PB}_5\text{H}_8\text{-B}$  (kindly recorded for us by Dr. R. E. Williams) showed the same triplet.

(7) A. B. Burg, K. K. Joshi, and J. F. Nixon, *J. Am. Chem. Soc.*, **88**, 31 (1966).



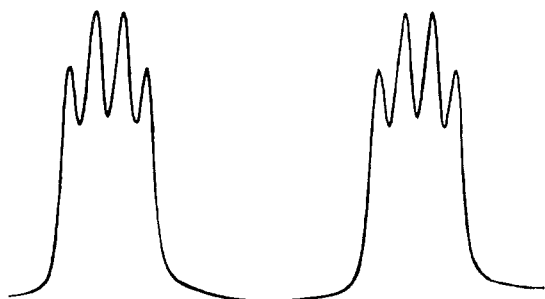


Figure 6.—Fluorine nmr spectrum of  $(\text{CF}_3)_2\text{PB}_3\text{H}_8$  at 94.1 Mc.

Where feasible, the relative intensities were determined as  $k = (100/PL) \log (I_0/I)$  for path length  $L$  and pressure  $P$  (reduced to  $25^\circ$ ), both in centimeters. These intensities are shown in parentheses after the corresponding frequencies ( $\text{cm}^{-1}$ ) in the following summary of results. The most detailed record was obtained for the relatively volatile isomer  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-B}$ , whereas the results for isomer A were limited by its low volatility and fairly rapid evolution of the vapor of isomer B from the liquid phase during infrared scans. Also the results for  $(\text{CH}_3)_2\text{PB}_3\text{H}_8$  were somewhat limited by the possibility of decomposition if one tried to develop a high vapor concentration.

Methyl-group stretching frequencies were recorded for  $(\text{CH}_3)_2\text{-PB}_3\text{H}_8$  at 2988 (0.8) and 2925 (0.8) and for  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-B}$  at 3017 (0.08) and 2943 (0.08). The higher frequencies and far lower intensities for the latter are in accord with much experience: C-H stretching loses intensity when C-F bonds are in the same molecule; and  $\text{CF}_3$  on P induces stronger bonding in adjacent groups. For  $(\text{CH}_3)_2\text{PB}_3\text{H}_8$ , the methyl-group deformations were normal at 1428 (0.9) and 1395 (0.6), but for  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-B}$  the same region was more confused by the skeletal modes included in the observed peaks 1427 (1.3), 1422 (1.1), 1403 (0.6), 1331 (0.7), 1308 (1.37), 1281 (1.43), and 1229 (0.93). For  $(\text{CF}_3)_2\text{-PB}_3\text{H}_8$ , there was continuous absorption in the same range (1540–1320), with peaks at 1475 (5) and 1443 (6). B-H-B bridging frequencies for  $(\text{CF}_3)_2\text{PB}_3\text{H}_8$  appeared at 1867 (5) and 1830 (2); and for the two kinds of B-H-B bridging in  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-B}$ , broad peaks were seen at 1975 (0.3), 1935 (0.3), 1875 (0.3), 1775 (0.2), and 1700 (0.1).

The B-H stretching frequencies correlate well with the structures of the various phosphinopentaboranes. For  $1\text{-(CF}_3)_2\text{PB}_3\text{H}_8$ , the four equal B-H bonds show one peak: 2630 (27). Contrastingly, the B-P-B bridged compounds would have 1-B, 2,3-B, and 4,5-B all different. Accordingly, for  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-A}$  we find peaks at 2610 (17), 2606 (17), and 2558 (17), all recorded before isomerization could occur; for  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-B}$ , 2615 (10), 1611 (12), 2609 (11), 2576 (8), and 2556 (13); and for  $(\text{CH}_3)_2\text{PB}_3\text{H}_8$ , 2607 (12), 2601 (16), 2596 (13), 2565 (7), and 2528 (16). The lowest frequencies here correlate with weaker-than-usual B-H bonding, suggested also by low B-H coupling constants, as will be discussed.

The C-F stretching region shows for  $(\text{CF}_3)_2\text{PB}_3\text{H}_8$  the peaks 1197 (45), 1155 (80), 1142 (78), and 1105 (48); for  $\text{CH}_3\text{CF}_3\text{-PB}_3\text{H}_8\text{-B}$ , 1187 (28) and 1142 (56); and for  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-A}$ , 1208 (18), 1190 (10), 1154 (25), and 1137 (30), some of which may be strongly affected by the presence of the more volatile isomer B.

Most of the remaining peaks are difficult to assign, for they are in regions appropriate for B-H bending,  $\text{CH}_3$  rocking and wagging,  $\text{CF}_3$  symmetric deformation, and possible skeletal modes. For  $(\text{CF}_3)_2\text{PB}_3\text{H}_8$  there is broad absorption from 930 to  $850\text{ cm}^{-1}$ , with a maximum at 885 (4), and peaks at 745 (0.9), 685 (2), 560 (1), and 460 (8). The others were not scanned below  $600\text{ cm}^{-1}$ . For  $(\text{CH}_3)_2\text{PB}_3\text{H}_8$ , peaks were found at 947 (8), 865 (1.3), 770 (3.6), and 742 (2); for  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-A}$ , at 976 (2?), 891 (36), and 879 sh (16); and for  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8\text{-B}$ , at 1110 (0.9), 1040 (0.9), 1014 (0.6), 971 (1.1), 937 (1.3), 916 R (9.3), 913 Q (10.7), 910 P (8.7), 860 (1.6), 776 (3), 772 (3), 758 (2), and 723 (3).

**Rate of Isomerization.**—The sharp and well-separated  $^{19}\text{F}$  nmr peaks for the  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8$  isomers were used to explore the rate of the  $\text{A} \rightarrow \text{B}$  conversion. A sample containing about 90% isomer A and 10% B was thermally equilibrated in the probe at  $35.0^\circ$  during 10 min. For each observation, both doublets were integrated at the same amplification level, usually within the same 40 sec. The mole fraction  $x$  of component A was calculated from the heights of both integrals.

In the range 70–23% A the rate was strictly first order:  $-\log x = 0.0240 + 0.002270t$  (with  $t$  meaning time in minutes and the constant 0.0240 describing the arbitrary starting point). The conformity to this equation, of points taken as regional averages of  $\log x$  and  $t$ , is shown in Table V.

TABLE V  
RATE DATA FOR  $\text{A} \rightarrow \text{B}$  ISOMERIZATION AT  $35.0^\circ$

| Time, min   | 65.51 | 88.32 | 126.05 | 142.92 | 189.90 | 246.25 |
|-------------|-------|-------|--------|--------|--------|--------|
| $x$ , obsd  | 0.669 | 0.595 | 0.492  | 0.448  | 0.351  | 0.259  |
| $x$ , calcd | 0.672 | 0.596 | 0.490  | 0.447  | 0.349  | 0.261  |

However, the rate constant for the first 0.5 hr was appreciably lower than for later stages. With every possible allowance for errors such as thermal lag and overestimation of the early small integrals for isomer B, the six points in the range 9–32 min determined the equation  $-\log x = 0.037 + 0.00220t$ . The difference probably is real but not easily explained.

A more extensive kinetic study of this isomerization, perhaps by vapor-phase infrared methods, might be informative.

### Structural Discussion

The bridging of two basal boron atoms (numbered 2 and 3) by phosphorus in  $(\text{CH}_3)_2\text{PB}_3\text{H}_8$  and  $\text{CH}_3\text{CF}_3\text{-PB}_3\text{H}_8$ —in contrast to the 1-B position of the  $(\text{CF}_3)_2\text{P}$  group in  $(\text{CF}_3)_2\text{PB}_3\text{H}_8$ —is verified beyond doubt by multiple aspects of the nmr spectra. Also, the close correspondence of the nmr spectra of the two  $\text{CH}_3\text{CF}_3\text{-PB}_3\text{H}_8$  species leaves no doubt of their isomeric relationship.

It is interesting that the three B-P-B bridged species all show 2,3-B-H coupling constants about one-third lower than is normal for pentaborane(9) derivatives (*ca.* 110 *vs.* normally 160 cps), suggesting that the B-H bonds adjacent to the B-P-B bridging have less  $\text{B}_{2s}$  and more  $\text{B}_{2p}$  character than usual. But hydrogen usually is more firmly bonded by more  $s$  character; hence one might expect these B-H bonds to be weaker than normal. In fact, all three of these species show some terminal B-H stretching frequencies decidedly lower than the usual  $2600\text{--}30\text{-cm}^{-1}$  range observed in most pentaborane(9) derivatives.

Another interesting item is the position—decidedly downfield relative to  $\text{B}_5\text{H}_9$ —of the 4,5-B nmr peaks, despite the expected electron-enriching effect of the electron-donor bridging phosphorus ligands: apparently the cross-ring boron atoms here go opposite to the situation for 4-B in  $2\text{-XB}_5\text{H}_8$  compounds, wherein the 4-B moves upfield in proportion to the availability of ligand  $\pi$  electrons.<sup>9</sup> In the present B-P-B bridged compounds, of course, there are no ligand  $\pi$  electrons.

However, a more important problem related to the nmr spectra is the structural decision concerning the  $\text{CH}_3\text{CF}_3\text{PB}_3\text{H}_8$  isomers. Assuming that the bridging

(9) A. B. Burg, *J. Am. Chem. Soc.*, **90**, 1407 (1968).

phosphorus atom is roughly tetrahedral and replaces a bridging proton insofar as direction from the B<sub>5</sub> skeleton is concerned, we would recognize that one R group on phosphorus has an "axial" position well below the B<sub>4</sub> plane and extending toward the 4,5-B atoms, while the other R group would be "equatorial," extending outward and almost coplanar with the basal boron atoms. Then space-induction effects might be used to decide which isomer has axial CF<sub>3</sub> and equatorial CH<sub>3</sub>, or *vice versa*.

By the simplest argument, a CF<sub>3</sub> group undergoing dipolar interaction with the especially electron-deficient 4,5-B atoms would have <sup>19</sup>F peaks more downfield and push the 4,5-B peaks upfield; and methyl groups far from the 4,5-B atoms would show relatively upfield proton peaks. All three expectations are met by the  $\delta$  values for isomer A. Accordingly, there is a temptation to conclude that isomer A has axial CF<sub>3</sub> with equatorial CH<sub>3</sub> and that isomer B has the opposite situation.

However, this conclusion would lead us to expect isomer A to be stabilized by internal dipolar interaction, whereas in fact it isomerizes quantitatively to isomer B. Also, we must not forget that isomer A is associated; and the only reasonable basis for the association would be an intermolecular dipolar interaction between CF<sub>3</sub> of one molecule and the 4,5-B atoms of another. Indeed, the association may amount to more than we have demonstrated, for the 50% too-high molecular weight was determined in

solution in ether, which could be expected to work against the postulated dipolar association. Attempts to determine the molecular weight in a more truly inert solvent, such as *n*-pentane, failed for lack of solubility.

Considering, then, that quite high actual molecular weights might prevail in the "neat" liquid isomer A, we may argue that this isomer could connect an equatorial CF<sub>3</sub> to the 4,5-B atoms of an adjacent molecule, with better space-inductive interaction (and more downfield chemical shift) than if an axial CF<sub>3</sub> were less effectively reaching toward 4,5-B in the same molecule. By this argument, isomer A would use equatorial CF<sub>3</sub> to maintain a weak dipolar polymer bonding, while axial CH<sub>3</sub> would be little affected by 4,5-B atoms which are receiving the charge effect of CF<sub>3</sub> from another molecule in a relatively effective manner. Then isomer B could be stabilized as a monomer by a dipolar action of axial CF<sub>3</sub> toward 4,5-B in the same molecule, while the equatorial methyl group would fail to develop effective polymer bonding to an adjacent molecule.

Both of the opposing ideas of the isomer structures have their merits, and we prefer not to choose between them at this time. Possibly the decision can be made later, on the basis of fuller information, including work on analogous compounds and more rigorously studied arguments.

Even less productive would be attempts to explain why isomer A is the exclusive product of the original synthesis.

CONTRIBUTION FROM THE AMES LABORATORY OF THE ATOMIC ENERGY COMMISSION  
AND THE DEPARTMENT OF CHEMISTRY, IOWA STATE UNIVERSITY, AMES, IOWA;  
MELLON INSTITUTE, PITTSBURGH, PENNSYLVANIA;  
THE DEPARTMENT OF CHEMISTRY, STATE UNIVERSITY OF NEW YORK, STONY BROOK, NEW YORK;  
AND THE DEPARTMENT OF CHEMISTRY, HARVARD UNIVERSITY, CAMBRIDGE, MASSACHUSETTS

## The Nuclear Magnetic Resonance Spectra of Aluminum Borohydride-Trimethylamine<sup>1a,b</sup>

By P. C. LAUTERBUR,<sup>1c</sup> R. C. HOPKINS, R. W. KING, O. V. ZIEBARTH, AND C. W. HEITSCH<sup>1d</sup>

Received October 9, 1967

The H<sup>1</sup>, B<sup>11</sup>, and Al<sup>27</sup> nmr spectra of aluminum borohydride and its trimethylamine adduct have been examined. On the basis of these spectra and cryoscopic data, the adduct is shown to be an undissociated monomer with magnetically equivalent borons and hydridic hydrogens. The adduct is unique in that it has a lower symmetry (C<sub>3</sub> or C<sub>3v</sub>) than the original aluminum borohydride (D<sub>3h</sub>) yet the electric field gradient at the Al<sup>27</sup> nucleus appears to be smaller in the adduct than in the free acceptor.

With their discovery of aluminum borohydride, Schlesinger, Sanderson, and Burg noted that it formed

1:1 adducts with trimethylamine, ammonia, and ether, all of which had some stability at room temperature.<sup>2</sup> Further addition of trimethylamine to its adduct resulted in the cleavage of the aluminum-boron framework to produce (CH<sub>3</sub>)<sub>3</sub>NBH<sub>3</sub>. More recently, it has been reported that a total of 4 equiv of amine will react with aluminum borohydride to degrade it to adducts of

(1) (a) Presented, in part, to the 74th Meeting of the Iowa Academy of Sciences, April 13, 1962, and, in part, at the 4th Omnibus Conference on Electronic and Nuclear Spectra, March 2, 1963, Pittsburgh, Pa. (b) Abstracted in part from a thesis submitted by O. V. Ziebarth in partial fulfillment of the requirements for the degree of Master of Science, Iowa State University, 1962, and in part from a thesis submitted by R. C. Hopkins in partial fulfillment of the requirements for the degree of Doctor of Philosophy, Harvard University, 1964. (c) Alfred P. Sloan Fellow. (d) Author to whom correspondence should be addressed: Monsanto Co., Inorganic Chemicals Division, St. Louis, Mo.

(2) H. I. Schlesinger, R. T. Sanderson, and A. B. Burg, *J. Am. Chem. Soc.*, **62**, 3421 (1940).