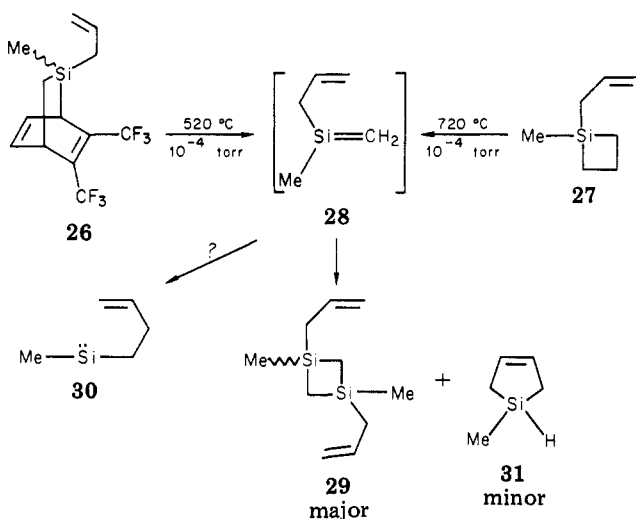


Scheme II



butadiene. After the hexafluoroxyene (74%), the major volatile product is silacyclohexene 24 (23%), the expected product of cycloaddition of 15 and the 1,3-diene. This result stands in stark contrast with the reported⁶ copyrolysis of silacyclobutane 14 and 1,3-butadiene (vide supra). The flow pyrolysis of 22 also affords a small amount (5%) of 2-methyl-3-(dimethylsilyl)methyl-1,3-butadiene (25). Although we have previously observed this type of linear adduct from the reactions of silenes and butadienes,¹¹ it was conceivable that 26 arose from a stepwise radical addition of dimethylsilylene to dimethylbutadiene followed by hydrogen migration from carbon to silicon. This interpretation is rendered highly unlikely since pyrolysis of 22 in an excess of triethylsilane produced no 2,2-dimethyl-1,1,1-triethylsilane, the normal product of silylene trapping by the Si-H bond.

Thus, as the evidence seems strong that pyrolysis of 22 does produce silene 15, the case for generation and rearrangement of 15 from silacyclobutane 14 is at best weakened. However, the picture is confused by the recent report of isomerization of 15 to Me_2Si : at 100 K.⁷ This amazingly facile hydrogen migration is consistent neither with our results for 22 nor with the recent calculations of Schaefer¹² which predict that the activation energy for the essentially thermoneutral rearrangement $\text{H}_2\text{Si}=\text{CH}_2 \rightarrow \text{H}-\text{Si}-\text{CH}_3$ is about 40 kcal/mol.

In the hopes of finding a silene particularly predisposed to rearrange to a silylene, we have generated allylmethylsilene 28. This was accomplished by the flash vacuum pyrolyses (10^{-4} torr) of both silabicyclo[2.2.2]octadiene 26 and 1-allyl-1-methyl-1-silacyclobutane (27) (Scheme II). Pyrolysis of 26 (520 °C) was extremely clean, affording only hexafluoroxyene (83%) and 1,3-diallyl-1,3-dimethyl-1,3-disilacyclobutane (28, 85%). Pyrolysis of 27 required ~ 200 °C more than for 26, but still produced 29 in 27% yield. Thus, although silene 28 seemed an excellent candidate for rearrangement to silylene 30 through a cyclic six-electron process, it appears that this does not happen, as 28 simply dimerizes in the normal head-to-tail fashion. However, in the pyrolysis of 27 there is a small amount (2.3%) of 4-methyl-4-silacyclopentene (31) formed which may well arise from the cyclization of silylene 30 followed by transannular hydrogen migration. This interpretation is not demanded as it is also possible

to imagine 31 arising from 28 through a di- π -methane sequence.

The many syntheses and structural proofs involved in this report will be incorporated with current work and complete details from ref 1 in a full paper.

Acknowledgment is made to the National Science Foundation and to Dow Corning Corporation for support of this work. We thank Procter and Gamble Co. for a fellowship to G.T.B. during this period. We are grateful to Professor Rob Conlin for sending us his report on the pyrolysis of 14 prior to publication.

Registry No. 2, 23290-58-6; 3, 76410-87-2; 5, 25261-26-1; 6, 865-76-9; 7, 3704-44-7; 8, 78697-99-1; 11 isomer 1, 78698-00-7; 11 isomer 2, 78698-01-8; 12, 78698-02-9; 13, 78698-03-0; 15, 38063-40-0; 21, 78698-04-1; 22, 78715-28-3; 23, 1628-01-9; 24, 78698-05-2; 26, 78698-06-3; 27, 3944-06-7; 28, 78698-07-4; 29, 78698-08-5; 2,3-dimethyl-1,3-butadiene, 513-81-5; $\text{F}_3\text{CC}\equiv\text{CCF}_3$, 692-50-2.

Rational Syntheses of Mixed Dialkylhaloboranes ($\text{R}^A\text{R}^B\text{BX}$) and Mixed Trialkylboranes ($\text{R}^A\text{R}^B\text{R}^C\text{B}$) via Stepwise Hydridation-Hydroboration of Alkyldihaloboranes

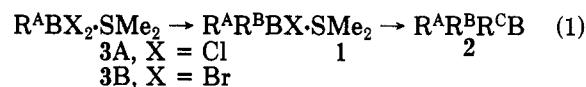
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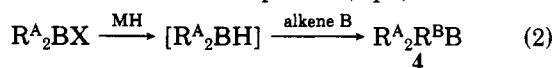
Received August 12, 1981

Summary: The stepwise hydridation of $\text{R}^A\text{BR}_2\cdot\text{SMe}_2$ with LiAlH_4 , followed by stepwise hydroboration of two different alkenes, provides a convenient route to the mixed organoboranes, $\text{R}^A\text{R}^B\text{R}^C\text{B}$, and to the products into which such mixed organoboranes can be transformed.

A rational synthesis of mixed dialkylhaloboranes ($\text{R}^A\text{R}^B\text{BX}$, 1) and trialkylboranes ($\text{R}^A\text{R}^B\text{R}^C\text{B}$, 2) has long eluded chemists. A convenient synthetic approach to such derivatives would greatly expand applications for the versatile organoboranes. We now wish to report the preparation of such mixed organoborane derivatives (1, 2) via the controlled hydridation of alkyldihaloboranes ($\text{R}^A\text{BX}_2\cdot\text{SMe}_2$, 3), followed by sequential hydroboration (eq 1).



The increasing importance of organoboranes¹ as synthetic intermediates reveals a need for mixed dialkylhaloboranes and mixed trialkylboranes. Partially mixed trialkylboranes, $\text{R}_2\text{R}^B\text{B}$ (4), have been prepared by various methods.²⁻⁶ Most of the earlier approaches involved use of the stable R_2BH or R^BBH_2 reagents.^{1,5} Recently, more general methods have been reported (eq 2).^{4,6}



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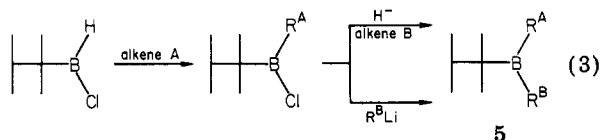
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Table I. Preparation of Mixed Organoboranes and Their Carbon Analogs

R ^A R ^B R ^C B substituents			yield, ^a %	
R ^A	R ^B	R ^C	ketone R ^A COR ^B	carbinol R ^A R ^B R ^C COH
<i>n</i> -C ₈ H ₁₁	<i>n</i> -C ₆ H ₁₃	Br	84 (70)	
<i>n</i> -C ₆ H ₁₃	<i>n</i> -C ₁₂ H ₂₅	Br	87 (72) ^b	
<i>n</i> -C ₆ H ₁₃	<i>n</i> -C ₈ H ₁₇	<i>n</i> -C ₈ H ₁₇		69 ^c

^a GC yields, determined by using a suitable internal standard. Isolated yields of >98% pure products are given in the parentheses. ^b Yield after double crystallization; >99% pure material, mp 47–48 °C. ^c Chemical purity 95%.

It is apparent that a general route to the totally mixed organoboranes, **2**, would require a practical synthesis of mixed dialkylhaloborane, **1**, or mixed dialkylboranes, R^AR^BBH. A general synthesis of totally mixed trialkylboranes, such as **5**, where R^C is the highly hindered 2,3-dimethyl-2-butyl (thexyl, Thx) group, has been reported recently (eq 3).^{7,8} However, our attempts to obtain

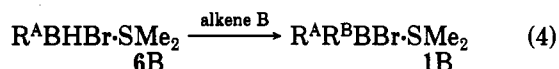


R^AR^BBH by the controlled elimination of 2,3-dimethyl-2-butene have thus far not been successful.^{9,10} Consequently, we undertook to search for a simple procedure for preparing the hitherto inaccessible mixed dialkylhaloboranes (**1**) and mixed trialkylboranes (**2**).

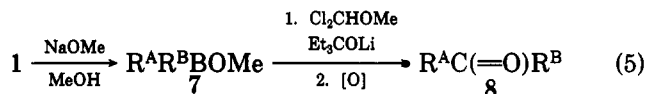
Alkyldihaloboranes (**3**) are readily prepared by hydroboration of alkenes with dihaloborane-methyl sulfide (HBX₂·SMe₂).^{11,12} The addition of LiAlH₄ to **3** in the presence of alkene B provides R^AR^BB (50%) and unreacted **3** (50%). Fortunately, the reaction of **3B** with the stoichiometric quantity of LiAlH₄ (borane:hydride ratio 1:1) in Et₂O affords the corresponding alkylboroborane, R^ABHBr·SMe₂ (**6B**).¹³ The ¹¹B NMR spectrum of **6B** exhibits a doublet around δ 0.7 (J_{BH} 136 Hz); heteronuclear proton decoupling shows a single resonance. The IR spectrum exhibits an absorption at 2450 cm⁻¹, but none in the region of 1550 cm⁻¹, indicating the presence of a nonbridged boron hydride species (this implies both the existence of **6B** as a SMe₂ complex and the absence of R^ABH₂ dimer). The hydridation is relatively slow, requiring 3 h at 0 °C, followed by 1 h at room temperature.

Terminal alkenes are readily hydroborated by **6B** at 0 °C to provide the corresponding mixed dialkylboroboranes (eq 4).¹⁴ This procedure provides a simple syn-

thetic route to the mixed dialkylboroboranes, R^AR^BBBr.

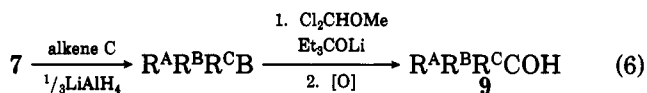


The absence of any significant disproportionation was established by converting the product, **1B**, into the borinate ester (R^AR^BBOMe, **7**) by treatment with methanol-sodium methoxide and converting the product into ketone, R^AR^BCO (**8**), via the DCME reaction (eq 5).¹⁵ Examination of the product by GC analysis revealed the absence of R^A₂CO and R^B₂CO impurities (Table I).



The reaction of **1** with 1 equiv of hydride (1/4 LiAlH₄) in the presence of alkene C failed to provide the desired mixed organoborane **2**. Instead, the DCME reaction¹⁶ revealed the presence of largely disproportionated boranes.¹⁷ Possibly the presence of the byproduct, LiAlBr₄, facilitates redistribution.

This difficulty was overcome by treating the reaction mixture containing **1** with methanol-sodium methoxide, when **1** was converted into **7** and the salts were precipitated as sodium bromide and aluminum methoxide. Finally, **7** was converted into the mixed borane by treatment with alkene C and LiAlH₄ (eq 6).² GC analysis of the product revealed the presence of a single component (**9**, 69% yield based on **3**) in 95% purity.



Consequently, the present procedures make available both mixed R^AR^BBX and mixed R^AR^BR^CB for the first time. The following procedures are representative.

To 3.18 g (10 mmol) of *n*-C₆H₁₃BBR₂·SMe₂¹¹ was added 2.2 mL (30 mmol) of SMe₂ and 15 mL of Et₂O at 0 °C, followed by a slow addition of LiAlH₄ in Et₂O (2.5 mmol) with stirring. The reaction was allowed to proceed for 3 h at 0 °C, followed by 1 h at room temperature, and 1.2 mL (11 mmol) of 1-pentene was added at 0 °C. After 3 h, 4.48-mL solution of NaOMe in MeOH (17.5 mmol, 3.9 M) was added slowly. The mixture was stirred for 0.5 h at room temperature; solvents were removed under the aspirator vacuum (15 mmHg) and extracted with 2 × 15 mL of a 1:1 mixture of pentane and Et₂O. The solvents were removed, 15 mL of THF and 1.3 mL (5 mmol) of *n*-tetradecane (internal standard for GC analysis) were added, and the DCME reaction was carried out as described elsewhere.¹⁵ The GC analysis on a 6 ft × 1/4 in. column packed with 10% SE-30 on Chromosorb W indicated 84% yield of 6-dodecanone. From a 20-mmol scale reaction, 97% pure ketone was isolated in 70% yield: bp 85–87 °C (0.9 mm); *n*_D²⁰ 1.4315 [lit.¹⁸ bp 125 °C (12 mm); *n*_D²⁰ 1.4339].

(14) A small amount of ether-cleaved product, R^AR^BBOEt (5–10%), was formed in this process. The ¹¹B NMR examination of the methanolysis product indicated the presence of >90% pure borinate ester **7** (δ 55), along with some boronate (δ 30, <5%) and borate (δ 18, <5%) esters as impurities.

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(17) The GC analysis revealed the formation of 5–10% R^AR^BCO (from R^AR^BBOEt) and several carbinols (from the corresponding redistributed boranes).

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(10) Monoalkylborane-amine complexes have been prepared by the displacement of 2,3-dimethyl-2-butene, with amines, from thexylalkylboranes (ThxBRH): Brown, H. C.; Negishi, E.; Katz, J.-J. *J. Am. Chem. Soc.* **1972**, *94*, 5893.

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(12) The hydridation-hydroboration reaction works with alkyldichloroboranes as well. But the yields obtained from bromoboranes are better than those from chloroboranes. Besides, the alkyldibromoborane-methyl sulfide complexes can be prepared and handled more conveniently. Therefore, we employed mainly the alkyldibromoboranes during the present study.

(13) Attempted hydridation of **3B** with potassium triisopropoxyborohydride in THF formed the boronate esters, R^AB(OR)₂.

For the preparation of **9**, the borinate ester **7** (10 mmol) was dissolved in 15 mL of THF and treated with 3.33 mmol of LiAlH_4 in the presence of 1.56 mL (10 mmol) of 1-octene at 0 °C. The reaction mixture was stirred for 3 h at room temperature, and the aluminum salts were removed by a slow addition of 0.65 g of H_2SO_4 (6.66 mmol) and 0.5 mL of water at 0 °C. The reaction mixture was extracted with pentane (2×20 mL), the combined organic extracts were dried over anhydrous K_2CO_3 , solvents were removed under vacuum, and the resulting **2** was subjected to the DCME reaction.¹⁶ GC analysis using *n*-dodecane as an internal standard showed a 69% yield of 7-pentylpentadecan-7-ol. Approximately 5% of impurities could be detected.

In summary, the bromine atoms of alkylidibromoboranes serve as effective and easily replaceable blocking groups for stepwise hydroboration. They can be sequentially replaced via the hydridation reaction.⁶ Subsequent stepwise hydroboration provides the mixed dialkylboroboranes and trialkylboranes. This process represents the first general synthesis for such hitherto inaccessible derivatives via the stepwise hydridation-hydroboration processes. Both the mixed organoboranes as well as the sequential hydridation-hydroboration process possess great synthetic potential, some of which we are presently exploring.

Acknowledgment. We thank the National Institutes of Health for the financial support through Grant GM 10937-19.

Registry No. **1B** ($R^A = n\text{-C}_5\text{H}_{11}$, $R^B = n\text{-C}_6\text{H}_{13}$), 79329-70-7; **1B** ($R^A = n\text{-C}_6\text{H}_{13}$, $R^B = n\text{-C}_{12}\text{H}_{25}$), 79357-03-2; **2** ($R^A = n\text{-C}_6\text{H}_{13}$, $R^B = n\text{-C}_6\text{H}_{11}$, $R^C = n\text{-C}_8\text{H}_{17}$), 79329-09-2; **3B** ($R^A = n\text{-C}_6\text{H}_{13}$), 79329-71-8; **6B** ($R^A = n\text{-C}_6\text{H}_{11}$), 79357-04-3; **6B** ($R^A = n\text{-C}_6\text{H}_{13}$), 79357-05-4; **7** ($R^A = n\text{-C}_5\text{H}_{11}$, $R^B = n\text{-C}_6\text{H}_{13}$), 79329-10-5; **7** ($R^A = n\text{-C}_6\text{H}_{13}$, $R^B = n\text{-C}_{12}\text{H}_{25}$), 79329-11-6; **8** ($R^A = n\text{-C}_6\text{H}_{11}$, $R^B = n\text{-C}_6\text{H}_{13}$), 6064-27-3; **8** ($R^A = n\text{-C}_6\text{H}_{13}$, $R^B = n\text{-C}_{12}\text{H}_{25}$), 79329-12-7; **9** ($R^A = n\text{-C}_6\text{H}_{13}$, $R^B = n\text{-C}_6\text{H}_{11}$, $R^C = n\text{-C}_8\text{H}_{17}$), 79329-13-8; 1-pentene, 109-67-1; 1-hexene, 592-41-6; 1-dodecene, 112-41-4; 1-octene, 111-66-0.

Conversion of a Bridging Methylene Ligand to a Ketenylidene Moiety. Synthesis and Reactivity of $\text{H}_2\text{Os}_3(\text{CO})_9(\text{CCO})$

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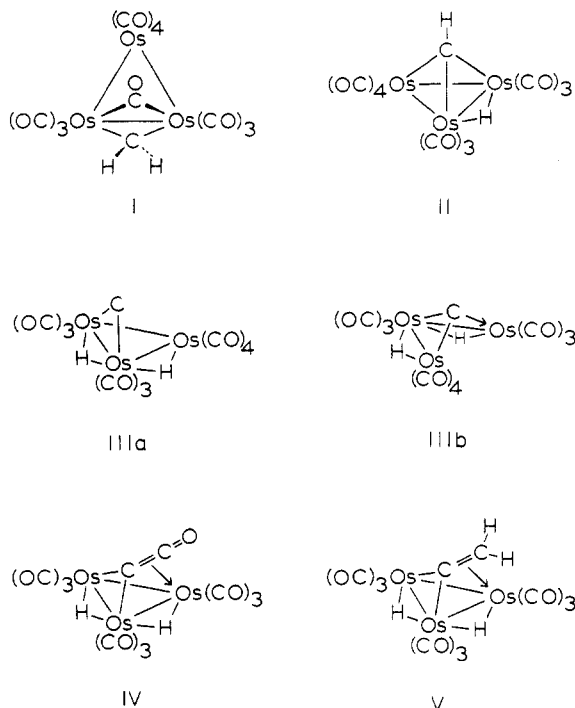
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Received June 3, 1981

Summary: The product formed by thermally induced loss of carbon monoxide from $\text{Os}_3(\text{CO})_{10}(\mu\text{-CO})(\mu\text{-CH}_2)$ is formulated as $\text{H}_2\text{Os}_3(\text{CO})_9(\mu_3\text{-}1,2\text{-}\eta^2\text{-CCO})$. The latter compound readily reacts with methanol to form the carbomethoxymethylidyne cluster, $\text{H}_3\text{Os}_3(\text{CO})_9(\mu_3\text{-CCO}_2\text{Me})$.

Recently a new triosmium methylene cluster, $\text{Os}_3(\text{CO})_{10}(\mu\text{-CO})(\mu\text{-CH}_2)$, was prepared by protonation of $[\text{Os}_3(\text{CO})_{11}\text{CHO}]^-$ and was proposed to have structure I based on infrared and ^{13}C and ^1H NMR spectroscopy.¹



The methylene cluster may also be obtained by treating $\text{Os}_3(\text{CO})_{11}\text{NCMe}$ with CH_2N_2 ;² an X-ray crystallographic study of the trimethylsilyl derivative $\text{Os}_3(\text{CO})_{10}(\mu\text{-CO})(\mu\text{-CHSiMe}_3)$ is consistent with structure I.² From hydrogeneration of $\text{Os}_3(\text{CO})_{10}(\mu\text{-CO})(\mu\text{-CH}_2)$ in refluxing benzene, a yellow byproduct having the stoichiometry " $\text{Os}_3(\text{CO})_{10}(\text{CH}_2)$ " was observed.¹ The same compound (mass spectrum, m/z 870 M^+ (^{192}Os); IR ($\nu(\text{CO})$, C_6H_{12}) 2121 cm^{-1} , 2086 cm^{-1} , 2064 cm^{-1} , 2055 cm^{-1} , 2034 cm^{-1} , 2006 cm^{-1} , 1994 cm^{-1} , 1984 cm^{-1}) is prepared in 84% yield by refluxing a nitrogen-purged toluene solution of $\text{Os}_3(\text{CO})_{10}(\mu\text{-CO})(\mu\text{-CH}_2)$.

Three formulations have been considered for " $\text{Os}_3(\text{CO})_{10}(\text{CH}_2)$ ". The prospect of $\text{HOs}_3(\text{CO})_{10}(\mu_3\text{-CH})$ (II), analogous to the conversion of $\text{H}_2\text{Os}_3(\text{CO})_{10}(\mu\text{-CH}_2)$ into $\text{H}_3\text{Os}_3(\text{CO})_9(\mu_3\text{-CH})$,³ is eliminated by the absence of a methyldyne ^1H NMR signal. A singlet hydride peak is seen at 25 °C (τ 29.73, CH_2Cl_2) and down to -80 °C. The $^{13}\text{C}\{^1\text{H}\}$ NMR spectrum of " $\text{Os}_3(\text{CO})_{10}(\text{CH}_2)$ " (90% ^{13}C enriched, from $\text{Os}_3(\text{CO})_{10}(\mu\text{-CO})(\mu\text{-}^{13}\text{CH}_2)$)² shows a singlet at δ 8.6 (CDCl_3 , 25 °C); off-resonance decoupling establishes that the labeled carbon is not hydrogen substituted (the τ 29.73 signal is now a doublet, $^2J(\text{C-H}) = 2.9$ Hz). A formula such as $\text{H}_2\text{Os}_3(\text{CO})_{10}(\text{C})$ seems unlikely, since compounds with either a divalent (IIIa) or a trivalent (IIIb) carbon atom are expected to be quite reactive and hence unstable. However, formulation of the yellow compound as a ketenylidene complex, $\text{H}_2\text{Os}_3(\text{CO})_9(\text{CCO})$ (IV), is consistent with all the available data. Thus, the IR spectrum (Nujol mull) shows a weak band at 1644 cm^{-1} which is assigned to $\nu(\text{CCO})$ of the ketenylidene moiety. Furthermore, the ^{13}C NMR spectrum of a sample prepared from $\text{Os}_3(\text{CO})_{10}(\mu\text{-CO})(\mu\text{-CH}_2)$ that was ca. 40% ^{13}C enriched at the carbonyls and 90% ^{13}C enriched at the methylene shows both the δ 8.6 singlet and a superimposed doublet ($^1J(\text{C-C}) = 86$ Hz) of ca. 40% total intensity. Simultaneously, a doublet with the same coupling constant, due to the ketenylidene carbonyl, appears at δ 160.3.

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