Chemistry and the Carbon Arc

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The composition and physical properties of carbon vapor have been widely investigated, but until the last decade its chemical properties had been largely neglected. This is true not only for carbon vapor but also for most other high-boiling elements (and compounds). Only recently have the techniques developed for studying the chemistry of carbon vapor been extended to other elements of the periodic table. This unexplored territory of 10 years ago has now been demonstrated to be a fruitful area of investigation.

High boiling point implies strong bonding between atoms. These strong bonds must be overcome for reactions to occur in which an atom is detached from its condensed-phase form. Only reagents which can engage the atom in a reaction sufficiently exothermic to overcome these forces can react. When the substance is vaporized these forces are overcome, and an atom unreactive in condensed phase becomes highly reactive. The higher the boiling point, the less will the chemical properties of the bulk material serve as a useful guide for the behavior of the free atoms.

Apparatus

The carbon arc is used as a source of carbon vapor which is produced as a result of the high temperature attained by the arc. Using the apparatus shown in Figure 1, carbon vapor is produced at low pressure (ca. 5 \times 10 ⁻⁵ Torr) by intermittent arcing (16 V ac) between two graphite electrodes and cocondensed on the liquid nitrogen cooled walls with a large excess of substrate.¹ The high vacuums employed insulate the high-temperature zone from the low-temperature zone, and the carbon vapor travels one mean free path or less to the reactor walls, thus eliminating gas-phase reactions. The high dilution of carbon species in the condensed phase is such that the thermodynamically favored association of carbon atoms to elemental carbon is minimized. Although it is difficult to know at which temperature the reactions occur, it is noteworthy that deoxygenation of ketones

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produces CO during the codeposition at -196° rather than upon warm-up.

Composition of Carbon Vapor

The arc produces a multitude of species, and the chemistry of C1, C2, C3, and C4 has been investigated. The relative abundances of the carbon species have been determined mass spectrometrically, but it was desirable for us to have a chemical determination also. The relative concentrations are obtained by analysis of the products from the reaction with chlorine. Thus, atomic carbon makes CCl₄, diatomic carbon yields C₂Cl₄ and C₂Cl₆, and C₃ produces perchloropropene and perchloropropane. The ratios change noticeably when the vapor is produced by Langmuir type of vaporization (resistive heating of graphite) as shown in Table I.^{2,3} Because of the geometry of our system, only 82% of the carbon emitted from the arc actually reaches the reaction zone. The remainder is deposited on mechanical obstructions in the reactor system. Thus yields are based on 82% of the weight lost from the carbon electrodes.

The carbon species produced in an arc are bombarded by free electrons such that there is a greater preponderance of excited species than would be expected for a Boltzmann distribution of species at arc temperatures ($\sim 2500^{\circ}$). The ¹S and ¹D excited states of atomic carbon are sufficiently long-lived to undergo collisionless flight to the walls (in *ca.* 10^{-5} sec) so that their chemistry, together with that of the ³P ground-state species, can be investigated.⁴⁻⁶ Higher excited states are lost during the flight; allowed optical transitions to one of the three states

Table I

	Relative molar concentrations ^a	
- Species	Arced carbon ²	Thermally vaporized carbon ³
C1	100	100
C_2	35 - 48	20
C ₃	7-10	61
C4	0.6-1.0	

 a These concentration ranges are determined from experiments in which at least 75% of the vaporized carbon is accounted for as products.

(1) P. S. Skell, L. D. Wescott, Jr., J. P. Goldstein, and R. R. Engel, J. Amer. Chem. Soc., 87, 2829 (1965).

(2) R. F. Harris, Ph.D. Thesis, The Pennsylvania State University, University Park, Pa., 1968.

(3) J. J. Havel, Ph.D. Thesis, The Pennsylvania State University, University Park, Pa., 1972.

(4) H. Yilmaz, Phys. Rev., 100, 1148 (1955).

(5) L. Ornstein, H. Brinkman, and A. Buenes, Z. Phys., 77, 72 (1932).

(6) R. C. Mason, Physica, 5, 777 (1938).

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Figure 1. Reaction chamber (immersed in liquid nitrogen).

are available. It is interesting to note that no current theory makes a satisfactory prediction of the proportions of ground state and metastable states which result from allowed decays from upper states.

Reactions of Atomic Carbon

Saturated hydrocarbons react with atomic carbon in low yields, giving a variety of products containing an extra carbon atom. Both triplet (ground state) and singlet (excited state) atoms insert into carbonhydrogen bonds to make respectively triplet and singlet carbenes. Triplet carbenes abstract two hydrogens to form a methyl group, whereas singlet carbenes react by migration of hydrogen atoms or alkyl groups to give olefins, and also by intramolecular insertions to produce cyclopropanes.⁷ Similar behavior has been noted for alkyl carbenes generated in other ways such as diazoalkane thermolyses.⁸ Scheme I il-



(7) P. S. Skell and R. R. Engel, J. Amer. Chem. Soc., 88, 4883 (1966).

lustrates the mode of product formation in the reaction of carbon atoms with *n*-butane.⁹ (Yields are based on C_1 produced by the arc, about 40% by weight of the carbon vaporized.)

Alcohols react with atomic carbon by insertion into C-H and O-H bonds and by abstraction of oxygen to make CO.¹⁰ All three possibilities occur when EtOH is the substrate; Scheme II outlines the proba-



ble pathways. (Yields are based on C_1 produced.) Further evidence for the proposed mechanisms is provided by the use of CD_3OH which yields products labeled as predicted for these pathways

$$CD_3OCH_2OCD_3 \xleftarrow{O-H \text{ insertion}} CD_3OH \xrightarrow{C-H \text{ insertion}} D \text{ migration}$$

 $CD_3HCDO \xleftarrow{CD_2=CDOH}$

CO (36%) and olefins are also observed, but the mechanism for their origin is still obscure.

With all alcohols studied, O-H insertion occurred five to eight times faster than C-H insertion per bond. The order of decreasing reactivity towards C-H insertion of C₁ is $CH_3-O > -CH_2-O > -CH_2 \sim -CH_3$.

Formation of C-CH₃ groups from C-H groups is attributable solely to insertion by ground-state (³P) carbon atoms followed by two hydrogen abstractions. Excited-state singlet carbon atoms insert to form singlet carbenes which can undergo rearrangements. Thus acetone yields methyl vinyl ketone (2.5%) and 2-butanone (3%), but when a radical trap such as butadiene is present the methyl vinyl ketone yield is unaltered but the 2-butanone decreases markedly. In contrast, when methanol, a singlet carbene trap, is present the CH₃-producing reaction to make 2-butanone is unaltered.¹¹

(8) W. Kirmse, "Carbene Chemistry," Academic Press, New York, N. Y., 1964.

(9) J. H. Plonka, Ph.D. Thesis, The Pennsylvania State University, University Park, Pa., 1970.

(10) P. S. Skell and R. F. Harris, J. Amer. Chem. Soc., 91, 4440 (1969).
(11) P. S. Skell, J. H. Plonka, and L. S. Wood, Chem. Commun, 710 (1970).



Alkyl halides react with atomic carbon to make olefins, cyclopropanes, and fragmentation products. CH_2Cl_2 and atomic carbon yields 1,1-dichloroethylene. This reaction could occur either via C-Cl insertion followed by Cl migration or via C-H insertion and then H migration.¹² This dichotomy is unresolvable except where each mechanism yields a unique product such as with 1,1,1-trichloroethane

$$CH_{3}CCl_{3} + C_{1} \not\subset CH_{3}CCl_{2}\dot{C}Cl \rightarrow CH_{3}CCl=CCl_{2}$$
$$CCl_{3}CH_{2}\dot{C}H \rightarrow CCl_{3}CH=CH_{2}$$

Experimentally, 1,1,2-trichloropropene is produced while 3,3,3-trichloropropene is not, indicating that C-Cl insertion is the predominant process. The less favored C-H insertion products are isolable from the 2-chloropropane reaction whereby not only the expected 1-chloro-2-methylpropene but also 3-chloro-1-butene is produced.¹²

$$(CH_3)_2CHCl \xrightarrow{C_1} (CH_3)_2CH\ddot{C}Cl \text{ and } CH_3CHClCH_2 = \ddot{C}H$$

$$\downarrow \qquad \qquad \downarrow$$

$$(CH_3)_2C = CHCl \qquad CH_3CHClCH = CH_2$$

Substrates possessing C-H bonds β to the halogen undergo cyclizations to cyclopropanes;¹³ γ insertion to yield cyclobutanes is not observed.



The acetylene yields from ethyl halides are much too high to be attributed solely to hydrogen abstractions by C₂ from the arc (*vide infra*); another precursor is clearly indicated. An insertion-fragmentation mechanism whereby the C₂H₂ comprises monoatomic carbon from the arc and the α carbon of the substrate is supported by labeling studies.¹²

$$CH_{3}^{14}CH_{2}X + C_{1} \longrightarrow CH_{3}^{14}CH \longrightarrow CH_{3} + X + H^{14}C = CH$$

Labeling of the β carbon produces essentially nonradioactive C₂H₂. This contrasts with the results using atomic carbon produced by nuclear recoil whereby 52% of the C₂H₂ was derived from the β carbon and 48% from the α carbon.¹⁴ The fragmen-

(13) P. S. Skell and R. F. Harris, J. Amer. Chem. Soc., 87, 5807 (1965).

tation processes exhibit a dependency on the dissociation energies of the bonds being cleaved and become increasingly important as the bonds decrease in strength; thus, C_2H_2 yields increase in the series $EtCl < EtBr < EtI.^{12}$

Olefins. Ethylene reacts with atomic carbon to form allene (25%) and propyne (8.8%);¹⁵ but these are also products of hydrogen abstraction by C_3 .¹⁶ Enrichment of the arced carbon with radioactive ¹⁴C showed that C_3 is the precursor of one-fifth of the allene and one-eleventh of the propyne and the remainder is derived from C_1 . These figures are obtained by comparing the relative molar activities of the products with that of 1-pentyne whose relative molar activity is taken as 3.0 since it is derived solely from C_3 and so contains three carbon atoms from the arc.¹⁵

The major products from the propene reaction are explicable by (a) addition to the double bond forming a cyclopropylidene carbene which collapses to an allene or (b) C-H insertion followed by either rearrangement or dehydrogenation.



As further evidence of the cyclopropylidene carbene intermediate, bromotrifluoroethylene yields not only the expected allene but also 1-bromo-2,3,3-trifluorocyclopropene¹⁷



The products containing one extra carbon atom are thought to be derived from excited-state carbon atoms (^{1}S or ^{1}D) since they are not formed when resistively heated graphite (a source of predominantly ground state atoms) is used.

Isobutylene reacts with carbon atoms, producing 1,1-dimethylallene and isoprene in 50 and 31% yields; an adduct consisting of one carbon atom and two olefin molecules is isolable in 3% yield.



cis- and trans-2-butenes react with carbon atoms

(15) P. S. Skell, J. E. Villaume, J. H. Plonka, and F. A. Fagone, J. Amer. Chem. Soc., 93, 2699 (1971).

(16) P. S. Skell and R. F. Harris, J. Amer. Chem. Soc., 91, 699 (1969).

(17) M. J. McGlinchey, T. Reynoldson, and F. G. A. Stone, Chem. Commun., 1264 (1970).

⁽¹²⁾ L. Eng, Ph.D. Thesis, The Pennsylvania State University, University Park, Pa., 1970.

⁽¹⁴⁾ H. J. Ache and A. P. Wolf, J. Amer. Chem. Soc., 88, 888 (1966).

to give 2,3-pentadiene (50-60%) and also a mixture of the 1,3-pentadienes. The latter are formed with predominant retention of stereochemistry about the double bond, but some cis-trans interconversion does occur. The intermediates leading to isomerization of the double bonds are thought to be vibrationally excited crotylcarbenes.¹⁵



Oxygen- or sulfur-containing compounds react with atomic carbon by abstraction to form carbon monoxide¹⁸ or carbon monosulfide;¹⁹ the active species in this process is the metastable singlet carbon atom since thermally vaporized carbon, which contains ground-state C_1 , C_2 , and C_3 , gives no deoxygenation. Deoxygenation of aldehydes or ketones of formula $C_nH_{2n}O$ produces carbenes which can isomerize to olefins or cycloalkanes, C_nH_{2n} , as typified.¹⁸



In these deoxygenation reactions in which the carbon monoxide and carbenes are produced at -196° , the product distributions are nearly identical with those from alkylcarbenes made at 160° by diazo thermolyses.²⁰

The intermediacy of carbenes in deoxygenations is demonstrable by olefin trapping in some cases. Phosgene and methyl formate react with carbon atoms to yield dichlorocarbene and methoxycarbene, respectively, and these may be trapped *via* stereospecific additions to olefins at -196° .²¹



In contrast, dimethylcarbene (from acetone) and methylmethoxycarbene (from methyl acetate) rearrange intramolecularly too fast for intermolecular addition to occur.²²

(18) P. S. Skell, J. H. Plonka, and R. R. Engel, J. Amer. Chem. Soc., 89, 1748 (1967).

- (19) P. S. Skell and K. J. Klabunde, J. Amer. Chem. Soc., 93, 3807 (1971).
 - (20) L. Friedman and H/Shecter, J. Amer. Chem. Soc., 81, 5512 (1959).
 - (21) P. S. Skell and J. H. Plonka, J. Amer. Chem. Soc., 92, 2160 (1970).
- (22) J. H. Plonka and P. S. Skell, Tetrahedron Lett., 4557 (1970).

$$\begin{array}{cccc} (\mathrm{CH}_2)_2\mathrm{C} = \mathrm{O} & \xrightarrow{:\mathrm{C}:} & (\mathrm{CH}_3)_2\mathrm{C}: & \longrightarrow & \mathrm{CH}_3\mathrm{CH} = \mathrm{CH}_2 + & \mathrm{CO} \\ \mathrm{CH}_3\mathrm{O}\mathrm{C}\mathrm{C}\mathrm{H}_3 & \xrightarrow{:\mathrm{C}:} & \mathrm{CH}_3\mathrm{O}\mathrm{C}\mathrm{H}_3 & \longrightarrow & \mathrm{CH}_3\mathrm{O}\mathrm{C}\mathrm{H} = \mathrm{CH}_2 + & \mathrm{CO} \\ & & \parallel & & \\ \mathrm{O} \end{array}$$

Deoxygenation of acetic acid produces hydroxycarbene; this unique carbene yields acetaldehyde. The mechanism of acetaldehyde formation has been elucidated using $CH_3-CO_2D.^{22}$



Atomic carbon also deoxygenates ethers. Diethyl ether gives CO, C_2H_4 (40%), C_2H_6 (35%), and traces of n- C_4H_{10} presumably, via formation and disproportionation of two ethyl radicals from each ether molecule. Tetrahydrofuran gives ethylene (27%, based on 2 mol of C_2H_4 per mol of C_1 reacted).¹⁸

$$\begin{array}{c} \swarrow \\ 0 \end{array} + :C: \longrightarrow CO + \swarrow \longrightarrow 2C_2H_4 \end{array}$$

Epoxides are deoxygenated in a nonstereospecific fashion, possibly via the formation and ring opening of an ylide.²³



A similar rationale can account for the nonstereospecificity of the deoxygenation of oxetanes to cyclopropanes whereby the intermediate diradical(s) is free to rotate before ring closure occurs.²⁴



On the basis of bond energies, deoxygenation is a highly exothermic process ($\sim 85-100$ kcal/mol),

⁽²³⁾ P. S. Skell and J. H. Plonka, Chem. Commun., 1108 (1970).
(24) P. S. Skell, J. H. Plonka, and K. J. Klabunde, Chem. Commun., 1109 (1970).

whereas desulfurization is much less so (~45 kcal/ mol).²⁵ One would thus anticipate less energetic intermediates in the latter process, and this is reflected in the increased stereospecificity of desulfurization of episulfides.¹⁹ Thus *cis*-1,2-dimethylethylene episulfide gives both *cis*- and *trans*-2-butene with a cis: trans ratio of 4.5; the corresponding epoxide deoxygenates, yielding both 2-butenes but with a cis: trans ratio of 1.5.²³ Furthermore, desulfurization of tetrahydrothiophene produces not only ethylene (91%) but also cyclobutane (9%).¹⁹ Deoxygenation of tetrahydrofuran presumably generates an energy-rich diradical which fragments rather than cyclizes. The intermediacy of carbon monosulfide in desulfurizations is demonstrable by trapping it with oxygen at -196° to form carbonyl sulfide.¹⁹

$$S = C = S + :C: \longrightarrow 2CS \xrightarrow{O_2} O = C = S$$

Singlet carbon atoms bring about decomposition of aziridines to olefins and HCN.²⁶ A scheme which accounts for the nonstereospecificity of the reaction is presented below.



Reactions of Diatomic Carbon

Mechanism of Acetylene Formation. Acetylene is a product from the reaction of arced carbon with most hydrogen-containing compounds. As was shown earlier,¹² acetylene can arise by an insertion-fragmentation reaction of atomic carbon with alkyl halides. For that case, using an arc enriched with ¹⁴C, it was shown that only one carbon in the acetylene was derived from the arc. However, cocondensation of ¹⁴C-enriched carbon vapor with a variety of hydrocarbons, alcohols, aldehydes, or ketones reveals that the acetylene (and also ethylene) has a relative molar activity of ~ 2 compared to that of a product unambiguously derived from $C_{1.27}$ Conversely, reaction of carbon vapor with ¹⁴C-enriched substrates produces acetylene and ethylene of low relative molar activity.² In these cases, both carbon atoms of the acetylene are arc derived.

The ground state of gaseous C_2 is the singlet $x \, {}^{1}\Sigma_{g}{}^{+}$ with a metastable triplex $x' \, {}^{3}II_{u}$ state lying only 1.74 kcal higher than the singlet ground state.^{28,29} The reaction of C_2 with an equimolar mixture of CH₃CHO and CD₃CDO gives C_2H_2 (46%), C_2HD (14%), and C_2D_2 (40%). If abstraction of hydrogen (or deuterium) occurred randomly in a two-step

(27) P. S. Skell and R. F. Harris, J. Amer. Chem. Soc., 88, 5933 (1966).

intermolecular process, one would expect C_2H_2 , C_2HD , and C_2D_2 in a 1:2:1 ratio (eq 2).

$$C = C \cdot \underbrace{CH_{3}CHO}_{CD_{3}CDO} + C = C \cdot \underbrace{CH_{3}CHO}_{CD_{3}CDO} + HC = CH \quad (25\%)$$

$$HC = C \cdot \underbrace{CH_{3}CHO}_{CD_{3}CDO} + HC = CD \quad (50\%) \quad (2)$$

$$DC = C \cdot \underbrace{CH_{3}CHO}_{CD_{3}CDO} + DC = CD \quad (25\%)$$

The C₂HD (14%) is attributed to such an intermolecular reaction; thus, 28% of the acetylene is produced by an intermolecular process. The remaining high yields of C₂H₂ and C₂D₂ clearly indicate that most of the acetylene (72%) is formed by an intramolecular diabstraction process.

Radicals are known to selectively abstract the acyl hydrogen from acetaldehyde.³⁰ Triplet C₂ would be expected to behave as a diradical, and thus the intermolecular reaction is attributed to this form of C₂. Determination of the intermolecularity of reactions of C₂ with a variety of labeled substrates shows that 25-30% of the C₂ derived from the 16-V ac arc is in the metastable triplet state.^{9,31}

 CD_3CHO yields acetylene of the following composition: C_2H_2 (28%), C_2HD (23%), and C_2D_2 (49%). C_2H_2 arises solely by successive abstraction of acyl hydrogens by triplet C_2 in a manner characteristic of a radical species (eq 3).

$$C = C \cdot (t) + CD_{3}CHO \longrightarrow CD_{3}\dot{C} = O + HC = C \cdot$$

$$C = CH + CD_{3}CHO \longrightarrow CD_{3}\dot{C} = O + HC = CH (28\%)$$
(3)

The singlet C_2 is shown to react by two intramolecular pathways, C_2D_2 (49%) is made by a 1,1-diabstraction while C_2HD is generated by a 1,2-diabstraction process. Similar conclusions may be drawn from the results using using the inversely labeled acetaldehyde-1-d.³²



The duality of mechanism for acetylene formation has also been demonstrated in the reaction of C_2 with CH_3COCD_3 , which produces C_2H_2 (42%), C_2HD (18%), and C_2D_2 (40%).³² The 1,1 diabstrac-

⁽²⁵⁾ Required bond dissociation energies were taken from B. deB. Darwent, Nat. Stand. Ref. Data Ser., Nat. Bur. Stand., No. 31 (1970).
(26) J. E. Villaume, Ph.D. Thesis, The Pennsylvania State University,

⁽²⁶⁾ J. E. Villaume, Ph.D. Thesis, The Pennsylvania State University, University Park, Pa., 1971.

 ⁽²⁸⁾ E. A. Ballik and D. A. Ramsay, J. Chem. Phys., 29, 1418 (1958).
 (29) E. A. Ballik and D. A. Ramsay, Astrophys. J., 137, 61, 84 (1963).

⁽³⁰⁾ C. Walling, "Free Radicals in Solution," Wiley, New York, N. Y., 1957, p 279.

⁽³¹⁾ P. S. Skell, J. H. Plonka, and R. F. Harris, Chem. Commun., 689 (1970).

⁽³²⁾ P. S. Skell and J. H. Plonka, J. Amer. Chem. Soc., 92, 5620 (1970).

tion of hydrogen from acetone by singlet C_2 produces acetylcarbene, which readily undergoes Wolff rearrangement to form methylketene. This ketene is trapped by methanol to make methyl propionate.

Singlet acetylcarbene generated by photolysis of dia-

zoacetone at -196° in a methanol matrix is known to

 $CH_3COCH_3 + C = C \rightarrow C_2H_2 + CH_3COCH$

vield only methyl propionate.¹¹

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possible mechanistic rationale is shown in Scheme III. Ethylene, in contrast, having no readily abstractable hydrogens, yields 1-butane, 1,2-butadiene, and 1-buten-3-yne in nearly equal amounts and a trace of methylenecyclopropane. Other products isolated are 1,5-hexadiyne, 1,2-hexadien-5-yne, and 1,2,4,5-hexatetraene, which all contain four atoms from the arc.

Scheme III



However, the yields are too high to be attributable to C₄, and it is postulated that they are derived from one molecule of ethylene and two C₂ moieties.²⁶

Reactions of Triatomic Carbon

Triatomic carbon was first recognized in the tail of a comet.³⁴ The ${}^{1}\Sigma_{g}$ ⁺ ground state of C₃ has been calculated to be a linear, symmetrical singlet;³⁵ an excited singlet state, ${}^{1}\Sigma_{u}{}^{-}$, and several metastable triplet states, ${}^{3}\Delta_{u}$, ${}^{3}\Sigma_{u}{}^{+}$, ${}^{3}II_{u}$, and ${}^{3}\Sigma_{u}{}^{-}$, have been postulated, 36 but none have been correlated with any of the known spectroscopic bands. The C3 emitted from the 16-V ac arc is 10-14% by weight of the carbon vapor, a mixture of ground-state singlet, triplet, and probably some metastable singlet,² but thermally vaporized carbon contains more than 33% C₃, exclusively ground-state singlet.³

 C_3 reacts with organic compounds in three ways: addition to carbon-carbon multiple bonds, insertion into O-H bonds, and hydrogen abstraction.

Bisethanoallenes from Olefins. Triatomic carbon adds to carbon-carbon multiple bonds as a dicarbene;³⁷ the product of the addition of one C₃ molecule to two olefin molecules is a bisethanoallene. Bisethanoallenes have two cyclopropane rings lying in orthogonal planes, so that, when viewed along the long axis of the molecule, the planes of the rings are represented by perpendicular lines and the substituents lie to either side of these lines (Scheme IV). For



⁽³⁴⁾ A. E. Douglas, Astrophys., J., 114, 466 (1951).

(35) K. Clusius and A. E. Douglas, Can. J. Phys., 32, 319 (1954)

 $CH_{3}CO\ddot{C}H \longrightarrow CH_{3}CH = C = 0 \xrightarrow{CH_{3}OH} CH_{3}CH_{3}CO_{7}CH_{3}$ Addition and Insertion Reactions. Although formation of acetylene is the major reaction of C_2 with most substrates, a number of addition reactions are known to occur in low yields. Alcohols yield ethynyl and vinyl ethers as well as small quantities of ethyl ethers; all were shown to contain two carbon atoms from a ¹⁴C-enriched arc.² The reaction probably proceeds via initial attack on the oxygen followed by



either hydrogen migration or abstraction.

C₂ undergoes insertion reactions with hydrogencontaining substrates to produce allenes and acetylenes.³³ Typically, isobutane gives 4-methyl-1,2-pentadiene, dimethyl ether yields methoxyallene and 3methoxypropyne,³³ and propane produces a mixture of allenes.

$$(CH_3)_3CH + C_2 \longrightarrow (CH_3)_2CHCH = C = CH_2 \quad (10\%)$$

$$CH_3OCH_3 + C_2 \longrightarrow CH_3OCH = C = CH_2 \quad (30\%)$$

$$CH_3CH_2CH_3 + C_2 \longrightarrow CH_3CH_2CH = C = CH_2 \quad (5\%) + (CH_3)_2C = C = CH_2 \quad (1\%)$$

Reaction of C_2 with a mixture of C_3H_8 and C_3D_8 showed that the allenes are formed intramolecularly,³³ and the postulated mechanism involves a 1,1 diabstraction of hydrogen by C_2 to produce an alkylcarbene and vinylidene. The latter can rearrange to acetylene or rotate through 180° (rate ~ 10^{11} sec⁻¹) and couple to produce the allenes. It is noteworthy that only by this fast-trapping method have carbenes with β hydrogens been trapped; in the presence of all other reagents they rearrange to olefins.

$$RCH_3 + C_2 \longrightarrow [RCH + H_2C=C:] \longrightarrow$$
$$[RCH + :C=CH_2] \longrightarrow RCH=C=CH_2$$

Labeling experiments with ¹⁴C show that C₂ adds to olefins, the course of the addition being largely dependent on the presence of allylic hydrogens in the substrate. Although the major C₂ product with propene is C_2H_2 , low yields of other products derived from C_2 and one olefin molecule are found, and a

(33) P. S. Skell, F. A. Fagone, and K. J. Klabunde, J. Amer. Chem. Soc., 94, 7862 (1972).

⁽³⁶⁾ K. S. Pitzer and E. Clementi, J. Amer. Chem. Soc., 81, 4477 (1959).

⁽³⁷⁾ P. S. Skell and L. D. Wescott, Jr., J. Amer. Chem. Soc., 85, 1023 (1963).

example, 1,1,1',1'-tetramethylbisethanoallene, the product from the reaction of C₃ with two molecules of isobutylene, is shown in eq 4.

$$C_3 + 2 \rightarrow \rightarrow \qquad (4)$$

The reaction of C_3 with propene produces three diastereomers of 1,1'-dimethylbisethanoallene, which have the ratio 1:2:1.¹ The eight possible configurations of this product are shown in Scheme V. Config-





urations $a \equiv d$ and $e \equiv h$ are an enantiomeric pair, b and g are a second enantiomeric pair, and c and f are a third such pair. Random additions account for the ratios of the three diastereomers observed, 1:2:1.

The reaction of C_3 from arced graphite (11 V ac) with *cis*-2-butene produces two bisethanoallenes, j and k, in the ratio $93:7.^1$ trans-2-Butene makes



three diastereomers, k, m, and n, in the ratio 6:47:47. When the triatomic carbon is obtained



from thermally vaporized graphite, product k is not found in the reaction of either *cis*- or *trans*-2-butene. Compound k is the only diastereomer which has one pair of methyl groups cis and the other trans.

Ground-state singlet C_3 , which is obtained from thermally vaporized graphite, adds stereospecifically to both *cis*- and *trans*-2-butene (eq 5). On the other



 hand, the mixed bisethanoallene can only arise from addition of the metastable triplet C_3 , which adds nonstereospecifically in its second addition step (eq 6). The amount of triplet C_3 is twice the percentage

$$\|c = c = c\| \longrightarrow c = c = c\| \rightarrow j(50\%) + k(50\%)$$

$$\|c = c = c\| \longrightarrow c = c = c\| \rightarrow (6)$$

k(50%) + m(25%) + n(25%)

of mixed bisethanoallene, k. The fraction of triplet C_3 from arced carbon increases from 14 to 60% as the arc voltage is increased from 11 to 14 V.

Alcohols react with C_3 to give as a major product the 3,3-dialkoxypropynes.¹⁶ The C_3 insertion probably takes the course shown in eq 7. None of the 1,3-

$$C_{3} + ROH \longrightarrow$$

$$[ROCH = C = C \leftrightarrow ROCH - C = \overline{C} \leftrightarrow ROC - C = CH]$$

$$\downarrow ROH \qquad (7)$$

$$(RO)_{0}CH - C = CH$$

dialkoxyallenes are detected, but lack of an authentic sample precluded demonstration that these compounds would survive work-up conditions.

Hydrogen Abstraction by C_3 . Products formed from arc-generated C₃ by hydrogen abstraction are mostly propyne, allene, propene, and traces of propane.¹⁶ These products are not found with thermally produced carbon vapor; this suggests a metastable excited-state C₃ as the precursor. Use of ¹⁴C-labeled electrodes using alcohols as substrates shows that nearly all of the propyne and allene and much of the propene is derived from C3. The hydrogen abstraction reactions may be further elucidated by the results of the reaction of C₃ with a 1:1 mixture of cyclohexane- d_0 and $-d_{12}$.² The major products are allene $(43\% d_0; 3\% d_1; 1\% d_2; 5\% d_3; 48\% d_4)$ and propyne (40% d₀; 12% d₁; 7% d₂; 20% d₃; 21% d₄). Similar deuterium distributions are found when 1:1 mixtures of other non- and perdeuterated substrates are used. For both allene and propyne, three hydrogen atoms (and to a large extent, a fourth) are abstracted from the same substrate molecule-an unexpected result. The possible precursor of allene is the metastable singlet, ${}^{1}\Sigma_{u}$, whereas a metastable triplet state could account for propyne formation.

Reactions of Tetraatomic Carbon

Tetraatomic carbon is predicted to have a triplet ground state with several excited singlet and triplet states.³⁸ The amount of C₄ produced by the 16-V arc is less than 2% by weight.² Tetraatomic carbon reacts with hydrogen-containing substrates to make ethylacetylene, methylallene, 1,3-butadiene, and vinylacetylene, with trace amounts of diacetylene.³⁹ These products were shown to originate from tetraatomic carbon by radioactive labeling experiments with carbon-14 vapor. Carbon vapor from thermally

(38) E. Clementi, J. Amer. Chem. Soc., 83, 4501 (1961).

⁽³⁹⁾ R. F. Harris and P. S. Skell, J. Amer. Chem. Soc., 90, 4172 (1968).

vaporized graphite produces none of these C_4 hydrocarbons.

Use of Inorganic Substrates

A number of laboratories, notably those of Schaeffer (Indiana) and Stone (Bristol), have extended the scope of carbon vaporizations to include reactions with inorganic substrates; in these cases, emphasis was placed more on synthetic utility than on elucidation of mechanisms.

Boron. Pentaborane-9 reacts with carbon vapor to yield carbahexaborane-7, B_5CH_7 , together with traces of $B_5C_2H_7$.⁴⁰ The yield of B_5CH_7 (3% based on substrate consumed) is more than 20 times greater than the yield by the previous synthesis.⁴¹



Reaction of carbon vapor with diboron tetrachloride produces mainly involatile polymers, but three moisture- and temperature-sensitive dichloroboryl derivatives of methane are isolable.⁴² They probably arise *via* insertion of carbon into a B-B bond to give a bis(dichloroboryl) carbene intermediate which can either abstract chlorines or further insert into B-B or B-Cl bonds as indicated.



Boron trichloride reacts with C_1 to give dichlorobis(dichloroboryl)methane, $(BCl_2)_2CCl_2$, by successive insertions into B-Cl bonds. BCl₃ is known to decompose to B₂Cl₄, and this may account for the isolation of $(BCl_2)_3CCl$ and B₂Cl₄ from this reaction. C_2 also reacts with BCl₃ to give 1,2-dichloro-1,2-dichloroborylethylene.

In contrast, diboron tetrafluoride shows products derived solely from insertion into the B-B bond.⁴² This reflects the difference in bond dissociation energies in B₂Cl₄ ($D_{B-B} = 59$ kcal/mol) and in B₂F₄ ($D_{B-B} = 36$ kcal/mol).²⁵ Thus, products are isolable from reaction of C₁, C₂, and C₃ with two molecules of B₂F₄. Analogous compounds of C₂ and C₃ are presumably formed in the B₂Cl₄ reaction, but their involatility precludes their isolation.

$$(BF_{2})_{4}C \xrightarrow{C_{1}} B_{2}F_{4} \xrightarrow{C_{2}} (BF_{2})_{2}C = C(BF_{2})_{3}$$
$$\downarrow C_{3}$$
$$(BF_{2})_{2}C = C = C(BF_{2})_{2}$$

The major products from the reaction with methylboron dichloride are C_2H_2 , CH_3Cl , and CH_2 =CHCl which are formed by abstraction processes. However, 2,2-bis(dichloroboryl)propane, $(CH_3)_2C(BCl_2)_2$, is formed in good yield, indicating that the B-C bond is attacked in preference to the B-Cl bond.⁴² A scheme showing the formation of some of the minor products is presented

$$\begin{array}{cccc} BCl_2CCH_3 & \xleftarrow{C_1} & BCl_2CH_3 & \xleftarrow{C_3} & \xleftarrow{CH_3} & C=C=C & \xleftarrow{CH_3} \\ & & \downarrow & \downarrow & c_2 & BCl_2 & C=C & \swarrow \\ BCl_2CHClCH_3 & CH_3(BCl_2)C=C: & & & & \\ & & & CH_4(BCl_2)C=CHCl & + & CH_4(BCl_2)C=CCl_3 & & & \\ \end{array}$$

Carbon vapor abstracts hydrogens from trimethylboron to give acetylene, ethylene, ethane, and diacetylene.⁴² Tetramethyldiborane is also formed and is thought to arise by an insertion-fragmentation process similar to that previously demonstrated for alkyl halides.¹²

Silicon. Silane and carbon vapor give the usual hydrogen abstraction products, and also methylsilane, disilane, and benzene. Methylsilane arises by Si-H insertion followed by hydrogen abstraction; disilane and benzene may be formed in gas-phase reactions since the high volatility of SiH₄ causes high pressures during arcing.⁴³

The major C_1 product from trimethylsilane is

⁽⁴⁰⁾ S. R. Prince and R. Schaeffer, Chem. Commun., 451 (1968).

⁽⁴¹⁾ T. Onak, R. Drake, and G. Dunks, J. Amer. Chem. Soc., 87, 2505 (1965).

⁽⁴²⁾ J. E. Dobson, P. M. Tucker, R. Schaeffer, and F. G. A. Stone, J. Chem. Soc. A, 1882 (1969).

⁽⁴³⁾ J. Binenboym and R. Schaeffer, Inorg. Chem., 9, 1578 (1970).

vinyldimethylsilane (11.8%); this could be formed via two mechanisms, silicon-hydrogen insertion followed by formation and rearrangement of a silacyclopropane or carbon-hydrogen insertion followed by hydrogen migration. Use of $(CH_3)_3SiD$ as substrate enables one to differentiate between these possibilities, since, in the former case, 25% of the deuterium label ends up on the silicon, and in the latter case, 100% of the deuterium label remains on the silicon.



Experimentally, over 98% of the deuterium was still attached to silicon. Therefore, most of the vinyldimethylsilane is produced by C-H insertion.⁴⁴ Trimethylsilane and carbon yield minor products from both C-H and Si-H insertion as shown in Scheme VI.

 C_2 reacts with a methyl group of trimethylsilane to produce dimethylsilylallene in a similar manner to that previously discussed³³ for alkanes. Ethynyl-,



(44) P. S. Skell and P. W. Owen, J. Amer. Chem. Soc., 94, 1578 (1972).

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vinyl-, and ethyltrimethyl-silanes are also formed. Also, di-insertion products of C_2 and C_3 have been observed.

$$2(CH_3)SiH \xrightarrow{C_2} (CH_3)_3SiCH = CHSi(CH_3)_3 + (CH_3)_3SiCH_2CH_2Si(CH_3)_3$$

Tetrachlorosilane reacts with C_2 to produce the novel compound chlorotrichlorosilylacetylene.⁴³

$$Cl_4Si + C_2 \longrightarrow Cl_3SiC(Cl) = C: \longrightarrow Cl_3SiC = CCl$$

Miscellaneous. Germanium tetrachloride and phosphorus trichloride react in a similar manner with carbon vapor. Monoatomic carbon inserts into the metalloid-chlorine bond to form an intermediate carbene which can abstract chlorines or reinsert into another substrate molecule.⁴⁵

$$\begin{aligned} \operatorname{GeCl}_{4} + \mathbb{C}_{1} &\longrightarrow \operatorname{Cl}_{3}\operatorname{GeCCl} &\longrightarrow \\ & \operatorname{Cl}_{3}\operatorname{GeCCl}_{3}\left(1\,\%\right) + (\operatorname{Cl}_{3}\operatorname{Ge})_{2}\operatorname{CCl}_{2}\left(10\,\%\right) \\ & \operatorname{PCl}_{3} + \mathbb{C}_{1} &\longrightarrow \operatorname{Cl}_{2}\operatorname{PCCl} &\longrightarrow \operatorname{Cl}_{2}\operatorname{PCCl}_{3}\left(5\,\%\right) + (\operatorname{Cl}_{2}\operatorname{P})_{2}\operatorname{CCl}_{2}\left(1\,\%\right) \end{aligned}$$

Products from C_2 and GeCl₄ or parallel those previously described for BCl₃.⁴²

$$GeCl_4 + C_2 \longrightarrow Cl_3GeC(Cl) = C: \longrightarrow Cl_3GeC(Cl) = CCl_2$$

Carbon atoms insert into an S-Cl bond of sulfur monochloride and can then either abstract chlorines or undergo intramolecular rearrangement with subsequent extrusion of sulfur to form thiophosgene.⁴⁵

$$\begin{array}{cccc} \text{CISSCI} + \text{C}_{1} \longrightarrow \text{Cl} \\ \text{CI} \\ \hline & \begin{array}{c} \text{CI} \\ \hline & \begin{array}{c} \text{CI} \\ \hline & \end{array} \\ \hline \end{array} \\ \hline \end{array} \\ \hline \end{array}$$

(45) M. J. McGlinchey, J. D. Odom, T. Reynoldson, and F. G. A. Stone, J. Chem. Soc. A, 31 (1970).