Unsaturated Carbenes

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Received October 17, 1977 (Revised Manuscript Received April 17, 1978)

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/. Introduction

The term "unsaturated carbene" properly refers to a reactive intermediate where the electron-deficient carbon is an integral part of an unsaturated system as in 1 and 2. To date, two main classes of such species are known: alkylidene carbenes (1) and vinylidene carbenes (2). Besides these two species, C_2 (3) and C_3 (4) belong to this class of intermediates. Despite the wide

$$
\begin{array}{c}\nR \\
R\n\end{array}\n\begin{array}{c}\nC = C: & C = C - C: & C = C - C:\n\end{array}
$$

occurrence of C_2 and C_3 in the Universe, 1 mostly in comets and carbon stars, their somewhat unusual terrestrial generation via flames and carbon arcs places them outside of every day organic chemistry. Furthermore, they have been recently reviewed,² and therefore this report will deal only with alkylidene and vinylidene carbenes.

Although unsaturated carbenes have not been seen as stable entities nor to date observed spectroscopically, their existence is clearly demonstrated by their chemical behavior. Pioneering work, primarily by Hartzler, Köbrich, Newman, and their col-

laborators, 3 which established unsaturated carbenes as viable reaction intermediates was carried out during the 1960s. The characterization of their exact nature and the full development of their chemistry and uses are still continuing. Compared to the activity and plethora of results in saturated carbenes,⁴ unsaturated carbene chemistry is still in its infancy. Hence, it is a particularly appropriate time for a critical review and perhaps some stimulus and influence on the future growth and development of this area of chemistry.

This review will treat in separate sections the generation, nature, and reactions of alkylidene (1) and vinylidene (2) carbenes, respectively. A separate part will deal with the increasing number of transition metal complexes of unsaturated carbenes, along with a brief summary and extrapolation into the future.

//. Generation of Alkylidene Carbenes

A. α -Elimination Methods

The most general method for the generation of 1 is α -elimination from a variety of vinylic progenitors. The most widely investigated of these reactions are α -eliminations of 1-haloolefins with bases. The base-promoted reaction of arylvinyl halides, the so-called Fritsch-Buttenberg-Wiechell rearrangement,⁵ used to produce the corresponding acetylenes has been known since the 1890s, although the exact nature of the reaction was not established until much later (see section IV.A).

 α -Elimination from a wide variety of primary alkylvinyl halides 5 and strong bases has been observed, usually in olefins that serve as both substrate and solvent. These reactions are gen-

erally accompanied by isomerization of the starting vinyl halides and tar formation with the carbene-derived methylenecyclopropane 6, products being formed to the extent of only 20-40%. These reactions proceed via carbenoids and organometallic intermediates rather than free carbenes (see section III.B).

In the absence of olefins or other suitable traps, butatriene products such as 7⁷ and 8⁸ have been observed. It is unlikely that

these products arose via dimerization⁹ of the respective carbenes. More likely is the interaction of two anions, 9, with for-

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 $(C_6H_5CH_2)_2C$ = CHCI

$$
\xrightarrow{\text{RLi}} (C_6H_5CH_2)_2C = C = C = C(CH_2C_6H_5)_2
$$

mation of 10 and subsequent loss of LiX; or trapping of the carbene by the anion, with formation of 11 and subsequent loss of $X^{-,9}$

Similar triene products have recently been observed in the interaction of phenyllithium with dialkylthioketenes¹⁰ 12 and thioalkynes¹¹ 13 with organocopper reagents. Although the possible involvement of alkylidene carbenes 1 in these reactions cannot be completely ruled out, other processes are more likely:

1,1-Dibromoolefins and alkyllithiums readily underwent halogen-metal exchange, α -elimination, and methylenecyclopropane formation:¹²

Nearly quantitative carbene-derived products were obtained by the use of primary vinyl trifluoromethanesulfonates¹³ (triflates) 14 and t-BuOK.¹⁴ Along with the olefin adducts, tert-butyl vinyl ethers 15 were observed as minor products.

Reactions were carried out under heterogeneous or homogeneous conditions by the addition of anhydrous glyme and/or crown ethers¹⁵ to solubilize the base. Primary vinyl triflates are readily available¹⁴ from aldehydes via their silyl enol ethers.¹⁶

Recently an interesting indirect α -elimination route to 1 was reported by fluoride ion initiated nucleophilic attack upon α chlorovinylsilane¹⁷ 16 and silylvinyl triflate¹⁸ 17. Reaction of 16

gave yields of up to 80% of carbene derived products at 25 $^{\circ}$ C in 48 h. Triflate 17 gave quantitative yields of carbene adducts with cyclohexene and ethyl vinyl ether respectively at —20 or 0 °C in 1-2 h with KF and crown ethers, with anhydrous $R_4N^+F^-$, or via KF and phase-transfer procedures.¹⁷ Chlorovinylsilanes such as 16 were prepared from vinyl halides and silylvinyl triflates such as 17 from the appropriate α -silyl ketones 18.¹⁹ lndirect evidence¹⁸ indicated a concerted nucleophilic attack upon Si and elimination of CF₃SO₃⁻ with 17 and a stepwise process in the case of 16. These two procedures are the best to date for the generation of alkylidene carbenes (1) under neutral and mild conditions.

B. Deamination and Related Procedures

Certain vinylamines such as 19, 20, 20 and 21 21 have been reported to give carbene-derived products upon reaction with alkyl nitrites:

$$
(\mathrm{C}_6\mathrm{H}_5)_2\mathrm{C}=\mathrm{CHNH}_2\overset{\mathrm{C}_6\mathrm{H}_{11} \mathrm{ONO}}{\longrightarrow} \mathrm{C}_6\mathrm{H}_5\mathrm{C} \equiv \mathrm{CC}_6\mathrm{H}_5
$$

19

These reactions proceed via the prior involvement of vinyldiazonium ions, 22. However, it has not been established

whether 22 first loses a proton to form unstable diazoethylenes 23 and subsequently loses N_2 to give carbene 1; or if 22 first loses N_2 to form a vinyl cation 24 and then a proton to give 1:

Although stable diazoethylenes 23 have not yet been reported, stable diazonium salts 22, such as 25 and 26, have been isolated and characterized.²² Their direct involvement in alkylidene carbene generation has not yet been investigated. Both routes

may be feasible, but formation of another high-energy reactive intermediate such as a primary vinyl cation²³ would, in this reviewer's opinion, seem less likely than involvement of diazoalkenes 23.

Since alkylvinylamines are not generally available, such deamination reactions are restricted to arylvinylamines. A more general process involving diazonium ions is lithium alkoxide initiated decomposition of W-nitrosooxazolidones 27, developed by Newman and co-workers.²⁴

Although the exact mechanism of carbene formation from 27 is not clearly understood,²⁵ the process is general for 5,5-disubstituted oxazolidones 27, yielding 40-60% of carbenederived products along with 40-60% of diols 29. Diols 29 presumably arise via 28.²⁴ To avoid such side products N-nitrosoacetylamino alcohols 30 were developed as carbene 1 progenitors.²⁶ Up to 83% of carbene-derived products have been obtained by reaction of 30 with NaOH under phase-transfer conditions.²⁶

Diazonium ions and diazo compounds have been invoked as intermediates, along with alkylidene carbenes 1 in the deamination of certain 2-(aminomethyl) aziridines 31:27a

C. Thermal and Photochemical Methods

Unfortunately there is no simple, general method for either the photochemical or thermal generation of unsaturated carbenes 1 or 2. The ideal progenitors for such photochemical alkylidene carbene generation would be diazoalkenes 32 or methylenediazirines 33. Neither of these two classes of com-

$$
(R)_2C = CN_2
$$

32
33
33

pounds has been isolated or reported,^{27b} although as mentioned in the preceding section diazoethylenes have been invoked as unstable intermediates in some alkylidene carbene formations.

A wide variety of precursors have been employed in the thermal and photochemical generation of alkylidene carbenes. The organomercurial 34 gave 20-25% of carbene derived products upon photolysis in cyclohexene or cyclohexane.²⁸

Photolysis of methylenecyclopropanes gave a methylenecyclopropane rearrangement reaction, ²⁹ and a cheletropic fragmentation with formation of small amounts of carbene derived products:³⁰

⊣₅)₂С — СН₂ + НС≡≡СН

Methylenecyclopropane itself gave ethylene and acetylene via mercury-sensitized photofragmentation.³¹ Difluorocarbene (35) has been generated by mercury-sensitized photolysis of either trifluoroethylene 32 (36) or perfluorovinylsilane 33 (37).

$$
\begin{array}{cccc}\nF_2C = C: & F_2C = CHF & CF_2 = CFSiF_3 \\
& 35 & 36 & 37\n\end{array}
$$

Alkynes, presumably via 1 (see section IV.A), were obtained in the thermolysis of a series of substituted tetrazoles 38:³⁴

Similarly, alkynes were observed in the thermal decomposition of aryl-substituted α -haloacrylate salts 39.^{35–37} The parent acrylate 40 gave acetylene and $CO₂$ as volatile products.³⁶ However, no carbene adduct was observed in the thermal reaction of 41 with diphenylethylene.³⁷

Azoethylene 42 gave N_2 , $CH_3C_6H_4SO_2H$, and diphenylacetylene in CCI₄ at room temperature.³⁸ Tosylazoalkenes 43 also gave N_2 and sulfinic acid along with 25-40% of carbenederived adducts when reacted with olefins at room temperature.¹⁸

Carbene adducts were also obtained by thermolysis of organomercurial 44³⁹ at 150 °C or silylsiloxyalkene¹⁸ 45 at 275 ^oC. The latter reaction presumably proceeds by deoxysilyla-

tion.⁴⁰ Excellent yields of arylacetylenes were obtained⁴¹ in the flash pyrolysis of isoxalones 46:

Arylacetylenes were also formed in the flash pyrolysis of a series of ring-substituted isopropylidene malonates 47:42

$$
\xrightarrow[0.02 \text{ mm}]{-600 \text{ C}} \text{XC}_6 H_4 C \equiv C H + C O_2 + C O + (CH_3)_2 CO
$$

64-98%

D. Miscellaneous Methods of Generation

New routes to alkylidene carbenes will undoubtedly be forthcoming. An interesting possibility on paper that does not seem to work in practice is deoxygenation of ketenes. Deoxygenation of isocyanates 48 to isonitriles 49, the chemical cousins of alkylidene carbenes, is a known process:⁴³

$$
RN = C = O + (EtO)3P \rightarrow R\ddot{N} = C: + (EtO)3PO
$$

48 49

Yet reaction of diphenylketene with triethyl phosphite gives a stable crystalline adduct 50, which decomposes to diphenyl-

$$
(C_6H_5)_2C = C = O + (EtO)_3P
$$
\n
$$
\longrightarrow \begin{array}{c}\nC_6H_5)_2C = C \\
\hline\nC_6H_5)_2C = C\n\end{array}
$$
\n
$$
\longrightarrow \begin{array}{c}\nC(C_6H_5)_2 \\
\hline\nD(OEt)_3 \\
\hline\n50\n\end{array}
$$
\n
$$
\xrightarrow{220°C} C_6H_5C = CC_6H_5 + (EtO)_3PO
$$

acetylene and triethyl phosphate only upon heating above 220 ^oC.^{44,45} Although 1 has been suggested as an intermediate in this reaction,⁴⁵ its actual involvement is uncertain in light of

results with dimethylketene.⁴⁶ Dimethylketene formed similar 2:1 phosphorane adducts 51 with a variety of trivalent phosphorus nucleophiles including $(CH_3O)_3P$. These adducts decomposed to lactone dimer 52 at various temperatures. No deoxygenation and no carbene products were observed.⁴⁶

///. Nature of Alkylidene Carbenes

One of the more interesting questions regarding any reactive intermediate is its exact nature. The nature of alkylidene carbenes may be defined in terms of spin multiplicity, free carbene or carbenoid character, electrophilicity or nucleophilicity, and steric factors involved in its reactions.

A. Spin Multiplicity

A priori, any carbene, including 1 and 2, may be in one of three low-energy states: a singlet state (S_0) in which one carbon orbital is empty and the second contains two unshared spin-paired electrons, 53; a singlet (S_1) with two singly occupied orbitals,

54; and the triplet state (T_1) in which two electrons with parallel spins singly occupy two carbon orbitals, 55. Methylene has been shown⁴⁷ to possess a triplet ground state, ³B₁, in agreement with theoretical calculations.⁴⁸ An energy vs. HCH bond angle profile for methylene shows⁴⁹ that the triplet state is 41.5 kcal/mol below the singlet state for a bond angle of 180°, 20.8 kcal/mol below that for an angle of 135° (the experimentally and theoretically predicted bond angle),^{47,48} and nearly degenerate for \sim 100°; for an HCH angle of \leq 90°, the singlet is more stable than the triplet. Since one can look upon 1 as a methylene with a very small bond angle, this analogy would predict the singlet to be the ground state for alkylidene 1, and for that matter vinylidene, 2, carbenes. Theoretical calculations of the Huckel type, 50 MINDO/2 procedures, 51 and most recently ab initio methods⁵² all predict a singlet ground state, 53, for alkylidene carbenes with a singlet triplet splitting of 20-46 kcal/mol. The $\frac{1}{2}$ calculated state of 1 (R = H) is calculated to be the $\frac{3}{2}$ state with an excitation energy of about 75 kcal/mol.⁵² Ab initio calwith an excitation energy or about 70 Adamnot. Ab finite data-
culations also predict a dipole moment of 2.23 D for the 14.7 (ground) state of 1 ($R = H$) in the direction of the lone pair electrons, a carbon-carbon bond dissociation energy of 162 kcal/ mol (compared with a dissociation energy at 169.9 kcal/mol for ϵ and ϵ and a ΔH^{298} (H $_{\rm c}$ CC:) = 100 kcal/mol.52 The C=C bond lengths are calculated to be 1.35 and 1.36 Å for the 14 bond lengins are calculated to be 1.55 and 1.56 A for the \mathcal{A}_1
pinglet ground state and ^{3B}, triplet state, respectively. The ionization potential for singlet 1 ($R = H$) is calculated to be 10.8 eV $(0,0,0)$ is contribution of the TT orbital); only 0.3 eV above the corresponding IP of out of the π

Experimentally, neither the ESR nor CINDP techniques commonly employed⁵³ to probe spin multiplicities of intermediates are compatible with the current methods of unsaturated carbene generation. The alternative technique, first proposed by Skell,⁵⁴ of examining the stereochemistry of carbene addition to olefins⁵⁵ has been employed. Reaction of isopropylidenecarbene (1, R = CH₃) with *cis*-2-butene was found to be $>$ 99.5% stereoselective and stereospecific with *trans*-2-butene, and more than 98 % stereoselective with each of the 2-methoxy-2-butenes.⁵⁶ Since the vinyl triflate produced carbene is known to be "free" rather than a carbenoid (vide infra), this datum suggests, with the usual caveat^{55,57} about relative life times of rotamer equilibration and spin relaxation in the triplet produced biradical 56, that the nascent carbene 1 is a singlet. The use of

an inert dilutent⁵⁸ (c-C₄F₈) did not alter this stereoselectivity, strongly suggesting⁵⁶ that the ground state for alkylidene carbene is the singlet, in accord with theoretical calculations.50-52 The slower reactivity⁵⁹ of 1,3-butadiene compared to 2-butene (see Table III) provides additional experimental evidence for the singlet nature of 1. With a triplet ground state, the diradical resulting from the stepwise addition to 1,3-butadiene 57 should be more stable and more readily accessible than the simple diradical 56 resulting from an olefin. Stereospecific addition of 1 ($R = CH₃$), as generated from the nitrosooxazolidone 27, to cis - and trans-4-methyl-2-pentene has also been observed.⁶⁰ There is little doubt that alkylidene carbenes 1 are singlets with a fairly sizable singlet-triplet energy difference.

B. Encumbrance

In discussions of carbene encumbrance one must consider three major species: organometallic reagents 58, carbenoids

59, and "free" carbenes 60. Species 58 is basically an organometallic compound where the metal and halogen are essentially covalently bound to carbon, whereas in a carbenoid⁶¹ either the leaving group or the metal of the base used, or both, is associated with the carbene intermediate 59. A free carbene, 60, is not burdened by any such encumbrance and is usually encountered only under dilute gas-phase conditions. An entire spectrum of intermediates, with a concomitantly wide range of chemical behavior, is possible between the two extremes of 58 and 60.

Experimentally, intermediate 58 may be distinguished from 59 and 60 by its anion-like nucleophilic behavior in contrast to the usual electrophilic properties of carbenes. The degree of encumbrance and the distinction between 59 and 60 in turn is generally accomplished by examination of the chemical behavior, usually stereochemistry and selectivity, of a particular carbene generated from as many different progenitors and by as many different processes as possible. Identical chemical behavior under a wide variety of conditions implies an unencumbered species 60.

From the extensive and elegant work of Köbrich and coworkers,⁶² there is no doubt that use of vinyl halides and alkyllithium and similar bases results in an organometallic species 58, which by α -elimination of LiX converts to 59. A large number of organometallic intermediates of type 58 have been formed and characterized at low temperature in THF.⁶³ These species undergo nucleophilic reactions typical of organometallic compounds and can be readily trapped with $CO₂$, CH₃I, COCI₂, Br₂, and other nucleophiles as well as salts of metals less electropositive than the alkali metals.^{63,64} Upon warming, these species, 58, readily react with olefins and other electrophiles presumably via carbenoids 59.

In contrast, the behavior of the species generated from primary vinyl triflates, 14, strongly suggests the intermediacy of

SCHEME I. Reaction of an Unsymetrical Carbene with an Unsymetrical Olefin

TABLE I. Selectivity of $(CH_3)_2C=$ C: toward Styrenes $(XC_6H_4CH=CH_2)$

an unencumbered carbene, 60. Specifically, the degree of encumbrance of alkylidene carbenes with the leaving group, presumably through the empty p orbital, has been determined by the following method. Addition of an unsymmetrical carbene 63 to an unsymmetrical olefin such as isobutylene results in two distinct adducts 64 and 65, as well as, in the case of the vinyl triflate derived carbene, in two distinct insertion products 66 and 67, as shown in Scheme I. It is then a necessary, albeit not sufficient, condition for the intermediacy of the free carbene (free of association with the leaving group) that both pure isomeric starting triflates 61 and 62 give an identical ratio of adducts and insertion products. To put it differently, if there is any association with the leaving group there should be a memory effect that reflects itself in the product ratio as a function of the stereochemistry of the isomeric carbene progenitors. As the data¹⁴ in Scheme I indicate within experimental error, the product ratios were identical in accord with an intermediate free of association with the leaving group.

Association with the metal may be established by selectivity studies under diverse conditions and with different precursors. This has been accomplished for alkylidene carbenes with the results summarized in Table I. The close agreement (probably the same within experimental error) in the selectivities of $(CH₃)₂$ C= C : as derived from **14, 17, and 43** (R = CH₃) strongly suggests an unencumbered carbene 60 as the intermediate in these reactions.⁶⁹ The results with the crown ether as well as with precursor 43 (columns 2 and 3 of Table I, respectively) are

particularly significant. In the case of crown ethers one would expect the 18-crown-6-to be a much more effective complexing agent with K⁺ than with a carbene. In fact, Moss and co-workers⁷⁰ have shown crown ethers to be an effective means of generating unencumbered carbenes, identical in their selectivities with the species derived from thermal or photochemical decomposition of diazo precursors and greatly differing from the carbenoid selectivities. With precursor 43 ($R = CH₃$) no metal or base as well as a different leaving group is involved in the alkylidene carbene generation, precluding the possibility of any such associations, yet the selectivities are identical with the triflate 14 derived results with or without crown ethers. In contrast the alkyl halide derived carbene selectivities differ markedly⁶⁸ from the others, perhaps suggesting the involvement of a carbenoid, 59, rather than the free carbene, 60. The oxazolidone 27 derived species seems to be between the vinyl halide and vinyl triflate 14 derived carbene selectivities.

The difference in behavior and the different nature of the intermediates as derived from vinyl halides (carbenoids 59) and vinyl triflates (unencumbered carbenes 60) may be accounted for by the vastly superior leaving ability of $CF_3SO_3^-$ compared to $X^{-13,71,72}$ The 10⁸-10⁹ times greater reactivity of $CF_3SO_3^$ compared to halogens would suggest that α -elimination of $CF₃SO₃H$ from vinyl triflates 14 by any base is a concerted process proceeding directly to a free carbene, 60, whereas HX elimination from vinyl halides is a stepwise process first forming 58 then 59 and perhaps in special cases 60. This hypothesis is

Figure 1. Hammett σ - ρ plot of (CH₃)₂C= \equiv C: addition to $X\check{C}_6H_4CH=CH_2.$

supported by the existence⁷³ of an anionic equilibrium and reisolation of deuterated starting vinyl halides in t-BuOD with $(C_6H_5)_2C$ CHBr and the lack¹⁴ of such equilibrium in the presence of base and *t*-BuOD with vinyl triflates 14.

Hence, analogous to normal carbenes,⁴ there is an entire spectrum of alkylidene carbene intermediates ranging from organometallic species 58 through carbenoids 59 to less encumbered if not "free" carbenes 60.

C. Electrophilicity

Alkylidene carbenes, like any carbene, being electron deficient would be expected to be electrophilic. A Hammett plot of the vinyl triflate data in Table I gives a good correlation with a $p = -0.75$ as shown in Figure 1. Other relevant carbene selectivities toward styrenes are summarized in Table II. It is evident from these data that alkylidene carbenes 1 are electrophilic, as are most carbenes, with the exception of cycloheptatrienylidene, 78 and initially interact with olefin π systems via their empty p orbitals. It is also obvious from the data in Table Il that the vinyl halide 5 and oxazolidone 27 derived species 1 are much more⁶⁸ electrophilic than the vinyl triflate derived carbene. It is possible that these large differences in selectivi t is e^{68} reflect differences in the nature of the intermediates with the halide and oxazolidone derived species being carbenoids 59 and the vinyl triflate derived species 60 being unencumbered. However, even considering such differences, the ρ values reported for the vinyl halide and oxazolidone derived species are the largest for any known carbenic additions to styrenes and suggest an unusually highly polarized transition state for such reactions. These ρ values are comparable to the values observed for bromination⁷⁹ ($\rho = -4.3$) and hydration⁸⁰ ($\rho = -4.0$) of styrenes with a nearly fully developed charge in the transition state. It is true that alkylidene carbenes 1 are isoelectronic with the corresponding vinyl cations but so are normal carbenes isoelectronic with carbonium ions. Yet saturated carbenes, including the : $CH₂$ carbenoid, seem to be much less selective (ρ $=$ -1.6) than the species derived from 5 and 27. An interesting rationalization has been provided for the high selectivity of these alkylidene carbenoids by invoking a less effective back-donation of electron density by the lone pair carbenic electrons into the empty olefin 7r* orbital from alkylidene carbenes than from empty olefin π^* orbital from alkylidene carbenes than from saturated carbenes.⁸¹

The selectivity of the vinyl triflate 14 derived carbene implies that unencumbered alkylidene carbenes 60 are mildly electrophilic with a concerted addition to olefins with but a slightly polar transition state. As the data in Table Il indicate, 60 seems about as electrophilic as the halocarbenes or cyclopentadienylidene. As Seyferth and co-workers⁷⁴ have pointed out, singlet-state divalent carbon species (carbenes) are also isoelectronic with

i able ii. Carbene selectivities toward styrenes

TABLE III. Relative Reactivity of (CH³ J2C=C: with Olefins

a Relative to cyclohexene.

trigonal boron compounds that have been shown to react with styrenes⁸² and other olefins⁸³ by means of relatively nonpolar π -complex-like transition states.

Finally, organometallic intermediates 58 that can serve as precursors to carbenoids 59 display nucleophilic behavior typical .
of anions^{65,64} (see section IIIB).

D. Steric and Other Factors

Further insight into the nature of alkylidene carbenes has been gained by studies of the relative reactivity and regioselectivity of these species toward various olefins. Relative reactivities are summarized in Table III.

The relative reactivity data in Table III indicate the considerable steric requirements of alkylidene carbene-olefin interactions. Unlike most saturated carbenes that react⁸⁴ faster with electron-rich olefins like tetramethylethylene than with less substituted alkenes, alkylidene carbenes react slower. It was to account for these steric effects as well as the observation of the thermodynamically less stable adduct 69 in the addition of carbene 68 to 1,1-diphenylethylene that Newman and Patrick^{60,85} intially postulated a two-step carbene addition with zwitterionic intermediates. Subsequently, it was recognized⁸⁶

TABLE IV. Regioselectivity of $R(CH_3)C=C$: Olefin Additions

that the stereochemical assignment of 69 was complicated by the known⁸⁷ thermal rearrangement of such methylenecyclopropanes to indenes 70. The Newman-Patrick pathway would

also require 1,1-disubstituted olefins to be more reactive than 1,2-disubstituted alkenes, contrary to recent experimental observations.⁵⁹ Such a sequential reaction would also suggest the formation of a bicyclic product 72 from the addition of such carbenes to cis, trans-1,5-cyclodecadiene (71), but only normal 1,2-addition products 73 and 74 were observed.⁸⁸

As Newman and Patrick^{60,85} recognized, the initial approach of the electrophilic carbene to the olefin π system occurs via the empty p orbital of the carbene, which is constrained to be in the same plane as the β -alkyl substituents, thereby causing steric congestion with the substituents on the olefin. However, this itself does not require sequential bond formation and zwitterionic intermediates.

The reactivities in Table III are the result of opposing effects of the electronic and steric factors in the olefin in its interaction

with the carbene. Electron-rich unhindered alkenes like ethyl vinyl ether react faster than alkyl-substituted olefins. More electron-rich disubstituted olefins react faster than nonsubstituted olefins. The extra electron density of additional substituents is outweighed by steric factors and tri- and tetrasubstituted olefins react slower than 1,2-disubstituted ones. Deactivated alkenes such as tetrachloroethylene do not react at all.

These steric effects are reaffirmed by the regioselectivity data⁸⁸ summarized in Table IV. There is a definite regioselectivity for addition of an unsymmetrical carbene to an unsymmetrical olefin with a preference for the thermodynamically more stable antiadducts. This preference is a function of both the structure of the carbenes as well as that of the olefin. Furthermore with large steric interactions such as in the addition of $(t-Bu)$ MeC= $-C$: to fert-butylethylene, a significant amount of intramolecular alkyl migration and $(CH₃)₃CC=CCH₃$ formation is observed as well.⁸⁸

It is obvious from the foregoing that alkylidene carbenes are singlets in their ground state, they are encumbered to various degrees depending upon their mode of generation, are electrophilic, and have definite steric requirements in their interactions with substrates. In the next section their reactions will be discussed with by and large little regard for their origin or nature.

IV. Reactions of Alkylidene Carbenes

A. Intramolecular Reactions

The fact that the β -substituents in alkylidene carbenes are constrained to lie in the same plane as the empty p orbital of the carbenic carbon (see 75) coupled with the shorter bond length

of the C_{sp}2-C_{sp} bond and the electron deficiency of divalent carbon makes intramolecular rearrangement and acetylene formation a particularly facile process for such species. As noted in section II.A the Fritsch-Buttenberg-Wiechell rearrangement⁸⁹ of β , β -diarylvinyl halides upon treatment with base has been known since the 1890s, with rearrangement occurring under a variety of conditions.⁹⁰ The elegant labeling studies of Bothner-By⁹¹ and Curtin and co-workers^{92a} established that the rearrangement is stereospecifically trans, thereby ruling out the involvement of the free carbene 60 and implicating an organometallic, 58, or perhaps carbenoid, 59, intermediate.^{92b}

Similar aryl migration and exclusive acetylene formation occurs in the base treatment of 1,1-dihalo-2,2-diarylethylenes, 93 5,5-diaryl-*N*-nitrosooxazolidones⁹⁴ (27, $R = C_6H_5$), as well as diarylvinyl triflates¹⁴ (14, R = C_6H_5).

Exclusive rearrangement occurs even with one β -aryl group such as in 1-halo-2-arylpropene⁹⁵ (76) and the corresponding triflates 77. Once again Köbrich and co-workers⁹⁶ have by means of stereochemical studies and low-temperature isolation established that the halides 76 react through organolithium or carbenoid intermediates. In contrast, the completely nonstereospecific rearrangement of the triflate 77 strongly suggests, but does not absolutely prove, the involvement of the free carbene 60 rather than a carbenoid 59 in this intramolecular reaction.⁹⁷

Acetylene formation in monosubstituted vinyl compounds can occur by one of two ways:

a classical bimolecular β -elimination or α -elimination, alkylidene carbene formation, and hydrogen migration. Calculations^{52,98} indicate that HC=CH is some 38-46 kcal/mol more stable than $H₂C=C$:, hence the rearrangements

B

$$
H_2C = C: \xrightarrow{a} CH \equiv CH \text{ and } R(H)C = C: \xrightarrow{a} RC \equiv CH
$$

should occur rapidly with little or no activation energy. Indeed, $H₂C$ = C: has only been trapped intermolecularly, at the expense of exclusive acetylene formation, by special techniques.^{2a,99} C_2 (3) was allowed to react with a mixture of C_3H_8 and C_3D_8 to yield allenes by 1,1-diabstraction of hydrogen by C_2 to produce alkylcarbene and vinylidene:

The vinylidene so formed could rearrange to acetylene or rotate through 180° (estimated rate \sim 10¹¹ s) and couple with the RCH to produce the observed allenes.⁹⁹ On the other hand, $F_2C=C$: has been calculated to have a minimum $E_a = 59$ kcal/mol for rearrangement to FC=CF although the latter is some 36 kcal/ mol more stable than the carbene.⁹⁸

The complete absence of a deuterium isotope effect as well as a lack of stereochemical preference in the reaction of **(E)-78** and (Z) -78 vinyl triflates with t-BuOK established an α -elimination and alkylidene carbene and hydrogen migration as the most likely reaction pathway.¹⁰⁰

An alkylidene carbene has also been suggested as an intermediate¹⁰¹ in the formation of phenylacetylene from β -styryl bromide and C_6H_5L i, a reaction first observed by Wittig and coworkers.¹⁰² However, more detailed stereochemical and deuterium isotope effect studies of Schlosser and Ladenberger¹⁰³ on the reaction of alkyllithiums with styryl chloride rule out the involvement of a free alkylidene carbene and suggest an E-2 type mechanism:

An alkylidene carbene **79** has also been suggested as an intermediate in the thermal scrambling of labeled phenylacetylene at 550–700 °C:¹⁰⁴

$$
C_6H_5C^*\overline{=}CH \rightleftarrows [C_6H_5C^*H\overline{=}C:] \rightleftarrows HC^*\overline{=}CC_6H_5
$$

Intramolecular alkyl migrations in alkylidene carbenes or carbenoids can also occur, but only under more vigorous conditions and in the absence of olefin or other traps. Intramolecular alkyl migration and acetylene formation was postulated in the reaction of bromocamphene (80) with t-BuOK in refluxing tolu-

ene.¹⁰⁵ Similarly, alkylidene carbenes followed by alkyl migration has been invoked in the formation of cycloalkynes 82 from bromomethylenecycloalkanes **81** upon reaction with f-BuOK. The smaller ring C_5 and C_6 cycloalkyne intermediate could be trapped as Diels-Alder adducts with 1,3-diphenylisobenzofuran. With C_8 and larger rings the acetylenes were isolated.¹⁰⁶ No alkylidenes are involved in the reaction of **83** with bases or the formation of 85 and 86 from 84 and t-BuOK.¹⁰⁷

Alkynes were also observed (besides insertion products) presumably via alkylidene carbenes in the reaction of acyclic vinyl halides 87, with *t*-BuOK at 240 °C:¹⁰⁸

(R)CH₃C=CHBr
$$
\xrightarrow{t-BuOK}
$$
 RC=CCH₃ + others
87 240 °C

Alkyl migration and acetylene formation compete with intermolecular trapping by olefin in the case of cyclopropyl-sub-

stituted alkylidene carbenes 88 as derived from the corresponding oxazolidone 27.¹⁰⁹ Dicyclopropylcarbenes (89) gave only acetylene products (90) and could not be intermolecularly trapped.^{109,110}

Rearrangement and alkyne formation also occurred on base treatment of 1-halo-1,3-dienes^{111a} and 1-halo-1-ene-3-alkynes:111b

It is obvious that alkylidene carbenes or carbenoids are involved in many if not all of the alkyne forming reactions mentioned in section Il even if the intermediates could not be intermolecularly trapped. It is also evident that intermolecular reactions with substrates can only be carried out with β -dialkyl substituted (and perhaps heteroatom substituted) alkylidene carbenes and not with aryl or hydrogen substituted ones.

B. Intermolecular Reactions

1. Insertions

Alkylidene carbenes readily insert into Si-H and 0-H bonds. Both vinyl triflate 14 and oxazolidone 27 derived carbenes gave high yields of vinylsilanes 91 by reaction with triethylsilane.^{14,112}

The carbene 68 from oxazolidone⁸⁵ or vinyl triflate¹¹³ gave

predominantly the thermodynamically less stable isomer 92, with only small amounts of 93 as product. A stepwise hydride removal, rather than a concerted insertion, has been postulated⁸⁵ to account for such contrathermodynamic behavior.

A large number of alkylidene carbenes from diverse progenitors have been observed to give vinyl ether products. Newman and Okorodudu^{24b} have investigated the interaction of a number of oxazolidone derived alkylidene carbenes with a variety of alcohols and found 60-90% yields of vinyl ether products:

$$
(R)_2C \longrightarrow C: + R'OH \longrightarrow (R)_2C \longrightarrow C
$$

\n
$$
R' = Me, Et, n-Pr, i-Pr, n-Bu, t-Bu, etc.
$$

The exact mechanism of formation of these vinyl ether products remains to be elucidated. It is not clear if reaction occurs by capture of the carbene by RO⁻, usually present, by ylide 94 formation followed by a hydrogen shift, direct insertion, or by yet a different pathway.

$$
R_2C = C: + R'O^- \longrightarrow R_2C = C \longrightarrow OR
$$

\n
$$
R_2C = C \longrightarrow OR
$$

\n
$$
R_2C = C: + R'O + \longrightarrow R_2C = \bar{C} \longrightarrow \bar{C}
$$

\n
$$
R_2C = \bar{C} \longrightarrow \bar{C}
$$

\n
$$
R_2C = C: + R'O + \longrightarrow R_2C = \bar{C} \longrightarrow \bar{C}
$$

\n
$$
R_2C = \
$$

Unlike "insertion" into Si-H and 0-H bonds by alkylidene carbenes and in contrast to the behavior of alkylcarbenes,⁴ intermolecular insertion into C-H bonds occurs much less readily. Only in special circumstances have C-H insertions by alkylidene carbenes been observed. Intermolecular C-H insertions have been observed only with photogenerated^{28,30} carbenes with but modest yields. No other alkylidene carbenes or carbenoids have

been reported to undergo intermolecular C-H insertions. It is possible that normal C-H bonds do not possess sufficient electron density to be attractive substrates for weakly electrophilic unsaturated carbenes.

A number of intramolecular C-H alkylidene carbene insertions have been reported. Walsh and Bottini¹¹⁴ reported the formation of dihydrofurans 96 and 3-pyrrolines 98 via intramolecular alkylidene carbene insertion, from halopropenyl ethers 95 and amines, 97, respectively.

Insertion to give cyclopentenes 101 and 102 was observed upon treatment of chlorodienes 99 and 100 with n-BuLi in THF, respectively.¹¹⁵

In a recent study, ¹⁰⁸ 1,5-carbon-hydrogen as well as 1,3carbon-hydrogen insertion was observed upon treatment of acyclic terminal vinyl bromides with *t*-BuOK, neat at 240 °C (see Scheme II). The reason for the preference for 1,5 and to some extent 1,3 insertion over 1,4 or 1,6 insertion is not clear. The

relative amount of 1,5-C-H insertion followed the order tertiary $>$ secondary, benzylic \gg primary.¹⁰⁸

1,5-C-H alkylidene carbene insertion was postulated to account for the formation of indene **(104)** from **103** upon flash pyrolysis at 700 °C.⁴²

No data are available on the mechanism of any of these alkylidene carbene C-H "insertions". It is not known if they occur by a concerted or a stepwise process nor are the stereochemical requirements of the reaction known.

2. Addition to Double Bonds

The preferred intermolecular reaction of alkylidene carbenes is methylenecyclopropane formation via addition to olefins. Such addition is commonly used to establish the intermediacy of alkylidene carbenes or carbenoids in a particular reaction; hence they were largely discussed in sections Il and III. This part will treat unusual additions to C=C bonds not previously mentioned, and interactions of alkylidene carbenes with heteroatom double bonds such as $C=O$ and $N=N$.

Alkylidene carbene **106,** derived from **105,** has been added to several cyclic dienes.¹¹⁶ Addition to cyclopentadiene gave adduct **108** via the intermediacy of **107** and a trimethylenemethane-type rearrangement.¹¹⁷ With cyclopentadiene only 1,2-adduct **107** was observed with no 1,4-alkylidene carbene addition products being observed. Since 1,4-alkylcarbene ad-

120 121 122 n-BuLi

CHCI $THF, -50°C$ **123**

124 (20-30%)

bornadiene to give **112** which upon flash thermolysis gave **113** at 400 ⁰C and a mixture of **113** and **114** at 600 ⁰C. Rearrangements giving **110** and **111** have been observed.¹²¹ $+$ (CH₃)₂C=C: $CCH₃$ ₂ 112 $CH(CH_3)_2$

1,4-cyclohexadiene gave the expected adduct **109.** Interaction of **106** with norbornadiene also resulted in a 1,2-addition product **110,** which upon heating at 150 ⁰C converted to **111** by a trimethylenemethane diradical path. The alternative cycloaddition

Similarly isopropylidenecarbene has been added¹²⁰ to nor-

process is symmetry forbidden.¹¹⁹

600 'C $CCH₃$ ₂ **113 114**

A trimethylenemethane-type diradical, **115,** was involved in the addition of isopropylidenecarbene to cyclobutene. The actual products observed consisted of dimers of **115,** namely **116- 119.**¹²² Trimethylenemethane **115** was an intermediate in the thermolysis or photolysis of appropriate azo precursors and was extensively investigated by Berson and collaborators.¹²³ An analogous diradical, **122,** was involved in an intramolecular addition of an unsaturated carbene, **120,** and dimer formation.¹²⁴ Adducts **124** and **126,** analogous to **121,** were isolated as stable compounds from the reactions of **123** and **125** via intramolecular carbene additions.^{125,126}

Alkylidene carbenes have also been added to carbonyls.¹²⁷ Carbene **127** from the deamination of 20 in the presence of ketones and aldehydes gave diadducts **129,** presumably via the intermediacy of allene expoxides **128,** although no such allene oxides¹²⁸ could actually be isolated.

Alkylidene carbenes have been reacted with azoarenes.¹²⁹ Good yields of 2-indazoles **131** were obtained, representing a simple, single step entry into this $10-\pi$ -electron heterocyclic system. A number of mechanisms are possible for the formation of **131,** but interaction of the carbene with the nitrogen lone pair to give ylide **130** and subsequent ring closure followed by hydrogen migration seem most reasonable. Analogous addition

of alkylcarbenes to azoarenes resulting in dihydroindazoles are known.¹³⁰

SCHEME III

3. Addition to Allenes, Alkynes, and Cummulenes

As first observed by Shechter, ¹³¹ alkylidene carbenes readily add to allenes to give 1:1 adducts **132.** The identical product was observed via the oxazolidone **27** route.⁶⁰

$$
(CH3)2 C = CHBr + (CH3)2 C = C = C(CH3)2 CH3 CH3
$$

Interaction of **27** derived carbene with ethoxyacetylene and ethoxypropyne, respectively, gave adducts **134** via the mechanism postulated in Scheme III.¹³² No methylenecyclopropene **133** could be isolated. Addition of vinyl triflate **14** derived carbene to 2-butyne or 3-hexyne has provided a ready route¹³⁴ into the in situ preparation of simple¹³⁵ alkyltrifulvenes **135,** as shown by trapping with cyclopentadiene or HCIO₄. Although triafulvenes **135,** could not be isolated, evidence¹³⁶ indicates that the difficulty is reaction with oxygen rather than just thermal sensitivity.

 $(CH₃)₂C = CHOTf + RC = CR$

Alkylidene carbenes gave both expected 1:1 adducts **136** and **137** with butatriene as well as some diaddition products.¹³⁷ Compound 137 is an interesting member of a class of strained¹³⁸ organic molecules known as radialenes. The parent radialene, **137, is known as well.**¹³⁹

4. Miscellaneous Reactions

Reaction of **105** under phase-transfer conditions in the presence of NaN3 and NaSCN yielded **138** and **139,** respectively, and reaction with $(EtO)_{3}P$ gave 140.¹⁴⁰ The involvement of alkylidene carbenes in these reactions has not been established except in the reaction with N_3 . Kirmse and Schnurr¹⁴¹ have

shown that the reaction of **105** derived diazonium ions with NaN₃ involves vinyl cations and alkylidene carbenes as well as pentazenes **141.** Similar intermediates may be involved in the other reactions as well.

$$
R_2C=CH\bar{N}-N=N-\bar{N}\equiv N
$$

141

Reaction of isonitriles with **17** derived alkylidene carbene gave amide 143 as product.^{142a} Such an amide could arise via the reaction of the initially formed alkylideneketenimine^{142b} adduct, **142,** with water.

Several potentially interesting reactions of alkylidene carbenes such as interaction with phosphines to form ylides **144,** then allenes **145,** have not been reported. Such a reaction would

$$
(R)_{2}C = C: + P(C_{6}H_{5})_{3} \rightarrow (R)_{2}C = C = P(C_{6}H_{5})_{3}
$$

144

$$
\xrightarrow{(R')_{2}CO} R_{2}C = C = C(R')_{2}
$$

145

represent an alternative entry into the novel cumulated phosphorus ylides investigated by Bestmann and co-workers.142b Neither have interactions of alkylidene carbenes with aromatic substrates nor the formation of heptafulvenes **146** via this route been reported, although the analogous addition of alkyl carbenes to aromatics and formation of norcardienes are well known.⁴

V. Generation of Vinylidene Carbenes

Vinylidene carbenes have mostly been generated by interaction of base with propargyl derivatives **147** or haloallenes **148.** It was to account for the slow reactivity in aqueous ethanol

(compared to tert-amyl chloride) and kinetic dependence on added NaOH of 1-chloro-3-methyl-1,2-butadiene and 3-chloro-3-methyl-1-butyne that such a carbene was first proposed by Hennion and Maloney.¹⁴³ A simple S_N1 reaction was ruled out by the effect of added NaOH on the reactivity of the terminal alkyne **147** and the lack of any effect on the methyl-substituted (2-pentyne analogue) halide.¹⁴⁴ Initially, Hennion and Maloney proposed a slow loss of H^+ and rapid subsequent loss of halide. .
Subsequent work of Shiner and co-workers¹⁴⁵ showing rapid deuterium exchange in deuterated solvent as well as common ion rate depression by added halide established that reaction occurs via a rapid preequilibrium and anion formation followed by a rate-determining loss of halide and carbene **149** formation (Scheme IV).

Carbene **149** can also be generated in aprotic hydrocarbon media from either 147¹⁴⁶ or 148^{147,148} and base, usually *t*-BuOK. Generally better yields of carbene-derived products are observed if the more recent phase transfer¹⁴⁹⁻¹⁵¹ or crown ether¹⁵² procedures are employed with either **147** or **148** for the generation of **149.**

Dihaloallenes 150^{153,154} and bromoalkyne¹⁵⁵ 151 have also been employed as carbene **149** progenitors. Cyclopropane¹⁵⁶

$$
\begin{array}{ccc}\n\text{(C}_6\text{H}_5)_2\text{C} = & \text{CBr}_2 & \text{r-C}_4\text{H}_9\text{CH}_2\text{C} = & \text{CBr} \\
\text{150} & \text{151}\n\end{array}
$$

152 as well as bis(nitrosourethane)¹⁵⁷**153** have been shown to give carbene-derived products (Scheme V). This latter reaction is the only one that might involve a diazoallene, **155,** in the generation of a vinylidene carbene. Unfortunately, the yields of carbene derived products are very low because of competing diphenyldiazopropanone **(157)** formation presumably via intermediates **154** and **156.** Vinylidene carbenes have also been implicated in the isomerization¹⁵⁸ of thioalkyne **158**, as well as in the reaction¹⁵⁹ of carbamate **159.**

Vinylidene carbenes are probably intermediates in the thermolysis of propynylaniline¹⁶⁰ **(160)** and the alkaline methanolic cleavage of propargylnitrosourea **(161)** and photolysis of tosylhydrazone¹⁶¹ **162.**

Other than the reactions of **162,** to date no procedures have been reported for the generation of vinylidene carbenes that do not involve some type of base initiated elimination. Other than

SCHEME V

$$
\begin{array}{cccc}\n & \text{CH}_3\text{NHCOC} & \text{C} \equiv \text{CCH} = \text{C} = \text{C(CH}_3)_2 \\
 & \times \text{CH}_3\text{NHCOC} & \text{C} \equiv \text{CCH} = \text{C} = \text{C(CH}_3)_2\n\end{array}
$$

$$
C_6H_5NHCH_2C \equiv CH \xrightarrow{100 \text{ GeV}} 149
$$

160
 $\rightarrow C_6H_5NH_2 + C_6H_5N(CH_2C \equiv CH)_2$

the photolysis of **162** there are also no photochemical processes

known for the generation of **149.**

 $HC = CCH₂NCNH₂$ NO 161 CH₃OH CH3ONa $149 \xrightarrow{CH_3OH} CH_2 = C = CHOCH_3$ $[HC = CCHN₂]$ h_v Na HC = $=$ CCH= $NNSO_2C_6H_4CH_3$ 162

 Ω

Species which are isomeric with vinylidene carbenes, namely alkynyl carbenes or propargylenes, **163,** should be briefly mentioned. The parent as well as substituted propargylenes

$$
\text{RCC} \equiv \text{CR} \stackrel{f'''}{\longleftarrow} \text{RC} \equiv \text{CCRN}_2
$$
\n
$$
\text{163} \qquad \text{164}
$$

have been prepared¹⁶² by photolysis of the appropriate diazoalkynes 164, and shown¹⁶³ to be triplets in accord with expectations.¹⁶⁴ Substituted propargylenes have also been produced by photolysis¹⁶⁵ of 165 and via the reaction¹⁶⁶ of 166 with f-BuOK.

Finally C₂O (168) has also been generated¹⁶⁷ by photolysis

$$
\rightarrow C_6H_5CH_2C \equiv CH + C_6H_5CH = C = CH_2
$$

 $C_6H_5C \equiv CCH_3$

$$
(C_6H_5C \equiv C\frac{1}{2}CHBr \xrightarrow{t-BuOK} [(C_6H_5C \equiv C\frac{1}{2}C]
$$

166

$$
\longrightarrow (C_6H_5C \equiv C\frac{1}{2}C \Longrightarrow C = C + C \quad CC_6H_5)_2 \rightarrow \text{others}
$$

of carbon suboxide **(167)** and its chemistry investigated.¹⁶⁸

$$
O=C=C=C=O \xrightarrow{hv} O=C=C:
$$

167 168

Vl. Nature of Vinylidene Carbenes

Unlike alkylidene carbenes (1), vinylidene carbenes (2) are hybrids of a zwitterionic and a neutral resonance form, **149.** The relative importance of the two resonance forms is strongly affected by the structure as well as the environment of the particular species. Electron-donating substituents such as alkyl and aryl groups should enhance the contribution of the zwitterionic form as should polar solvents such as aqueous ethanol and ethers capable of strong solvation. In contrast nonpolar solvents like hydrocarbons should favor the neutral carbenic resonance form. There is also good evidence from measurement of activation volumes that in aqueous protic solvents the intermediate from reaction of either **147** of **148** ($R = CH_3$) is a halide asso- $\frac{1}{2}$ circle is that the free carbene.¹⁶⁹ In contrast, relative reactivity data summarized in Table V strongly suggest that there is very little carbene encumbrance in the transition states of olefin additions. The similarity in olefin selectivity of vinylidene carbenes **149** regardless of progenitor not only suggests a free carbene but also shows very little steric requirement for reaction. The dimethyl-, di-tert-butyl-, and diphenylcarbenes react similarly with each olefin. This is as expected since in vinylidene carbenes, unlike alkylidene carbenes 75, the substituents are perpendicular and one carbon removed from the reactive site at the empty carbenic orbital in **169.** Faster reaction with more substituted, and hence electron-rich olefins, further indicates that vinylidene carbenes like alkylidene carbenes are electrophilic. This conclusion is supported by the ρ value of -0.95 observed in the interaction of $(CH_3)_2C=CC$: with various ring-substituted styrenes.⁶⁷

havior characteristic of such species.^{172,173} This suggests that

perhaps a rapid equilibrium exists between carbenoid and free carbene, even in nonsolvating solvents like olefins, with the free carbene reacting faster than the carbenoid. Such an equilibrium would account for all of the behavior of vinylidene carbenes **149,** both in protic polar solvents and in nonpolar olefin solvents (i.e., the similar selectivity of Table V).

By arguments similar to those applied to alkylidene carbenes in section III.A, vinylidene carbenes would be expected to be singlets. The addition of $(CH_3)_2C=CC=CC$: to cis - and trans-2butene was found to be stereospecific.¹⁷⁰ This implies that at least the nascent carbene is a singlet. Recent theoretical calculations¹⁷⁴ indicate that vinylidene carbenes 2 possess a singlet ground state, with a singlet triplet splitting of about 50 kcal/ mol.

Vinylidene carbenes 2, like their alkylidene counterparts 1, are singlets and electrophilic. They exhibit behavior characteristic of both carbenoids and free carbenes. Unlike alkylidene carbenes there is very little steric requirement for addition to olefins, but there are definite steric requirements for reaction with nucleophiles (vide infra).

VII. Reactions of Vinylidene Carbenes

Unlike alkylidene carbenes the substituents in vinylidene carbenes are not predisposed to migration by virtue of their location as shown in **169** and hence no intramolecular rearrangements are possible or known.

A. Insertions

Vinylidene carbenes readily insert into Si-H bonds to give silylallenes.¹⁷⁵ No data are available on the mechanism of such

$$
(R)_2C=C=C: + Et_3SH \longrightarrow (R)_2C=C=C\begin{matrix} R \\ S_1Et_3 \end{matrix}
$$

 \overline{H}

insertions. Insertion into C-H bonds α to oxygen has been observed with alcohols and 2,5-dimethyltetrahydrofuran.^{175,176}

Prior hydrogen bonding between the 0-H and the carbene lone pair orienting the empty p orbital in close proximity to the α -C-H bond has been proposed¹⁷⁶ to explain the specificity of C-H insertion. An alternative possibility of an anion carbene pair formed between alkoxide ion and the C-3 orbital was considered less likely.¹⁷⁶ No insertion was observed into cumene, THF, or glyme.

Nucleophilic character was attributed to vinylidene carbenes in their insertion into the benzylic C-H bond in a series of ringsubstituted benzyl alcohols, to explain the positive ρ value of 0.52 that was observed.¹⁷⁷ The reaction was considered to proceed by a concerted process due to the small value of ρ . since a full benzyl anion 178 has a Hammett ρ value of 4.0. Hydrogen bonding followed by ylide formation and collapse to product was proposed as a reaction pathway.¹⁷⁷

$$
XC_{6}H_{4}CH_{2}OH + (CH_{3})_{2}C = C = C:
$$
\n
$$
\longrightarrow XC_{6}H_{4}CH_{2}OH \longrightarrow XC_{6}H_{4}CHOH
$$
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\n $$

B. Addition Reactions

1. With Nucleophiles

A large number of nucleophiles react with vinylidene carbenes **149,** generally derived from the corresponding propargyl halides. Barring steric complications in protic polar media, reaction generally occurs at the more electrophilic tertiary carbon resulting in propargylic products. Steric factors, solvent effects, and nucleophilicity affect these reactions although in ways not clearly understood.¹⁷⁹

In contrast, in aprotic (hydrocarbon) media, reactions seem to be more straightforward with involvement of the carbene form rather than the zwitterionic form **149.** Hexapentaenes **172** can be formed by interaction of the initially formed acetylide with the

carbene.^{146,154,180-182} Similarly, butatrienes were obtained upon reaction of **149** with diazo compounds.¹⁸³

Alcohols readily react with **149** to give, in the absence of steric hindrance, propargyl ethers.^{143,184} Work of Hennion and

$$
(R)_2C \longrightarrow C \longrightarrow CH + R'OH \xrightarrow{base} (R)_2C \longrightarrow C \longrightarrow CH
$$

\n
$$
\downarrow
$$

\n
$$
OR'
$$

Motier¹⁷⁹ indicates that reaction probably proceeds via the alkoxide rather than the alcohol itself. In t -BuOH:H₂O only propargyl alcohol and ene-yne were observed, whereas in aprotic media the *tert-*butoxy propargyl ether was observed.¹⁸⁵ In contrast, with steric hindrance in **149** ($R = t$ -Bu) mixtures of allenyl and propargyl ethers were observed.¹⁸⁶

A large variety of amines readily react with propargyl halides via **149** to give propargyl amines.187-191 The reaction appears

$$
(R)_2CC \equiv CH + (R')_2NH \longrightarrow 149 \longrightarrow (R)_2CC \equiv CH
$$

\n
$$
\downarrow
$$

\n
$$
N(R')_2
$$

to be catalyzed by trace amounts of cuprous salts¹⁸⁹ and less affected by steric factors than reaction with alcohols. Tertiary amines were found to give mixtures of allenic and alkynyl amine

$$
(R)_{2}CC \equiv CH + (R')_{3}N \longrightarrow 149
$$
\n
$$
\times
$$
\n
$$
CI^{-}
$$
\n
$$
\longrightarrow (R)_{2}C \equiv C \equiv CHN(R')_{3} + (R)_{2}C \quad \text{C} \equiv CH
$$
\n
$$
Cl^{-}
$$
\n
$$
Cl^{-}
$$
\n
$$
Cl^{-}
$$
\n
$$
N(R')_{3}
$$

salts.¹⁹² Alkynyl amines were also obtained from allenyl bromides **173** and **174** and amines.¹⁹³

$$
ROCH2CH = C = CHBr + R1R2NH \longrightarrow 149
$$

\n173
\n
$$
\longrightarrow ROCH2CHC = CH
$$

\n
$$
NR1R2
$$

\n(R)₂NCH₂CH = C = CHBr + R₁R₂NH \longrightarrow 149
\n174
\n
$$
\longrightarrow (R)2NCH2CHC = CH
$$

 NR_1R_2

Enamines and propargyl chlorides gave after hydrolysis acetylenic ketones 175.¹⁹⁴ Reaction presumably occurred via C-alkylation with 149 although the nature of the initial product is unknown.¹⁹⁴

Sulfur nucleophiles such as thiophenoxide gave mixtures of allenyl and acetylenic thioethers with an identical product ratio from either haloallene or acetylene.^{145b} Vinyl sulfide 176 gave

sulfide 178 upon reaction with 149 via a [2,3]-sigmatropic shift of ylide 177.¹⁹⁵ Such [2,3]-sigmatropic shifts of sulfur ylides are well known.¹⁹⁶ Subsequent hydrolysis of 178 in the presence of HgCI₂ gave artemesia ketone 179 in good yield.¹⁹⁵

Various substituted diethyl sodiomalonates have also been employed as nucleophilic traps for 149. Sodio unsubstituted diethyl malonate gave¹⁹⁷ 180 whereas substituted diethyl sodiomalonates such as diethyl sodiomethylmalonate¹⁹⁸ and diethyl

sodioformylaminomalonate¹⁹⁹ gave both acetylenic and allenic products 181 and 182. Similar products were obtained with sodioacetoacetate as nucleophile.²⁰⁰ Grignard reagents have also been used as nucleophiles.²⁰¹

$$
149 + NaCH(CO_2Et)_2 \longrightarrow (R)_2CCH(CO_2Et)_2
$$
\n
$$
\begin{array}{ccc}\n & & \\
 \downarrow & & \\
 \downarrow
$$

2. With π Bonds

Similar to alkylidene carbenes, the major reaction of vinylidene carbenes with π bonds is addition to olefins, to give alkenylidene cyclopropanes 183. Alkenylidene cyclopropanes in

turn have been extensively employed as substrates in cycloaddition reactions²⁰² and in thermolysis.²⁰³ Addition to tetramethylallene gave 184 in 35% yield.¹³¹ Addition to skatole

heating readily underwent rearrangement to 188 whereas the methylindene addition product resulted in a different rearrangement product, 189.²⁰⁵ Addition to dihydronaphthalenes

resulted in similar products.²⁰⁵ Vinylidene carbenes have been employed in an elegant synthesis of chrysanthemic acid.²⁰⁶ Addition of dimethylvinylidenecarbene to allyl alcohol 190 resulted in a 45% yield of adduct 191, which upon reduction with Na in liquid ammonia gave a 90% yield of a 3:1 trans:cis chrysanthemyl alcohol 192, which was readily oxidized to frans-chrysanthemic acid 193.

There is only one report on the addition of vinylidene carbenes to heteronuclear bonds. Azobenzene gives benzimidazole 195 via adduct 194.207

VIII. Transition Metal Complexes of Unsaturated Carbenes

An enormous amount of work has been done on metal-carbene complexes²⁰⁸ of the type 196. The simplest and most common entry into these complexes is the procedure²⁰⁸ de-

veloped by Fischer and co-workers involving metal carbonyls and alkyllithium:

$$
M(CO)_x + RLi \begin{bmatrix} O \\ \vdots \\ (OC)_{x-1}MCR \end{bmatrix} Li^+ \xrightarrow{Me_3OBF_4} (OC)_{x-1}MCR
$$

Unfortunately, to date, no such general procedure exists for the preparation of analogous alkylidene 197 or vinylidene 198 complexes, and little is known about these unsaturated carbene complexes. UV irradiation of diphenylketene and Fe(CO)₅ in benzene results in the bridged complex 199.²⁰⁹ Unambiguous

(C6Hg) ²C=C=O + Fe(CO)⁵ C«H^R (OC)4Fe.. (OC)4Fe .C=C(C6H⁵) ² 199

evidence for 199 was obtained by x-ray analysis that indicated a Fe-C (carbene) distance of 1.98 A, Fe-Fe of 2.64 A, and a C=C of 1.33 Å with a nearly planar arrangement of the two iron and two double-bonded carbon atoms.²⁰⁹ Complex 199 could also be obtained by treatment of diphenylketene with $Fe₂(CO)₉$ in the absence of any irradiation, strongly suggesting that it was not formed by direct trapping of the preformed diphenylcarbene.

The manganese complexes 200 and 201 of $C_6H_5CH=C$: were obtained in 7 and 2% yields, respectively, by treatment of phenylacetylene with $\mathsf{C}_5\mathsf{H}_5\mathsf{Mn}(\mathsf{CO})_2(\mathsf{THF})$.²¹⁰ X-ray analysis was employed to unambiguously establish the structures of 200 and 201. In 200 the Mn-C (carbene) distance was found to be 1.68 Å, which is considerably shorter than the Mn–C alkyl σ bond of 2.16 A and the Mn-C bond of 1.88 A in a saturated carbene-Mn complex, indicating double bond character between the carbenic

$$
C_{5}H_{5}Mn(CO)_{2}(THF) + C_{6}H_{5}C = CH \xrightarrow{fN}
$$
\n
$$
C_{5}H_{5}Mn(CO)_{2}(C = CHC_{6}H_{5}) + \sum_{(OC)_{2}C_{5}H_{5}Mn}C = CHC_{6}H_{5}
$$
\n
$$
200 \qquad (OC)_{2}C_{5}H_{5}Mn
$$
\n
$$
201
$$

carbon and manganese. 210 The C= C distance was found to be 1.34 A. A nearly linear bond angle of 174° was observed around the Mn=C—C moiety. In complex **201,** the Mn-Mn bond was found to be 2.73 A and the Mn-C (carbene) to be 1.94 A, respectively, with an Mn-C-Mn angle of 88.0°. At the moment nothing is known about the chemistry of complexes **199- 201.**

The only systematic investigation of alkylidene carbene-metal complexes to date has been carried out by King and co-workers²¹¹ on complexes of dicyanovinylidene **(202).** Treatment of 1,1-dichloro-2,2-dicyanoethylene with the highly nucleophilic anion $[C_5H_5Fe(CO)_2]$ ⁻ resulted in the bridging cis and trans

$(NC)_2$ C $=$ C: **202**

complexes **203.**²¹² Once again x-ray crystallography unambiguously established²¹³ the structure of **203.** The Fe–Fe bond was found to be 2.5 Å and the Fe-C (carbene) 1.87 Å which is

considerably shorter than the Fe-C 1.94 A for the bridging $C = 0.213$

Subsequently, **202** was also observed as a terminal ligand in complexes of the type **205** obtained by reaction of **204** with various phosphine nucleophiles.²¹⁴ X-ray data on **206** estab-

$$
(NC)2C = C
$$

\n
$$
M_0C_5H_5(CO)3
$$

\n
$$
204
$$

\n
$$
(NC)2C = C;]M_0C_5H_5[P(R)3]2(Cl)
$$

205

lished²¹⁵ the structure to be a distorted square pyramid around the metal with a Mo-C (carbene) bond of 1.83 A which is much shorter than the Mo-C of 1.93-2.06 Å for $C=O$. The shorter M-C (carbene) bond distance in these complexes compared to the analogous M-C(CO) distances along with other spectral data

indicate that dicyanovinylidene 202 is a stronger π acceptor than $C = 0$ in metal complexes.²¹⁶ A number of related complexes of **202** of the general structure **207** may be prepared by analogous reactions.²¹¹

Finally, although not isolated, the parent carbene $(CH_2=CC)$ complex has been implicated as an intermediate by Fischer and co-workers:²¹⁷

No reports have yet appeared that would indicate that any of the above metal complexes of alkylidene carbenes would serve as progenitor of the carbene itself, or that they might serve as transfer agents of the carbene to olefins or other substrates.

Most recently two reports have appeared on the preparation of vinylidene transition metal complexes. Reaction of carbene complex **208** with a Lewis acid followed by a weak base gave complex 209.²¹⁸ X-ray analysis established that the heterocu-

mulene moiety in **209** is nearly linear with a M-C (carbene) bond order of between 1 and 2. This, along with the short $C_1 - C_2$ and C_3 -N bond lengths and the observation of two distinct CH₃ groups in the proton NMR indicate a considerable contribution

from the mesomeric ionic form in **209.** A similar complex, **211,** was obtained²¹⁹ by treatment of complex 210 with t-BuLi. The structure of 211 was assigned by spectral means.²¹⁹ To date no chemistry has been reported with these novel complexes.

IX. Conclusion and Prognosis

There is no doubt that both alkylidene and vinylidene carbenes are viable reactive intermediates. Alkylidene carbenes may be generated by a variety of techniques from diverse progenitors, whereas vinylidene carbenes have been generated by baseinitiated eliminations. Both types of unsaturated carbenes are singlets and electrophilic, and their degree of encumbrance varies as a function of media and progenitor. Both types of carbenes readily add to alkenes to give alkylidene and alkenylidene cyclopropanes, respectively. Interaction with a variety of other substrates occurs as well. Transition metal complexes of both alkylidene and vinylidene carbenes have been reported.

Yet, a lot remains to be done. Clearly, better and more general methods of photochemical generation of unsaturated carbenes would be highly desirable. Such a mode of generation might allow matrix isolation and better physical characterization of these novel intermediates.

Little is known about heteroatom (O, N, S, Si, halogens, etc.) substituted unsaturated carbenes. These would be of interest not only from a mechanistic point of view and their possible contrasting behavior to their alkyl substituted counterparts, but also from a synthetic point of view. Since they would possess added functionality imparted by the heteroatom, they have the potential for further synthetic manipulation.

There will undoubtedly be more work on the synthetic application and uses of unsaturated carbenes, particularly in the area of small ring and strained compounds.

More work needs to be done in deciphering the nature and mechanism of unsaturated carbene "insertion" reactions, along with elucidating the exact profile of olefin-carbene interactions.

The reported synthesis of the transition metal complexes is but a start. General methods of preparation of unsaturated carbene transition metal complexes need to be developed along with exploring the chemistry of such complexes.

Extended unsaturated carbenes such as **212, 213,** and **214** might be observed. Such species would be of interest not only

from a mechanistic point of view but also from a synthetic point of view, for they might provide ready entry into some interesting cumulene chemistry. The recent observation²²⁰ of vinylvinylidene **(215)** might serve as a model for the generation of such extended carbenes via 216, 217,²²¹ and 218, respectively.

Acknowledgments. Financial support by Public Health Service Research Grants l-RO-l-CA 16903-01 and -02 from the National Cancer Institute and the Petroleum Research Fund administered by the American Chemical Society is appreciated. Helpful discussions and exchange of information with Professors W. A. Goddard III, T. B. Patrick, and J. P. Simons were valuable. I also wish to thank the Alexander-von-Humboldt Foundation for a "Senior U.S. Scientist Award", and Professor M. Hanack and the lnstitut fur Organische Chemie der Universitat Tubingen for their kind hospitality during 1977/78 and the preparation of this manuscript.

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