# **Molecular Photochemistry of Alkanones in Solution:** a-Cleavage, Hydrogen Abstraction, Cycloaddition, and **Sensitization Reactions**<sup>1</sup>

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Excitation by absorption of ultraviolet or visible light provides a convenient, but by no means unique, method for generating electronically excited molecules in the laboratory.<sup>3</sup> Following absorption, two electrons become orbitally decoupled. Generally, the spins of these electrons remain antiparallel, i.e., absorption of ultraviolet or visible light by an organic molecule will generally produce an electronic isomer of the ground state. This electronic isomer is called an excited singlet state, which we shall hereafter designate as  $S_1$ , in contradistinction to  $S_0$ , the electronic ground state. A "spin isomer" of  $S_1$  results if the spins of the orbitally decoupled electrons are parallel. This triplet state we shall designate as  $T_1$ ; it is an electronic and spin isomer of  $S_0$ . In general, because of quantum mechanical constraints (the Pauli principle), the  $T_1$  state is more stable than  $S_1$ .

In formaldehyde,<sup>4</sup> the  $S_1$  and  $T_1$  states are well characterized and are saliently different from the ground state (Figure 1). The electronic isomers of formaldehyde are pyramidal species possessing a very long, essentially single, CO bond and a substantially reduced dipole moment.<sup>5</sup> Indeed, H<sub>2</sub>CO electronic isomers undergo rapid inversion (at about  $10^{10} \text{ sec}^{-1}$ ) through a mirror plane, analogous to the behavior of the ammonia molecule.

This beautifully detailed description of  $S_1$  and  $T_1$  in  $H_2CO$  is possible because of its amenability to spectroscopic analysis. Although  $S_1$  and  $T_1$  possess about 80 kcal of energy above that of the ground-state formaldehyde, these species last long enough to be characterized by their spectra. The  $S_1$  and  $T_1$  states are generally the only photochemically reactive states (Kasha's rule) which need be considered in solution.<sup>6</sup>

The diffuse absorption and emission spectra of alkanones, on the other hand, are not amenable to such detailed analysis. As a result, we must employ a strongly coupled combination of spectroscopic and chemical evidence in order to infer the nature and behavior of the electronic isomers of the alkanones at the molecular level.

An abundance of theoretical and experimental evidence leads to the working hypothesis that the configuration of S<sub>1</sub> and T<sub>1</sub> in alkanones is  $(n_0)^1(\pi^*)^1$ , where n is an orbital more or less localized on the carbonyl oxygen atom and  $\pi^*$  is an orbital delocalized over both the carbonyl atoms.<sup>7</sup> We shall designate both  $S_1$  and  $T_1$  as "n,  $\pi^*$  states," thereby emphasizing the approximate electronic configuration of these states. This model<sup>8</sup> suggests that both states will be amphoteric reagents, i.e., electrophilic and radical-like in the vicinity of the half-vacant n orbital on oxygen, and nucleophilic (and possibly radical-like) above and below the carbonyl faces because of the presence of a  $\pi^*$  electron (Figure 2).

In this Account, we test the usefulness and validity of the above model as a means of predicting and understanding how structural changes influence the reactivity, specificity, and efficiency of photoreactions of alkanones. We shall attempt to show how studies of quantum yields and photochemical kinetics have led to a considerable increase in our confidence in the understanding of the behavior of electronically excited alkanones. In addition, we shall show how biradical intermediates, concerted rearrangements, cycloadditions, and other topics of wide interest to chemists are involved in the photochemistry of alkanones.

#### Chemistry of the $S_1$ and $T_1$ States of Alkanones<sup>9a,b</sup>

The overwhelming majority of known photochemical reactions of alkanones falls in three typical classes:

(1) Molecular Photochemistry. L. Paper XLIX: A. Yekta and N. J. Turro, Mol. Photochem., 3, 307 (1972). (b) This work was generously supported by the Air Force Office of Scientific Research by Grant AFOSR-70-1848.

(2) (a) NIH Predoctoral Fellow; (b) NATO Postdoctoral Fellow,
(c) NSF Predoctoral Fellow.

(3) For example, electronically excited alkanones are produced by thermal decomposition of 1,2-dioxetanes: E. H.White, J. Wiecko, and D. F. Roswell, J. Amer. Chem. Soc., 91, 5194 (1969).

(4) J. C. D. Brand and D. G. Williamson, Advan. Phys. Org. Chem., 1, 365 (1963).

(5) D. E. Freeman and W. Klemperer, J. Chem. Phys., 45, 52 (1966).

(6) M. Kasha, Discuss. Faraday Soc., 9, 14 (1950).
(7) M. Kasha in "Light and Life," W. D. McElroy and B. Glass, Ed., Johns Hopkins University Press, Baltimore, Md., 1961, p 31.

(8) H. E. Zimmerman, Advan. Photochem., 1, 183 (1963). (9) For recent reviews see (a) A. A. Lamola and N. J. Turro, "Energy Transfer and Organic Photochemistry," Interscience, New York, N. Y., 1969, p 191 ff; (b) J. C. Dalton and N. J. Turro, Annu. Rev. Phys. Chem., 21, 499 (1970); (c) P. J. Wagner and R. W. Spoerke, J. Amer. Chem. Soc., 91, 4437 (1969).

Nicholas Turro has authored several books in the general area of photochemistry. He received his Ph.D. in 1963 from Caltech, and after a year's postdoctoral work, joined the staff at Columbia University. Professor Turro supervises a large and active research group between handball games. Several of the coauthors of this Account have received or are working on their Ph.D.'s with him: Douglas R. Morton, Jr. (now a postdoctoral associate with William Johnson at Stanford University), J. Christopher Dalton (now Assistant Professor at the University of Rochester), George L. Farrington, Neil E. Schore, and Mark Ρ. Niemczyk. Keith Dawes (Ph.D. from University of Manchester, England, and presently at Oxford University) and Richard R. Hautala (Ph.D. from Northwestern) were postdoctoral fellows in his group,

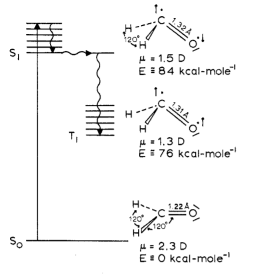


Figure 1. Properties of formaldehyde in its thermally equilibrated ground state, lowest excited singlet state, and lowest triplet state. It should be noted that, although the ground state is planar, the excited singlet possesses an angle of  $20^{\circ}$  intersecting the HHC plane and the CO axis. The triplet state is even more puckered, with a corresponding angle of  $35^{\circ}$ .

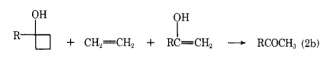
(a) cleavage of the C—C=O bond, known as  $\alpha$  cleavage or type I cleavage (eq 1); (b) abstraction of a hy-

$$\operatorname{RCOR} \xrightarrow{h\nu} \operatorname{R\dot{C}O} + \operatorname{R} \cdot \tag{1}$$

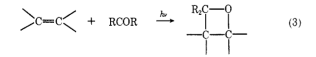
drogen atom by the carbonyl oxygen atom, which may occur intermolecularly (eq 2a) or intramolecularly (eq 2b); the latter case is known as type II reaction;

$$\begin{array}{ccc} & & & & \\ & & & & \\ RCOR + R'H & \xrightarrow{h\nu} & RCR + R' \cdot \end{array}$$
 (2a)

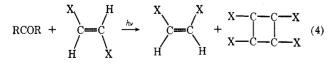
RCOCH<sub>2</sub>CH<sub>2</sub>CH<sub>3</sub> -



(c) cycloaddition to an unsaturated carbon-carbon linkage (eq 3). Reactions associated with (c) are



sensitized isomerization and dimerization of ethylenes (eq 4).



We shall consider each of the above processes, emphasizing recent contributions from work done at Columbia and other laboratories. First, we must briefly review some of the important characteristics of photochemical reactions which will be crucial for a valid

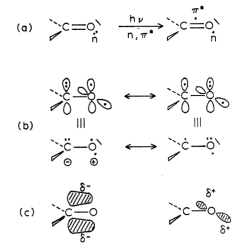


Figure 2. Description of an  $n, \pi^*$  excitation (a), the electronic distribution in an  $n, \pi^*$  S<sub>1</sub> or T<sub>1</sub> state (b), and the amphoteric nature of an  $n, \pi^*$  S<sub>1</sub> and T<sub>1</sub> state (c). Key concepts of the approximate model (which cannot differentiate between properties of S<sub>1</sub> and T<sub>1</sub> or predict their nuclear configurations) are (1) the nucleophilic character of the  $\pi$  faces, (2) the electrophilic character at the edges of the carbonyl oxygen atom, and (3) the radical-like character of both the  $\pi$  faces and the carbonyl edges.

and convincing extrapolation from the realm of laboratory observations into the realm of intellectual constructs of what occurs at the molecular level.

# Efficiency and Yields of Photoreactions: Probability That an Absorbed Photon Will Result in the Formation of a Molecule of Product<sup>10</sup>

Simple and convenient methods are available for measuring the number of photons absorbed by an organic molecule. The efficiency of photoreaction, *i.e.*, the number of molecules of photoproduct formed divided by the number of photons absorbed, is known as the quantum yield,  $\phi$ , of product formation. The "chemical yield" of a photoreaction depends upon the relative efficiencies of competing irreversible reactions which occur after photon absorption. The reactivity of S<sub>1</sub> and T<sub>1</sub> toward a given process is properly associated with the rate constant, k, for the process. The quantum yield is basically a measure of the rate of a process (from S<sub>1</sub> or T<sub>1</sub>) divided by the rate of light absorption ( $I_a$ ) leading to the reactive state, *i.e.* 

$$\phi = \frac{\text{molecules reacting/sec}}{\text{photons producing reactive state/sec}} = \frac{k[M^*]}{I_a} \quad (5)$$

Thus, in order to measure k we must have information in addition to  $\phi$ . In general,  $\phi$  is a poor guide to reactivity, even in closely related cases, because of the nature of competing photochemical processes which determine  $\phi$ .<sup>11</sup> As we shall see, however, measurement of  $\phi$  will be useful in establishing certain aspects of photochemical reactions.

Knowledge of both efficiencies and reactivities of  $S_1$  and/or  $T_1$  allows us to make important conclusions

<sup>(10)</sup> N. J. Turro, J. Chem. Educ., 44, 536 (1967).

<sup>(11)</sup> For example, see P. J. Wagner and H. N. Schott, J. Amer. Chem. Soc., 91, 5383 (1969).

about "invisible chemical or physical processes," namely radiationless energy loss mechanisms which lead to no net chemical change.

## Energetics and Dynamics of Alkanone S<sub>1</sub> and T<sub>1</sub> States

The chemically significant processes which interconnect  $S_0$ ,  $S_1$ , and  $T_1$  of acetone in an inert solvent provide a concise and informative model of the energetics and dynamics which will be crucial in understanding the photochemical reactions of alkanones. Excitation of acetone with light of wavelength 260-300 nm produces (after vibrational relaxation)  $S_1$ , the  $n, \pi^*$ singlet state, in quantitative yield. The important features of  $S_1$  for acetone are: (1) it possesses  $\sim 84$ - $\rm kcal/mole^{12}$  excess electronic energy over S\_0, acetone's ground state; (2) in the absence of external quenchers of  $S_1$ , only two important processes deactivate  $S_1$ : (a) intersystem crossing to  $T_1$  and (b) fluorescence to yield  $S_0$ ; (3) the rate constant for intersystem crossing  $(k_{ST})$ is  $5 \times 10^8 \, {\rm sec^{-1}}$  while the rate constant for fluorescence  $(k_{\rm F})$  is only  $\sim 10^6 \ {\rm sec^{-1}}$ ; (4) both S<sub>1</sub> and T<sub>1</sub> are n, $\pi^*$ states.

The lifetime of  $S_1(\tau_s)$  is defined as the reciprocal of the rates of deactivation of  $S_1$  (eq 6) and is inherently limited by the rate of intersystem crossing. Although the quantum yield of fluorescence from  $S_1$  for acetone

$$\tau_{\rm S} = (k_{\rm F} + k_{\rm ST})^{-1} = k_{\rm ST}^{-1} \tag{6}$$

is quite low ( $\sim 0.001$ ), this emission is easily detected by sensitive fluorescence techniques. Fluorescence is a particularly versatile and valuable probe for quantitative study of the photoreactions of  $S_1$  of alkanones.

If  $T_1$  is produced, its major mode of deactivation is intersystem crossing to  $S_{0.13}$  The latter process has a rate constant  $(k_{\rm D})$  which is considerably smaller (~10<sup>6</sup> sec<sup>-1</sup>) than the intersystem crossing process  $(k_{\rm ST} \sim 5 \times$  $10^{8}$  sec<sup>-1</sup>), which serves as the major deactivation for S<sub>1</sub>. The lifetime of T<sub>1</sub> ( $\tau_{\rm T}$ ) is thus given by  $\tau_{\rm T}$  =  $(k_{\rm D})^{-1}$ .

Although the  $T_1$  state possesses about the same amount of excess electronic energy ( $\sim$ 78 kcal/mole) as  $S_1$ , quenching by electronic energy transfer to a qualified energy acceptor is generally more important for  $T_1$ than for  $S_1$  because of the longer lifetime of  $T_1$ .

#### Measurement of the Reactivity of $S_1$ and $T_1$

 $S_1$  and  $T_1$  of alkanones have certain "inherent" unimolecular paths for deactivation. If a molecule, Q, is capable of efficiently "robbing"  $S_1$  or  $T_1$  of its electronic excitation, then Q is called a "quencher." Experimentally, if the efficiency of a decay path of  $S_1$  (e.g., fluorescence) or  $T_1$  (e.g., photoreaction) is measured as a function of [Q], then one has obtained, in principle, information on the relative rates of quenching and excited-state deactivation. Equation 7, called the Stern-Volmer equation, is a convenient expression for relating the deactivation of  $S_1$  and/or  $T_1$  to a quenching process. In this expression,  $\phi^0$  is the quantum yield for a photoprocess (from either  $S_1$  or  $T_1$ ) in the absence

$$\frac{\phi^0}{\phi} = 1 + k_q[Q]r \tag{7}$$

of Q,  $\phi$  is the quantum yield for the same process in the presence of Q,  $\tau$  is the lifetime of the reactive state in the absence of Q,  $k_{q}$  is the bimolecular rate constant for quenching of the reactive state, and [Q] is the concentration of quencher. Thus, the slope of an experimentally linear plot of  $\phi^0/\phi$  vs. [Q] yields  $k_{\rm q}\tau$ , which is simply  $k_{\rm q}/k_{\rm d}$  where  $k_{\rm d}$  is the "inherent" rate constant for deactivation of the excited state;  $k_d$  is generally directly relatable to the reactivity of  $S_1$  and/or  $T_1$ . Finally, if a plot of  $\phi^0/\phi$  vs. [Q] is not linear (and experimental artifacts for the nonlinearity can be reasonably eliminated), then either the reaction or the quenching must be occurring from more than one excited state.<sup>14</sup>

Fluorescence quenching is an exceedingly specific and convenient technique for measuring the reactivity of alkanones toward various primary processes. In effect, the absorbed photon simultaneously "labels" and activates the alkanone  $S_1$  state, while the emitted fluorescence photon serves as a built-in "clock" to tell us how fast our labeled ketone reacts or is quenched. We can use eq 7 by measuring the quenching of fluorescence as a probe of the reactivity of  $S_1 (k_q = k_q^F)$ .

Unfortunately, since phosphorescence from alkanones is generally too weak to be measured in fluid solution at room temperature, an indirect method, usually quenching of  $T_1$  by 1,3-dienes, must be employed to monitor the reactivity of alkanone triplets.<sup>15</sup> 1,3-Dienes are probably diffusion-controlled quenchers of  $T_1$ . Primary photochemical processes from  $T_1$  can be made to compete with bimolecular diene quenching, and if Stern-Volmer kinetics are followed, the slope of the Stern-Volmer plot yields information on the reactivity of T<sub>1</sub>.

### The $\alpha$ -Cleavage (Type I) Reaction

Although irradiation of acetone in "inert" organic solvents does not lead to efficient photoreaction associated with  $\alpha$  cleavage,<sup>16a</sup> methyl *tert*-butyl ketone and di-tert-butyl ketone undergo photolysis to hydrocarbon and carbonyl fragments.<sup>16b</sup>

Nearly all cyclopentanones and cyclohexanones have a tendency to photorearrange to an unsaturated aldehyde and/or a ketene (Scheme I).<sup>17a,b</sup> These reactions are nicely correlated by the following simple working hypothesis: photon absorption is followed by homolytic cleavage of a bond  $\alpha$  to the excited carbonyl function. In the case of acyclic ketones, the result of  $\alpha$  cleavage is

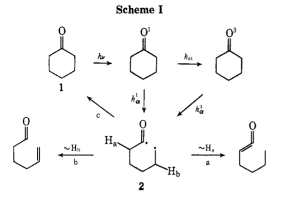
<sup>(12)</sup> The energies of  $S_1$  and  $T_1$  are only approximate, since structure is essentially absent in the electronic spectra of acetone.

<sup>(13)</sup> R. F. Borkman and D. R. Kearns, J. Amer. Chem. Soc., 88, 3467 (1966).

<sup>(14)</sup> J. C. Dalton and N. J. Turro, Mol. Photochem., 2, 133 (1970).

<sup>(15)</sup> For a recent review see ref 9a, p 96 ff.
(16) (a) N. C. Yang in "Reactivity of the Photoexcited Organic Molecule," Interscience, New York, N. Y., 1967, p 145; (b) N. C. Yang and E. D. Feit, J. Amer. Chem. Soc., 90, 504 (1968).

<sup>(17)</sup> For reviews of recent work on the photochemistry of cyclic ketones see: (a) N. J. Turro, et al., Annu. Surv. Photochem., 2, 1970; (b) P. Yates, Pure Appl. Chem., 16, 93 (1968).



production of a radical pair in a solvent cage. The observed products then follow naturally from the expected chemistry (in and out of the solvent cage) of an acylalkyl radical pair: disproportionation, decarbonylation, and hydrogen abstraction from solvent. For 5- and 6-ring ketones, a 1,5- and 1,6-biradical, respectively, are produced, and intramolecular disproportionation yields the observed predominant products.

Let us now ask the question: In which state or states,  $S_1$  and/or  $T_1$ , does  $\alpha$  cleavage occur? In order to answer questions of this type there is a standard technique employed which takes advantage of the very fast rate of quenching of alkanone triplet states by 1,3-dienes. One measures the efficiency of formation of  $\alpha$ -cleavage products as a function of 1,3-diene concentration. Since unimolecular  $\alpha$  cleavage from T<sub>1</sub> will compete with bimolecular quenching by 1.3-diene (to regenerate  $S_0$  of the alkanone), a Stern-Volmer plot, which obeys eq 7, will be observed experimentally if the reaction occurs only from  $T_1$  and only  $T_1$  is quenched by the diene. Results for methyl tert-butyl ketone<sup>16b</sup> and for cyclohexanone<sup>9c</sup> (1) reveal that, while the plot for cyclohexanone does indeed follow Stern-Volmer kinetics, the plot for methyl *tert*-butyl ketone is distinctly nonlinear, decreasing in slope at higher concentrations of 1,3-diene. This latter behavior is prima facie evidence for reaction from two different alkanone excited states.<sup>14</sup> In addition, the quenching which occurs at high diene concentration is paralleled by quenching of ketone fluorescence.<sup>18,19</sup> We can conclude that quenching of  $S_1$  is occurring at high concentrations of diene, while at low concentrations of diene predominantly  $T_1$  is being quenched. The data allow us to estimate the reactivity of both  $S_1$  and  $T_1$  of 2 toward  $\alpha$  cleavage, while for 1 we can only estimate the reactivity for  $T_1$  because no reaction from  $S_1$  could be detected by diene quenching. However, since the lifetime of  $S_1$  can be measured directly,<sup>20</sup> the *total* rate of deactivation of  $S_1$  is available and an *upper* limit for the reactivity of  $S_1$  toward  $\alpha$  cleavage can be obtained.<sup>21</sup>

Analysis of such data<sup>9,16,20</sup> has led to the following generalizations concerning the  $\alpha$ -cleavage process: (1) there is a startling difference in reactivity toward  $\alpha$ cleavage between  $T_1$  and  $S_1$  for alkanones,  $T_1$  being generally more reactive; (2) substitution of  $\alpha$  hydrogens by  $\alpha$ -alkyl groups increases T<sub>1</sub> reactivity toward cleavage; (3) cleavage of the more substituted bond (which parallels radical stability) generally determines product structure; (4) a "loss" mechanism exists for dissipating the energy of absorbed photons without net product formation.

We can summarize the general pattern with cyclohexanone as an example (Scheme I). Path c in Scheme 1 is included in order to provide a "loss" mechanism for photon energy. This choice, rather than a more general "radiationless deactivation" of  $T_1$  (no or very feeble phosphorescence is generally observed for alkanones in fluid solution), is preferred. A loss mechanism is required to explain that, although a relative reactivity difference of 100-1000 is observed experimentally,9,16,20 the efficiencies of net reaction are always significantly Thus, the proposal that reactivity less than unity. measures the rate of formation of an intermediate, and the product formation efficiency measures the stabilization of the intermediate, is an attractive model to help us understand the available data. The partitioning of the radical pair produced by  $\alpha$  cleavage will depend on the ease of formation of transition states for transfer of H<sub>a</sub>, H<sub>b</sub>, and recyclization. Use of molecular models shows that intermediates such as 2 can be used to predict variations in the relative rates of paths a and b as a function of structure.<sup>22</sup> Finally, strong support for the existence of an intermediate such as 2 and a general loss mechanism as suggested by path c comes from the observation that cis- and trans-2,3-dimethylcyclohexanones<sup>23</sup> undergo photoepimerization concomitant with photoisomerization by way of  $\alpha$  cleavage.

The photochemistry of cyclobutanones differs significantly from that of most alkanones.<sup>24</sup> For example, three major products which do not have a general analogy in cyclopentanone or cyclohexanone photochemistry are generally formed. Indeed, product formation is both stereospecific and stereoselective (eq 8-10), and no photoepimerization occurs during the photolyses.<sup>25</sup> Furthermore, in contrast to the relatively facile quenching of cyclopentanone photochemistry by 1,3-dienes, even high concentrations (>1 M) of the latter do not measurably<sup>25</sup> inhibit cyclobutanone reactions. One can conceive of an intermediate, formed by  $\alpha$  cleavage, which can be stabilized by reasonable paths which will lead to the observed products. However, one must impose a special quality on such an intermediate-

<sup>(18)</sup> N. C. Yang, E. D. Feit, M. H. Hui, N. J. Turro, and J. C. Dalton, J. Amer. Chem. Soc., 92, 6974 (1970).

<sup>(19)</sup> F. S. Wettack, G. D. Renkes, M. G. Renkly, N. J. Turro, and J. C. Dalton, *ibid.*, 92, 1793 (1970).

<sup>(20)</sup> D.S. Weiss, N. J. Turro. and J. C. Dalton, Mol. Photochem., 2, 91 (1970).

<sup>(21)</sup> J. C. Dalton, D. M. Pond, D. S. Weiss, F. D. Lewis, and N. J. Turro, J. Amer. Chem. Soc., 92, 2564 (1970).

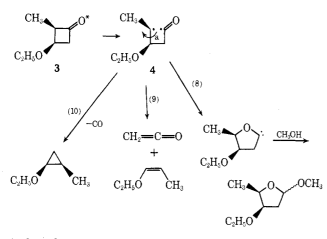
<sup>(22)</sup> G. Quinkert, Angew. Chem., Int. Ed. Engl., 4, 211 (1965); Pure Appl. Chem., 9, 607 (1964).

<sup>(23)</sup> J. A. Barltrop and J. D. Coyle, Chem. Commun., 1081 (1969).

 <sup>(24) (</sup>a) N. J. Turro and R. M. Southam, Tetrahedron Lett., 545
 (1967); (b) N. J. Turro, E. Lee-Ruff, D. R. Morton, and J. M. Conia. ibid., 2991 (1969); (c) D. R. Morton, E. Lee-Ruff, R. Southam, and

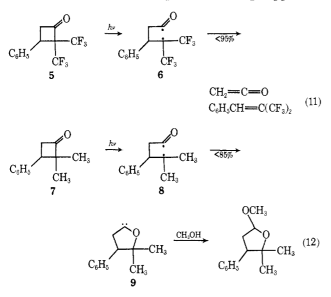
N. J. Turro, J. Amer. Chem. Soc., 92, 4349 (1970).
 (25) N. J. Turro and D. M. McDaniel, *ibid.*, 92, 5727 (1970).

it must collapse to products without loss of stereochemistry. Thus, if we consider the "biradical" 4 as a species formed by  $\alpha$  cleavage, 4 must undergo reaction via paths 8, 9, and 10 faster than rotation about bond "a."



Although such rapid unimolecular stabilization is conceivable, calling 4 a "biradical" does not sit well, if we expect such species to behave like two loosely intramolecularly connected monoradicals.<sup>26</sup> At any rate, we feel that either 4 exists briefly but then rapidly collapses to products or the transition states for product formation all have considerable biradical character. Since 1,3-dienes fail to quench the photoreactions of cyclobutanones, the reactive state might be  $S_1$  or an unquenchable  $T_1$  (or some mixture of both), although the stereospecificity might make a singlet reaction more attractive.25,27

The striking contrast<sup>28</sup> in the photochemical behavior of 5 and 7 (eq 11 and 12) provides strong support for



an intermediate or transition state such as 6 and/or 8. Since the most substituted  $\alpha$  bond breaks in both cases, in spite of considerably different electronic den-

(28) D. R. Morton, Columbia University, unpublished results.

sities of the  $\alpha$  bond which is cleaved, it appears that the  $\alpha$  bond must be nearly completely cleaved before the product-determining step occurs. On the basis of hypotheses of transition states or intermediates such as 6 and 8 we can understand why (a) the most substituted bond cleaves and (b) why the more nucleophilic alkyl carbon of 8 migrates preferentially relative to the electron-poor free-radical carbon of 6.

## Hydrogen Abstraction Reactions of Alkanones

Only a handful of reactions concerned with intermolecular hydrogen abstraction reactions of S<sub>1</sub> and/or T<sub>1</sub> states of alkanones have been reported.<sup>29</sup> A report has appeared<sup>29a</sup> in which evidence for a large reactivity difference between  $T_1$  and  $S_1$  states toward hydrogen abstraction from (n-Bu)<sub>3</sub>SnH was presented. Later work, however, showed that complications,<sup>27</sup> *i.e.*, possible "quenching" of ketyl radicals by dienes and reactions of tin radicals with dienes,<sup>30</sup> can vitiate standard kinetic analyses. 2-Adamantanone fluorescence<sup>31</sup>  $(i.e., S_1)$  is quenched by  $(n-Bu)_3SnH$  with a rate constant of 5  $\times$  10<sup>8</sup>  $M^{-1}$  sec<sup>-1</sup>. This number is quite comparable to the reported rate ( $\sim 8 \times 10^8 M^{-1} \, \mathrm{sec^{-1}}$ ) for hydrogen abstraction from (n-Bu)<sub>3</sub>SnH by T<sub>1</sub> of acetone.29a

The intramolecular analog (type II abstraction) has been the focus of considerable recent investigation.<sup>32</sup> Although here also complications of kinetic analyses have been found, several reactivity and efficiency trends appear to be clear-cut.

Since intermolecular hydrogen abstraction seems to clearly involve the intermediacy of ketyl radicals, an analogous model may explain the intramolecular counterpart. We shall assume a model in which reaction from both  $S_1$  and  $T_1$  of alkanones structurally capable of undergoing type II reactions proceeds with biradicallike transition states and then ask whether evidence exists to suggest that such a transition state will lead to actual biradical intermediates such as 11 (Scheme II). We will try also to inquire about the spin states of such intermediates.

It is generally found that  $k_d$  (which is the measure of all deactivation paths of  $S_1$  and  $T_1$ ) increases as the abstracted hydrogen goes from primary to secondary to tertiary in keeping with expectations of the effect of bond strengths of the  $\gamma$  C-H bonds.<sup>32</sup> But again an inefficiency is noted, as measured by  $\phi_{II}$ , which is maintained throughout the series even though rates go up by an order of magnitude.

Available data suggest that the  $T_1$  reaction very probably proceeds via a relatively long-lived biradical; e.g., racemization of the  $\gamma$  carbon of 10 (R<sub>1</sub> = CH<sub>3</sub>;  $R_2 = C_2H_5$ ) (Scheme II) appears to occur from  $T_1$ .<sup>33</sup> Thus, the triplet must survive long enough to undergo

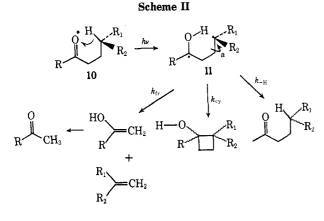
(29) (a) P. J. Wagner, J. Amer. Chem. Soc., 88, 5672 (1966); (b) R. Simonaitis, G. W. Cowell, and J. N. Pitts, Jr., Tetrahedron Lett., 3751 (1967).
(30) N. J. Albert, et al., Chem. Ber., 103, 1372 (1970).

(31) G. Farrington and R. Hautala, Columbia University, unpublished results

(32) P. J. Wagner, Accounts Chem. Res., 4, 168 (1971).

<sup>(26)</sup> For interesting discussions of the biradical problem see (a) R. Freeman, Can. J. Chem., 44, 245 (1966); (b) L. M. Stephenson and J. I. Brauman, J. Amer. Chem. Soc., 93, 1988 (1971). (27) N. J. Turro and D. M. McDaniel, Mol. Photochem., 2, 98

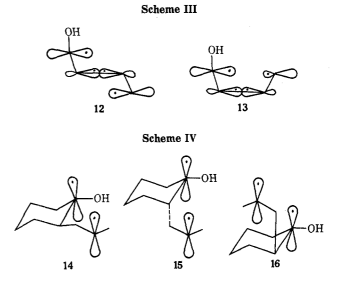
<sup>(1970)</sup> 



rotations about bond a and *then disproportionate* by hydrogen return to yield the optical enantiomer of the starting material.<sup>33</sup> This result is also consonant with the observation that the efficiency of the type II reaction from  $T_1$  is enhanced as solvent polarity increases, if one assumes that the intermediate biradical **11** survives long enough to be solvated and that intermolecular hydrogen bonding interferes to a greater extent with hydrogen reversal than with the fragmentation and cyclization.

On the other hand, the  $S_1$  state apparently does not yield a biradical which survives long enough to have its chemistry affected by solvent. In addition, the  $S_1$ reactions of 10 do not result in racemization and are not totally efficient, thereby again demanding a "loss mechanism." Although hydrogen reversal would explain the inefficiency, *i.e.*, from a "singlet biradical," we must again impose requirements on the rates of internal molecular motions of this species, *i.e.*, rotations about bond a must be slow relative to hydrogen reversal, in order to explain the lack of racemization from  $S_1$ . This assumption implies that an ethylene (in a properly labeled case) should be formed stereospecifically from  $S_1$ , as has been found.<sup>26b</sup>

Measurements of reactivity and efficiency can thus help reveal "invisible" radiationless paths which regenerate the starting materials. For example, quenching data suggest that type II abstraction determines the triplet lifetime of 2-n-propylcyclohexanone  $(1/\tau_{\rm T} = 7 \times 10^8 \text{ sec}^{-1}; 1/\tau_{\rm T}$  for 2-methylcyclohexanone is  $3 \times 10^8 \text{ sec}^{-1}$ ), yet only a very small percentage of the T1 states undergo net type II fragmentation or cyclization.<sup>20,34</sup> Thus, while kinetic analyses suggest that a major deactivation path of  $T_1$  is type II abstraction, efficiency measurements demand that hardly any net type II reaction occurs from  $T_1$ . Kinetic and efficiency analyses of the  $S_1$  state lead to the conclusion that the rate of type II abstraction from  $S_1$  is  $\sim 10^9$  sec<sup>-1</sup>, with an efficiency of ca. 10%. Interestingly then, the  $T_1$  and  $S_1$  states are of comparable reactivity, but  $S_1$  is more efficient at going to products,



in striking contrast to acyclic alkanones whose  $T_1$  state goes to products with up to 80--100% efficiency in some cases.  $^{33-35}$ 

How can we understand these results? Let us assume that (a) a biradical intermediate is produced by type II abstraction from both  $S_1$  and  $T_1$  and that (b) it can undergo fragmentation, cyclization, or hydrogen return. Hydrogen return to regenerate the starting ketone is apparently much faster than fragmentation or cyclization. This derives naturally from consideration of the principle of maximum overlap of developing orbitals and the shape of the lowest energy transition states for cleavage and cyclization of a 1,4-biradical. Transition states similar to 12 and 13, in which the p orbitals at the free-radical centers can be parallel to the  $\beta$  bond, should be favored for cleavage (Scheme III), since they allow maximum continuing overlap of the developing  $\pi$  orbitals of the enol and alkene. Now, in the case of 2-n-propylcyclohexanone, species 14 is produced by intramolecular  $\gamma$ -hydrogen abstraction. Molecular models confirm what Scheme IV suggests: for 14 a transition state analogous to 15 is not attainable without severe molecular distortion, and a transition state like 16 requires either flipping the a higher energy shape, *i.e.*, one with an axial alkyl group, or a bond rotation to a twisted chair conformation. These unfavorable shapes may also be expected to resemble the best transition state for cyclization, thereby explaining the inefficiency of both cyclization and fragmentation. On the other hand, hydrogen reversal does not suffer from such orbital restrictions since only one new  $\pi$  system is to be developed and, indeed, the reversal becomes highly favored relative to the other two processes.

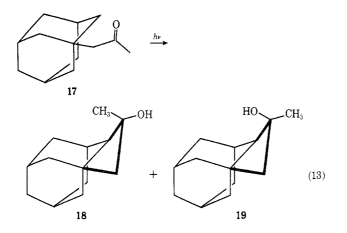
The 2-adamantyl-acetone system dramatically illustrates the effect of solvent polarity on type II reactions from  $S_1$  and  $T_1$ . Irradiation of 17 in benzene results in a quantitative yield<sup>36</sup> of the alcohols 18 and 19. In

<sup>(33) (</sup>a) N. C. Yang and S. P. Elliott, J. Amer. Chem. Soc., 91, 7550 (1969);
(b) N. C. Yang, S. P. Elliott, and B. Kim, *ibid.*, 91, 7551 (1969).

<sup>(34)</sup> D. S. Weiss and K. Dawes, Columbia University, unpublished results.

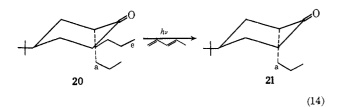
<sup>(35)</sup> P.J. Wagner, Tetrahedron Lett., 5385 (1968).

<sup>(36)</sup> R. B. Gagosian, J. C. Dalton, and N. J. Turro, J. Amer. Chem. Soc., 92, 4752 (1970).



MeOH the efficiency of formation of 18 and 19 from the  $S_1$  state is the same as in benzene, but the efficiency of formation of 18 and 19 from  $T_1$  is increased by a factor of 10. As expected,<sup>35</sup> the more polar solvent inhibits hydrogen back-transfer in the biradical produced from  $T_1$  only, and allows cyclization to compete more effectively with regeneration of starting ketone.

The proposed model for the  $n, \pi^*$  state suggests that type II abstraction from a side chain for which cyclic interaction of a  $\gamma$  hydrogen with the half-vacant n orbital is geometrically feasible should be facile, relative to abstraction of a  $\gamma$  hydrogen which is constrained by molecular geometry to approach the carbonyl oxygen atom from one of the  $\pi$  faces. This conclusion derives from the electrophilic nature of hydrogen abstraction by ketone  $n, \pi^*$  states. A prediction, therefore, for **20** is that only the equatorial group should be cleaved photochemically, thereby forming the less stable 2-*n*-propyl epimer,<sup>37</sup> and this is indeed found to be the result. In this case, 1,3pentadiene is a convenient solvent to quench  $\alpha$ cleavage reactions originating from T<sub>1</sub>.



## Photocycloaddition of Alkanones to Ethylenes and Photosensitized Reactions of Ethylenes

Let us now consider the interaction of the  $S_1$  and  $T_1$ states of alkanones with the following classes of ethylenes: (a)  $\alpha,\beta$ -unsaturated nitriles<sup>38</sup> and (b) enol ethers.<sup>39</sup> Equations 15–18 summarize the types of reactions which are observed in such systems. Clearly, there exists a striking contrast in the behavior of these systems with respect to stereospecificity, regioselec-

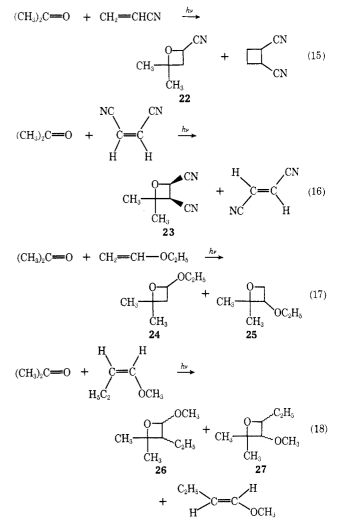
 Table I

 Relative Rates for Quenching of Alkanone Fluorescence for Some

  $\alpha,\beta$ -Unsaturated Nitriles and Some Enol Ethers<sup>a</sup>

	t-NCCH=	$CH_2 = $	$CH_2 = C$	CH₃CH==
Alkanone	CHCN	CHCN	(CH₃)CN	CHCN
Acetone	1.08	0.046	0.020	<0.01
Adamantanone	1.0°	0.020	0.011	0.001
	Enol Ethers			
	c-EtOCH=	CH2=C-	$CH_2 == C-$	$CH_2 = $
	CHOEt	$(OEt)_2$	(CH <sub>3</sub> )OCH <sub>3</sub>	CHOEt
Acetone	1.0ª	0.42	$\sim 0.11$	<0.032
Adamantanone	1.0	0.13	$\sim 0.017$	0.007

<sup>a</sup> N. J. Turro, N. Schore, C. G. Lee, J. A. Barltrop, and H. A. J. Carless, J. Amer. Chem. Soc., **93**, 3079 (1971). Acetonitrile solution. Error limits  $\pm 20\%$ . <sup>b</sup> The absolute quenching rate constant is  $2.5 \times 10^9 M^{-1} \sec^{-1}$ . <sup>c</sup> The absolute quenching rate constant is  $5.0 \times 10^9 M^{-1} \sec^{-1}$ . <sup>d</sup> The absolute quenching rate constant is  $2.0 \times 10^9 M^{-1} \sec^{-1}$ . <sup>e</sup> The absolute quenching rate constant is  $1.1 \times 10^9 M^{-1} \sec^{-1}$ .



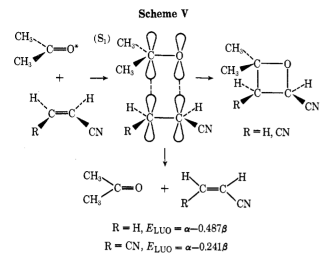
tivity, and sensitization of ethylene reactions. With such a wide variation in behavior one is faced with a challenging task to present a unifying theory, consistent with the previously discussed behavior of alkanone excited states.

Acetone fluorescence quenching by ethylenes (Table I) offers some real insight into and places severe restrictions on a unifying explanation of how the excited

<sup>(37)</sup> K. Dawes, J. C. Dalton, and N. J. Turro, Mol. Photochem., 3, 71 (1971).

<sup>(38) (</sup>a) N. J. Turro, P. A. Wriede, and J. C. Dalton, J. Amer. Chem. Soc., 90, 3274 (1968); (b) J. J. Beereboom and M. von Witterman, J. Org. Chem., 30, 1231 (1965).

<sup>(39) (</sup>a) S. H. Schroeter and C. M. Orlando, *ibid.*, 34, 1181 (1969);
(b) N. J. Turro and P. A. Wriede, *ibid.*, 34, 3562 (1969).



alkanone-ethylene interaction occurs. We see that the reactivity of  $S_1$  toward unsaturated nitriles, which should behave as electrophilic (electron-poor) substrates, increases roughly as the "electron poorness" of the C=C bond increases. On the other hand, the reactivity of  $S_1$  toward enol ethers, which should behave as nucleophilic (electron-rich) substrates, increases roughly as the "electron richness" of the C=C bond increases. Remember that  $S_1$  is an amphoteric reagent (in the sense of Lewis acids and bases). In fact, the simple model of an  $n, \pi^*$  state has precisely the properties desired to explain the amphoteric nature of the  $S_1$  state of alkanones: (a) an electrophilic region, due to a half-vacant n orbital, about the "edges" of the carbonyl oxygen, and (b) a nucleophilic region of space, due to the presence of a  $\pi^*$  electron, above and below the carbonyl "faces" (see Figure 2). The precise geometry of the S<sub>1</sub> state is deduced to be somewhat puckered on the basis of spectroscopic analyses,<sup>40</sup> although rapid inversion of one puckered shape into another is also expected. Thus, we anticipate that the interaction between the  $S_1$  state and the nucleophilic cyanoethylenes will be most favorable when the reagent and substrate approach each other so that the  $\pi^*$  electron of S<sub>1</sub> overlaps most effectively with the lowest unoccupied orbital (LUO) of the cyanoethylenes. Such an expectation allows us to write down an interaction which nicely rationalizes the stereospecificity, regioselectivity, and reactivity of  $S_1$  of alkanones and cyanoethylenes: the geometry for the most favored interaction transforms smoothly into the geometry of the product, and, therefore, stereospecificity is a natural consequence of the  $\pi^*$ -LUO interaction (Scheme V).

Indeed, the regioselectivity also nicely derives from such an interaction since the  $\pi^*$  orbital of the S<sub>1</sub> state is expected to have a coulombic negative character near the carbonyl carbon and a coulombic positive character near the carbonyl oxygen. Therefore, the sole formation of **22** is easily understandable on the basis of stabilization of plus-minus forces. Finally, the energy of the LUO of CH<sub>2</sub>=CHCN is much higher than that

(40) H. E. Howard-Lock and G. W. King. J. Mol. Spectrosc., 36, 53 (1970).

Table IIQuenching of the Fluorescence of Norcamphors $^{a,b}$ 

Alkanone	$k_q^f(t\text{-}DCE)$	$k_q^f(c\text{-}\text{DEE})$	
	5.3	1.3	
$\mathcal{A}^{\circ}$	2.1	0.16	
	1.0	1.5	
A~°	5.7	1.4	

<sup>a</sup> Reference 43a and unpublished results. <sup>b</sup> 0.1 M solutions of ketones in acetonitrile.  $k_q^f$  values  $\times 10^{-9} M^{-1} \sec^{-1}$ .

of NCCH=CHCN; therefore, the rate constant for quenching by the latter should be faster, and is indeed found to be, in spite of increased steric hindrance to the approach of the interacting centers.

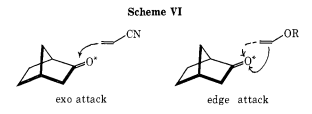
The maximum efficiency of oxetane formation from  $S_1$  is only about 10%. Thus, 90% of the photon energy is lost and does not result in isomerization or dimerization (at least in the case of *trans*-1,2-dicyanoethylene, DCE).<sup>41</sup> This result leads us to propose that the interaction between  $S_1$  and DCE yields an exciplex,<sup>42</sup> *i.e.*, an aggregate of  $S_1$  and DCE which possesses some measure of stability and for which electronic excitation is a property of the aggregate rather than being essentially localized on either component. The exciplex can then (a) stereospecifically dissociate into  $S_0$  of the alkanone and  $S_0$  of DCE with the electronic energy being dissipated as increased translational or vibrational energy of the fragments and solvent, or (b) stereospecifically collapse into oxetane (Scheme V).

A compelling confirmation of the strong directional specificity of quenching of  $S_1$  by DCE was obtained by a study of fluorescence quenching<sup>43</sup> of norcamphor and some of its alkyl derivatives (bicyclo[2.2.1]-2-heptanones) by DCE (Table II). The results convincingly demonstrate a preference for quenching by DCE above and below the carbonyl faces, *i.e.*, the region of space expected to be nucleophilic according to the  $n,\pi^*$  state model. Furthermore, 7-methyl groups have the strongest effect on inhibiting the rate of quenching by DCE. It should be noted that the response of  $k_q^t$  (eq 7, fluorescence quenching) to ketone structure shows a strong resemblance to the reactivity for attack of nucleophiles on the ground states of bicyclo[2.2.1]-2-

<sup>(41)</sup> J. C. Dalton, P. A. Wriede and N. J. Turro, J. Amer. Chem. Soc., 92, 1318 (1970).

<sup>(42)</sup> T. Forster, Angew. Chem., Int. Ed., Engl., 8, 333 (1969); exciplexes are molecular aggregates defined as being stable in an electronically excited state but not in their ground states.

<sup>(43) (</sup>a) N. J. Turro, J. C. Dalton, G. Farrington, M. Niemczyk, and D. M. Pond, J. Amer. Chem. Soc., 92, 6978 (1970); for related studies see (b) J. C. Dalton D. M. Pond, and N. J. Turro, *ibid.*, 92, 2173 (1970); (c) N. J. Turro, M. Niemczyk, and D. M. Pond, Mol. Photochem., 2, 345 (1970).



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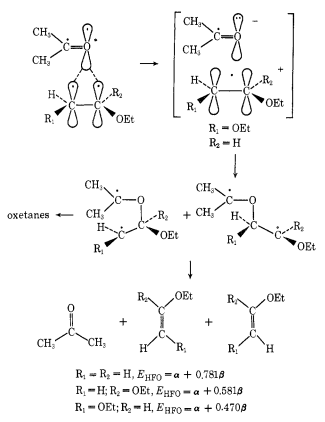
heptanones; *i.e.*, attack on the *exo* face of the carbonyl group is favored (Scheme VI).

In striking contrast to the situation for unsaturated nitriles, the formation of oxetanes from enol ethers is neither stereospecific nor regioselective. Furthermore, the reactivity pattern resembles that expected for attack by an electrophilic reagent on the nucleophilic enol ethers. Our model for the  $n, \pi^*$  state suggests that the edges of the carbonyl oxygen atom attack the highest filled orbital (HFO) of the enol ethers, *i.e.*, the reactivity increase indicated in Table I parallels the increase in energy of the electrons in the enol ether's HFO. This, in turn, suggests a transition state for quenching which possesses considerable charge-transfer (CT) character. In fact, if the transition states for product formation also are assumed to possess CT character, then the lack of stereospecificity and regiospecificity can be explained in the following way. Suppose the interaction of, e.g.,  $S_1$  with an enol ether is dominated by the overlap of the half-filled n orbital with the HFO of the enol ether to form either a radical anion-radical cation pair or to lead to a transition state for quenching very similar to such a pair, *i.e.*, as shown in Scheme VII. The charge densities of  $C_1$  and  $C_2$  of a radical cation of an enol ether turn out to be very similar, so that the collapse of a transition state possessing considerable CT character (or an actual radical cation-radical anion pair) might well be very unselective with respect to the regiospecificity of C-O bond formation. The loss of stereospecificity could come about in one of two ways: the radical cation may have a sufficiently low bond order that it may isomerize prior to oxetane formation or capture of an electron to regenerate an enol ether; the second possibility is that collapse of the radical cation-radical anion pair leads to biradicals which themselves lose stereochemistry and collapse to oxetanes or ground-state alkanone and enol ether. Calculations<sup>44</sup> indicate that the bond order in the radical cations of enol ethers is too high to permit facile bond rotation.

Another interesting aspect of oxetane formation and olefin isomerization is revealed by quenching studies with 1,3-pentadiene. At a fixed concentration of 1methoxybutene (0.05 M), linear Stern–Volmer kinetics for quenching of T<sub>1</sub> of acetone are observed at low concentrations of diene (<0.05 M), while at higher concentrations of diene (>0.1 M) the Stern–Volmer plot shows clear-cut curvature downwards, until at ~0.4 M a residual unquenchable reaction remains.<sup>45</sup>

(44) N. Schore, Columbia University, unpublished results. For parameters see: B. Pullman and A. Pullman, *Rev. Mod. Phys.*, **32**, 428 (1960).



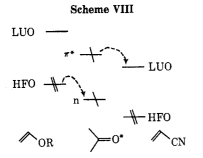


Thus, we conclude that at higher concentrations of diene  $S_1$  is reacting with methoxybutene to yield oxetane and to isomerize the enol ether, while at lower concentrations of diene both  $S_1$  and  $T_1$  attack the enol ether. Importantly, (a) there is greater oxetane-forming stereospecificity in the  $S_1$  reaction relative to  $T_1$  reaction, (b) the efficiency of isomerization by  $S_1$  appears to be less than that of  $T_1$ , (c) the rate constants for attack by  $S_1$  and  $T_1$  are comparable, and (d) the regiospecificity for oxetane formation is essentially the same for attack of  $S_1$  or  $T_1$  on 1-methoxybutene.<sup>45</sup>

Points a and b relate, we believe, to the spin state immediately preceding the intermediates (charge radical pairs or 1,4-biradicals) which lose stereochemistry, *i.e.*, the singlet intermediates tend to cyclize to oxetanes or fragment to starting alkanone and enol ether favorably relative to loss of stereochemistry, However, the triplet intermediates may only rotate about single bonds, thereby losing initial stereochemical memory, until a spin flip occurs. At that point the newly formed singlet biradical can collapse to stable products. Indeed, as measured by stereochemical loss in oxetane formation, the triplet intermediate(s) undergo many bond rotations before cyclization occurs.

Now let us consider the alkanone photosensitization<sup>46</sup> of ethylene dimerization and isomerization and ask whether dimerizations and cis-trans isomerizations are mechanistically connected to photocycloaddition,

(45) N. J. Turro and P. A. Wriede, J. Amer. Chem. Soc., 92, 320
(1970); 90, 6863 (1968).
(46) N. J. Turro, Photochem. Photobiol., 9, 555 (1969).



or are they unrelated side reactions?<sup>11</sup> The isomerization and dimerization of the unsaturated nitriles probably result solely as a result of triplet energy transfer from T<sub>1</sub> to the ethylene. The triplet energy ( $E_3$ ) of CH<sub>2</sub>=CHCN is estimated to be ~60 kcal/mole above its ground state.<sup>47</sup> Thus, T<sub>1</sub> of alkanones ( $E_3 \sim 78$ kcal/mole) should transfer excitation to  $\alpha_{,\beta}$ -unsaturated nitriles at close to the diffusion-controlled rate. Whether dimerization occurs or not is probably a function of unsaturated nitrile structure, with 2-substituents reducing the efficiency of triplet ethylene coupling to form dimers, relative to simple deactivation of triplet ethylene by rotation about the excited C=C bond.

Scheme VIII summarizes our view of the leading interactions between the  $n,\pi^*$  states of alkanones and ethylenes. This formulation emphasizes donor-acceptor interactions in the transition state for quenching. Thus, a charge transfer from the HFO of an enol ether to the electrophilic n orbital in the one case, and a charge transfer from the  $\pi^*$  orbital to the LUO of the unsaturated nitrile, appear to be the most important electronic factors in the quenching step.

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<sup>(47) (</sup>a) D. M. Gale, J. Org. Chem., 35, 970 (1970); (b) R. S. H.
Liu and D. M. Gale, J. Amer. Chem. Soc., 90, 1897 (1968); (c) M.
Herberhold and G. S. Hammond, Ber. Bunsenges. Phys. Chem., 72, 309 (1968).