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Mechanistic Probes
of Organic
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ROBERT C. NEUMAN, JR.

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Pressure Effects as Mechanistic Probes of Organic Radical Reactions

Robert C. Neuman, Jr.

Department of Chemistry, University of California, Riverside, California 92502 Received March 29, 1972

Activation parameters are useful in understanding

Robert Neuman received the Ph.D. from Cal Tech under the direction of George Hammond and then spent the 1962–1963 academic year as an NSF postdoctoral fellow with Cheves Walling at Columbia University. His active interest in high-pressure studies developed during this latter period. Besides the topic of this Account, another research area thas been nmr studies of hindered rotation reactions. He has been at University of California at Riverside since 1963 and is now Professor of Chemistry.

chemical reactions since they provide information about the nature of the activated complexes formed from the reactants. Enthalpies and entropies of activation $(\Delta H^*$ and ΔS^*) are routinely determined because they are easily obtained via the temperature dependence of rate constants. The volume change of activation (ΔV^*) is much less frequently reported, but in principle

should provide equally valuable mechanistic information. It reflects the change in volume of the reacting system on passing from reactants to transition state. Its infrequent appearance in the literature is the result of experimental difficulties associated with its determination.

The activation volume is obtained from the pressure dependence of a reaction rate constant (eq 1), and its

$$\partial \ln k / \partial P = -\Delta V^* / RT \tag{1}$$

magnitude determines how much the rate constant changes with pressure. While a change in reaction rate by a factor of 2 to 4 can be anticipated for a change in temperature of 10° , much greater pressure changes are required to bring about the same effect even for reactions with the relatively large activation volume of ± 40 cm³/mole (Table I). The special equipment re-

Table I
Pressure Required to Change a Reaction Rate by a Factor of 3

ΔV^* , cm ³ /mole	Pressure, atm
5	5800
10	2900
25	1200
40	700

quired to obtain the thousands of atmospheres of pressure needed has limited these studies to a few laboratories.

Rapid development of mechanistic organic chemistry occurred at a time when pressure equipment was not readily available to workers in the field. As a result, pressure studies have largely played a confirmatory role with respect to major mechanistic discoveries. However, pressure effects on reactions have often permitted choices between mechanisms and have also enabled substantial refinement of mechanistic detail. This is illustrated by our pressure studies on free-radical initiator decompositions which are reviewed here.

Besides presenting specific details about radical reactions, we hope that this discussion will also provide the reader with a basis for the general understanding of pressure effects as probes of reaction mechanism. The following section contains a brief general review of pressure effects on organic reactions. Extensive reviews have been published.¹

Background

Pressure accelerates reactions which are characterized by a volume shrinkage in passing from reactants through transition state (negative ΔV^*) and retards those with a volume expansion (positive ΔV^*). These volume changes arise from two sources: (1) making and breaking of chemical bonds (molecular reorganization); and (2) interactions of the reactants and acti-

Table II Volume Changes for Some Equilibria

Reaction	ΔV_0 , cm ³ /mole
$c\text{-ClCH} = \text{CHCl} \Rightarrow t\text{-ClCH} = \text{CHCl}$	+2
$NO_2 + NO_2 \rightleftharpoons N_2O_4$	-23
$\mathrm{CH_3CO_2H} + \mathrm{H_2O} \rightleftharpoons \mathrm{CH_3CO_2}^- + \mathrm{H_3O}^+$	-12
$CH_3NH_2 + H_2O \rightleftharpoons CH_3NH_3^+ + OH^-$	-27

vated complex with the medium. They are illustrated by the volume changes associated with the chemical equilibria shown in Table II. The first two entries are relatively free of solvent effects. The large volume change for dimerization of NO₂ reflects the expected decrease in volume of the system due to a decrease in the number of solute molecules. The small volume change for isomerization of 1,2-dichloroethylene is consistent with there being no major change in molecular structure and no change in the number of solute species. While the third and fourth reactions formally involve no change in numbers of solute species, the formation of ions and the resulting solvation lead to large volume decreases.

Observations on chemical equilibria provide a foundation for interpreting kinetic data. Some representative activation volumes are shown in Table III.¹ The first

Table III
Activation Volumes for Some Reactions

Reaction	ΔV^* , cm ³ /mole
$\begin{array}{c} PhCMe_2N{=\!\!\!\!\!=}NCMe_2Ph \to homolytic\\ scission \end{array}$	+5
$0 \longrightarrow 0$ + $\longrightarrow 0$ dimerization	-40
$n\text{-}\mathrm{C}_{6}\mathrm{H}_{13}\mathrm{SH} + \mathrm{DPPH}^{a} \rightarrow \mathrm{hydrogen}$ abstraction	-17
t -BuCl + H ₂ O \rightarrow solvolysis	-15
MeBr + OH ⁻ → displacement	-8
$Me_3S^+ + PhO^- \rightarrow displacement$ $(Me_2S + PhOMe)$	+12

^a DPPH is diphenylpicrylhydrazyl.

three reactions are nonionic and reflect changes in volume due to molecular reorganization in the transition states, the first involving volume expansion, and the second and third, volume contractions. The fourth reaction involves the creation of ions leading to increased solvation. The fifth and sixth reactions are both bimolecular displacements. The former involves no change in the number of charges, and solvation effects on ΔV^* are minimized, but the latter is characterized by charge neutralization in the transition state. The volume expansion in this last case clearly demonstrates the general trend that volume changes arising from creation or destruction of charge dominate those arising from molecular reorganization.

Activation Volumes and Homolytic Scission Mechanism

When we began pressure studies of homolytic scission reactions we were struck by the relatively wide range of values of ΔV^*_{obsd} for these processes. Literature

⁽¹⁾ Some reviews of pressure effects on chemical systems are: (a) S. D. Hamann in "High Pressure Physics and Chemistry," R. S. Bradley, Ed., Vol. II, Academic Press, New York, N. Y., 1963, Chapters 7ii and 8; (b) E. Whalley, Advan. Phys. Org. Chem., 2, 93, (1964); (c) W. J. le Noble, Progr. Phys. Org. Chem., 5, 207 (1967); (d) C. Eckert, Annu. Rev. Phys. Chem., in press.

data available at that time are shown in Table IV,2-4

Table IV Early Literature Data for Homolytic Scission Reactions⁵

Initiator	$\Delta V *_{\mathrm{obsd}}$, cm ³ /mole
Benzoyl peroxide	+10
tert-Butyl peroxide	+5 to +13
Pentaphenylethane	+13
Azoisobutyronitrile (AIBN)	+4

and our first result was the very small value of about +1 cm³/mole (cumene, solvent) for decomposition of tert-butyl phenylperacetate (eq 2).5,6

$$PhCH_2CO_2OCMe_3 \longrightarrow PhCH_2 \cdot CO_2 \cdot OCMe_3$$
 (2)

Consideration of these data led us to conclude that values of ΔV^*_{obsd} for decomposition of radical initiators do not necessarily reflect the volume change during homolytic scission. We proposed that values of ΔV^*_{obsd} for initiators which broke at least two bonds simultaneously in the primary decomposition process would reflect the characteristics of the homolytic scission step, but that this would not necessarily be true for those in which only one bond was broken (e.g., decomposition of diacyl peroxides, eq 3). In these latter

$$RCO_2O_2CR \Longrightarrow RCO_2 \cdot \cdot O_2CR$$
 (3)

cases recombination of the initial geminate radicals could lead to re-formation of starting material, a process unlikely in multiple bond scission systems (e.g., eq 2).

From a general mechanism including "return" of the initial geminate radicals (eq 4) it can be seen that the

initiator
$$\underset{k_{-1}}{\overset{k_1}{\rightleftharpoons}}$$
 radical pair $\underset{k_d}{\overset{k}{\rightleftharpoons}}$ cage reactions (4)

observed decomposition rate constant could depend not only on k_1 but also on k_{-1} , k_d , and k (eq 5).

$$k_{\text{obsd}} = k_1 \left[\frac{k_{\text{d}} + k}{k_{-1} + k_{\text{d}} + k} \right]$$
 (5)

pressure dependence of k_{obsd} would depend on that of this complex quantity, and observed activation volumes would be described by eq 6. We suggested that

$$\Delta V^*_{\text{obsd}} = \Delta V_1^* + RT \, \partial \ln \left[1 + k_{-1} / (k_d + k) \right] / \partial P$$
 (6)

 ΔV^*_{obsd} would usually be greater than the actual volume change for scission (ΔV_1^*) because it seemed to us that the ratio $k_{-1}/(k_d + k)$ should generally increase with pressure; radical recombination (k_{-1}) would be pressure accelerated, while separative diffusion (k_d) would be retarded with increasing pressure.6

(2) (a) A. E. Nicholson and R. G. W. Norrish, *Discuss. Faraday Soc.*, 22, 97, 104 (1956).
(3) C. Walling and G. Metzger, *J. Amer. Chem. Soc.*, 81, 5365

(4) A. H. Ewald, Discuss. Faraday Soc., 22, 138 (1956).

(5) P. D. Bartlett and C. Rüchardt, J. Amer. Chem. Soc., 82, 1756 (1960).

(6) R. C. Neuman, Jr., and J. V. Behar, ibid., 89, 4549 (1967).

(7) J. W. Taylor and J. C. Martin, ibid., 88, 3650 (1966).

We have examined pressure effects on decomposition of a variety of two-bond scission initiators (Table V).8-12 Their decomposition activation volumes of

Table V **Activation Volumes for Homolytic Scission** of Two-Bond Initiators in Cumene

Initiator	$\Delta V *_{\rm obsd}$, cm ³ /mole
$\mathrm{Me_{2}C(CN)N_{2}C(CN)Me_{2}}$	+4
$Me_2C(Ph)N_2C(Ph)Me_2$	+5
$Me_2C(p-MePh)N_2(p-MePh)Me_2$	+4
$Me_3CON_2OCMe_3^a$	+4
$\mathrm{C_6H_{11}CO_2OCMe_3}$	+4

a n-Octane.

+4 to +5 cm³/mole are substantially smaller than those for the one-bond scission initiators given in Table IV (note that azoisobutyronitrile, AIBN, decomposes via two-bond scission). In further agreement, the values of ΔV^*_{obsd} for tert-butyl perbenzoate decomposition, a one-bond scission process (eq 7), are relatively large (+11 cm³/mole; cumene, solvent).9

$$C_6H_5CO_2OC(CH_3)_3 \longrightarrow C_6H_5CO_2 \cdot \cdot \cdot OC(CH_3)_3$$
 (7)

In the following sections more examples are presented which demonstrate that one-bond scission initiators give values of ΔV^*_{obsd} which are abnormally large. Data will also be presented which indicate that the actual volume change of activation for homolytic scission is generally +4 to +5 cm³/mole for single or multiple bond scission processes. For molar volumes between 150 to 250 cm³/mole, these values of ΔV^*_{obsd} indicate that homolytic scission transition states are on the order of 2 to 3% larger in volume than the groundstate initiators.

Polar Effects

Several two-bond scission peresters have given values of ΔV^*_{obsd} (cumene, solvent) substantially less than those obtained for other systems (Table V) and AIBN (Table IV). These include ring-substituted tert-butyl phenylperacetates ($\Delta V^*_{\rm obsd} < 2 \, {\rm cm}^3/{\rm mole}$)¹³ and tertbutyl perpivalate ($\Delta V^*_{\rm obsd}$ ca. +1 cm³/mole).¹⁴ The former group has long been thought to decompose via polar transition states (eq 8),5 and we believe that this

$$\begin{bmatrix} \delta^{+} & \delta^{-} \\ ArCH_{2} - CO_{2} - OCMe_{2} \end{bmatrix}^{\pm} \longrightarrow ArCH_{2} \cdot CO_{2} \cdot OCMe_{3} \quad (8)$$

is the explanation for the results. If ΔV^*_{obsd} reflects both molecular reorganization (ΔV^*_{bond}) and solvation $(\Delta V^*_{\text{solv}})$ (eq 9), small negative values of the solvation

(9) R. C. Neuman, Jr., and J. V. Behar, J. Amer. Chem. Soc., 91, 6024 (1969).

(10) R. C. Neuman, Jr., Intra-Sci. Chem. Rep., 3, 269 (1969).

(11) R. C. Neuman, Jr., and R. J. Bussey, J. Amer. Chem. Soc., 92, 2440 (1970).

(12) R. C. Neuman, Jr., G. D. Lockyer, Jr., and M. J. Amrich, Tetrahedron Lett., 1221 (1972).

(13) R. C. Neuman, Jr., and J. V. Behar, J. Org. Chem., 36, 654

(14) R. C. Neuman, Jr., and R. P. Pankratz, unpublished results.

⁽⁸⁾ R. C. Neuman, Jr., and J. V. Behar, Tetrahedron Lett., 3281 (1968).

$$\Delta V^*_{\text{obsd}} = \Delta V^*_{\text{bond}} + \Delta V^*_{\text{solv}} \tag{9}$$

term could effectively reduce a positive contribution of +4 to +5 cm³/mole for $\Delta V^*_{\rm bond}$ to the small positive values observed.¹³

Polar character in the decomposition transition state for *tert*-butyl perpivalate (eq 10; R = Me₃C) has been RCO₂OCMe₃ ->

$$\begin{bmatrix} \delta^{+} & \delta^{-} \\ R-CO_{2}-OCMe_{3} \end{bmatrix}^{\pm} \longrightarrow R \cdot CO_{2} \cdot OCMe_{3} \quad (10)$$

considered, but evidence has been weak.¹⁵ In contrast to its low value of +1 cm³/mole, $\Delta V^*_{\rm obsd}$ for decomposition of *tert*-butyl perisobutyrate (R = Me₂CH) is +3 cm³/mole, a value close to that expected when solvation effects are absent. All of the low values of $\Delta V^*_{\rm obsd}$ are found for peresters whose R groups (eq 10) are particularly good at stabilizing electron deficiency.

Cage Reactions

The pressure dependence of product ratios from decomposition of several two-bond scission initiators has permitted us to probe the relative effects of pressure on the rapid competitive reactions available to geminate radical pairs. These processes most commonly include combination, disproportionation, and separative diffusion.

Two cases in which only combination and diffusion compete are those shown in Schemes I and II. Partitioning of the initial cages from ring-substituted tertbutyl phenylperacetate decomposition gives ethers via combination, and benzyl radical products and tertbutyl alcohol subsequent to diffusion (Scheme I).¹⁶

Scheme I

$$X$$
 $CH_2OC(CH_3)_3$
 $\xrightarrow{k_c}$
 X
 $CH_2 \cdot CO_2 \cdot OC(CH_3)_3$
 $\xrightarrow{k_d}$
 $CH_3 \cdot COH_3 \cdot$

The pressure dependence of $k_{\rm e}/k_{\rm d}$ can be determined from the ether/alcohol product ratio. Similarly, $k_{\rm e}/k_{\rm d}$ for geminate *tert*-butoxy radicals from di-*tert*-butyl hyponitrite (DBH) (Scheme II) can be equated to the

Scheme II

$$(CH_3)_3CON = NOC(CH_3)_3$$

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

$$(CH_3)_3COOC(CH_3)_3 \xrightarrow{k_c} \overline{(CH_3)_3CO \cdot N_2 \cdot OC(CH_3)_3} \xrightarrow{k_d}$$

$$2(CH_3)_3COH$$

peroxide/alcohol ratio. $^{11,17-19}$ In both cases $k_{\rm e}/k_{\rm d}$ increases with pressure (Figure 1).

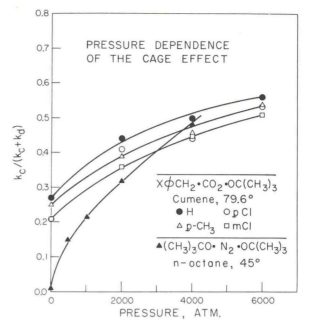


Figure 1.

We have suggested that radical combination should be pressure accelerated while diffusion is retarded. The qualitative trends in these results are thus consistent with expectation. However, in contrast we found that the "cage effect" from decomposition of azocumene (Scheme III) was relatively pressure in-

$\begin{array}{c} \textbf{Scheme III} \\ PhCMe_2N = NCMe_2Ph \\ \downarrow \\ \text{bicumyl} + \\ \text{cumene} + \\ \star \\ \textbf{Styrene} \end{array} \xrightarrow{k_c \ + \ k_a} \frac{\downarrow}{Ph\dot{C}Me_2 \ N_2 \cdot CMe_2Ph} \xrightarrow{k_d} 2Ph\dot{C}Me_2$

sensitive, although it showed an initial increase (Figure 2).²⁰ In this case combination of the cumyl radicals to form bicumyl (k_c) and disproportionation to form cumene and styrene (k_a) are competitive with diffusion (k_d) . These results prompted us to examine closely the available data, and we concluded that the pressure-induced increases in k_c/k_d for the phenylperacetates and DBH were smaller than might be expected just from the anticipated pressure retardation of k_d .¹⁶ In other words, in all of these cases combination (and disproportionation) seemed to be somewhat retarded by pressure.

We have rationalized this by pointing out that bimolecular combination and disproportionation reactions require that the inert gas molecules (nitrogen or carbon dioxide) evacuate the region between the radicals (Scheme IV). 16 Such a process is akin to diffusion and might be expected to be pressure retarded. The effect is most dramatic for the azocumene system

⁽¹⁵⁾ T. Koenig and R. Wolf, J. Amer. Chem. Soc., 91, 2574 (1969).
(16) R. C. Neuman, Jr., and J. V. Behar, J. Org. Chem., 36, 657 (1971).

⁽¹⁷⁾ H. Kiefer and T. Traylor, J. Amer. Chem. Soc., 89, 6667 (1967)

⁽¹⁸⁾ R. C. Neuman, Jr., and R. J. Bussey, *Tetrahedron Lett.*, 5859 (1968)

⁽¹⁹⁾ See also R. C. Neuman, Jr., J. Org. Chem., 37, 495 (1972).
(20) M. Amrich, Ph.D. Dissertation, University of California, Riverside, 1971.

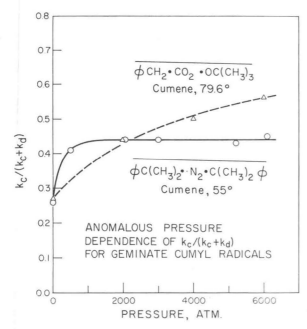
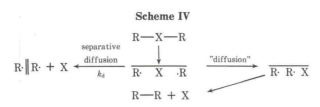


Figure 2.



(Scheme III) where the radicals are particularly bulky and might make escape of nitrogen especially difficult. This analysis suggests that pressure effects on cage return for one-bond scission initiators (e.g., $k_{-1}/(k+k_{\rm d})$; eq 4) could be substantially greater than had been predicted using the $k_{\rm c}/k_{\rm d}$ data from two-bond scission initiators.

Increasing pressure seems to favor disproportionation of geminate radicals over combination. The disproportionation to combination ratio (k_a/k_c) for cumyl radicals from azocumene is derived from the relative yields of cumene and bicumyl. Our results indicate that k_a/k_c for these radical pairs increases from about 0.1 at atmospheric pressure to about 0.5 at 6000 atm.²⁰ Similar results have been observed for α -cyanocyclohexyl radical pairs from azocyanocyclohexane (see below).²¹ It has been shown that the medium can have a dramatic effect on disproportionation/combination ratios,²² and we ascribe these changes to pressure effects on the rotational motions of radicals within the solvent cage.²³

Isomeric Vinyl Peresters

Most of our data for homolytic scission had been obtained from studies of two-bond scission initiators. To extend our knowledge of one-bond initiators we

(22) J. M. McBride, *ibid.*, 93, 6302 (1971).
(23) Recently we learned that similar conclusions have been reached by V. M. Zhulin and M. G. Gonikberg, *Izv. Akad. Nauk SSSR*, Ser. Khim., 2, 331 (1972).

undertook a kinetic and product study of a pair of isomeric tert-butyl vinyl peresters.²⁴ The results of this

study are of particular interest because of the mechanistic information they provided.

The values of ΔV^*_{obsd} for decomposition of the cis and trans peresters (cis, $+6.8 \pm 0.4 \text{ cm}^3/\text{mole}$; trans $+9.0 \pm 1.0 \text{ cm}^3/\text{mole}$) (cumene) were greater than those for two-bond scission initiators, thus supporting a one-bond scission mechanism with internal return (Scheme V).

Scheme V

$$\begin{array}{c} \text{vinyl perester} \\ & \stackrel{k_1 \downarrow \uparrow k_{-1}}{\longleftarrow} \\ \text{EtC} = \text{CCH}_2\text{Et} & \stackrel{k_e}{\longleftarrow} \text{EtCH} = \text{C} < \stackrel{\text{CO}_2}{\longleftarrow} \\ + \text{HOCMe}_3 \\ + \text{CO}_2 & \downarrow & \stackrel{k'_e}{\longleftarrow} \\ \text{EtCH} = \text{C} < \stackrel{\text{CO}_2}{\longleftarrow} \\ \text{CH}_2\text{Et} & + \text{OCMe}_3 & + \text{HOCMe}_3 \\ & + \text{CO}_2 & \\ \end{array}$$

The remainder of Scheme V was supported by product data. In particular the data indicated that the acetylene and allene were formed only from the initial cage.

We were surprised that the values of ΔV^*_{obsd} were smaller than those observed for other one-bond initiators and particularly that they were different for the two isomers. We concluded that this difference was due to a difference in reactivity of the first formed isomeric vinyl carboxyl radicals. All data suggested that the cis radical was more susceptible to cage bimolecular elimination of CO_2 to form the acetylene. A possible difference in k_{-1} could not be probed. The relatively low values of ΔV^*_{obsd} indicated that k_e and k_e' (processes competing with k_{-1}) were pressure accelerated.

An Activation Volume for Homolytic Scission

A study of the thermal decomposition of a ketenimine (eq 11) has provided evidence for our proposals about

magnitudes of initiator decomposition activation volumes. In the absence of radical scavengers, decomposition is described by the mechanism in Scheme VI where

Scheme VI

$$RR' \xrightarrow[k_{-1}]{k_1} \overline{R} \cdot \overline{R} \cdot \xrightarrow{k_d} RR + R \cdot + R \cdot$$

$$\downarrow k_d \qquad RH + R(-H)$$

⁽²¹⁾ R. C. Neuman, Jr., and M. Amrich, J. Amer. Chem. Soc., 94, 2730 (1972).

⁽²⁴⁾ R. C. Neuman, Jr., and G. D. Holmes, J. Amer. Chem. Soc., 93, 4242 (1971).

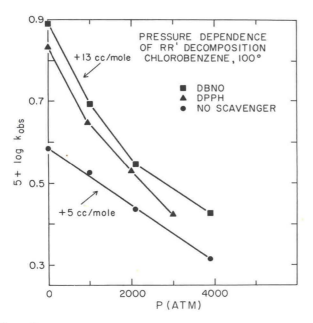


Figure 3.

RR' is the ketenimine, RR is the symmetrical dinitrile coupling product, and RH and R(-H) are the disproportionation products, cyanocyclohexane and cyanocyclohexene, respectively.²⁵

$$\begin{array}{c|c}
 & \text{NC} \\
\hline
\text{CN} \\
\text{RR}
\end{array}$$

$$\begin{array}{c|c}
 & \text{CN} \\
\text{RH}
\end{array}$$

$$\begin{array}{c|c}
 & \text{CN} \\
\text{R(-H)}
\end{array}$$

Because cyanoalkyl radicals are relatively stable, those which have separated by diffusion ultimately return to radical pairs. As a result, separative diffusion is not kinetically visible, and the observed rate of decomposition (and the observed decomposition activation volume) would *not* depend on separative diffusion (eq. 12), behavior substantially different from that

$$k_{\text{obsd}} = k_1 \left[\frac{k_{\text{o}} + k_{\text{a}}}{k_{-1} + k_{\text{c}} + k_{\text{a}}} \right]$$
 (12)

(eq 6) of other one-bond initiators previously discussed. However, in the presence of scavengers, diffusion would once again be a destructive process (eq 13) and k_d would

$$\overline{R} \cdot R \cdot \xrightarrow{kd} R \cdot + R \cdot \xrightarrow{2S} 2RS \tag{13}$$

return to the kinetic expression (eq 6 and 14). The

$$k_{\text{obsd}} = k_{\text{I}} \left[\frac{k_{\text{c}} + k_{\text{a}} + k_{\text{d}}}{k_{-1} + k_{\text{c}} + k_{\text{a}} + k_{\text{d}}} \right]$$
 (14)

experimental results (Figure 3) indicated that the pressure dependence of $k_{\rm d}$ had a big effect on $\Delta V^*_{\rm obsd}$. While this quantity in the absence of scavengers was on the order of +5 cm/mole, it increased to at least +13 cm³/mole in the presence of either diphenylpicrylhydrazyl (DPPH) or di-tert-butyl nitroxide (DBNO).

In the absence of scavengers, product data suggested that the pressure dependence of the rate constant ratio

(25) H. P. Waits and G. S. Hammond, J. Amer. Chem. Soc., 86, 1911 (1964). $(k_c + k_a)/(k_{-1} + k_c + k_a)$ was very small. Thus, under the no-scavenger conditions the observed activation volume (+5 cm³/mole) may be a good approximation to ΔV^* ₁ (eq 15), the activation volume for

$$\Delta V^*_{\text{obsd}} = \Delta V^*_1 + RT \, \delta \ln \left[1 + k_{-1}/(k_a + k_c) \right] / \delta P \approx \Delta V^*_1 \quad (15)$$

one-bond scission of the C-N bond in the ketenimine. Its approximate magnitude of +5 cm³/mole is suggestively similar to many of those for two-bond scission which we have observed.

The Viscosity Test

Our studies and results have complemented a growing literature describing the use of atmospheric pressure variation of solvent viscosity to probe free-radical reaction mechanisms. This "viscosity test" has proven to be valuable in demonstrating both the presence and extent of kinetically invisible cage return. Our conclusions have generally, but not always, been consistent with results of these studies. 13

One of the first systems studied by the "viscosity test" was p-nitrophenylazotriphenylmethane (NAT) and its unsubstituted isomer phenylazotriphenylmethane (PAT). The decomposition rates of both decreased with increasing solvent viscosity, and it was proposed that they decomposed via one-bond scission (Scheme VII) with substantial return (k_{-1}) . Before

Scheme VII

this study, little evidence was available supporting such a decomposition mode for azo systems. We have carried out pressure variation studies on both systems, and our data completely agree with these conclusions. The observed activation volumes determined in several solvents fall within the range of +15 to $+20~\rm cm^3/mole$, the largest reported values for homolytic scission to our knowledge. They seem to demand a substantial dependence of $k_{\rm obsd}$ on separative diffusion.

Cyclic Systems

The viscosity test cannot be applied to studies of homolytic scission of cyclic systems which might produce diradicals (Scheme VIII). Since reactions of the diradical do not include separative diffusion, the observed decomposition rate constant should be viscosity insensitive whether or not return (k_{-1}) occurs in competition

(26) See, for example, W. A. Pryor and W. K. Smith, ibid., 92, 5403 (1970).

Scheme VIII

$$A - B$$
 $\stackrel{k_1}{\underset{k_{-1}}{\longleftrightarrow}}$ $A \cdot \cdot B$ $\stackrel{k}{\longrightarrow}$

with the other reactions (k). However, the possibility that pressure effects on k_{obsd} might give clues concerning detailed mechanism (see eq 16) has led us to study a

$$\Delta V^*_{\text{obsd}} = \Delta V^*_1 + RT \, \delta \ln \left(1 + k_{-1}/k \right) / \delta P \quad (16)$$

series of cyclic azo compounds.

Kinetic and product studies of the six- through eightmembered ring systems shown below are in progress.²⁷

The values of $\Delta V^*_{\rm obsd}$ for decomposition leading to expulsion of nitrogen for the six- and eight-membered ring systems are +5 and +7 cm³/mole, respectively. Together with product data, the results suggest that mechanistic differences may be present which can be probed using pressure studies.

Summary

Effects of pressure on the decomposition rates and (27) R. C. Neuman, Jr., and E. W. Ertley, *Tetrahedron Lett.*, 1225 (1972).

products of radical initiators have provided detailed mechanistic information about these systems. Decomposition activation volumes for homolytic scission appear to be ca. +4 to +5 cm³/mole. Larger values indicate the presence of cage return regenerating the initiator, and smaller values can reflect polar effects in the decomposition transition state.

Cage reactions of radical pairs such as combination and disproportionation are generally favored over separative diffusion by increasing pressure. However, in spite of this, these cage bimolecular processes (combination and disproportionation) appear to be retarded by pressure because of its effect on the prerequisite rotational diffusion processes within the initial cages.

Pressure studies complement the use of solvent viscosity as a probe of initiator decomposition mechanism and provide information in cases where the latter technique cannot be used. Since diffusion rates can be altered by pressure without changes in temperature or medium, data derived from these studies may be of use in probing microscopic features of solvent structure and its interaction with solutes.

The work reported here was carried out by my able collaborators Drs. Michael Amrich, Joseph Behar, Robert Bussey, Gary Holmes, and Messrs. Ernest Ertley, George Lockyer, and Richard Pankratz. We thank the National Science Foundation for its continuing support of these studies. This account was written during the author's tenure as an NIH Special Research Fellow at Princeton University, 1971–1972.

