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# Cyclopropylcarbinyl Radicals in the Photochemistry of $\beta, \gamma$ -Cyclopropyl Ketones

#### Ioannis M. Takakis and William C. Agosta\*

Contribution from the Laboratories of The Rockefeller University, New York, New York 10021. Received October 3, 1978

Abstract: Photolysis of tricyclic ketones 8b-d, 9b, and 10 leads to the products collected in Table I. These results, along with earlier findings with related ketones 6 and 7, are interpreted as strong evidence for the stepwise mechanism of eq 1, in which opening of the three-membered ring occurs by way of  $\alpha$ -cleavage, rearrangement of the initial cyclopropylcarbinyl radical to a homoallyl radical under stereoelectronic control, and then product formation.

One of the principal modes of photochemical reaction of  $\beta,\gamma$ -cyclopropyl ketones can be explained as an  $\alpha$ -cleavage, rearrangement of the cyclopropylcarbinyl radical to a homoallyl radical, and then product formation through disproportionation, coupling, or some more complex transformation of the resulting radical pair or biradical (eq 1). This intuitively



reasonable, stepwise mechanism, first advanced over a decade ago, <sup>1</sup> satisfactorily accounts for a variety of rearrangements,<sup>2,3</sup> although there is at least one observation suggesting that some refinement of it may be necessary. This is the report<sup>3</sup> that photolysis of both cis- and trans-4-caranone (1) leads selectively to products explainable by way of eq 1, as particularized in eq 2. There is no concomitant epimerization of the methyl group, and less than 1% of product attributable to  $\alpha$ -cleavage toward the methyl group (as 3) is found. This result contrasts with the behavior of typical 2-alkylcyclohexanones, which



undergo  $\alpha$ -cleavage predominantly or solely on the more substituted side; the products from 2-methylcyclohexanone (4), for example, arise only from biradical 5.4 Thus, if the photochemistry of the 4-caranones (1) follows eq 1, it is necessary to specify that the  $\beta$ ,  $\gamma$ -cyclopropane ring controls the site of  $\alpha$ -cleavage. As originally suggested,<sup>3</sup> the cyclopropane ring appears to weaken the appropriate bond  $\alpha$  to the carbonyl group by conjugative or inductive effects. Although the authors did not carry the argument so far, at the extreme this observation might mean that no discrete cyclopropylcarbinyl intermediate is involved and that  $\alpha$ -cleavage and fragmentation are concerted. To our knowledge there is no other evidence directly concerning this possible alternative to eq 1. However, there is evidence that, in the gas phase at least, rupture of the three-membered ring of cyclopropylacetone occurs without prior or concomitant  $\alpha$ -cleavage.<sup>1</sup>

If photochemical reactions of  $\beta$ , $\gamma$ -cyclopropyl ketones do involve a discrete cyclopropylcarbinyl radical intermediate, its rearrangement should be under stereoelectronic control, with preferential cleavage of the  $\beta$ , $\gamma$ -cyclopropane bond that can better overlap the orbital on C( $\alpha$ ) containing the odd electron.<sup>5,6</sup> The studies noted above,<sup>1-3</sup> however, dealt only with ketones that would lead to cyclopropylcarbinyl radicals in which rotation about the original C( $\alpha$ )-C( $\beta$ ) bond is unconstrained, so that the question of stereoelectronic control could not be answered. As we discuss below, a previous brief study demonstrated that the photochemistry of tricyclic ketones **6** and **7** is in fact consistent with such stereoelectronic



control.<sup>7</sup> We have now studied the photochemistry of several additional tricyclic  $\beta$ , $\gamma$ -cyclopropyl ketones related to **6** and **7**. This series of compounds has the advantage that, if discrete cyclopropylcarbinyl radicals are involved, the site of stereo-electronically controlled cleavage can be predicted and is expected to change along the series.

The entire set of compounds comprises seven tricyclic ketones: **8a-d**, in which the bridge of methylene groups ostensibly



uninvolved in reaction is progressively increased from one to four members; **9a,b**, in which the cyclopropane stereochemistry of **8a,b** is inverted; and **10**, in which the carbonyl group of **8c** has been formally transposed to the three-carbon bridge. Application of eq 1 to these compounds leads to eq 3, and we note



that the site of cyclopropylcarbinyl rupture (a or b) in biradical 11 should be largely a function of the value of n. Both from previous work with related radicals<sup>6</sup> and from examination of molecular models, one can conclude that, when n is 1 or 2, the

orbital containing the odd electron can efficiently overlap only with external bond a. However, when n is 3, the flexibility of 11 is materially increased, and conformations permitting orbital overlap of the radical center with either bond a or bond b are reasonable. This will also be true for larger values of n, of course, but study of space-filling models suggests that, when n is 4, the molecular motions necessary to achieve these favorable conformations may be hindered by transannular interactions of the hydrogen atoms in the eight-membered ring. These predictions are summarized in structures 11a-d. Several



points are noteworthy: (1) The conformational changes that favor facile cleavage at a and/or b can take place only in the bicyclic biradicals 11 and cannot occur in the tricyclic starting ketones 8-10. Thus if  $\alpha$ -cleavage is not the first step in these reactions, no simple relationship between n and the site of opening of the cyclopropane is obvious. (2) The stereochemistry of the acyl side chain relative to the three-membered ring should be irrelevant, so that isomers from series 8 and 9 should fragment similarly. (3) Lengthening the acyl side chain should be irrelevant, so that 10 should open like 9b. (4) Ketones 6 and 7 are 8a and 9a, respectively, and their photochemistry follows these predictions. The only pertinent products from their photolysis are the bicyclooctenones cis- and trans-13, respectively, and these are regarded as arising through cleavage a in **11a** and subsequent coupling of homoallyl biradical **12**.<sup>7</sup> Guided by these considerations we turned to examination of the photochemistry of the remaining ketones with a particular interest in any difference between the behavior of 8b and 8c.

Ketones **8b** and **9b** are well-known and were prepared from **14**.<sup>8</sup> Hydrogenation gave **9b**, as previously reported, <sup>9</sup> while Wolff-Kishner reduction<sup>10</sup> furnished hydrocarbon **15**, which has been prepared in other ways in the past;<sup>9,11</sup> hydroboration<sup>9</sup> and subsequent oxidation<sup>9,12</sup> then gave **8b**. Properties of the products and intermediates were in agreement with those already on record.<sup>8,9,11,12</sup> Ketone **10** was available through hydrogenation of the known unsaturated ketone **16**,<sup>13</sup> and we



have described elsewhere the preparation of  $8c^{13}$  and  $8d.^{14}$ 

Solutions of **8b-d**, **9b**, and **10** (0.025 M) in benzene containing 3.0% methanol (v/v, 0.74 M) were irradiated through Pyrex ( $\lambda > 2800$  Å) using a 450-W medium-pressure mercury lamp. Under these conditions ketene products are trapped as the related photochemically unreactive methyl esters, and all isolated esters are presumed to have arisen in this fashion. Products and unreacted starting ketones were isolated by preparative vapor phase chromatography (VPC). The results are presented in Table I. Yields are based on converted starting material and generally were determined by calibrated VPC



Table I. Products of Photolysis of Ketones 8b-d, 9b, and 10

<sup>a</sup> Characterization incomplete; see text.

measurements; they varied somewhat from run to run, and those given are typical.

With the two exceptions noted, the products listed were completely characterized. For compounds not discussed further here, structural assignments rest on spectroscopic data given in the Experimental Section. The structures of 17 and 19 were substantiated by hydrogenation to yield *cis*- and *trans*-2hydrindanone (34 and 35), respectively, which had IR and



NMR spectra that agreed with published values.<sup>15</sup> Similarly, reduction of 23 furnished 36, which was identified by comparison of melting points of the derived semicarbazone and 2,4-dinitrophenylhydrazone with those reported,<sup>16</sup> and reduction of 28 gave trans-2-decalone (37), the properties of which matched those of an authentic sample. Cycloheptatriene (22) was identical with the commercially available material. Hydrocarbon 26 was indistinguishable from samples prepared from either 38<sup>17</sup> or 39<sup>18</sup> by base-catalyzed equilibration, <sup>18</sup> and cis, cis-1,4-cyclooctadiene (27) was identical with an authentic sample prepared as described by Moon and Ganz.<sup>19</sup> The bicyclodecene 31 appears to be previously unknown but was identical with the product of Simmons-Smith monocyclopropanation<sup>20</sup> of *cis, cis*-1,4-cyclononadiene.<sup>21,22</sup> The structural assignment for the corresponding trans olefin 32 is tentative and is based only on the similarity in IR spectra and VPC retention times of 31 and 32 and the fact that formation of this product is reasonable (see below). The assignment for **30** is

fully supported by IR and NMR spectra, but the small amount available, along with persistent impurities, prevented our obtaining an analytically pure sample. The incomplete characterization of **30** and **32** in no way affects the analysis of our results given below.

Considering first the isomeric ketones 8b and 9b, we note that products 17-20, in which the cyclopropane ring has opened, can all be accounted for by  $\alpha$ -cleavage to 11b, scission of bond *a* to form 40, and either collapse to 17 and 19 or dis-



proportionation to 18 and 20. The latter process involves a different 1,4 hydrogen shift in each case, a result indicating that steric and electronic factors favor the cis transition states 41 (from 8b) and 42 (from 9b). The aldehyde 21 is explained as the product of competitive hydrogen transfer in 11b without opening of the three-membered ring. Since a separate experiment showed that irradiation of 21 efficiently yields cycloheptatriene (22), the small amount of 22 formed on photolysis of 9b is believed to arise in this way. The pertinent reactions of 8b and 9b thus involve only cleavage of bond a, in agreement with stereoelectronic control in opening of 11b.

From our earlier discussion the key compound in this study should be 8c, and it is obvious from Table I that the behavior of this substrate differs from that of its lower homologues. Ketone 23 and ester 24 are analogous to 17 and 18 and can be accounted for by collapse or disproportionation of biradical 43. However, the isolation of two products with eight-membered rings strongly implies concomitant cleavage of 11c at bond b to furnish the isomeric species 44. Straightforward



fragmentation of this 1,4 biradical could then vield diene 27. while decarbonylation and ring closure could lead to 26. From molecular models it is clear that generation of the observed cis disubstituted double bond of 27 and cis disubstituted cyclopropane of 26 should be energetically preferred in these reactions of 44. Since hydrocarbon 26 is the dominant product from **8c**, its derivation by a pathway requiring scission of bond b is particularly significant. However, the mechanism suggested for its formation is unusual (but not unique<sup>2a</sup>) in requiring the uneconomical opening of one cyclopropane ring followed by the closure of another. Also, any mechanism leading to 26 involves a decarbonylation that is remarkably efficient for a photochemical reaction in solution at  $\sim 30$  °C.<sup>2a</sup> For these reasons we investigated the origin of 26 through preparation and photolysis of deuterated ketones 46-48. It is convenient to discuss this work before considering the remaining entries in Table I.



Treatment of **8c** with sodium carbonate in hot ethanol-O-d yielded the dideuterated ketone **46**. The two monodeuterated isomers were available from the related olefin **49**.<sup>13</sup> Deuter-ioboration<sup>22</sup> gave a mixture of the epimeric alcohols **50** and **51**, which could be separated by preparative VPC of their trimethylsilyl ethers and identified by comparison with their known<sup>13</sup> undeuterated analogues. The stereochemistry of the deuterium substituents was confirmed through lanthanide-shifted NMR spectra of the deuterated and undeuterated species. Oxidation of **50** and **51** with ruthenium tetroxide<sup>24</sup> in carbon tetrachloride solution then gave **47** and **48**.

Photolysis of ketone **46** furnished a deuterated hydrocarbon that could be assigned structure **52** from its NMR spectrum. This result substantiates the general features of the mechanism suggested above, which requires that the new cyclopropane methylene group of **26** (or **52**) arise from the  $\alpha$ -methylene group of **8c** (or **46**). Thus scission of bond b in this transformation is assured.

We were interested in the photolysis of 47 and 48 because of a recent suggestion made by Weiss, Haslanger, and Lawton. On the basis of studies in another system, in which a 1,6 acyl alkyl biradical undergoes stereospecific decarbonylation with carbon-carbon bond formation, these investigators postulated that in conformationally favorable cases such decarbonylation of the acyl radical may be facilitated through backside assistance by the alkyl radical.<sup>25</sup> Assistance of this sort in decarbonylation of 44 could account for the efficiency of this reaction and would presumably lead to generation of 26 with inversion at the methylene carbon atom. However, photolysis of 47 and 48 gave in each case a nearly 1:1 mixture, as indicated by NMR measurements, of the deuterated hydrocarbons 53 and 54, indicating a virtually complete loss of stereochemistry at this center. Thus there is no stereochemical evidence for assisted decarbonylation of 8c, and stepwise reaction by way of 45 cannot be dismissed.

An alternative route to 26 and 27 entails collapse of 44 to the cyclobutanone 55, followed by secondary photolysis. This



would offer a convenient explanation for the efficient decarbonylation, since cyclobutanones undergo exceptionally easy loss of carbon monoxide in solution.<sup>26</sup> We consider this pathway unlikely, however, for several reasons. (1) We found no evidence for **55** in the early stage of reaction of **8c**. (2) The decarbonylation of cyclobutanones is known to be a stereospecific reaction,<sup>26</sup> but our findings with **47** and **48** indicate a loss of stereochemistry in formation of **26**. (3) The photochemistry of the closely related *cis*- and *trans*-bicyclodecanones **56** is on record.<sup>27</sup> Not only does each ketone give cyclopropane and olefin with predominant retention of stereochemistry, but much more olefin than cyclopropane is formed from *cis*-**56**, both in benzene (70:30) and in methanol (88:12). In contrast, from photolysis of **8c** in benzene-methanol the ratio of olefin **27** to cyclopropane **26** is 7:93.

Thus a convincing rationalization for the decarbonylation of **8c** awaits further investigation, but we may conclude for present purposes that this reaction does involve cleavage at bond b. Judged by the products, opening at b is twice as important as opening at a in **8c**. We considered the possibility that this change on passing from **8b** and **9b** to **8c** was attributable simply to the increased molecular flexibility of the tricyclo[3.3.2.0<sup>2,4</sup>]decane carbon skeleton of **8c** relative to that of the lower homologues. This possibility led us to investigation of the photochemistry of **10**, a ketone that has the same carbon skeleton as **8c**. As we pointed out above,  $\alpha$ -cleavage in **10** would furnish a species (see **57**) which should behave like **11b**. In fact



the photochemistry of 10 parallels that of 8b and 9b, and both 28 and 29 are products of the opening of bond a. It is interesting to note that, although 9b and 10 have the same stereochemistry, 1,5 hydrogen transfer in 58 leads to ketene (ester) and therefore occurs in the direction opposite to the 1,4 transfer in 9b (see 42).

Finally, we prepared and photolyzed 8d in the hope of finding reactivity similar to that of 8c. This effort met with only limited success. The hydrocarbon products 31 and 32 are indeed analogous to their lower homologue 26, and thus their presence presumably does signal opening of bond b in 11d. However, the striking result with 8d is that opening of the cyclopropane ring is such a minor process. Under our standard conditions photolysis of 8d is considerably slower than that of the lower members of the series, and the major product 33 is a compound in which the original three-membered ring remains intact. Essentially this same distribution of products resulted from photolysis of 8d in refluxing benzene-methanol or on irradiation through quartz with either benzene-methanol or methanol as solvent. We ascribe this divergent behavior of 8d to hindered rotation in 11d due to nonbonded transannular interactions that should lead to preferential collapse of the biradical back to starting ketone. In models these appear to interfere severely with the motions necessary to attain any of the conformations favoring opening of a or b with generation of either a cis- or a trans-homoallyl radical. As a consequence there is reduced fragmentation of the three-membered ring, and simple disproportionation becomes the predominant isomerization of 11d.

In summary the photochemistry of tricyclic ketones 6, 7, 8b-d, 9b, and 10 is in accord with the involvement of discrete cyclopropylcarbinyl radicals that open under stereoelectronic control. This single assumption provides a straightforward explanation for the relatively simple behavior of 6, 7, 8b, 9b, and 10, a basis for the otherwise unexpectedly complex photochemistry of 8c, and a rationale for the failure of 8d to undergo significant cleavage of the cyclopropane ring at all. These results then provide strong support for the stepwise mechanism of eq 1.

#### **Experimental Section**

Materials and Equipment. These have been previously described.<sup>13</sup> In the present work, the following VPC columns were used: A, 29 g of 20% DEGS, 9.7 ft; B, 16 g of 20% Carbowax 20M, 5 ft. Columns were constructed of standard aluminum tubing having  $\frac{3}{16}$ -in. i.d., and Chromosorb P, mesh 60/80, was used as the solid support. An ultra-

violet (UV) spectrum was recorded on a Cary Model 14 PM spectrophotometer. Infrared (IR) spectra were obtained on a Perkin-Elmer Model 621 (PE-621), or on a Perkin-Elmer Model 237B grating infrared spectrophotometer. Exceptions to the above are noted.

General Procedure for Photolysis and VPC Analysis of the Photolysates. Photolyses were carried out in benzene (distilled, J. T. Baker) solution containing 3.0% (by volume) methanol through a double Pyrex filter ( $\lambda > 2800$  Å) using a 450-W Hanovia lamp. Details of the procedure are described elsewhere.<sup>28</sup> Yields for the products from **8b,c** and **9b** were ascertained by the internal standard technique. For those from **10** VPC calibration curves were constructed, and yields from **8d** were determined by collecting and weighing VPC fractions. All yields are based on converted starting material. Exceptions to these procedures are noted.

endo-Tricyclo[3.3.2.0<sup>2,4</sup>]nonan-6-one (8b). Ketone 14<sup>8</sup> (4.86 g, 36.2 mmol) was treated with 97% hydrazine (21.4 g) and KOH (28.5 g) in diethylene glycol (86 mL) according to a modified Wolff-Kishner procedure<sup>10</sup> to give 4.1 g (34 mmol) of endo-tricyclo[3.2.2.0<sup>2,4</sup>]non-6-ene (15). Crude 15 was hydroborated with disiamylborane (34 mmol) in THF following a standard procedure<sup>23</sup> to obtain 3.2 g of endo-tricyclo[3.2.2.0<sup>2,4</sup>]nonan-exo-6-ol. A sample purified by VPC on column B gave mp 166–168 °C (sealed tube) (lit.<sup>9</sup> 165–168 °C). Oxidation<sup>29</sup> and purification by VPC on column A gave 2.93 g of 8b as a white solid (59% based on 14). Analytical samples of 8b, 14, and 15 had the same physical and spectroscopic properties as those reported.<sup>8,9,11,12</sup>

**exo-Tricyclo[3.3.2.0<sup>2,4</sup>]decan-6-one** (10). A mixture of *exo*-tricyclo[3.3.2.0<sup>2,4</sup>]dec-9-en-6-one (16,<sup>13</sup> 476 mg, 3.21 mmol) and 5% Pd/C (77 mg) in methanol (20 mL) was hydrogenated at atmospheric pressure. Purification by VPC on column B gave 397 mg (82%) of a white solid: mp 116–117 °C (sealed tube); IR (PE-621) 3389 (w), 3079 (w), 3012 (m), 2943 (s), 2926 (s), 2885 (m), 2864 (m), 1704 (s), 1476 (m), 1454 (m), 1427 (m), 1285 (w), 1177 (m), 1160 (m), 1105 (w), 1026 (m), 1014 (w) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  0.36–1.28 (m, 4 H), 1.28–2.06 with major absorption at 1.56 (m, 6 H), 2.06–2.79 (m, 4 H).

Anal. Calcd for C<sub>10</sub>H<sub>14</sub>O: C, 79.95; H, 9.39. Found: C, 80.19; H, 9.55.

cis-(Z)-Bicyclo[7.1.0]dec-3-ene (31). cis,cis-1,4-Cyclononadiene (62 mg, 0.50 mmol)<sup>21</sup> was cyclopropanated for 18 h with diiodomethane (405 mg, 3 equiv) and zinc-copper couple (198 mg, 6 equiv) in the presence of a catalytic amount of iodine in ether (10 mL).<sup>9,13</sup> Preparative VPC on column A (75 °C, 78 mL/min) gave starting diene (22 min, 16 mg, 74% conversion) and a colorless oil identified as **31** (36 min, 18 mg, 35% based on converted diene): IR 3050 (w), 3000 (m), 2985 (m), 2955 (m), 2930 (s), 2860 (m), 1470 (m), 1435 (w), 1270 (w), 1015 (w), 970 (w), 840 (w), 715 (w), 690 (w) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  –0.22 (m, 1 H), 0.20–2.80 (m, 13 H), 4.93–5.87 (m, 2 H).

Anal. Calcd for  $C_{10}H_{16}$ : C, 88.16; H, 11.84. Found: C, 88.24; H, 11.77.

A biscyclopropanated product was also isolated and tentatively assigned as  $1\alpha,3\beta,5\beta,10\alpha$ -tricyclo[8.1.0.0<sup>3,5</sup>]undecane (57 min, 6.4 mg, 11% based on converted diene): IR 3055 (m), 2985 (s), 2950 (m), 2930 (s), 2850 (m), 1470 (m), 1440 (w), 1310 (w), 1020 (m), 990 (w), 840 (m) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  -0.38 (m, 2 H), 0.083-1.22 with peak maximum of 0.68 (m, 9 H), 1.22-2.45 with peak maximum at 1.63 (m, 7 H).

Anal. Calcd for  $C_{11}H_{18}$ : C, 87.92; H, 12.08. Found: C, 87.92; H, 12.06.

**Deuterated Ketone 46.** A solution of **8c** (207 mg, 1.38 mmol), anhydrous sodium carbonate (1.46 g, 13.8 mmol), and ethanol-*O-d* was heated at reflux under nitrogen for 43 h. After workup, the crude product was purified by VPC on column B to give 127 mg (60%) of **46.** The NMR spectrum (220 MHz) indicated 79  $\pm$  3% of two deuterium atoms at C(10).

exo-Tricyclo[3.3.2. $0^{2,4}$ ]decan-anti-9-ol-anti-10-d and exo-Tricyclo[3.3.2. $0^{2,4}$ ]decan-syn-9-ol-syn-10-d (50 and 51). Deuterioboration of 49 (478 mg, 3.56 mmol)<sup>13</sup> was accomplished with B<sub>2</sub>D<sub>6</sub> as previously described<sup>13</sup> for hydroboration of this olefin to give a quantitative yield of a white solid identified as a mixture of 50 (80%) and 51 (20%) (after separation via their trimethylsilyl ethers<sup>13</sup>) by comparison of the NMR and IR spectra of each with those of the protio analogues.<sup>13</sup> The stereochemistry of deuterium in each alcohol was confirmed by comparison of their Eu(f0)<sub>3</sub><sup>30</sup> shifted NMR spectra (60 MHz) with those of the undeuterated alcohols. exo-Tricyclo[3.3.2.0<sup>2,4</sup>]decan-9-one-anti-10-d and -syn-10-d (47 and 48). Oxidation of 50 and 51 with ruthenium tetroxide in CCl<sub>4</sub> solution<sup>24</sup> (10 min) followed by VPC purification on column B gave these desired ketones 47 and 48, respectively. Photolysis of endo-Tricyclo[3.2.2.0<sup>2,4</sup>]nonan-6-one (8b). Irradiation

Photolysis of endo-Tricyclo[3.2.2.0<sup>2,4</sup>]nonan-6-one (8b). Irradiation of 8b (197 mg, 1.45 mmol, 0.0290 M, 50.0 mL solution) for 8.00 h followed by VPC analysis on column A (120 °C, 69 mL/min) and using o-methylacetophenone (100 min) as internal area standard indicated some starting material (159 min, 90% conversion) and two products (80%) identified as 17 and 18.

Methyl *cis*-2-methylcyclohex-3-en-1-acetate (**18**, 65 min, 17%): IR (PE-621) 3021 (m), 2960 (s), 2929 (s), 2876 (m), 2842 (m), i738 (s), 1648 (w), 1435 (s), 1276 (s), 1254 (m), 1199 (m), 1172 (s), 1147 (s), 701 (w), 672 (w), 666 (w) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  0.81 (d, *J* = 6.5 Hz, 3 H), 0.95-2.35 with major absorptions at 1.42, 2.07 (m, 8 H), 3.43 (s, 3 H), 5.24 (m, 2 H). Irradiation at  $\delta$  2.00 caused collapse of d into br s.

Anal. Calcd for  $C_{10}H_{16}O_2$ : C, 71.39; H, 9.59. Found: C, 71.22; H, 9.49.

*cis*-Bicyclo[4.3.0]non-2-en-8-one (**17**, 149 min, 63%): IR (PE-621) 3470 (w), 3024 (m), 2925 (s), 2860 (m), 2843 (m), 1742 (s), 1650 (w), 1406 (m), 1161 (m), 1146 (m), 700 (m), 666 (w), 647 (w) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  0.81–2.88 (m, 10 H), 5.37 (m, 2 H).

Anal. Calcd for  $C_9H_{12}O$ : C, 79.37; H, 8.88. Found: C, 79.25; H, 8.89.

Hydrogenation of 17 (31 mg, 0.23 mmol) was carried out as described below for 19. VPC purification gave 19 mg (61%) of a colorless liquid the infrared and NMR spectra of which were identical with those of *cis*-bicyclo[4.3.0]nonan-8-one (34).<sup>15</sup>

**Photolysis of endo-Tricyclo**[3.3.2.0<sup>2,4</sup>]decan-9-one (8c). Ketone 8c (191 mg, 1.27 mmol, 0.0254 M, 50.0 mL solution)<sup>13</sup> was irradiated for 24.0 h. VPC analysis on column A (initial column temperature 65 °C, raised to 135 °C after 51 min, 69 mL/min) using *p*-xylene (34 min) and valerophenone (167 min) as internal area standards indicated some starting 8c (184 min, 88% conversion) and seven additional fractions (76%). Infrared and NMR spectroscopy indicated that two of these fractions with retention times of 112 (2.8%) and 119 (3.4%) min were mixtures of two or more products each. The remaining five fractions were identified as 23-27.

1,4-Cyclooctadiene (27, 23 min, 3.2%): This was identical in all respects with an authentic sample prepared by the method of Moon and Ganz.<sup>19</sup>

cis-(Z)-Bicyclo[6.1.0]non-3-ene (26, 43 min, 40%): This was identical in all respects with an authentic sample prepared by equilibration of 38<sup>17</sup> or by equilibration of 39 with potassium *tert*-butoxide in Me<sub>2</sub>SO.<sup>18</sup>

Anal. Calcd for C<sub>9</sub>H<sub>14</sub>: C, 88.45; H, 11.55. Found: C, 88.31; H, 11.38.

Methyl *cis*-2-methylcyclohept-3-en-1-acetate (**24**, 105 min, 4.7%): IR (PE-621) 3018 (m), 2954 (m), 2929 (s), 2858 (m), 1740 (s), 1648 (w), 1450 (m), 1438 (m), 1259 (m), 1195 (m), 1159 (m), 718 (w), 678 (w) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  1.02 (d, J = 7 Hz, 3 H), 1.15–2.88 with major absorptions at 1.63, 2.17 (m, 10 H), 3.60 (s, 3 H), 5.02–5.98 (m, 2 H).

Anal. Calcd for  $C_{11}H_{18}O_2$ : C, 72.49; H, 9.96. Found: C, 72.70; H, 10.00.

Methyl *cis-endo*-bicyclo[5.1.0]octan-2-acetate (**25**, 132 min, 3.0%): IR (PE-621) 3073 (w), 2994 (m), 2919 (s), 2850 (m), 1740 (s), 1467 (m), 1450 (m), 1435 (m), 1353 (m), 1288 (m), 1273 (m), 1255 (m), 1243 (m), 1213 (m), 1192 (m), 1178 (m), 1148 (m), 1024 (m) cm<sup>-1</sup>; NMR (60 MHz, CHCl<sub>3</sub> as internal reference)  $\delta$  0–2.62 with major absorptions at 0.35, 0.98, 1.60, 2.35 (m, 15 H), 3.68 (s, 3 H).

Anal. Calcd for C<sub>11</sub>H<sub>18</sub>O<sub>2</sub>: C, 72.49; H, 9.96. Found: C, 72.40; H, 10.00.

cis-Bicyclo[5.3.0]dec-2-en-9-one (23, 196 min, 19%): IR (PE-621) 3473 (w), 3014 (m), 2928 (s), 2869 (m), 1748 (s), 1450 (m), 1410 (m), 1154 (m), 716 (w), 688 (m) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  1.01-2.75 with major absorptions at 1.71, 2.15 (m, 11 H), 3.08 (m, 1 H), 5.48 (m, 2 H). Irradiation at  $\delta$  3.08 simplified m at  $\delta$  5.48.

Anal. Calcd for C<sub>10</sub>H<sub>14</sub>O: C, 79.95; H, 9.39. Found: C, 79.75; H, 9.34.

Hydrogenation of 23 (85 mg, 0.57 mmol) as for 19 afforded 47 mg (55%) of a colorless liquid identified as 36: semicarbazone, mp 197.5–198.5 °C dec (lit.<sup>16</sup> 198–199 °C); 2,4-dinitrophenylhydrazone, mp 122–123 °C (lit.<sup>16</sup> 125–126 °C); IR (PE-621) 3465 (w), 2920 (s), 2854 (s), 1742 (s), 1460 (m), 1454 (m), 1442 (m), 1404 (m), 1265

(w), 1240 (w), 1212 (w), 1163 (m)  $cm^{-1}$ .

Photolysis of Ketone 46. The dideuterated ketone 46 prepared above was irradiated (94 mg, 0.62 mmol, 0.025 M) as described for 8c. The major product was isolated as before and identified as 52. The NMR spectrum (220 MHz, CHCl<sub>3</sub> as internal standard) showed the following differences from that of 26:  $\delta$  -0.084, 0.1 H rather than 1.0 H; 0.60, 1.2 H rather than 2.0 H.

Photolysis of Ketones 47 and 48. The monodeuterated ketone 47 (65 mg, 0.43 mmol, 0.017 M) was irradiated as described for 8c. The major product was identified as a mixture of 53 and 54. The NMR spectrum (220 MHz, CHCl<sub>3</sub> as internal standard) showed the following differences from that of 26:  $\delta$  -0.084, 0.59 H rather than 1.0 H; 0.60, 1.4 H rather than 2.0 H. Similar results were obtained with 48.

Photolysis of exo-Tricyclo[4.3.2.0<sup>7,9</sup>]undecan-10-one (8d). Ketone 8d (86 mg, 0.52 mmol, 0.021 M, 25.0 mL solution) was irradiated for 41.0 h. Preparative VPC of the concentrated photolysate on column A (initial column temperature 70 °C, raised to 150 °C after 46 min, 78 mL/min) afforded starting 8d (129 min, 36 mg, 58% conversion) and three products (50 mg, 84% based on unrecovered 8d) which were identified as 31-33.

trans-Bicyclo[7.1.0]dec-3-ene (32, 35 min, ~1 mg, ~2%): IR 3050 (w), 3005 (m), 2970 (m), 2920 (s), 2845 (m), 1455 (m), 1440 (m), 1100 (m), 1090 (m), 1070 (m), 1020 (w), 900 (w), 690 (m) cm<sup>-1</sup>. This identification is tentative; see text.

cis-(Z)-Bicyclo[7.1.0]dec-3-ene (31, 43 min, ~1 mg, ~2%): This was identical in all respects with an authentic sample prepared as described above

Methyl cis-bicyclo[6.1.0]nonan-2-acetate 33, 91 min, 48 mg, 80%): IR 3050 (w), 2980 (m), 2915 (s), 2855 (m), 2840 (m), 1740 (s), 1470 (m), 1450 (m), 1440 (m), 1430 (m), 1350 (w), 1280 (m), 1255 (m), 1245 (m), 1185 (m), 1155 (s), 1130 (m), 1020 (m), 985 (w) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  -0.22 (m, 1 H), 0.23-2.80 with maxima at 1.57, 2.28 (m, 16 H), 3.62 (s, 3 H).

Anal. Calcd for C12H20O2: C, 73.43; H, 10.27. Found: C, 73.37; H. 10.21.

Photolysis of 8d under the following modified conditions gave essentially the same distribution of products as above: (1) Ketone 8d (27.2 mg) in 10.0 mL of benzene-3% methanol at reflux for 21.1 h. (2) Ketone 8d (11.4 mg) in 10 mL of benzene-3% methanol through quartz for 17.4 h (100% conversion). (3) Ketone 8d (22.9 mg) in 12.0 mL of methanol through quartz for 5.0 h.

Photolysis of exo-Tricyclo[3.2.2.0<sup>2,4</sup>]nonan-6-one (9b). Ketone 9b (197 mg, 1.45 mmol, 0.0290 M, 50.0 mL solution) was irradiated for 5.00 h. VPC analysis on column A (initial column temperature 65 °C, raised to 120 °C after 40 min, and raised again to 150 °C after 156 min, 72 mL/min), using p-xylene (33 min) and o-methylacetophenone (133 min) as internal area standards, indicated some starting 9b (179 min, 96% conversion) and four products (95%) which were identified as 19-22.

Cycloheptatriene (22, 22 min, 6.7%) was identical in all respects with an authentic sample of cycloheptatriene (Aldrich Chemical Co.).

2-Methylenecyclohex-3-en-1-acetaldehyde (20, 113 min, 23%): UV (absolute EtOH)  $\lambda_{max}$  ( $\epsilon$ ) 231 nm (27 300); IR (PE-621) 3080 (w), 3030 (m), 2927 (s), 2862 (m), 2840 (m), 2716 (m), 1726 (s), 1639 (m), 1600 (m), 1451 (m), 1435 (m), 887 (s), 587 (m) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  1.0–3.1 with major absorptions at 1.62, 2.01, 2.39, 2.74 (m, 7 H), 4.56 (m, 2 H), 5.20–5.96 (m, 2 H), 9.27 (t, J = 2 Hz, 1 H).

Anal. Calcd for C<sub>9</sub>H<sub>12</sub>O: C, 79.37; H, 8.88. Found: C, 79.17; H, 8.81.

endo-Bicyclo[4.1.0]hept-4-en-2-acetaldehyde (21, 120 min, 9.6%): IR (PE-621) 3076 (w), 3037 (m), 3005 (m), 2925 (m), 2917 (m), 2897 (m), 2880 (m), 2859 (m), 2818 (m), 2716 (m), 1725 (s), 1639 (w), 1455 (m), 1401 (m), 1024 (m), 697 (m)  $cm^{-1}$ ; NMR (60 MHz)  $\delta$  0.50–1.33 (m, 4 H), 1.33–2.83 with major absorptions at 1.78, 2.28 (m, 5 H), 4.97 (m, 1 H), 5.68 (m, 1 H), 9.30 (t, J = 1.5 Hz, 1 H).

Anal. Calcd for C<sub>9</sub>H<sub>12</sub>O: C, 79.37; H, 8.88. Found: C, 79.42; H, 8.90

Irradiation of 21 (19.7 mg, 0.145 mmol, 0.0145 M, 10.0 mL solution) for 16.1 h followed by VPC analysis indicated 21 (42%) and cycloheptatriene (22, 58%).

trans-Bicyclo[4.3.0]non-2-en-8-one (19, 144 min, 56%); IR (PE-621) 3475 (w), 3025 (m), 2964 (m), 2923 (m), 2895 (m), 2864 (m), 2840 (m), 1748 (s), 1635 (w), 1412 (m), 1183 (m), 1179 (m), 1128 (m), 1118 (m), 680 (m) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  1.92 (m, 10 H), 5.37 (m, 2 H).

Anal. Calcd for C<sub>9</sub>H<sub>12</sub>O: C, 79.37; H, 8.88. Found: C, 79.06; H, 8.96.

Hydrogenation of 19 (73 mg, 0.54 mmol) with 5% Pd/C in methanol at  $\sim$ 1 atm followed by VPC isolation gave 44 mg (59%) of a colorless liquid which had identical infrared and NMR spectra with those of trans-bicyclo[4.3.0]nonan-8-one (35).15

Photolysis of exo-Tricyclo[3.3.2.0<sup>2,4</sup>]decan-6-one (10). Ketone 10 (97.2 mg, 0.647 mmol, 0.0259 M, 25.0 mL solution) was irradiated for 8.00 h. VPC analysis of the photolysate was carried out on column A (140 °C, 109 mL/min) and yields were determined from calibrated chromatograms. The longest retention time fraction was identified as starting 10 (71 min, 96% conversion) with the other three fractions (68%) identified as 28-30.

Methyl trans-2-methylcyclohex-3-ene-1-propanoate (29, 22 min, 26%): IR (PE-621) 3060 (w), 3019 (s), 2952 (s), 2927 (s), 2874 (s), 2841 (s), 1739 (s), 1647 (w), 1459 (s), 1451 (s), 1434 (s), 1418 (m), 1370 (m), 1315 (m), 1300 (m), 1257 (s), 1198 (s), 1170 (s), 1083 (w), 1016 (m), 880 (w), 705 (m), 684 (s) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$ 0.83-2.50 with d, J = 7 Hz, at 1.03 (m, 13 H), 3.62 (s, 3 H), 5.47 (m, 2 H). Irradiation at  $\delta$  1.97 caused collapse of d at  $\delta$  1.03 into br s.

Anal. Calcd for C<sub>11</sub>H<sub>18</sub>O<sub>2</sub>: C, 72.49; H, 9.96. Found: C, 72.45; H, 9.95

exo-Bicyclo[4.1.0]hept-4-en-2-propanal (30, 33 min, 7.9%): IR (PE-621) 3073 (w), 3033 (m), 3006 (m), 2929 (s), 2893 (m), 2858 (m), 2816 (m), 2714 (m), 1790 (w), 1727 (s), 1637 (w), 1451 (m), 1409 (w), 1388 (w), 1169 (w), 1020 (m), 970 (w), 949 (w), 724 (w), 692 (m) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$  0.42–2.80 with major absorptions at 0.65, 1.07, 1.87 and with dt, J = 1, 7 Hz, at 2.42 (m, 11 H), 5.22 (m, 1 H), 5.96 (m, 1 H), 9.74 (t, J = 1 Hz, 1 H).<sup>31</sup>

trans-Bicyclo[4.4.0]dec-2-en-9-one (28, 45 min, 34%): IR (PE-621) 3055 (w), 3024 (m), 2923 (s), 2863 (s), 2842 (m), 1789 (w), 1716 (s), 1676 (w), 1644 (w), 1455 (m), 1447 (w), 1434 (m), 1428 (m), 1419 (m), 1336 (m), 1315 (m), 1253 (m), 1228 (m), 1213 (m), 1166 (m), 1157 (w), 694 (m), 664 (w), 650 (m) cm<sup>-1</sup>; NMR (60 MHz)  $\delta$ 0.95-2.68 with peak maxima at 1.23, 1.48, 1.65, 1.92, 2.12, 2.22, 2.35, 2.42 (m, 12 H), 5.48 (m, 2 H).

Anal. Calcd for C<sub>10</sub>H<sub>14</sub>O: C, 79.95; H, 9.39. Found: C, 79.94; H, 9.50

Hydrogenation of 28 (76 mg, 0.51 mmol) as described above for 19 gave 59 mg (77%) of a colorless liquid identified as trans-2decalone (37) from comparison of its infrared and NMR spectra with those of an authentic commercial sample.

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## Competitive Condensation and Proton-Transfer Reactions. Temperature and Pressure Effects and the Detailed Mechanism

### Michael Meot-Ner (Mautner)

Contribution from The Rockefeller University, New York, New York 10021. Received June 12, 1978

Abstract: At high pressures ( $P \gtrsim 1$  Torr) carbonium ions (R<sup>+</sup>) and amines (A) undergo proton-transfer reactions yielding AH<sup>+</sup> in competition with the formation of condensation products RA<sup>+</sup>. For example,  $t-C_4H_9^+$  reacts with NH<sub>3</sub> to give NH<sub>4</sub><sup>+</sup> in competition with the formation of  $t-C_4H_9NH_3^+$ ;  $i-C_3H_7^+$  reacts with  $C_2H_5NH_2$  to give  $C_2H_5NH_3^+$  in competition with  $i-C_3H_7(C_2H_5)NH_2^+$ , etc. The product distribution ratio  $I_{RA+}/I_{AH+}$  increases linearly with third body pressure at low pressures, but levels off to small finite values at high pressures. The product distribution ratio does not vary significantly with temperature between 200 and 500 K in any of five reactions whose temperature dependence was measured. Displacement-exchange reactions such as  $i-C_3H_7^+ + C_2H_5NH_2 \rightarrow i-C_3H_7NH_3^+ + C_2H_6$  do not take place even when those reactions would be more exothermic than the proton-transfer reactions. The pressure effects and the absence of exothermic displacement reactions suggest that the major channel for proton transfer does not proceed through the excited condensation-product-like  $\sigma$ -bonded complexes (RA<sup>+\*</sup>)<sub>tight</sub>. Rather, the data is consistent with a two-stage mechanism (see Scheme II) in which  $(RA^{+*})_{tight}$  and  $AH^+$  are formed from a common precursor. It is proposed that the collision between  $R^+$  and A forms first a loose, electrostatically bonded complex  $(R^+A)*_{loose}$  in which  $R^+$  and A undergo multiple internal collisions, some of which may produce AH+ or RA+\* depending on the geometry of the intracomplex collision. It is further proposed that a similar twostage mechanism can account for some unexpected pressure and temperature effects in other ion-molecule reactions.

The theoretical treatment of bimolecular and higher order reactions in general, and ion-molecule reactions in particular, should be greatly facilitated if the reactions proceed through intermediates which possess well-defined structures and whose internal energy is distributed statistically among the internal degrees of freedom. The decomposition rates of such intermediates to products or to reactants may then be calculated using unimolecular decomposition theory.<sup>1</sup> However, in order properly to evaluate the role of the intermediate in the overall reaction, the detailed mechanism of the reaction must be known. The present work deals with the detailed mechanism of competitive condensation and proton-transfer reactions between carbonium ions and alkylamines as model ion-molecule processes.

Several authors have applied unimolecular theory to the decomposition of intermediates in ion-molecule reactions. For example, Buttrill<sup>2</sup> applied statistical unimolecular theory to calculate product distributions in the decomposition of the  $C_4H_8^{+*}$  and  $C_4H_6^{+*}$  intermediates in the reaction of  $C_2H_4^{+}$ with C<sub>2</sub>H<sub>4</sub> and C<sub>2</sub>H<sub>2</sub>. Several workers used statistical unimolecular theory to interpret temperature and pressure effects in clustering reactions of protonated amines.<sup>3-5</sup> Olmstead and Brauman<sup>6</sup> used RRKM theory to calculate reaction efficiencies of nucleophilic displacement reactions involving negative ions. Su and Bowers<sup>7</sup> also considered that such an approach may be applicable to the decomposition of the reaction intermediate in proton transfer from  $t-C_4H_9^+$  to ammonia. In these treatments it was generally assumed that a specifically bonded complex with randomized internal energy is formed in every capture collision of the reactants.

The present work deals with reactions between carbonium ions (R<sup>+</sup>) and amines (A). At high pressures ( $P \gtrsim 1$  Torr) these reactants yield condensation products RA+, in competition with the major product AH<sup>+</sup> which results from proton transfer. It is safe to assume that the condensation product RA<sup>+</sup> is preceded by a  $\sigma$ -bonded excited species RA<sup>+</sup>\* whose structure is identical with that of RA<sup>+</sup>. This intermediate could then be treated by statistical unimolecular theory. The objective of the present study is to use competition kinetics to examine whether proton transfer also proceeds through this