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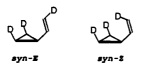
Stereochemistry of the Thermal Isomerization of Vinylcyclopropane to Cyclopentene

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Received August 29, 1994

The stereochemistry of the thermal isomerization of vinylcyclopropane to cyclopentene¹ has long been recognized as a mechanistic key for this simplest known [1,3] carbon sigmatropic pericyclic reaction, 2,3 yet experiments to determine reaction stereochemistry-the relative importance of the four stereochemical modes potentially available for the process—have not been reported. Such experiments have appeared most daunting since deuterium-labeled vinylcyclopropanes are known to suffer thermal stereomutations much faster than they isomerize to labeled cyclopentenes.⁴ To get past this awkward circumstance, one would need to prepare suitably labeled vinylcyclopropanes, heat them to cause both stereomutations and structural isomerizations, analyze recovered starting material and product mixtures at such short reaction times that substantial stereochemical integrity remained in the vinylcyclopropanes, and calculate the relative rate constants for the si, ar, sr, and ai paths characterized by suprafacial or antarafacial allylic participation and retention or inversion at the migrating carbon atom. This project has now been accomplished. Vinylcyclopropane-2,3,2' d_3 isomers syn-E and syn-Z were prepared⁵ and heated in the gas phase at 300 °C. The overall structural isomerizations were followed by capillary GC; the rate constant for isomerizations to cyclopentenes was found to be $3.4 \times 10^{-6} \text{ s}^{-1.6}$



Vinylcyclopropanes recovered from product mixtures were analyzed by ¹H NMR to determine the extent of equilibration of *exo*-C(2,3)-H with *endo*-C(2,3)-H absorption intensities

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(5) Baldwin, J. E.; Villarica, K. A. J. Org. Chem. submitted for publication.

(6) Followed over several half-lives, the isomerizations exhibit a diastereotopically distinct k_H/k_D effect; the Z isomers rearrange somewhat faster than the E isomers.

Table 1. Absorption Frequencies and Spin-Spin Coupling Constants for the Three Isomers of Cyclopentene $-3,4,5-d_3$ at Room Temperature

isomer	ν of C(4)—H (Hz at 500 MHz) ^a	C(4)-H vicinal J values (Hz)
сс	2.58	9.37, 9.37
rac	0.00	9.41, 5.58
tt	-2.72	5.79, 5.79

^a Relative to $\nu(rac) = 0$, in CD₂Cl₂.

Table 2. Cyclopentene-3,4,5- d_3 Isomers from Thermal Gas Phase Reactions of syn-E and syn-E at 300 °C

time (min)	all (%)	all (calcd)a	cc (rel %)	rac (rel %)	tt (rel %)
$61 (E)^{b}$	1.3	1.2	16	60	24
102 (E)	2.0	2.1	18	56	26
130 (E)	2.7	2.6	18	56	26
48 (Z)	1.0	1.0	23	41	36
88 (Z)	1.8	1.8	25	42	35

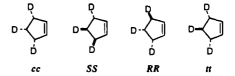
^a From $k = 3.4 \times 10^{-6} \text{ s}^{-1}$. ^b From syn-E as starting material.

associated with (SS-E + RR-E) = rac-E and anti-E (or the corresponding Z isomers).^{4,7} Rate constants for stereomutations



of starting materials, $k_i(syn-E) = 1.7 \times 10^{-4} \text{ s}^{-1}$ and $k_i(syn-Z) = 2.1 \times 10^{-4} \text{ s}^{-1}$, were based on ¹H NMR spectroscopic analyses of GC-purified vinylcyclopropanes recovered from thermal reaction mixtures (five time, absorption intensity ratio values for each starting material).

The three cyclopentene-3,4,5- d_3 isomers cc, (SS + RR) = rac, and tt were quantified through high-resolution ${}^{1}H\{{}^{2}H\}$ NMR spectroscopy, taking advantage of the expected stere-ochemically sensitive isotope-induced differences in chemical shift for the C(4)-H absorptions in each. The parameters giving



the best fit to the 12 C(4)—H NMR transitions seen for the three isomers are summarized in Table 1 and are consistent with previous data.^{8,9} The cyclopentene- d_3 isomers in product mixtures are recorded in Table 2.

Starting from syn-E, the time-dependent fractional concentrations of vinylcyclopropanes are given by $[syn-E] = \exp(-kt)(1 + 3\exp(-k_it))/4$, $[rac-E] = \exp(-kt)(1 - \exp(-k_it)/2)$, and $[anti-E] = \exp(-kt)(1 - \exp(-k_it))/4$. With these and analogous expressions for the Z isomers, one may calculate the average concentration of each isomer over a given reaction time (Table 3).

The relative magnitudes of the rate constants for the four paths for the vinylcyclopropane to cyclopentene isomerization $(k_{si},$

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⁽⁷⁾ The relationship between k_i for approach to equilibrium and conventional stereomutation rate constants is $k_i = 4(k_2 + k_{12}) \approx 2(k_1 + k_2 + k_{12} + k_{23})$; [rac-E] is always 2[anti-E].⁴

⁽⁸⁾ Anet, F. A. L.; Leyendecker, F. J. Am. Chem. Soc. 1973, 95, 156-

⁽⁹⁾ Details of the NMR spectroscopic work, including temperature-dependent studies, will be published separately.

Table 3. Calculated Average Concentrations of Vinylcyclopropane- d_3 Isomers over Short Reaction Time Intervals at 300 °C

time (min)	e (min) syn-E or -Z ra		anti-E or -Z	
61 (E)a	80.4	12.6	6.3	
102(E)	71.0	18.6	9.3	
130 (E)	65.8	22.0	11.0	
48 (Z)	80.9	12.4	6.2	
88 (Z)	69.8	19.6	9.8	

^a From syn-E starting material.

 $k_{\rm ar}$, $k_{\rm sr}$, $k_{\rm ai}$) follow from the relevant linear equations, such as eqs 1-3.

$$k_{ar}[syn-E] + (k_{si} + k_{ai})[rac-E]/2 + k_{sr}[anti-E] = [cc]$$
 (1)

$$(k_{\rm si} + k_{\rm sr})[syn-E] + [rac-E]/2 + (k_{\rm ar} + k_{\rm ai})[anti-E] = [rac]$$

(2)

$$k_{ai}[syn-E] + (k_{sr} + k_{ar})[rac-E]/2 + k_{si}[anti-E] = [tt]$$
 (3)

All vinylcyclopropane concentration terms on the left of eqs 1-3 are averages over a reaction time (Table 3); the relative product concentrations on the right are those observed at that time (Table 2), and $(k_{\rm si} + k_{\rm ar} + k_{\rm sr} + k_{\rm ai}) = 1$. From each

Table 4. Experimental Relative Rate Constants for the Four Stereochemically Distinct Paths for the Isomerizations of Vinylcyclopropane-2,3,2'-d₃ to Cyclopentene-3,4,5-d₃ at 300 °C

starting isomer	time (min)	$k_{ m si}$	$k_{\rm sr}$	$k_{\rm ar}$	k_{ai}
syn-E	61	0.64 0.61 0.62		0.13	0.24
	102			0.14	0.27
	130			0.13	0.27
syn-Z	48	0.40 0.22 0.42 0.25		0.38	
	88			0.37	

product mixture from *anti-E*, one may measure $(k_{\rm si}+k_{\rm sr})$, $k_{\rm ar}$, and $k_{\rm ai}$, while each derived from syn-Z gives $k_{\rm sr}$, $k_{\rm si}$, and $(k_{\rm ar}+k_{\rm ai})$. These experimental relative rate constants are summarized in Table 4.

The five kinetic runs give quite consistent information: all four paths are kinetically competitive; the relative magnitudes of the rate constants (in percentage terms, $\pm 2-3\%$) are $k_{\rm si}=40\%$, $k_{\rm sr}=23\%$, $k_{\rm ar}=13\%$, and $k_{\rm ai}=24\%$. Thus there is no significant kinetic preference for the orbital symmetry-allowed $k_{\rm si}$ and $k_{\rm ar}$ paths, there in no "energy of concert" in evidence, and diradical-mediated paths seem mechanistically essential.

Acknowledgment. We thank the National Science Foundation for support of this work at Syracuse University through CHE 9100246 and at UCLA through the Instrumentation Program.

⁽¹⁰⁾ The average of f(t) on (0,t) is $\int_0^t f(t)dt/t$.