Compd	$\nu_2 - \nu_3{}^a$	$\nu_2 - \nu_7$	$\nu_3 - \nu_7$	$D_{23}{}^a$	D 24	D_{25}	D_{34}	D_{27}	D_{37}	$D_{77'}$	Δf^b	c, mol %
1 2	$\begin{array}{r} 0.7 \\ -17.2^{d} \\ -21.7^{e} \\ -17.3^{e} \\ -11.9^{e} \end{array}$	346.5	347.2	-450.4 -393.5 -393.8 -393.4 -393.5	-51.6 14.1 14.2 14.1 14.1	-20.4 36.6 37.2 36.5 36.5	-183.6330.1329.7330.5330.0	-50.1 -48.6	- 79.5 - 48.2	1795.0 455.1	0.65 0.63	16 13

^a Probable error of parameters <0.8 Hz. ^b Root-mean-square error for calculated and observed line positions. ^c Concentration. ^d Complete analysis. . Subspectral analysis.

Table II. r_{ij} Ratios and Orientation Parameters for 1 and 2

Compd	r_{23}/r_{34}	r_{24}/r_{34}	r_{25}/r_{34}	r_{27}/r_{34}	r_{37}/r_{34}	r777'/r34	S_{xx^a}	$S_{yy}{}^a$	S_{zz}^{a}
1	1.028	1.771	2.081	1.669	2.214	0.760	0.0233	-0.1003	0.0770
	± 0.020	± 0.018	± 0.028	± 0.038	± 0.051	± 0.013	± 0.0001	± 0.0061	± 0.0061
2	1.024	1.769	2.081	1.741	2.293	0.879	-0.0419	-0.0487	0.0906
	± 0.006	± 0.006	± 0.015	± 0.028	± 0.030	± 0.055	± 0.0001	± 0.0021	± 0.0021
Benzene	1.000	1.732	2.000						

^a Based on an assumed H₃-H₄ distance of 2.48 A.

been analyzed separately. The results for both compounds are collected in Table I.

Orientation parameters S_{kl} and the ratios of the various H,H, H,F, and F,F distances r_{ij} were then calculated from the experimental data of Table I using an iterative procedure.⁹ Since both compounds possess C_{2v} symmetry,¹⁰ two independent parameters are sufficient to characterize their orientation. For these calculations, the distance H_3 - H_4 was held constant. The r_{ii} ratios are insensitive to this limitation.¹ To check the results obtained in this way, the ratios of the H,H distances in the six-membered ring of 1 and 2 have also been derived from explicit expressions given for the oriented AA'BB' system.¹¹ They were identical with those found by the iterative method. The results, together with the orientation parameters and the relevant r_{ii} ratios of benzene, are presented in Table II. From these values those based on H,F- and F,F-coupling constants in 2, *i.e.*, r_{27}/r_{34} , r_{37}/r_{34} , and $r_{77'}/r_{34}$, are less reliable, since the experimental $D_{\rm HF}$ and $D_{\rm FF}$ values contain unknown contributions from the anisotropy of the indirect spin-spin coupling.¹²

From the negative signs of the dipolar coupling constants it follows that 1, as other aromatics,¹ orients with its x, z plane preferentially parallel to the external magnetic field. For 2, however, a different orientation is indicated by the positive sign found for D_{24} , D_{34} , and D_{25} and, more directly, by the different S_{xx} value. In contrast to 1, the y as well as the x axis of the molecular coordinate system orient preferentially perpendicular to the magnetic field direction.

Within the limits of error, our measurements yield identical proton geometry for the six-membered ring of 1 and 2. Compared to benzene, all r_{ij}/r_{34} ratios have increased. This seems to indicate that the benzocyclopropene system suffers a distortion of type 3 with positive changes δ for the CCC bond angles at C_3 and C_4 and negative changes for these parameters

at C_2 and C_5 . Similar angle distortions have been measured for 7,8-dichlorobenzocyclobutene13 and ben-

zocyclobutene-7,8-dione,14 where the strain effects should be comparable to those operating in 1 and 2. Attempts to obtain the "best" carbon skeleton which fits into the geometry determined by the proton positions in both compounds using "molecular mechanics"¹⁵ are at present under way.

Acknowledgments. We wish to thank Dr. D. Schäpel and Professor Dr. E. Vogel, Cologne, for samples of 1 and 2, Professor Dr. P. Diehl, Basel, for a copy of his program LAOCOONOR, and one of the referees for a helpful comment. This work was supported by the Deutsche Forschungsgemeinschaft and by the Fonds der Chemischen Industrie.

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Pyrolysis of Cyclopropyl Azides. A Route to 1-Azetines¹

Sir:

Photolytic or pyrolytic decomposition of cyclopropyl azides with ring enlargement or substituent migration presents a potential route to 1-azetines or N-substituted cyclopropylimines. Neither the latter nor 2-alkyl- (or aryl-) 1-azetines have heretofore been reported.² In

⁽⁹⁾ Details of the method will be given in the full paper. (10) The magnetic equivalence of X_7 and X_7' in 1 and 2 demonstrates C_{2v} symmetry only on the nmr time scale, but it seems reasonable to assume that both compounds are planar.

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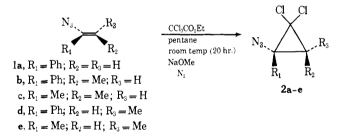
⁽²⁾ The only 1-azetines in the literature have ether or thioether linkages at the 2 position and are obtained from β -lactams. See, for

Azide	% yield (isolated)	Ri		R ₃	Bp, °C (mm)	Pyrolysis rate, $k \times 10^{-4}$ sec ⁻¹
2a	42	2.60 (s)	7.85, (d, J = 8.0)	8.09 (d, J = 8.0)	83-85 (0.1)	4.6
cis-2b	53	2.57 (s)	8.80 (d, J = 6.5)	7.99(q, J = 6.5)	90 (0, 2)	14.7
cis -2c	25	8.50 (s)	8.81 (d, $J = 6.5$)	8.30(q, J = 6.5)	64 (7)	8.2
trans-2d	66	2.57 (s)	7.89(q, J = 6.5)	8.60 (d, J = 6.5)	82 (0.1)	2.9
trans-2e	52	8.27 (s)	8.61 (q, J = 4.5)	8.80 (d, J = 4.5)	65 (7)	1.8

a s = singlet, d = doublet, q = quartet; J values in hertz.

fact, since cyclopropyl halides are neither readily susceptible to displacement reactions nor to solvolysis without rearrangement, cyclopropyl azides have remained a virtually unknown class of compounds.³

Our recent discovery of a general synthesis of vinyl azides from olefins⁴ suggested their reaction with dichlorocarbene as a route to 2. N-Octyl azide has been reported to react with dichlorocarbene to give the corresponding isocyanide dichloride.⁵ In sharp contrast, the vinyl azides 1 gave solely the corresponding 1-azido-2,2-dichlorocyclopropanes 2 in moderate yields (see Table I). This is in analogy with recent studies which showed that vinyl azides react at carbon with other electrophiles such as acid⁶ and bromine.⁷ In a typical experiment 21.9 g (0.115 mol) of ethyl trichloroacetate was added dropwise over a period of 30 min to a solution of 0.1 mol of the vinyl azide and 12.8 g (0.2 mol)mol) of sodium methoxide in 100 ml of pentane at 0° under nitrogen. After stirring for 20 hr at room temperature the mixture was filtered through Celite. Removal of the pentane in vacuo left an oil containing starting material and azide. Purification of the latter was accomplished by careful chromatography followed by distillation below 100° at reduced pressure. (Cau*tion:* these compounds explode on heating at ca. 110°.)



Azides 2a-c decompose smoothly between 105 and 125° in a highly regiospecific⁸ manner to give the corresponding 3,3-dichloroazetines 3 in 50-75% isolated yields (see Table II). No spectral evidence for formation of 4 or 5 was observed. The primary side

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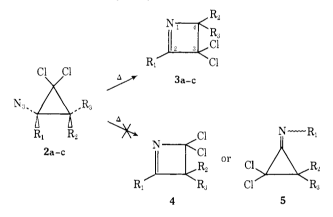
Table II. Yields and Nmr Data of Azetines 3

Azetine	% yield	Nmr spectra, τ^a
3a	55	5.49 (s, 2 H)
3b	72	2.51, 2.05 (2 m, 5 H) 8.49 (d, 3 H, $J = 7$)
		5.40 (q, 1 H, $J = 71$) 2.60, 2.03 (2 m, 5 H)
3c	75	8 59 (d, 3 H, $J = 6$ 5) 7 91 (d, 3 H, $J = 1$ 6)
		5.56 (q of q, 1 H, $J = 1.6$ and 6.5)

n - 1

a s = singlet, d = doublet, q = quartet, m = multiplet; J valuesin hertz.

products appear to be the nitrile 6 and the olefin 7 (ca. 5%), possibly resulting from the fragmentation of the azetine ring. Typically, 5 g of the azide was



refluxed in 200 ml of toluene for 9 hr, the toluene removed in vacuo, and the resultant azetine chromatographed on silica gel.

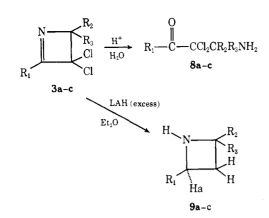
$$R_{1} \xrightarrow{N} Cl \xrightarrow{R_{2}} R_{3} \xrightarrow{mass} R_{1} \xrightarrow{-CN} + R_{2}R_{3}C \xrightarrow{-CCl_{2}} Cl_{2}$$

The structures of **3a-c** were indicated by ir, nmr, and mass spectra. The mass spectra showed strong peaks corresponding to the molecular ion, the nitriles 6, and the olefins 7. Conclusive evidence for the structures came from acid hydrolysis and LiAlH₄ reduction to give the analogous β -amino- α , α -dichloroketones 8 and azetidines 9, respectively.

The structures of the amino ketones 8a-c were indicated by spectral data, the mass spectra being especially helpful since they showed the expected amine α cleavage corresponding to $R_2R_3C{=\!\!=}NH_2{}^+$ and no α cleavage at m/e 98, 100, and 102, corresponding to $CCl_2 = NH_2^+$. The crude nmr spectra of **8a-c** indicated

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example, G. Pifferi, P. Consonni, G. Pelizza, and E. Testa, J. Heterocycl. Chem., 4, 619 (1967); D. Bormann, Justus Lieblgs Ann. Chem., 752, 124 (1969). For another recent example, see A. Hassner, J. O. Currie, A. S. Steinfeld, and R. F. Atkinson, Angew. Chem., Int. Ed. Engl., 9, 731 (1970).



no other products present. The structures of azetidines 9a-c were obvious from their ir, nmr, and mass spectra. That the geminal hydrogens were in the 3 position in **9a** and **b** was indicated by the low-field (τ 5.4) broad triplet for H_a. In 9c H_a could not be resolved, but the mass spectra showed the expected α cleavage at m/e 43 (CH₃CH==NH·+) and no peak at m/e 30 $(CH_2 = NH \cdot +).$

The regioselectivity in the formation of 3 over 4 and 5 cannot be equated with conformational effects⁹ governing a concerted loss of N₂ with ring enlargement, since the conformers of 2 are expected to be of nearly equal stability and should produce a mixture of 3, 4, and **5**.

The driving force for ring expansion in preference to formation of a cyclopropylimine 5 (Ph migration) must be relief of ring strain in going from a threeto a four-membered ring. Apparently, the reaction involves selective migration of the electron richer CH₂ (or CR_2) rather than the CCl_2 ring residue to the electron-deficient nitrogen atom.

Rate data on the pyrolysis of cyclopropyl azides 2 (Table I) indicate that in general cis substituents (see 2b and 2c) accelerate the reaction. This implies that the rate-determining step cannot be loss of N₂ unless the reaction is concerted. A concerted ring expansion with loss of nitrogen is in fact suggested by the negative entropy of activation ($\Delta S^{\pm} = -18$ eu for $2c \rightarrow 3c$) and is in accord with data on pyrolysis of simple alkyl azides.10

Acknowledgment. Support of this work by a grant (CA 04474) from the National Cancer Institute (Public Health Service) is gratefully acknowledged.

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Effect of CH Scalar Coupling on ¹³C Transverse **Relaxation Times**

Sir:

In view of the increasing interest in ¹³C nmr spectroscopy, we wish to report some preliminary relaxation studies of ¹³C-enriched compounds. We find that the transverse relaxation times, T_{2C} ($\equiv R_{2C}^{-1}$), are usually shorter than the longitudinal relaxation times, T_{1C} $(\equiv R_{1C}^{-1})$. This result influences the sensitivity and resolution in both conventional and Fourier transform (FT) ¹³C spectra and, in particular, affects the signal enhancement obtainable by the use of refocusing schemes such as driven equilibrium and spin-echo Fourier transform (DEFT and SEFT) techniques.¹⁻³

The T_{2C} measurements were performed at 15.08 MHz by the usual Meiboom-Gill⁴-modified Carr-Purcell⁵ technique on a high-resolution magnet with external fluorine lock. The π pulse width was 6 μ sec and the inhomogeneity over the sample gave a free-induction decay constant $T_{2}^{*} \sim 100$ msec. Degassed and sealed sample tubes were constricted below the liquid surface to minimize interchange between liquid and vapor. T_{1C} and T_{1H} values were measured by $\pi - \tau - \pi/2$ pulse sequences, T_{1C} 's at 15.08 MHz, T_{1H} 's at 23.2 MHz. The T_1 and T_2 values were obtained by the usual leastsquares fitting procedures.

Table I shows that T_{2C} is substantially shorter than

Table I. Relaxation Times^a for 60% ¹³C-Enriched Compounds at 25°

	T_{1C} , sec	T_{2C} , sec	$T_{1\mathrm{H}},^{b}$ sec
¹³ CH ₃ I	13.4	3.9	10.9°
$^{13}CH_{3}COOCD_{3}^{d}$	19.2	6,1	12.5
$^{13}CS_{2}$	48e	38	

^a The experimental error is estimated to be $\sim 5\%$. ^b T_{1H} for 100% enriched compounds will be less than these values. CT_{1H} for ¹²CH₃I at 25° is reported [M. D. Zeidler, Ber. Bunsenges. Phys. *Chem.*, **69**, 659 (1965)] to be 12.5 sec. ^{*d*} Deuterium enrichment is 99%. $^{\circ}T_{1C} = 30$ sec (15 MHz, 26–28°) was measured by rapid adiabatic passage: A. Olivson, E. Lippmaa, and J. Past, Eesti NSV Tead. Akad. Toim., 16 390 (1967).

 T_{1C}^{6} for methyl iodide and methyl- d_{3} acetate-¹³C. We would like to point out that short T_{2C} 's occur when relaxing spin- $1/_2$ nuclei, e.g., protons, are coupled to 1^{3} C such that

$$A \equiv 2\pi J \gg R_{1\rm H} \tag{1}$$

where $R_{1H} \equiv 1/T_{1H}$. Gutowsky, Vold, and Wells⁸ have pointed out the analogy between chemically exchanging, equally populated, uncoupled two-site systems and relaxing, scalar-coupled AX systems. They suggest that the two cases are related by the simple substitution of $\Omega \to A/2$ and $k \to R_{1X}$, where Ω is half the chemical shift between the two sites, k the exchange rate, and R_{1X} the spin-lattice relaxation rate of X.

Scalar contributions to R_{2A} for a general AX_n system originate from relaxation processes within the X manifold which preserve the spin state of A while changing the scalar coupling energy by a nonzero multiple of J. These processes randomly alter the precession frequency

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