Microwave spectrum, dipole moment, structure, and internal rotation of the cyclopropane-sulfur dioxide van der Waals complex

Anne M. Andrews, Kurt W. Hillig II, and Robert L. Kuczkowski Department of Chemistry, University of Michigan, Ann Arbor, Michigan 48109-1055

(Received 29 July 1991; accepted 23 October 1991)

The rotational spectrum of the cyclopropane-sulfur dioxide complex was observed by Fourier transform microwave spectroscopy. The spectrum exhibited a- and c-dipole selection rules with the c-dipole transitions split into doublets of unequal intensity separated by about 150 kHz. The structure has C_s symmetry with the sulfur and carbon atoms all lying in the ac plane; the oxygen and hydrogen atoms straddle the plane. The sulfur dioxide plane is nearly parallel to a C-C bond edge. The distance from the center of mass of the SO₂ to the C-C bond center is 3.295 Å. The dipole moment of the complex is 1.681(1) D, with components $\mu_a = 0.815(1)$ D and $\mu_c = 1.470(1)$ D. The splittings in the spectrum arise from an internal rotation of the cyclopropane subunit about its local C_2 axis which lies nearly along the line connecting the centers of mass.

I. INTRODUCTION

Complexes of HF, HCl, and HCN with the hydrocarbon series ethylene, 1-3 acetylene, 4-6 and cyclopropane 7-9 have been studied recently. In each complex, the acid is hydrogen bonded to the π (ethylene and acetylene) or pseudo- π (cyclopropane) system. 10 It has been noted that for each HX series there is a decrease in hydrogen-bond length, an increase in pseudodiatomic stretching force constant, and an increase in induced dipole moment from ethylene to acetylene to cyclopropane. 11,12 As these properties are often correlated with the strength of the interaction, this has led to a discussion that the pseudo- π system of cyclopropane forms stronger hydrogen bonds than the classical π systems of ethylene and acetylene. Legon and Millen, studying the pseudodiatomic force constants of the HCl and HCN series, have assigned nucleophilicities to C₂H₄ (4.7), C₂H₂ (5.1), and C_3H_6 (6.4).¹²

While these trends are well documented for the hydrocarbon-acid complexes, which are hydrogen bonded, there is less data for this hydrocarbon series complexed to a nonhydrogen-bonding partner. Recently in our lab the complexes of ethylene and acetylene with sulfur dioxide have been observed. 13,14 In both cases, the complexes have a stacked structure with the C_2 axis of the SO_2 crossed at 90° to the C = C or C = C bond. The sulfur of the SO_2 apparently interacts with the π system of the ethylene and acetylene. Like the acid complexes, it was observed that the interaction distance, as measured from the C-C bond center to the sulfur, is shorter for $C_2H_2 \cdot SO_2$ (3.359 Å) than $C_2H_4 \cdot SO_2$ (3.446 Å) and that the induced dipole moment is greater for $C_2H_2 \cdot SO_2$. The force constants for the stretching vibration between the hydrocarbon and the SO₂ are not easily compared due to the effect of internal rotation in C₂H₄·SO₂. This is discussed in Ref. 14.

Here we report on the cyclopropane \cdot SO₂ (C_3 H₆ \cdot SO₂) complex. The sulfur atom interacts with the pseudo- π system as expected, however, the structure is slightly different from the C_2 H₄ \cdot SO₂ and C_2 H₂ \cdot SO₂ complexes in that the C_2 axis of the SO₂ is nearly parallel to the C-C bond, rather than crossed at 90°. The interaction distance (S to C-C bond

center) of 3.203 Å is shorter than in $C_2H_2 \cdot SO_2$. Moreover, the induced dipole moment and the pseudodiatomic force constant are both greater.

II. EXPERIMENT

A. Spectrometer

The spectrum was observed in a Fourier transform microwave spectrometer of the Balle-Flygare type which has been described previously. The molecular beam was generated with a modified Bosch fuel injector. Linewidths were typically 20–30 kHz full width at half maximum (FWHM) and center frequencies were estimated to be accurate to \pm 2–3 kHz. For deuterated isotopomers, transitions were broadened to 100 kHz or more from unresolved nuclear quadrupole hyperfine structure and line centers were accurate to \pm 20–30 kHz. Stark effects were measured by applying up to 10 000 V with opposite polarities to two parallel steel mesh plates separated by about 30 cm.

B. Samples

The spectrum of $C_3H_6 \cdot SO_2$ was observed with a mixture of approximately 1% each of C_3H_6 (Aldrich) and SO_2 (Matheson) in Ar at a total pressure of 1.5 atm. $S^{18}O_2$ (98% $^{18}O_3$) was purchased from Alfa Products and used without dilution to observe the $C_3H_6 \cdot S^{18}O_2$ spectrum. A 50:50 mixture of $S^{18}O_2$ and $S^{16}O_2$ was used to produce the $C_3H_6 \cdot S^{18}O^{16}O$ spectrum; the samples exchanged immediately upon mixing. The $C_3H_6 \cdot ^{34}SO_2$ spectrum was observed in natural abundance (4% $^{34}S)$. C_3D_6 (98% D) and 1,1- $C_3H_4D_2$ (98%) were purchased from MSD isotopes.

 $\rm C_3\,H_5\,D$ was synthesized in poor yield as follows. Cyclopropyl Grignard reagent was produced by reacting cyclopropyl bromide (Aldrich) with Mg (Baker) in dry diethyl ether in the usual manner. The flask containing the Grignard reagent was then placed in line with a trap cooled with a $\rm CCl_4$ slush ($-25\,^{\circ}\rm C$) followed by a liquid-nitrogen trap ($-196\,^{\circ}\rm C$) which was isolated from the atmosphere by a mercury bubbler. Nitrogen gas was passed through the apparatus while $\rm D_2O$ (Cambridge Isotope Lab) was slowly

TABLE I. Observed transitions in MHz of C₃H₆·SO₂.

$J'_{K_pK_0}$	$J_{K_pK_0}^{"}$	A_1/A_2^a	$J'_{K_PK_0}$	$J_{K_pK_0}^{\prime\prime}$	A_1/A_2
3 ₀₃	202	7 311.753	523	4 ₂₂	12 207.788
322	221	7 315.089	514	4 ₁₃	12 383.484
321	220	7 318.850	6,6	5 ₁₅	14 387.127
3,2	211	7 432.386	606	5 ₀₅	14 597.283
414	313	9 595.002	615	514	14 856.965
404	303	9 744.449	7,,	616	16 780.978
423	3,2	9 752.487	707	606	17 015.605
432	331	9 754.511	716	615	17 382.667
431	330	9 754.584	221	211	14 754.456
422	321	9 761.865	220	212	14 990.582
413	312	9 908.517	321	313	15 112.216
515	414	11 991.722	422	414	15 279.080
505	404	12 173.271	5 ₂₃	515	15 495.147
524	423	12 189.058			
		A_2 (strong)		A_1 (weak	()
110	000	7 435.084	·,·····	7 435.24	15
2,,	101	9 951.926		9 952.07	70
3,2	202	12 508.184		12 508.34	16
4,3	303	15 104.949		15 105.10)4
6 ₀₆	514	9 026.565		9 026.41	12
707	615	11 185.201		11 185.03	39

^{*}Symmetry designation of tunneling doublets. A_1/A_2 indicates that the transition was unsplit.

added dropwise to the cyclopropyl Grignard. Excess D_2O and ether were trapped in the CCl_4 trap and a small amount of C_1H_5D was collected in the liquid-nitrogen trap.

III. RESULTS AND ANALYSIS

A. Spectrum

The spectrum of $C_3H_6 \cdot SO_2$ exhibited a- and c-dipole selection rules. The c-type R-branch transitions were split into doublets of unequal intensity and the strong and weak ctype transitions were each fit independently with the a-type transitions to a Watson S-reduced Hamiltonian. 17 The observed transitions are listed in Table I and the derived constants are shown in Table II. The splitting of the c-type transitions arises from an internal rotation of the cyclopropane subunit exchanging three pairs of protons (see internal rotation section below) and the A_1 and A_2 symmetry labels of the states are taken by analogy to ethylene SO2 which has a similar tunneling path. 13 For C₃H₆·SO₂, the A₁ label corresponds to the weaker transitions and the A_2 to the stronger. Additional effects of the internal rotation are seen in the difference in the sign of the D_K distortion constant between the A_1 and A_2 states. A similar effect was observed in the ethylene · SO₂ complex and it is believed to arise from neglect of an internal rotation term in the Hamiltonian.

The spectra of $C_3H_6 \cdot {}^{34}SO_2$, $C_3H_6 \cdot S^{18}O_2$, and $C_3H_6 \cdot S^{18}O^{16}O$ were also all split into doublets, with the same relative intensity pattern as the normal isotopic species. For $C_3H_6 \cdot S^{18}O_2$ and $C_3H_6 \cdot S^{18}O^{16}O$, the magnitude of the splitting of the $2_{11} \cdot 1_{01}$ transitions decreased by 30% and 10%, respectively. For $C_3H_6 \cdot {}^{34}SO_2$, the splitting increased

by 10%. In the $C_3D_6 \cdot SO_2$, the doublets were unresolved due to deuterium nuclear quadrupole broadening. For $C_3H_5D\cdot SO_2$ the transitions were unsplit. Two different spectra were observed for the 1,1- $C_3H_4D_2 \cdot SO_2$ isotopic species; one was split into doublets (labeled *apical*) and the other was unsplit (labeled *basal*). These labels will be dis-

TABLE II. Spectroscopic constants (Watson S reduction; I' representation) of C_3H_6 ·SO₂.

	$A_2^{\ a}$	A_1
A/MHz	6176.635(5) ^b	6176.828(5)
B/MHz	1258.500(1)	1258.500(1)
C/MHz	1180.101(1)	1180.101(1)
D_J/kHz	1.958(5)	1.959(6)
D_{JK}/kHz	15.5(1)	15.4(2)
D_K/kHz	-18.1(9)	20.5(11)
d ₁ /kHz	-0.078(4)	= 0.077(5)
d_2/kHz	0.179(6)	0.178(7)
H_{KI}/kHz	0.9(1)	0.8(2)
h ₃ /kHz	0.04(1)	0.04(1)
n ^c	33	33
$\Delta v_{\rm rms}^{\rm d}/{\rm kHz}$	6	7
μ_a/D	0.815(1)	
μ_c/D	1.470(1)	
μ_T/D	1.681(1)	

a Symmetry label of tunneling state (see text).

 $^{^{\}rm b}$ Uncertainties represent 1σ in the least-squares fit.

^c Number of transitions in the fit.

 $^{^{\}rm d}\Delta v = v_{\rm obs} - v_{\rm calc}$

TABLE III. Observed transitions of cyclopropane-SO₂ isotopic species (in MHz).

J'K, K J'K, K.	C,H, 390,	%	O,O,O,,S.*Y.S	0,0,	0 C,H,S ¹ O,	S ¹⁸ O ₂		C ₂ H ₄ D ₂ ·SO ₂ capical ^b	C, D, -80,	C2H,D·SO2	I,D·SO ₂ C,H,D,·SO ₂
	A2.c	Ā	4,	¥	A2	4	A,	¥.	A_1/A_2^d		
1,0-0m	7403.587	7403.741								7311.752	
211-101	9894.945	9895.104	9676.028	9676.158	9407.874	9407.978	9589.050	9589.150	8851.200	9809.891	710.7756
312-202	12 424.588	12 424.744	12 205.618	12 205.749	11 912.787	11 912.891	11 985.335	11 985.441	11 168.850		12 093.137
413-303	14 993.558	14 993.719	14 778.921	14 779.036	14 464.672	14 464.763	14 419.172	14 419.282	13 517.715		14 646.758
514-404									15 899.180		
220-212											14 072.658
321-313									13 108.825		14 185,448
303-202	7242.123	123	7204.747	747							
322-221			7208.844	#					12 740.400	7263.862	
321-220			7213.358	358							
312-211	7359.271	171	7334.698	869	7242.826	.826				7380.262	
414-313	9506.431	131	9441.707	707	9295.386	.386	806	9000.566	8781.476	9536.547	9480.090
404-303	9651.854	3 5	9600.896	968	9464.759	.759	913	9139.714	8898.074	9680.765	9618.737
423-322			9610.684	684	9476.560	.560	880	8903.725	9146.734		9626.160
422-321			9621.962	962	9490.029	.029	891	8910.253	9154.891		9634.778
413-312	9811.094	\$	9778.040	040	9655.286	.286	626	9291.601	9024.846	9839.058	9770.806
515-414	11 881.103	103	11 799.390	390	11 616.492	(492	11.24	11 248.950	10 975.387	11 918.715	11 848.225
S ₀₅ -4 ₀₄	12 057.907	707	11 992.417	.417	11 820.635	1,635	1141	11 418.305	11 117.481	12 093.932	12 016.714
524-423							11 43	11 432.093	11 128.502		12 031.243
523-422							11.4	11 448.382	11 141.545		
514-413	12 261.818	318	12 220.021	.021	12 066.153	153	1161	11 615.640	11 279.513	12 296.731	12 211.498
614-513			14 155.851	.851					13 168.348		14 215.132
6 ₀₆ -5 ₀₅	14 459.410	011	14 378.225	225	14 169.756	1.756	13 69	13 692.702	13 333.516		14 410.259
625-524							13 71	13 716.568	13 352.544		14 435.352
624-523							13.74	13 745.006	13 375.327		
615-514									13 533.130		14 650.816
717-616									15 360.197		
70,-606									15 545.550		

 $^{\bullet}D_{1}$ or D_{2} substitution at the C atom closest to the SO₂ (see Fig. 2). Only a singlet was observed.

 $^{^{}b}D_{2}$ substitution at the C atom most distant from the SO₂ (see Fig. 2).

^eSymmetry label of tunneling state (see text).

^dTunneling doublets not resolved.

TABLE IV. Spectroscopic constants for C₁H₆·SO₂ isotopic species.

	Symmetry*	A/MHz	B/MHz	C/MHz	n^{b}	$\Delta u_{ m rms}^{\ \ c}$
C ₃ H ₆ ·SO ₂	A ₁	6176.635(5) ^d	1258.500(1)	1180.101(1)	33	6
	A_2	6176.828(5)	1258.500(1)	1180.101(1)	33	7
C ₃ H ₆ ·34SO ₂	A_1	6157.911(7)	1245.746(2)	1169.572(2)	13	9
	A_2	6158.066(7)	1245.745(2)	1169.571(2)	13	10
C3H6.S18O2	A_1	5718.558(2)	1229.851(1)	1139.843(1)	13	11
	A_2	5718.664(4)	1229.850(1)	1139.845(1)	13	3
C3H6.S18O16O	A_1	5945.214(3)	1243.685(1)	1159.500(1)	17	3
	A_2	5945.345(3)	1243.684(1)	1159.501(1)	17	2
1,1-C ₃ H ₄ D ₂ ·SO ₂	-					
(apical)f	A_1	6049.426(5)	1179.925(1)	1107.162(1)	16	6
	A_2	6049.523(5)	1179.926(1)	1107.162(1)	15	5
$C_3D_6 \cdot SO_2$ $C_3H_5D \cdot SO_3$	A_1/A_2 °	5420.840(57)	1143.558(1)	1082.701(1)	30	6
(basal) ² 1,1-C ₃ H ₄ D ₃ ·SO ₃		6062.694(14)	1249.091(3)	1173.458(4)	10	18
(basal)f,8		5857.749(6)	1239.833(1)	1167.135(1)	19	4

^{*}Symmetry label of tunneling state (see text).

cussed in Sec. III C. For the spectrum of the apical species both tunneling doublets were very weak, making it difficult to assign strong and weak components. Therefore, the symmetry labels were assigned by higher/lower frequency to be consistent with the other isotopic species. The magnitude of the splitting decreased by 30%. The transitions for the isotopomers are listed in Table III and the rotational constants are given in Table IV; because the fitted distortion constants were similar to those for the normal isotopic form, only A, B, and C are reported for the isotopomers.

B. Dipole moments

The dipole moment of the complex was measured by tracking the Stark effect of seven M components from four transitions with increasing electric field. The electric field was calibrated using the $J=1\leftarrow0$, $M=0\leftarrow0$ transition of OCS at 12 162.980 MHz. The observed Stark effects were least-squares fit to dipole components $\mu_a=0.815(1)$ D and $\mu_c=1.470(1)$ D resulting in $\mu_T=1.681(1)$ D. When μ_b was included in the fit, the value for μ_b^2 was 0.030(50) D, indicating that μ_b was zero. The dipole moment components of $basal\text{-C}_3\text{H}_4\text{D}_2\cdot\text{SO}_2$ were also determined as

 $\mu_a = 0.803(3)$ D and $\mu_c = 1.482(3)$ D from six M components from two transitions.

C. Structure

It was assumed in the structural analysis that the geometries of the cyclopropane and sulfur dioxide were not changed upon complexation from their free gas-phase structures. ^{19,20} The moments of inertia and planar second moments of the stronger set of transitions were used for the isotopic species which exhibited splittings. Because the difference in rotational constants is very small (< 200 kHz), this choice will not markedly affect the determination of the structure. The planar moments for $C_3H_6 \cdot SO_2$, $C_3H_6 \cdot ^{34}SO_2$, cyclopropane, and sulfur dioxide are listed in Table V.

The a- and c-type selection rules and the absence of evidence for a μ_b dipole component suggest that the complex has an ac symmetry plane. Comparison of the planar moment P_{bb} of $C_3H_6 \cdot SO_2$ (54.2495 amu Ų) with P_{bb} of $C_3H_6 \cdot ^{34}SO_2$ (54.2460 amu Ų) indicates that the S atom lies in this plane. P_{bb} for the complex is also equal to the sum of P_{aa} of free SO_2 and P_{cc} of free cyclopropane, placing the

TABLE V. Planar second moments $[P_{aa} = 0.5 (I_c + I_b - I_a) = \sum m_i a_i^2$ and similarly for P_{bb} , P_{cc} of cyclopropane-SO₂, cyclopropane and sulfur dioxide.

	$C_3H_6\cdot SO_2$	$C_3H_6\cdot {}^{34}SO_2$	Cyclopropane	SO ₂
P _{aa} ∕amu Ų	374.0012	377.8599	20.1254	48.7679
P _{bb} ∕amu Ų	54.2495	54.2460	20.1254	8.3574
P _∞ /amu Ų	27.5716	27.8239	5.0262	0.0

^b Number of transitions in fit.

 $^{^{}c}\Delta v = v_{\rm obs} - v_{\rm calc}$.

^d Uncertainty is 1σ.

^eTunneling doublets not resolved for $C_3D_6 \cdot SO_2$.

See footnote to Table III for explanation of apical and basal labels for 1,1-C₃H₄D₂·SO₂.

^{*} No tunneling doublets observed for these isotopomers.

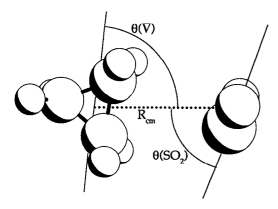


FIG. 1. Definition of structural parameters for cyclopropane \cdot SO₂. $R_{c.m.}$ is the distance from between the centers of mass, $\theta(\nabla)$ is the angle between $R_{c.m.}$ and the line parallel to the bonded C–C bond running through the center of mass of cyclopropane, and $\theta(SO_2)$ is the angle between the C_2 axis of SO₂ and $R_{c.m.}$.

 C_2 axis of SO₂ and the ab plane of cyclopropane in the ac plane of the complex. This orientation is shown in Fig. 1 with the S and C atoms in the ac symmetry plane and the O and H atoms straddling it. Once this is established, the geometry of the complex can be described by the three coordinates in Fig. 1: $R_{\rm c.m.}$, the distance between the centers of mass of the SO₂ and the cyclopropane; $\theta({\rm SO}_2)$, the angle between the C_2 axis of SO₂ and $R_{\rm c.m.}$; $\theta(\nabla)$, the angle subtended by $R_{\rm c.m.}$ and a line through the center of mass of cyclopropane parallel to the C-C bond to which the SO₂ is bonded. These angles define the tilt of the cyclopropane or the SO₂ from perpendicular to $R_{\rm c.m.}$, i.e., $\theta(\nabla) = \theta({\rm SO}_2) = 90^\circ$ corresponds to no tilt.

With the symmetry of the molecule deduced, a brief comment about the spectra of the deuterated isotopes is appropriate before proceeding with a discussion of the structure. The relative orientation of the cyclopropane and the SO₂ permits three different isomers to exist for both $C_3H_5D \cdot SO_2$ and $C_3H_4D_2 \cdot SO_2$. In each case the isotopic substitution could be on either of the CH₂ groups in the C-C bond which is closest to the SO₂ (labeled basal species), or on the CH₂ group opposite that bond (apical). In the case of $C_3H_5D\cdot SO_2$, due to difficulties with the synthesis, only one spectrum was observed before the sample was depleted. The rotational constants indicated that it was one of the basal isomers. For the $C_3H_4D_2\cdot SO_2$, two spectra were observed: one was consistent with the D₂ in the apical position and the other with D₂ in one of the basal positions. Although efforts were made to find the spectrum of the third species, it was not observed.

Little information could be gleaned from the moments of inertia of the normal isotopic species alone. Because the b coordinates of all the atoms are fixed by symmetry and the geometries of the monomers, only the P_{aa} and P_{cc} moments of inertia are useful for structure determination. The result is that $R_{c.m.}$ can be determined from I_b as approximately 3.73 Å but a series of correlated values for $\theta(SO_2)$ and $\theta(\nabla)$ are obtained from a single isotopic species.

In determining the structure by least-squares fitting of

the moments of inertia of all the isotopic species, a choice must be made about assignment of the $C_3H_4D_2\cdot SO_2$ isotopic spectra. The assignment of one spectrum to the species substituted at the *apical* position and one substituted at the *basal* position was unambiguous based on the isotope shifts. However, the latter could be assigned to the CD_2 group either at the S or O atom side of the SO_2 (see Fig. 1). The same ambiguity occurs in the location of the *basal* deuterium in the $C_3H_5D\cdot SO_2$. Both assignments were tried and they resulted in the two fits shown in Table VI. It was evident that the CH_2-CD_2 or CH_2-CHD bond is tilted slightly to the SO_2 and that the D_2 (or D_1) substitution occurs at the carbon closer to the SO_2 . The quality of the fits is similar because the coordinates of the deuterium are nearly identical in both structures.

Kraitchman's equations were used to calculate the positions of the substituted atoms. ²¹ They are compared with the values from the least-squared fit in Table VI. The coordinates determined for the $C_3H_5D\cdot SO_2$ species and the basal- $C_3H_4D_2\cdot SO_2$ species are the same and indicate the basal substitution with the CD_2 group nearer the SO_2 . The coordinates determined for the apical- $C_3H_4D_2\cdot SO_2$ species place the hydrogen atom near to the a axis and much further from the SO_2 . The substitution coordinates do not, however, distinguish between the two structures. The S and O coordinates from Kraitchman's equations are included in Table VI for completeness.

A similar ambiguity about the sign of an angle was encountered in the $C_2H_4 \cdot SO_2$ complex, where the tilt angle of the ethylene was difficult to determine.¹³ This was resolved by examining the change in the dipole moment projections

TABLE VI. Structural parameters and atomic coordinates obtained from least-squares fitting of moments of inertia and Kraitchman equations.

		Fit 1*	Fit 2 ^a	Kraitchman
R _{cm} /Å ^b		3.729(1)	3.729(1)	
$\theta(SO_2)/de$	g	73.2(1.7)	73.3(1.7)	
$\theta(\nabla)/\deg$		83.3(2.4)	96.8(2.4)	
$\Delta I_{\rm rms}$ /amu	$\mathring{\mathbf{A}}^2$	0.52	0.52	
S	a	1.37 Å	1.38 Å	1.40 Å
	b	0.0	0.0	0.0
	c	0.36	0.36	0.36
O	a	1.58	1.58	1.52
	b	1.24	1.24	1.24
	c	0.38	0.38	0.32
H ^c (basal)	а	1.38	1.39	1.26°
	b	0.91	0.91	0.89
	c	1.17	1.15	1.21
Hd (apical)) a	3.69	3.69	3.67°
	b	0.91	0.91	0.89
	c	0.19	0.15	0.26

^a Least-squares fit of 24 moments of inertia (A₂ symmetry state) from the eight isotopic species in Table IV. Fit 1 is preferred by the authors (see text).

^b See Fig. 1 for definition of coordinates.

^eH (basal) is the hydrogen at the carbon position in the CH₂-CH₂ bond which is closest to the SO₂ (see Fig. 2).

^dH (apical) is the hydrogen at the carbon most distant from the SO₂ (see Fig. 2).

Calculated from the 1,1-C₃H₄D₂·SO₂ species.

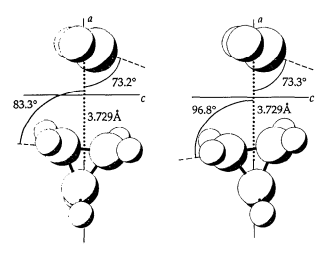


FIG. 2. Two structures of cyclopropane·SO₂ determined from least-squares fitting of the moments of inertia. The structure on the left with $\theta(\nabla) = 83.3^{\circ}$ is preferred.

upon isotopic substitution. A similar analysis was employed here. When a molecule is isotopically substituted, its principal inertial axes translate and rotate. The result of the rotation is a small change in the projections of the dipole moment on the principal axes. It is assumed that the change in the total dipole moment upon isotopic substitution is negligible.

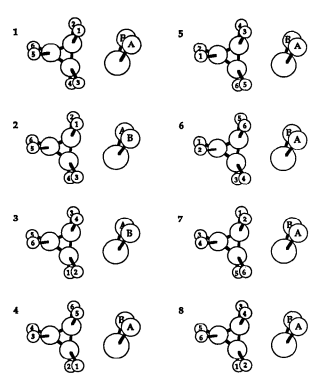


FIG. 3. Feasible permutation of nuclei which were considered to account for tunneling splittings in the spectrum of $C_3H_6 \cdot SO_2$. Permutations and likely pathways are $1 \rightarrow 2$ (ab), rotation of SO_2 about its local C_2 axis; $1 \rightarrow 3$ (ab)(14)(23)(56), inversion of SO_2 ; $1 \rightarrow 4$ (135)(246) and $1 \rightarrow 5$ (153)(264), rotation of cyclopropane about its C_3 axis; $1 \rightarrow 6$ (34)(16)(25), $1 \rightarrow 7$ (12)(36)(45), and $1 \rightarrow 8$ (14)(23)(56), rotations of cyclopropane about each of its C_2 axes.

TABLE VII. Predicted and observed dipole moments for basal-1,1- $C_3H_4D_2 \cdot SO_2$ for the two structures with $\theta(\nabla) = 83.3^\circ$ and $\theta(\nabla) = 96.8^\circ$.

	$\theta(\nabla) = 83.3^{\circ}$	$\theta(\nabla) = 96.8^{\circ}$	Obs.
μ_a/D	0.802	0.829	0.803(3)
$\mu_c/{ m D}$	1.478	1.463	1.482(3)

This should be a reasonable assumption for $C_3 H_4 D_2 \cdot SO_2$. The dipole moment of 1,1-C₃H₄D₂ has been measured as 0.011(5) D,²² and the total dipole moments of the $C_3H_4D_2 \cdot SO_2$ and $C_3H_6 \cdot SO_2$ differ by only 0.006(4) D. It is also assumed that the effect of vibrational averaging on the momomer and induced dipole moments is not dependent on the direction of $\theta(\nabla)$. A quantitative estimate of these effects is difficult. However, the success of the method in the $C_2H_4 \cdot SO_2$ problem, where several cross checks of the structure were available, suggests that averaging effects on the dipole moments can be neglected. The two options for the assignment of the basal CD2 spectrum rotate the axes in opposite directions. For both rotations the a- and c-dipole components were predicted and they are shown in Table VII. The direction of the dipole moment of the complex is selected such that it is dominated by the permanent dipole moment of SO₂.²³ The observed dipole components for basal-C₃H₄D₂·SO₂ indicate $\theta(\nabla) = 83.3^{\circ}$. The uncertainties for $R_{c.m.}$ and the tilt angles in Table VI are the statistical uncertainties arising from the fitting process. The structural parameters are the so-called r_0 values.²⁴ It is difficult to estimate how closely they approximate the equilibrium values due to the large amplitude vibrational motions in such complexes; it is probably reasonable to expect these values to be within ± 0.03 Å for $R_{c.m.}$ and $\pm 5^{\circ}$ for the tilt angles.

D. Internal rotation

The splitting in the c-type transitions signifies that a tunneling motion occurs between two or more equivalent configurations in the complex. In an attempt to determine the tunneling path, the feasible permutations of identical nuclei shown in Fig. 3 were considered. Possible tunneling paths for the permutations are the following: $1 \rightarrow 2$, rotation of SO_2 about its C_2 axis; $1 \rightarrow 3$, inversion of SO_2 through a C_{2v} intermediate structure; $1 \rightarrow 4$ and $1 \rightarrow 5$, rotation of cyclopropane about its C_3 axis; $1 \rightarrow 6$ and $1 \rightarrow 7$, rotation of cyclopropane about either of its C_2 axes including the bond CH_2 groups (the symmetry arguments are the same for both); $1 \rightarrow 8$, rotation of cyclopropane about its C_2 axis including the apical CH_2 group.

The $1\rightarrow 2$ and $1\rightarrow 3$ motions can be eliminated as the source of the tunneling doublets based on the spectrum of the normal isotopic species. If the tunneling path were $1\rightarrow 2$, two identical oxygen atoms (I=0) would be exchanged and half the levels would have zero nuclear spin weight. For the $1\rightarrow 3$ motion, the direction of the c-dipole moment reverses, resulting in c-dipole selection rules between the A_1 and A_2 symmetry states. Since the two sets of c-type transitions may be fit separately, the observed selection rules are inconsistent with this tunneling path.

The remaining tunneling paths involve the cyclopropane subunit, therefore, the deuterated isotopomers were instrumental in exploring them. From structural considerations alone, three isomers would be expected for the $1,1-C_3H_4D_2\cdot SO_2$: one with the CD₂ group in the basal position at the S of the SO₂, a second in the basal position at the O of the SO₂, and the third in the apical position. Referring to framework 1 in Fig. 3, these correspond to substitution in positions 3 and 4, positions 1 and 2, and positions 5 and 6, respectively. The tunneling paths under consideration, however, would result in different splitting patterns for the different isomers. For the $1 \rightarrow 4$ and $1 \rightarrow 5$ motions, rotation of cyclopropane about its C_3 axis, none of the $C_3H_4D_2 \cdot SO_2$ spectra should be split. This path would exchange the CD₂ group among three structurally inequivalent frameworks which have different moments of inertia. With the exception of H35Cl·H37Cl, tunneling doublets are generally not observed under those conditions. ²⁵ The $1 \rightarrow 6$ and $1 \rightarrow 7$ motions would produce a split spectrum for the basal CD2 group on the C_2 axis about which the cyclopropane rotates as this would exchange identical nuclei. However, unsplit spectra would be expected for the the CD₂ group at the other basal and the apical positions. For the $1 \rightarrow 8$ path, the CD₂ group in the apical position results in the exchange of identical atoms and tunneling doublets, while both the basal CD₂ groups are distinct and would be unsplit. The observation of splittings in the spectrum of the CD₂ group in the apical position and an unsplit spectrum for the CD₂ group in the basal position then indicates that the $1 \rightarrow 8$ motion is the correct tunneling path. This would produce nuclear spin statistical weights of approximately 1:1.3 in the normal isotopic form and, although the relative intensities of the two states could not be measured, they are estimated to be between 1:1 and 1:2.

It should be noted that the $1\rightarrow 8$ path is described as rotation of cyclopropane about its C_2 axis. This facilitates the discussion of the symmetry but does not necessarily imply that the SO_2 is a fixed framework on which the cyclopropane rotates. Although this division is common and a good description when the masses of the two parts are very different, such a separation is not obvious for $C_3H_6\cdot SO_2$. Perhaps a geared rotation of the two subunits against one another would be a more appropriate description. There is some suggestion of this since the magnitude of the tunneling splitting

is affected not only by isotopic substitution on the cyclopropane, but also on the sulfur dioxide. It is not trivial to estimate the barrier from the observed splittings and this was not attempted.

Finally, it is interesting that out of three possible structural isomers for $1,1-C_3H_4D_2\cdot SO_2$, only two were observed. While it is not uncommon under supersonically cooled beam conditions for only the isotopomer with the lowest zeropoint energy to be populated, 26 the observation of two out of three isomers is puzzling. The tunneling path seems to shed some light on this. Based on statistical arguments, it is equally likely that the SO₂ will bond to any of the three C-C bonds in the 1,1-C₃H₄D₂ species when the complexes are formed in the nozzle. If the two bond positions have different zeropoint energies, a path exists for cooling to the lower energy basal position through the tunneling coordinate, i.e., the internal rotation pathway is a means for equilibrating between the two CD₂ bond isomers. However, there is no ready pathway for cooling between the basal and apical positions if the barrier to exchange for this motion is very high, and both of these forms remain populated at the level determined by the beam kinetics.

IV. DISCUSSION

The structure of $C_3H_6 \cdot SO_2$ is similar to the structures of C₂H₄·SO₂ and C₂H₂·SO₂ in that the S atom is closest to the pseudo- π system of the cyclopropane. The symmetry, however, is different with the dihedral angle (α) between the C_2 axis of the SO_2 and the C-C bond equal to 0° for $C_3H_6 \cdot SO_2$ compared to 90° for $C_2H_4 \cdot SO_2$ and $C_2H_2 \cdot SO_2$. To explore whether this could be attributed to electrostatic considerations alone, the distributed multipole (DMA) model of Buckingham and Fowler was employed.²⁷ Distributed multipoles for SO₂ were taken directly from Buckingham and Fowler, while those for cyclopropane were calculated using the CADPAC program with a DZP (double zeta plus polarization) basis set.²⁸ The coordinates and multipoles are listed in Table VIII. The DMA gives a minimum of energy at the $\alpha = 0^{\circ}$ geometry with $\theta(SO_2) = 90^{\circ}$ and $\theta(\nabla)$ = 75°, which are considerably different from the experimental angles. The well seems to be very shallow, however, and the experimental geometry $\theta(SO_2) = 74^{\circ}$, $\theta(\nabla) = 84^{\circ}$ is only 13 cm⁻¹ higher in energy. The lowest-energy structure for $\alpha = 90^{\circ}$ is 75 cm⁻¹ higher in energy.

TABLE VIII. Cyclopropane distributed multipole moments in atomic units.

Site	x	y	z	\boldsymbol{q}	$\mu_x^{\ a}$	$\mu_{_{y}}$	μ_z	$\theta_{\star\star}$	$ heta_{yy}$	$ heta_{zz}$	$ heta_{yz}$
C1	0.0	0.827	1.432	0.0845	0.0	0.0908	- 0.1711	- 0.1331	0.4772	- 0.3441	0.711
C2	0.0	0.827	-1.432	0.0845	0.0	-0.0908	0.1711	-0.1331	0.4772	- 0.3441	- 0.711
C3	0.0	- 1.654	0.0	0.0845	0.0	0.1975	0.0	- 0.7547	0.8879	-0.1331	0.0
H1	1.724	1.371	2.375	-0.0423	0.1487	0.0525	0.0909	0.0226	- 0.0130	- 0.0096	0.0
H2	1.724	1.371	-2.375	-0.0423	0.1487	0.0525	- 0.0909	0.0226	-0.0130	- 0.0096	0.0
H3	— 1.724	- 2.742	0.0	0.0423	-0.1487	- 0.1050	0.0	0.0226	- 0.0109	- 0.0116	0.0
H4	-1.724	1.371	2.375	-0.0423	- 0.1487	0.0525	0.0909	0.0226	- 0.0130	0.0096	0.0
H5	— 1.724	1.371	-2.375	0.0423	-0.1487	0.0525	0.0909	0.0226	- 0.0130	- 0.0096	0.0
H6	1.724	- 2.742	0.0	-0.0423	0.1487	-0.1050	0.0	0.0226	- 0.0109	- 0.0116	0.0

^a Dipole moment directions are from regions of negative to positive charge.

TABLE IX. Comparison of the complexes of cyclopropane, acetylene, and ethylene.

		Cyclopropane	Acetylene	Ethylene
SO ₂	R/Ų	3.203	3.359	3.446
2	k/mdyne Å - 1b	0.059	0.047	0.057
	€/cm - le	652	390	490
	μ_a (ind)/ D^d	0.349	0.326	0.289
	$\mu_{\rm c}({\rm ind})/{\rm D}^{\rm d}$	0.095	0.063	0.072
	$\mu_I(\text{ind})/D^d$	0.361	0.332	0.298
	Reference		14	13
HCl	R/Å	3.567	3.699	3.724
.,	k ∕mdyne Å - ¹	0.087	0.067	0.061
	€/cm ⁻¹	959	614	575
	Reference	8	5	2
HCN	R/Å	3.475	3.656	3.711
	k ∕mdyne Å ^{- 1}	0.062	0.053	0.046
	€/cm ⁻¹	862	642	575
	Reference	9	6	3
HF	R/Å	3.021	3.121	3.143
	$\mu_{\scriptscriptstyle ind}/\mathrm{D}$	0.78	0.65	0.67
	Reference	7	4	1

^{*} R is the distance from hydrocarbon C-C bond to nearest heavy atom.

The appropriate hydrocarbon X distances, stretching force constants, well depths, and induced dipole moments for the cyclopropane, ethylene, and acetylene complexes with SO2, HCl, HCN, and HF are shown in Table IX. The pseudodiatomic stretching force constant for C₃H₆·SO₂ was calculated using Millen's model²⁹ as 0.059 mdyne/Å and from this the binding energy was estimated as $650 \, \text{cm}^{-1}$. The induced dipole moments for the HF complexes were taken from Nelson, Fraser, and Klemperer¹¹ $\mu_{\text{ind}} = \mu - \langle \cos \theta \rangle \mu_{\text{HF}}$, where $\langle \cos \theta \rangle$ is approximated by $\langle \cos^2 \theta \rangle^{1/2}$ which is determined from hyperfine interaction constants. For the SO₂ complexes an estimate of the averaging effects from bending motions on the dipole moments is not so straightforward as there are no hyperfine interactions giving information on the SO₂ bending vibration. These effects were neglected and the induced moments were taken as the difference between the observed dipole components and the projections of the SO₂ permanent dipole moment on the principal axes of the complex.

The SO₂ complexes exhibit the same decrease in sulfur to the C-C bond center distance previously noted for the hydrogen-bonded complexes. There is an increase in induced dipole moment in the series ethylene, acetylene, and cyclopropane, while in the HF complexes, the induced dipole moment is about the same for $C_2H_2 \cdot HF$ and $C_2H_4 \cdot HF$ and larger for C₃H₆·HF. The force constants and binding energies are also greater for $C_3H_6 \cdot SO_2$ than $C_2H_2 \cdot SO_2$. The values for C₂H₄·SO₂ are anomalously high, most probably because of perturbations in the spectrum due to a tunneling motion which contaminates the distortion constants. The difficulty with the C₂H₄·SO₂ distortion constants aside, it is reasonable to conclude from the other data that $C_3H_6 \cdot SO_2$ is the most strongly bound of the three, consistent with the finding of Legon and Millen that C₃H₆ is the best nucleophile.12b

Legon and Millen's model for determining nucleophilicities and electrophilicities has been successful in interpreting hydrogen-bonded complexes, but it has not yet been applied to other weak complexes. Because many SO_2 complexes have been characterized in recent years, the $C_xH_y\cdot SO_2$ complexes seemed an ideal situation to test it. The formula for relating nucleophilicities and electrophilicities to the pseudodiatomic stretching force constant is 12a

$$k_{\sigma} = cNE$$

where k_{σ} is the force constant, c is a proportionality constant (equal to about 0.25), N is the nucleophilicity of the nucleophile, and E is the electrophilicity of the electrophile. Using N=6.4 for cyclopropane as determined by Legon and Millen, E for SO_2 was calculated as 3.7. This was then used to predict k_{σ} for a number of SO_2 complexes with molecules for which the nucleophilicities have been calculated. These are shown in Table X along with the experimentally deter-

TABLE X. Pseudodiatomic stretching force constants for SO₂ containing complexes.

	$k_a/10^{-2}$ r			
	Predicted ^a	Observed ^b	Reference	
C ₃ H ₆ ·SO ₂ ^c	5.9	5.9		
$C_2H_2 \cdot SO_2$	4.7	4.7	14	
H ₂ O·SO ₂	9.3	8.4	32	
H ₂ S·SO ₂	4.4	5.3	33	
HCN·SO ₂	6.7	2.7	34	
(CH ₃) ₂ O·SO ₂	10.4	6.8	35	

^aPredicted using nucleophilicity from Ref. 12(b) and electrophilicity of $SO_2 = 3.7$ (see text).

b k is the pseudodiatomic stretching force constant.

 $[\]epsilon$ is the pseudodiatomic well depth.

d Induced dipole moment/D (see text).

^b Obtained from D_J and pseudodiatomic approximation (Ref. 29).

 $^{^{\}circ}$ C₃H₆ $^{\circ}$ SO₂ was used to determine the electrophilicity of SO₂; therefore, the match is required to be exact.

mined k_{σ} . There is excellent agreement for $C_2H_2 \cdot SO_2$ and quite good agreement for $SO_2 \cdot H_2O$ (Ref. 30) and $H_2S \cdot SO_2$ (Ref. 31). The agreement for $HCN \cdot SO_2$ (Ref. 32) and $(CH_3)_2O \cdot SO_2$ (Ref. 33) is rather poor. These complexes, however, are unusual in that the two subunits are somewhat closer than the sum of their van der Waals radii. In the case of $(CH_3)_2O \cdot SO_2$ the interaction may involve some charge-transfer interaction. This raises questions about the validity of the psuedodiatomic approximation, but also suggests that the nucleophilicity model is perhaps only applicable to complexes which are bound by primarily electrostatic forces. A larger data set is needed to test whether these complexes represent an anomaly or whether the model does not readily transfer to non-hydrogen-bonded weak complexes.

ACKNOWLEDGMENTS

This work was supported by a grant from the National Science Foundation. We are grateful for time granted at the San Diego Supercomputing Center. A.M.A. acknowledges the support of a Regents-Baer Fellowship at the University of Michigan.

- ¹ J. A. Shea and W. H. Flygare, J. Chem. Phys. 76, 4857 (1982).
- ²S. G. Kukolich, P. D. Aldrich, W. G. Read, and E. J. Campbell, Chem. Phys. Lett. **90**, 329 (1982).
- ³S. G. Kukolich, W. G. Read, and P. D. Aldrich, J. Chem. Phys. 78, 3552 (1983).
- ⁴ W. G. Read and W. H. Flygare, J. Chem. Phys. 76, 2238 (1982).
- ⁵ A. C. Legon, P. D. Aldrich, and W. H. Flygare, J. Chem. Phys. 75, 625 (1981).
- ⁶P. D. Aldrich, S. G. Kukolich, and E. J. Campbell, J. Chem. Phys. 78, 3521 (1983).
- ⁷L. W. Buxton, P. D. Aldrich, J. A. Shea, A. C. Legon, and W. H. Flygare, J. Chem. Phys. **75**, 2681 (1981).
- ⁸ A. C. Legon, P. D. Aldrich, and W. H. Flygare, J. Amer. Chem. Soc. **104**, 1486 (1982); P. D. Aldrich, S. G. Kukolich, E. J. Campbell, and W. G. Read, *ibid.* **105**, 5569 (1983).
- ⁹S. G. Kukolich, J. Chem. Phys. 78, 4832 (1983).

- ¹⁰ C. A. Coulson and W. E. Moffit, J. Chem. Phys. 15, 151 (1947); C. A. Coulson and W. E. Moffit, Philos. Mag. 40, 1 (1949).
- ¹¹ D. D. Nelson, Jr, G. T. Fraser, and W. Klemperer, J. Chem. Phys. 82, 4483 (1985).
- (a) A. C. Legon and D. J. Millen, J. Amer. Chem. Soc. 109, 356 (1987);
 (b) A. C. Legon and D. J. Millen, J. Chem. Soc. Chem. Comm. 986 (1987).
- ¹³ A. M. Andrews, A. Taleb-Bendiab, M. S. LaBarge, K. W. Hillig II, and R. L. Kuczkowski, J. Chem. Phys. 93, 7030 (1990).
- ¹⁴ A. M. Andrews, K. W. Hillig, R. L. Kuczkowski, A. C. Legon, and N. W. Howard, J. Chem. Phys. 94, 6946 (1991).
- ¹⁵T. J. Balle and W. H. Flygare, Rev. Sci. Instrum. 52, 33 (1981).
- ¹⁶ K. W. Hillig II, J. Matos, A. Scioli, and R. L. Kuczkowski, Chem. Phys. Lett. 133, 359 (1987).
- ¹⁷ J. K. G. Watson, J. Chem. Phys. 46, 1935 (1967).
- ¹⁸ K. Tanaka, H. Ito, K. Harada, and T. Tanaka, J. Phys. Chem. 80, 5893 (1984).
- ¹⁹ Y. Endo, M. C. Chang, and E. Hirota, J. Mol. Spec. 126, 63 (1987).
- ²⁰ M. D. Harmony, V. W. Laurie, R. L. Kuczkowski, R. H. Schwendeman, D. A. Ramsey, F. J. Lovas, W. J. Lafferty, and A. G. Maki, J. Chem. Phys. Ref. Data 8, 619 (1979).
- ²¹ J. Kraitchman, Am J. Phys. 21, 17 (1953); A. Chutjian, J. Mol. Spectrosc. 14, 361 (1964).
- ²²O. Böttcher, N. Heineking, and D. H. Sutter, J. Mol. Spec. 139, 236 (1990).
- ²³ F. J. Lovas, J. Chem. Phys. Ref. Data, 14, 395 (1985).
- ²⁴ R. H. Schwendeman, Critical Evaluation of Chemical and Physical Structural Information, edited by D. R. Lide and M. A. Paul (National Academy of Sciences, Washington, D. C., 1974), pp. 74-115.
- ²⁵ N. Ohashi and A. S. Pine, J. Chem. Phys. 81, 73 (1984).
- ²⁶ T. R. Dyke, B. J. Howard, and W. Klemperer, J. Chem. Phys. **56**, 2442 (1972); H. S. Gutowsky, C. Chuang, J. D. Keen, T. D. Klots, and T. Emilsson, J. Chem. Phys. **83**, 2070 (1985).
- ²⁷ A. D. Buckingham and P. W. Fowler, Can. J. Chem. 63, 2018 (1985).
- ²⁸ R. D. Amos and J. E. Rice, Cambridge Analytical Derivatives Package, Issue 4.0, Cambridge, 1987.
- ²⁹ D. J. Millen, Can. J. Chem. 63, 1477 (1985).
- ³⁰ K. Matsumura, F. J. Lovas, and R. D. Suenram, J. Chem. Phys. 91, 5887 (1989).
- ³¹ R. E. Bumgarner, D. J. Pauley, and S. G. Kukolich, J. Chem. Phys. 87, 3749 (1987); D. J. Pauley, and S. G. Kukolich, *ibid.* 93, 1487 (1990).
- ³² E. J. Goodwin and A. C. Legon, J. Chem. Phys. 85, 6828 (1986).
- ³³ J. J. Oh, K. W. Hillig II, and R. L. Kuczkowski, Inorg. Chem. (to be published).