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Infrared Spectra of Molecules and Materials

of Astrophysical Interest

Grant NGL-41-002-003 Quarterly Progress Report Report Number 36 June 1976

Office of Grants and Research Contracts Office of Space Science and Applications National Aeronautics and Space Administration Washington, D. C. 20546

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SUMMARY OF PROGRESS

We have been studying the vibrational spectra from 4000 to 33 cm⁻¹ of several molecules which may be present in the atmosphere of the Jovian planets or exist in outer space. These studies have been made to provide vibrational frequencies which can be used to: (1) determine the composition of the cloud covers of several of the planets, (2) provide structural information under favorable circumstances, (3) provide necessary data from which accurate thermodynamic data can be calculated, and (4) furnish information as to the nature of the potential energy function of the molecules and forces acting within them.

Some of the molecules which we have studied can be produced photochemically from methane, ammonia, and hydrogen sulfide which are thought to be constituents of the planets with reducing atmospheres. Some of the compounds will polymerize under ultraviolet radiation and drop out of the atmospheres. However, planets with a hot base, like that of Jupiter, may rebuild molecules destroyed photochemically. Therefore, we have used these criteria in selecting the compounds which we have studied.

Gerald P. Kuiper¹ has pointed out that the Jovian atmosphere is expected to contain H₂, He, N₂, H₂O, NH₃, CH₃, Ar and possibly SiH₄. He has also listed a number of other gases that should be considered because they are composed of fairly abundant atomic species and have boiling points below 120°C (see Table 8, pg. 349-350 of reference 1). He has also pointed out that until more is known about the atmospheres of the planets, it is useful to keep a fairly large number of possible constituents in mind in planning further spectroscopic work.

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In our initial work on the vibrational spectra of molecules of astrophysical interest, we studied hydrazine,² methylamine,³ as well as several substituted hydrazines.⁴⁻⁸ Recently both ethane and acetylene have been found in the atmosphere of Jupiter.⁹ It is expected that substituted ethanes will also be eventually found in some of the planetary atmospheres. In fact, ethanol has been found in the Sagittarius Bl2 cloud of dust and gas which is near the center of the Milky Way.¹⁰ In addition, molecules such as acetaldehyde (CH₃CHO), methanol (CH₃OH), dimethylether (CH₃)₂O, formic acid (HCOOH), have been identified in outer space. As a continuation of our earlier studies on the low frequency torsional motions of dimethylether¹¹ we have recorded the high resolution microwave spectra of this molecule along with the two isotopic derivatives, CH₃OCD₃ and (CD₃)₂O and reported this study in the last progress report. We have now completed the relatively high resolution far infrared study along with a study of the low frequency Raman spectra of the gas.

Several years ago the gas phase far infrared spectra of molecules of the type $(CH_3)_2X$ and $(CD_3)_2X$ were published. Some of these spectra showed a very complex pattern in the region where the methyl torsional transitions are expected. Tuazon and Fateley¹² reported in 1971 the far infrared spectrum of CD_3OCH_3 along with the already measured species $(CH_3)_2O$ and $(CD_3)_2O$. The spectrum of this compound is similarly complex. These authors took the most likely band in the spectra to obtain an estimate of the barrier height, but detailed interpretation of the complex features were not presented. To our knowledge, only one Raman spectrum of gaseous CH_3OCH_3 has been reported. In 1957, Taylor and Vidale¹³ reported two Raman bands at 480 and 450 cm⁻¹ as overtones of both torsional modes.

In order to get better barrier information on these isotopic species of dimethylether and eventually to explain the complicated structure of their

far infrared spectra, the Raman spectra of these isotopes were measured in the gas phase on a Cary Model 82 instrument with a Spectra Physics 171 argon ion laser. For the same purpose, the far infrared spectra were remeasured with 0.5 cm⁻¹ resolution with a Digilab FTS 15B interferometer. In Figure 1 are shown the far infrared spectrum of the normal isotope. The band at 240 $\rm cm^{-1}$ is clearly asymmetric, and the band at 223 cm^{-1} is split into at least three components at 223.2, 224.8 and 226.0 cm⁻¹. Minor bands occur around 200 cm⁻¹. The 240 cm⁻¹ band has been interpreted previously as the $0 \rightarrow 1$ transition of the infrared active torsion. But the 223 cm⁻¹ band is not the $1 \rightarrow 2$ transition of the same mode, as one might believe. If this were the case, the overtone band should occur at 463 cm⁻¹. The Raman spectrum in Figure 2 clearly shows that there is no band at this frequency. There are bands at 480 and 450 $\rm cm^{-1}$, as previously reported, in addition to the bending mode at 412 cm⁻¹. Knowing the far infrared spectrum, the 480 $\rm cm^{-1}$ band may really be interpreted as the overtone of the infrared active torsion, assuming that the 240 cm⁻¹ band consists of at least two torsional transitions. The infrared inactive mode was estimated by Fateley and Miller¹⁴ and several normal coordinate calculations to be at about 200 cm⁻¹. Therefore, the 395 cm⁻¹ band could well be the overtone of this torsion. All Raman bands are strongly polarized. In the torsional region, only a very broad, very weak Raman signal with no structure was observed.

In order to check this preliminary assignment and to determine a better potential function, a computer program has been developed. It solves the eigenvalue problem of the internal Hamiltonian of any model with two internal rotors of C_{3v} symmetry (see Table I). Since the models for dimethylether d_0 , d_3 and d_6 have higher symmetry than the most general model, some of the terms in this operator vanish or have equal coefficients. The program is set up in the direct product basis of two exponential basis sets and uses a diagonalization subroutine for complex Hermitian matrices. It first solves the eigenvalue

problems of the single top models (first and second line, respectively) and transforms the matrices of the operators \hat{P}_i , cos $3\tau_i$ and sin $3\tau_i$ into their eigenbasis. Using direct product calculus, the Hamiltonian matrix is set up for the coupled problem and diagonalized. A standard iteration routine allows the fitting of observed transition frequencies and determination of the potential coefficients. It further calculates the squares of the matrix elements of $\cos 3\tau_i$ and $\sin 3\tau_i$ and weights them with the Boltzman population factor. These numbers are used to identify the energy levels with respect to the symmetry of the two top model and to obtain rough estimates of relative intensities of the transitions. With this program, the fit in Table II was obtained. It looks quite reasonable. However, it does not explain the weak Raman band at 363 cm^{-1} . Another weak Raman band at 381 cm^{-1} may eventually be interpreted as a line belonging to a different torsional substate of the 385 $\rm cm^{-1}$ band, although the calculated splitting is not larger than 2 cm^{-1} . Another point which is not quite satisfactory is the fact that the intensity ratio of the 450 and 480 ${\rm cm}^{-1}$ bands is about 1:2, the calculated ratio, however, is about 1:10. For the present, this fit seems to be the best we can find and the corresponding potential coefficients are given in the lower half of Table II. For comparison, the results of a recent microwave investigation of ground state and two torsionally excited states splittings are given. Instead of using $V_{30} = V_{03}$, V_{33} and V_{33}^{i} to define the potential function, an identical function may be written as

 $2V = V_{30}(1-c3\tau_0)(1-c3\tau_1) + V_{eff}(1-c3\tau_0c3\tau_1) + V_{33}^{i}s3\tau_0s3\tau_1$. The coefficients in this equation have lower dispersions and are much less correlated to each other. They are also given in Table II.

In Figure 3 is shown the far infrared spectrum of CD_3OCH_3 . Due to the higher resolution, it shows a lot more features than the previously published spectrum. Note the band series at 162.0, 154.5, 145.2 and 133.2 cm⁻¹. They

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may be assigned to the $0 \rightarrow 1$, $1 \rightarrow 2$, $2 \rightarrow 3$ and $3 \rightarrow 4$ transitions of the CD₃ torsional mode. In the region of the "light" methyl torsion, the interpretation is not straightforward. The main features are at 224.0, 222.0, 218.2, 210.7 and 204.9 $\rm cm^{-1}$, all but the first two are rather broad and probably consisting of more than one torsional transition. The Raman spectrum, shown in Figure 4 is similarly complex in the overtone region. The bending mode at 373 $\rm cm^{-1}$ is relatively weak compared with the light isotope. Bands of similar intensities appear at 384 and 380 cm^{-1} , the former being assigned to the lowest combination transition of both torsions. The weak band at 316.5 cm^{-1} is assigned as the $0 \rightarrow 2$ transition of the CD_3 torsion, since its frequency is the sum of 162.0 and 154.5 cm^{-1} which corresponds to two infrared transitions. The 442 cm^{-1} band is the sum of 224 and 218 cm⁻¹; therefore, this band is assigned to the $0 \rightarrow 2$ transition of the light methyl torsion. We suspect Fermi resonance type interactions to be responsible for the particular intensity pattern in the 380 ${
m cm}^{-1}$ region, although a good frequency fit could be obtained without this assumption. The results of the fit are represented in Table III. This fit again is reasonable, although it does not explain everything, for instance, the infrared absorption at 210 cm⁻¹. Again, the potential function can be expressed by different coefficients. One choice is to use V_s , V_d and V_{seff} in the form

 $2V = V_s(1-c3\tau_0)(1-c3\tau_1) + V_d(cos3\tau_2-c3\tau_1) + V_{seff}(1-c3\tau_0c3\tau_1) + V_{33}s3\tau_0s3\tau_1$. The coefficients are also given in this Table III, the correlations are weaker than in the original set and they are better determined.

The Raman spectrum of CD₃OCD₃ in Figure 5 shows, besides the bending mode, at least 15 different polarized lines. At present, no interpretation can be given since the analysis has not been completed yet.

The agreement between the potential coefficients of different isotopes is not very exciting. Similarly, there are certain discrepancies with the results of the microwave splittings analysis. Several reasons may be responsible for

these facts. The present assignment may not be correct since, as already pointed out, several features of the spectra can not be explained, although we tried ruite a number of possiblities. The microwave analysis was performed using splittings in the ground state and two excited states. We think that from this information, at most, three and probably only two potential coefficients can be determined. The torsional analysis, however, includes transitions involving energy levels almost to the top of the effective barrier, so that the information about the higher parts of the potential surface should be much better than in the microwave investigation. Despite these doubts, the cos-cos coupling term is definitely a large number and it has definitely no relation at all with the sin-sin coefficient which has been proposed in some earlier work.

Table I

Hamiltonian and Basis Functions

$$\begin{split} \hat{H} &= \frac{1}{29} g^{44} \hat{p}_0^2 + \frac{1}{2} [V_{30}(1 - \cos 3\tau_0) + V_{60}(1 - \cos 6\tau_0) + V_{60}' \sin 6\tau_0] \\ &+ \frac{1}{29} g^{55} \hat{p}_1^2 + \frac{1}{2} [V_{03}(1 - \cos 3\tau_1) + V_{06}(1 - \cos 6\tau_1) + V_{06}' \sin 6\tau_1] \\ &+ g^{45} \hat{p}_0 \hat{p}_1 + \frac{1}{2} [V_{33}(\cos 3\tau_0 \cos 3\tau_1 - 1) + V_{33}' \sin 3\tau_0 \sin 3\tau_1] \\ &+ V_{33}' \sin 3\tau_0 \cos 3\tau_1 + V_{33}'' \cos 3\tau_0 \sin 3\tau_1] \end{split}$$

$$\{ \Phi_{mn}(\tau_0, \tau_1) \} = \{ \phi_m(\tau_0) \} \times \{ \phi_n(\tau_1) \}$$
$$\phi_{\ell}(\tau_j) = (2\pi)^{-\frac{1}{2}} \exp(i\ell\tau_j)$$

(b) and a set of the set of th

$$\begin{array}{c} CH_{3}OCH_{3} \\ CD_{3}OCD_{3} \end{array} \begin{cases} g^{44} = g^{55}; V_{30} = V_{03}; V_{60} = V_{06} \\ V_{60}^{\prime} = V_{06}^{\prime} = V_{33}^{\prime\prime} = V_{33}^{\prime\prime\prime} = 0 \\ CD_{3}OCH_{3} \\ V_{60}^{\prime} = V_{06}^{\prime} = V_{33}^{\prime\prime} = V_{33}^{\prime\prime\prime} = 0 \end{array}$$

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			Tab	le II			·
		Fit	of Torsional Fr	equencies for (CH ₃ OCH ₃		
		g ⁴⁴ = g	⁵⁵ = 13.5644 cm	-7 g ⁴⁵ = -2	2.7381 cm ⁻¹		
Transition	Obs(cm ^{-l})	Calc(cm ⁻¹) Obs-Calc	Transition	Obs(cm ⁻¹)	Calc(cm ⁻¹)	Obs-Calc
1 → 3	241.0	213.8	-2.8	1→6	481.3	484.2	-2.9
3→6	239.9	240.4	-0.5	2 → .9	450.2	446.9	3.3
2 + 5	223.2	222.7	0.5] → 4	395.5	397.3	-1.8
5 → 9	226.0	224.2	1.8	2 → 7	385.2	383.8	1.4
4 → 8	200.7	199.7	1.0				
5 → 9	224.8	223.4	1.4 j The	ese transitions	; were not fi	itted. They	are assigned
		198.1		different tors			
Standard deviation of frequencies 2.51 cm ⁻¹							
Potential Coefficients (cm ⁻¹) ^a			Microwave	Derived Pote	ential Coeffi	cients	Microwave ^b
V ₃₀ =V ₀₃ 1132.91±37.52		909.05±0.25	5 V _{eff} = V ₃₀ -V ₃₃ 918.66±10.95 90		909.05±0.25		
V ₂₂	214.25±47.09						
۷ ² 3	20.92 ± 8.05		10.12±1.60				
V ₆₀ =V ₀₆	20.92±8.05 0(not varied)		O(not varied)				

 $p_{1,1} = p_{1,1} + p_{1,2} + p_{2,1} + p_{2,2} + p_{3,2} + p_{3$

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^a Error limit = dispersion of parameter

^b J. R. Durig, Y. S. Li and P. Groner, accepted for publication (J. Mol. Spectry).

Table III

Fit of Torsional Frequencies for CD30CH3

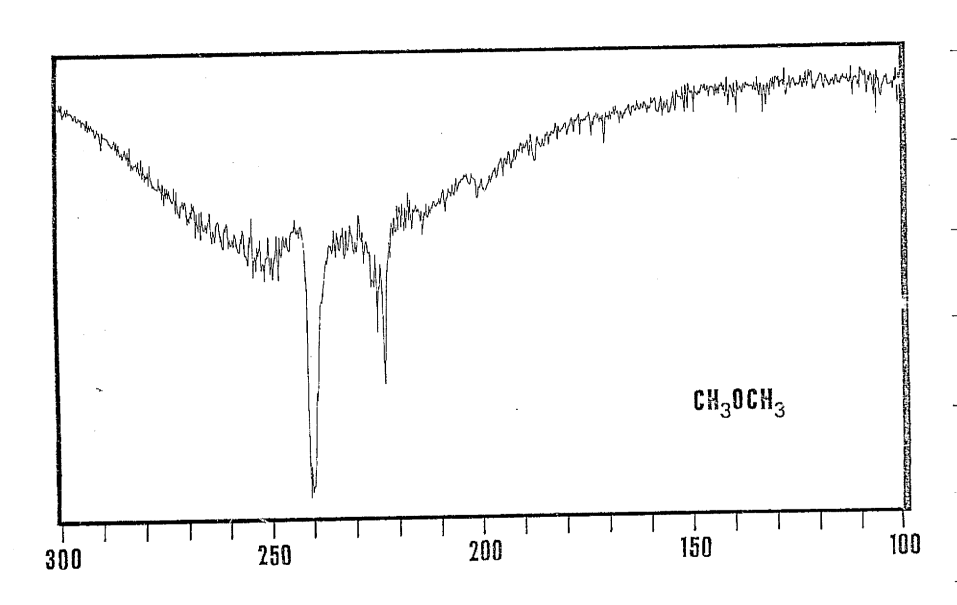
	g ⁴⁴ =	= 13.0973 cm	-1 g ⁵⁵ = 8.	.1872 cm ⁻¹	$g^{45} = -2.476$	64 cm ⁻¹	
Transition	Obs(cm ⁻¹)	Calc(cm ⁻¹)	Obs-Calc	Transition	Obs(cm ⁻¹)	Calc(cm ⁻¹)	Obs-Calc
l → 2	162.0	161.8	0.2	1 → 3	224.0	225.9	-1.9
2 → 4	154.5	155.3	-0.8	3 → 6	218.2	218.3	-0.1
4 → 7	145.2	145.6	-0.4	1→6	442.0	444.2	-2.2
7 → 10	133.0	131.8	1.2	3 -> 11	435.0	433.6	1.4
1 → 4	316.5	317.2	-0.7	2 → 5	222.0	221.9	١.0
1 + 5	384.0	383.7	0.3	5 → 9	204.9	202.2	2.7
2 → 8	380.0	381.1	-1.1	6 → 11 [*]	215.8 sh	215.3	0.5
3 → 9	361.5	360.0	1.5	* not fi	tted		
Standard	deviation of	f frequencies	1.55 cm ⁻¹				
Potential Coefficients (cm ⁻¹) ^a M			Microwave ^b	Derived Potential Coefficients			Microwave ^b
۷ ₃₀	1313.52±30.31		10.45±0.17	$V_{s} = \frac{1}{2}(V_{30} + V_{03})$ 1285.97±30.08			903.81±0.65
V ₀₃	1258.41±30.45		97.18±1.21	$V_d = \frac{1}{2}(V_{30} - V_{03})$ 27.55 ± 4.24		5±4.24	6.64±0.56
V ₃₃	417.11±35.54 0		(not varied)	$V_{seff} = V_s$	-V ₃₃ 868.8	5±6.14	903.81±0.65
V'33	34.22 ± 7.99 0		(not varied)	$V_{30eff} = V_3$		1±6.99	910.45±0.17
V ₀₆ =V ₆₀	0(not vari	ied) O	(not varied)	$V_{03eff} = V_{0}$		0 ± 7.92	897.18±1.21

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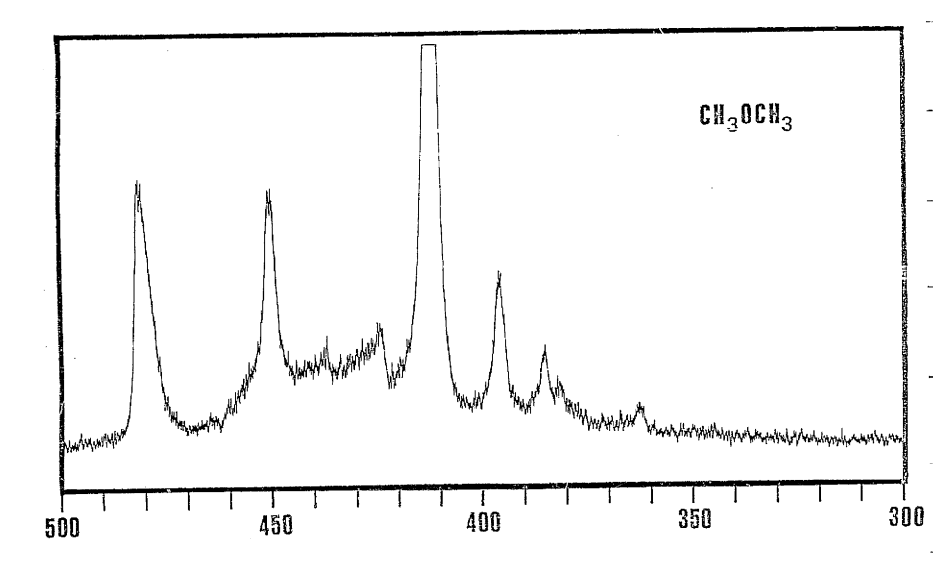
^a Error limit = dispersion of parameter

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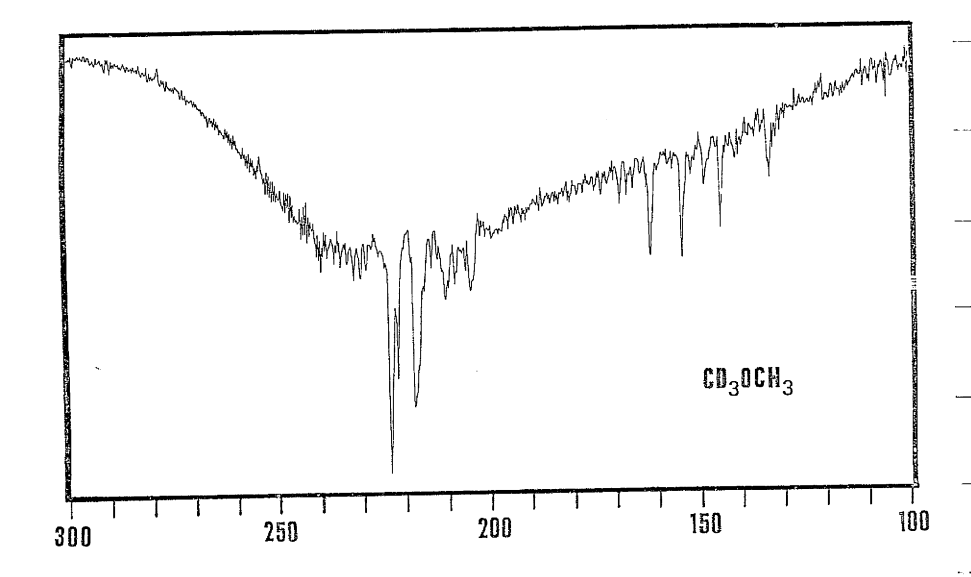
^b J. R. Durig, Y. S. Li and P. Groner, accepted for publication (J. Mol. Spectry).



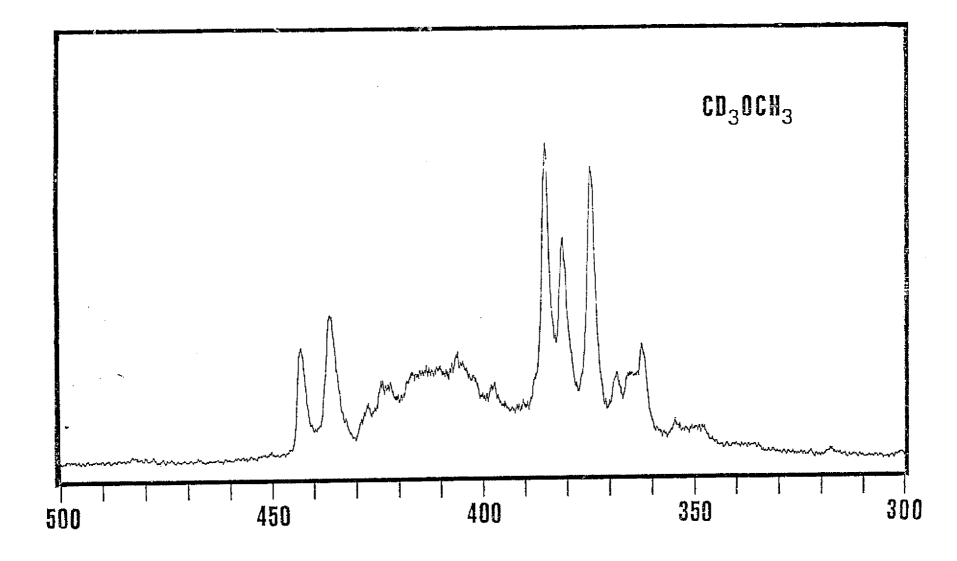
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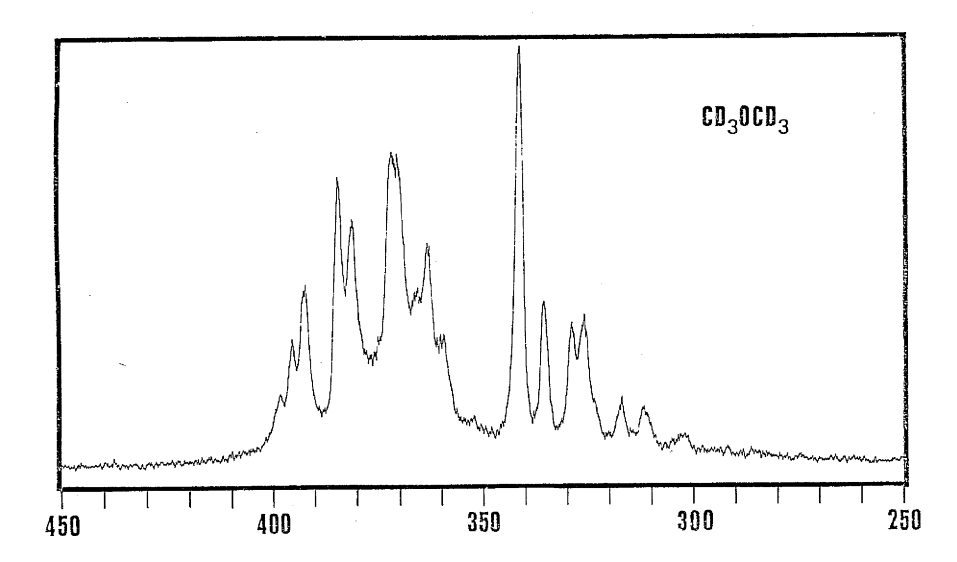
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Although this study is not yet complete, we should be able to provide a complete manuscript by the next progress report time.

For several progress reports we have mentioned our work on methylchloroformate and anticipated that this work would soon be finished. We are pleased to report that a manuscript has been submitted to the Journal of Molecular Spectroscopy on this molecule.

The structure of methylchloroformate has been the subject of considerable interest. Four dipole moment studies have been reported. $^{15-18}$ From two of these 15,17 the results were interpreted in terms of a planar <u>s-cis</u> structure. The data from the other dipole moment studies 16,18 were interpreted in terms of a planar <u>s-trans</u> structure.

Three analyses of the vibrational spectra have been reported. ¹⁹⁻²¹ These analyses were all in terms of a molecular model of C_s symmetry and were consistent with an <u>s-trans</u> structure. An electron diffraction study of the vapor of methylchloroformate was performed.²² The data yielded only an <u>s-trans</u> structure that was slightly non-planar with the methyl group approximately 20° out of the plane of the other heavy atoms.

An investigation of the infrared spectrum and of the proton NMR spectrum of methylchloroformate in solution was reported.²³ The authors found evidence for the coexistence of both the <u>s-cis</u> and the <u>s-trans</u> conformations and indicated that the <u>s-cis</u> was more stable. From a study comparing observed and calculated ³⁵Cl nuclear quadrupole resonance frequencies the authors²⁴ concluded that the s-trans was the most likely stable structure in the solid.

A microwave spectroscopic investigation²⁵ of natural abundance 35 Cl and 37 Cl isotopic species of methylchloroformate showed that the planar <u>s-trans</u> molecular structure was most consistent with the data. Small amounts of a second rotamer, however, could not be precluded.

Therefore, in an effort to clear up the ambiguities in the conformational preferences of methylchloroformate and to provide more complete structural data, the investigation of the microwave, Raman and far infrared spectra of the vapor were undertaken. The results of this study can be summarized by the abstract which follows of the paper that has been submitted for publication. Abstract: The microwave spectra of six isotopic species of methylchloroformate, $C1C0_2CH_3$, have been recorded from 18.0 to 40.0 GHz. Structural parameters have been determined and it is shown that the only stable conformer at ambient temperature is the s-trans. From the Stark effect, the dipole moment components were determined to be $|\mu_a|$ = 1.7 ± 0.2, $|\mu_b|$ = 1.0 \pm 0.2, and $|\mu_t|$ = 2.0 \pm 0.2D. The Raman and far infrared spectra of the vapor are reported. Four cases of Fermi resonance have been observed in the Raman effect. Both the methyl and methoxy torsions have been observed in the far infrared and the methyl barrier to internal rotation has been determined to be 1.15 kcal/mole (1.19 kcal/mole for the CD₃ rotor) which is lower than the 1.23 kcal/mole obtained from the microwave splitting method. It is shown from both the ${}^{13}C$ and ${}^{1}H$ NMR spectra that only one conformer exists which is contrary to what was previously reported. The vibrational spectrum of the solid is also reported and discussed.

For the complete paper, see appendix I of this report.

In addition to our studies of carbonyl containing compounds, we have been pursuing investigations on the vibrational and rotational spectra of a number of nitrogen and phosphorus containing compounds which are likely to be found in the atmospheres of the Jovian planets. In earlier progress reports we reported the vibrational spectra and structure of dimethylphosphine, ethylphosphine and isopropylphosphine. As a continuation of these studies, we have investigated the vibrational spectrum of tertiary-butylphosphine.

Vibrational spectroscopy has been shown to be a valuable tool in the determination of intramolecular potential functions governing internal rotation about single bonds.^{26,27} This method has recently been extended to the study of asymmetric tops rotating against a molecular frame, as opposed to the more familiar symmetric tops such as $-CH_3$ and $-SiH_3$.²⁸ If the molecular frame is asymmetric as well, it is possible for there to be spectroscopically distinct rotational isomers or conformers. Both ethylphosphine ($CH_3CH_2PH_2$) and isopropylphosphine $[(CH_3)_2 CHPH_2]$ are examples of this type of system since the PH₂ group constitutes an asymmetric internal rotor. The internal rotation barrier for the PH₂ top has been determined for both of these molecules. 29,30 If, however, the molecular frame is symmetric, as is the case in tertiarybutylphosphine, $[(CH_3)_3CPH_2]$, only one conformation is possible and the potential function governing internal rotation is considerably simplified, becoming threefold symmetric as in the case of methylphosphine (CH_3PH_2). Thus, the determination of this potential function for t-butylphosphine is a natural extension of the prior studies of ethyl- 29 and isopropylphosphine. 30 Also, it would be of interest to determine the barriers to internal rotation of the methyl groups in this molecule, as part of continuing studies^{31,32} of the top-top interactions in such "three-top" molecules.

Finally, the assignment of the vibrational spectrum of t-butylphosphine would finish the series of singly alkyl-substituted phosphines: methyl, ethyl

and isopropy]. Group frequencies determined for the PH₂ moiety from these studies could be expected to apply to almost any hydrocarbon derivative. The results of this study have been submitted for publication in the Journal of Molecular Structure and can be summarized by the abstract of the paper (see appendix II for the complete paper).

ABSTRACT: The infrared spectra of gaseous and solid tertiary-butylphosphine, $[(CH_3)_3CPH_2]$, have been recorded from 50 cm⁻¹ to 3500 cm⁻¹. The Raman spectra of gaseous, liquid and solid $(CH_3)_3CPH_2$ have been recorded from 10 to 3500 cm⁻¹. A vibrational assignment of the 42 normal modes has been made. A harmonic approximation of the methyl torsional barrier from observed transitions in the solid state gave a result of 4.22 kcal/mole and 3.91 kcal/mole in the gaseous state. Hot band transitions for the phosphino torsional mode have been observed. The potential function for internal rotation about the C-P bond has been calculated. The two potential constants were determined to be:

 $V_3 = 2.79 \pm 0.01$ kcal/mole and $V_6 = 0.07 \pm 0.01$ kcal/mole.

FUTURE WORK

Studies are nearly complete on $(CH_3)_2NH$ and the closely related silane, $(CH_3)_2NSiH_3$. The papers are currently being written on both of these compounds. For the dimethylamine work we have written a new computer program for the interpretation of the torsional data for the two-top case. We have also undertaken a study of the vibrational spectrum of isopropyl amine. Studies have also been initiated on the low frequency spectra of dimethylphosphine. Vibrational work is also being done on propylaldehyde. We shall also continue our current work on several compounds which have potential interest as planetary atmosphere materials.

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APPENDIX I

THE MICROWAVE, RAMAN, FAR INFRARED AND NMR SPECTRA, STRUCTURE AND DIPOLE MOMENT OF METHYLCHLOROFORMATE

Abstract

The microwave spectra of six isotopic species of methylchloroformate, $C1CO_2CH_3$, have been recorded from 18.0 to 40.0 GHz. Structural parameters have been determined and it is shown that the only stable conformer at ambient temperature is the <u>s-trans</u>. From the Stark effect the dipole moment components were determined to be $|\mu_a| = 1.7 \pm 0.2$, $|\mu_b| = 1.0 \pm 0.2$, and $|\mu_t| = 2.0 \pm 0.2D$. The Raman and far infrared spectra of the vapor are reported. Four cases of Fermi resonance have been observed in the Raman effect. Both the methyl and methoxy torsions have been observed in the far infrared and the methyl barrier to internal rotation has been determined to be 1.15 kcal/mole (1.19 kcal/mole for the CD₃ rotor) which is lower than the 1.23 kcal/mole obtained from the microwave splitting method. It is shown from both the ¹³C and ¹H NMR spectra that only one conformer exists which is contrary to what was previously reported. The vibrational spectrum of the solid is also reported and discussed.

INTRODUCTION

The structure of methylchloroformate has been the subject of considerable interest. Four dipole moment studies have been reported.¹⁻⁴ From two of these^{1,3} the results were interpreted in terms of a planar <u>s-cis</u> strucutre. The data from the other dipole moment studies^{2,4} were interpreted in terms of a planar s-trans structure (Fig. 1).

Three analyses of the vibrational spectra have been reported.⁵⁻⁷ These analyses were all in terms of a molecular model of C_s symmetry and were consistent with an <u>s-trans</u> structure. An electron diffraction study of the vapor of methylchloroformate was performed.⁸ The data yielded only an <u>s-trans</u> structure that was slightly non-planar with the methyl group approximately 20° out of the plane of the other heavy atoms.

An investigation of the infrared spectrum and of the proton NMR spectrum of methylchloroformate in solution was reported.⁹ The authors found evidence for the coexistence of both the <u>s-cis</u> and the <u>s-trans</u> conformations and indicated that the <u>s-cis</u> was more stable. From a study comparing observed and calculated ³⁵Cl nuclear quadrupole resonance frequencies the authors¹⁰ concluded that the <u>s-trans</u> was the most likely stable structure in the solid.

A microwave spectroscopic investigation¹¹ of natural abundance 35 Cl and 37 Cl isotopic species of methylchloroformate showed that the planar <u>s-trans</u> molecular strucutre was most consistent with the data. Small amounts of a second rotamer, however, could not be precluded.

Therefore, in an effort to clear up the ambiguities in the conformational preferences of methylchloroformate and to provide more complete structural data, the investigation of the microwave, Raman and far infrared spectra of the vapor were undertaken.

EXPERIMENTAL

Practical grade methylchloroformate was purchased from Aldrich Chemical Company, Incorporated and was fractionally distilled before use. The isotopic derivatives were all prepared by the reaction of the appropriate isotopic form of methanol with phosgene.¹² The CD₃OD and CH₂DOH were obtained from Merck, Sharpe and Dohme, Canada Limited. The ¹³CH₃OH was purchased from Stohler Isotope Chemicals, Inc. The methanols were all used without further purification (Stated purities: CD₃OD and CH₂DOH > 95%; ¹³CH₃OH 60-70%). The phosgene was obtained from Matheson and was used without further purification. The 1,2-dibromo-1,1,2,2-tetrafluoroethane was purchased from Columbia Organic Chemical Co. and was fractionally distilled before use.

The microwave spectra were recorded with a Hewlett-Packard 8460A MRR spectrometer in the 12.4-40.0 GH_{z} region. Samples were run both with the waveguide at room temperature and with the waveguide packed in dry ice. Sample pressures were 100 microns for survey spectra and typically 4-10 microns for measurement of the quadrupole hyperfine components. In general, the frequency accuracy should be no worse than \pm 0.05 MH_z.

The Raman spectra were recorded on a Cary Model 82 spectrophotometer with a Spectra Physics Model 171 argon ion laser. The 5145Å line was used for excitation of the samples. The gas was contained at its room temperature vapor pressure in a quartz cell and the laser beam was multipassed through the cell with the standard Cary accessory. Laser power for the gas sample was 5.5 W measured at the laser head. Laser power for the solid sample was 2.0 W measured at the laser head.

The cryostat used was a Spectrim Spectrometric Sample Conditioner controlled by a Lake Shore Electronics, Inc. Cryogenic Temperature Indicator/ Controller. The temperature of the sample was maintained at 18.7 ± 0.1° K

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and was measured with a calibrated Silicon Diode Sensor obtained from Cryogenic Technology, Inc.

The far infrared spectra of the vapor were recorded on a Digilab Model 15B Fourier Transform Interferometer with a 12.5 micron Mylar beamsplitter. The vapor was contained in a 10 cm. glass cell with polyethylene windows at its room temperature vapor pressure. Resolution was equal to 0.5 cm⁻¹.

All NMR measurements were made on a Varian XL-100-15 NMR spectrometer equipped with a 16K (1K = 1024 words) 620I computer and a 2 million word disc (VDM-36). Temperatures were measured with a thermocouple. All spectra were obtained in the pulse Fourier Transform mode.

The ¹³C spectra were obtained using broadband noise-modulated proton decoupling with the spectrometer locked to the ¹⁹F resonance of an external sample of C_6F_6 . Samples for ¹³C measurements were 50% CH_3 -O-C-C1, 50% CS_2^{-} . Proton spectra were obtained with the spectrometer locked to the ²H resonance of internal CD_2Cl_2 . The sample used for proton measurements was approximately 5 mM in CH_3 -O-C-C1 in a 50/50 mixture of CS_2 and CD_2Cl_2 . Proton T_1 measurements were obtained using the Freeman-Hill¹³ inversion recovery pulse sequence. Peak heights were employed to determine the T_1 values.

MICROWAVE RESULTS Rotational Spectra

The rotational spectra of all the isotopic species of methylchloroformate were first assigned on the basis of the known spectral pattern of a prolate asymmetric ($\kappa \approx -0.86$) rotor compared to high pressure (~ 100 Torr) survey spectra. Individual lines were then remeasured at a pressure (4-10 Torr) suitable for resolution of the hyperfine splittings that result from the nuclear quadrupole coupling. The quadrupole coupling constants were then fit to the observed splitting. The positions of all observed low pressure

transitions were then corrected for quadrupole shift and used to fit the rotational constants of a rigid rotor. Initial fits were made with a-type R branch transitions and were used to approximately locate the b-type R branches. The b-type R branches were then identified by their predicted quadrupole hyperfine splittings and were used to refine the final fit of the rotational constants. The frequencies from the low pressure measurements for all the isotrpic species of methylchloroformate studied here are listed in Table I. The rotational constants, moments of inertia, inertial defect, and nuclear quadrupole coupling constants for these isotopic species are listed in Table II.

STRUCTURE

The coordinates of the atoms for which isotopic substitutions were made have been calculated using the Kraitchman equations¹⁴ and are listed in Table III. For a non-planar asymmetric top only N-3 substitutions are needed to determine the structure. This is true because there are nine moment relations:

> $Ixx = \sum_{i} m_{i} (y_{i}^{2} + z_{i}^{2})$ (1) $Ixy = -Iyx = -\sum_{i} m_{i} x_{i} y_{i}$ (2) $\sum_{i} m_{i} x_{i} = 0$ (3)

that may be solved simultaneously. The inertial defect (Table II) for all isotopic species is indicative of an essentially planar molecule with only two hydrogens symmetrically out of the plane. For that reason the z coordinate of the atoms in the plane has been taken as zero. The nine moment relations then yield six non-trivial equations of which only five are independent. To completely solve the structure we need to calculate the six in-plane coordinates of the three unsubstituted atoms. It is, therefore, possible to calculate

the complete structure assuming only one in-plane coordinate of one of the three unsubstituted atoms.

The electron diffraction study of methylchloroformate⁸ placed the carbonyl bond length at 1.19Å. From microwave studies of some related structures $r_{c=0}$ values have been calculated for COH₂ as 1.20¹⁵, for COCl₂ as 1.17¹⁶, for HCO₂CH₃ as 1.20¹⁷, for HCO₂H as 1.20¹⁸, for NH₂COH as 1.19¹⁹, and for HFCO as 1.18²⁰. Consideration of the carbonyl bond length listed strongly suggests that $r_{c=0}$ for methylchloroformate should be between 1.18Å and 1.20Å.

In Table IV are listed the coordinates of the three unsubstituted atoms calculated from the moment relations with the x-coordinate of the carbonyl carbon chosen so that the carbonyl bond length equals 1.19\AA . The coordinates listed in Table III and Table IV reproduce the observed rotational constants in Table II to within ± 0.02%. The structural parameters calculated from these coordinates are listed in Table V. The structure of methylchloroformate is depicted in Fig. 2.

NUCLEAR QUADRUPOLE COUPLING CONSTANT

The first order chlorine nuclear quadrupole coupling constants were evaluated by a least squares fit of the observed hyperfine splittings and are listed in Table II. The value of the coupling constant along the bond axis may be calculated if the angle, θ , between the carbon-chlorine bond and the A principal axis is known using the equations:

$$\chi_{xx} \cos^2 \theta + \chi_{yy} \sin^2 \theta = \chi_{aa}$$
(4)
$$\chi_{xx} \sin^2 \theta + \chi_{yy} \cos^2 \theta = \chi_{bb}$$
(5)

where the x axis lies strong the carbon chlorine bond and y is perpendicular to it in the plane of the molecule and $\chi_{cc} = \chi_{zz}$. This transormation yields

 $\chi_{xx} = -73.7$ MHz, $\chi_{yy} = 45.1$ MHz, and $\chi_{zz} = 28.6$ for the parent molecule with $\theta = 20^{\circ}$.

These coupling constants along the bond may then be interpreted to give some information about the nature of the hybridization of the carbon-chlorine bond.²¹ In the earlier microwave study of methylchloroformate the authors¹¹ presented such a discussion and that discussion is not significantly altered by the data presented here.

BARRIER TO INTERNAL ROTATION

The barrier to internal rotation of the methyl group was computed from the observed internal rotation splittings of the a-type and b-type R branches. The same method as adopted previously²² was used to determine the barrier. The observed internal rotation splittings are listed in Table VI with the parameters calculated from the structure listed in Table III and IV. The value for V₃ calculated here is within the error limits of the previously determined V₃¹¹ value and is consistent with that found for other substituted methylformates.²³

DIPOLE MOMENT

The second order Stark effects were observed and measured for the following components: |M| = 0,3,4 of the $5_{05} + 4_{04}$ transition. Field calibration was accomplished by measuring the M=O component of OCS for the J = 2+1 transition.²⁴ The observed and calculated Stark coefficients for 35 Cl 12 Cl 16 O₂ 12 CH₃ are listed in Table VII. The microwave spectrum associated with this molecule is quite rich. The Stark components that were tracked could be followed as a function of voltage for only 2-3 MHz before being lost in other unidentified Stark component. Considerable effort was expended in an attempt to find other suitable |M| components

to track; none were found. The error limits on the dipole components are, therefore, quite high.

VIBRATIONAL RESULTS

The Raman spectra of methylchloroformate as a vapor at room temperature and as a solid at 18.7°K are shown in Fig. 3. The far infrared spectra of gaseous methylchloroformate and methylchloroformate-d₃ are presented in Fig. 4. Transitions marked with an arrow correspond to a small amount of an impurity (HCl). The vibrational data are listed in Table VIII with their assignments.

The Raman spectrum of gaseous methylchloroformate (Fig. 4A) is largely interpretable in terms of a previously reported assignment.⁷ The features at 2905 and 2845 cm⁻¹ correspond to transitions observed in the Raman spectrum of the liquid⁷ at 2897 and 2844 cm⁻¹. They are assigned as the overtones of the antisymmetric, v_4/v_{14} , and the symmetric, v_5 , methyl deformations in Fermi resonance with the symmetric CH stretch, v_2 .

The carbonyl stretch, v_2 , is assigned at 1801 cm⁻¹ because it is the stronger of the two Q-branches in the region; the other Q branch (1792 cm⁻¹) is ascribed to Fermi resonance of the combination of v_8 , the COC symmetric stretch, and v_9 , the carbon chlorine stretch, (970 + 823 = 1793) with v_2 the carbonyl stretch.

The result of another Fermi resonance interaction is apparent in the pair of Q branches at 977/962 cm⁻¹. One of the Q branches may be assigned to v_8 , the COC symmetric stretch and the other Q branch is the overtone of v_{10} , an in-plane skeletal bend (2 X 487 = 974). The last feature of the Raman spectrum of the vapor of methylchloroformate is the Q branch series starting at 487 cm⁻¹. The relative intensity of the Q branches at 487, 480, and 472 cm⁻¹ indicates they are hot bands of v_{12} , the COC in-plane bend.

The far infrared spectra of methylchloroformate-d₀ and d₃ (Fig. 4) cover the region expected for v_{17} and v_{18} , the methoxy and the methyl torsions, respectively. The methoxy torsion is assigned to the Q branches at 163.2 cm⁻¹ for the light compound and to the doublet at 148.8/148.2 cm⁻¹ for the perdeutero compound. The apparent splitting of the Q branch for the perdeutero compound remains unexplained.

The methyl torsion, v_{18} , had been reported⁷ as occurring at 62 cm⁻¹ in the liquid phase. The three-fold barrier and the structure obtained from the microwave data indicate that it should be observed near 134 cm⁻¹ in the spectra of the vapor. It is observed at 129.2 cm⁻¹ in the spectrum of the "light" vapor (Fig. 4A) and at 99.0 cm⁻¹ in the spectra of the perdeutero compound (Fig. 4B). The shift factor, $v_{\rm H}/v_{\rm D}$ = 1.305 is lower than should be expected if the torsion was not mixed with other normal modes.

The barrier to internal rotation of the methyl group, V_3 , was calculated using a computer program similar to one described by Lewis <u>et al.</u>²⁶ The barrier for the CH₃ rotor is 1.15 kcal/mole and for the CD₂ rotor it is 1.19 kcal/mole. Both barriers are lower than that calculated by the microwave splitting method.

The Raman spectrum of methylchloroformate solid at 18.7°K is quite rich (Fig. 3B). It had been suggested previously⁷ that the unit cell of methylchloroformate contained one molecule or that if it contained two molecules it should be considered centrosymmetric. Eight sharp lattice modes are apparent in the Raman spectrum below the lowest fundamental $(v_{18}, \text{ methyl torsion, 135 cm^{-1}})$. This number of lattice modes rules out the possibility of there being only one molecule per unit cell and also rules out the possibility of two molecules in a centrosymmetric unit cell. Crystal structure data remain unavailable for methylchloroformate but such

data would be of interest.

There are several apparent examples of crystal splitting evident in the spectrum, some of them quite striking. The pair of bands at 3028/3016 cm⁻¹ had also been reported in the infrared study of the solid.⁷ The authors reported an isotopic dilution study of this band pair from which they concluded the pair did not result from crystal splitting. Further examples in the Raman spectrum that apparently result from crystal splitting are v_3 , the carbonyl stretch (1774/1757/1752 cm⁻¹), v_{12} , the COC in-plane bend (283/277 cm⁻¹), v_{15} , the CH₃ rock (1152/1147 cm⁻¹) and v_{17} , the methoxy torsion (194/182 cm⁻¹). These apparent examples of crystal splitting are quite reasonable in view of the crystal structure required by the large number of lattice modes evident in the Raman spectrum.

NUCLEAR MAGNETIC RESONANCE RESULTS

The proton decoupled ¹³C NMR spectrum of $C1CO_2^{13}CH_3$ in 1,2-dibromo,tetrafluoroethane at 29°C consists of a single resonance with a half height linewidth (v_{l_2}) of 0.8 Hz. The spectrum was studied as a function of temperature to -105°C. Throughout the temperature range covered, there was no significant line-broadening observed.

The ¹H NMR spectrum of methylchloroformate in CS_2/CD_2Cl_2 (50/50) at 26°C consists of a single resonance with $v_{l_2} = 0.25$ Hz. T_1 was measured to be 15 seconds. The spectrum was again studied as a function of temperature to -100°C. At -100°C, $v_{l_2} = 0.55$ Hz and $T_1 = 2.36$ seconds. The small amount of apparent line-broadening observed at the lower temperature can be completely attributed to the change in T_1 and the fact that the magnetic field was more difficult to shim at -100°C because of the ²H lock utilized on the Varian model XL-100. For these reasons the small apparent line-broadening in the ¹H NMR spectrum at -100°C should not be ascribed to any chemical exchange processes.⁹

CONCLUSIONS

The microwave spectrum of methylchloroformate has been thoroughly reinvestigated. The data are interpreted to indicate that only one form of this molecule exists through, given the richness of the spectrum, small amounts of a second rotamer cannot positively be ruled out with these data alone. Further the microwave data are conclusively interpreted to demonstrate that the stable form of methylchloroformate is the <u>s-trans</u> form not the <u>s-cis</u> form as had been previously reported.⁹ From this microwave study it has been possible to obtain the structural parameters of this stable form along with the dipole moment components.

The vibrational data include the first direct observation of the methyl torsion of gaseous methylchloroformate and that observation is consistent with the structure and barrier calculated from the rotational data. The Raman spectrum of the solid provides more specific data on which to base generalizations as to the crystal structure of methylchloroformate than the infrared spectrum of the solid alone⁷ and again confirms the complimentary nature of the two sources of vibrational data.

The 13 C NMR temperature study is interpreted to be much more strongly suggestive of the likelihood that methylchloroformate has only one stable molecular conformation (<u>s-trans</u>) at room temperature and below. The ¹H NMR temperature study provided data that is felt to demonstrate the need for care in the performance and interpretation of such experiments.⁹

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Table I. The Microwave Spectra of Methylchloroformate

³⁵C1¹²C¹⁶O2¹²CH3

$J'_{\frac{j'_{1}}{k_{1}}} \leftarrow J_{\frac{k_{1}}{k_{1}}}$	3 Observed	F'	÷	F	Calculated	Obs-Calc	Unperturbed Level	A or E
$3_{03} + 2_{02}$	14366.90	9/2	-+-	7/2	14366.76	0.14	14366.21	A
00 02	14363.24	5/2		3/2	14363.12	0.12		A
$3_{13} + 2_{12}$	13642.09	9/2	~-	7/2	13641.94	0.15	13640.52	A
	13638.40	7/2	~-	5/2	13638.22	0.18		A
³ ₁₂ ← 2 ₁₁	15303.35	9/2	~-	7/2	15303.29	0.06	15301.74	A
	15299.62	7/2	÷	5/2	15299.57	0.05		A
4 ₀₄ ← 3 ₀₃	19014.28	11/2+9	/2 ←	9/2+7/2	19014.14	0.14	19013.84	A
04 05	19012.57	7/2+5	/2 ←	5/2+3/2	19012.39	0.18		A
⁴ 14 ^{← 3} 13	18154.10	11/2		9/2	18153.98	0.12	18153.33	А
14 10	18152.70	9/2	÷	7/2	18152.51	0.19		А
	18152.15	7/2		5/2	18151.77	0.38		А
	18153.40	5/2	÷	3/2	18153.24	0.16		Å
4 ₁₃ ← 3 ₁₂	20364.25	11/2	÷	9/2	20363.94	0.31	20363.35	A
10 ,	20362.57	9/2	÷	7/2	20362.48	0.09		А
	20362.11	7/2	÷	5/2	20361.65	0.46		A
4 ₂₂	19605.73	11/2	~-	9/2	19605.76	-0.03	19103.48	А
		9/2	÷	7/2	19599.69			А
	19601.73	7/2	4	5/2	19601.80	-0.07		А
	19607.78	5/2	÷	3/2	19607.87	-0.09		A
	19596.54	11/2	- 4 -	9/2			19594.42	E
	19590.72	9/2	~-	7/2			10001110	E
	19592.80	7/2	÷	5/2				Ε
-		5/2	+	3/2				Е

	Tab	le	Ι	Continued
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	с ¹⁶ 02 ¹² СН							Unperturbed	
<u>k'jk+j</u>	^{← J} , <u>k_ןk+</u> ן	Observed	F'	 -	F	Calculated	Obs-Calc	Level	Aorl
⁵ 05 ≁	4 ₀₄	23553.51	13/2	4-	11/2	23553.40	0.11	23553.25	А
		23552.58	9/2	*	7/2	23552.48	0.10		А
⁵ 15 [←]	4 ₁₄	22640.36	13/2	*	11/2	22640.40	-0.04	22640.02	A
• -		22639.83	11/2	~ -	9/2	22639.68	0.15		А
		22639.41	9/2	4	7/2	22639.03	0.38		A
⁵ 14 ←	4 ₁₃	25387.83	13/2	÷	11/2	25388.02	-0.19	25387.24	A
•••		25387.02	11/2	÷	9/2	25387.32	-0.30		А
5 ₂₄ ←	⁴ 23	24082.85	13/2	÷	11/2	24082.75	0.10	24081.70	A
	2.	24080.12	11/2	÷	9/2	24079.78	0.34		А
		24084.85	13/2	~	11/2			24083.74	E
		24082.19	11/2	÷	9/2				Ε
5 ₂₃ ←	⁴ 22	24680.69	13/2	÷	11/2	24680.41	0.28	24679.45	A
2.5		24677.75	11/2	. (-	9/2	24677.32	0.43		А
		24678.69	13/2	*	11/2			24677.54	Ε
		24675.94	11/2		9/2				Ε
5 4	5	27981.37	15/2	4	13/2	27981.36	0.01	27981.18	A
⁶ 06 ←	⁹ 05	27981.37	-			27980.78	-0.01	27501.10	A
	-	07000 04	7.5.10		10/0	87000 00	0.01	07007 00	a
⁶ 16 ⁺	⁵ 15	27098.04				27098.08	-0.01	27097.80	A
		27097.79				27097.68	0.11		A
		27097.33	11/2	+	9/2	27097.18	0.15		A
6 ₁₅ ←	5 ₁₄	30361.85	15/2	÷	13/2	30361.83	0.02	30361.54	А
-		30361.49	13/2	÷	11/2	30361.46	0.03		А
		30361.08	11/2		9/2	30360.92	0.16		А

			•					
35 _{C1} 12 _C 16 ₀₂ 12 _{CH}								
$\frac{J'_{k_1k_1} \leftarrow J_{k_1k_1}}{k_1k_1}$	-	F۱		F	Calculated	Obs-Calc	Unperturbed Level	A or E
⁶ 25 ^{← 5} 24	28841.62	15/2	+	13/2	28841.50	0.12	28840.82	A
	28839.95	13/2	~-	11/2	28839.82	0.13		А
	28842.29	15/2	(13/2			28841.49	Е
	28840.62	13/2	÷	11/2				E
6 ₂₄	29840.51	15/2	+	13/2	29840.42	0.09	29839.65	A
	29838.72	13/2	÷	11/2	29838.09	0.03		А
	29839.98	15/2	≁ -	13/2			29839.14	E
	29838.23	13/2	*	11/2				E
7 ← 6	32313.35	17/2	4	15/2	32313.31	0.04	32313.17	А
7 ₀₇	32312.99	13/2		11/2	32312.92	0.07	52515.17	A
	32312.66		÷	9/2	32312.81	-0.15		A
7 ₁₇ ← 6 ₁₆	31526.02	17/2	~-	15/2	31526.06	-0.04	31525.78	A
	31525.75	15/2	+	13/2	31525.82	-0.07		А
	31525.48	13/2	÷	11/2	31525.43	0.05		А
7 ₁₆	35269.20	17/2	*	15/2	35269.35	-0.15		А
	35269.01	15/2	~-	13/2	35269.15	-0.14	35269.03	А
	35268.82	13/2	-	11/2	35268.93	-0.11		А
⁷ 26 ^{← 6} 25	33529.92	17/2	4	15/2	33569.88	0.04	33569.38	A
	33568.91	15/2		13/2	33568.85	0.06		A
7 _{.25} ← 6 ₂₄	35064.18	·		15/2	35064.29	-0.11	35063.55	А
	35062.98	15/2	- - -	13/2	35063.15	-0.17		A
7 ₃₄ + 6 ₃₃	34143.18			15/2	34143.41	-0.23	34142.13	A
	34140.89	-		-	34140.96	-0.07		A
	34143.55	(1/2	÷	9/2	34143.73	-0.18		А

Table I Continued

Table I Continued

³⁵C1¹²C¹⁶O2¹²CH3

	-						Unperturbed	
$\frac{J'_{k'_1k_{+1}} + J_{k_{-1}k_{+1}}}{k_{-1}k_{+1}}$	Observed	F'	+	F	Calculated	Obs-Calc	Level	A or E
⁸ 08 ⁺ ⁷ 07	36578.20	19/2		17/2	36578.36	-0.16	36578.11	A
	36577.99	15/2	÷	13/2	56578.07	-0.08		А
	36577.79	13/2	*	11/2	36577.99	-0.20		A
8 ₁₈ ← 7 ₁₇	35924.89	19/2	÷	17/2	35925.15	-0.26	35924.70	A
	35924.75	17/2	 -	15/2	35924.96	-0.21		А
⁸ 17 ÷ 7 ₁₆	40092.32	19/2	~	17/2	40092.53	-0.21	40092.09	А
	40092.07	17/2	4	15/2	40092.44	-0.37		А
	40091.87	15/2	- (13/2	40092.10	-0.23		А
⁸ 27 ← ⁷ 26	38263.35	19/2	÷	17/2	38263.48	-0.13	38263.00	А
2. 2.	3826 2.7 7	17/2	~ -	15/2	38262.81	-0.04		A
6 4 5	31148.83	15/2	~ -	13/2	31148.73	0.10	31149.07	А
⁶ 16 ^{÷ 5} 05	31150.00			11/2	31149.86	0.14		A
	31147.90	9/2		7/2	31147.80	0.10		A
	31146.81	•		13/2			31147.00	E
	31147.90			11/2		,		Ε
	31145.80	9/2		7/2				E
7.6	24602 44	17/2	,	15/2	34693.41	0.03	34693.55	A
⁷ 17 ^{← 6} 06	34693.44 34694.18	-		13/2	34694.19	-0.01	04050.00	A
	34694.18 34692.74	15/2		9/2	34692.78	-0.01		A
	34692.74 34691.56			9/2 15/2	J+V37/O	-0.04	34691.72	E
	34691.36			13/2			0-001.72	E
	34692.30			9/2				E

³⁵ c1 ¹² c ¹⁶ 02 ¹² c	CH ₃						llopentushed	
$J_{k_1k_1} \leftarrow J_{k_1k_1}$	Observed	F'	4 -	F	Calculated	Obs-Calc	Unperturbed Level	A or E
⁸ 18 + ⁷ 07	38305.20	19/2	*	17/2	38305.24	-0.04	38305.27	A
10 0,	38305.68	17/2		15/2	38305.77	-0.09		А
	38304.74	13/2		11/2	38304.79	-0.05		А
	38303.58	19/2	÷	17/2			38303.61	E
	38304.01	17/2		15/2				Έ
	38303.06	13/2	÷	11/2				E
³⁷ _{C1} ¹² C ¹⁶ 02 ¹² C	CH ₃							
3 ₀₃ ← 2 ₀₂	14034.80	9/2+7	/2+	- 7/2+5/2	14034.77	0.03	14034.24	A
03 02	14031.90	5/2+3	8/2+	- 3/2+1/2	14031.87	0.03		А
6 ₀₆ ← 5 ₀₅	27377.36	13/2	÷	11/2	27377.32	0.04	27377.03	A
	27377.07	15/2	÷	13/2	27377.18	-0.09		A
⁶ 16 ^{+ 5} 15	26502.95	15/2	~	13/2	26502.90	0.05	26502.68	А
10 13	26502.64	13/2	÷	11/2	26502.60	0.04		A
⁶ 15 ^{← 5} 14	29630.17	15/2+13	3/2⊀	-13/2+11/	2 29630.10	0.07	29630.03	A
⁶ 24 ^{+ 5} 23	29085.75	15/2	- 4-	13/2	29085.63	0.12	29085.07	A
	29084.33	13/2		11/2	29084.19	0.14		А
	29085.10	15/2		13/2			29084.50	Е
	29083.84	13/2	÷	11/2				E
7 ₀₇ ← ⁶ 06	31629.18	15/2	÷	13/2	31629.10	0.08	31628.94	A
′0/ ` °06	31629.00	-			31628.97	0.03		A.
7 ₁₇ ← ⁶ 16	30838.25	17/2+1	5/24	-15/2+13/	2 30838.27	-0.02	30838.15	A

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³⁷ с1 ¹² с ¹⁶ 02 ¹² сн	3							
$J_{k'_{1}k'_{1}} \stackrel{\leftarrow}{=} J_{k_{1}k'_{1}}$	Observed	F'	+	F (Calculated	Obs-Calc	Unperturbed Level	A or E
7 ₁₆ ← 6 ₁₅	34430.71	17/2+15	i/2+	15/2+13/2	34430.70	0.01	34430.59	A
7 ₂₅ ← 6 ₂₄	34170.53	17/2	~ -	15/2	34170.54	-0.01	34170.04	A
	34169.60	19/2	4-	17/2	34169.62	-0.02		A
8 ₀₈ ← 7 ₀₇	35813.41	15/2	÷	13/2	35813.40	0.01	35813.22	A
	35813.24	17/2	÷	15/2	35813.28	-0.04		А
⁸ 18 + 7 ₁₇	35146.57	19/2	*	17/2	35146.64	-0.07	35146.44	А
	35146.48	17/2	~-	15/2	35146.54	-0.06		А
⁸ 17 ^{← 7} 16	39155.34	19/2+17	/2+	-17/2+15/2	39155.41	-0.07	39155.24	A
⁸ 27 ← ⁷ 26	37382.34	19/2	 -	17/2	37382.34	0.00	37382.06	А
	37381.88	17/2	-t	15/2	37381.83	0.05		А
⁸ 26 ^{~~7} 25	39287.67	19/2	-1-	17/2	39287.78	-0.11	39287.31	А
	39287.10	17/2	~ -	15/2	39287.16	-0.06		А
³⁵ c1 ¹² c ¹⁶ 02 ¹² cc) ₃							
⁶ 06 ^{← 5} 05	25500.25	15/2		13/2	25500.18	0.07	25500.09	A
	25499.72	11/2	÷	9/2	25499.63	0.09		А
6 ₁₅ ^{← 5} 14	27398.40	15/2	~-	13/2	27398.34	0.06	27398.08	A
13 11	27397.99				27397.90	0.09		A
	27397.59	11/2	÷	9/2	27397.46	0.13		A
6 ₂₅ ← 5 ₂₄	26128.59	15/2	÷	13/2	26128.51	0.09	26127.84	A
	26127.04	13/2+1	1/2-	⊷11/2+9/2	26126.91	0.11		A

Table I Continued

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³⁵_{C1}¹²C¹⁶O₂¹²CD₃

$J_{k_{1}k_{1}}^{i} \leftarrow J_{k_{1}k_{1}}^{i}$	3 Observed	۴	÷	F (Calculated	Obs-Calc	Unperturbed Level	A or E
$6_{24} \leftarrow 5_{23}$	26855,95	15/2	+	13/2	26855.88	0.07	26855.12	A
24 23	26854.23	13/2	÷	11/2	26854.18	0.05		A
7 ₀₇ ← 6 ₀₆	29494.20	17/2	« -	15/2	29494.17	0.03	29494.02	A
0, 00	29493.77	15/2+13	8/2≁	-13/2+11/2	29493.63	0.14		А
7 ₁₇ ← 6 ₁₅	31858.88	17/2	÷	15/2	31858.92	-0.04	31858.67	А
17 10	31858.70	15/2	~ ~-	13/2	31858.72	-0.02		А
	31858.53	11/2	~	9/2	31858.52	0.01		А
·	31858.32	13/2	~ -	11/2	31858.32	0.00		А
7 ₂₆	30426.94	17/2	4	15/2	30427.02	-0.08	30426.42	A
20 23	30425.91	15/2+13	3/2-	413/2+11/2	30425.98	-0.07		A
7 ₂₅ ← 6 ₂₄	31530.17	17/2		15/2	31530.23	-0.06	31529.60	A
20 24	31529.03	15/2+13	3/2-	+13/2+11/2	31529.08	-0.05		А
7 ₃₄ ← 6 ₃₃	30829.38	17/2	~ -	15/2	30829.49	-0.11	30828.38	A
54 55	30827.08	15/2	+	13/2	30827.18	-0.10		А
6	28881.94	13/2	÷	11/2	28881.95	-0.01	28881.03	A
⁶ 16 ^{≁ 5} 05	28880.78			13/2	28880.79	-0.01		А
³⁵ C1 ¹² C ¹⁶ O2 ¹³ CF	1 ₃							
$4_{04} + 3_{03}$	18566.43	11/2+9	/2 -	+9/2+7/2	18566.42	0.01	18565.98	А
	18564.72	7/2+5	/2 -	←5/2+3/2	18564,74	-0.02		A
⁵ 05 ^{← 4} 04	23011.96	i 13/2+1	1/2	←]]/2+9/2	23012.01	-0.05	23011.65	A
ບວ U4					23011.01	0.02		А

Table I Continued

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$\frac{K_{1}K_{1}}{6}_{25} \leftarrow \frac{5}{24}$ 28	bserved	F'	-					
⁶ 25 ^{← 5} 24 ²⁸	81/10 Q5		*	F	Calculated	Obs-Calc	Unperturbed Level	A or E
	0140.30	15/2+9/	2 +13	/2+7/2	28140.90	0.05	28140.16	А
20	8139.31	13/2+11	/2-11	/2+9/2	28139.30	0.01		А
28	8141.67	15/2+9/	2 ~13	3/2+7/2			28140.88	Е
28	8140.02	13/2+11	/2-4]1	/2+9/2				Е
₆₂₄ ← ⁵ 23 25	9054.61	15/2+9/	'2 + 13	3/2+7/2	29054.53	0.08	29053.74	А
	9052.79	13/2+11	/2+11	/2+9/2	29052.78	0.01		А
2'	9053.99	15/2+9/	2 ← 13	3/2+7/2			29053.16	E
2'	9052.26	13/2+13	/2 + 11	/2+9/2				Ε
$7_{26} + 6_{25} = 3$	2759.63	17/2+1	1/2+1	5/2+9/2	32759.64	-0.01	32759.19	А
	82758.69	15/2+13	3/2+13	3/2+11/2	2 32758.66	0.03		А
7 ₂₅ ← 6 ₂₄ 3	34133.36	17/2+1	1/2+1	5/2+9/2	34133.46	-0.10	34132.88	А
	34132.30	15/2+1	3/2+1∶	3/2+11/3	34132.33	-0.03		A
								_
-18 07	37646.46				37646.63	-0.17	37646.14	A
3	37646.12	19/2+1	5/2+1	7/2+13/2	2 37646.10	0.02		A
3	37645.69	13/2	+	11/2	37645.55	0.14		A
35 _{C1} 12 _C 16 ₀₂ 12 _{CH2}	D-00P							
3 ÷ 2	13923.95	9/2+7	/2+7	/2+5/2	13923.88	0.07	13923.22	Α
-03 02	13920.26			/2+1/2	13920.24	0.02		А
$3_{12} \leftarrow 2_{11}$	14783.69	9/2+3	¦/2+7	/2+1/2	14783.50	0.19	14782.06	А
· [Z]]	14780.00			5/2+3/2	14799.86	0.14		А

Table 1 Continued

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³⁵ с1 ¹² с ¹⁶ 02 ¹² сн	2 ^{D-00P}							
]' k <u>'</u> 1k'+1 ← J k_1k+1	Observed	F۱		F	Calculated	Obs-Calc	Unperturbed Level	Aori
4 ₀₄	13923.95	9/2+7/2	≁7	/2+5/2	13923.88	0.07	13923.22	А
04 00	18440.64	7/2+5/2	∻-5	/2+3/2	18440.64	0.00		A
4 ₁₄ + ³ 13	17632.27	11/2	-	9/2	17632.31	-0.04	17631.45	A
,	17631.54	5/2	+	3/2	17631.55	-0.01		А
	17630.83	9/2	~-	7/2	17630.86	-0.03		А
	17630.19	7/2	÷	5/2	17630.10	0.09		А
⁴ 13 ^{+ 3} 12	19676.46	11/2	÷	9/2	19676.52	-0.06	19675.65	A
	19675.62	5/2	÷	3/2	19675.74	-0.12		A
	19675.13	9/2	÷	7/2	19675.08	0.05		А
	19674.41	7/2	~	5/2	19674.30	0.11		A
⁵ 05 ^{← 4} 04	22865.86	13/2+11/	2 <i>+</i> 1	1/2+9/2	22865.86	0.00	22865.56	A
05 04	22864.88	9/2+11/2		7/2+5/2	22864.84	0.04		А
5 ₁₄ ← 4 ₁₃	24538.20	13/2	÷	11/2	24538.28	-0.08	24537.72	А
,	24537.62	11/2+7/2	. +	9/2+5/2	24537.60	0.02		А
	24536.92	9/2	~	7/2	24536.91	0.01		А
⁵ 24 ^{← 4} 23	23325.99	13/2+7/2	T	11/2+5/2	23326.04	-0.05	23324.54	А
2	23323.10	11/2+9/2	<u>}</u> +	9/2+7/2	23323.12	-0.02		А
0 7	27/ 71 70	17/9		15/2	37471.73	0.05	37471.34	A
⁸ 18 ^{← 7} 07	37471.78				37471.73	0.03	J/7/1.JT	A
	37471.29	-		17/2				A
	37470.81	13/2		11/2	37470.85	-0.04		ч

Table I Continued

³⁵c1¹²c¹⁶02¹²CH2D-IP

^j k'_k, ^{+ ,j} k_k	Observed	F' ← F	Calculated	Obs-Calcd	Unperturbed Level	A or E
³ ₁₂ ← ² ₁₁	14686.41	9/2 ← 7/2	14686.29	0.12	14684.90	A
14 11	14682.92	7/2+5/2+5/2+3/2	14682.61	0.31		А
⁶ 15 ^{+ 5} 14	29144.92	15/2 ← 13/2	29144.83	0.09	29144.66	A
15 14	29144.56	1 [°] 3/2 ← 11/2	29144.44	0.12		А
	29144.23	11/2 + 9/2	29143.93	0.30		А
⁶ 24 ^{+ 5} 23	28628.59	5/2+9/2+3/2+7/2	28628.54	0.05	28627.74	А
24 23	28626.87	13/2+11/2+11/2+9/2	28626.81	0.06		A
⁶ 25 ^{≁ 5} 24	27693.97	15/2+9/2 ←1 3/2+7/2	27694.00	-0.03	27693.20	А
20 24	27692.38	13/2+11/2+11/2+9/2	27692.34	0.04		А
7 ₂₅ + 6 ₂₄	33637.46	17/2+11/2+15/2+9/3	33637.45	0.01	33636.96	A
20 24	33636.40	15/2+13/2+13/2+11/3	33636.37	0.03		А
7 ₂₆	32236.10	17/2+11/2+15/2+9/2	32236.12	-0.02	32235.60	А
20 25		15/2+13/2+13/2+11/2	32235.11	0.06		А
8 ₂₆ ← 7 ₂₅	38675.55	19/2+13/2≁17/2+11/2	38675.60	-0.05	38675.24	А
20 20	38674.86	17/2+15/2+(15/2+13/	2 38674.92	-0.06		А
⁸ 27 ^{← 7} 26	36745.89	19/2+13/2+13/2+11/2	36745.98	-0.09	36745.57	А
27 20	36745.19	17/2+15/2+15/2+13/3	36745.30	-0.11		А
$7_{17} \div 6_{06}$	33426.72	15/2 + 13/2	33426.76	-0.07	33426.23	A
17 00	33426.19	17/2 + 15/2	33426.11	0.08		А
8 ₁₈ + 7 ₀₇	36889.70	17/2 + 15/2	36889.72	-0.02	36889.31	А
18 U7	36889.25	19/2+15/2+17/2+13/2	36889.29	-0.04		А
	36888.81	13/2 + 11/2	36888.86	-0.05		А

	³⁵ C1 ¹² C ¹⁶ 02 ¹² CH3	³⁷ C1 ¹² C ¹⁶ 02 ¹² CH ₃	³⁵ _{C1} ¹² C ¹⁶ O2 ¹² CD3	³⁵ c1 ¹² c ¹⁶ 02 ¹³ CH ₃	³⁵ C1 ¹² C ¹⁶ O2 ¹² CH2D-00P	³⁵ C1 ¹² C ¹⁶ 02 ¹² CH2 ^{D-IP}
д	9765.82 ± 0.20	9738.39 ± 0.29	9172.25 ± 0.18	9738.40 ± 0.30	9619.16 ± 0.15	9465.36 ± 0.30
3	2692.17 ± 0.01	2623.00 ± 0.01	2413.68 ± 0.01	2620.09 ± 0.01	2594.66 ± 0.01	2582.49 ± 0.01
C	2138.03 ± 0.01	2092.92 ± 0.01	1957.65 ± 0.01	2091.18 ± 0.01	2082.27 ± 0.01	2054.86 ± 0.01
`а	51.7498 ± 0.0011	51.8955 ± 0.0016	55.0987 ± 0.0011	51.8955 ± 0.0016	52.5388 ± 0.0009	53.3925 ± 0.001c
I _b	187.722 ± 0.001	192.671 ± 0.001	209.381 ± 0.001	192.886 ± 0.001	194.777 ± 0.001	195.694 ± 0.001
I c	236.376 ± 0.001	241.471 ± 0.001	258.156 ± 0.001	241.672 ± 0.001	242.706 ± 0.001	245.943 ± 0.001
7	3.096	3.097	6.324	3.110	4.610	
(aa	-59.8 ± 0.5	046.4 ± 0.5	-56.7 ± 0.5	-57.2 ± 0.5	-58.6 ± 0.5	-56.2 ± 0.5
aa ⁽ bb	31.2 ± 1.0	25.4 ± 1.0	29.8 ± 1.0	30.9 ± 1.0	29.7 ± 1.0	29.6 ± 1.0
cc	28.6 ± 1.0	21.0 ± 1.0	26.9 ± 1.0	26.3 ± 1.0	28.9 ± 1.0	26.6 ± 1.0

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TABLE II. Rotational constants, moments of inertia and nuclear quadrupole coupling constants for metnylchloroformate

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TABLE III

	Coord	inates of the atoms	for which isotopic sub	stitutions were made"
	Chlorine	Carbon (methyl)	Hydrogen (in plane)	Hydrogen (out of plane)
Х	-1.590	2.278	2.808	2.510
Y	-0.278	-0.381	-1.312	0.182
Z	0.014 ^b	0,084 ^b	0.161 ^b	±0.887

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 $^{\rm a} Error$ limits on the coordinates are never worse than $\pm 0.005.$

^bThe Z-coordinate of these atoms was taken as 0.0 for subsequent calculations.

TABLE IV

Coordinate: of the unsubstituted atoms calculated from the moment relations

	Carbon (carbonyl)	Oxygen (carbonyl)	Oxygen (ether)
Х	0.034 ^a	0.353	0.895
Y	0.312	1,458	-0.740
Z	0.0 ^b	0.0 ^b	0.0 ^b

^aThis value was chosen so that the carbonyl bond length would equal 1.190 Å. ^bThese values were assumed based on a consideration of the inertial defect.

TABLE V

Structural parameters for methylchloroformate^a

	Bond Lengths (Å)	Bond Angles (°)	
C1-C	1.73	C1-C=0 126	
C=0	1.19	0=C-0 125	
(0=)C-0	1.36	C-O-C 115	
0-C(H ₃)	1.43	0-C-H(IP) 105	
C-H(IP)	1.07	0-C-H(00P) 110	
C-H(OOP)	1.08	H-C-H(OOP-OOP) 111	
		H-C-H(IP-00P) 110	

^aCalculated from the coordinates in Table III and IV. The errors associated with these parameters because of the assumptions involved in their calculation are probably no worse than ± 0.02 Å for the bond lengths and $\pm 2^{\circ}$ for the bond angles.

TABLE VI	
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Internal rotation splitting (MHz) and parameters of methylchloroformate

	³⁵ с1	¹² c ¹⁶ 02 ¹² CH3	37 _{Cl} 12	² с ¹⁶ 02 ¹² СН ₃	³⁵ cī ¹² c	¹⁶ 02 ¹³ CH3	
Transition	^v e ^{-v} A	V ₃ (cal/mol)	ve-va	V ₃ (cal/mol)	`Έ ΥΑ	V ₃ (cal/mol)	
4 ₂₂ ← 3 ₂₁	-9.06	1190					
⁵ 24 + ⁴ 23	2.04	1230					
⁵ 23 ^{← 4} 22	-1.91	1240					
⁶ 25 ÷ ⁵ 24	0.67	1230			0.72	1230	
6 ₂₄ ← ⁵ 23	-0.51	1270	-0.57	1270	-0.58	1270	
⁶ 16 ^{← 5} 05	-2.09	1220					
7 ₁₇ ← ⁶ 06	-1.83	1220					
8 ₁₈ + 7 ₀₇	-1.66	1220					
I		2.954 amu • Å ²	2.	954 amu • Ų	2.955	amu · Å2	
Ι _α F		171.1 GH ₃	17	'1.1 GH ₃	171.0	GH3	
α		0.05598	0.	05945	0.05926		
β		0.00432	0.	00439	0.00456		
γ		0.0	С.	0	0.0		
٧ ₃		1230 ± 30 cal/mol					

	Т	AB	LE	۷	I	Ī
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Stark coefficients [(MHz cm^2)/V²] and dipole moments of methylchloroformate^a

Transition	[M]	$\Delta v/E^2$ ((x10 ⁵)	
	• ••••••••••••••••••••••••••••••••••••	Obsd.	Calcd.	
⁵ 05 ^{←4} 04	0	0.09	0.07	
05 04	3	0.28	0.41	
	4	0.59	0.67	
$ \mu_{a} = 1.7 \pm 0.2$ $ \mu_{b} = 1.0 \pm 0.2$ $ \mu_{c} = 0.0^{a}$ $ \mu_{t} = 2.0 \pm 0.2$				

a.) By symmetry

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	Vapor				<u>-</u>	Sol	lid			Assignment		
	Infrared	à	Raman		Infrared ^{a,b} Raman				Raman ^C			
, 			·			A'	<u>A"</u>		-		_	
Cm	-1 t	/ре	∆cm ^{−1}	Rel Int	cm	Rel Int		Rel Int	∆cm ⁻¹	Rel Int		
					3976	vvw ^d	3976	W			Α" ν ₈ + ν ₁₃ (3970)	
					3904	VW					Α' ν ₂ + ν ₈ (3914)	
3571					3509	W					A' 2v ₃ (3500)	
3569	י א	∖B			3491	m						
,	,								3251	VVW	Α" ν _] + ν _{]7} (3253)	
									3232	VVW	A'/A" v ₃ + v ₄ /v ₁₄ (3232)	
									3207	VVW	$A'/A'' v_3 + v_4/v_{14}$ (3207)	
•									3072	vvw	3 4, 14 (0-0)	
3057)		3047	w,p	3048	W	3048	vvw	3055	m	Α' ν _ן CH ₃ asymmetric stretch	
3033			∿3025	w,dp	3025	w	3025	S	3028	m		
3016	<i>۲</i>	IC	00020	տ,սի	3016		3016		3018		Α" ν ₁₃ CH ₃ asymmetric stretch	
					3010	S	3010	S	3010	W		
3005)				2978	sh						
2979												
2973	> P	k	2962	vs,p	2961	S	2961	w	2963	vvs	A' ν ₂ CH ₃ symmetric stretch	
2967			2934	p							2 3 5	
2 920				L.	2919	W			2914			
			2905	р	2902	W			to	vw.h	Α' 2(ν ₄ /ν ₁₄) F.R. ŵ ν ₂	
				F		.,	2881	W	2880			
					2879	W	2001					
2863					2869							
2855	م <		2845	n	2846	W	2846	L.f	2845			
2849	ſ	L	2043	р	2040	ms	2040	W	2040	m	A' 2 v_5 F.R. $\bar{w} v_2$	

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Table VIII. The Vibrational Spectra and Assignment of Methylchloroformate

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	Vapor					Solid				Assignment		
	Infrared ^a	Rama	ın		I A'	nfrared ^{a,b} A"		Raman	2			
cm ⁻¹	type	∆cm ⁻¹	Rel Int.	cm ⁻¹	Rel	Int cm ⁻¹	Re1	Int ∆cm ⁻¹	Rel Int			
· · · · · · · · · · · · · · · · · · ·								2640	VVW			
2610	?			2601	W			∿2603	VVW	Α' ν ₃ + ν ₉ (2599)		
				2582	W			∿2592	VVW			
				2570	W	2570	W	2577	VW	Α' 2ν ₆ (2578)		
				2418	W							
2402				2406	W					A' 2v ₇ (2408)		
				2371	٧W					A'		
				2316	m	2316	m			Α' 2ν ₁₅ (2304)		
								2297	VW	A' 2015 (2296)		
								2273	VW	$A'/A" v_4/v_{14} + v_9$ (2271)		
								2178	W	A' v ₃ + v ₁₁ (2183)		
				2142	W							
								2120	VVW	Α' ν ₇ + ν ₈ (2116)		
				2095	W			~2100	VVVW	A" v ₈ + v ₁₅ (2099)		
				2089	W			2079	VVW			
2027				2025	vw					A' א _ד י א _ק (2025)		
2022												
1973	?			1960	٧W					A" v ₃ + v ₁₇ (1956)		
1924				1905	W					· · ·		
				1890	W					A' 2v ₈ (1894)		
				1882	ms			∿1880	vvw	~		

Table VIII. Con't.

				Ta	ble VII	I. Con't.					
·		Vapor			S	olid			Assignment		
	Infrared ^a	Rama	n		In	frared ^{a,b}		Raman ^C			
cm ⁻¹	type	∆cm ⁻¹	Rel Int	cm ⁻¹	A' Rel I	A" nt_cm ⁻¹	Re1	Int ∆cm ⁻¹	Rel In	t_	
1806		1801	p	1789)				1774	w)		
1794	В	1792	p	to >	S			1757	mw >	ν ₃ C=O stretch	
1788			·	1755				ì752	m (5	
,				1725	m				j		
				1716	s			1717	VW	Α' ν ₁₂ + ν ₅ (1717)	
				1646	mw			1652	VW	A' 20g (1642)	
				1622	mw			∿1625	VW	A' v ₁₄ + v ₁₇ (1622)	
1464		1466	p								
1459	>?							1458	m		
1448		1459	dp	1448.6	S	1449.2	`S	1450	mw {	A'/A" م ₄ /م ₁₄ CH ₃ asymmetric deformation	
1441 (1433 (?										
,				1431	S	1431	sh	1432	mw	A' v ₅ CH ₃ symmetric deforma- tion	
1396								~1405	VVW		
1382	?	1389	р	1387	s	1387	٧W			A' 2v ₁₆ (1378)	
			·	1326	mw					A' v ₁₅ + v ₁₇ (1322)	
1313											
1308	> AB	-		1306	S	1306	W			$A' v_9 + v_{10}$ (1302)	
1300				1297	S					A' v ₉ + v ₁₀ (1296)	

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	Vapor				Solid					Assignment		
	Infrared ^a	Rama	'n		Infr	ared ^{a,h}		Raman ^C				
					<u>A'</u>	<u>A"</u>						
1	type	∆cm ⁻¹	Rel Int	1	Rel Int	cm ⁻¹	Rel In	t ∆cm ⁻¹	Rel Int			
1233				1250)		1250	W					
1210	?			to >	S	1226	VW			-		
1202	·	1157	р	1210		1211	٧w	1204	тw	A' ν ₆ COC asymmetric stretch		
1196				-						5		
				1186	S	1180	W			_		
1162	AB			1174	S			1169	mw	A' ν ₇ CH ₃ r.		
1154		1124	dp	1152	sh	1152	ms	1152	W	$A'' v_{15} CH_3 r.$		
				1142	m	1142	W	1148	W	A" $v_{9} + v_{17}$ (1141)		
								1144	sh			
1060								1087	VVVW			
								1056	VVVW			
1036								1046	VVVW			
1007										-		
								988	W	^{A'} v16 ⁺ v17 ⁽⁹⁸⁵⁾		
				961	W			972	VW	A' 2010 (972)		
				958	W			960	vw	A' 2 ₁₀ (958)		
971		977	р					951	sh	Α" ν ₉ + ν ₁₈ (956)		
966 >	?	962	р	947	S	947	W	947	m	A' v8 COC symmetric stretch		
957								932	VVW	Ç		
,								924	VVW			
				920	W			918	VVW			
892				889	mw			∿893	WVV	A' v ₁₀ + v ₁₁ (895)		

Table VIII. Con't.

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		Solid					Assignment				
	Infrared ^a	Rama	n		Infra	red ^{a,b}		Raman ^C			
					Α'	Α"					
cm ⁻¹	type	∆cm ⁻¹	Rel Int		Rel Int		Rel I	nt_∆cm ⁻¹	Rel Int		
882				881	W						
830]											
824 >	А	823	þ	825	s	825	W				
816				821	S	821	W	821	ms	A' ν _g C-Cl stretch	
2				817	ms					-	
				758	W			766	VW		
706											
694 >	С			689	W	689	S	702	VVVW	Α" ν ₁₆ C-Cl bend	
677				672	W	672	m				
668											
542											
527								518	VVVW		
494]											
488 >	A	487	p	481	S	481	٧W	486	vs	A' v ₁₀ OCO bend	
477		480	р	477	m	477	vw	479	ms∫	10	
		472	р								
413]											
405 >	AB	406	р	407	S	407	W	409	ms	A' v _{ll} C-Cl bend	
397											
-								392	VVW	A' 2v ₁₇ (388)	
354								367	VVW	A' 2017 (364)	

Table VIII. Con't.

	Vapor				Solid				Assignment	
I	nfrared ^a	Rama	In	A۱	Inf	rared ^{a,b} A"		Raman ^C		
cm ⁻¹	type	∆cm ^{-]}	Rel Int	cm ⁻¹	Rel Int	1	Rel Int	∆cm ⁻¹	Rel Int	-
325								341	vvvw	
314	?									
304						·		283	w l	
279 268	?	274	р	270	S			277	w	A' v_{12} COC bend
		Methylch	loroformate-d	1,						
			ared Vapor	3				202	٧₩	
163.2	S	148.8	S					194	w l	
		148.2	s					182	w	Α" ν ₁₇ Methoxy torsion
129.2	W	99.0	W					135	VVW	A" v ₁₈ Methyl torsion
								122	ms	Lattice mode
								100	sh	Lattice mode
								94	s	Lattice mode
								81	W	Lattice mode
								67	mw	Lattice mode
								60	W	Lattice mode
								52	mw	Lattice mode
a.) Taken from Ref. 2° b.) Recorded at ∿77°K c.) Recorded at 187°K d.) vw = very weak, w = weak, m = medium, s = strong vs = very strong, b = broad, p = polarized, dp =							larized	45	m	Lattice mode

Table VIII. Con't.

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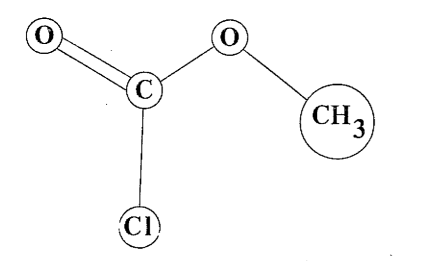
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FIGURE CEPTIONS

- Fig. 1 Possible conformations of methylchloroformate.
- Fig. 2 The structure of methylchloroformate in the principal axis system.
- Fig. 3 The Raman spectra of methylchloroformate. [A] The room temperature vapor. [B] The solid at 18.7°K.
- Fig. 4 The far infrared spectra of [A] methylchloroformate and [B] methylchloroformate-d₃. Transitions marked with arrows correspond to a small amount of HCl impurity.



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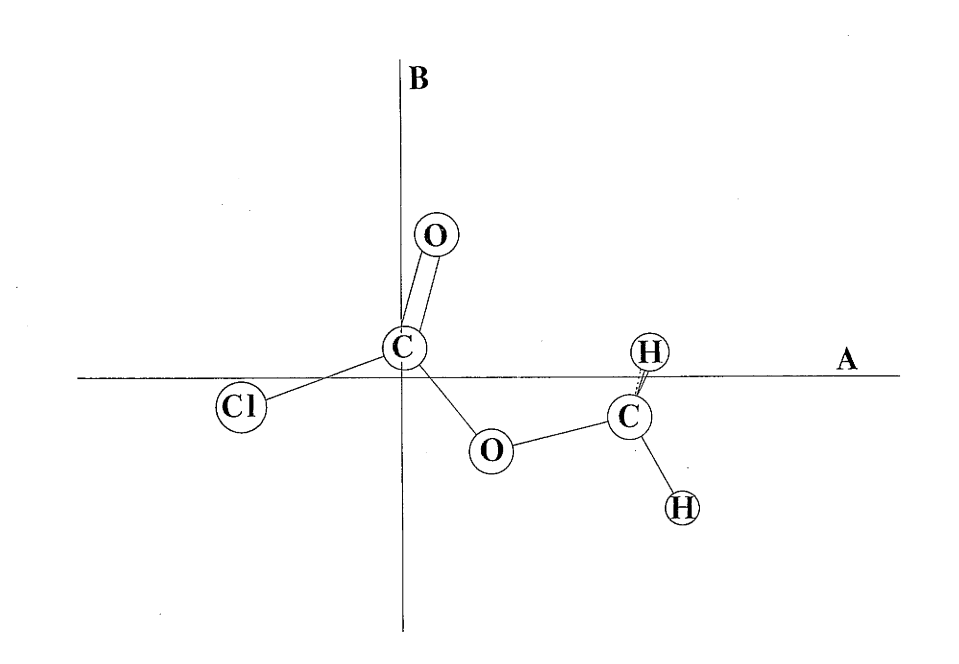
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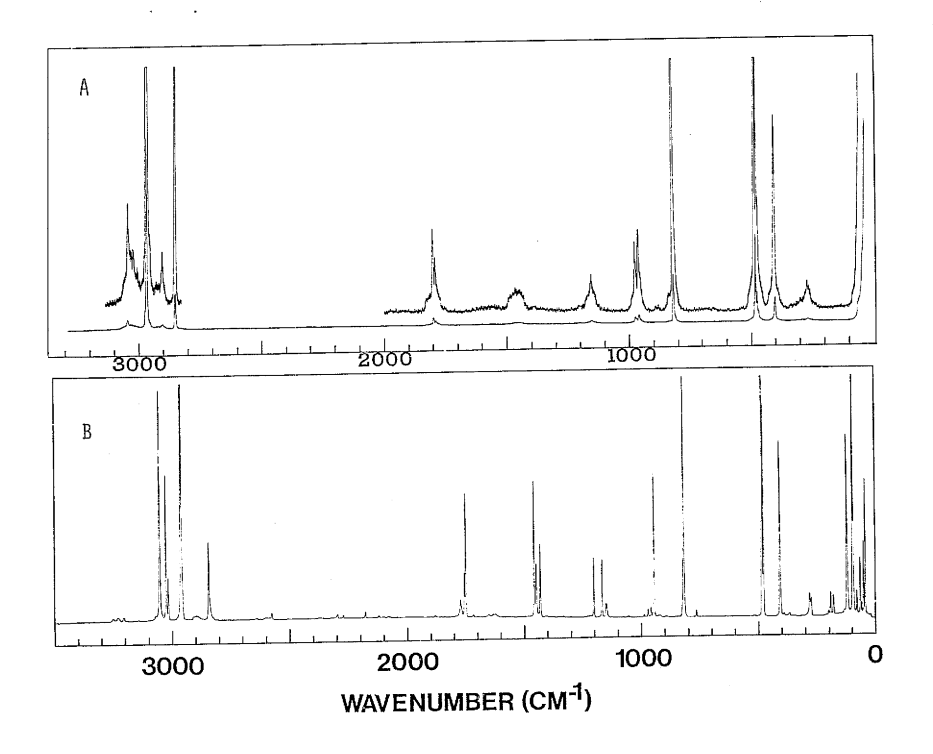
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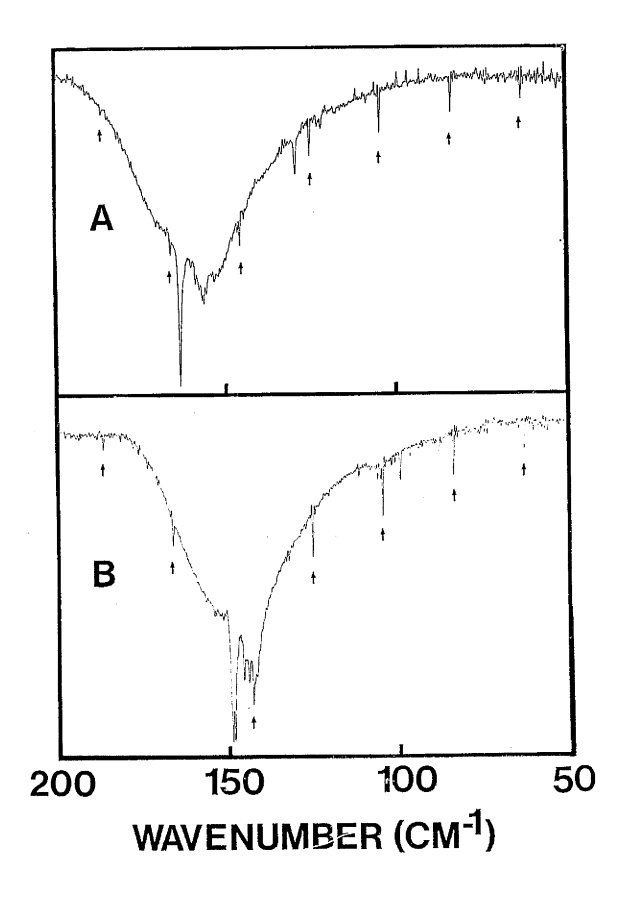
S-TRANS

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APPENDIX II

SPECTRA AND STRUCTURE OF ORGANOPHOSPHORUS COMPOUNDS. XVII. INFRARED AND RAMAN SPECTRA, VIBRATIONAL ASSIGNMENT AND BARRIERS TO INTERNAL ROTATION FOR t-BUTYLPHOSPHINE

Abstract

The infrared spectra of gaseous and solid tertiary-butylphosphine, $[(CH_3)_3CPH_2]$, have been recorded from 50 cm⁻¹ to 3500 cm⁻¹. The Raman spectra of gaseous, liquid and solid $(CH_3)_3CPH_2$ have been recorded from 10 to 3500 cm⁻¹. A vibrational assignment of the 42 normal modes has been made. A harmonic approximation of the methyl torsional barrier from observed transitions in the solid state gave a result of 4.22 kcal/mole and 3.81 kcal/mole in the gaseous state. Hot band transitions for the phosphino torsional mode have been observed. The potential function for internal rotation about the C-P bond has been calculated. The two potential constants were determined to be:

 $V_3 = 2.79 \pm 0.01$ kcal/mole and $V_6 = 0.07 \pm 0.01$ kcal/mole.

INTRODUCTION

Vibrational spectrosocopy has been shown to be a valuable tool in the determination of intramolecular potential functions governing internal rotation about single bonds.^{1, 2} This method has recently been extended to the study of asymmetric tops rotating against a molecular frame, as opposed to the more familiar symmetric tops such as $-CH_3$ and $-SiH_3$.³ If the molecular frame is asymmetric as well, it is possible for there to be spectroscopically distinct rotational isomers or conformers. Both ethylphosphine ($CH_3CH_2PH_2$) and isopropylphosphine $[(CH_3)_2CHPH_2]$ are examples of this type of system since the PH₂ group constitutes an asymmetric internal rotor. The internal rotation barrier for the PH2 top has been determined for both of these If, however, the molecular frame is symmetric, as is molecules. 4, 5 the case in tertiary-butylphosphine, $[(CH_3)_3CPH_2]$, only one conformation is possible and the potential function governing internal rotation is considerably simplified, becoming three-fold symmetric as in the case of methylphosphine (CH_3PH_2). Thus, the determination of this potential function for t-butylphosphine is a natural extension of the prior studies of ethyl- 4 and isopropylphosphine. 5 Also, it would be of interest to determine the barriers to internal rotation of the methyl groups in this molecule, as part of continuing studies ⁶, ⁷ of the top-top interactions in such "three-top" molecules.

Finally, the assignment of the vibrational spectrum of t-butylphosphine would finish the series of singly alkyl-substituted phosphines: methyl, ethyl and isopropyl. Group frequencies determined for the PH₂ moiety from these studies could be expected to apply to almost any hydrocarbon derivative.

EXPERIMENTAL

In general, all sample preparations and manipulations were performed using standard high vacuum techniques where possible. This was done in view of the relative instability of trivalent phosphorous compounds with respect to oxidation as well as to minimize exposure to the compounds, which are probably quite toxic and undeniably unpleasant smelling. Sample purifications were performed using fractional condensation (trap-to-trap) methods or more generally a low temperature, low pressure fractionating column.

The sample of $(CH_3)_3CPH_2$ was prepared by the reduction of $(CH_3)_2CHPCl_2$ with LiAlH₄ in di-n-butyl ether. After low temperature distillation, the purity of the $(CH_3)_3CPH_2$ was checked by mass spectrometry and ¹H, ¹³C and ³¹P NMR.

Mid-infrared spectra of $(CH_3)_3 CPH_2$ were obtained using a Digilab FTS-15B Fourier transform interferometric spectrometer. A germanium beamsplitter on a KBr substrate gave spectral coverage in the range from 3800 cm⁻¹ to 400 cm⁻¹. Spectra of the gaseous phase were obtained by using a 10 cm cell with KBr windows and typically less than 10 torr of sample pressure.

Atmospheric water vapor was removed from the spectrometer housing py purging with dry nitrogen. Since the FTS system uses a He-Ne laser as a frequency reference of relatively high stability, calibration is achieved by adjusting a software parameter, the laser wave length, to reproduce an observed vibrational transition of a suitable standard at its correct frequency. As a result, the accuracy of a measured frequency can be expected to be at least 0.1 cm⁻¹ throughout the spectrum.

Both apodization function and resolution were varied for each sample to attain the best quality spectrum, but a typical experiment would be run at an effective resolution of better than 0.5 cm⁻¹ using a modified "boxcar" apodization function for photometric accuracy.

Far infrared spectra were obtained by using the FTS-15B and substituting a mylar beamsplitter for the germanium one to obtain a spectral range of 400 to 80 cm⁻¹. A high pressure Hg arc lamp source was substituted for the glower used in the mid-infrared, but otherwise the operation of the instrument was the same as described above. Spectra of compounds in the gas phase were obtained by using 10 cm cells equipped with polyethylene windows. Typical sample pressures ranged from 100 to 300 torr, and spectra were taken at various resolutions betwhen 0.5 and 2.0 cm⁻¹. The spectrum of the solid phase sample was obtained by condensing the sample onto a wedged silicon plate cooled to \sim 15°K by a closed cycle He refrigerator.

Raman spectra were recorded to 4000 cm⁻¹ using a Cary 82 spectrometer equipped with either a Spectra Physics model 171 or a Coherent Radiation Labs model 53A argon ion laser operating on the 5145 Å line. Spectra of the gaseous phase were obtained using the Cary multipass accessory at sample pressures of \sim 300 torr. The spectrum of the liquid was recorded from samples in sealed glass capillaries. The spectrum of the solid was obtained by condensing the sample on a copper block maintained at \sim -190°C by boiling nitrogen. Polarization measurements were made using the standard Cary accessories. Frequencies measured for sharp, resolvable bands are expected to be accurate to at least ± 2 cm⁻¹.

VIBRATIONAL ASSIGNMENT

Tertiary-butylphosphine was assumed to adopt a fully staggered molecular conformation, and as such would possess a single plane of symmetry and belong to the point group C_s . However, if one ignores the hydrogen atoms attached to the phosphorus atom, the molecular symmetry would be C_{3v} . If the major portion of the molecule follows this "local" symmetry, the assignment of the vibrational spectra would be somewhat simplified. Pritchard and Nelson 8 were able to analyze the spectrum of t-butanol using the higher symmetry model for all but a few vibrational modes. Hence for (CH₃)₃CP there would be 24 normal modes, distributed $8A_1$, $4A_2$ and 12E. Under C_s symmetry, $(CH_3)_3CPH_2$ would exhibit 23A' and 19A" vibrational modes. Spectra taken of $(CH_3)_3 CPH_2$ in the fluid phases might be expected then to show groups of unresolved bands. However, spectra taken of the sample in the solid phase should allow observation of all 42 fundamentals since in the crystal closely spaced fundamentals should be resolvable. Where applicable, the origin of the vibrational mode under the hypothetical C_{3v} symmetry will be indicated in parenthesis as a guide to the assignment. To aid in the analysis of gas phase band contours, the principal moments of inertia for (CH₃)₃CPH₂ were calculated using a reasonable assumed structure. These calculations indicate that the molecule is very nearly a symmetric top, with the A and C inertial axes in the symmetry plane. Thus the A' vibrations are expected to show A/C

hybrid band contours with a sharp Q branch, and A" vibrations B type bands with no central maximum. The vibrational frequencies given in Table I for the A" modes in the infrared spectra of gaseous $(CH_3)_3CPH_2$ therefore reflect estimates of the band center. The vibrational assignments were guided by those of Pritchard and Nelson⁸ for $(CH_3)_3COH$ and those of Evans and Lo⁹ for $(CH_3)_3CC1$.

The Raman spectra of $(CH_3)_3 CPH_2$ in the gaseous, liquid and solid phases are shown in Fig. 1, and the mid-infrared spectra of $(CH_3)_3$ CPH_2 in the gaseous and solid phases are shown in Fig. 2. Fig. 3 depicts the far-infrared spectra of gaseous and solid $(CH_3)_3 CPH_2$. The observed frequencies and their assignments are given in Table I.

The C-H stretching region, 2870 to 3000 cm^{-1} , showed a dense cluster of bands in all sample phases in both the Raman and infrared spectra. In addition to the 9 fundamentals expected in this region, Durig, <u>et al.</u>, ¹⁰ have pointed out that additional strong bands may be expected to occur in this region, attributable to overtones of the antisymmetric methyl deformations in Fermi resonance with the stretching fundamentals. The bands in this region were not resolved even in the spectra of the solid phase, and no additional isotopic species were available for study. In view of these difficulties, attempts to assign this portion of the vibrational spectra were abandoned and the observed band maxima tabulated.

The two very strong bands at 2293 cm⁻¹ and 2278 cm⁻¹ in the spectra of the solid sample are undoubtedly the antisymmetric and symmetric P-H stretches, v_{28} and v_6 , respectively. As previous experience has indicated, ^{4, 5} only the A' mode is seen in the Raman spectra of the gas phase.

The advantage of recording spectra of the solid phase is easily seen in the analysis of the CH₃ antisymmetric deformations v_{29} , v_7 (E), v_{30} , v_8 (E), v_9 (A₁) and v_{31} (A₂). The Raman spectrum of gaseous (CH₃)₃CPH₂ shows only a broad, featureless band centered near 1450, but the Raman spectrum of the solid clearly shows all six fundamentals. The assignment of the CH₃ symmetric deformations, v_{10} (A₁), v_{11} and v_{32} (E) is also straightforward, although these modes are not seen in the Raman spectra of the fluid phases.

Assignment of the vibrational spectrum in the region between 1200 cm^{-1} and 800 cm^{-1} is clearly the most difficult. Twelve fundamental vibrations; 6 methyl rocks, 3 C-C stretches and the 3 PH_2 deformations, are all expected to occur in this region. In addition, severe mixing of several vibrational modes, especially the C-C stretches and methyl rocking modes, could make relative intensities unreliable as assignment aids. Three fundamentals of this group should fall considerably lower than the rest; v_{17} (A₁), the C-C symmetric stretch should be below 900 cm⁻¹, as should v_{37} , the PH₂ twist, and v_{18} , the PH₂ wag. Only v_{17} and $\nu^{}_{18}$ should exhibit sharp Q branches in the gaseous phase Raman spectrum, and only three bands are apparent in this region in the Raman spectrum of the solid phase. Thus, v_{17} , v_{37} and v_{18} are assigned to the bands at 836 cm^{-1} , 818 cm^{-1} and 795 cm^{-1} in the Raman spectrum of the solid sample. The frequency of v_{15} , the PH₂ scissoring mode, is well established 4, 5, 11 and is assigned to the band at 1074 cm⁻¹ in the mid-infrared spectrum of solid (CH₃)₃CPH₂. The C-C antisymmetric stretches, v_{33} and v_{12} (E), occurred near 1240 cm⁻¹ in both (CH₃)CC1⁹ and $(CH_3)COH;^8$ however, there are no bands in this region for $(CH_3)_3CPH_2$.

There are two strong bands at 1211 cm^{-1} and 1192 cm^{-1} in the Raman spectrum of the solid, and have been assigned to v_{33} and v_{12} , respectively. The slightly lower, weaker band at 1170 is assigned to v_{13} (A $_1$), a CH $_3$ rocking mode. The complex series of bands seen on the low frequency side of $\nu^{}_{13}$ in the gas phase spectra are most probably hot bands due to the methyl torsions or phosphino torsion. Although this assignment of $\boldsymbol{\nu}_{1,3}$ may seem to place this rocking mode quite high, the corresponding modes in $(CH_3)_3COH^8$ and $(CH_3)_3CC1^9$ were assigned to bands at 1189 cm⁻¹ and 1115 cm⁻¹, respectively. The sharp, polarized peak seen at 1029 $\rm cm^{-1}$ in the Raman spectra of the gaseous and liquid phases must be due to the A' methyl rocking mode, v_{16} . The sharp, moderately intense Q branch observed at 1090 cm^{-1} in the mid-infrared spectrum of gaseous $(CH_3)_3 CPH_2$ must then belong to the remaining A' methyl rocking mode, v_{14} . In terms of the C $_{3v}$ model symmetry, v_{14} and v_{34} are derived from a degenerate E vibration, as are v_{16} and v_{35} . Therefore v_{34} and v_{35} which would be expected to be clearly discernable only in the solid phase spectra, should fall near their respective A' modes, v_{14} and v_{16} . On this basis, v_{34} is assigned to a band seen at 1065 ${\rm cm}^{-1}$ in the spectra of the solid phase, and ${\rm v}_{35}$ to the band at 1036 cm⁻¹. The remaining rocking mode v_{36} (A₂) can then be assigned to the broad, depolarized band seen near 940 $\rm cm^{-1}$.

The assignment of v_{19} , the C-P stretch, is unambiguous, as it is usually $^{4, 5}$ the most intense band in the Raman spectrum below 2000 cm⁻¹. In contrast, this band, located near 590 cm⁻¹, is quite weak in the mid-infrared spectrum.

In the region below 400 cm^{-1} there should be 5 skeletal deformations

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as well as the 4 torsional vibrations. In the far infrared spectrum of gaseous $(CH_3)_3 CPH_2$ there are apparently two sharp Q branches at 352 cm⁻¹ and 346 cm^{-1} , also seen in the Raman spectrum of the gas. However, this structure disappears upon condensation, and only a single, strong, sharp band is seen in the solid phase spectra at 353 cm⁻¹. This band has been assigned to v_{21} (A₁), the CCC symmetric deformation, with the 346 cm^{-1} band assigned as a hot band. The other two CCC deformations, v_{38} and v_{20} (E), are assigned to the bands at 394 cm⁻¹ and 389 cm⁻¹, respectively, in the Raman spectrum of the solid. The CCP deformational modes, v_{39} and v_{22} (E), have been assigned to the bands at 297 cm⁻¹ and 295 cm⁻¹, respectively, in the Raman spectrum of the solid. The anomalously low intensity of v_{38} , v_{20} , v_{39} and v_{22} in the far infrared spectrum of solid (CH₃)CPH₂ is possibly attributable to crystal effects. Following the assignments of Durig, $\underline{et \ al.}$, for a series of t-butyl compounds, the bands at 278 cm^{-1} , 273 cm^{-1} , and 248 cm^{-1} in the far infrared spectrum of the solid have been assigned to the methyl torsional modes, v_{23} , v_{40} (E) and v_{41} (A₂). The strong band system centered near 176 $\rm cm^{-1}$ in the far infrared spectrum of the gas is undoubtedly due to the fundamental PH_2 torsion, v_{42} , and its hot bands. This band becomes a single sharp band centered at 200 cm⁻¹ upon solidification.

DETERMINATION OF THE TORSIONAL BARRIERS

In order to treat the problem of the barriers to internal rotation of the methyl groups in t-butylphosphine, the assumption was made that, so far as the methyl torsions are concerned, the molecule has C_{3v}

symmetry. Lide and Mann¹² have expressed the internal potential energy of such a system in a three dimensional Fourier expansion in terms of $3n\phi_1$, $3n\phi_2$, and $3n\phi_3$, where ϕ_j is the internal rotation angle of the jth top. If terms with n>l are ignored, the potential energy becomes: $2V = V_0 - V_1 \sum_{i=1}^{3} \cos 3\phi_i - V_2 \sum_{i>j} \cos 3\phi_i \cos 3\phi_j - V_3 \cos 3\phi_1 \cos 3\phi_2 \cos 4\phi_1 \cos 3\phi_2 \cos 4\phi_1 \cos 3\phi_2 \cos 4\phi_1 \cos 3\phi_1 \sin 3\phi_1 \sin 3\phi_j + V_5 \sum_{i\neq j\neq k} \cos 3\phi_i \sin 3\phi_j \sin 3\phi_k$ with the inclusion of the proper kinetic energy terms, the torsional

frequencies are given in the harmonic approximation by:¹²

$$v_{A} = [2(F-F^{-})(K + 2L)]^{\frac{1}{2}}$$

 $v_{E} = [2(F + 2F^{-})(K - L)]^{\frac{1}{2}}$

where

$$K = (9/2) (V_1 + 2V_2 + V_3)$$

L = (9/2)(V_A + V_5)

The kinetic terms for the present case were obtained by averaging the terms for each top, which were only slightly different. The results of a calculation using this approximation are shown in Table II for both the solid phase and gas phase data. If the interaction constant L is interpreted as a measure of the forces between hydrogen atoms on different CH_3 groups, then the negative sign found for L indicates that the net force is repulsive. Thus, the A_2 - type motion, in which the three tops move in phase, is more effective in keeping the hydrogen atoms apart than the E torsion, where one top is out of phase. In view of the small value found for L, the cosine interaction terms are probably negligible also and (2/9)K is a good approximation of the total barrier.

To determine the barrier to internal rotation of the PH_2 top, a computer program was used which is similar to one described by Lewis, <u>et al.</u>¹³ It is assumed that the torsional potential can be represented as a Fourier cosine series in the internal rotation angle:

$$V(\phi) = \left(\frac{1}{2n}\right) V_n(1 - \cos \phi).$$

Since the potential function must be three-fold symmetric, only terms where n=3 and 6 were used. The band centers of the $1 \div 0$ and $2 \div 0$ phosphino torsions were estimated from the observed far infrared spectrum of the gaseous phase. At this point, the program was allowed to iterate on the two potential constants, and a prediction made for the location of the $3 \div 2$ transition using the resulting potential function. A slight change in the band contour at 162 cm^{-1} had been noted previously and coincided reasonably well with the prediction. This band was then included in the program input and the potential function re-evaluated. The calculated $4 \div 3$ transition splits by $\simeq 3 \text{ cm}^{-1}$ as the energy levels near the top of the well, due to quantum mechanical tunnelling. ¹⁴ This predicted transition roughly coincided with another contour change seen at about 151 cm^{-1} ; however, this transition was not included in the fit. The results of this calculation are shown in Table III.

DISCUSSION

With regard to the vibrational assignment, several small points are probably ambiguous, but overall it is felt that the given assignment is quite reasonable. A vibrational study of the $(CH_3)_2 CPD_2$ isotope would confirm the assignment of PH₂ deformations, and might clarify the

assignment of the skeletal deformations if vibrational coupling of these modes is sufficient to cause an appreciable frequency shift.

The methyl barrier potential constants determined in this work may be compared with those found by Durig, $\underline{et \ al}$.,⁷ for some similar molecules observed in the solid state. The (2/9)K term for $(CH_3)_3COH$ of 4.13 kcal/mole goes to 4.41 kcal/mole in $(CH_3)_3CSH$, while the (2/9)L terms are 0.44 and 0.49 kcal/mole, respectively. In comparison, the (2/9)K term for $(CH_3)_3CNH_2$, is 4.28 kcal/mole and 4.22 kcal/mole for $(CH_3)_3CPH_2$, with the (2/9)L terms 0.70 and 0.15 kcal/mole, respectively. The general agreement in the magnitude of (2/9)K is reassuring; however, the trends in the magnitude of (2/9)L for the amine and phosphine are not clearly understood. The magnitude of the (2/9)L term is derived from the frequency splitting of the pseudo-E torsional mode and the pseudo-A₂ torsional mode. Differing degrees of intermolecular effects in the solid, such as hydrogen bonding, might be expected to produce some variation in this interaction term, but the trend should then also be apparent in the $(CH_3)_3COH \rightarrow (CH_3)_3CCH$ series. Similar arguments concerning lone pair effects or lengths of the tertiary carbon to unique substituent bond should be evident in the $(CH_3)_3COH \rightarrow (CH_3)_3CSH$ series, also. The most probable explanation is that the methyl torsional modes in $(CH_3)_3CNH_2$ are severly coupled with the amine torsional vibration which falls between the two methyl torsional regions. Thus the large interaction term seen for $(CH_3)_3 CNH_2$ is due in part to interactions of the methyl torsions with the NH₂ torsion.

The barrier to internal rotation of the PH_2 group in $(CH_3)_3CPH_2$, 2.79 kcal/mole, is somewhat higher than that determined by Tsuboi,

<u>et al.</u>,¹⁵ for $(CH_3)_3CNH_2$, 2.36 kcal/mole. The barrier for t-butylphosphine is nearly a kilocalorie/mole higher than the barrier for CH_3PH_2 ,¹¹ determined to be 1.96 kcal/mole. Similarly the barrier in CH_3NH_2 is 1.95 kcal/mole.¹⁶ From these results, one might infer that electrostatic interactions of the PH bonds and phosphorus lone pair are greater with CC bonds than with CH bonds, if these interactions are predominantly repulsive.

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Infrared gas	Rel. Int.	Infrared <u>so</u> lid	Re]. Int.	Raman gas	Rel. Int.	Raman Liquid	Rel. Int.	Raman Solid	Rel. Int.	Assignment
2967 2960 2954	VS VS VS							2959	s,sh	
		2949	٧s	2961	s p	2953	s p	2949	vvs	
2946	vs,sh			2947	s,sh p	2936	s,sh p	2943	s,sh	
		2937	٧s							
				2934	vs p	2925	vs p	2930	s,sh	
		2924	S							v1.22,v3,v4,v5 vC−H sym. 24,v25,v26,v27 vC−H ssym.
								2910	m,sn	J243V25,V26,V27 VC-H asy
		2892	S					2894	vvs	
2906	s			2907	vvs p					
		2877	s					2878	s,sh	
		2857	vs					2861	s	
2873	s			2875	vs p	2865	vs p			
								2850	s,sh]	
				2786	m p	2767	m p	2769	m	20 ₈ ?
				2724	s p	2715	m p	2707	m	2010?
		2279	vvs			2284		2293	vvs	ν₂ в νP−H
2292	٧VS	2293	vvs	2292	vvs p			2278	vvs	v_6

Table I Vibrational Frequencies (cm⁻¹) and their Assignments for t-Butylphosphine.^a

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Infrared gas	Rel. Int.	Infrared solid	Rel. Int.	Raman gas	Rel. Int.	Raman Liguid	Rel. Int.	Raman Solid	Rel. Int.	Assignment
		1467	m,sh]	· ·	ſ		1473	W	v ₂ 9
1476	s,sh							1463	m	V7
		1462	٧S					1462	m,sh	
				1450	w,brd * dp?	1458	w,brd dp?			≻δCH₃ asym.
1466	s	1457	s,sh		մի։			1457	s	ve ·
1456	s,sh	1442	m,sh			1445	w ,br d	1442	S	ν ₉
		1436	w,sh			Į		1438	m,sh	V31
1370	s	1368	m,sh	,		-				ر ۱۳
		1363	s,sh					1363	vw	vıı ≽õCH₃ sym.
1361	s,sh									Çeng 25m
		1361	S					1361	vw,sh	V32
1204	m	1211	w,sh	1205	w,sh			1211	S	vaa Vee asym
1198	Ш	1192	m	1199	m p	1192	m,brd p	1192	s	v ₁₂ ∫ ⁵ C-C abj
1192	m	1169	W	1192	m p	1173	w,sh p	1170	w,sh	ν ₁₃ ρCH ₃
1183 1172	m W			1184 1173	w,sh w,sh					"hot" bands of v
1090	M	1096	s			1089	vw,sh p	1092	VW	ν ₁₄ ρCH ₃
1074	m	1074	m	1080	vw,brd p	1072	ww,brd p	1075	m	$v_{1s} \rho PH_2 sic$
		1065	w,sh					1065	w,sh	V34]
		1036	w,sh	חמתו	MI-1 - 5	1029	10.4 5	1036	w,sh	ν ₃₅ }ρCH ₃
1029	т	1029	m	1029	лм b	1029	vw p	1032	W	Vie
940	W	938	W	940	vw dp	938	vw,brd dp	937	m	V3E

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Table 1 (conclused)	Table I	(continued)
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nfrared gas	Rel. Int.	Infrared gas	Rel. Int <u>.</u>	Raman gas	Rel. Int.	Raman Liquid		Raman Solid	Rel. Int.	As	signment
834	m	834	W	835	m p	833	w p	836	vs	V17	νc-c ^{sym.}
826	S	816	m	825	w dp	820	vw,sh dp?	818	m	V37	δPH_2 twist
799	s	793	vs	800	s p	798	m p	795	m	ν _{1 в}	δPH_2 wag
589 ^b	m	592	W	590	vs p	589	vs p	593	vs	ν19	^v с-Р
394	W							394	vw,sh	V36	ôCCC asym.
		383 ^b	vw	382	vw,sh	386	vw,sh				
389	w							389	m	V20	δCCC asym.
352	5	354	S	354	s p	354	m p	353	s	V ₂₁	δCCC sym.
346	m			347	w,sh p					"ho	t" band of v_2
								2 97	w,sh	ν39	δCCP sym.
285	W	295	VW			287	vw,brd dp?				500D
		270	a ch	280	w,brd			295	W	V22 V23	δCCP asym. τCH ₃
266	w	278	s,sh	280	w,bru	272	vw,sh p?	276	W	v ₂ 3	10113
200		273	s							V40	τCH ₃
227	W	248	w,brd			239	vvw,brd dp	248	٨M	v_{41}	⊤CH₃
181	S	200	vs	172	vvw,brd d	p 182	vvw,brd dp	199	VW	V42	τPH₂
171	s									_ "hot	"band of v_{42}
								77			
								63		[
								50		Lat م	tice modes
								42			
								32		<u> </u>	

^aAbbreviations used: s, strong; m, medium; w, weak; v, very; brd, broad; sh, shoulder; p, polarized, dp, depolarized. ^bThis and lower frequencies obtained from the far infrared spectra.

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TABLE II METHYL TORSIONAL FREQUENCIES AND POTENTIAL CONSTANTS FOR T-BUTYLPHOSPHINE

-	Solid	Gas
$v_{\rm E}^{\rm a}({\rm cm}^{-1})$	275.5	266
ν _A (cm ⁻¹)	248	227
F(cm ⁻¹)	5.3559	5.3559
$F(cm^{-1})$	0.0278	0.0278
2/9K(kcal/mole)	4.22	3.81
2/9L(kcal/mole)	-0.15	-0,20

a) $\frac{1}{2}(A^{1} + A^{n})$

	frequency							
Transition	obs.	calc.	obscalc.					
1~0	181	180.9	0.1					
2+1	172	172.2	-0.2					
3+2	162	161.8	0.2					
4⊷3¯		146.1	4.9					
4←3 ⁻ 4←3 ⁺	(151) ²	149.1	0.9					
۷ ₃ ۷ ₆	976 ± 4 ^b -25 ± 2	1	± 0.01 kca1/mole ± 0.01 kcal/mole					

TABLE III PHOSPHINO TORSIONAL FREQUENCIES (cm⁻¹) AND POTENTIAL CONSTANTS

a) estimatedb) quoted error limits represent one standard deviation

FIGURE CAPTIONS

Figure 1 Raman spectra of t-butylphosphine

- A) gaseous
- B) liquid
- C) solid

Figure 2 Mid-infrared spectra of t-butylphosphine

- A) gaseous, approximately 2 torr sample pressure
- B) gaseous, approximately 40 torr sample pressure
- C) solid
- D) solid, more sample deposited

Figure 3 Far infrared spectra of t-butylphosphine

- A) gaseous, approximately 400 torr
- B) solid^a
- a) pronounced baseline "roll" is due to interference fringes from the sample film

