

Tables of Molecular Vibrational Frequencies

Consolidated Volume II

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The compilations of fundamental vibrational frequencies of molecules previously published as Tables of Molecular Vibrational Frequencies Part 5, Part 6, Part 7, and Part 8, which appeared in the Journal of Physical and Chemical Reference Data in 1972, 1973, and 1974, have been revised. This Consolidated Volume II includes data on a total of 212 molecules in addition to those on 223 molecules included in Volume I (NSRDS-NBS 39). Selected values of the fundamental vibrational frequencies are given for each molecule, together with observed infrared and Raman spectral data and citations to the original literature. The selection of vibrational fundamentals has been based on careful studies of the spectral data and comprehensive normal-coordinate analyses. An estimate of the accuracy of the selected values is included. The tables provide a convenient source of information for those who require vibrational energy levels and related properties in molecular spectroscopy, thermodynamics, analytical chemistry, and other fields of physics and chemistry.

Key words. Fundamental frequencies; infrared spectra; polyatomic molecules; Raman spectra; vibrational frequencies.

1. Introduction

Establishing the assignment of molecular vibrational frequencies has fundamental importance in elucidating various problems in physics and chemistry. The information concerning the force field and motion of atoms in a molecule can be most directly derived from its vibrational frequencies. If all the vibrational frequencies of a molecule are known, as well as the molecular structure, thermodynamic quantities can be easily computed on the ideal gas model. Thus, the need for a tabulation of evaluated reference data on molecular vibrational frequencies has often been felt by many investigators. In 1964 a project for producing such tables was initiated at the University of Tokyo in cooperation with the National Standard Reference Data System of the National Bureau of Standards. The evaluated data resulting from this project were first published in the three parts of Tables of Molecular Vibrational Frequencies [1]¹. A Consolidated Volume [2] of these tables appeared in 1972 which includes revised versions of all the tables in [1] plus tables for 52 additional molecules (a total of 223 molecules). Tables covering 160 more molecules have appeared as Parts 5, 6, 7, and 8 of the series in earlier issues [3, 4, 5, 6] of this journal.

The present volume consists of Parts 5, 6, 7, and 8 after extensive revision in the light of new experimental data.

2. Molecules Selected and Their Ordering

The present volume contains tables of fundamental vibrational frequencies for 212 molecules. The molecules were selected from basic organic and inorganic molecules for which the vibrational assignments have been established with little

ambiguity. The effort of extending the tables to many other important molecules is continuing in this laboratory. Diatomic molecules and electronically excited species are not included in this volume, since [7, 8, 9] contain good compilations of data for them. Rotational isomers are treated as independent molecular species, and a separate table is made for each of the isomers. When the gas and liquid state spectra are significantly different from each other, they are tabulated separately.

The molecules are ordered according to the following rules:

(a) Number of carbon atoms.

(b) Total number of atoms.

(c) Molecular shape: linear, planar, and non-planar.

(d) Molecular symmetry, in descending order of the number of symmetry elements. Isotopically substituted molecules directly follow the normal species regardless of their symmetry.

(e) Atomic number of main atoms.

(f) Atomic number of the other atoms.

Molecules are first divided into groups by the items (a) and (b) and the ordering of molecules in each group is given by the items (c), (d), (e), and (f). A complete list in the order presented is given at the beginning of the tables. Indices by compound name and empirical formula follow the tables.

3. Description of the Tables

3.1. Symmetry

The symmetry (point group) of each molecule is given by the Schoenflies notation. Detailed discussions of symmetry properties will be found in [10] and [11].

3.2. Symmetry Number

The symmetry number, σ , is used in the calculation of thermodynamic quantities. It is the number of indistinguishable positions into which the molecule can be transformed by simple rigid rotations. A general discussion and pertinent formulas may be found in [11], page 508.

¹ Figures in brackets indicate the literature references in section 5.

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3.3. Symmetry Species

In the table the normal modes are divided into the symmetry species of the point group to which the molecule belongs. The ordering of species in each point group is given in table I, which is a summary of tables 12–30 of [11]. When a molecule has two or three planes of symmetry, the relationship between the vibrational modes and symmetry species cannot be defined uniquely. In such cases we generally follow the notation adopted in [11].

TABLE I. Ordering of symmetry species

(In the present article small letters are used to designate the species of fundamental frequencies)

Point group	Symmetry species
C_2	A, B
C_s	A', A''
C_i	A_g, A_u
C_{2v}	A_1, A_2, B_1, B_2
C_{2h}	A_g, A_u, B_g, B_u
D_2	A, B ₁ , B ₂ , B ₃
D_{2h}	$A_g, A_u, B_{1g}, B_{1u}, B_{2g}, B_{2u}, B_{3g}, B_{3u}$
C_{3v}	A_1, A_2, E
D_3	A_1, A_2, E
C_{3h}	A_1, A_2, E_1, E_2
$C_{\infty v}$	$\Sigma^+, \Sigma^-, \pi, \Delta, \Phi, \dots$
C_{4v}, D_4, D_{2d}	A_1, A_2, B_1, B_2, E
C_{6v}, D_6	$A_1, A_2, B_1, B_2, E_1, E_2$
D_{3d}	$A_{1g}, A_{1u}, A_{2g}, A_{2u}, E_g, E_u$
D_{4d}	$A_1, A_2, B_1, B_2, E_1, E_2, E_3$
D_{3h}	$A_1', A_1'', A_2', A_2'', E', E''$
D_{6h}	$A_1', A_1'', A_2', A_2'', E_1', E_1'', E_2', E_2''$
D_{4h}	$A_{1g}, A_{1u}, A_{2g}, A_{2u}, B_{1g}, B_{1u}, B_{2g}, B_{2u}, E_g, E_u$
D_{6h}	$A_{1g}, A_{1u}, A_{2g}, A_{2u}, B_{1g}, B_{1u}, B_{2g}, B_{2u}, E_{1g}, E_{1u}, E_{2g}, E_{2u}$
$D_{\infty h}$	$\Sigma_g^+, \Sigma_u^+, \Sigma_g^-, \Sigma_u^-, \pi_g, \pi_u, \Delta_g, \Delta_u, \Phi_g, \Phi_u, \dots$
C_3	A, E
C_6	A, B, E ₁ , E ₂
S_6	A_g, A_u, E_g, E_u
C_{3h}	A', A'', E', E''
C_{4h}	$A_g, A_u, B_g, B_u, E_g, E_u$
C_{6h}	$A_g, A_u, B_g, B_u, E_{1g}, E_{1u}, E_{2g}, E_{2u}$
T_d, O	A_1, A_2, E, F_1, F_2
O_h	$A_{1g}, A_{1u}, A_{2g}, A_{2u}, E_g, E_u, F_{1g}, F_{1u}, F_{2g}, F_{2u}$
T	A, E, F

3.4. Numbering of Frequencies

The numbering is indicated by ν_i given in the second column of each table. The normal modes are first grouped into symmetry species, and then those in each species are ordered from higher to lower values of the frequency. However, we

always denote the bending vibration of a linear triatomic molecule as ν_2 , following the widely accepted nomenclature [12]. For the C_{2v} type of molecule we adopt the numbering given in [11], although it is based on D_{3h} symmetry. For some deuterated compounds the frequencies are arranged so that the same ν_i numbering is given to the corresponding vibrational modes of deuterated and normal compounds.

3.5. Approximate Type of Mode

The approximate type of mode given in the third column of each table is the local symmetry coordinate which makes the maximum contribution to the normal mode. Local symmetry coordinates are defined for several chemical groups in table II. It should be emphasized that two or more local symmetry coordinates are often coupled strongly in a normal coordinate, and the approximate type of mode given in the table has only limited significance in such a case.

TABLE II. Definition of local symmetry coordinates

(a) Local symmetry coordinates for the CH ₃ group (see fig. 1a)	CH ₃ symmetrical stretching: $(\Delta r_1 + \Delta r_2 + \Delta r_3) / \sqrt{3}$
	CH ₃ degenerate stretching: $(2\Delta r_1 - \Delta r_2 - \Delta r_3) / \sqrt{6}$
	$(\Delta r_2 - \Delta r_3) / \sqrt{2}$
	CH ₃ symmetrical deformation:
	$(\Delta \alpha_{23} + \Delta \alpha_{31} + \Delta \alpha_{12} - \Delta \beta_1 - \Delta \beta_2 - \Delta \beta_3) / \sqrt{6}$
	CH ₃ degenerate deformation: $(2\Delta \alpha_{23} - \Delta \alpha_{31} - \Delta \alpha_{12}) / \sqrt{6}$
	$(\Delta \alpha_{31} - \Delta \alpha_{12}) / \sqrt{2}$
	CH ₃ rocking: $(2\Delta \beta_1 - \Delta \beta_2 - \Delta \beta_3) / \sqrt{6}$
	$(\Delta \beta_2 - \Delta \beta_3) / \sqrt{2}$
(b) Local symmetry coordinates for the CH ₂ group (see fig. 1b)	CH ₂ symmetrical stretching: $(\Delta r_1 + \Delta r_2) / \sqrt{2}$
	antisymmetrical stretching: $(\Delta r_1 - \Delta r_2) / \sqrt{2}$
	CH ₂ scissors: $(4\Delta \alpha - \Delta \beta_{1x} - \Delta \beta_{2x} - \Delta \beta_{2y}) / \sqrt{20}$
	CH ₂ wagging: $(\Delta \beta_{1x} + \Delta \beta_{2x} - \Delta \beta_{1y} - \Delta \beta_{2y}) / 2$
	CH ₂ twisting: $(\Delta \beta_{1x} - \Delta \beta_{2x} - \Delta \beta_{1y} + \Delta \beta_{2y}) / 2$
	CH ₂ rocking: $(\Delta \beta_{1x} - \Delta \beta_{2x} + \Delta \beta_{1y} - \Delta \beta_{2y}) / 2$
(c) Local symmetry coordinates for the CH group (see fig. 1c)	CH stretching: Δr_{CH}
	CH bending: $(2\Delta \beta_{HX} - \Delta \beta_{HY} - \Delta \beta_{HZ}) / \sqrt{6}$
	$(\Delta \beta_{HY} - \Delta \beta_{HZ}) / \sqrt{2}$
(d) Local symmetry coordinates for the planar CH ₂ group (see fig. 1d)	CH ₂ symmetrical stretching: $(\Delta r_1 + \Delta r_2) / \sqrt{2}$
	CH ₂ antisymmetrical stretching: $(\Delta r_1 - \Delta r_2) / \sqrt{2}$
	CH ₂ scissors: $(2\Delta \alpha - \Delta \beta_1 - \Delta \beta_2) / \sqrt{6}$
	CH ₂ rocking: $(\Delta \beta_1 - \Delta \beta_2) / \sqrt{2}$
	CH ₂ wagging: $\Delta \theta \cdot \sin \alpha$
(e) Local symmetry coordinates for the planar CH group (see fig. 1e)	CH stretching: Δr_{CH}
	in-plane CH bending: $(\Delta \beta_{HX} - \Delta \beta_{HY}) / \sqrt{2}$
	out-of-plane CH bending: $\Delta \theta_H \cdot \sin \gamma_{XY}$

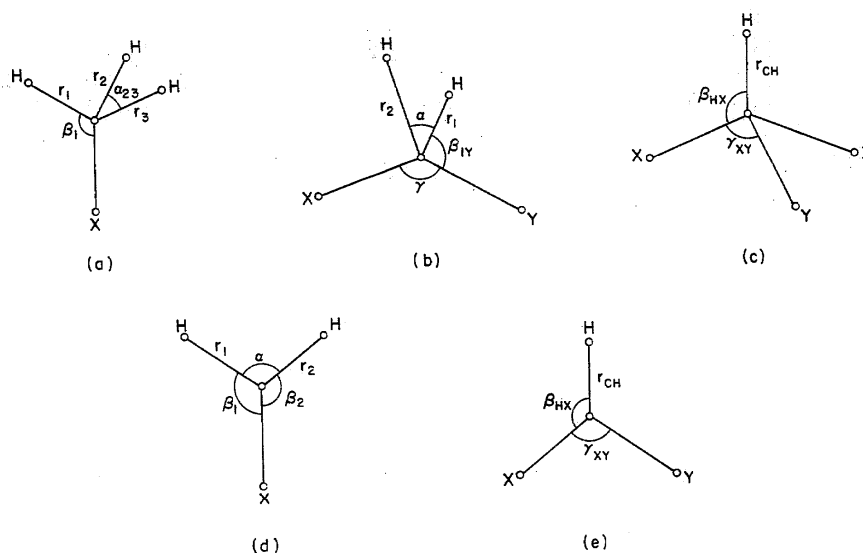


FIGURE 1. Parameters of methyl, methylene, and methine groups.

The following abbreviations are used for the type of mode:

stretch.	stretching
deform.	deformation
rock.	rocking
twist.	twisting
wag.	wagging
scis.	scissors
bend.	bending
sym. or s-	symmetrical
anti. or a-	antisymmetrical
deg. or d-	degenerate
ip-	in-plane
op-	out-of-plane

The plane to which the in-plane and out-of-plane expressions refer is the molecular plane of a planar molecule or the symmetry plane of a general molecule belonging to point group C_s . Local symmetry coordinates of the CX_3 groups attached to a relatively large molecule are designated as s-stretch, s-deform., d-stretch., and d-deform. In such a molecule with low symmetry none of the normal vibrations are genuinely "symmetrical" or "degenerate" with respect to the three-fold symmetry axis of the CX_3 group. However, the

notation is retained because it is convenient for indicating the correspondence between similar modes in large and small molecules.

3.6. Selected Value of Frequency²

The fundamental frequency ν_i is defined as the difference between the term values $G(\nu_i = 1, \text{ all other } \nu_j = 0)$ and $G(\nu_i = 0, \text{ and other } \nu_j = 0)$ expressed in cm^{-1} . Fundamental frequencies rather than harmonic frequencies (ω_j) are listed in the table. Although harmonic frequencies are of greater physical significance, they are accurately known only for a small number of polyatomic molecules. The selected values are rounded to the nearest 1 cm^{-1} .

The letter code, A, B, C, D, or E following the selected value of frequency indicates the evaluator's judgment of the accuracy of the value. The basis for estimating accuracy of an observed frequency is given in table III, together with the range of uncertainty in cm^{-1} for each grade.

² In keeping with the universally accepted convention in molecular spectroscopy, the fundamental frequencies are expressed in their wavenumber (cm^{-1}) equivalents. The actual frequency in units of hertz may be obtained by multiplying the numbers in these tables by the speed of light expressed in centimeters per second.

TABLE III. Uncertainty code for the selected values of frequencies

Notation	Uncertainty	Basis ^a
	cm ⁻¹	
A	0 ~ 1	(i) Gas, grating spectrometer, rotational fine structure accurately analyzed. (ii) Gas, grating spectrometer, a sharp <i>Q</i> branch.
B	1 ~ 3	(i) Gas, grating spectrometer, rotational fine structure partly analyzed. (ii) Gas, prism spectrometer, fairly high resolution (e.g., 700 ~ 1000 cm ⁻¹ for NaCl prism).
C	3 ~ 6	(i) Gas, prism spectrometer, low resolution (e.g., 1000 ~ 2000 cm ⁻¹ for NaCl prism). (ii) Solid, liquid or solution, accurate measurement.
D	6 ~ 15	(i) Gas, prism spectrometer, very low resolution (e.g., > 2000 cm ⁻¹ for NaCl prism). (ii) Solid, liquid or solution, inaccurate measurement.
E	15 ~ 30	(i) Value estimated from Fermi resonance doublet. (ii) Value estimated from overtone or combination tone. (iii) Calculated frequency.

^a The uncertainty assigned here to each method of measurement is a typical value; greater accuracy is often achieved with some of the methods.

Frequencies derived from infrared and Raman measurements in the gaseous state are chosen unless otherwise mentioned. When a detailed analysis of the rotational fine structure of an infrared band is available, the band center ν_0 is chosen as the fundamental frequency and given the uncertainty code A (see below). For a well-analyzed perpendicular band of a symmetric top molecule, the frequency listed contains the nonvibrational part $A' \zeta^2$, where A' is the rotational constant of the vibrational level and ζ is the Coriolis coupling constant. This is in accord with the definition of ν_0 given in [11], page 404 and equation (IV, 60).

When the spectra in the gaseous state are not available, the frequencies observed in the liquid or solid state are listed. When no spectral data have been obtained, the results of normal vibration calculations or of some other methods of estimating frequencies are listed with the grade D or E.

The torsional frequency may be calculated using the barrier height and reduced moment derived from microwave spectroscopy. The value obtained in this way is given as MW (frequency in cm⁻¹) in the "Comments" column or as a footnote for comparison with the value observed or calculated by the normal coordinate treatment. Microwave data are taken from [13] unless otherwise noted.

For many molecules the assignments given in the literature have been checked by normal vibration calculations carried out in this laboratory as part of the project. Revisions in some assignments have been made as a result of these calculations. The details of the normal coordinate treatment and evaluation of force constants may be found in [14].

Thermodynamic quantities may be computed in most cases by employing the harmonic oscillator partition function and by assuming that the harmonic frequencies are not much different from the fundamental frequencies given here. Such an approximation is not adequate, however, for molecules with highly anharmonic motions such as internal rotation, inversion, and ring-puckering. The vibrational partition function

should be formed for these molecules by summing the terms due to the individual energy levels.

3.7. Infrared and Raman Spectra

The observed infrared and Raman frequencies are given in the fifth and sixth columns of each table. Rough estimates of relative intensities, band shapes, and polarization characteristics are also given. An additional significant figure is included here when warranted. The abbreviations used here are as follows:

VS	very strong
S	strong
M	medium
W	weak
VW	very weak
ia	inactive
b	broad
vb	very broad
sh	shoulder
p	polarized
dp	depolarized

For some molecules the relative intensities of Raman lines are indicated by the numbers from one to ten in accordance with the tradition widely used. These estimates of intensity are taken from the original references without any attempt at critical evaluation.

3.8. Comments

In the last column of each table brief comments are added to give special information which is not indicated in the preceding columns. The abbreviations used in this column are as follows:

FR	Fermi resonance with an overtone or a combination tone indicated in the parentheses.
OC	Frequency estimated from an overtone or a combination tone indicated in the parentheses.
CF	Calculated frequency.
SF	Calculation shows that frequency approximately equals that of the vibration indicated in the parentheses.
OV	Overlapped by the band indicated in the parentheses.
MW	Torsional frequency calculated from microwave spectroscopic data.
RP	Frequency determined by the Ritz principle.
EL	Derived from analysis of electronic transitions.

3.9. Footnotes and References

The footnote is used to supply other necessary information which cannot be placed simply in the column of Comments. The references accompanying the table are not comprehensive. Only the papers relevant to the present tabulation are cited. The abbreviations IR, R, MW, and Th stand for infrared, Raman, microwave, and theoretical, respectively.

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5. References

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List of Tables of Molecular Vibrational Frequencies

C ₀ -triatomic molecules			No.		
No. 1	Krypton difluoride	KrF ₂	23	Nitrosyl bromide	¹⁶ O ¹⁴ NBr
No. 2	Nitrogen dioxide	¹⁴ NO ₂	24	Nitrosyl bromide	¹⁶ O ¹⁵ NBr
No. 3	Nitrogen dioxide	¹⁵ NO ₂	25	Nitrosyl bromide	¹⁸ O ¹⁵ NBr
No. 4	Ozone	¹⁶ O ₃	26	Hypofluorous acid	HOF
No. 5	Ozone	¹⁸ O ₃	27	Hypochlorous acid	HOCl
No. 6	Water	H ₂ ¹⁸ O	28	Hypochlorous acid-d	DOCl
No. 7	Sulfur dioxide	S ¹⁸ O ₂	29	Thionitrosyl-S-fluoride	NSF
No. 8	Sulfur dichloride	SCl ₂	30	Thionitrosyl-S-chloride	NSCl
No. 9	Germanium (II) chloride	GeCl ₂	C ₀ -four-atomic molecules		
No. 10	Tin (II) fluoride	SnF ₂	No. 31	Boron trifluoride	¹⁰ BF ₃
No. 11	Tin (II) chloride	SnCl ₂	No. 32	Boron trifluoride	¹¹ BF ₃
No. 12	Tin (II) bromide	SnBr ₂	No. 33	Sulfur trioxide	SO ₃
No. 13	Lead (II) fluoride	PbF ₂	No. 34	Ammonia-t ₃	NT ₃
No. 14	Lead (II) chloride	PbCl ₂	No. 35	Nitrogen trifluoride	¹⁵ NF ₃
No. 15	Nitrosyl fluoride	¹⁶ O ¹⁴ NF	No. 36	Aluminum trichloride	AlCl ₃
No. 16	Nitrosyl fluoride	¹⁶ O ¹⁵ NF	No. 37	Phosphorus triiodide	PI ₃
No. 17	Nitrosyl fluoride	¹⁸ O ¹⁴ NF	No. 38	Arsenic trifluoride	AsF ₃
No. 18	Nitrosyl fluoride	¹⁸ O ¹⁵ NF	No. 39	Arsenic triiodide	AsI ₃
No. 19	Nitrosyl chloride	¹⁶ O ¹⁴ NCl	No. 40	trans-1,2-Difluorodiazine	N ₂ F ₂
No. 20	Nitrosyl chloride	¹⁶ O ¹⁵ NCl	No. 41	Nitryl fluoride	F ¹⁴ NO ₂
No. 21	Nitrosyl chloride	¹⁸ O ¹⁴ NCl	No. 42	Nitryl fluoride	F ¹⁵ NO ₂
No. 22	Nitrosyl chloride	¹⁸ O ¹⁵ NCl	No. 43	Nitryl chloride	Cl ¹⁴ NO ₂
			No. 44	Nitryl chloride	Cl ¹⁵ NO ₂

No. 45	Chlorine trifluoride	ClF_3	No. 103	Nitric acid-d	D^{14}NO_3
No. 46	Bromine trifluoride	BrF_3	No. 104	Nitric acid	H^{15}NO_3
No. 47	Hydrogen peroxide	H_2O_2	No. 105	Nitric acid-d	D^{15}NO_3
No. 48	Hydrogen persulfide	H_2S_2	No. 106	Fluoronitrate	$\text{FO}^{14}\text{NO}_2$
No. 49	Disfluoro disulphane	F_2S_2	No. 107	Fluoronitrate	$\text{FO}^{15}\text{NO}_2$
No. 50	Hydrazoic acid	HN_3	No. 108	Chloronitrate	$\text{ClO}^{14}\text{NO}_2$
No. 51	Hydrazoic acid	DN_3	No. 109	Chloronitrate	$\text{ClO}^{15}\text{NO}_2$
No. 52	Disfluoroamine	NF_2H	No. 110	Phosphoryl dichlorofluoride	POCl_2F
No. 53	Chlorodifluoroammonia	NClF_2	No. 111	Phosphoryl dibromofluoride	POBr_2F
No. 54	Dichlorofluoroammonia	NCl_2F	No. 112	Thiophosphoryl chlorodifluoride	PSClF_2
No. 55	Thionyl fluoride	SOF_2	No. 113	Thiophosphoryl dichlorofluoride	PSCl_2F
No. 56	Thionyl chloride	SOCl_2			
No. 57	Thionyl bromide	SOBr_2			
C₀-five-atomic molecules					
No. 58	Titanium tetrachloride	TiCl_4	No. 114	Phosphorus pentafluoride	PF_5
No. 59	Titanium tetrabromide	TiBr_4	No. 115	Phosphorus trichloride difluoride	PCl_3F_2
No. 60	Titanium tetraiodide	TiI_4	No. 116	Phosphorus pentachloride	PCl_5
No. 61	Zirconium tetrachloride	ZrCl_4	No. 117	Vanadium pentafluoride	VF_5
No. 62	Zirconium tetrabromide	ZrBr_4	No. 118	Arsenic pentafluoride	AsF_5
No. 63	Zirconium tetraiodide	ZrI_4	No. 119	Antimony pentachloride	SbCl_5
No. 64	Ruthenium tetroxide	RuO_4	No. 120		
No. 65	Hafnium tetrachloride	HfCl_4	No. 121	Tellurium hexafluoride	TeF_6
No. 66	Hafnium tetrabromide	HfBr_4	No. 122	Iridium(VI)fluoride	IrF_6
No. 67	Hafnium tetraiodide	HfI_4	No. 123	Sulfur chloride pentafluoride	SClF_5
No. 68	Osmium tetroxide	Os^{16}O_4	No. 124	Tungsten chloride pentafluoride	WClF_5
No. 69	Osmium tetroxide	Os^{18}O_4	No. 125		
No. 70	Thionitrosyl trifluoride	NSF_3	No. 126		
No. 71	Trifluorosilane	SiHF_3	No. 127		
No. 72	Trifluorosilane-d	SiDF_3	No. 128		
No. 73	Trichlorosilane	SiHCl_3			
No. 74	Trichlorosilane-d	SiDCl_3	No. 129		
No. 75	Tribromosilane	SiHBr_3	No. 130	Carbonyl fluoride	COF_2
No. 76	Phosphoryl fluoride	POF_3	No. 131	Carbonyl chloride	COCl_2
No. 77	Phosphoryl chloride	POCl_3	No. 132	Carbonyl bromide	COBr_2
No. 78	Phosphoryl bromide	POBr_3	No. 133	Thiocarbonyl fluoride	SCF_2
No. 79	Thiophosphoryl trifluoride	PSF_3	No. 134	Thiocarbonyl chloride	CSCl_2
No. 80	Thiophosphoryl trichloride	PSCl_3	No. 135	Isocyanic acid	HNCO
No. 81	Perchloryl fluoride	$^{35}\text{ClO}_3\text{F}$	No. 136	Isocyanic acid-d	DNCO
No. 82	Vanadyl(V) chloride	VOCl_3	No. 137	Carbonyl chlorofluoride	COCIF
No. 83	Permanganyl fluoride	MnO_3F	No. 138	Carbonyl bromochloride	COBrCl
No. 84	Germyl fluoride	GeH_3F	No. 139	Thiocarbonyl bromochloride	CSBrCl
No. 85	Germyl fluoride-d ₃	GeD_3F			
No. 86	Germyl chloride	GeH_3Cl	No. 139	C₁-five-atomic molecules	
No. 87	Germyl chloride-d ₃	GeD_3Cl	No. 140	Trifluoromethane-d	CDF_3
No. 88	Germyl bromide	GeH_3Br	No. 141	Chlorotrifluoromethane	CClF_3
No. 89	Germyl bromide-d ₃	GeD_3Br	No. 142	Bromotrifluoromethane	CBrF_3
No. 90	Germyl iodide	GeH_3I	No. 143	Trifluoroiodomethane	CF_3I
No. 91	Germyl iodide-d ₃	GeD_3I	No. 144	Trichlorofluoromethane	CCl_3F
No. 92	Trichlorogermane	GeHCl_3	No. 145	Trichloroiodomethane	CCl_3I
No. 93	Tribromogermane	GeHBr_3	No. 146	Tribromofluoromethane	CBr_3F
No. 94	Rhenium (VII) oxide chloride	ReO_3Cl	No. 147	Dichlorodifluoromethane	CCl_2F_2
No. 95	Rhenium (VII) oxide bromide	ReO_3Br	No. 148	Dibromodifluoromethane	CBr_2F_2
No. 96	Sulfuryl fluoride	SO_2F_2	No. 149	Bromodichlorofluoromethane	CBrCl_2F
No. 97	Sulfuryl chloride	SO_2Cl_2		Dibromochlorofluoromethane	CBr_2ClF
No. 98	Dichlorosilane	SiH_2Cl_2	No. 150		
No. 99	Dichlorosilane-d ₂	SiD_2Cl_2	No. 151	C₁-six-atomic molecules	
No. 100	Dibromosilane	SiH_2Br_2	No. 152	Borine Carbonyl	$^{10}\text{BH}_3\text{CO}$
No. 101	Selenium dioxide difluoride	SeO_2F_2	No. 153	Borine Carbonyl-d ₃	$^{10}\text{BD}_3\text{CO}$
No. 102	Nitric acid	H^{14}NO_3	No. 154	Borine Carbonyl	$^{11}\text{BH}_3\text{CO}$
				Borine Carbonyl-d ₃	$^{11}\text{BD}_3\text{CO}$
				Methylmercuric iodide	CH_3HgI

C₁-eight-atomic molecules		C₂-ten-atomic molecules	
No. 155	Methylsilane CH ₃ SiH ₃	No. 185	Azomethane CH ₃ NNCH ₃
No. 156	Methylsilane-d ₃ CH ₃ SiD ₃	No. 186	Azomethane-d ₆ CD ₃ NNCD ₃
No. 157	Methylgermane CH ₃ GeH ₃	C₃-five-atomic molecules	
No. 158	Methyl-d ₃ -germane CD ₃ GeH ₃	No. 187	Carbon suboxide C ₃ O ₂
No. 159	Methyltriiodogermane CH ₃ GeI ₃	No. 188	Carbon subsulfide C ₃ S ₂
No. 160	Methylstannane CH ₃ SnH ₃	No. 189	Chloro cyanoacetylene ClCCCN
No. 161	Methylstannane-d ₃ CH ₃ SnD ₃	No. 190	Bromo cyanoacetylene BrCCCN
No. 162	Methyl-d ₃ -stannane CD ₃ SnH ₃	No. 191	Iodo cyanoacetylene ICCCN
C₂-four-atomic molecules		C₃-seven-atomic molecules	
No. 163	Cyanogen C ₂ N ₂	No. 192	Chloropropadiene CH ₂ CCHCl
No. 164	Dichloroacetylene C ₂ Cl ₂	No. 193	Bromopropadiene CH ₂ CCHBr
No. 165	Dibromoacetylene C ₂ Br ₂	No. 194	Iodopropadiene CH ₂ CCHI
No. 166	Diiodoacetylene C ₂ I ₂	No. 195	Trifluoropropyne CF ₃ CCH
No. 167	Bromochloroacetylene C ₂ BrCl	No. 196	Trifluoropropyne-d CF ₃ CCD
No. 168	Chloroiodoacetylene C ₂ ClI	No. 197	1-Chloro-3,3,3-trifluoropropyne CF ₃ CCCl
No. 169	Bromoiodoacetylene C ₂ BrI	No. 198	1-Bromo-3,3,3-trifluoropropyne CF ₃ CCBr
C₂-six-atomic molecules		No. 199	1-Iodo-3,3,3-trifluoropropyne CF ₃ CCI
No. 170	Trifluoroacetonitrile CF ₃ CN	No. 200	Propargyl fluoride CHCCH ₂ F
No. 171	trans-1,2-Difluoroethylene CHFCHF	No. 201	Propargyl chloride CHCCH ₂ Cl
No. 172	trans-1,2-Difluoroethylene-d ₁ CHFCDF	No. 202	Propargyl bromide CHCCH ₂ Br
No. 173	trans-1,2-Difluoroethylene-d ₂ CDFCDF	No. 203	Propargyl iodide CHCCH ₂ I
No. 174	Glyoxal C ₂ H ₂ O ₂	C₄-six-atomic molecules	
No. 175	Glyoxal-d ₁ C ₂ HDO ₂	No. 204	Chlorodiacetylene ClCCCH
No. 176	Glyoxal-d ₂ C ₂ D ₂ O ₂	No. 205	Bromodiacetylene BrCCCH
C₂-seven-atomic molecules		No. 206	Iododiacetylene ICCCH
No. 177	Acetyl fluoride CH ₃ COF	C₄-fourteen-atomic molecules	
No. 178	Acetyl fluoride-d ₃ CD ₃ COF	No. 207	1,4-Dioxane C ₄ H ₈ O ₂
C₂-nine-atomic molecules		C₅-eight-atomic molecules	
No. 179	Dimethylzinc CH ₃ ZnCH ₃	No. 208	Diethynyl ketone (HC≡C) ₂ CO
No. 180	Dimethylzinc-d ₆ CD ₃ ZnCD ₃	C₆-eight-atomic molecules	
No. 181	Dimethylcadmium CH ₃ CdCH ₃	No. 209	Dicyanodiacetylene NCCCCCN
No. 182	Dimethylcadmium-d ₆ CD ₃ CdCD ₃	C₆-thirteen-atomic molecules	
No. 183	Dimethylmercury CH ₃ HgCH ₃	No. 210	Hexacarbonylchromium(0) Cr(¹² C ¹⁶ O) ₆
No. 184	Dimethylmercury-d ₆ CD ₃ HgCD ₃	No. 211	Hexacarbonylmolybdenum(0) Mo(¹² C ¹⁶ O) ₆
		No. 212	Hexacarbonyltungsten(0) W(¹² C ¹⁶ O) ₆

No. 1		Krypton difluoride		KrF_2		Symmetry number $\sigma = 2$	
Symmetry D_{2h}							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)		
σ_g^+	ν_1	Sym. stretch.	449 B	ia	449		
π_u	ν_2	Bend.	233 B	232.6 S	ia		
σ_u^+	ν_3	Antisym. stretch.	590 A	589.89 VS	ia		

References

- [1] IR.R. H. H. Classen, G. L. Goodman, J. G. Malm, and F. Schreiner, *J. Chem. Phys.* **42**, 1229 (1965).
 [2] IR. C. Murchison, S. Reichman, D. Anderson, J. Overend, and F. Schreiner, *J. Amer. Chem. Soc.* **90**, 5690 (1968).

No. 2		Nitrogen dioxide		$^{14}\text{NO}_2$		Symmetry number $\sigma = 2$	
Symmetry C_{2v}							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}		
a_1	ν_1	Sym. stretch.	1318 B	1318 W			
	ν_2	Bend.	750 A	749.650 S			
b_1	ν_3	Antisym. stretch.	1618 B	1617.8 VS			

References

- [1] IR. E. T. Arakawa and A. H. Nielsen, *J. Mol. Spectry.* **2**, 413 (1958).
 [2] IR. J. W. Keller and A. H. Nielsen, *J. Chem. Phys.* **29**, 252 (1958).
 [3] IR. S. C. Hurlock, K. N. Rao, L. A. Weller, and P. K. L. Yin, *J. Mol. Spectry.* **48**, 372 (1973).

No. 3		Nitrogen dioxide		$^{15}\text{NO}_2$		Symmetry number $\sigma = 2$	
Symmetry C_{2v}							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}		
a_1	ν_1	Sym. stretch.	1306 B	1306 W			
	ν_2	Bend.	740 B	740.2 S			
b_1	ν_3	Antisym. stretch.	1580 B	1580.3 VS			

Reference

- [1] IR. E. T. Arakawa and A. H. Nielsen, *J. Mol. Spectry.* **2**, 413 (1958).

No. 4 Ozone $^{16}\text{O}_3$
Symmetry C_{2v}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a_1	ν_1	Sym. stretch.	1103 A	1103.157 VW		
	ν_2	Bend.	701 B	701.42 W		
b_1	ν_3	Antisym. stretch.	1042 A	1042.096 S		

References

- [1] IR. S. A. Clough and F. X. Kneizys, *J. Chem. Phys.* **44**, 1855 (1966).
 [2] IR. D. J. McCaa and J. H. Shaw, *J. Mol. Spectry.* **25**, 374 (1968).
 [3] IR. J. L. Brumant, A. Barve, and P. Jouve, *Comptes Rendus* **B268**, 459 (1969).

No. 5 Ozone $^{18}\text{O}_3$
Symmetry C_{2v}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Xe Matrix)	cm ⁻¹	
a_1	ν_1	Sym. stretch.	1038 C	1038 W		
	ν_2	Bend.	660 C	660 W		
b_1	ν_3	Antisym. stretch.	974 C	974 S		

References

- [1] IR. L. Brewer and J. Ling-Fai Wang, *J. Chem. Phys.* **56**, 759 (1972).
 [2] IR.Th. M. Spoliti, S. N. Cesaro, and B. Mariti, *J. Chem. Phys.* **59**, 985 (1973).

No. 6 Water H_2^{18}O
Symmetry C_{2v}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a_1	ν_1	Sym. stretch.	3650 A	3649.68		
	ν_2	Bend	1588 A	1588.275		
b_1	ν_3	Antisym. stretch.	3742 A	3741.58		

References

- [1] IR. W. S. Benedict, *Mem. Soc. Roy. Sci. Liege*, **18**, 557 (1957).
 [2] IR. P. E. Fraley, K. N. Rao, and L. H. Jones, *J. Mol. Spectry.* **29**, 312 (1969).
 [3] IR. J. G. Williamson, K. N. Rao, and L. H. Jones, *J. Mol. Spectry.* **40**, 372 (1971).

No. 7 Sulfur dioxide S¹⁸O₂
Symmetry C_{2v} **Symmetry number $\sigma = 2$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
a ₁	ν_1	Sym. stretch.	1101 B	1100.65		
	ν_2	Bend.	497 B	496.7		
b ₁	ν_3	Antisym. stretch.	1318 B	1317.9		

Reference

- [1] IR. A. Barbe and P. Jauve, *J. Mol. Spectry.* **38**, 273 (1971).

No. 8 Sulfur dichloride SCl₂
Symmetry C_{2v} **Symmetry number $\sigma = 2$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
a ₁	ν_1	Sym. stretch.	525 D	525 VS	514 (10), p	
	ν_2	Bend.	208 B		208 (5), p	
b ₁	ν_3	Antisym. stretch.	535 D		535 (1)	

References

- [1] R. H. Stammerich, R. Forneris, and K. Sone, *J. Chem. Phys.* **23**, 972 (1955).
 [2] IR. G. M. Barrow, *Can. J. Chem.* **59**, 987 (1955).
 [3] IR. Y. Morino, Y. Murata, T. Ito, and J. Nakamura, *J. Phys. Soc. Japan* **17**, Supplement B-II, 37 (1962)

No. 9 Germanium (II) chloride GeCl₂
Symmetry C_{2v} **Symmetry number $\sigma = 2$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
a ₁	ν_1	Sym. stretch.	399 C	399	399 p	
	ν_2	Bend.	159 C		159 p	
b ₁	ν_3	Antisym. stretch.	374 D	374	362 dp	
				(Ar Matrix)	(Ar Matrix)	

References

- [1] R. I. R. Beattie and R. O. Perry, *J. Chem. Soc.* **A1970**, 2429.
 [2] IR. L. Andrews and D. L. Frederick, *J. Amer. Chem. Soc.* **92**, 775 (1970).
 [3] R. C. A. Ozin and A. Vander Voet, *J. Chem. Phys.* **56**, 4768 (1972).

No. 10		Tin (II) fluoride	SnF ₂	Symmetry number $\sigma = 2$		
Symmetry C _{2v}						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Ar Matrix)		
a ₁	ν_1	Sym. stretch.	593 D	592.7		
	ν_2	Bend.	197 D	197		
b ₁	ν_3	Antisym. stretch.	571 D	570.9		

Reference

[1] IR. R. H. Hauge, J. W. Hastie, and J. L. Margrave, J. Mol. Spectry. **45**, 420 (1973).

No. 11		Tin (II) chloride	SnCl ₂	Symmetry number $\sigma = 2$		
Symmetry C _{2v}						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Ar Matrix)	(Gas)	
a ₁	ν_1	Sym. stretch.	352 C	354	352 p	
	ν_2	Bend.	120 C		120 p	
b ₁	ν_3	Antisym. stretch.	334 D	334	332	
					(Ar Matrix)	

References

See No. 9(GeCl₂).

No. 12		Tin (II) bromide	SnBr ₂	Symmetry number $\sigma = 2$		
Symmetry C _{2v}						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
					(Gas)	
a ₁	ν_1	Sym. stretch.	244 D		244	
	ν_2	Bend.	80 C		(Ar Matrix)	
b	ν_3	Antisym. stretch.	231 D		80	
					231	
					(Ar Matrix)	

References

See No. 9(GeCl₂).

No. 13 Lead (II) fluoride PbF_2
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Ar Matrix)	cm^{-1}	
a_1	ν_1	Sym. stretch.	531 D	531.2		
	ν_2	Bend.	165 D	165		
b_1	ν_3	Antisym. stretch.	507 D	507.2		

ReferenceSee No. 10(SnF_2).

No. 14 Lead (II) chloride PbCl_2
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Ar Matrix)	cm^{-1} (Gas)	
a_1	ν_1	Sym. stretch.	314 C	322	314 p	
	ν_2	Bend.	99 C		99 p	
b_1	ν_3	Antisym. stretch.	299 D	299	300 dp (Ar Matrix)	

ReferencesSee No. 9(GeCl_2).

No. 15 Nitrosyl fluoride $^{16}\text{O}^{14}\text{NF}$
Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a'	ν_1	NO stretch.	1844 B	1843.5 VS		
	ν_2	NF stretch.	766 B	765.8 VS		
	ν_3	Bend.	520 C	519.9 VS		

References

- [1] IR. P. J. H. Woltz, E. A. Jones, and A. H. Nielsen, *J. Chem. Phys.* **20**, 378 (1952).
 [2] IR. D. W. Magnuson, *J. Chem. Phys.* **20**, 380 (1952).
 [3] IR. L. H. Jones, L. B. Asprey, and R. R. Ryan, *J. Chem. Phys.* **47**, 3371 (1967).

No. 16 Nitrosyl fluoride			$^{16}\text{O}^{15}\text{NF}$		Symmetry number $\sigma = 1$	
Symmetry C_s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a'	ν_1	NO stretch.	1812 C	1811.8 VS		
	ν_2	NF stretch.	749 C	748.7 VS		
	ν_3	Bend.	517 C	517.4 VS		

Reference

[1] IR. L. H. Jones, L. B. Asprey, and R. R. Ryan, J. Chem. Phys. **47**, 3371 (1967).

No. 17 Nitrosyl fluoride			$^{18}\text{O}^{14}\text{NF}$		Symmetry number $\sigma = 1$	
Symmetry C_s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a'	ν_1	NO stretch.	1796 C	1795.6 VS		
	ν_2	NF stretch.	758 C	757.8 VS		
	ν_3	Bend.	511 C	511.3 VS		

Reference

See No. 16($^{16}\text{O}^{15}\text{NF}$).

No. 18 Nitrosyl fluoride			$^{18}\text{O}^{15}\text{NF}$		Symmetry number $\sigma = 1$	
Symmetry C_s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a'	ν_1	NO stretch.	1763 C	1763.0 VS		
	ν_2	NF stretch.	740 C	740.4 VS		
	ν_3	Bend.	509 C	509.3 VS		

Reference

See No. 16($^{16}\text{O}^{15}\text{NF}$).

No. 19		Nitrosyl chloride		$^{16}\text{O}^{14}\text{NCl}$		Symmetry number $\sigma = 1$	
Symmetry C_s							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹	cm ⁻¹		
				(Gas)			
a'	ν_1	NO stretch.	1800 C	1799.7 VS			
	ν_2	NCl stretch.	596 C	595.8 VS			
	ν_3	Bend.	332 C	331.9 VS			

References

- [1] IR. W. G. Burns, and H. J. Bernstein, J. Chem. Phys. **18**, 1669 (1950).
 [2] IR. A. G. Pulford and A. Walsh, Trans. Faraday Soc. **47**, 347 (1951).
 [3] IR. W. H. Eberhardt and T. G. Burke, J. Chem. Phys. **20**, 529 (1952).
 [4] IR. L. Landau and W. H. Fletcher, J. Mol. Spectry. **4**, 276 (1960).
 [5] IR. L. H. Jones, R. R. Ryan, and L. B. Asprey, J. Chem. Phys. **49**, 581 (1968).

No. 20		Nitrosyl chloride		$^{16}\text{O}^{15}\text{NCl}$		Symmetry number $\sigma = 1$	
Symmetry C_s							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹	cm ⁻¹		
				(Gas)			
a'	ν_1	NO stretch.	1769 C	1768.9 VS			
	ν_2	NCl stretch.	582 C	581.7 VS			
	ν_3	Bend.	330 C	329.9 VS			

References

- [1] IR. L. Landau and W. H. Fletcher, J. Mol. Spectry. **4**, 276 (1960).
 [2] IR. L. H. Jones, R. R. Ryan, and L. B. Asprey, J. Chem. Phys. **49**, 581 (1968).

No. 21		Nitrosyl chloride		$^{18}\text{O}^{14}\text{NCl}$		Symmetry number $\sigma = 1$	
Symmetry C_s							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹	cm ⁻¹		
				(Gas)			
a'	ν_1	NO stretch.	1751 C	1752.7 VS 1749.0 VS		FR($3\nu_2$)	
	ν_2	NCl stretch.	588 C	588.4 VS			
	ν_3	Bend.	325 C	324.9 VS			

Reference

- [1] IR. L. H. Jones, R. R. Ryan, and L. B. Asprey, J. Chem. Phys. **49**, 581 (1968).

No. 22		Nitrosyl chloride		$^{18}\text{O}^{15}\text{NCl}$		Symmetry number $\sigma = 1$	
Symmetry C_s							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}		
a'	ν_1	NO stretch.	1721 C	1720.6 VS			
	ν_2	NCl stretch.	574 C	573.9 VS			
	ν_3	Bend.	323 C	323.2 VS			

ReferenceSee No. 21($^{18}\text{O}^{14}\text{NCl}$)

No. 23		Nitrosyl bromide		$^{16}\text{O}^{14}\text{NBr}$		Symmetry number $\sigma = 1$	
Symmetry C_s							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}		
a'	ν_1	NO stretch.	1799 C	1799.0 VS			
	ν_2	NBr stretch.	542 C	542.0 VS			
	ν_3	Bend.	266 C	266.4 S			

References

- [1] IR. W. G. Burns and H. J. Bernstein, J. Chem. Phys. **18**, 1669 (1950).
 [2] IR. J. Laane, L. H. Jones, R. R. Ryan, and L. B. Asprey, J. Mol. Spectry. **30**, 485 (1969).

No. 24		Nitrosyl bromide		$^{16}\text{O}^{15}\text{NBr}$		Symmetry number $\sigma = 1$	
Symmetry C_s							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}		
a'	ν_1	NO stretch.	1768 C	1768.3 VS			
	ν_2	NBr stretch.	528 D	527.8 VS			
	ν_3	Bend.	264 C	264 S			

Reference

- [1] IR. J. Laane, L. H. Jones, R. R. Ryan, and L. B. Asprey, J. Mol. Spectry. **30**, 485 (1969).

No. 25		Nitrosyl bromide		$^{18}\text{O}^{15}\text{NBr}$	Symmetry number $\sigma = 1$	
		Symmetry C_s				
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a'	ν_1	NO stretch.	1720 C	1719.5 VS		
	ν_2	NBr stretch.	522 C	521.5 VS		
	ν_3	Bend.	257 C	257.0 S		

ReferenceSee No. 24($^{16}\text{O}^{15}\text{NBr}$).

No. 26		Hypofluorous acid		HOF	Symmetry number $\sigma = 1$	
		Symmetry C_s				
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (N_2 Matrix)	cm^{-1}	
a'	ν_1	OH stretch.	3537 C	3537.1		
	ν_2	OF stretch.	1393 C	1393.0		
	ν_3	Bend.	886 C	886.0		

References

- [1] IR. P. N. Noble and G. C. Pimentel, *Spectrochim. Acta* **24A**, 797 (1968).
 [2] IR. J. A. Golb, H. H. Claassen, M. H. Studier, and E. H. Appklman, *Spectrochim. Acta* **28A**, 65 (1972).

No. 27		Hypochlorous acid		HOCl	Symmetry number $\sigma = 1$	
		Symmetry C_s				
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a'	ν_1	OH stretch.	3609 A	3609 M		
	ν_2	Bend.	1242 C	1242 S		
	ν_3	OCl stretch.	725 C	725 M		

References

- [1] IR. K. Hedberg and R. M. Badger, *J. Chem. Phys.* **19**, 508 (1951).
 [2] IR. R. A. Ashby, *J. Mol. Spectry.* **23**, 439 (1967).
 [3] IR. I. Schwager and A. Arkell, *J. Amer. Chem. Soc.* **89**, 6006 (1967).
 [4] IR. R. A. Ashby, *J. Mol. Spectry.* **40**, 639 (1971).

No. 28 Hypochlorous acid-d DOCl
Symmetry C_s

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a'</i>	ν_1	OD stretch.	2666 B	2666 M		
	ν_2	Bend.	911 C	911 S		
	ν_3	OCl stretch.	739 C	739 M		

References

- [1] IR. K. Hedberg and R. M. Badger, *J. Chem. Phys.* **19**, 508 (1951).
 [2] IR. R. A. Ashby, *J. Mol. Spectry.* **23**, 439 (1967).

No. 29 Thionitrosyl-S-fluoride NSF
Symmetry C_s

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a'</i>	ν_1	NS stretch.	1372 C	1372 VS		
	ν_2	SF stretch.	640 C	640 VS		
	ν_3	Bend.	366 C	366 S		

References

- [1] IR. H. Richert and O. Glemser, *Z. Anorg. Alleg. Chem.* **307**, 328 (1961).
 [2] IR. I. Schwager and A. Arkell, *J. Amer. Chem. Soc.* **89**, 6006 (1967).

No. 30 Thionitrosyl-S-chloride NSCl
Symmetry C_s

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a'</i>	ν_1	NS stretch.	1325 C	1325		
	ν_2	SCl stretch.	414 C	414		
	ν_3	Bend.	273 D			OC($\nu_2 + \nu_3$, $2\nu_3$).

Reference

- [1] IR. A. Müller, G. Nagarajan, O. Glemser, S. F. Cyvin, and J. Wegener, *Spectrochim. Acta* **23A**, 2683 (1967).

No. 31 Boron trifluoride $^{10}\text{BF}_3$
Symmetry D_{3h} Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1'	ν_1	Sym. stretch.	888 B	ia	888 VS	
a_2''	ν_2	Op-deform.	719 A	719.3 S	ia	
e'	ν_3	Deg. stretch.	1503 A	1502.5 S	1481 VW	
	ν_4	Deg. deform.	482 B	482.0 M	482 W, b	

References

- [1] R. D. M. Yost, D. DeVault, T. F. Anderson, E. N. Lassette, J. Chem. Phys. **6**, 424 (1938).
 [2] IR. L. P. Lindeman and M. K. Wilson, J. Chem. Phys. **24**, 242 (1956).
 [3] IR. M. N. Dreska, K. N. Rao, and L. H. Jones, J. Mol. Spectry. **18**, 404 (1965).
 [4] IR. S. W. Ginn, J. K. Kenney, and J. Overend, J. Chem. Phys. **48**, 1571 (1968).
 [5] IR. S. W. Ginn, C. W. Brown, J. K. Kenney, and J. Overend, J. Mol. Spectry. **28**, 509 (1968).

No. 32 Boron trifluoride $^{11}\text{BF}_3$
Symmetry D_{3h} Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1'	ν_1	Sym. stretch.	888 B	ia	888 VS	
a_2''	ν_2	Op-deform.	691 A	691.2 S	ia	
e'	ν_3	Deg. stretch.	1449 B	1449 S	1448 VW	
	ν_4	Deg. deform.	480 B	480.4 M	482 b	

References

See No. 31($^{10}\text{BF}_3$).

No. 33 Sulfur trioxide SO₃
Symmetry D_{3h} **Symmetry number $\sigma = 6$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Gas)	
a ₁ '	ν_1	Sym. stretch.	1065 B	ia	1065 VS, p	
a ₂ ''	ν_2	Op deform.	498 B	497.55 M	ia	
e'	ν_3	Deg. stretch.	1391 A	1391.3 VS	1390 W	
	ν_4	Deg. deform.	530 B	530.2 S	528 W	

References

- [1] R. H. Gerding and W. J. Nijverd, *Rec. Trav. Chim.* **59**, 1206 (1940).
 [2] IR. R. W. Lovejoy, J. H. Colwell, D. F. Eggers, and G. D. Halsey, *J. Chem. Phys.* **36**, 612 (1962).
 [3] IR. R. Bent and W. R. Lander, *Spectrochim. Acta* **19**, 931 (1963).
 [4] R. K. Stopperka, *Z. Anorg. Allegem. Chem.* **345**, 277 (1966).
 [5] IR. J. B. Miline and A. Ruoff, *J. Mol. Spectry.* **23**, 408 (1967).
 [6] IR. R. K. Thomas and H. Thompson, *Proc. Roy. Soc. London* **A314**, 329 (1970).
 [7] IR.R.Th. A. Kaldar, A. G. Maki, A. J. Dorney, and L. M. Mills, *J. Mol. Spectry.* **45**, 247 (1973).

No. 34 Ammonia-t₃ NT₃
Symmetry C_{3v} **Symmetry number $\sigma = 3$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)		
a ₁	ν_1	Sym. stretch.	2014 A	2014.1		
	ν_2	Sym. deform.	657 B	656.4 s ^a 657.2 a ^a		
e	ν_3	Deg. stretch.	2185 A	2184.8		
	ν_4	Deg. deform.	996 A	996.3		

^a"s" and "a" refer to symmetric and antisymmetric levels [1].

References

- [1] IR. K. N. Rao, W. W. Brim, J. M. Hoffman, L. H. Jones, and R. S. McDowell, *J. Mol. Spectry.* **7**, 362 (1961).
 [2] IR. L. H. Jones, W. W. Brim, and K. N. Rao, *J. Mol. Spectry.* **11**, 389 (1963).

No. 35 Nitrogen trifluoride $^{15}\text{NF}_3$
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a_1	ν_1	Sym. stretch.	1009 A	1008.93		
	ν_2	Sym. deform.	645 B	644.84		
e	ν_3	Deg. stretch.	886 A	886.34		
	ν_4	Deg. deform.	492 B	492.02		

References

- [1] MW. M. Otake, C. Matsumura, and Y. Morino, J. Mol. Spectry. **28**, 316 (1968).
 [2] IR.MW.Th. M. Otake, E. Hirota, and Y. Morino, J. Mol. Spectry. **28**, 325 (1968).
 [3] IR.Th. A. Allan, J. L. Duncan, J. H. Holloway, and D. C. Mckean, J. Mol. Spectry. **31**, 368 (1969).

No. 36 Aluminum trichloride AlCl_3
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Ar Matrix)	cm ⁻¹ (Gas)	
a_1	ν_1	Sym. stretch.	375 C	380.2	375 p	
	ν_2	Sym. deform.	183 C	182.8		
e	ν_3	Deg. stretch.	595 C	594.7		
	ν_4	Deg. deform.	150 C	149.2	150	

References

- [1] IR. W. Klemperer, J. Chem. Phys. **24**, 353 (1956).
 [2] R. I. R. Beatie and J. R. Horder, J. Chem. Soc. **B1969**, 2655.
 [3] IR. M. L. Lesiecki and J. S. Shirk, J. Chem. Phys. **56**, 4171 (1972).

No. 37 Phosphorus triiodide PI₃
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(CS ₂ , C ₆ H ₆ , CH ₂ Cl ₂ soln.)	C ₆ H ₆ , CCl ₄ soln.)	
a ₁	ν_1	Sym. stretch.	303 D	306 M	303 (3) p	
	ν_2	Sym. deform.	111 D	112 VW	111 (7) p	
e	ν_3	Deg. stretch.	325 D	328 VS	325 (1b) dp	
	ν_4	Deg. deform.	79 D	80 VW, b	79 (10) dp	

References

- [1] R. R. H. Stammreich, R. Forneris, and Y. Tavares, *J. Chem. Phys.* **25**, 580 (1956).
 [2] IR. S. G. Frankiss, F. A. Miller, H. Stammreich and Th. T. Sans, *Spectrochim. Acta* **23A**, 543 (1967).

No. 38 Arsenic trifluoride AsF₃
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a ₁	ν_1	Sym. stretch.	741 A	740.5 S	707 (10)	
	ν_2	Sym. deform.	337 B	336.5 M	341 (2)	
e	ν_3	Deg. stretch.	702 B	702.2 S	644 (9)	
	ν_4	Deg. deform.	262 B	262.3 M	274 (4)	

References

- [1] R. D. M. Yost and J. E. Sherborne, *J. Chem. Phys.* **2**, 125 (1934).
 [2] IR. L. C. Hoskins and R. C. Lord, *J. Chem. Phys.* **43**, 155 (1965).
 [3] IR.Th. I. W. Levin and S. Abramowitz, *J. Chem. Phys.* **44**, 2562 (1966).
 [4] IR.Th. L. C. Hoskins, *J. Chem. Phys.* **45**, 4594 (1966).
 [5] Th. A. M. Mirri, *J. Chem. Phys.* **47**, 2823 (1967).
 [6] IR. S. Reichman and J. Overend, *Spectrochim. Acta* **26A**, 379 (1970).
 [7] Th. S. Reichman, D. F. Smith, Jr. and J. Overend, *Spectrochim. Acta* **26A**, 927 (1970).

No. 39		Arsenic triiodide		AsI ₃		Symmetry number $\sigma = 3$	
Symmetry C _{3v}							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹ (Solid)	cm ⁻¹ (CHBr ₃ , CS ₂ soln.)		
a ₁	ν_1	Sym. stretch.	219 C	225.7	218.5 p		
	ν_2	Sym. deform.	94 C	101.6	94 p		
e	ν_3	Deg. stretch.	224 C	201.2 VVS,b	223.5 dp		
	ν_4	Deg. deform.	71 C	73.6	71 dp		

References

- [1] R. R. Stammreich, R. Forneris, and Y. Tavares, *J. Chem. Phys.* **25**, 580 (1956).
 [2] IR. T. R. Marley and D. A. Williams, *Spectrochim. Acta* **21**, 1773 (1968).
 [3] R. M. A. Hooper and D. N. James, *Austral. J. Chem.* **21**, 2379 (1968).
 [4] R. W. Kiefer, *Z. Naturforsch.* **A25**, 1101 (1970).

No. 40		trans-1,2-Difluorodiazine		N ₂ F ₂		Symmetry number $\sigma = 2$	
Symmetry C _{2h}							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Gas)		
a _g	ν_1	NN stretch.	1523 C	ia	1523 (2)		
	ν_2	NF stretch.	1018 C	ia	1018 (5)		
	ν_3	FNN deform.	603 C	ia	603 (10)		
a _u	ν_4	Torsion	364 B	363.5 M	ia		
b _u	ν_5	NF stretch.	991 A	991.01 VS	ia		
	ν_6	FNN deform.	423 C	423 M	ia		

References

- [1] IR. R. H. Sanborn, *J. Chem. Phys.* **33**, 1855 (1960).
 [2] IR.R. S-T. King and J. Overend, *Spectrochim. Acta* **22**, 689 (1966).
 [3] IR. S-T. King and J. Overend, *Spectrochim. Acta* **23A**, 2875 (1967).
 [4] IR.R. J. Shamir and H. H. Hyman, *Spectrochim. Acta*, **23A**, 1191 (1967).

No. 41 Nitryl fluoride F¹⁴NO₂
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	NO ₂ s-stretch.	1310 B	1309.6 VS		
	ν_2	NF stretch.	822 B	822.4 VS		
	ν_3	NO ₂ scis.	568 C	567.8 S		
b ₁	ν_4	NO ₂ a-stretch.	1792 B	1791.5 VS		
	ν_5	NO ₂ rock.	560 C	559.6 S		
b ₂	ν_6	Op-deform.	742 B	742.0 M		

References

- [1] IR. R. E. Dodd, J. A. Roufe, and L. A. Woodward, *Trans. Farad. Soc.* **52**, 145 (1956).
 [2] IR. D. L. Bernitt, R. H. Miller, and I. C. Hisatsune, *Spectrochim. Acta* **22A**, 237 (1967).
 [3] Th. T. Tanaka and Y. Morino, *J. Mol. Spectry.* **32**, 430 (1969).

No. 42 Nitryl fluoride F¹⁵NO₂
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	NO ₂ s-stretch.	1298 B	1298.0 VS		
	ν_2	NF stretch.	809 B	808.5 VS		
	ν_3	NO ₂ scis.	567 C	567.1 S		
b ₁	ν_4	NO ₂ a-stretch.	1751 B	1750.5 VS		
	ν_5	NO ₂ rock.	559 C	558.9 S		
b ₂	ν_6	Op-deform.	722 B	722.0 M		

References

See No. 41(F¹⁴NO₂).

No. 43 Nitryl chloride $\text{Cl}^{14}\text{NO}_2$
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a_1	ν_1	NO_2 s-stretch.	1286 C	1318.5 S 1267.1 VS		FR($2\nu_6$).
	ν_2	NCl stretch.	793 B	792.6 VS		
	ν_3	NO_2 scis.	370 C	369.6 VS		
b_1	ν_4	NO_2 a-stretch.	1685 B	1684.6 VS		
	ν_5	NO_2 rock.	408 C	408.1 VW		
b_2	ν_6	Op-deform.	652 C	652.3 M		

References

- [1] IR. R. Ryason and M. K. Wilson, J. Chem. Phys. **22**, 2000 (1954).
 [2] IR. D. L. Bernitt, R. H. Miller, and I. C. Hisatsune, Spectrochim. Acta **23A**, 237 (1967).

No. 44 Nitryl chloride $\text{Cl}^{15}\text{NO}_2$
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a_1	ν_1	NO_2 s-stretch.	1286 D	1318.5 S 1267.1 VS		FR($2\nu_6$).
	ν_2	NCl stretch.	793 B	792.6 VS		
	ν_3	NO_2 scis.	370 C	369.6 VS		
b_1	ν_4	NO_2 a-stretch.	1685 B	1684.6 VS		
	ν_5	NO_2 rock.	408 C	408.1 VW		
b_2	ν_6	Op-deform.	652 C	652.3 M		

References

See No. 43($\text{Cl}^{14}\text{NO}_2$).

No. 45 Chlorine trifluoride ClF_3
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	ClF stretch.	752 C	751 S	752.1 S, p	
	ν_2	ClF_2 s-stretch.	529 C	530 M	529.3 VS, p	
	ν_3	ClF_2 ip-deform.	328 C	328 S	337 W, p 321	OV(ν_6).
b_1	ν_4	ClF_2 a-stretch.	702 C	702 VS		
	ν_5	ClF_2 ip-deform.	442 C	442 W	431 W, dp	
b_2	ν_6	ClF_2 op-deform.	328 D	328 S		OV(ν_3).

References

- [1] IR.R. H. Selig, H. H. Claassen, and J. H. Holloway, J. Chem. Phys. **52**, 3517 (1970).
 [2] Th. K. Ramaswamy and P. Muthusbramanian, J. Mol. Structure **9**, 193 (1971).

No. 46 Bromine trifluoride BrF_3
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	BrF stretch.	675 C	675 S	675 S, p	
	ν_2	BrF_2 s-stretch.	552 C	552 W	552 VS, p	
	ν_3	BrF_2 ip-deform.	242 C	242 S	233 W, p	OV(ν_6).
b_1	ν_4	BrF_2 a-stretch.	614 C	614 VS	612 VW	
	ν_5	BrF_2 ip-deform.	350 C	350 VW		
b_2	ν_6	BrF_2 op-deform.	242 D	242 S		OV(ν_3).

References

- [1] IR.R. H. Selig, H. H. Claassen, and J. H. Holloway, J. Chem. Phys. **52**, 3517 (1970).
 [2] IR. K. O. Christe, E. C. Curtis, and D. Pilipovich, Spectrochim. Acta **27A**, 931 (1971).
 [3] Th. K. Ramaswamy and P. Muthusbramanian, J. Mol. Structure **9**, 193 (1971).

No. 47 Hydrogen peroxide H_2O_2 Symmetry C_2 Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
<i>a</i>	ν_1	OH stretch.	3599 B	3599.2	3364 p	
	ν_2	OH bend.	1402 E		1402 p	
	ν_3	OO stretch.	877 C	877	880 p	
	ν_4	Torsion	371 B	370.7		
			243 C	242.8		
<i>b</i>	ν_5	OH stretch.	3608 B	3607.9		
	ν_6	OH bend.	1266 D	1266		

References

- [1] IR. P. A. Giguère, J. Chem. Phys. **18**, 88 (1950).
 [2] IR. P. A. Giguère and O. Bain, J. Phys. Chem. **56**, 340 (1952).
 [3] IR. O. Bain and P. A. Giguère, Can. J. Chem. **33**, 527 (1956).
 [4] R. R. C. Tylor and P. C. Cross, J. Chem. Phys. **24**, 41 (1956).
 [5] IR. R. L. Redington, W. B. Olson, and P. C. Cross, J. Chem. Phys. **36**, 1311 (1962).
 [6] IR. R. H. Hunt, R. A. Leacock, C. W. Peters and K. T. Hecht, J. Chem. Phys. **42**, 1931 (1965).

No. 48 Hydrogen persulfide H_2S_2 Symmetry C_2 Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
<i>a</i>	ν_1	SH stretch.	2556 A	2555.78	2509 (2), dp	
	ν_2	SH bend.	883 D		883 (2), dp	
	ν_3	SS stretch.	509 D		509 (9)	
	ν_4	Torsion	416 B	416		
<i>b</i>	ν_5	SH stretch.	2559 A	2558.64		
	ν_6	SH bend.	886 C	886		

References

- [1] IR. M. K. Wilson and R. M. Badger, J. Chem. Phys. **17**, 1232 (1949).
 [2] R. F. Fehér, W. Laue, and G. Winkhaus, Z. Anorg. Allg. Chem. **288**, 113 (1956).
 [3] IR. R. L. Redington, J. Mol. Spectry. **9**, 469 (1962).
 [4] IR. B. P. Winnwischer, J. Mol. Spectry. **36**, 414 (1970).

No. 49 **Difluoro disulphane** F_2S_2
 Symmetry C_2 Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
<i>a</i>	ν_1	SF stretch.	717 B	717.0		
	ν_2	SS stretch.	615 B	614.6	623 p	
	ν_3	SF bend.	320 B	319.8	322 p	
	ν_4	Torsion	183 C	182.5	193 p	
<i>b</i>	ν_5	SF stretch.	681 B	680.8	683	
	ν_6	SF bend.	301 C	301	297 dp	

References

- [1] IR. F. Seel and R. Budenz, Chem. Ber. **98**, 251 (1965).
 [2] IR.R. R. D. Brown and G. P. Pez, Spectrochim. Acta **26A**, 1375 (1970).

No. 50 **Hydrazoic acid** HN_3
 Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
<i>a'</i>	ν_1	NH stretch.	3336 B	3336 M		
	ν_2	$N\equiv N$ stretch.	2140 B	2139.8 VS		
	ν_3	NH bend.	1264 B	1263.7 M		
	ν_4	N-N stretch.	1151 B	1150.5 VS		
	ν_5	NNN bend.	534 B	534.2 W		
<i>a''</i>	ν_6	NNN bend.	607 B	607.0 VW		

References

- [1] IR. D. A. Dows and G. C. Pimentel, J. Chem. Phys. **23**, 1258 (1955).
 [2] IR. G. C. Pimentel, S. W. Charles, and K. J. Rosengren, J. Chem. Phys. **44**, 3029 (1966).
 [3] IR. G. B. Moore and K. J. Rosengren, J. Chem. Phys. **44**, 4108 (1966).
 [4] IR. D. M. Levine and D. A. Dows, J. Chem. Phys. **46**, 1168 (1967).

No. 51 Hydrazoic acid DN₃
Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a'</i>	ν_1	ND stretch.	2478 B	2478 M		
	ν_2	N≡N stretch.	2112 C	2112 VS		FR($\nu_3 + \nu_4$).
	ν_3	ND bend.	954 B	953.8 S		
	ν_4	N-N stretch.	1184 B	1183.7 M		
	ν_5	NNN bend.	492 B	492.2 W		
<i>a''</i>	ν_6	NNN bend.	588 B	588.4 VW		

ReferencesSee No. 50(NH₃).

No. 52 Difluoroamine NF₂H
Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a'</i>	ν_1	NH stretch.	3193 C	3193 W		
	ν_2	NH bend.	1307 B	1307 S		
	ν_3	NF ₂ s-stretch.	972 B	972 S		
	ν_4	NF ₂ scis.	500 C	500 M		
<i>a''</i>	ν_5	NH bend.	1424 B	1424 S		
	ν_6	NF ₂ a-stretch.	888 C	888 VS		

Reference

[1] IR. J. J. Comeford, D. E. Mann, J. L. Schoen, and D. R. Lide, Jr.,
 J. Chem. Phys. **38**, 461 (1963).

No. 53 Chlorodifluoroammonia NClF₂
Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a'</i>	ν_1	NF ₂ s-stretch.	930 C	930.2		
	ν_2	NCl stretch.	697 C	696.9		
	ν_3	NF ₂ deform.	556 C	555.5		
	ν_4	NClF s-deform.	377 C	377		
<i>a''</i>	ν_5	NF ₂ a-stretch.	855 C	855.4		
	ν_6	NClF a-deform.	382 D	382 ^a		

^a An average of two splitting bands.**References**

[1] IR. R. Ettinger, J. Chem. Phys. **38**, 2427 (1963).
 [2] IR. J. J. Comeford, J. Chem. Phys. **45**, 3463 (1966).

No. 54 Dichlorofluoroammonia NCl_2F Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)		
a'	ν_1	NF stretch.	825 C	825 S		
	ν_2	NCl_2 s-stretch.	615 C	615 M		
	ν_3	NFCl s-deform.	409 C	409 W		
	ν_4	NCl_2 deform.	274 C	274 W		
a''	ν_5	NCl_2 a-stretch.	692 C	692 S		
	ν_6	NFCl a-deform.	344 C	344 M		

References

- [1] IR. D. E. Milligan, NBS Report 8149.
 [2] IR. R. P. Hirschmann, L. R. Anderson, D. F. Harnish, and W. B. Fox, Spectrochim. Acta **24A**, 1267 (1968).

No. 55 Thionyl fluoride SOF_2 Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Gas)	
a'	ν_1	SO stretch.	1333 D	1340.8 S	1339.3 VS	
	ν_2	SF_2 s-stretch.	808 B	1330.9 S	1329.9 VS	FR($\nu_2 + \nu_3$).
	ν_3	OSF deform.	530 B	808.2 VS	808.3 VS	
	ν_4	SF_2 scis.	378 D	530.4 M	529.6 S	
				377.8 W	379.5 W	
					(liquid)	
a''	ν_5	SF_2 a-stretch.	747 C	747.0 VS	746.8 W	
	ν_6	OSF deform.	393 D	392.5 W	398.6 M	
					(liquid)	

References

- [1] IR. J. K. O'Loane and M. K. Wilson, J. Chem. Phys. **23**, 1313 (1955).
 [2] R. P. Bender and J. M. Wood, Jr., J. Chem. Phys. **23**, 1316 (1955).
 [3] IR. F. Seel and R. Budenz, Chem. Ber. **98**, 251 (1965).
 [4] IR.R. E. L. Pace and H. V. Samuelson, J. Chem. Phys. **44**, 3682 (1966).

No. 56 Thionyl chloride SOCl₂
 Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a'	ν_1	SO stretch.	1251 C	1251 VS	1230 W, p	
	ν_2	SCl ₂ s-stretch.	492 C	492 VS	490 M, p	
	ν_3	OSCl deform.	344 D		344 S, p	
	ν_4	SCl ₂ scis.	194 D		194 M, p	
a''	ν_5	SCl ₂ a-stretch.	455 C	455 VS	445 W, dp	
	ν_6	OSCl deform.	284 D		284 M, dp	

References

- [1] R. R. Vogel-Hogler, *Acta. Phys. Austriaca* **1**, 323 (1948).
 [2] R. C. A. McDowell, *Trans. Faraday Soc.* **49**, 371 (1953).
 [3] IR. D. E. Martz and R. T. Lagemann, *J. Chem. Phys.* **22**, 1193 (1954).
 [4] R. G. Allen and C. A. McDowell, *J. Chem. Phys.* **23**, 209 (1955).
 [5] Th. A. Mueller and G. Nagarajan, *Z. Phys. Chem. (Leipzig)* **235**, 57 (1967).
 [6] Th. K. Ramaswamy and S. Jayaraman, *J. Mol. Structure* **7**, 470 (1971).

No. 57 Thionyl bromide SOBr₂
 Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹ (Liquid)	
a'	ν_1	SO stretch.	1121 D	1121 (2) p		
	ν_2	SBr s-stretch.	405 D		405 (7) p	
	ν_3	OSBr deform.	267 D		267 (10) p	
	ν_4	SBr ₂ scis.	120 D		120 (10) p	
a''	ν_5	SBr ₂ a-stretch.	379 D		379 (5) dp	
	ν_6	OSBr deform.	223 D		223 (7) dp	

Reference

- [1] R. H. Stammereich, R. Forneris, and Y. Tavares, *J. Chem. Phys.* **25**, 1277 (1956).

No. 58 Titanium tetrachloride TiCl_4
Symmetry T_d

Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	Sym. stretch.	389 B	ia, 388 ^a	389	
e	ν_2	Deg. deform.	114 B	ia, 119 ^a	114	
f_2	ν_3	Deg. stretch.	498 C	498.5	498	
	ν_4	Deg. deform.	136 C	136	136	

^a Estimated from combination bands.**References**

- [1] IR. N. J. Hawkins and D. R. Carpenter, J. Chem. Phys. **23**, 1700 (1955).
 [2] IR. H. Bürger and A. Ruoff, Spectrochim. Acta **24A**, 1863 (1968).
 [3] R. R. J. H. Clark, B. K. Hunter, and D. M. Rippon, Inorg. Chem. **11**, 56 (1972).

No. 59 Titanium tetrabromide TiBr_4
Symmetry T_d

Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Liquid)	cm^{-1} (Gas)	
a_1	ν_1	Sym. stretch.	232 B	ia	231.5	
e	ν_2	Deg. deform.	69 B	ia	68.5	
f_2	ν_3	Deg. stretch.	393 C	383 VS	393	
	ν_4	Deg. deform.	88 C		88	

References

- [1] IR. R. F. A. Miller and G. L. Carlson, Spectrochim. Acta **16**, 6 (1960).
 [2] R. R. J. H. Clark, B. K. Hunter, and D. M. Rippon, Inorg. Chem. **11**, 56 (1972).

No. 60 Titanium tetraiodide TiI_4
Symmetry T_d

Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (C_6H_{12} soln.)	cm^{-1} (C_6H_{12} soln.)	
a_1	ν_1	Sym. stretch.	162 D	ia	162 VS, p	
e	ν_2	Deg. deform.	51 D	ia	51 M, dp	
f_2	ν_3	Deg. stretch.	322 D	322 VS	324 M, dp	
	ν_4	Deg. deform.	67 D	67 M		

Reference

- [1] IR. R. R. J. H. Clark and C. J. Willis, J. Chem. Soc. A **1971**, 838.

No. 61 Zirconium tetrachloride $ZrCl_4$ Symmetry T_d Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	Sym. stretch.	377 B	ia, 388 ^a	377 VS, p	
e	ν_2	Deg. deform.	98 B	ia, 102 ^a	98	
f_2	ν_3	Deg. stretch.	418 C	421	418 VW	
	ν_4	Deg. deform.	113 C	112 ^a	113	

^a Estimated from combination bands.**References**

- [1] IR. J. K. Wilmshurst, *J. Mol. Spectry.* **5**, 343 (1960).
 [2] IR. A. Büchler, J. B. Berkowitz-Mattuck, and D. H. Dugre, *J. Chem. Phys.* **34**, 2202 (1961).
 [3] R. R. J. H. Clark, B. K. Hunter, and D. M. Rippon, *Inorg. Chem.* **11**, 56 (1972).

No. 62 Zirconium tetrabromide $ZrBr_4$ Symmetry T_d Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1} (Gas)	
a_1	ν_1	Sym. stretch.	223 B	ia	222.5	
e	ν_2	Deg. deform.	60 B	ia	60	
f_2	ν_3	Deg. stretch.	315 C		315	
	ν_4	Deg. deform.	72 C		72	

Reference

- [1] R. R. J. H. Clark, B. K. Hunter, and D. M. Rippon, *Inorg. Chem.* **11**, 56 (1972).

No. 63 Zirconium tetraiodide ZrI_4 Symmetry T_d Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1} (Gas)	
a_1	ν_1	Sym. stretch.	158 B	ia	158	
e	ν_2	Deg. deform.	43 B	ia	43	
f_2	ν_3	Deg. stretch.	254 C		254	
	ν_4	Deg. deform.	55 C		55	

ReferenceSee No. 62($ZrBr_4$).

No. 64 Ruthenium tetroxide RuO₄
 Symmetry T_d
Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a ₁	ν_1	Sym. stretch.	885 C	ia	885.3 S	
e	ν_2	Deg. deform.	322 C	ia	322.4 M	
f ₂	ν_3	Deg. stretch.	921 C	921.0	912.9 W	
	ν_4	Deg. deform.	336 C	336.0	333 W	

References

- [1] IR. R. E. Dodd, *Trans. Faraday Soc.* **55**, 1480 (1959).
 [2] IR. M. H. Ortner, *J. Chem. Phys.* **34**, 559 (1961).
 [3] R. G. Davidson, N. Logan, and A. Morris, *Chem. Commun.* **1968**, 1044.
 [4] R. W. P. Griffith, *J. Chem. Soc.* **A1968**, 1663.
 [5] IR. A. Müller and B. Krebs, *J. Mol. Spectry.* **26**, 136 (1968).
 [6] IR.R. I. W. Levin and S. Abramowitz, *J. Chem. Phys.* **50**, 4860 (1969).
 [7] Th. R. S. McDowell, *J. Chem. Phys.* **53**, 4407 (1970).
 [8] IR.R. R. S. McDowell, L. B. Asperry and L. C. Hoskins, *J. Chem. Phys.* **56**, 5712 (1972).

No. 65 Hafnium tetrachloride HfCl₄
 Symmetry T_d
Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Gas)	
a ₁	ν_1	Sym. stretch.	382 B	ia	382	
e	ν_2	Deg. deform.	102 B	ia	101.5	
f ₂	ν_3	Deg. stretch.	390 C	393	390	
	ν_4	Deg. deform.	112 C		112	

References

- [1] IR. A. Büchler, J. B. Berkowitz-Mattuck, and D. H. Dugre, *J. Chem. Phys.* **34**, 2202 (1961).
 [2] R. R. J. H. Clark, B. K. Hunter, and D. M. Rippon, *Inorg. Chem.* **11**, 56 (1972).

No. 66 Hafnium tetrabromide HfBr₄
 Symmetry T_d
Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
					(Gas)	
a ₁	ν_1	Sym. stretch.	236 B	ia	235.5	
e	ν_2	Deg. deform.	63 B	ia	63	
f ₂	ν_3	Deg. stretch.	273 C		273	
	ν_4	Deg. deform.	71 C		71	

ReferenceSee No. 62(ZrBr₄).

No. 67 Hafnium tetraiodide HfI_4
 Symmetry T_d Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
					(Gas)	
a_1	ν_1	Sym. stretch.	158 B	ia	158	
e	ν_2	Deg. deform.	55 C	ia	55	
f_2	ν_3	Deg. stretch.	224 C		224	
	ν_4	Deg. deform.	63 C		63	

ReferenceSee No. 62(ZrBr_4).

No. 68 Osmium tetroxide Os^{16}O_4
 Symmetry T_d Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
					(Gas)	
a_1	ν_1	Sym. stretch.	965 B	ia	965.2	
e	ν_2	Deg. deform.	333 B	ia	333.1	
f_2	ν_3	Deg. stretch.	960 B	960.5	960.1	
	ν_4	Deg. deform.	329 B	329.0	322.7	

References

- [1] IR. N. J. Hawkins and W. W. Sabal, *J. Chem. Phys.* **25**, 775 (1956).
 [2] IR. R. E. Dodd, *Trans. Faraday Soc.* **55**, 1480 (1959).
 [3] IR. I. W. Levin and S. Abramowitz, *Inorg. Chem.* **5**, 2024 (1966).
 [4] IR. R. S. McDowell, *Inorg. Chem.* **6**, 1759 (1967).
 [5] R. W. P. Griffith, *J. Chem. Soc. A* **1968**, 1663.
 [6] R. C. Davidson, N. Logan, and A. Morris, *Chem. Commun.* **1968**, 1044.
 [7] R. I. W. Levin, *Inorg. Chem.* **8**, 1018 (1969).
 [8] R. J. L. Huston and H. H. Claassen, *J. Chem. Phys.* **52**, 5646 (1970).
 [9] IR. C. G. Barraclough and M. M. Sinclair, *Spectrochim. Acta* **26A**, 207 (1970).
 [10] IR.R. R. S. McDowell and M. Goldblatt, *Inorg. Chem.* **10**, 625 (1971).

No. 69 Osmium tetroxide Os^{18}O_4
 Symmetry T_d Symmetry number $\sigma = 12$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
					(CCl_4 soln.)	
a_1	ν_1	Sym. stretch.	910 C	ia	909.7	
e	ν_2	Deg. deform.	317 C	ia	316.6	
f_2	ν_3	Deg. stretch.	912 B	911.8		
	ν_4	Deg. deform.	313 B	312.7		

References

- [1] IR. C. G. Barraclough and M. M. Sinclair, *Spectrochim. Acta* **26A**, 207 (1970).
 [2] IR.R. R. S. McDowell and M. Goldblatt, *Inorg. Chem.* **10**, 625 (1971).

No. 70 Thionitrosyl trifluoride NSF₃
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a ₁	ν_1	NS stretch.	1515 B	1515	1512 p	
	ν_2	NF ₃ s-stretch.	775 B	775	768 p	
	ν_3	NF ₃ s-deform.	521 B	521	520 dp	
e	ν_4	NF ₃ d-stretch.	811 B	811	812 dp	
	ν_5	NF ₃ d-deform.	429 B	429	430 dp	
	ν_6	NF ₃ rock.	342 B	342	340 dp	

References

- [1] IR. H. Richert and O. Glemser, Z. Anorg. Alleg. Chem. **307**, 328 (1961).
 [2] R. A. Müller, A. Ruoff, B. Krebs, O. Glemser, and W. Koch, Spectrochim. Acta **25A**, 199 (1969).

No. 71 Trifluorosilane SiHF₃
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	SiH stretch.	2316 B	2315.6 M		
	ν_2	SiF ₃ s-stretch.	858 B	858.25 S		
	ν_3	SiF ₃ s-deform.	425 B	425.2 M		
e	ν_4	SiF ₃ d-stretch.	998 B	997.83 VS		
	ν_5	SiH bend.	844 B	843.6 S		
	ν_6	SiF ₃ d-deform.	306 B	306.2 M		

References

- [1] IR. C. Newman, S. R. Polo, and M. K. Wilson, Spectrochim. Acta **15**, 793 (1959).
 [2] IR. H. Bürger und S. Biedermann, Spectrochim. Acta **27A**, 1687 (1971)

No. 72 Trifluorosilane-d SiDF₃
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	SiD stretch.	1691 B	1690.72		
	ν_2	SiF ₃ s-stretch.	854 B	853.84 S		
	ν_3	SiF ₃ s-deform.	422 B	422.1 M		
e	ν_4	SiF ₃ d-stretch.	992 B	991.76 VS		
	ν_5	SiD bend.	628 B	628.41 M		
	ν_6	SiF ₃ d-deform.	303 B	302.8 M		

References

See No. 71(SiHF₃).

No. 73 Trichlorosilane SiHCl_3
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_1	ν_1	SiH stretch.	2261 B	2260.9 S	2258 p	
	ν_2	SiCl_3 s-stretch.	499 B	498.6 S	489 p	
	ν_3	SiCl_3 s-deform.	254 B	253.7 M	250 p	
e	ν_4	SiH bend.	811 B	810.8 VS	799 dp	
	ν_5	SiCl_3 d-stretch.	600 B	600.1 VS	587 dp	
	ν_6	SiCl_3 d-deform.	176 B	175.5 M	179 dp	

References

- [1] R. M. L. Delwaille and M. F. Francois, *Comptes Rendus* **228**, 1007 (1949).
 [2] IR.Th. H. Bürger and A. Ruoff, *Spectrochim. Acta* **26**, 1449 (1970).

No. 74 Trichlorosilane-d SiDCl_3
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a_1	ν_1	SiD stretch.	1647 B	1647.1 S		
	ν_2	SiCl_3 s-stretch.	496 B	495.9 S		
	ν_3	SiCl_3 s-deform.	252 B	252.0 M		
e	ν_4	SiD bend.	634 B	633.8 VS		
	ν_5	SiCl_3 d-stretch.	550 B	550.0 VS		
	ν_6	SiCl_3 d-deform.	175 B	174.5 M		

Reference

- [1] IR.Th. H. Bürger and A. Ruoff, *Spectrochim. Acta* **26**, 1449 (1970).

No. 75 Tribromosilane SiHBr_3
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_1	ν_1	SiH stretch.	2239 B	2239.2 S	2236 p	
	ν_2	SiBr_3 s-stretch.	362 C	362 M, b	362 p	
	ν_3	SiBr_3 s-deform.	168 B	168.5 M	166 p	
e	ν_4	SiH bend.	774 C	774 VS	770 dp	
	ν_5	SiBr_3 d-stretch.	484 C	483.5 VS	470 dp	
	ν_6	SiBr_3 d-deform.	117 C	116.8 W	115 dp	

References

- [1] R. M. L. Delwaille and M. F. Francois, *Comptes Rendus* **230**, 743 (1950).
 [2] IR.Th. H. Bürger und J. Cichon, *Spectrochim. Acta* **27A**, 2191 (1971).

No. 76 Phosphoryl fluoride POF₃
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Gas)	
a ₁	ν_1	PO stretch.	1417 B	1415 M	1416.8 p	
	ν_2	PF ₃ s-stretch.	873 B	873 M	872.7 p	
	ν_3	PF ₃ s-deform.	472 B	473 S	472.4	OV(5) in Raman.
e	ν_4	PF ₃ d-stretch.	991 C	990 VS	991.0 dp	
	ν_5	PF ₃ d-deform.	482 C	485 M	482.0 dp	
	ν_6	PO bend.	336 C		335.5 dp	

References

- [1] R. M. L. Delwaulle and M. F. François, *Comptes Rendus* **222**, 550 (1946).
 [2] IR. H. S. Gutowsky and A. D. Liehr, *J. Chem. Phys.* **20**, 1652 (1952).
 [3] R.Th. R. J. H. Clark and D. M. Rippon, *Mol. Phys.* **28**, 305 (1974).

No. 77 Phosphoryl chloride POCl₃
Symmetry C₂ Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹ (Gas)	
a ₁	ν_1	PO stretch.	1322 C		1321.5	
	ν_2	PCl ₃ s-stretch.	481 C		480.5	
	ν_3	PCl ₃ s-deform.	266 C		265.5	
e	ν_4	PCl ₃ d-stretch.	590 C		589.5	
	ν_5	PO bend.	333 C		332.5	
	ν_6	PCl ₃ d-deform.	187 C		187.0	

References

- [1] R. M. L. Delwaulle and M. F. François, *J. Chim. Physique* **45**, 50 (1948).
 [2] R.Th. R. J. H. Clark and D. M. Rippon, *Mol. Phys.* **28**, 305 (1974).

No. 78 Phosphoryl bromide POBr₃
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹ (Liquid)	
a ₁	ν_1	PO stretch.	1261 D		1261	
	ν_2	PBr ₃ s-stretch.	340 D		340	
	ν_3	PBr ₃ s-deform.	173 D		173	
e	ν_4	PBr ₃ d-stretch.	488 D		488	
	ν_5	PO bend.	267 D		267	
	ν_6	PBr ₃ d-deform.	118 D		118	

Reference

See No. 77(POCl₃).

No. 79 Thiophosphoryl trifluoride PSF_3 Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
a_1	ν_1	PF_3 s-stretch.	981 B	981 VS	969 (4.5) p	
	ν_2	PS stretch.	695 B	695 M	696 (100) p	
	ν_3	PF_3 s-deform.	440 B	440 M	441 (20) p	
e	ν_4	PF_3 d-stretch.	945 B	945 VS	937 (6.0) dp	
	ν_5	PF_3 d-deform.	402 C	402 M	404 (13) dp	
	ν_6	PF_3 rock.	275 B	275 W	276 (37) dp	

References

- [1] R. M. L. Delwaulle and F. François, *Comptes Rendus* **226**, 894 (1948).
 [2] R. M. L. Delwaulle and F. François, *J. Chim. Phys.* **46**, 87 (1949).
 [3] IR. R. G. Cavell, *Spectrochim. Acta* **23A**, 249 (1968).
 [4] IR.R. J. R. Durig and J. W. Clark, *J. Chem. Phys.* **46**, 3057 (1967).

No. 80 Thiophosphoryl trichloride PSCl_3 Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Gas)	
a_1	ν_1	PS stretch.	767 B	770	767.2 p	
	ν_2	PCl_3 s-stretch.	432 B	431	431.6 p	
	ν_3	PCl_3 s-deform.	246 C	252	246.0 dp	OV(ν_5).
e	ν_4	PCl_3 d-stretch.	544 C	547	543.7 dp	
	ν_5	PCl_3 d-deform.	246 C	252	246 dp	OV(ν_3).
	ν_6	PCl_3 rock.	167 C	174	167.3 dp	
				(liquid)		

References

- [1] IR. G. Cilento, D. A. Ramsay, and R. N. Jones *J. Amer. Chem. Soc.* **71**, 2753 (1949).
 [2] IR.R. J. R. Durig and J. W. Clark, *J. Chem. Phys.* **46**, 3057 (1967).
 [3] R.Th. R. J. H. Clark and D. M. Rippon, *Mol. Phys.* **28**, 305 (1974).

No. 81 Perchlorylfluoride $^{35}\text{ClO}_3\text{F}$
 Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	ClO_3 s-stretch.	1063 B	1061 S	1062.5 VS	
	ν_2	ClF stretch.	716 B	714 S	716.0 S	
	ν_3	ClO_3 s-deform.	550 B	549 W	549.5 M	
e	ν_4	ClO_3 d-stretch.	1318 B	1315 VS	1317.5 W	
	ν_5	ClO_3 d-deform.	590 B	590.55 M	589.5 W	
	ν_6	ClO_3 rock.	405 B	405 W	404.5 W	

References

- [1] IR. R. P. Madden and W. S. Benedict, J. Chem. Phys. **25**, 594 (1956).
 [2] IR. D. R. Lide, Jr. and D. E. Mann, J. Chem. Phys. **25**, 1128 (1956).
 [3] R. F. X. Powell and E. R. Lippincott, J. Chem. Phys. **32**, 1883 (1960).
 [4] R. H. H. Claassen and E. H. Appleman, Inorg. Chem. **9**, 622 (1970).
 [5] R.Th. R. J. H. Clark and D. M. Rippon, Mol. Phys. **28**, 305 (1974).

No. 82 Vanadyl(V) chloride VOCl_3
 Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Liquid)	cm^{-1} (Gas)	
a_1	ν_1	VO stretch.	1042 C	1035 VS	1042.5 (1.8) p	
	ν_2	VCl_3 s-stretch.	408 C	408 S	409.5 (10.0) p	
	ν_3	VCl_3 s-deform.	163 C		163.0 (0.8)	
e	ν_4	VCl_3 d-stretch.	502 C	502 VS	503 (20) dp	
	ν_5	VCl_3 d-deform.	246 C		246.0 (1.5) dp	
	ν_6	VCl_3 rock.	125 C		124.5 (3.8)	

References

- [1] R. H. J. Eichoff and F. Weigel, Z. Allg. Chem. **275**, 267 (1954).
 [2] IR.R. F. A. Miller and L. R. Cousins, J. Chem. Phys. **26**, 329 (1957).
 [3] R. J. R. Beattie, K. M. S. Livingston, D. J. Reynolds, and G. A. Ozin, J. Chem. Soc. **1970**, 1210 (1970).
 [4] R.Th. R. J. H. Clark and D. M. Rippon, Mol. Phys. **28**, 305 (1974).

No. 83 Permanganyl fluoride MnO_3F
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a_1	ν_1	MnO_3 s-stretch.	905 B	905.2 M		
	ν_2	MnF stretch.	721 B	720.7 M		
	ν_3	MnO_3 s-deform.	338 B	337.7 W		
e	ν_4	MnO_3 d-stretch.	953 B	952.5 S		
	ν_5	MnO_3 d-deform.	374 B	373.9 W		
	ν_6	MnF bend.	264 B	264.3 W		

Reference

[1] IR. M. J. Reisfeld, L. B. Asprey, and N. A. Matuiyoff, *Spectrochim. Acta* **27A**, 765 (1971).

No. 84 Germyl fluoride GeH_3F
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a_1	ν_1	GeH_3 s-stretch.	2121 C	2120.6 S		
	ν_2	GeH_3 s-deform.	859 C	859.0 VS		
	ν_3	GeF stretch.	689 B	689.1 S		
e	ν_4	GeH_3 d-stretch.	2132 B	2131.7 S		
	ν_5	GeH_3 d-deform.	874 B	874.0 S		
	ν_6	GeH_3 rock.	643 B	642.6 M		

References

[1] IR. J. E. Griffiths, T. N. Srivastava, and M. Onyszchuk, *Can. J. Chem.* **40**, 579 (1962).

[2] IR. D. E. Freeman, K. H. Rhee, and M. K. Wilson, *J. Chem. Phys.* **39**, 2908 (1963).

[3] IR. K. H. Rhee and M. K. Wilson, *J. Chem. Phys.* **43**, 331 (1965).

No. 85 Germyl fluoride- d_3 GeD_3F
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a_1	ν_1	GeD_3 s-stretch.	1525 C	1525		
	ν_2	GeD_3 s-deform.	696 B	696		
	ν_3	GeF stretch.	618 C	618		
e	ν_4	GeD_3 d-stretch.	1535 B	1535		
	ν_5	GeD_3 d-deform.	635 C	635		
	ν_6	GeD_3 rock.	466 C	466		

Reference

[1] IR. D. E. Freeman, K. H. Rhee, and M. K. Wilson, *J. Chem. Phys.* **39**, 2908 (1963).

No. 86 **Germyl chloride** **GeH₃Cl**
 Symmetry C_{3v}

 Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	GeH ₃ s-stretch.	2120 C	2119.9		
	ν_2	GeH ₃ s-deform.	848 B	847.5		
	ν_3	GeCl stretch.	422 C	421.7		
e	ν_4	GeH ₃ d-stretch.	2129 B	2128.9		
	ν_5	GeH ₃ d-deform.	874 B	874.1		
	ν_6	GeH ₃ rock.	602 B	602.2		

References

- [1] IR. R. C. Lord, Jr., and C. M. Steese, J. Chem. Phys. **22**, 542 (1954).
 [2] IR. D. E. Freeman, K. H. Rhee, and M. K. Wilson, J. Chem. Phys. **39**, 2908 (1963).
 [3] IR. K. H. Rhee and M. K. Wilson, J. Chem. Phys. **43**, 331 (1965).

No. 87 **Germyl chloride-d₃** **GeD₃Cl**
 Symmetry C_{3v}

 Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
				(Gas)		
a ₁	ν_1	GeD ₃ s-stretch.	1518 C	1518		
	ν_2	GeD ₃ s-deform.	609 C	609		
	ν_3	GeCl stretch.	422 C	422		
e	ν_4	GeD ₃ d-stretch.	1530 C	1530		
	ν_5	GeD ₃ d-deform.	630 C	630		
	ν_6	GeD ₃ rock.	434 C	434		

Reference

- [1] IR. D. E. Freeman, K. H. Rhee, and M. K. Wilson, J. Chem. Phys. **39**, 2908 (1963).

No. 88 **Germyl bromide** **GeH₃Br**
 Symmetry C_{3v}

 Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	GeH ₃ s-stretch.	2115 C	2115.2		
	ν_2	GeH ₃ s-deform.	833 C	832.7		
	ν_3	GeBr stretch.	308 C	307.7		
e	ν_4	GeH ₃ d-stretch.	2127 B	2126.7		
	ν_5	GeH ₃ d-deform.	871 B	870.9		
	ν_6	GeH ₃ rock.	578 B	578.2		

References

See No. 84(GeH₃F).

No. 89 **Germyl bromide-d₃ GeD₃Br**
Symmetry C_{3v} **Symmetry number $\sigma = 3$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a</i> ₁	ν_1	GeD ₃ s-stretch.	1514 C	1514		
	ν_2	GeD ₃ s-deform.	596 C	596		
	ν_3	GeBr stretch.	305 C	305		
<i>e</i>	ν_4	GeD ₃ d-stretch.	1530 C	1530		
	ν_5	GeD ₃ d-deform.	621 C	621		
	ν_6	GeD ₃ rock.	420 C	420		

Reference

[1] IR. D. E. Freeman, K. H. Rhee, and M. K. Wilson, J. Chem. Phys. **39**, 2908 (1963).

No. 90 **Germyl iodide GeH₃I**
Symmetry C_{3v} **Symmetry number $\sigma = 3$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a</i> ₁	ν_1	GeH ₃ s-stretch.	2110 C	2110.1		
	ν_2	GeH ₃ s-deform.	808 C	808.2		
	ν_3	GeI stretch.	248 C	248.2		
<i>e</i>	ν_4	GeH ₃ d-stretch.	2121 B	2120.8		
	ν_5	GeH ₃ d-deform.	854 B	853.6		
	ν_6	GeH ₃ rock.	558 B	558.3		

References

[1] IR. D. E. Freeman, K. H. Rhee, and M. K. Wilson, J. Chem. Phys. **39**, 2908 (1963).

[2] IR. K. H. Rhee and M. K. Wilson, J. Chem. Phys. **43**, 331 (1965).

No. 91 **Germyl iodide-d₃ GeD₃I**
Symmetry C_{3v} **Symmetry number $\sigma = 3$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a</i> ₁	ν_1	GeD ₃ s-stretch.	1508 C	1508		
	ν_2	GeD ₃ s-deform.	582 C	582		
	ν_3	GeI stretch.	249 C	249		
<i>e</i>	ν_4	GeD ₃ d-stretch.	1525 C	1525		
	ν_5	GeD ₃ d-deform.	603 C	603		
	ν_6	GeD ₃ rock.	404 C	404		

Reference

[1] IR. D. E. Freeman, K. H. Rhee, and M. K. Wilson, J. Chem. Phys. **39**, 2908 (1963).

No. 92 Trichlorogermane GeHCl₃Symmetry C_{3v}Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
a ₁	ν_1	GeH stretch.	2159 D		(Liquid) 2159 p	
	ν_2	GeCl ₃ s-stretch.	409 D		409 p	
	ν_3	GeCl ₃ s-deform.	181 D		181 p	
e	ν_4	GeH bend.	699 D		699 dp	
	ν_5	GeCl ₃ d-stretch.	438 D		438 dp	
	ν_6	GeCl ₃ d-deform.	149 D		149 dp	

Reference[1] R. M. L. Delwaille and M. F. François, *Comptes Rendus* **228**, 1007 (1949).**No. 93 Tribromogermane GeHBr₃**Symmetry C_{3v}Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
a ₁	ν_1	GeH stretch.	2116 D		(Liquid) 2116 p	
	ν_2	GeBr ₃ s-stretch.	273 D		273 p	
	ν_3	GeBr ₃ s-deform.	128 D		128 p	
e	ν_4	Ge H bend.	674 D		674 dp	
	ν_5	GeBr ₃ d-stretch.	325 D		325 dp	
	ν_6	GeBr ₃ d-deform.	95 E		95 dp	

Reference[1] R. M. L. Delwaille and M. F. François, *Comptes Rendus* **230**, 743 (1950).**No. 94 Rhenium (VII) oxide chloride ReO₃Cl**Symmetry C_{3v}Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
a ₁	ν_1	ReO ₃ s-stretch.	1001 C	(Liquid) 1001.5 VS	(Liquid) 1001 S, p	
	ν_2	ReO ₃ s-deform.	434 C	434 VS	435 M, p	
	ν_3	ReCl stretch.	293 C	293 S		
e	ν_4	ReO ₃ d-stretch.	960 C	960 VS	962.5 M, dp	
	ν_5	ReO ₃ d-deform.	344 C	343.5 S	344 M, dp	
	ν_6	ReCl bend.	196 C	196 M	196 S, dp	

References

- [1] R. H. J. Eichhoff and F. Weigel, *Z. Anorg. Allg. Chem.* **275**, 267 (1954).
 [2] IR.R. F. A. Miller and G. L. Carlson, *Spectrochim. Acta* **16**, 1148 (1960).
 [3] Th. A. Müller, B. Krebs, and W. Höltje, *Spectrochim. Acta* **23A**, 2753 (1967).

No. 95 Rhenium (VII) oxide bromide ReO_3Br Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (CCl_4 soln.)	cm^{-1}	
a_1	ν_1	ReO_3 s-stretch.	997 C	997 S		
	ν_2	ReO_3 s-deform.	350 C	350 M		
	ν_3	ReBr stretch.	195 C	195 M		
e	ν_4	ReO_3 d-stretch.	963 C	963 VS		
	ν_5	ReO_3 d-deform.	332 C	332 S		
	ν_6	ReBr bend.	168 C	168 M		

References

- [1] IR.R. F. A. Miller and G. L. Carlson, *Spectrochim. Acta* **16**, 1148 (1960).
 [2] Th. A. Müller, B. Krebs, and W. Höltje, *Spectrochim. Acta* **23A**, 2753 (1967).

No. 96 Sulfuryl fluoride SO_2F_2 Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	SO_2 s-stretch.	1269 C	1269 S	1270 M	
	ν_2	SF_2 s-stretch.	848 C	848 S	847 S	
	ν_3	SO_2 scis.	544 D	544.3 M	543 M	
	ν_4	SF_2 scis.	385 D	384.5 VW		
a_2	ν_5	SF_2 twist.	388 C	ia	388 W	
b_1	ν_6	SO_2 a-stretch.	1502 C	1502 S	1502 VW	
	ν_7	SO_2 rock.	553 D	552.8 M		
	ν_8	SF_2 a-stretch.	885 C	885 S	883 VW	
	ν_9	SO_2 wag.	539 D	539.4 M		

References

- [1] IR. W. D. Perkins and M. K. Wilson, *J. Chem. Phys.* **20**, 1791 (1952).
 [2] R. P. Bender and J. M. Wood, Jr., *J. Chem. Phys.* **23**, 1316 (1955).
 [3] Th. G. R. Hunt and M. K. Wilson, *Spectrochim. Acta* **18**, 959 (1962).
 [4] IR. D. R. Lide, Jr., D. E. Mann, and J. J. Comeford, *Spectrochim. Acta* **21**, 497 (1965).
 [5] Th. H. Toyuki and K. Shimizu, *Bull. Chem. Soc. Japan*, **39**, 2364 (1966).
 [6] IR. A. J. Sumodi and E. L. Pace, *Spectrochim. Acta* **28A**, 1129 (1972).

No. 97		Sulfuryl chloride		SO ₂ Cl ₂		Symmetry number $\sigma = 2$	
Symmetry C _{2v}							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹	cm ⁻¹		
				(Gas)	(Liquid)		
a ₁	ν_1	SO ₂ s-stretch.	1205 C	1205 S	1182 S, p		
	ν_2	SO ₂ scis.	577 D	577 VS	560 VS, p		
	ν_3	SO ₂ s-stretch.	408 D		408 VS, p		
	ν_4	SO ₂ scis.	218 D		218 VS, p		
a ₂	ν_5	SO ₂ twist.	282 D	ia	282 S, dp		
b ₁	ν_6	SO ₂ a-stretch.	1434 C	1434 S	1414 M, dp		
	ν_7	SO ₂ rock.	388 D		388 S, dp		
b ₂	ν_8	SO ₂ a-stretch.	586 D	586 VS	580 VW, dp		
	ν_9	SO ₂ wag.	362 D		362 sh, dp		

References

- [1] R. R. Vogel-Hogler, *Acta. Phys. Austriaca*, **1**, 323 (1948).
 [2] IR. D. E. Martz and R. T. Lagemann, *J. Chem. Phys.* **22**, 1193 (1954).
 [3] R. J. Gillespie and E. A. Robinson, *Can. J. Chem. Phys.* **39**, 2171 (1961).
 [4] Th. C. R. Hunt and M. K. Wilson, *Spectrochim. Acta* **18**, 959 (1962).
 [5] Th. H. Toyuki and K. Shimizu, *Bull. Chem. Soc. Japan* **39**, 2364 (1966).

No. 98		Dichlorosilane		SiH ₂ Cl ₂		Symmetry number $\sigma = 2$	
Symmetry C _{2v}							
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹	cm ⁻¹		
				(Gas)	(Liquid)		
a ₁	ν_1	SiH ₂ s-stretch.	2224 C	2224 S	2221 S, p		
	ν_2	SiH ₂ scis.	954 C	954 S	942 W, dp		
	ν_3	SiCl ₂ s-stretch.	527 C	527 W	514 S, p		
	ν_4	SiCl ₂ scis.	188 D		188 M, dp		
a ₂	ν_5	SiH ₂ twist.	710 D	ia	710 W, dp		
b ₁	ν_6	SiH ₂ a-stretch.	2237 B	2237 S			
	ν_7	SiH ₂ rock.	602 C	602 W			
b ₂	ν_8	SiH ₂ wag.	876 B	876 VS	868 VW, dp		
	ν_9	SiCl ₂ a-stretch.	590 C	590 S	566 VW, dp		

References

- [1] IR. R. J. A. Hawkins and M. K. Wilson, *J. Chem. Phys.* **21**, 360 (1953).
 [2] IR. J. A. Hawkins, S. R. Polo, and M. K. Wilson, *J. Chem. Phys.* **21**, 1122 (1953).
 [3] IR. D. H. Christensen and O. F. Nielsen, *J. Mol. Spectry.* **27**, 489 (1968).
 [4] Th. D. H. Christensen and O. F. Nielsen, *J. Mol. Spectry.* **33**, 425 (1970).

No. 99 **Dichlorosilane-d₂** **SiD₂Cl₂**
 Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	SiD ₂ s-stretch.	1608 C	1608 S		
	ν_2	SiD ₂ scis.	695 C	695 S		
	ν_3	SiCl ₂ s-stretch.	519 C	519 W		
	ν_4	SiCl ₂ scis.	187 D	187 W		
a ₂	ν_5	SiD ₂ twist.	506 D	ia		CF.
b ₁	ν_6	SiD ₂ a-stretch.	1637 C	1637 S		
	ν_7	SiD ₂ rock.	466 B	466 W		
b ₂	ν_8	SiD ₂ wag.	663 B	663 VS		
	ν_9	SiCl ₂ a-stretch.	566 C	566 W		

References

- [1] IR. D. H. Christensen and O. F. Nielsen, J. Mol. Spectry. **27**, 489 (1968).
 [2] Th. D. H. Christensen and O. F. Nielsen, J. Mol. Spectry. **33**, 425 (1970).

No. 100 **Dibromosilane** **SiH₂Br₂**
 Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a ₁	ν_1	SiH ₂ s-stretch.	2206 D	2200 S	2206 S, p	
	ν_2	SiH ₂ scis.	942 C	942 VS	925 W, dp	
	ν_3	SiBr ₂ s-stretch.	407 C	407 M	393 S, p	
	ν_4	SiBr ₂ scis.	122 D		122 M, p	
a ₂	ν_5	SiH ₂ twist.	688 D	ia	688 W, dp	
b ₁	ν_6	SiH ₂ a-stretch.	2232 D	2200 S	2232 W, dp	
	ν_7	SiH ₂ rock.	556 C	556 S		
b ₂	ν_8	SiH ₂ wag.	843 C	843 VS	828 VW, dp	
	ν_9	SiBr ₂ a-stretch.	471 C	471 S	456 W, dp	

References

- [1] R. F. Francois and M. Buisset, Comptes Rendus **230**, 1946 (1950).
 [2] IR. D. W. Mayo, H. E. Opitz, and J. S. Peake, J. Chem. Phys. **23**, 1344 (1955).

No. 101 Selenium dioxide difluoride SeO_2F_2 Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
a_1	ν_1	SeO_2 s-stretch.	973 C	973 S	969 S, p	
	ν_2	SeF_2 s-stretch.	700 C	700 S	702 S, p	
	ν_3	SeO_2 scis.	360 C	360 S	357 M	
	ν_4	SeF_2 scis.	284 C	284 W	280 M	
a_2	ν_5	SeF_2 twist.	272 E			OV($2\nu_5$).
b_1	ν_6	SeO_2 a-stretch.	1059 C	1059 S	1059 W	
	ν_7	SeO_2 rock.	335 D	335 S	334 M	OV(ν_9).
b_2	ν_8	SeF_2 s-stretch.	756 C	756 W		
	ν_9	SeO_2 wag.	335 D	335 S	334 M	OV(ν_7).

References

- [1] R. Paetzold and K. H. Ziegenbalg, Z. Chem. **4**, 461 (1964).
 [2] IR.R. T. Birchall and R. J. Gillespie, Spectrochim. Acta **22**, 681 (1966).

No. 102 Nitric acid H^{14}NO_3 Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)		
a'	ν_1	OH stretch.	3550 B	3550.0 M		
	ν_2	NO_2 a-stretch.	1708 B	1708.2 VS		
	ν_3	OH bend.	1331 B	1330.7 S		
	ν_4	NO_2 s-stretch.	1325 B	1324.9 VS		
	ν_5	NO_2 scis.	879 B	878.6 S		
	ν_6	NO stretch.	647 B	646.6 W		
	ν_7	NO_2 rock.	579 B	579.0 W		
a''	ν_8	NO_2 wag.	762 B	762.2 S		
	ν_9	Torsion	456 B	455.8 M		

Reference

- [1] IR. G. E. McGraw, D. L. Bernitt, and I. C. Hisatsune, J. Chem. Phys. **42**, 237 (1965).

No. 103 Nitric acid-d D¹⁴NO₃
Symmetry C_s

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a'	ν_1	OD stretch.	2622 B	2621.5 M		
	ν_2	NO ₂ a-stretch.	1687 B	1687.0 VS		
	ν_3	NO ₂ s-stretch.	1308 B	1308.4 VS		
	ν_4	OD bend.	1014 B	1013.6 M		
	ν_5	NO ₂ scis.	888 B	888.0 S		
	ν_6	NO stretch.	641 B	641.0 S		
	ν_7	NO ₂ rock.	541 B	541.0 W		
a''	ν_8	NO ₂ wag.	763 B	762.8 S		
	ν_9	Torsion	342 B	342.0 M		

ReferenceSee No. 102(H¹⁴NO₃).

No. 104 Nitric acid H¹⁵NO₃
Symmetry C_s

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a'	ν_1	OH stretch.	3550 B	3550.0 M		
	ν_2	NO ₂ a-stretch.	1672 B	1672.0 VS		
	ν_3	OH bend.	1327 B	1327.0 S		
	ν_4	NO ₂ s-stretch.	1321 B	1320.6 S		
	ν_5	NO ₂ scis.	871 B	870.8 S		
	ν_6	NO stretch.	647 B	646.6 W		
	ν_7	NO ₂ rock.	578 B	578.0 W		
a''	ν_8	NO ₂ wag.	744 B	743.6 S		
	ν_9	Torsion	456 B	455.9 M		

ReferenceSee No. 102(H¹⁴NO₃).

No. 105		Nitric acid-d	D ¹⁵ NO ₃		Symmetry number $\sigma = 1$	
Symmetry C _s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
				(Gas)		
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
a'	ν_1	OD stretch.	2622 B	2621.9 M		
	ν_2	NO ₂ a-stretch.	1656 B	1655.5 VS		
	ν_3	NO ₂ s-stretch.	1291 B	1290.7 S		
	ν_4	OD bend.	1012 B	1011.5 M		
	ν_5	NO ₂ scis.	876 B	876.3 S		
	ν_6	NO stretch.	641 B	640.9 S		
	ν_7	NO ₂ rock.	541 B	541.0 VW		
a''	ν_8	NO ₂ wag.	743 B	743.4 S		
	ν_9	Torsion	345 D			OV($\nu_7 + \nu_9$, $\nu_8 + \nu_9$).

ReferenceSee No. 102(H¹⁴NO₃).

No. 106		Fluoronitrate	FO ¹⁴ NO ₂		Symmetry number $\sigma = 1$	
Symmetry C _s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)		
a'	ν_1	NO ₂ a-stretch.	1759 C	1759.1 VS		
	ν_2	NO ₂ s-stretch.	1301 B	1300.9 VS		
	ν_3	OF stretch.	928 B	927.7 S		
	ν_4	NO ₂ scis.	804 B	803.7 S		
	ν_5	NO stretch.	633 B	633.0 VW		
	ν_6	NO ₂ rock.	455 B	454.5 MS		
	ν_7	OF bend.	303 B	302.6 VW		
a''	ν_8	NO ₂ wag.	709 C	708.5 M		
	ν_9	OF torsion	152 B	151.6 VVW		

References

- [2] IR. A. J. Arvia, L. F. R. Cafferata, and H. J. Schumacher, Chem. Ber. **96**, 1187 (1963).
 [2] IR. R. H. Miller, P. L. Bernitt, and I. C. Hisatsune, Spectrochim. Acta **23A**, 223 (1967).

No. 107		Fluoronitrate	FO ¹⁵ NO ₂		Symmetry number $\sigma = 1$	
Symmetry C _s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a'</i>	ν_1	NO a-stretch.	1720 C	1719.8 VS		
	ν_2	NO ₂ s-stretch.	1291 B	1290.5 VS		
	ν_3	OF stretch.	928 B	927.5 S		
	ν_4	NO ₂ scis.	792 B	791.8 S		
	ν_5	NO stretch.	632 B	631.5 VW		
	ν_6	NO ₂ rock.	455 B	454.4 MS		
	ν_7	OF bend.	302 B	301.6 VW		
<i>a''</i>	ν_8	NO ₂ wag.	690 C	690.2 M		
	ν_9	OF torsion	151 D	151.2		OC($\nu_1 + \nu_9$, $\nu_1 - \nu_9$, $\nu_3 + \nu_9$).

References

See No. 106(FO¹⁴NO₂).

No. 108		Chloronitrate	ClO ¹⁴ NO ₂		Symmetry number $\sigma = 1$	
Symmetry C _s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
<i>a'</i>	ν_1	NO ₂ a-stretch.	1735 B	1735.4 VS		
	ν_2	NO ₂ s-stretch.	1292 B	1291.9 VS		
	ν_3	OCl stretch.	809 C	809.3 S		
	ν_4	NO ₂ scis.	780 C	780.2 MS		
	ν_5	NO stretch.	560 B	560.0 S		
	ν_6	NO ₂ rock.	434 B	434.1 M		
	ν_7	OCl bend.	270 C	270 VVW		
<i>a''</i>	ν_8	NO ₂ wag.	711 B	710.8 W		
	ν_9	OCl torsion	121 E	121		OC($\nu_5 + \nu_7 - \nu_9$).

References

- [1] IR. K. Brandle, M. Schmeisser, and W. Luttko. Chem. Ber. **93**, 2300 (1960).
 [2] IR. A. J. Arvia, L. F. R. Cafferata, and H. J. Schumacher, Chem. Ber. **96**, 1187 (1963).
 [3] IR. R. H. Miller, P. L. Bernitt, and I. C. Hisatsune, Spectrochim. Acta **23A**, 223 (1967).

No. 109		Chloronitrate	ClO ¹⁵ NO ₂		Symmetry number $\sigma = 1$	
Symmetry C _s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a'	ν_1	NO ₂ a-stretch.	1694 B	1694.0 VS		
	ν_2	NO ₂ s-stretch.	1280 B	1280.2 VS		
	ν_3	OCl stretch.	805 C	804.7 MS		
	ν_4	NO ₂ scis.	773 C	772.9 S		
	ν_5	NO stretch.	557 B	556.8 S		
	ν_6	NO ₂ rock.	432 B	432.2 M		
	ν_7	OCl bend.	263 D	263		OC($\nu_5 + \nu_7$).
a''	ν_8	NO ₂ wag.	694 B	693.7 W		
	ν_9	OCl torsion	122 E	122		OC($\nu_5 + \nu_7 - \nu_9$).

References

- [1] IR. K. Brandle, M. Schmeisser, and W. Lutke, Chem. Ber. **93**, 2300 (1960).
 [2] IR. A. J. Arvia, L. F. R. Cafferata, and H. J. Schumacher, Chem. Ber. **96**, 1187 (1963).
 [3] IR. R. H. Miller, P. L. Bernitt, and I. C. Hisatsune, Spectrochim. Acta **23A**, 223 (1967).

No. 110		Phosphoryl dichlorofluoride	POCl ₂ F		Symmetry number $\sigma = 1$	
Symmetry C _s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹ (Liquid)	
a'	ν_1	PO stretch.	1331 D		1331 p	
	ν_2	PF stretch.	894 D		894 p	
	ν_3	PCl ₂ s-stretch.	547 D		547 p	
	ν_4	PO ip-bend.	386 D		386 p	
	ν_5	PF bend.	330 D		330 p	
	ν_6	PCl ₂ scis.	207 D		207 p	
a''	ν_7	PCl ₂ a-stretch.	620 D		620 dp	
	ν_8	PO op-bend.	372 D		372 dp	
	ν_9	PCl ₂ rock.	254 D		254 dp	

Reference

- [1] R. M. L. Delwaulle and F. François, Comptes Rendus **222**, 550 (1946).

No. 111 Phosphoryl dibromofluoride POBr_2F Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
a'	ν_1	PO stretch.	1303 D		(Liquid) 1303 p	
	ν_2	PF stretch.	880 D		880 p	
	ν_3	PBr_2 s-stretch.	466 D		466 p	
	ν_4	PO ip-bend.	306 D		306 p	
	ν_5	PF bend.	273 D		273 p	
	ν_6	PBr_2 scis.	134 D		134 p	
a''	ν_7	PBr_2 a-stretch.	538 D		538 dp	
	ν_8	PO op-bend.	291 D		291 dp	
	ν_9	PBr_2 rock.	220 D		220 dp	

Reference

See No. 110(POCl_2F).No. 112 Thiophosphoryl chlorodifluoride PSClF_2 Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
a'	ν_1	PF_2 s-stretch.	946 B	(Gas) 946 VS	(Liquid) 939 (3.7) p	
	ν_2	PS stretch.	738 B	738 VS	727 (50) p	
	ν_3	PCI stretch.	541 B	541 S	536 (100) p	
	ν_4	PCI bend.	395 B	395 M	394 (31) p	
	ν_5	PF_2 scis.	361 B	361 M	359 (38) p	
	ν_6	PS ip-bend.	207 C	198	207 (65) p	
a''	ν_7	PF_2 s-stretch.	920 B	920 VS	913 (6.5) dp	
	ν_8	PF_2 rock.	317 B	317 M	314 (14) dp	
	ν_9	PS op-bend.	251 C		251 (14) dp	

Reference

[1] IR.R. J. R. Durig and J. W. Clark, J. Chem. Phys. **46**, 3057 (1967).

No. 113 Thiophosphoryl dichlorofluoride PSCl_2F Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a'	ν_1	PF stretch.	912 B	912 VS	902 (2) p	
	ν_2	PS stretch.	753 B	753 VS	737 (17) p	
	ν_3	PCl_2 s-stretch.	478 B	478 S	474 (100) p	
	ν_4	PF bend.	331 B	331 M	327 (13) dp	
	ν_5	PS ip-bend.	268 B	268 M	267 (32) p	
	ν_6	PCl_2 scis.	192 D	192 W	193 (65) dp	$\text{OV}(\nu_9)$.
a''	ν_7	PCl_2 a-stretch.	575 C	575 S	567 (6) dp	
	ν_8	PS op-bend.	319 B	319 M	317 (19) dp	
	ν_9	PCl_2 rock.	192 D	192 W	193 (65) dp	$\text{OV}(\nu_6)$.

Reference

See No. 112(PSClF_2).No. 114 Phosphorus pentafluoride PF_5 Symmetry D_{3h} Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1'	ν_1	PF_3 s-stretch.	816 B		816 (10) p	
	ν_2	PF_2 s-stretch.	648 C		648 (1b) p	
a_2''	ν_3	PF_2 a-stretch.	947 B	946.6 VS		
	ν_4	PF_3 op-deform.	575 B	575.1 M		
e'	ν_5	PF_3 d-stretch.	1024 B	1024 VS	1029 (1b)	
	ν_6	PF_3 d-deform.	533 B	532.5 M	535 (1sh) dp	
	ν_7	PF bend.	174 C		174 (1b) dp	
e''	ν_8	PF bend.	520 C		520 (1b) dp	

References

- [1] IR. J. E. Griffiths, R. P. Carter, and R. R. Holmes, *J. Chem. Phys.* **41**, 863 (1964).
- [2] IR. L. C. Hoskins, *J. Chem. Phys.* **42**, 2631 (1965).
- [3] IR. J. E. Griffiths, *J. Chem. Phys.* **42**, 2632 (1965).
- [4] IR.R. L. C. Hoskins and R. C. Lord, *J. Chem. Phys.* **46**, 2402 (1967).
- [5] IR. R. M. Deiters and R. R. Holmes, *J. Chem. Phys.* **48**, 4796 (1968).
- [6] R. I. W. Levin, *J. Chem. Phys.* **50**, 1031 (1969).
- [7] Th. I. W. Levin, *J. Mol. Spectry.* **33**, 61 (1970).
- [8] R. F. A. Miller and R. J. Capwell, *Spectrochim. Acta* **27A**, 125 (1971).

No. 115 Phosphorus trichloride difluoride PCl_3F_2
Symmetry D_{3h} Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_1'	ν_1	PF_2 s-stretch.	633 C		633 M, p	
	ν_2	PCl_3 s-stretch.	387 C		387 S, p	
a_2''	ν_3	PF_2 a-stretch.	867 C	867 VS		
	ν_4	PCl_3 op-deform.	328 C	328 M		
e'	ν_5	PCl_3 d-stretch.	625 C	625 VS	609 VW, dp	
	ν_6	PCl_3 d-deform.	404 C	404 VS	408 M, dp	
	ν_7	PF bend.	122 C		122 S, dp	
e''	ν_8	PF bend.	357 C		357 W, dp	

References

- [1] IR.R. J. E. Griffiths, R. P. Carter, Jr., and R. R. Holmes, *J. Chem. Phys.* **41**, 863 (1964).
 [2] Th. R. R. Holmes, R. M. Deiters, and J. A. Golen, *Inorg. Chem.* **8**, 2612 (1969).
 [3] Th. I. R. Beattie, K. M. S. Livingston, and D. J. Reynolds, *J. Chem. Phys.* **51**, 4269 (1969).

No. 116 Phosphorus pentachloride PCl_5
Symmetry D_{3h} Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (C_6H_6 or CS_2 soln.)	
a_1'	ν_1	PCl_3 s-stretch.	395 D		395 VS, p	
	ν_2	PCl_2 s-stretch.	370 D		370 VW	
a_2''	ν_3	PCl_2 a-stretch.	465 C	465 VS	441 VW	
	ν_4	PCl_3 op-deform.	299 D	299 S (CS_2 soln.)	301	
e'	ν_5	PCl_3 d-stretch.	592 C	592 VS	581 W, p	
	ν_6	PCl_3 d-deform.	273 D	273 S (CS_2 soln.)	281 W, dp	
	ν_7	PCl bend.	100 C	100 W	100 M, dp	
e''	ν_8	PCl bend.	261 D		261 W, dp	

References

- [1] R. H. Moureu, P. Sue, and M. Magat, *Contribution a l' étude de la structure moleculaire.* Vol. commem. Victor Henni, p.125 Maison Desoerg, Liege (1947/48).
 [2] IR.R. J. K. Wilmschurst and H. J. Bernstein, *J. Chem. Phys.* **27**, 661 (1957).
 [3] IR.R. G. L. Carlson, *Spectrochim. Acta* **19**, 1291 (1963).
 [4] R. M. J. Taylor and L. A. Woodward, *J. Chem. Soc.* **1963**, 4670 (1963).
 [5] Th. R. R. Holmes, R. M. Deiters, and J. A. Golen, *Inorg. Chem.* **8**, 2612 (1969).
 [6] Th. R. R. Holmes and R. M. Deiters, *J. Chem. Phys.* **51**, 4043 (1969).

No. 117 Vanadium pentafluoride VF_5
Symmetry D_{3h}
Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1'	ν_1	VF_3 s-stretch.	718 B		718 VS	
	ν_2	VF_2 s-stretch.	608 B		608 M	
a_2''	ν_3	VF_2 a-stretch.	784 B	784 S		
	ν_4	VF_3 op-deform.	331 B	331 W		
e'	ν_5	VF_3 d-stretch.	810 C	810 M	812 W	
	ν_6	VF_3 d-deform.	282 C	282 M	286 VW	
	ν_7	VF bend.	110 D	109.5 M	99 W	
e''	ν_8	VF bend.	336 B		336 M	

References

- [1] IR. R. G. Cavell and H. C. Clark, *Inorg. Chem.* **3**, 1789 (1964).
 [2] IR.R. H. H. Claassen and H. Selig, *J. Chem. Phys.* **44**, 4039 (1966).
 [3] IR.R. I. R. Beattie, K. M. S. Livingston, G. A. Ozin, and D. J. Reynolds, *J. Chem. Soc. A* **1969**, 958.
 [4] IR.R. H. Selig, J. H. Holloway, J. Tyson, and H. H. Claassen, *J. Chem. Phys.* **53**, 2559 (1970).

No. 118 Arsenic pentafluoride AsF_5
Symmetry D_{3h}
Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1'	ν_1	AsF_3 s-stretch.	734 C		734.3 VS, p	
	ν_2	AsF_2 s-stretch.	644 C		644 M, p?	
a_2''	ν_3	AsF_2 a-stretch.	787 B	787.4 VS		
	ν_4	AsF_3 op-deform.	400 B	400.4 S		
e'	ν_5	AsF_3 d-stretch.	811 B	811.4 VS	813 M, dp	
	ν_6	AsF_3 d-deform.	372 C	372 S	366 (Liquid)	
	ν_7	AsF bend.	123 C	123 W	130 M, dp	
e''	ν_8	AsF bend.	386 C		386 M, dp	

References

- [1] IR.R. L. C. Hoskins and R. C. Lord, *J. Chem. Phys.* **46**, 2402 (1967).
 [2] Th. I. W. Levin, *J. Mol. Spectry.* **33**, 61 (1970).
 [3] R. H. Selig, J. H. Holloway, J. Tyson, and H. H. Claassen, *J. Chem. Phys.* **53**, 2559 (1970).
 [4] IR. L. C. Hoskins and C. N. Perng, *J. Chem. Phys.* **55**, 5063 (1971).

No. 119 Antimony pentachloride SbCl_5
Symmetry D_{3h} **Symmetry number $\sigma = 6$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
a_1'	ν_1	SbCl_3 s-stretch.	357 C		357 (10) p	
	ν_2	SbCl_2 s-stretch.	307 C	306 W (liquid)	307 (4) dp	
a_2''	ν_3	SbCl_2 a-stretch.	384 C	384 VS		
	ν_4	SbCl_3 op-deform.	154 D	154 (CCl_4 soln.)		
e'	ν_5	SbCl_3 d-stretch.	398 C	398 VS	397 (2)	
	ν_6	SbCl_3 d-deform.	177 C	172 S (CCl_4 soln.)	177 (4) dp	
	ν_7	SbCl bend.	72 D		72 dp	
e''	ν_8	SbCl bend.	165 C		165 (2) dp	

References

- [1] IR. J. K. Wilmschurst, *J. Mol. Spectry.* **5**, 343 (1960).
 [2] IR.R. G. L. Carlson, *Spectrochim. Acta* **19**, 1291 (1963).
 [3] Th. R. R. Holmes, R. M. Deiters, and J. A. Golen, *Inorg. Chem.* **8**, 2612 (1969).
 [4] Th. R. R. Holmes and R. M. Deiters, *J. Chem. Phys.* **51**, 4043 (1969).

No. 120 Tellurium hexafluoride TeF_6
Symmetry O_h **Symmetry number $\sigma = 24$**

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Gas)	
a_{2g}	ν_1	Sym. stretch.	697 B	ia	697.1 VS	
e_g	ν_2	Deg. stretch.	670 C	ia	670.3 W	
f_{1u}	ν_3	Deg. stretch.	751 C	751.0	ia	
	ν_4	Deg. deform.	327 C	326.5	ia	
f_{2g}	ν_5	Deg. deform.	314 C	ia	314 W	
f_{2u}	ν_6	Deg. deform.	197 E	ia	ia	OC($2\nu_6$) [3].

References

- [1] IR.R. B. Weinstock and Goodman, *Advan. Chem. Phys.* **9**, 169 (1966), and references cited there.
 [2] IR. S. Abramowitz and I. W. Levin, *J. Chem. Phys.* **44**, 3353 (1966).
 [3] R. H. H. Claassen, G. L. Goodman, J. H. Holloway, and H. Selig, *J. Chem. Phys.* **53**, 341 (1970).

No. 121 Iridium(VI)fluoride IrF₆
 Symmetry O_h
Symmetry number $\sigma = 24$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Gas)	
a _{1g}	ν_1	Sym. stretch.	702 B	ia	701.7 VS, p	
e _g	ν_2	Deg. stretch.	645 C	ia	645 W, dp	
f _{1u}	ν_3	Deg. stretch.	720 B	719.8 S	ia	
	ν_4	Deg. deform.	276 B	276.0 S	ia	
f _{2g}	ν_5	Deg. deform.	267 C	ia	267 W, dp	
f _{2u}	ν_6	Deg. deform.	206 D	ia	ia	OC($\nu_2+\nu_6, \nu_2-\nu_6, \nu_5+\nu_6$). [1].

References

- [1] IR.R. B. Weinstock and G. L. Goodman, *Advan. Chem. Phys.* **9**, 169 (1966), and references cited there.
 [2] R. H. H. Claassen and H. Selig, *Israel J. Chem.* **7**, 449 (1969).
 [3] IR. H. Kim, P. A. Souder, and H. H. Claassen, *J. Mol. Spectry.* **26**, 46 (1968).

No. 122 Sulfur chloride pentafluoride SCIF₅
 Symmetry C_{4v}
Symmetry number $\sigma = 4$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a ₁	ν_1	SF stretch.	855 B	854.6 VS	833 W, p	
	ν_2	SF ₄ s-stretch.	707 B	707.2 VS	704 S, p	
	ν_3	SF ₄ op-deform.	602 B	601.9 VS	603 W, p	
	ν_4	SCl stretch.	402 B	401.7 VS	403 VS, p	
b ₁	ν_5	SF ₄ a-stretch.	625 C	ia	625 M, dp	
	ν_6	SF ₄ op-deform.	271 C	ia	271 M, dp	
b ₂	ν_7	SF ₄ ip-deform.	505 C	ia	505 W, dp	
e	ν_8	SF ₄ d-stretch.	909 B	909.0 VS	927 W, dp	
	ν_9	SF bend.	579 B	579.0 M	584 VW, dp	
	ν_{10}	SF ₄ ip-deform.	441 B	441.0 S	442 M, dp	
	ν_{11}	SCl bend.	397 B	396.5 S	396 S, dp	

References

- [1] IR.R. L. H. Cross, H. L. Roberts, P. Goggin, and L. A. Woodward, *Trans. Faraday Soc.* **56**, 945 (1960).
 [2] IR.R. J. E. Griffiths, *Spectrochim. Acta* **23A**, 2145 (1967).

No. 123 Tungsten chloride pentafluoride $WClF_5$
Symmetry C_{4v}
Symmetry number $\sigma = 4$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a_1	ν_1	WF stretch.	743 C	743 M	744 VS, p	
	ν_2	WF ₄ s-stretch.	703 C	703 VS	703 M, p	
	ν_3	WCl stretch.	400 C	400 VS	407 S, p	
	ν_4	WF ₄ op-deform.	254 C	254 VS	257 W	
b_1	ν_5	WF ₄ a-stretch.	644 D	ia	644 W	
	ν_6	WF ₄ op-deform.	182 D	ia	182 W	
b_2	ν_7	WF ₄ ip-deform.	377 D	ia	377 M, dp	
e	ν_8	WF ₄ d-stretch.	671 C	671 S	661 M, dp	
	ν_9	WF bend.	302 C	302 M	307 M, dp	
	ν_{10}	WF ₄ ip-deform.	278 C	278 S	290 W	
	ν_{11}	WCl bend.	228 C	228 S	227 W	

Reference

- [1] IR.R. D. M. Adams, G. W. Fraser, D. M. Morris, and R. D. Peacock, *J. Chem. Soc. A* **1968**, 1131.

No. 124 Iodine heptafluoride IF_7
Symmetry D_{5h}
Symmetry number $\sigma = 10$

Sym. class	No.	Approximate type of mode ^a	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Gas)	
a_1'	ν_1	IF _{ax} s-stretch.	676 C	ia	676 W, p	
	ν_2	IF _{eq} s-stretch.	635 C	ia	635 S, p	
a_2''	ν_3	IF _{ax} a-stretch.	672 C	672 VS	ia	
	ν_4	F _{eq} IF _{ax} deform.	257 C	257 W	ia	
e_1'	ν_5	IF _{eq} a-stretch.	746 C	746 S, b	ia	
	ν_6	F _{eq} IF _{eq} deform.	425 C	425 VS	ia	
	ν_7	F _{ax} IF _{ax} deform.	363 C	363 S	ia	
e_1''	ν_8	F _{eq} IF _{ax} deform.	310 C	ia	310 W, dp	
e_2'	ν_9	IF _{eq} a-stretch.	510 C	ia	510 W, dp	
	ν_{10}	F _{eq} IF _{eq} deform.	352 C	ia	352 W, dp	
e_2''	ν_{11}	F _{eq} IF _{ax} deform.	200 D	ia	ia	CF[2]. OC($\nu_2 + \nu_{11}$, $\nu_5 + \nu_{11}$).

^a F_{ax} and F_{eq} represent the axial and equatorial fluorine atoms, respectively.

References

- [1] IR.R. H. H. Claassen, E. L. Gasner, and H. Salig, *J. Chem. Phys.* **49**, 1803 (1968).
 [2] IR.R.Th. H. H. Eysel and K. Seppelt, *J. Chem. Phys.* **56**, 5081 (1972).
 [3] Th. E. Wendling and S. Rahmondi, *Bull. Soc. Chim. (France)* **1**, 33 (1972).

No. 125 Digermane GeH_3GeH_3
Symmetry D_{3d}
Symmetry number $\sigma = 6$

-sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_{1g}	ν_1	GeH_3 s-stretch.	2068 C	ia	2068 S, p	
	ν_2	GeH_3 s-deform.	832 C	ia	832 VW, p	
	ν_3	GeGe stretch.	268 C	ia	268.4 S, p	
a_{1u}	ν_4	Torsion	146 E ^a	ia	ia	
a_{2u}	ν_5	GeH_3 s-stretch.	2077 B	2077.0 VS	ia	
	ν_6	GeH_3 s-deform.	756 B	756.0 VS	ia	
e_u	ν_7	GeH_3 d-stretch.	2091 B	2090.7 S	ia	
	ν_8	GeH_3 d-deform.	879 B	879.0 S	ia	
	ν_9	GeH_3 rock.	370 B	370.3 M	ia	
e_g	ν_{10}	GeH_3 d-stretch.	2081 E	ia		CF[4].
	ν_{11}	GeH_3 d-deform.	880 C	ia	879.6 M, dp	
	ν_{12}	GeH_3 rock.	567 C	ia	566.6 W, dp	

^a Estimated from combination bands.**References**

- [1] IR. D. A. Dows and R. M. Hexter, *J. Chem. Phys.* **24**, 1029 (1956).
 [2] IR.R. V. A. Crawford, K. H. Rhee, and M. K. Wilson, *J. Chem. Phys.* **37**, 2377 (1962).
 [3] IR.R. J. E. Griffiths and G. E. Walraffen, *J. Chem. Phys.* **40**, 321 (1964).
 [4] Th. E. A. Clark and A. Weber, *J. Chem. Phys.* **45**, 1759 (1966).

No. 126 Digermane- d_6 GeD_3GeD_3
Symmetry D_{3d}
Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_{1g}	ν_1	GeD_3 s-stretch.	1484 C	ia	1483.7 S, p	
	ν_2	GeD_3 s-deform.	600 C	ia	600 VW, p	
	ν_3	GeGe stretch.	264 C	ia	264 S, p	CF[3].
a_{1u}	ν_4	Torsion	106 E ^a	ia	ia	
a_{2u}	ν_5	GeD_3 s-stretch.	1490 B	1489.9 VS	ia	
	ν_6	GeD_3 s-deform.	546 C	546.2 S	ia	
e_u	ν_8	GeD_3 d-deform.	630 B	629.7 M	ia	
	ν_7	GeD_3 d-stretch.	1511 C	1511 S	ia	
	ν_9	GeD_3 rock.	262 C	262 S	ia	
e_g	ν_{10}	GeD_3 d-stretch.	1501 D	ia	1501	
	ν_{11}	GeD_3 d-deform.	626 C	ia	626.1 M, dp	
	ν_{12}	GeD_3 rock.	412 C	ia	412.0 W, dp	

^a From the product rule.**References**

- [1] IR.R. V. A. Crawford, K. H. Rhee, and M. K. Wilson, *J. Chem. Phys.* **37**, 2377 (1962).
 [2] IR.R. J. E. Griffiths and G. E. Walraffen, *J. Chem. Phys.* **40**, 321 (1964).
 [3] Th. E. A. Clark and A. Weber, *J. Chem. Phys.* **45**, 1759 (1966).

No. 127 Cyanogen fluoride FCN
Symmetry $C_{\infty v}$ Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
σ^+	ν_1	CN stretch.	2323 C	2323		
π	ν_2	Deform.	451 A	451.32		
σ^+	ν_3	CF stretch.	1077 A	1076.52		

References

- [1] IR. R. E. Dodd and R. Little, *Spectrochim. Acta* **16**, 1083 (1960).
 [2] IR. A. R. H. Cole, L. Isaacson, and R. C. Lord, *Spectrochim. Acta* **23**, 86 (1967).
 [3] Th. A. Ruoff, *Spectrochim. Acta* **26A**, 545 (1970).

No. 128 Cyanogen iodide ICN
Symmetry $C_{\infty v}$ Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
σ^+	ν_1	CN stretch.	2188 C	2188.0		
π	ν_2	Deform.	305 C	304.5		
σ^-	ν_3	CI stretch.	486 C	485.8		

References

- [1] IR. S. Hemple and E. R. Nixon, *J. Chem. Phys.* **47**, 4273 (1967).
 [2] Th. A. Ruoff, *Spectrochim. Acta* **26A**, 545 (1970).

No. 129 Carbonyl fluoride COF₂
Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a_1	ν_1	CO stretch.	1928 C	1928 VS	1944 VW	
	ν_2	CF ₂ s-stretch.	965 B	965 VS	965 VS	
	ν_3	CF ₂ deform.	584 C	584 M	571 W	
b_1	ν_4	CF ₂ a-stretch.	1249 B	1249 VS	1238 VW	
	ν_5	CO deform.	626 C	626 M	620 M	
b_2	ν_6	Op-deform.	774 B	774 M	771 VW	

References

- [1] IR.R. A. H. Nielsen, T. G. Burke, P. J. H. Woltz, and E. A. Jones, *J. Chem. Phys.* **20**, 596 (1952).
 [2] Th. J. Overend and J. R. Scherer, *J. Chem. Phys.* **32**, 1296 (1960).

No. 130 **Carbonyl chloride** **COCl₂**
 Symmetry C_{2v}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a ₁	ν_1	CO stretch.	1827 B	1827 VS	1807 M	
	ν_2	CCl ₂ s-stretch.	567 C	567 M	573 VS	
	ν_3	CCl ₂ deform.	285 C	285 W	302 S	
b ₁	ν_4	CCl ₂ a-stretch.	849 B	849 S	832 VW	
	ν_5	CO deform.	440 C	440 M	442 M	
b ₂	ν_6	Op-deform.	580 C	580 M		

References

- [1] IR.R. A. H. Nielsen, T. G. Burke, P. J. H. Woltz, and E. A. Jones, J. Chem. Phys. **20**, 596 (1952).
 [2] IR. E. Catalano and K. S. Pitzer, J. Amer. Chem. Soc. **80**, 1054 (1958).
 [3] IR.R. J. Overend and J. C. Evans, Trans. Faraday Soc. **55**, 1817 (1959).
 [4] Th. J. Overend and J. R. Scherer, J. Chem. Phys. **32**, 1296 (1960).

No. 131 **Carbonyl bromide** **COBr₂**
 Symmetry C_{2v}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a ₁	ν_1	CO stretch.	1828 B	1828 VS		
	ν_2	CBr ₂ s-stretch.	425 C	425 M	429 S	
	ν_3	CBr ₂ deform.	181 D		181 S	
b ₁	ν_4	CBr ₂ a-stretch.	757 C	787 VS		
	ν_5	CO deform.	350 C	747 VS		
b ₂	ν_6	Op-deform.	512 B	350 VW	350 M, b	FR($\nu_2 + \nu_5$).
				512 M		

References

- [1] IR.R. J. Overend and J. C. Evans, Trans. Faraday Soc. **15**, 1817 (1959).
 [2] Th. J. Overend and J. R. Scherer, J. Chem. Phys. **32**, 1296 (1960).

No. 132 Thiocarbonyl fluoride SCF₂
Symmetry C_{2v}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	CS stretch.	1368 C	1368 VS		
	ν_2	CF ₂ s-stretch.	787 C	787 M		
	ν_3	CF ₂ scis.	526 C	526 M		
b ₁	ν_4	CF ₂ a-stretch.	1189 C	1189 S		
	ν_5	CF ₂ rock.	417 C	417 VW		
b ₂	ν_6	CF ₂ wag.	622 C	622 W		

References

- [1] IR. A. J. Downs, Spectrochim. Acta **19**, 1165 (1963).
 [2] IR. M. J. Hopper, J. W. Russel, and J. Overend, Spectrochim. Acta **28A**, 1215 (1972).

No. 133 Thiocarbonyl chloride CSCI₂
Symmetry C_{2v}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a ₁	ν_1	CS stretch.	1137 C	1137 VS	1121 (10)	
	ν_2	CCl ₂ s-stretch.	505 C	505 M	496 (5)	
	ν_3	CCl ₂ scis.	220 D	220 VW, b	200 (1)	
b ₁	ν_4	CCl ₂ a-stretch.	816 C	816 VS		
	ν_5	CS deform.	294 C	294	287 (3)	
b ₂	ν_6	op-Bend.	473 C	473 W		

References

- [1] R. H. W. Thompson, J. Chem. Phys. **6**, 748 (1938).
 [2] IR. A. J. Downs, Spectrochim. Acta **19**, 1165 (1963).
 [3] IR. M. J. Hopper, J. W. Russell, and J. Overend, Spectrochim. Acta **28A**, 1215 (1972).

No. 134 Isocyanic acid HNCO
Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a'	ν_1	NH stretch.	3531 C	3531 S	3410	
	ν_2	CO stretch.	2274 C	2274 VS		
	ν_3	CN stretch.	1327 C	1327 W	1318	
	ν_4	NH bend.	762 B ^a	777.1 S		
	ν_5	NCO deform.	643 B ^a	659.8 M		
a''	ν_6	NCO deform.	610 B ^a	577.5 M		

^a Three fundamentals, ν_4 , ν_5 , and ν_6 , are strongly coupled through Coriolis interaction. The unperturbed frequencies are given in this column [3].

References

- [1] R. G. Herzberg and C. Reid, Disc. Faraday Soc. **9**, 92 (1950).
 [2] IR. C. Reid, J. Chem. Phys. **18**, 1544 (1954).
 [3] IR. R. A. Ashby and R. L. Werner, J. Mol. Spectry. **18**, 184 (1965).

No. 135 Isocyanic acid-d DNCO
Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a'	ν_1	ND stretch.	2621 B	2620.7		
	ν_2	CO stretch.	2235 C	2235		
	ν_3	CN stretch.	1310 C	1310		
	ν_4	ND bend.	758 C ^a	766.8		
	ν_5	CNO deform.	458 C ^a	460		
a''	ν_6	CNO deform.	603 B ^a	602.9		

^a See footnote of HNCO.

References

- [1] IR. R. A. Ashby and R. L. Werner, Spectrochim. Acta **22**, 1345 (1966).
 [2] IR. W. D. Sheasley, C. W. Mithews, E. L. Ferretti, K. N. Rao, J. Mol. Spectry. **37**, 377 (1971).

No. 136 Carbonyl chlorofluoride COClF

Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a'	ν_1	CO stretch.	1868 C	1876 VS	1858 M	FR($\nu_2 + \nu_3$).
	ν_2	CF stretch.	1095 B	1847 VS	1832 M	
	ν_3	CCl stretch.	776 C	1095 S	1085 VW	
	ν_4	CO deform.	501 C	776 M	765 VS	
	ν_5	CClF deform.	415 C	501 W	506 S	
a''	ν_6	Op-deform.	667 B	415 VW	410 M	
				667 M	665 VW	

References

- [1] IR.R. A. H. Nielsen, T. G. Burke, P. J. H. Woltz, and E. A. Jones, J. Chem. Phys. **20**, 596 (1952).
 [2] Th. J. Overend and J. R. Scherer, J. Chem. Phys. **32**, 1296 (1960).

No. 137 Carbonyl bromochloride COBrCl

Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a'	ν_1	CO stretch.	1828 B	1828 VS		
	ν_2	CCl stretch.	806 C	806 VS		
	ν_3	CBr stretch.	517 C	517 M	518 M	
	ν_4	CO deform.	374 C	374 W	372 M	
	ν_5	CBrCl deform.	240 D		240 S	
a''	ν_6	Op-deform.	547 B	547 W		

References

- [1] IR.R. J. Overend and J. C. Evans, Trans. Faraday Soc. **55**, 1817 (1959).
 [2] Th. J. Overend and J. R. Scherer, J. Chem. Phys. **32**, 1296 (1960).

No. 138 Thiocarbonyl bromochloride CSBrCl

Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a'	ν_1	CS stretch.	1130 C	1130 S	1125 W, p	
	ν_2	CBr stretch.	764 C	764 S	761 W	
	ν_3	CBr stretch.	438 C	438 M	437 S, p	
	ν_4	CBrCl deform.	256 D	256 W	257 M, p	
	ν_5	CS deform.	222 D	222 VW (liquid)	222 M, p	
a''	ν_6	op-Bend.	405 E			CF [1].

Reference

[1] IR.R.Th. J. L. Brema and D. C. Moule, Spectrochim. Acta **28A**, 809 (1972).No. 139 Trifluoromethane-d CDF₃Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)		
a_1	ν_1	CD stretch.	2261 B	2261.0 S		
	ν_2	CF ₃ s-stretch.	1111 B	1110.6 M		
	ν_3	CF ₃ s-deform.	694 B	694.2 M		
e	ν_4	CD bend.	1202 D	1202.2 M		FR($\nu_3 + \nu_6$).
	ν_5	CF ₃ d-stretch.	975 B	975.1 S		
	ν_6	CF ₃ d-deform.	502 B	502.4 M		

References

- [1] IR. S. R. Polo and M. K. Wilson, J. Chem. Phys. **21**, 1129 (1953).
 [2] IR. C. C. Costain, J. Mol. Spectry. **9**, 317 (1962).
 [3] IR.Th. A. Ruoff, H. Bürger, and S. Biedermann, Spectrochim. Acta **27A**, 1359 (1971).

No. 140 Chlorotrifluoromethane CClF_3
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	CF_3 s-stretch.	1105 C	1105 VS	1106 W, p	
	ν_2	CCl stretch.	781 C	781 S	781.7 VS, p	
	ν_3	CF_3 s-deform.	476 C		475.8 S, p	
e	ν_4	CF_3 d-stretch.	1212 C	1212 VS	1217 W, dp	
	ν_5	CF_3 d-deform.	563 C	563 M	560 W, dp	
	ν_6	CCl bend.	350 C		350 M, dp	

References

- [1] IR. H. W. Thompson and R. B. Temple, J. Chem. Soc. **1948**, 1422.
 [2] IR. E. K. Plyler and W. S. Benedict, J. Res. NBS **47**, 202 (1951).
 [3] R. H. H. Claassen, J. Chem. Phys. **22**, 50 (1954).
 [4] R. W. Holzer and H. Moser, J. Mol. Spectry. **20**, 188 (1966).
 [5] Th. L. H. Ngai and R. H. Mann, J. Mol. Spectry. **38**, 322 (1971).

No. 141 Bromotrifluoromethane CBrF_3
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	CF_3 s-stretch.	1089 C	1089 VS	1082 (8) p	
	ν_2	CF_3 s-deform.	760 C	760 VS	761 (9) p	
	ν_3	CBr stretch.	349 C	350*	349 (7) p	
e	ν_4	CF_3 d-stretch.	1210 C	1210 VS	1207 (8) dp	
	ν_5	CF_3 d-deform.	547 C	547 M	541 (2) dp	
	ν_6	CBr bend.	306 C	297*	306 (2) dp	

* Estimated from overtone and combination bands.

References

- [1] IR.R. W. F. Edgell and C. E. May, J. Chem. Phys. **20**, 1822 (1952).
 [2] IR. E. K. Plyler and N. Acquista, J. Res. NBS **48**, 92 (1952).
 [3] IR. P. R. McGee, F. F. Cleveland, A. G. Meister, C. E. Decker, and S. I. Miller, J. Chem. Phys. **21**, 242 (1953).
 [4] R. W. Holzer, J. Mol. Spectry. **25**, 123 (1968).
 [5] Th. L. H. Ngai and R. H. Mann, J. Mol. Spectry. **38**, 322 (1971).

No. 142 Trifluoroiodomethane CF₃ISymmetry C_{3v}Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a ₁	ν_1	CF ₃ s-stretch.	1080 C	1080 VS	1056	
	ν_2	CF ₃ s-deform.	742 C	742 VS	741	
	ν_3	CI stretch.	286 E	286 ^a	284 ^a	
e	ν_4	CF ₃ d-stretch.	1187 C	1187 VS	1168	
	ν_5	CF ₃ d-deform.	537 C	537	537	
	ν_6	CI bend.	260 E	260 ^a	260 ^a	

^a Estimated from overtone and combination bands.

References

- [1] IR.R. W. F. Edgell and C. E. May, J. Chem. Phys. **20**, 1822 (1952).
 [2] IR. E. K. Plyler and N. Acquista, J. Res. NBS **48**, 92 (1952).
 [3] IR. P. R. McGee, F. F. Cleveland, A. G. Meister, C. E. Decker, and S. I. Miller, J. Chem. Phys. **21**, 242 (1953).
 [4] Th. L. H. Ngai and R. H. Mann, J. Mol. Spectry. **38**, 322 (1971).

No. 143 Trichlorofluoromethane CCl₃FSymmetry C_{3v}Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Gas)	
a ₁	ν_1	CF stretch.	1085 C	1085 S	1090 VW, p	
	ν_2	CCl ₃ s-stretch.	535 C	535 M	535 VS, p	
	ν_3	CCl ₃ s-deform.	350 C	350 VS	349.5 S, p	
e	ν_4	CCl ₃ d-deform.	847 C	847 VS	847 M, dp	
	ν_5	CF bend.	394 C	401 VW	394 S, dp	
	ν_6	CCl ₃ d-deform.	241 C		241 S, dp	

References

- [1] IR. H. W. Thompson and R. B. Temple, J. Chem. Soc. **1948**, 1422.
 [2] R. J. P. Zietlow, F. F. Cleveland, and A. G. Meister, J. Chem. Phys. **18**, 1076 (1950).
 [3] IR. E. K. Plyler and W. S. Benedict, J. Res. NBS **47**, 202 (1951).
 [4] IR. J. P. Zietlow and F. F. Cleveland, J. Chem. Phys. **21**, 1778 (1951).
 [5] R. H. H. Claassen, J. Chem. Phys. **22**, 50 (1954).
 [6] R. W. Holzer and H. Moser, J. Mol. Spectry. **20**, 188 (1966).
 [7] Th. L. H. Ngai and R. H. Mann, J. Mol. Spectry. **38**, 322 (1971).

No. 144 Trichloriodomethane CCl₃I
Symmetry C_{3v}

Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Liquid)	(Liquid)	
a ₁	ν_1	CCl ₃ s-stretch.	684 D	684 VS	687 W, b, p	
	ν_2	CI stretch.	390 D	390 M	405 W, p	
	ν_3	CCl ₃ s-deform.	224 D	224 W	224 W	
e	ν_4	CCl ₃ d-stretch.	755 D	755 VS		
	ν_5	CCl ₃ d-deform.	284 D	284 W	288 W, dp	
	ν_6	CI bend.	188 D	188 M		

Reference[1] IR.R. R. H. Mann and P. M. Manis, J. Mol. Spectry. **45**, 65 (1973).

No. 145 Tribromofluoromethane CBr₃F
Symmetry C_{3v}

Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
					(Liquid)	
a ₁	ν_1	CF stretch.	1069 D		1069 p	
	ν_2	CBr ₃ s-stretch.	398 D		398 p	
	ν_3	CBr ₃ s-deform.	218 D		218 p	
e	ν_4	CBr ₃ d-stretch.	743 D		743 dp	
	ν_5	CF bend.	306 D		306 dp	
	ν_6	CBr ₃ d-deform.	150 D		150 dp	

References

- [1] R. M. L. Delwaille and M. F. Francois, Comptes Rendus **214**, 828 (1942).
 [2] R. M. L. Delwaille and M. F. Francois, J. Phys. (Paris) **7**, 15 (1946).
 [3] Th. A. G. Meister, S. E. Rosson, and F. F. Cleveland, J. Chem. Phys. **18**, 346 (1950).
 [4] Th. L. H. Ngai and R. H. Mann, J. Mol. Spectry. **38**, 322 (1971).

No. 146 Dichlorodifluoromethane CCl_2F_2 Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Gas)	
a_1	ν_1	CF_2 s-stretch.	1101 C	1101 S	1098 M	
	ν_2	CCl_2 s-stretch.	667 C	667 S	667.2 S, p	
	ν_3	CF_2 scis.	458 D		457.5 S	
	ν_4	CCl_2 scis.	262 C		261.5 S, dp	
a_2	ν_5	CF_2 twist.	322 C	ia	322 W, dp	
b_1	ν_6	CF_2 a-stretch.	1159 C	1159 S	1167 W, dp	
	ν_7	CF_2 rock.	446 C	446 W		
b_2	ν_8	CCl_2 a-stretch.	902 E	922 VS	923 W, dp	
				882 VS		FR($\nu_3 + \nu_9$).
	ν_9	CF_2 wag.	437 C	437 W	433 M	

References

See No. 143(CCl_3F).No. 147 Dibromodifluoromethane CBr_2F_2 Symmetry C_{2v} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_1	ν_1	CF_2 s-stretch.	1090 C	1090	1077 W, p	
	ν_2	CF_2 scis.	623 C	623	624 W, p	
	ν_3	CBr_2 s-stretch.	340 D		340 S, p	
	ν_4	CBr_2 scis.	168 D		168 M, p	
a_2	ν_5	CF_2 twist.	281 D	ia	281 W, dp	
b_1	ν_6	CF_2 a-stretch.	1153 C	1153 VS	1141 VW, dp	
	ν_7	CF_2 rock.	369 D		369 VW, dp	
b_2	ν_8	CBr_2 a-stretch.	831 C	831 VS	816 W, dp	
	ν_9	CF_2 wag.	325 D		325 VW, dp	

References

- [1] R. G. Glockler and G. R. Leader, *J. Chem. Phys.* **7**, 553 (1939).
 [2] IR. E. K. Plyler and N. Acquista, *J. Res. NBS* **48**, 92 (1952).
 [3] IR. C. E. Decker and F. F. Cleveland, *J. Chem. Phys.* **21**, 189 (1953).
 [4] IR.R. C. E. Decker, A. G. Meister, F. F. Cleveland, and R. B. Bernstein, *J. Chem. Phys.* **21**, 1781 (1953).
 [5] Th. L. H. Ngai and R. H. Mann, *J. Mol. Spectry.* **38**, 322 (1971).

No. 148 Bromodichlorofluoromethane CBrCl₂F
 Symmetry C_s
Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
<i>a'</i>	ν_1	CF stretch.	1080 C	1080 VS	1070 (3) p	
	ν_2	CCl ₂ s-stretch.	796 C	796 VS	783 (12) p	
	ν_3	CBr stretch.	502 C	502 M	504 (100) p	
				(liquid)		
	ν_4	CCl ₂ scis.	339 D		339 (14) p	
	ν_5	CBrF scis.	306 D		306 (98) p	
	ν_6	CCl ₂ wag.	218 D		218 (53) p	
<i>a''</i>	ν_7	CCl ₂ a-stretch.	838 C	838 VS	831 (5) dp	
	ν_8	CCl ₂ twist.	392 D		392 (10) dp	
	ν_9	CCl ₂ rock.	204 D		204 (34) dp	

References

- [1] R. M. L. Delwaille and F. François, *Comptes Rendus* **214**, 828 (1942).
 [2] IR.R. R. L. Gilbert, E. A. Piotrowski, J. M. Dowing, and F. F. Cleveland, *J. Chem. Phys.* **31**, 1633 (1959).
 [3] Th. L. H. Ngai and R. H. Mann, *J. Mol. Spectry.* **38**, 322 (1971).

No. 149 Dibromochlorofluoromethane CBr₂ClF
 Symmetry C_s
Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
<i>a'</i>	ν_1	CF stretch.	1075 C	1075 VS	1059 (2) p	
	ν_2	CCl stretch.	806 E	795 VS	793 (4) p	
				814 VS	815 (5) p	
	ν_3	CBr ₂ s-stretch.	460 C	460 W	464 (58) p	FR($\nu_3 + \nu_4$).
				(liquid)		
	ν_4	CClF scis.	341 D		341 (17) p	
	ν_5	CBr ₂ scis.	268 D		268 (100) p	
	ν_6	CBr ₂ wag.	162 D		162 (50) p	
<i>a''</i>	ν_7	CBr ₂ a-stretch.	754 C	754 VS	742 (11) dp	
	ν_8	CBr ₂ twist.	308 D		308 (6) dp	
	ν_9	CBr ₂ rock.	196 D		196 (20) dp	

ReferencesSee No. 148(CBrCl₂F).

No. 150 Borine Carbonyl $^{10}\text{BH}_3\text{CO}$
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
a_1	ν_1	BH_3 s-stretch.	2387 D	2387 M (Gas)		
	ν_2	CO stretch.	2166 D	2166.0 VS (solid)		
	ν_3	BH_3 s-deform.	1083 C	1083.1 S		
	ν_4	BC stretch.	707 B	707.0 S		
e	ν_5	BH_3 d-stretch.	2456 D	2456 VS		
	ν_6	BH_3 d-deform.	1115 E	1114.8 S		
	ν_7	BH_3 rock.	819 B	818.8 M		
	ν_8	BCO bend.	314 B	313.7 S		

References

- [1] IR. R. D. Cowan, J. Chem. Phys. **18**, 1101 (1950).
 [2] IR.Th. G. W. Bethke and M. K. Wilson, J. Chem. Phys. **26**, 1118 (1957).

No. 151 Borine Carbonyl- d_3 $^{10}\text{BD}_3\text{CO}$
Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
a_1	ν_1	BD_3 s-stretch.	1695 C	1694.8 M (Gas)		
	ν_2	CO stretch.	2169 D	2169.0 VS		
	ν_3	BD_3 s-deform.	888 D	888.4 W (solid)		
	ν_4	BC stretch.	630 B	629.5 S		
e	ν_5	BD_3 d-stretch.	1852 C	1852 S		
	ν_6	BD_3 d-deform.	802 B	801.5 W		
	ν_7	BD_3 rock.	718 B	718.0 S		
	ν_8	BCO bend.	266 B	266.0 S		

References

See No. 150($^{10}\text{BH}_3\text{CO}$).

No. 152 Borine Carbonyl $^{11}\text{BH}_3\text{CO}$
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
a_1	ν_1	BH_3 s-stretch.	2380 C	2379 M (Gas) (solid)	2380 S, p (Liquid)	
	ν_2	CO stretch.	2165 D	2164.7 VS	2169 S, p	
	ν_3	BH_3 s-deform.	1073 C	1073.4 S	1073 S, p	
	ν_4	BC stretch.	691 B	691.4 S	692 W, p	
e	ν_5	BH_3 d-stretch.	2444 D	2444 VS	2434 S	
	ν_6	BH_3 d-deform.	1106 E	1105.8 S	1101 M	
	ν_7	BH_3 rock.	809 B	809.3 M	816 W	
	ν_8	BCO bend.	313 B	313.2 S	317 M	

References

- [1] IR. R. D. Cowan, J. Chem. Phys. **18**, 1101 (1950).
 [2] IR.Th. G. W. Bethke and M. K. Wilson, J. Chem. Phys. **26**, 1118 (1957).
 [3] R.Th. R. C. Taylor, J. Chem. Phys. **26**, 1131 (1957).

No. 153 Borine Carbonyl- d_3 $^{11}\text{BD}_3\text{CO}$
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
a_1	ν_1	BD_3 s-stretch.	1679 C	1679.0 M (Gas)	1678 S (Liquid)	
	ν_2	CO stretch.	2169 D	2168.5 VS	2169 S	
	ν_3	BD_3 s-deform.	860 C	867.7 W (solid)	860 M	
	ν_4	BC stretch.	625 B	624.8 S	619 M	
e	ν_5	BD_3 d-stretch.	1840 C	1840 S	1825 S	
	ν_6	BD_3 d-deform.	801 B	801.3 W	808 M	
	ν_7	BD_3 rock.	709 B	709.3 S	706 W	
	ν_8	BCO bend.	266 B	266.0 S	264 W	

ReferencesSee No. 152($^{11}\text{BH}_3\text{CO}$).

No. 154 Methylmercuric iodide CH_3HgI Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected frequency value of cm^{-1}	Infrared cm^{-1} (Solid)	Raman cm^{-1} (CH_3NO_2 soln.)	Comments
a_1	ν_1	CH_3 s-stretch.	2920 D	2920 M	2914 (Solid)	
	ν_2	CH_3 s-deform.	1180 D	1180 M	1182 p	
	ν_3	CHg stretch.	531 D	531 M	538 p	
	ν_4	HgI stretch.	184 D	184 M	180 p	
e	ν_5	CH_3 d-stretch.	3008 D	3008 M		
	ν_6	CH_3 d-deform.	1400 D	1400 W		
	ν_7	CH_3 rock.	781 D	781 S		
	ν_8	CHgI bend.	64 D	64 M		

References

- [1] R. F. Fehér, W. Kolb and L. Leverenz, *Z. Naturforsch.*, A 2, 454 (1947).
 [2] IR.R. P. L. Goggin and L. A. Woodward, *Trans. Faraday Soc.* 62, 1423 (1966).
 [3] IR.Th. J. H. S. Green, *Spectrochim. Acta* 24A, 863 (1968).

No. 155 Methylsilane CH_3SiH_3 Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected frequency value of cm^{-1}	Infrared cm^{-1} (Gas)	Raman cm^{-1}	Comments
a_1	ν_1	CH_3 s-stretch.	2898 E	2928.8 M 2867.4 M		FR($\nu_2 + \nu_5$).
	ν_2	SiH_3 s-stretch.	2169 C	2169 S		
	ν_3	CH_3 s-deform.	1260 C	1260 M		
	ν_4	SiH_3 s-deform.	940 C	940 VS		
	ν_5	CSi stretch.	700 C	700 S		
a_2	ν_6	Torsion	187 D	ia		187.6(A). MW. 186.9(E).
e	ν_7	CH_3 d-stretch.	2982 A	2981.6 M		
	ν_8	SiH_3 d-stretch.	2166 A	2965.7 S		
	ν_9	CH_3 d-deform.	1403 C	1403 M		
	ν_{10}	SiH_3 d-deform.	980 D	980 ^a		
	ν_{11}	CH_3 rock.	868 B	867.5 S		
	ν_{12}	SiH_3 rock.	540 C	540 M		

^a The band origin is not determined clearly.

References

- [1] IR. S. Kaye and S. Tannenbaum, *J. Org. Chem.* 18, 1750 (1953).
 [2] IR. D. F. Ball, P. L. Goggin, D. C. McKean, and L. A. Woodward, *Spectrochim. Acta* 16, 1358 (1960).
 [3] IR. R. E. Wilde, *J. Mol. Spectry.* 8, 427 (1962).
 [4] IR. M. Randie, *Spectrochim. Acta* 18, 115 (1962).
 [5] Th. E. A. Clark and A. Weber, *J. Chem. Phys.* 45, 1759 (1966).

No. 156 Methylsilane-d₃ CH₃SiD₃
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency		Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)			
<i>a</i> ₁	ν_1	CH ₃ s-stretch.	2923 A	2923.4 M			
	ν_2	SiD ₃ s-stretch.	1558 C	1558 M			
	ν_3	CH ₃ s-deform.	1262 C	1262 S			
	ν_4	CSi stretch.	741 C	741 VS			
	ν_5	SiD ₃ s-deform.	652 C	652 S			
<i>a</i> ₂	ν_6	Torsion	172 D		ia		172.1(A) MW. 171.9(E)
<i>e</i>	ν_7	CH ₃ d-stretch.	2982 A	2981.8 M			
	ν_8	SiD ₃ d-stretch.	1577 C	1577 VS			
	ν_9	CH ₃ d-deform.	1401 C	1401 M			
	ν_{10}	CH ₃ rock.	825 C	825 S			
	ν_{11}	SiD ₃ d-deform.	668 C	668 S			
	ν_{12}	SiD ₃ rock.	433 C	433			

References

- [1] IR. D. E. Ball, P. L. Goggin, D. C. McKean, and L. A. Woodward, *Spectrochim. Acta* **16**, 1358 (1960).
 [2] IR. R. E. Wilde, *J. Mol. Spectry.* **8**, 427 (1962).
 [3] Th. E. A. Clark and A. Weber, *J. Chem. Phys.* **45**, 1759 (1966).

No. 157 Methylgermane CH₃GeH₃
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency		Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)			
<i>a</i> ₁	ν_1	CH ₃ s-stretch.	2938 B	2937.8 S			
	ν_2	GeH ₃ s-stretch.	2085 B	2084.8 S			
	ν_3	CH ₃ s-deform.	1254 B	1254.1 M			
	ν_4	GeH ₃ s-deform.	843 B	842.5 VS			
	ν_5	CGe stretch.	602 B	601.6 VS			
<i>a</i> ₂	ν_6	Torsion	157 D		ia		158.1(A) MW. 156.1(E)
<i>e</i>	ν_7	CH ₃ d-stretch.	2997 B	2997.2 S			
	ν_8	GeH ₃ d-stretch.	2084 B	2084.3 S			
	ν_9	CH ₃ d-deform.	1428 C	1427.7 W			
	ν_{10}	GeH ₃ d-deform.	900 C	900.4 S			
	ν_{11}	CH ₃ rock.	848 C	847.5 S			
	ν_{12}	GeH ₃ rock.	506 C	505.9 S			

References

- [1] IR. J. E. Griffiths, *J. Chem. Phys.* **38**, 2879 (1963).
 [2] Th. E. A. Clark and A. Weber, *J. Chem. Phys.* **45**, 1759 (1966).

No. 158 Methyl-d₃-germane CD₃GeH₃
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	CD ₃ s-stretch.	2090 C	2091.0 S		
	ν_2	GeH ₃ s-stretch.	2073 C	2073 S		
	ν_3	CD ₃ s-deform.	978 C	977.5 S		
	ν_4	GeH ₃ s-deform.	844 C	844.3 S		
	ν_5	CGe stretch.	550 B	549.6 S		
a ₂	ν_6	Torsion	132 D	ia		131.7(A) MW. 131.5(E)
e	ν_7	CD ₃ d-stretch.	2248 B	2247.5 S		
	ν_8	GeH ₃ d-stretch.	2089 B	2089.0 S		
	ν_9	CD ₃ d-deform.	1032 B	1032.0 M		
	ν_{10}	GeH ₃ d-deform.	904 B	903.9 S		
	ν_{11}	CD ₃ rock.	713 C	713 S		
	ν_{12}	GeH ₃ rock.	452 B	451.8 S		

ReferencesSee No. 157(CH₃GeH₃).
No. 159 Methyltriiodogermane CH₃GeI₃
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Liquid)	cm ⁻¹ (Liquid)	
a ₁	ν_1	CH ₃ s-stretch.	2911 C	2911 M	2909 W	
	ν_2	CH ₃ s-deform.	1225 C	1225 M	1222 W	
	ν_3	CGe stretch.	597 C	597 S	596 M	
	ν_4	GeI ₃ s-stretch.	195 C	196 M	195 S	
	ν_5	GeI ₃ s-deform.	92 C	96 M	92 S	
a ₂	ν_6	Torsion	123 D	123 Vw	123 Vw	
e	ν_7	CH ₃ d-stretch.	2999 C	2999 M	2996 W	
	ν_8	CH ₃ d-deform.	1392 C	1392 S	1394	
	ν_9	CH ₃ rock.	810 C	810 S	809	
	ν_{10}	GeI ₃ d-stretch.	252 C	252 S	251 M	
	ν_{11}	GeI ₃ rock.	147 C		147 M	
	ν_{12}	GeI ₃ d-deform.	67 C		67 S	

References

- [1] IR. R. J. Cross and F. Glockling, *J. Organometal. Chem.* **3**, 146 (1965).
 [2] IR.R.Th. J. R. Durig, C. F. Jumper, and J. N. Willis, Jr., *J. Mol. Spectry.* **37**, 260 (1971).

No. 160 Methylstannane CH_3SnH_3
Symmetry C_{3v}

Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a_1	ν_1	CH_3 s-stretch.	2933 B	2932.5 W		
	ν_3	CH_3 s-deform.	1209 B	1209.3 W		
	ν_2	SnH_3 s-stretch.	1875 D	1874.5 S		OV(ν_8).
	ν_4	SnH_3 s-deform.	695 B	694.5 S		
	ν_5	SnC stretch.	527 B	526.9 M		
a_2	ν_6	Torsion	109 C			MW[1].
e	ν_7	CH_3 d-stretch.	3005 B	3005.4 W		
	ν_8	SnH_3 d-stretch.	1875 B	1874.5 S		OV(ν_2).
	ν_9	CH_3 d-deform.	1417 B	1417.0 W		
	ν_{10}	CH_3 rock.	774 C	774.1 M		
	ν_{11}	SnH_3 d-deform.	741 C	741.3 M		
	ν_{12}	SnH_3 rock.	416 B	416.3 M		

References

- [1] MW. P. Cahill and S. Butcher, J. Chem. Phys. **35**, 2255 (1961).
 [2] IR. H. Kimmel and C. R. Dillard, Spectrochim. Acta **24A**, 909 (1968).

No. 161 Methylstannane- d_3 CH_3SnD_3
Symmetry C_{3v}

Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1}	
a_1	ν_1	CH_3 s-stretch.	2930 D	2930 M		
	ν_2	SnD_3 s-stretch.	1352 D	1352.0 S		OV(ν_9).
	ν_3	CH_3 s-deform.	1205 C	1204.5 S		
	ν_4	SnC stretch.	509 C	509.1 M		
	ν_5	SnD_3 s-deform.	493 C	493.0 S		
a_2	ν_6	Torsion	101 C			MW[1].
e	ν_7	CH_3 d-stretch.	3000 D	3000 W		
	ν_8	CH_3 d-deform.	1400 D	1400 W		
	ν_9	SnD_3 d-stretch.	1352 C	1352.0 S		OV(ν_2).
	ν_{10}	CH_3 rock.	765 D	765 W		
	ν_{11}	SnD_3 d-deform.	503 C	502.5 S		
	ν_{12}	SnD_3 rock.	317 D	316.6 M		

ReferencesSee No. 160(CH_3SnH_3).

No. 162 Methyl-d₃-stannane CD₃SnH₃
Symmetry C_{3v}

Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a ₁	ν_1	CD ₃ s-stretch.	2144 B	2144.3 M		
	ν_2	SnH ₃ s-stretch.	1889 D	1889.0 S		OV(ν_8).
	ν_3	CD ₃ s-deform.	920 B	920.2 M		
	ν_4	SnH ₃ s-deform.	704 B	703.5 S		
	ν_5	SnC stretch.	478 B	478.0 M		
a ₂	ν_6	Torsion	88 C			MW[1].
e	ν_7	CD ₃ d-stretch.	2255 C	2254.5 M		
	ν_8	SnH ₃ d-stretch.	1889 B	1889.0 S		OV(ν_2).
	ν_9	CD ₃ d-deform.	1017 C	1017.1 W		
	ν_{10}	SnH ₃ d-deform.	738 B	738.1 M		
	ν_{11}	CD ₃ rock.	628 B	628.4 S		
	ν_{12}	SnH ₃ rock.	392 C	392.4 W		

ReferencesSee No. 160(CH₃SnH₃).

No. 163 Cyanogen C₂N₂
Symmetry D_{∞h}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Gas)	
σ_g^+	ν_1	CN s-stretch.	2330 B	2339.9 ^a	2330.5	
	ν_2	NN stretch.	846 C	ia	845.5	
σ_u^+	ν_3	CN a-stretch.	2158 A	2157.83	ia	
π_g	ν_4	CCN bend.	503 C	ia	502.8	
π_u	ν_5	CCN bend.	234 B	233.1 ^a	233.7	

^a Determined from difference bands[3].**References**

- [1] R. A. Langseth and C. K. Moller, Acta Chem. Scand. **4**, 725 (1950).
 [2] IR. G. D. Craine and H. W. Thompson, Trans. Faraday Soc. **49**, 1273 (1953).
 [3] IR. A. G. Maki, J. Chem. Phys. **43**, 3193 (1965).
 [4] Th. W. Sawodnt and A. Ruoff, J. Mol. Spectry. **34**, 173 (1970).
 [5] R.IR.Th. L. J. Jones, J. Mol. Spectry. **45**, 55 (1973).
 [6] IR. M. A. Picard, Spectrochim. Acta **29**, 423 (1973).
 [7] R.Th. L. H. Jones, J. Mol. Spectry. **49**, 82 (1974).

No. 164 Dichloroacetylene C_2Cl_2
 Symmetry D_{2h}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
σ_g^+	ν_1	CC stretch.	2234 D	ia	2234 S, p	
	ν_2	CCl stretch.	477 D	ia	477 M, p	
σ_u^+	ν_3	CCl stretch.	988 C	988 VS	ia	
π_g	ν_4	CCCl deform.	333 D	ia	333 VS, dp	
π_u	ν_5	CCCl deform.	172 C	172 S	ia	

Reference[1] IR.R. P. Klaboe and E. Kloster-Jensen, *Spectrochim. Acta* **26A**, 1567 (1970).
No. 165 Dibromoacetylene C_2Br_2
 Symmetry D_{2h}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (C_6H_6 soln.)	
σ_g^+	ν_1	CC stretch.	2185 D	ia	2185 VS, p	
	ν_2	CBr stretch.	267 D	ia	267 M, p	
σ_u^+	ν_3	CBr stretch.	832 C	832 VS	ia	
π_g	ν_4	CCBr deform.	311 D	ia	311 VS, dp	
π_u	ν_5	CCBr deform.	137 C	137 S	ia	

ReferenceSee No. 164(C_2Cl_2).
No. 166 Diiodoacetylene C_2I_2
 Symmetry D_{2h}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1} (C_6H_6 soln.)	
σ_g^+	ν_1	CC stretch.	2118 D	ia	2118 VS, p	
	ν_2	CI stretch.	190 D	ia	190 S, p	
σ_u^+	ν_3	CI stretch.	720 D	720 VS (CS_2 soln.)	ia	
π_g	ν_4	CCI deform.	296 D	ia	296 VS, dp	
π_u	ν_5	CCI deform.	132 D	132 M (C_6H_6 soln.)	ia	

References[1] IR.R. A. G. Meister and F. F. Cleveland, *J. Chem. Phys.* **17**, 212 (1949).[2] IR.R. P. Klaboe and E. Kloster-Jensen, *Spectrochim. Acta* **26A**, 1567 (1970).

No. 167 Bromochloroacetylene C₂BrClSymmetry C_{2v}Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (C ₆ H ₆ soln.)	
σ^+	ν_1	CC stretch.	2223 C	2223 VS	2205 M, p	
	ν_2	CCl stretch.	923 C	923 VS	917 VS	
	ν_3	CBr stretch.	389 C	389 VW	388 S, p	
					(CCl ₄ soln.)	
π	ν_4	CCCl deform.	326 D		326 VS, dp	
					(CCl ₄ soln.)	
	ν_5	CCBr deform.	152 C	152 S	165 M	

ReferenceSee No. 164(C₂Cl₂).**No. 168 Chloriodoacetylene C₂ClI**Symmetry C_{2v}Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (C ₆ H ₆ soln.)	
σ^+	ν_1	CC stretch.	2191 C	2191 VS	2178 VS, p	
	ν_2	CCl stretch.	886 C	886 VS	880 VW	
	ν_3	CI stretch.	276 D		276 M, p	
π	ν_4	CCCl deform.	325 D		325 VS, dp	
	ν_5	CCI deform.	135 C	135 S	145 VW	
					(CCl ₄ soln.)	

ReferenceSee No. 167(C₂ClBr).

No. 169 Bromoiodoacetylene C_2BrI
 Symmetry C_{2v}
Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (CS_2 soln.)	
σ^+	ν_1	CC stretch.	2166 B	2166 S	2153 VS	
	ν_2	CBr stretch.	782 C	782 M	775 VW	
	ν_3	CI stretch.	222 D		222 M, p	
π	ν_4	CBr bend.	304 D		304 VS, dp	
	ν_5	CI bend.	122 B	122 S		

Reference

- [1] IR.R. D. H. Christensen, T. Stroger-Hansen, P. Klaboe, E. Kloster-Jensen, and E. E. Tucker, Spectrochim. Acta **28A**, 939 (1972).

No. 170 Trifluoroacetonitrile CF_3CN
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_1	ν_1	CN stretch.	2275 B	2275.0	2274 S, p	
	ν_2	CF_3 s-stretch.	1227 B	1227.2	1222 VW	
	ν_3	CC stretch.	802 B	801.7	818 M, p	
	ν_4	CF_3 s-deform.	522 B	521.8	521 M, p	
e	ν_5	CF_3 d-stretch.	1214 B	1214.3	1192 W	
	ν_6	CF_3 d-deform.	618 B	618.3	620 W, dp	
	ν_7	CF_3 rock.	463 B	462.7	463 W, dp	
	ν_8	CCN deform.	196 B	196.0	192 S, dp	

References

- [1] IR.R. W. F. Edgell and R. M. Potter, J. Chem. Phys. **24**, 80 (1956).
 [2] IR. J. A. Faniran and H. F. Shurvell, Spectrochim. Acta **26A**, 1459 (1970).
 [3] IR.Th. J. A. Faniran and H. F. Shurvell, Spectrochim. Acta **27A**, 1945 (1971).

No. 171 *trans*-1,2-Difluoroethylene CHFCHFSymmetry C_{2h} Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a_g	ν_1	CH stretch.	3111 C	ia	3111 VS, p	
	ν_2	CC stretch.	1694 C	ia	1694 VS, p	
	ν_3	CH bend.	1286 C	ia	1286 S, p	
	ν_4	CF stretch.	1123 C	ia	1123 M, p	
	ν_5	CCF deform.	548 C	ia	548 S, p	
a_u	ν_6	CH bend.	875 B	875 S	ia	
	ν_7	Torsion	329 D	333 M (Xe Matrix) 325 M	ia	
				(Xe Matrix)		
b_g	ν_8	CH bend.	788 C	ia	788 S, dp	
b_u	ν_9	CH stretch.	3114 C	3114 M	ia	
	ν_{10}	CH bend.	1274 C	1274 M	ia	
	ν_{11}	CF stretch.	1159 C	1159 VS	ia	
	ν_{12}	CCF deform.	341 D	341 M (Xe Matrix)	ia	

Reference

[1] IR.R. N. C. Craig and J. Overend, J. Chem. Phys. **51**, 1127 (1969).No. 172 *trans*-1,2-Difluoroethylene-d₁ CHFCD_FSymmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a'	ν_1	CH stretch.	3110 C	3110 M	3112 S, p	
	ν_2	CD stretch.	2335 C	2335 M	2333 M, p	
	ν_3	CC stretch.	1674 C		1674 S, p	
	ν_4	CH bend.	1274 C	1274 M	1274 M, p	
	ν_5	CF stretch.	1166 C	1166 VS	1155 VW	
	ν_6	CF stretch.	1138 C	1138 M	1119 M, p	
	ν_7	CD bend.	940 C	940 M	941 M, dp	
	ν_8	CCF deform.	542 C		542 S, p	
	ν_9	CCF deform.	332 D	332 M		
a''	ν_{10}	CH bend.	828 B	828 S	829 M, dp	
	ν_{11}	CD bend.	673 B	673 M	673 M, dp	
	ν_{12}	Torsion	316 C	316 S		

Reference

See No. 171(CHFCHF).

No. 173 **trans-1,2-Difluoroethylene-d₂** CDFCDFSymmetry C_{2h}Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a _g	ν_1	CD stretch.	2355 C	ia	2355 S, p	
	ν_2	CC stretch.	1642 C	ia	1642 VS, p	
	ν_3	CF stretch.	1109 C	ia	1109 S, p	
	ν_4	CD bend.	935 C	ia	935 M, dp	
	ν_5	CCF deform.	538 C	ia	538 S, p	
a _u	ν_6	CD bend.	651 B	651 S	ia	
	ν_7	Torsion	309 C	309 S	ia	
b _g	ν_8	CD bend.	685 C	ia	685 S, dp	
b _u	ν_9	CD stretch.	2312 C	2312 M	ia	
	ν_{10}	CF stretch.	1173 C	1173 VS	ia	
	ν_{11}	CD bend.	942 C	942 M	ia	
	ν_{12}	CCF deform.	324 D	324 M	ia	

References

- [1] R. N. C. Craig and J. Overend, *Spectrochim. Acta* **20**, 1561 (1964).
 [2] IR.R. N. C. Craig and J. Overend, *J. Chem. Phys.* **51**, 1127 (1969).

No. 174 **Glyoxal** C₂H₂O₂Symmetry C_{2h}Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)		
a _g	ν_1	CH stretch.	2843 B	ia		EL[4].
	ν_2	CO stretch.	1745 B	ia		EL[4].
	ν_3	CH bend.	1338 D	ia		OC[1].
	ν_4	CC stretch.	1065 B	ia		EL[4].
	ν_5	CCO deform.	551 B	ia		EL[3].
a _u	ν_6	CH bend.	801 B	801.36 M		
	ν_7	Torsion	127 B	126.5 W		EL[3].
b _g	ν_8	CH bend.	1048 B	ia		[5].
b _u	ν_9	CH stretch.	2835 B	2835.07 VS		
	ν_{10}	CO stretch.	1732 C	1732 VS		
	ν_{11}	CH bend.	1312 B	1312.38 S		
	ν_{12}	CCO deform.	339 B	338.55 S		

References

- [1] IR.R. R. K. Harris, *Spectrochim. Acta* **20**, 1129 (1964).
 [2] IR. A. R. H. Cole and G. A. Osborne, *J. Mol. Spectry.* **36**, 276 (1970).
 [3] EL. F. W. Birss, J. M. Brown, A. R. H. Cole, A. Loftus, S. L. N. G. Krishnamachari, G. A. Osborne, J. Puldus, D. A. Ramsay, and L. Watmann, *Can. J. Phys.* **48**, 1230 (1970).
 [4] EL. W. Holzer and D. A. Ramsay, *Can. J. Phys.* **48**, 1759 (1970).
 [5] IR. A. R. H. Cole and G. A. Osborne, *Spectrochim. Acta* **27A**, 2461 (1971).

No. 175 Glyoxal-d₁ C₂HDO₂
 Symmetry C_s
Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a'	ν_1	CH stretch.	2835 B	3835.2 S		
	ν_2	CD stretch.	2130 B	2130.2 S		
	ν_3	CO s-stretch.	1735 D	1735 M		
	ν_4	CO a-stretch.	1717 C	1717 VS		
	ν_5	CH bend.	1335 B	1335.0 W		
	ν_6	CC stretch.	1104 E			CF[1].
	ν_7	CD bend.	972 B	971.6 M		
	ν_8	CCO s-deform.	542 D	542 ^a		EL[2].
	ν_9	CCO a-deform.	323 B	323.7 S		
a''	ν_{10}	CH bend.	999 D	999 W		
	ν_{11}	CO bend.	688 D			CF. ^b
	ν_{12}	Torsion	124 B	123.9 W		

^a From the analyses of electronic transitions.^b From the product rule.**References**

- [1] Th. T. Fukuyama, Ph. D. Thesis (University of Tokyo, 1970).
 [2] IR. A. R. H. Cole and G. A. Osborne, *Spectrochim. Acta* **27A**, 2461 (1971).

No. 176 Glyoxal-d₂ C₂D₂O₂
 Symmetry C_{2h}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
a _g	ν_1	CD stretch.	2138 D	ia		OC[5].
	ν_2	CO stretch.	1722 C	ia		OC[5].
	ν_3	CD bend.	1130 D	ia		OC[5].
	ν_4	CC stretch.	915 E	ia		CF[3].
	ν_5	CCO deform.	537 B	ia		EL[2].
a _u	ν_6	CD bend.	630 D			CF[1]. ^a
	ν_7	Torsion	118 B	118.2 W		EL[2].
b _g	ν_8	CD bend.	911 D	ia		CF[1]. ^a
b _u	ν_9	CD stretch.	2130 C	2130 S		
	ν_{10}	CO stretch.	1710 C	1710 VS		
	ν_{11}	CH bend.	1010 B	1010.12 M		
	ν_{12}	CCO deform.	311 B	311.05 M		

^a From the product rule**References**

- [1] IR. J. C. D. Brand and G. J. Minkoff, *J. Chem. Soc.* **1954**, 2970.
 [2] EL. J. C. D. Brand, *Trans. Faraday Soc.* **50**, 431 (1954).
 [3] Th. T. Fukuyama, K. Kuchitsu, and Y. Morino, *Bull. Chem. Soc. Japan*, **41**, 3019 (1968).
 [4] EL. D. M. Agar, E. J. Bair, F. W. Birss, P. Bovrell, P. C. Chen, C. N. Gurrie, A. J. Mchugh, B. J. Orr, D. A. Ramsay, and J. Y. Roncin, *Can. J. Phys.* **49**, 323 (1971).
 [5] IR. A. R. H. Cole and G. A. Osborne, *Spectrochim. Acta* **27A**, 2461 (1971).

No. 177		Acetyl fluoride	CH ₃ COF	Symmetry number $\sigma = 1$		
Symmetry C _s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a'	ν_1	CH ₃ d-stretch.	3043 C	3043 W	3043 VW	
	ν_2	CH ₃ s-stretch.	2955 C	2955 VW	2952 S	
	ν_3	C=O stretch.	1870 C	1870 VS	1847 S	
	ν_4	CH ₃ d-deform.	1440 C	1440 W	1440 W	
	ν_5	CH ₃ s-deform.	1378 C	1378 M	1379 W	
	ν_6	CF stretch.	1188 C	1188 VS	1178 VW	
	ν_7	CH ₃ rock.	1000 C	1000 M	1003 M	
	ν_8	CC stretch.	826 C	826 S	822 S	
	ν_9	OCF deform.	598 C	598 W	602 M	
	ν_{10}	OCF deform.	420 D	420 VW	428 W	
a''	ν_{11}	CH ₃ d-stretch.	3004 C	3004 W	3004 VW	
	ν_{12}	CH ₃ d-deform.	1437 D	1437 W	1440 W	
	ν_{13}	CH ₃ rock.	1054 C	1054 M		
	ν_{14}	C=O op-bend.	567 C	567 W	573 W	
	ν_{15}	CH ₃ torsion	123 E			CF[1].

References

- [1] MW. L. Pierce and L. C. Krisher, *J. Chem. Phys.* **31**, 875 (1959).
 [2] R. H. Seewan-Albert and L. Kahovec, *Acta Phys. Austriaca* **1**, 352 (1948).
 [3] IR. J. A. Ramsey and J. A. Ladd, *J. Chem. Soc.* **B1968**, 118.
 [4] IR.R. C. V. Berney and A. D. Cormier, *Spectrochim. Acta* **28A**, 1813 (1972).

No. 178		Acetyl fluoride-d ₃	CD ₃ COF	Symmetry number $\sigma = 1$		
Symmetry C _s						
Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a'	ν_1	CD ₃ d-stretch.	2274 D	2286 M	2272 W	
	ν_2	CD ₃ s-stretch.	2144 C	2144 W	2140 S	
	ν_3	C=O stretch.	1869 C	1869 VS	1849 M	
	ν_4	CF stretch.	1204 C	1204 VS	1196 VW	
	ν_5	CD ₃ s-deform.	1149 C	1149 W		
	ν_6	CD ₃ d-deform.	1030 C	1030 S	1030 W	
	ν_7	CD ₃ rock.	839 C	839 M	845 M	
	ν_8	CC stretch.	778 C	778 S	774 M	
	ν_9	OCF deform.	575 C	575 M	578 M	
	ν_{10}	OCF deform.	395 D	395 VW	378 W	
a''	ν_{11}	CD ₃ d-stretch.	2242 C	2242 W	2250 W	
	ν_{12}	CD ₃ d-deform.	1057 C	1057 M	1053 W	
	ν_{15}	CD ₃ rock.	915 C	915 M		
	ν_{14}	C=O op-bend.	491 C	491 M	495 W	
	ν_{15}	CD ₃ torsion	93 E			CF[1].

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- [1] MW. L. Pierce and L. C. Krisher, *J. Chem. Phys.* **31**, 875 (1959).
 [2] IR. C. V. Berney, R. L. Redington and K. C. Lin, *J. Chem. Phys.* **53**, 1713 (1970).
 [3] IR.R. C. V. Berney and A. D. Cormier, *Spectrochim. Acta* **28A**, 1813 (1972).

No. 179 Dimethylzinc CH_3ZnCH_3 Symmetry $D_{3h}(G^+_{36})^a$ Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
a_1'	ν_1	CH_3 s-stretch.	2900 C	ia	2900 S, p	
	ν_2	CH_3 s-deform.	1157 C	ia	1157 S, p	
	ν_3	CZn s-stretch.	503 C	ia	503 VS, p	
a_1''	ν_4	Torsion		ia	ia	
a_2''	ν_5	CH_3 s-stretch.	2915 C	2915 S	ia	
	ν_6	CH_3 s-deform.	1183 C	1183 M	ia	
e'	ν_7	CZn a-stretch.	613 C	613	ia	
	ν_8	CH_3 d stretch.	2966 C	2966 S	2947	
	ν_9	CH_3 d-deform.	1301 C	1301 M	1302 W	
	ν_{10}	CH_3 rock.	704 C	704 S		
e''	ν_{11}	CZnC deform.	134 D		134 sh	
	ν_{12}	CH_3 d-stretch.	2843 C	2843	2830	
	ν_{13}	CH_3 d-deform.	1434 C		1434 W	
	ν_{14}	CH_3 rock.	620 C		620 M	

^a Free rotation [3,5].

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- [1] IR.R. H. S. Gutowsky, J. Chem. Phys. **17**, 128 (1949).
 [2] Th.IR. D. R. J. Boyd, R. L. Williams, and H. W. Thompson, Nature **167**, 766 (1951).
 [3] IR.R. J.-L. Bribes, Ph. D. Thesis, (Universite des Sciences et Techniques du Languedoc 1971).
 [4] Th. A. M. W. Bakke, J. Mol. Spectry. **41**, 1 (1972).
 [5] IR.R. J. R. Durig and S. C. Brown, J. Mol. Spectry. **45**, 338 (1973).

No. 180 Dimethylzinc- d_6 CD_3ZnCD_3 Symmetry $D_{3h}(G^+_{36})^a$ Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
a_1'	ν_1	CD_3 s-stretch.	2109 C	ia	2109 S, p	
	ν_2	CD_3 s-deform.	898 C	ia	898 S, p	
	ν_3	CZn s-stretch.	458 C	ia	458 S, p	
a_1''	ν_4	Torsion		ia	ia	
a_2''	ν_5	CD_3 s-stretch.	2117 C	2117 M	ia	
	ν_6	CD_3 s-deform.	934 C	934 M	ia	
e'	ν_7	CZn a-stretch.	554 C	554 S	ia	
	ν_8	CD_3 d-stretch.	2219 C	2219 M	2206 M	
	ν_9	CD_3 d-deform.	960 C	960 W		
	ν_{10}	CD_3 rock.	585 D	585 sh		
e''	ν_{11}	CZnC deform.	96 D		96 sh	
	ν_{12}	CD_3 d-stretch.	2073 C	2073 W	2075 sh	
	ν_{13}	CD_3 d-deform.	1006 C	1006 W		
	ν_{14}	CD_3 rock.	495 C		495 W, p	

^a Free rotation [1].

Reference

- [1] IR.R. J. R. Durig and S. C. Brown, J. Mol. Spectry. **45**, 338 (1973).

No. 181 Dimethylcadmium CH_3CdCH_3 Symmetry $D_{3h}(G_{36})^a$ Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
a_1'	ν_1	CH_3 s-stretch.	2903 C	ia	2903 S, p	
	ν_2	CH_3 s-deform.	1127 C	ia	1127 S, p	
	ν_3	CCd s-stretch.	459 C	ia	459 S, p	
a_1''	ν_4	Torsion		ia	ia	
a_2''	ν_5	CH_3 s-stretch.	2923 C	2923 S	ia	
	ν_6	CH_3 s-deform.	1136 C	1136 M	ia	
	ν_7	CCd a-stretch.	535 C	535 S	ia	
e'	ν_8	CH_3 d-stretch.	2980 C	2980 VS		
	ν_9	CH_3 d-deform.	1315 C	1315	1324	
	ν_{10}	CH_3 rock.	700 C	700 S		
	ν_{11}	CCdC deform.	124 C	124 W	120 W, b	
e''	ν_{12}	CH_3 d-stretch.	2859 C	2859	2834 M	
	ν_{13}	CH_3 d-deform.	1427 C		1427 W	
	ν_{14}	CH_3 rock.	634 C		634 M, dp	

^a Free rotation [3,5].

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No. 182 Dimethylcadmium- d_6 CD_3CdCD_3 Symmetry $D_{3h}(G_{36})^a$ Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
a_1'	ν_1	CD_3 s-stretch.	2112 C	ia	2112 S, p	
	ν_2	CD_3 s-deform.	873 C	ia	873 S, p	
	ν_3	CCd s-stretch.	419 C	ia	419 S, p	
a_1''	ν_4	Torsion		ia	ia	
a_2''	ν_5	CD_3 s-stretch.	2120 C	2120 S	ia	
	ν_6	CD_3 s-deform.	900 C	900 W	ia	
	ν_7	CCd a-stretch.	492 C	492 S	ia	
e'	ν_8	CD_3 d-stretch.	2229 C	2229 S	2214 M, dp	
	ν_9	CD_3 d-deform.	1047 D		1047 W	
	ν_{10}	CD_3 rock.	538 C	538 S		
	ν_{11}	CCdC deform.	109 D		109 sh	
e''	ν_{12}	CD_3 d-stretch.	2080 D	2080	2086 sh	
	ν_{13}	CD_3 d-deform.	1126 C		1126	
	ν_{14}	CD_3 rock.	476 C		476 M, dp	

^a Free rotation [1].

Reference

See No. 180(CD_3ZnCD_3).

No. 183 Dimethylmercury CH_3HgCH_3 Symmetry $D_{3h}(G^+_{3h})^a$ Symmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1}	cm^{-1}	
				(Gas)	(Liquid)	
a_1'	ν_1	CH_3 s-stretch.	2911 C	ia	2911 VS, p	
	ν_2	CH_3 s-deform.	1182 C	ia	1182 VS, p	
	ν_3	CHg s-stretch.	515 C	ia	515 VS, p	
a_1''	ν_4	Torsion		ia	ia	
a_2''	ν_5	CH_3 s-stretch.	2925 D	2925 b	ia	
	ν_6	CH_3 s-deform.	1191 C	1191 M	ia	
	ν_7	CHg a-stretch.	540 C	540 VS	ia	
e'	ν_8	CH_3 d-stretch.	2962 C	2962 S		
	ν_9	CH_3 d-deform.	1397 C		1397 W, dp	
	ν_{10}	CH_3 rock.	780 C	780 VS	779 VW, dp	
	ν_{11}	CHgC deform.	161 C		161 M, dp	
e''	ν_{12}	CH_3 d-stretch.	2874 C		2874 S, p	
	ν_{13}	CH_3 d-deform.	1442 C		1442 W, dp	
	ν_{14}	CH_3 rock.	699 C	700 sh	699 M, dp	

^a Free rotation [3,6].

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No. 184 Dimethylmercury-d₆ CD₃HgCD₃Symmetry D_{3h}(G⁺₃₆)^aSymmetry number $\sigma = 6$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)	
a ₁ '	ν_1	CD ₃ s-stretch.	2111 C	ia	2111 M, p	
	ν_2	CD ₃ s-deform.	909 C	ia	909 VS, p	
	ν_3	CHg s-stretch.	471 C	ia	471	
a ₁ "	ν_4	Torsion		ia	ia	
a ₂ "	ν_5	CD ₃ s-stretch.	2114 C	2114 S	ia	
	ν_6	CD ₃ s-deform.	931 C	931 M	ia	
	ν_7	CHg a-stretch.	491 C	491 VS	ia	
e'	ν_8	CD ₃ d-stretch.	2224 C	2224	2224 S, dp	
	ν_9	CD ₃ d-deform.	1030 C	1030 M	1030 W, dp	
	ν_{10}	CD ₃ rock.	598 C	598 VS		
	ν_{11}	CHgC deform.	141 C		141 S, dp	
e''	ν_{12}	CD ₃ d-stretch.	2039 C	2039	2044 M, p	
	ν_{13}	CD ₃ d-deform.	1050 C		1050 VW, p	
	ν_{14}	CD ₃ rock.	525 C		525 M, sh, dp	

^a Free rotation [2,4].

References

- [1] IR.R. J.-L. Briber et R. Gaufres, *J. Chim. Phys.* **67**, 1168 (1970).
 [2] Th. J.-L. Briber et R. Gaufres, *J. Mol. Structure* **9**, 423 (1971).
 [3] Th. A. M. W. Bakke, *J. Mol. Spectry.* **41**, 1 (1972).
 [4] Th. J. R. Durig and S. C. Brown, *J. Mol. Spectry.* **45**, 338 (1973).

No. 185 Azomethane CH_3NNCH_3
Symmetry C_{2h}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_g	ν_1	CH_3 d-stretch.	2977 C	ia	2977 M, dp	
	ν_2	CH_3 s-stretch.	2916 C	ia	2916 S, p	
	ν_3	NN stretch.	1580 C	ia	1580 VW, p	
	ν_4	CH_3 d-deform.	1434 C	ia	1434 M	
	ν_5	CH_3 s-deform.	1380 C	ia	1380 W	
	ν_6	CN stretch.	1176 C	ia	1176 VW	
	ν_7	CH_3 rock.	916 C	ia	916 M	
	ν_8	CCN bend.	589 C	ia	589 VS, p	
a_u	ν_9	CH_3 d-stretch.	2966 D	2966 W, sh (solid)	ia	
	ν_{10}	CH_3 d-deform.	1438 C	1438	ia	
	ν_{11}	CH_3 rock.	1109 C	1109 W	ia	
	ν_{12}	CCN bend.	312 D	312 S (solid)	ia	
	ν_{13}	CH_3 torsion	222 D	222 VW	ia	
b_g	ν_{14}	CH_3 d-stretch.	2982 D	ia	2982 VS (solid)	
	ν_{15}	CH_3 d-deform.	1447 D	ia	1447 S (solid)	
	ν_{16}	CH_3 rock.	1010 D	ia	1010 VW	
	ν_{17}	CH_3 torsion	223 D	ia	223 W (solid)	
b_u	ν_{18}	CH_3 d-stretch.	2982 C	2982 VS	ia	
	ν_{19}	CH_3 s-stretch.	2926 C	2926 VS	ia	
	ν_{20}	CH_3 d-deform.	1445 C	1445 S	ia	
	ν_{21}	CH_3 s-deform.	1393 C	1393 M	ia	
	ν_{22}	CN stretch.	1300 C	1300 VW	ia	
	ν_{23}	CH_3 rock.	1009 C	1009 W	ia	
	ν_{24}	CCN bend.	352 C	352 M	ia	

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 [2] R.Th. R. A. R. Pearce, I. W. Lewin, and W. C. Harris, J. Chem. Phys. **59**, 1209 (1973).

No. 186 Azomethane-d₆ CD₃NNCD₃
 Symmetry C_{2h}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹	cm ⁻¹		
				(Gas)	(Liquid)		
<i>a_g</i>	ν_1	CD ₃ d-stretch.	2227 C	ia	2227 M		
	ν_2	CD ₃ s-stretch.	2117 C	ia	2117 S, p		
	ν_3	NN stretch.	1563 C	ia	1563 VW, dp		
	ν_4	CN stretch.	1125 C	ia	1125 W, p		
	ν_5	CD ₃ d-deform.	1042 C	ia	1042 M		
	ν_6	CD ₃ s-deform.	1035 D	ia	1035		
					(solid)		
		ν_7	CD ₃ rock.	764 C	ia	764 M, p	
<i>a_u</i>		ν_8	CNN bend.	524 C	ia	524 VS, p	
		ν_9	CD ₃ d-stretch.	2178 C	2178 VW	ia	
		ν_{10}	CD ₃ d-deform.	1043 D	1043 S	ia	
					(solid)		
		ν_{11}	CD ₃ rock.	894 D	894 S, b	ia	OV(ν_{23}).
		ν_{12}	CCN bend.	273 C	273 M	ia	
		ν_{13}	CD ₃ torsion	166 D	166 VW	ia	
<i>b_g</i>				(solid)			
		ν_{14}	CD ₃ d-stretch.	2250 C	ia	2250 M	
		ν_{15}	CD ₃ d-deform.	1057 C	ia	1057 M	
					(solid)		
		ν_{16}	CD ₃ rock.	792 D	ia	792 W	
<i>b_u</i>		ν_{17}	CD ₃ torsion	183 D	ia	183 W	
					(solid)		
		ν_{18}	CD ₃ d-stretch.	2239 C	2239 VS	ia	
		ν_{19}	CD ₃ s-stretch.	2111 C	2111 M	ia	
		ν_{20}	CN stretch.	1116 D	1116 VW	ia	
					(solid)		
		ν_{21}	CD ₃ d-deform.	1047 C	1047 M	ia	
		ν_{22}	CD ₃ s-deform.	1029 C	1029 sh	ia	
	ν_{23}	CD ₃ rock.	894 D	894 S, b	ia	OV(ν_{11}).	
	ν_{24}	CNN bend.	304 C	304 M	ia		

References

See No. 185(CH₃NNCH₃)

No. 187 Carbon suboxide C₃O₂
 Symmetry D_{∞h}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Gas)	
σ_g^+	ν_1	CO s-stretch.	2196 B	ia	2196.5 W	
	ν_2	CC s-stretch.	786 B	ia	786.1 W, p	
σ_u^+	ν_3	CO a-stretch.	2258 B	2258 S		
	ν_4	CC a-stretch.	1573 B	1573 S		
π_g	ν_5	CCO bend.	573 C	ia	573.0 W	
π_u	ν_6	CCO bend.	550 B	550 S		
	ν_7	CCC bend.	61 B	61		

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No. 188 Carbon subsulfide C₃S₂
 Symmetry D_{∞h}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (CS ₂ soln.)	
σ_g^+	ν_1	CC s-stretch.	1663 D	ia	1663 S	
	ν_2	CS s-stretch.	485 D	ia	485 S, p	
σ_u^+	ν_3	CC a-stretch.	2089 B	2088.5 VS	ia	
	ν_4	CS a-stretch.	1030 B	1029.8 S	ia	
π_g	ν_5	CCS bend.	470 D	ia	470 M	
π_u	ν_6	CCS bend.	502 C	502 VW	ia	
	ν_7	CCC bend.	94 C	94 W	ia	

References

- [1] IR.R. W. H. Smith and G. E. Leroi, J. Chem. Phys. **45**, 1778 (1966).
 [2] Th. W. H. Smith and G. E. Leroi, J. Chem. Phys. **45**, 1784 (1966).
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No. 189 Chloro cyanoacetylene ClCCCN
Symmetry C_{2v}

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (C_6H_6 soln.)	
σ^+	ν_1	CN stretch.	2297 C	2297 VS	2290 VS, p	
	ν_2	$C\equiv C$ stretch.	2194 B	2194 W	2196 M, p	
	ν_3	C-C stretch.	1093 C	1093 S	1103 W, p	
	ν_4	C-Cl stretch.	527 B	527 M	530 M, p	
π	ν_5	CCN bend.	483 B	483 S	487 S	
	ν_6	CCC bend.	333 C	333 M	338 M	
	ν_7	CCCl bend.	145 D		145 VW	

References

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 [2] IR.R. P. Klaboe and E. Kloster-Jensen, *Spectrochim. Acta* **23A**, 1981 (1967).

No. 190 Bromo cyanoacetylene BrCCCN
Symmetry C_{2v}

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (C_6H_6 soln.)	
σ^+	ν_1	CN stretch.	2292 C	2292 VS	2276 VS, p	
	ν_2	$C\equiv C$ stretch.	2150 C	2150 S	2123 W	
	ν_3	C-C stretch.	1054 B	1054 VW	1052 W	
	ν_4	CBr stretch.	419 B	419 W	418 M, p	
π	ν_5	CCN bend.	485 B	485 S	489 S	
	ν_6	CCC bend.	312 B	312 M	317 S	
	ν_7	CCBr bend.	142 D		142 VW	

References

See No. 189(ClCCCN).

No. 191 Iodo cyanoacetylene ICCCN

Symmetry C_{2v} Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (C_6H_6 soln.)	
σ^+	ν_1	CN stretch.	2270 C	2270 S	2265 VS, p	
	ν_2	$C\equiv C$ stretch.	2131 C	2131 M	2125 M, p	
	ν_3	C-C stretch.	1031 C	1031 S	1034 W, p	
	ν_4	CI stretch.	364 B	364 W	356 M	
π	ν_5	CCN bend.	496 B	496 M	497 S, dp	
	ν_6	CCC bend.	309 C	309 W	315 M, dp	
	ν_7	CCI bend.	130 D		130 VW	

References

See No. 189(CICCCN).

No. 192 Chloropropadiene CH_2CCHCl Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a'	ν_1	CH stretch.	3079 C	3079 S	3062 S, p	OV(ν_{11}).
	ν_2	CH_2 s-stretch.	3009 C	3009 M	2990 S, p	
	ν_3	CCC a-stretch.	1963 C	1963 M	1951 W, dp	
	ν_4	CH_2 scis.	1435 C	1435 S	1419 S, p	
	ν_5	CH bend.	1256 C	1256 VS	1244 W, p	
	ν_6	CCC s-stretch.	1101 C	1101 S	1095 VS, p	
	ν_7	CH_2 wag.	875 C	875 VS	877 W, dp	
	ν_8	CCl stretch.	767 C	767 VS	751 W, dp	
	ν_9	CCC deform.	592 C	592 W		
	ν_{10}	CCCl deform.	494 C	494 M	490 VS, p	
a''	ν_{11}	CH_2 a-stretch.	3079 C	3079 S	3062 S, p	OV(ν_{11}).
	ν_{12}	CH_2 rock.	999 B	999 W	995 W, dp	
	ν_{13}	CH bend.	822 B	822 S	815 W, dp	
	ν_{14}	CCC deform.	548 B	548 M	546 W, dp	
	ν_{15}	CCCl deform.	184 C		184 S, dp	

Reference

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No. 193 Bromopropadiene CH₂CCHBr
 Symmetry C_s
Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a'	ν_1	CH stretch.	3080 C	3080 M	3060 (46), p	OV(ν_{11}).
	ν_2	CH ₂ s-stretch.	3005 C	3005 M	2985 (63), p	
	ν_3	CCC a-stretch.	1961 C	1961 W	1954 (19), p	
	ν_4	CH ₂ scis.	1432 B	1432 S	1422	
	ν_5	CH bend.	1217 C	1217 VS	1209 (12)	
	ν_6	CCC s-stretch.	1078 C	1078 W	1086 (29), p	
	ν_7	CH ₂ wag.	862 C	862 VS	873 (3), dp	
	ν_8	CBr stretch.	681 C	681 VS	667 (31), p	
	ν_9	CCC deform.	603 C	603 W		
	ν_{10}	CCBr deform.	423 C	423 VW	426 (33), p	
				(CS ₂ soln.)		
a''	ν_{11}	CH ₂ a-stretch.	3080 C	3080 M	3060 (46), p	OV(ν_{11}).
	ν_{12}	CH ₂ rock.	1000 B	1000 W		
	ν_{13}	CH bend.	812 B	812 S	806 (3), dp	
	ν_{14}	CCC deform.	519 B	519 M		
	ν_{15}	CCCl deform.	169 C		169 (43b), dp	

ReferenceSee No. 192(CH₂CCHCl).
No. 194 Iodopropadiene CH₂CCHI
 Symmetry C_s
Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a'	ν_1	CH stretch.	3070 C	3070 M	3057 M	OV(ν_{11}).
	ν_3	CCC a-stretch.	1953 C	1953 M	1947 VS	
	ν_2	CH ₂ s-stretch.	3004 C	3004 W	2978 VS	
	ν_4	CH ₂ scis.	1425 B	1425 M	1412 VS	
	ν_5	CH bend.	1178 C	1178 VS	1174 S, p	
	ν_6	CCC s-stretch.	1076 C	1076 M	1076 VS, p	
	ν_7	CH ₂ wag.	854 C	854 S		
	ν_8	CCC deform.	625 C	625 S	635 W	
	ν_9	CI stretch.	609 C	609 S		
	ν_{10}	CCI deform.	387 C		387 S, p	
a''	ν_{11}	CH ₂ a-stretch.	3070 C	3070 M	3057 M	OV(ν_{11}).
	ν_{12}	CH ₂ rock.	995 B	995 W		
	ν_{13}	CH bend.	807 B	807 S		
	ν_{14}	CCC deform.	485 B	485 W		
	ν_{15}	CCI deform.	154 C		154 W	

ReferenceSee No. 192(CH₂CCHCl).

No. 195 Trifluoropropyne CF₃CCH
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)	(Liquid)	
a ₁	ν_1	CH stretch.	3327 B	3327.3 S	3316 (9) p	
	ν_2	C \equiv C stretch.	2165 C	2165.4 S	2156 (80) p	FR($\nu_4 + \nu_6 + \nu_{10}$).
	ν_3	CF ₃ s-stretch.	1253 B	1253.2 VS	1250 (1b)	
	ν_4	C-C stretch.	812 B	811.9 W	810 (50) p	
	ν_5	CF ₃ s-deform.	536 B	536.1 M	537 (11) p	
e	ν_6	CF ₃ d-stretch.	1179 B	1179.2 VS	1155 (5b) dp	
	ν_7	CH bend.	686 B	685.5 S	696 (6) dp	
	ν_8	CF ₃ d-deform.	612 B	611.9 M	606 (4) dp	
	ν_9	CF ₃ rock.	453 B	453.0 M	445 (6) dp	
	ν_{10}	CCC bend.	171 B	171 M	170 (100) dp	

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No. 196 Trifluoropropyne-d CF₃CCD
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹	cm ⁻¹	
				(Gas)		
a ₁	ν_1	CD stretch.	2626 B	2626.0 M		
	ν_2	C \equiv C stretch.	2014 B	2013.9 S		
	ν_3	CF ₃ s-stretch.	1250 B	1249.7 VS		
	ν_4	C-C stretch.	808 B	808.4 W		
	ν_5	CF ₃ s-deform.	529 B	528.9 M		
e	ν_6	CF ₃ d-stretch.	1179 B	1179.0 VS		
	ν_8	CF ₃ d-deform.	611 B	611.3 M		
	ν_7	CD bend.	539 B	538.5 S		
	ν_9	CF ₃ rock.	456 B	455.5 M		
	ν_{10}	CCC bend.	163 B	163 M		

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- [1] IR.R. C. V. Berney, L. R. Cousins, and F. A. Miller, *Spectrochim. Acta* **19**, 2019 (1963).
 [2] IR. R. H. Sanborn, *Spectrochim. Acta* **23A**, 1999 (1967).

No. 197 1-Chloro-3,3,3-trifluoropropyne CF_3CCCl
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_1	ν_1	$\text{C}\equiv\text{C}$ stretch.	2270 C	2270 VS	2266 S, p	
	ν_2	CF_3 s-stretch.	1276 C	1276 VS	1272 W	
	ν_3	C-C stretch.	937 C	937 S	933 S, p	
	ν_4	CCl stretch.	723 C	723 S	723 S, p	
	ν_5	CF_3 s-deform.	377 C	377 W	374 S, p	
e	ν_6	CF_3 d-stretch.	1174 C	1174 VS	1153 W, b	
	ν_7	CF_3 d-deform.	606 C	606 M	603 W	
	ν_8	CF_3 rock.	448 D		448 M, dp	
	ν_9	CCCCl deform.	311 C	311 W	316 VS, dp	
	ν_{10}	CCCCl deform.	97 D		97 VS, dp	

Reference

[1] IR.R. E. Augdahl, E. Kloster-Jensen, V. Devarajan and S. J. Cyvin, Spectrochim. Acta **29A**, 1329 (1973).

No. 198 1-Bromo-3,3,3-trifluoropropyne CF_3CCBr
 Symmetry C_{3v}
Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_1	ν_1	$\text{C}\equiv\text{C}$ stretch.	2243 C	2243 S	2236 M, p	
	ν_2	CF_3 s-stretch.	1275 C	1275 VS	1267 W	
	ν_3	C-C stretch.	883 C	883 S	879 S, p	
	ν_4	CBr stretch.	687 C	687 S	684 M, p	
	ν_5	CF_3 s-deform.	285 D	285 M	292 VS	
e	ν_6	CF_3 d-stretch.	1176 C	1176 VS	1155 W, b	
	ν_7	CF_3 d-deform.	607 C	607 M	606 W	
	ν_8	CF_3 rock.	455 D	455 VW	452 S	
	ν_9	CCCBBr deform.	278 D	278 VW, sh		
	ν_{10}	CCCBBr deform.	90 D		90 VS, dp	

Reference

See No. 197(CF_3CCCl).

No. 199 1-Iodo-3,3,3-trifluoropropyne CF_3CCI Symmetry C_{3v} Symmetry number $\sigma = 3$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_1	ν_1	$\text{C}\equiv\text{C}$ stretch.	2218 C	2218 S	2207 S, p	
	ν_2	CF_3 s-stretch.	1264 C	1264 VS	1253 W	
	ν_3	$\text{C}-\text{C}$ stretch.	857 C	857 M	857 S, p	
	ν_4	CI stretch.	661 C	661 S	658 M	
	ν_5	CF_3 s-deform.	250 D	250 W, sh	254 VS, p	
e	ν_6	CF_3 d-stretch.	1174 C	1174 VS	1150 W, b	
	ν_7	CF_3 d-deform.	608 C	608 W	605 W	
	ν_8	CF_3 rock.	450 C	450 W	449 M, dp	
	ν_9	CCCI deform.	263 C	263 M	267 VS, dp	
	ν_{10}	CCCI deform.	85 D		85 VS, dp	

Reference

See No. 197(CF_3CCCl).No. 200 Propargyl fluoride CHCCH_2F Symmetry C_s Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a'	ν_1	CH stretch.	3338 C	3338 S	3330 (4), p	
	ν_2	CH_2 s-stretch.	2955 C	2955 M	2960 (22), p	
	ν_3	$\text{C}=\text{C}$ stretch.	2150 C	2150 W	2135 (43), p	
	ν_4	CH_2 scis.	1465 C	1465 VW	1458 (3), dp	
	ν_5	CH_2 wag.	1381 C	1381 S	1374 (3), dp	
	ν_6	CF stretch.	1039 C	1039 VS		
	ν_7	$\text{C}-\text{C}$ stretch.	940 C	940 M	935 (6), p	
	ν_8	CH bend.	675 C	675 S	692 (1), dp	
a''	ν_9	CCF deform.	539 C	539 W	544 (3), p	
	ν_{10}	CCC deform.	211 C		211 (10), dp	
	ν_{11}	CH_2 a-stretch.	2972 C	2972 M	2986 (7), dp	
	ν_{12}	CH_2 twist.	1242 D	1240 VW	1242 (1), dp	
	ν_{13}	CH_2 rock.	1018 D	1018 W	1012 (3b), p	
	ν_{14}	CH bend.	635 C	635 S	646 (1b), dp	
	ν_{15}	CCC deform.	310 C		310 (4), dp	

Reference

[1] IR.R. J. C. Evans and R. A. Nyquist, Spectrochim. Acta **19**, 1153 (1963).

No. 201 Propargyl chloride CHCCH_2Cl
 Symmetry C_s
Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a'	ν_1	CH stretch.	3335 C	3335 VS	3305 (2), p	
	ν_2	CH_2 s-stretch.	2968 C	2968 M	2957 (29), p	
	ν_3	$\text{C}\equiv\text{C}$ stretch.	2147 C	2147 W	2130 (35), p	
	ν_4	CH_2 scis.	1441 C	1441 M	1432 (2), p	
	ν_5	CH_2 wag.	1271 C	1271 VS	1267 (5), p	
	ν_6	C-C stretch.	960 C	960 S	961 (2), dp	
	ν_7	CCl stretch.	725 C	725 VS	713 (12), p	
	ν_8	CH bend.	650 D	650 S	650 (2b)	
	ν_9	CCCl deform.	451 C	451 (CS_2 soln.)	452 (5), p	
	ν_{10}	CCC deform.	186 D		186 (10), dp	
a''	ν_{11}	CH_2 a-stretch.	3002 C	3002 M	2995 (4), dp	
	ν_{12}	CH_2 twist.	1179 B	1179 W	1174 (1), dp	
	ν_{13}	CH_2 rock.	908 D		908 (1), dp	
	ν_{14}	CH bend.	637 D	637 M	650 (2b), dp	
	ν_{15}	CCC deform.	311 C		311 (4), dp	

ReferenceSee No. 200(CHCCH_2F).
No. 202 Propargyl bromide CHCCH_2Br
 Symmetry C_s
Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a'	ν_1	CH stretch.	3335 C	3335 S	3305 (2), p	
	ν_2	CH_2 s-stretch.	2976 C	2976 W	2958 (20), p	
	ν_3	$\text{C}\equiv\text{C}$ stretch.	2138 D	2138 VW	2125 (36), p	
	ν_4	CH_2 scis.	1431 C	1431 W	1425 (2), dp	
	ν_5	CH_2 wag.	1218 C	1218 S	1214 (9), p	
	ν_6	C-C stretch.	961 C	961 M	962 (3), p	
	ν_7	CH bend.	652 C	652 VS		
	ν_8	CBr stretch.	621 C	621 S	618 (22), p	
	ν_9	CCBr deform.	399 C	399 (CS_2 soln.) (soln.)	399 (9), p	
	ν_{10}	CCC deform.	168 C		168 (10), p	
a''	ν_{11}	CH_2 a-stretch.	3006 C	3006 (CS_2 soln.)	3008 (4), p	
	ν_{12}	CH_2 twist.	1152 D	1152 VW	1146 (1)	
	ν_{13}	CH_2 rock.	866 D		866 (1), dp	
	ν_{14}	CH bend.	637 C	637 S		
	ν_{15}	CCC deform.	314 C		314 (3), dp	

ReferenceSee No. 200(CHCCH_2F).

No. 203		Propargyl iodide		CHCCH ₂ I		Symmetry number $\sigma = 1$	
Symmetry C _s							
Sym. class	No.	Approximate type of mode	Selected value of frequency cm ⁻¹	Infrared cm ⁻¹	Raman cm ⁻¹	Comments	
a'	ν_1	CH stretch.	3335 C	(Gas) 3335 S	(Liquid)		
	ν_2	CH ₂ s-stretch.	2958 C	2958	2957 VS		
	ν_3	C≡C stretch.	2130 C	(CS ₂ soln.) 2130 C	2128 VS		
	ν_4	CH ₂ scis.	1423 C	(CCl ₄ soln.) 1423 W	1414		
	ν_5	CH ₂ wag.	1160 C	1160 M	1160 VS		
	ν_6	C-C stretch.	959 C	959 W	964 S		
	ν_7	CH bend.	640 C	640 S	650 VW,b	OV(ν_{14}).	
	ν_8	CI stretch.	570 C	570 W	567 VS		
	ν_9	CCI deform.	364 C		364 S		
	ν_{10}	CCC deform.	157 C		157 M		
a''	ν_{11}	CH ₂ a-stretch.	3008 D	3008	2990 b		
	ν_{12}	CH ₂ twist.	1116 D	(CS ₂ soln.) 1116			
	ν_{13}	CH ₂ rock.	810 E ^a				
	ν_{14}	CH bend.	640 D	640 S	650 VW, b	OV(ν_7).	
	ν_{15}	CCC deform.	314 C		314 VW		

^a Estimated from the corresponding frequencies of other propargyl halides.

Reference

See No. 200(CHCCH₂F).

No. 204 Chlorodiacetylene C₂ClCCCH
Symmetry C_{2v} Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency cm ⁻¹	Infrared cm ⁻¹ (Gas)	Raman cm ⁻¹	Comments
σ^-	ν_1	CH stretch.	3327 C	3327 VS		
	ν_2	C≡C a-stretch.	2252 C	2252 VS		
	ν_3	C≡C s-stretch.	2071 C	2071 M		
	ν_4	C-C stretch.	1093 C	1093 S		
	ν_5	CCl stretch.	525 C	525 S		
π	ν_6	CH bend.	621 B	621 VS		
	ν_7	CCCC a-bend.	463 B	463 S		
	ν_8	CCCC s-bend.	335 C	335 M		
	ν_9	CCCl bend.	133 E	133		OC($\nu_7 + \nu_9$, $\nu_6 + \nu_9$, $\nu_7 + \nu_9$).

Reference

[1] IR. P. Klaboe and E. Kloster-Jensen, Spectrochim. Acta **23A**, 2733 (1967).

No. 205 Bromodiacetylene BrCCCCH
Symmetry C_{2v}

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
σ^+	ν_1	CH stretch.	3335 C	3335 VS		
	ν_2	C≡C a-stretch.	2237 C	2237 S		
	ν_3	C≡C s-stretch.	2095 C	2095 W		
	ν_4	C-C stretch.	1046 C	1046 M		
	ν_5	CBr stretch.	425 C	425 M		
π	ν_6	CH bend.	623 C	623 VS		
	ν_7	CCCC a-bend.	470 B	470 S		
	ν_8	CCCC s-bend.	355 B	355 M		
	ν_9	CCBr bend.	118 E			OC($\nu_1+\nu_9, \nu_7+\nu_9$).

Reference

See No. 204(CICCCCH).

No. 206 Iododiacetylene ICCCH
Symmetry C_{2v}

Symmetry number $\sigma = 1$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹	
σ^-	ν_1	CH stretch.	3332 C	3332 VS		
	ν_2	C≡C a-stretch.	2211 C	2211 S		
	ν_3	C≡C s-stretch.	2060 C	2060 VW		
	ν_4	C-C stretch.	1025 D	1025 VW (CS ₂ soln.)		
π	ν_5	CI stretch.	362 C	362 M		
	ν_6	CH bend.	623 B	623 VS		
	ν_7	CCCC a-bend.	473 B	473 M		
	ν_8	CCCC s-bend.	357 C	357 W		
	ν_9	CCI bend.	110 E	110		OC($\nu_8+\nu_9, \nu_7+\nu_9$).

Reference

See No. 204(CICCCCH).

No. 207 1,4-Dioxane $C_4H_8O_2$
Symmetry C_{2h}

Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (Liquid)	
a_g	ν_1	CH_2 a-stretch.	2968 C	ia	2968 S, p	
	ν_2	CH_2 s-stretch.	2856 C	ia	2856 S, p	
	ν_3	CH_2 scis.	1444 C	ia	1444 M, p	
	ν_4	CH_2 wag.	1397 C	ia	1397 W, p	
	ν_5	CH_2 twist.	1305 C	ia	1305 S, p	
	ν_6	CH_2 rock.	1128 C	ia	1128 M, p	
	ν_7	CC stretch.	1015 C	ia	1015 S, p	
	ν_8	CO stretch.	837 C	ia	837 VS, p	
	ν_9	OCC deform.	435 C	ia	435 M	
	ν_{10}	COC deform.	424 C	ia	424 M, p	
a_u	ν_{11}	CH_2 a-stretch.	2970 C	2970 M	ia	
	ν_{12}	CH_2 s-stretch.	2863 C	2863 VS	ia	
	ν_{13}	CH_2 scis.	1449 C	1449 S	ia	
	ν_{14}	CH_2 wag.	1369 C	1369 S	ia	
	ν_{15}	CH_2 twist.	1256 C	1256 S	ia	
	ν_{16}	CO stretch.	1136 C	1136 VS	ia	
	ν_{17}	CH_2 rock.	1086 C	1086 S	ia	
	ν_{18}	CC stretch.	881 C	881 S	ia	
	ν_{19}	OCC deform.	288 C	288 W,sh	ia	
	b_g	ν_{20}	CH_2 a-stretch.	2968 C	ia	2968 S, p
ν_{21}		CH_2 s-stretch.	2856 C	ia	2856 S, p	SF(ν_2).
ν_{22}		CH_2 scis.	1459 C	ia	1459 M, dp	
ν_{23}		CH_2 wag.	1335 C	ia	1335 W	
ν_{24}		CH_2 twist.	1217 C	ia	1217 M, dp	
ν_{25}		CO stretch.	1110 C	ia	1110 M, dp	
ν_{26}		CH_2 rock.	853 C	ia	853 M,sh, dp	
ν_{27}		OCC deform.	490 C	ia	490 M, dp	
b_u	ν_{28}	CH_2 a-stretch.	2970 C	2970 M	ia	SF(ν_{11}).
	ν_{29}	CH_2 a-stretch.	2863 C	2863 VS	ia	SF(ν_{12}).
	ν_{30}	CH_2 scis.	1457 C	1457 S	ia	
	ν_{31}	CH_2 wag.	1378 C	1378 M	ia	
	ν_{32}	CH_2 twist.	1291 C	1291 S	ia	
	ν_{33}	CH_2 rock.	1052 C	1052 S	ia	
	ν_{34}	CO stretch.	889 C	889 S	ia	
	ν_{35}	OCC deform.	610 C	610 S	ia	
	ν_{36}	COC deform.	274 C	274 S	ia	

References

- [1] IR.R. F. E. Malherbe and H. J. Bernstein, *J. Amer. Chem. Soc.* **74**, 4408 (1952), and references cited there.
 [2] IR.Th. R. G. Snyder and G. Zerbi, *Spectrochim. Acta* **23A**, 391 (1967).
 [3] IR.R.Th. O. H. Ellested and P. Klabo, *Spectrochim. Acta* **27A**, 1025 (1971).

No. 208 Diethynyl ketone (HC≡C)₂CO
 Symmetry C_{2v}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments	
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (Liquid)		
a ₁	ν_1	CH stretch.	3325 C	3325 S	3275 W, b	OV(ν_{13}).	
	ν_2	C≡C stretch.	2097 C	2098 VS, sh (liquid)	2097 VS, p		
	ν_3	CO stretch.	1669 C	1669 S	1636 S, p		
	ν_4	C-C stretch.	739 C	739 VW	749 M, p		
	ν_5	CH bend.	648 C	648 S			
	ν_6	C-C-C deform.	571 C		571 W		
	ν_7	C=C-C deform.	122 C	122 VW			
a ₂	ν_8	CH bend.	712 C	ia	712 VW, dp		
	ν_9	C≡C-C deform.	268 C	ia	268 M		
b ₁	ν_{10}	CH bend.	729 C	729 S	733 VW (solid)		
	ν_{11}	CO op-bend.	688 C	688 S	689 VW		
	ν_{12}	C=C-C deform.	190 C	190 VW	200 W		
	b ₂	ν_{13}	CH stretch.	3325 C	3325 S	3275 VW, b	OV(ν_1).
		ν_{14}	C≡C stretch.	2115 C	2115 S	2107 M	
ν_{15}		C-C stretch.	1144 C	1144 VS			
ν_{16}		CH bend.	682 C	682 M			
	ν_{17}	CO ip-bend.	548 C	548 M			
	ν_{18}	C=C-C deform.	229 C	229 VW	239 M		

Reference[1] IR.R. F. A. Miller, B. M. Harney, and J. Tyrrell, Spectrochim. Acta **27A**, 1003 (1971).
No. 209 Dicyanodiacetylene NCCCCCN
 Symmetry D_{2h}
Symmetry number $\sigma = 2$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (CCl ₄ soln.)	
σ_g^+	ν_1	CN s-stretch.	2235 C	ia	2235 VS, p	
	ν_2	C≡C s-stretch.	2183 C	ia	2183 M, p	
	ν_3	C-C s-stretch.	1288 C	ia	1287.5 W, p	
	ν_4	C-C stretch.	571	ia	571 W, dp	
σ_u^+	ν_5	CN a-stretch.	2266 B	2266 S	ia	
	ν_6	C=C a-stretch.	2097 B	2097 M	ia	
	ν_7	C-C a-stretch.	717 B	717 S	ia	
π_g	ν_8	Bend.	501 C	ia	501 M, dp	
	ν_9	Bend.	455 C	ia	455 W, dp (C ₆ H ₁₄ soln.)	
π_u	ν_{10}	Bend.	170 D	ia	170 W, b, dp	
	ν_{11}	Bend.	491 B	490.5 VS	ia	
	ν_{12}	Bend.	276 B	276 VS	ia	
	ν_{13}	Bend.	62 B	61.5 VS	ia	

Reference[1] IR.R. F. A. Miller and D. H. Lemmon, Spectrochim. Acta **23A**, 1415 (1967).

No. 210 Hexacarbonylchromium(0) $\text{Cr}^{(12}\text{C}^{16}\text{O})_6$
Symmetry O_h

Symmetry number $\sigma = 24$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (CCl_4 soln.)	
a_{1g}	ν_1	CO stretch.	2112 C	ia	2112.4 M	
	ν_2	CrC stretch.	381 C	ia	381.2 S	
e_g	ν_3	CO stretch.	2018 C	ia	2018.4 M	
	ν_4	CrC stretch.	394 D	ia	394 W	
f_{1g}	ν_5	CrCO bend.	364 D	ia	ia	$\text{OC}(\nu_5 + \nu_7)$.
f_{1u}	ν_6	CO stretch.	2000 B	2000.4 VS	ia	
	ν_7	CrC stretch.	668 B	668.1 VS	ia	
	ν_8	CrCO bend.	441 B	440.5 S	ia	
	ν_9	CCrC deform.	98 B	97.8 M	ia	
f_{2g}	ν_{10}	CrCO bend.	436 D	ia	436.4 (liquid)	
	ν_{11}	CCrC deform.	95 D	ia	94.8 S (liquid)	
f_{2u}	ν_{12}	CrCO bend.	511 D	ia	ia	$\text{OC}(\nu_{10} + \nu_{12}, \nu_5 + \nu_{12})$.
	ν_{13}	CCrC deform.	68 E	ia	ia	$\text{OC}(\nu_3 + \nu_{13})$.

References

- [1] IR. N. J. Hawkins, H. C. Matraw, W. W. Sabol, and D. R. Carpenter, J. Chem. Phys. **23**, 2422 (1955).
 [2] Th. H. Murata and K. Kawai, J. Chem. Phys. **27**, 605 (1957).
 [3] IR.Th. L. H. Jones, Spectrochim. Acta **19**, 329 (1963).
 [4] IR.Th. J. M. Smith and L. H. Jones, J. Mol. Spectry. **20**, 248 (1966).
 [5] IR.R.Th. L. H. Jones, R. S. McDowell, and M. Goldblatt, Inorg. Chem. **8**, 2349 (1969).
 [6] R.Th. R. Pince et R. Poilblanc, Spectrochim. Acta **28A**, 907 (1972).

No. 211 Hexacarbonylmolybdenum(0) $\text{Mo}(^{12}\text{C}^{16}\text{O})_6$
Symmetry O_h

Symmetry number $\sigma = 24$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm^{-1}	cm^{-1} (Gas)	cm^{-1} (CCl_4 soln.)	
a_{1g}	ν_1	CO stretch.	2117 C	ia	2116.7 M	
	ν_2	MoC stretch.	402 C	ia	402.2 S	
e_g	ν_3	CO stretch.	2019 C	ia	2018.8 M	
	ν_4	MoC stretch.	392 C	ia	392 W	
f_{1g}	ν_5	MoCO bend.	342 D	ia	ia	$OC(\nu_5 + \nu_7)$.
f_{1u}	ν_6	CO stretch.	2003 B	2003.0 VS	ia	
	ν_7	MoC stretch.	596 B	595.6 VS	ia	
	ν_8	MoCO bend.	367 B	367.2 S	ia	
	ν_9	CMoC deform.	82 B	81.6 M	ia	
f_{2g}	ν_{10}	MoCO bend.	448 D	ia	448.3 (liquid)	
	ν_{11}	CMoC deform.	86 D	ia	86.0 S (liquid)	
f_{2u}	ν_{12}	MoCO bend.	507 D	ia	ia	$OC(\nu_5 + \nu_{12}, \nu_{10} + \nu_{12})$
	ν_{13}	CMoC deform.	60 E	ia	ia	$OC(\nu_3 + \nu_{13}, \nu_{10} + \nu_{13})$.

References

- [1] IR. N. J. Hawkins, H. C. Mattraw, W. W. Sabol, and D. R. Carpenter, J. Chem. Phys. **23**, 2422 (1955).
 [2] Th. H. Murata and K. Kawai, J. Chem. Phys. **27**, 605 (1957).
 [3] IR. L. H. Jones, J. Chem. Phys. **36**, 2375 (1962).
 [4] IR.Th. L. H. Jones, Spectrochim. Acta **19**, 329 (1963).
 [5] IR.Th. J. M. Smith and L. H. Jones, J. Mol. Spectry. **20**, 248 (1966).
 [6] IR.R.Th. L. H. Jones, R. S. McDowell, and M. Goldblatt, Inorg. Chem. **8**, 2349 (1969).
 [7] R.Th. R. Pince et R. Poilblanc, Spectrochim. Acta **28A**, 907 (1972).

No. 212 Hexacarbonyltungsten(0) $W(^{12}C^{16}O)_6$ Symmetry O_h Symmetry number $\sigma = 24$

Sym. class	No.	Approximate type of mode	Selected value of frequency	Infrared	Raman	Comments
			cm ⁻¹	cm ⁻¹ (Gas)	cm ⁻¹ (CS ₂ soln.)	
a_{1g}	ν_1	CO stretch.	2117 C	ia	2116.6 M	
	ν_2	WC stretch.	427 C	ia	427.1 S	
e_g	ν_3	CO stretch.	2010 C	ia	2009.8 M	
	ν_4	WC stretch.	412 C	ia	412 W	
f_{1g}	ν_5	WCO bend.	362 D	ia	ia	OC($\nu_5 + \nu_7$).
f_{1u}	ν_6	CO stretch.	1998 B	1997.6 VS	ia	
	ν_7	WC stretch.	587 B	586.6 VS	ia	
	ν_8	WCO bend.	374 B	374.4 S	ia	
	ν_9	CWC deform.	82 C	82.0 M	ia	
f_{2g}	ν_{10}	WCO bend.	458 D	ia	458.3 (liquid)	
	ν_{11}	CWC deform.	88 D	ia	88.3 S (liquid)	
f_{2u}	ν_{12}	WCO bend.	521 D	ia	ia	OC($\nu_5 + \nu_7$, $\nu_{10} + \nu_{12}$).
	ν_{13}	CWC deform.	61 E	ia	ia	OC($\nu_3 + \nu_{13}$).

References

- [1] IR.Th. L. H. Jones, Spectrochim. Acta **19**, 329 (1963).
 [2] IR.Th. J. M. Smith and L. H. Jones, J. Mol. Spectry. **20**, 248 (1966).
 [3] IR.R.Th. L. H. Jones, R. S. McDowell, and M. Goldblatt, Inorg. Chem. **8**, 2349 (1969).
 [4] R.Th. R. Pince et R. Poilblanc, Spectrochim. Acta **28A**, 907 (1972).

Empirical Formula Index for Consolidated Volumes I and II

In this index molecules are divided into two groups: (a) those containing no carbon atoms, which are arranged with the elemental symbols of the empirical formulas in alphabetical order and are listed alphabetically, and in ascending order of the empirical formula subscripts; (b) molecules containing carbon, which are ordered in the same way except that carbon is listed first and hydrogen second. No distinction is made for isotopic species in the empirical formula; thus deuterium is listed as H.

As an assistance to users of these data, this Index identifies the molecules treated in the present Consolidated Volume II and also those in the earlier Consolidated Volume I, which is available as part of a previous publication series, the NSRDS-NBS series distributed by the U.S. Government Printing Office. It can be ordered as Tables of Molecular Vibrational Frequencies, Consolidated Volume I, (SD Catalog No. C13.48:39), stock number 003-003-00845-1, price \$5.10, from the Superintendent of Documents, U.S. Government Printing Office, Washington, D.C. 20402.

Compounds Not Containing Carbon

Empirical Formula	Name	Volume and Number	Empirical Formula	Name	Volume and Number
AlCl ₃	Aluminum trichloride	II-36	Br ₃ ClSi	Tribromochlorosilane	I-47
AsF ₃	Arsenic trifluoride	II-38	Br ₃ GeH	Tribromogermane	II-93
AsF ₅	Arsenic pentafluoride	II-118	Br ₃ HSi	Tribromosilane	II-75
AsH ₃	Arsine	I-21	Br ₃ OP	Phosphoryl bromide	II-78
AsH ₃	Arsine-d ₃	I-22	Br ₄ Ge	Germanium tetrabromide	I-39
AsI ₃	Arsenic triiodide	II-39	Br ₄ Hf	Hafnium tetrabromide	II-66
BF ₃	Boron trifluoride- ¹⁰ B ¹¹ F ₃	II-31	Br ₄ Si	Silicon tetrabromide	I-31
BF ₃	Boron trifluoride- ¹¹ B ¹¹ F ₃	II-22	Br ₄ Sn	Tin (IV) bromide	I-41
B ₂ H ₆	Diborane- ¹⁰ B ₂ H ₆	I-55	Br ₄ Ti	Titanium tetrabromide	II-59
B ₂ H ₆	Diborane- ¹¹ B ₂ D ₆	I-56	Br ₄ Zn	Zirconium tetrabromide	II-62
BrCl ₃ Si	Bromotrichlorosilane	I-45	ClFO ₃	Perchloryl fluoride	II 81
BrF ₃	Bromine trifluoride	II-46	ClF ₂ N	Chlorodifluoroammonia	II-53
BrGeH ₃	Germyl bromide	II-88	ClF ₂ PS	Thiophosphoryl chlorodifluoride	II-112
BrGeH ₃	Germyl bromide-d ₃	II-89	ClF ₃	Chlorine trifluoride	II-45
BrH ₃ Si	Silyl bromide	I-44	ClF ₅ S	Sulfur chloride pentafluoride	II-122
BrNO	Nitrosyl bromide- ¹⁶ O ¹⁴ NBr	II-23	ClF ₅ W	Tungsten chloride pentafluoride	II-123
BrNO	Nitrosyl bromide- ¹⁶ O ¹⁵ NBr	II-24	ClGeH ₃	Germyl chloride	II-86
BrNO	Nitrosyl bromide- ¹⁸ O ¹⁵ NBr	II-25	ClGeH ₃ -d ₃	Germyl chloride-d ₃	II-87
BrO ₃ Re	Rhenium (VII) oxide bromide	II-95	ClHO	Hypochlorous acid	II-27
Br ₂ Cl ₂ Si	Dibromodichlorosilane	I-49	ClHO	Hypochlorous acid-d	II-28
Br ₂ FOP	Phosphoryl dibromofluoride	II-111	ClH ₃ Si	Silyl chloride	I-43
Br ₂ H ₂ Si	Dibromosilane	II-100	ClI ₃ Si	Chlorotriiodosilane	I-48
Br ₂ OS	Thionyl bromide	II-57	ClNO	Nitrosyl chloride- ¹⁶ O ¹⁴ NO	II-19
Br ₂ Sn	Tin (II) bromide	II-12	ClNO	Nitrosyl chloride- ¹⁶ O ¹⁵ NCl	II-20
			ClNO	Nitrosyl chloride- ¹⁸ O ¹⁴ NCl	II-21
			ClNO	Nitrosyl chloride- ¹⁸ O ¹⁵ NCl	II-22
			ClNO ₂	Nitryl chloride-Cl ¹⁴ NO ₂	II-43
			ClNO ₂	Nitryl chloride-Cl ¹⁵ NO ₂	II-44
			ClNO ₃	Chloronitrate-ClO ¹⁴ NO ₂	II-108
			ClNO ₃	Chloronitrate-ClO ¹⁵ NO ₂	II-109
			CINS	Thionitrosyl-S-chloride	II-30
			ClO ₃ Re	Rhenium (VII) oxide chloride	II-94
			Cl ₂ FN	Dichlorofluoroammonia	II-54
			Cl ₂ FOP	Phosphoryl dichlorofluoride	II-110
			Cl ₂ FPS	Thiophosphoryl dichlorofluoride	II-113
			Cl ₂ Ge	Germanium (II) chloride	II-9
			Cl ₂ H ₂ Si	Dichlorosilane	II-98
			Cl ₂ H ₂ Si	Dichlorosilane-d ₂	II 99
			Cl ₂ O	Oxygen dichloride	I-8
			Cl ₂ OS	Thionyl chloride	II-56
			Cl ₂ O ₂ S	Sulfuryl chloride	II-97
			Cl ₂ Pb	Lead (II) chloride	II-14
			Cl ₂ S	Sulfur dichloride	II-8
			Cl ₂ Sn	Tin (II) chloride	II-11
			Cl ₃ F ₂ P	Phosphorus trichloride difluoride	II-115
			Cl ₃ GeH	Trichlorogermane	II-92
			Cl ₃ HSi	Trichlorosilane	II-73
			Cl ₃ HSi	Trichlorosilane-d	II-74
			Cl ₃ ISi	Trichloroiodosilane	I-46
			Cl ₃ OP	Phosphoryl chloride	II-77
			Cl ₃ OV	Vanadyl (V) chloride	II-82

Empirical Formula	Name	Volume and Number	Empirical Formula	Name	Volume and Number
Cl ₃ P	Phosphorus trichloride	I-20	F ₆ Te	Tellurium hexafluoride	II-120
Cl ₃ PS	Thiophosphoryl trichloride	II-80	F ₆ U	Uranium (VI) fluoride	I-54
			F ₆ W	Tungsten (VI) fluoride	I-53
Cl ₄ Ge	Germanium tetrachloride	I-38			
Cl ₄ Hf	Hafnium tetrachloride	II-65	GeH ₃ I	Germyl iodide	II-90
Cl ₄ Si	Silicon tetrachloride	I-30	GeH ₃ I	Germyl iodide-d ₃	II-91
Cl ₄ Sn	Tin (IV) chloride	I-40	GeH ₄	Germane	I-33
Cl ₄ Ti	Titanium tetrachloride	II-58	GeH ₄	Germane-d ₁	I-34
Cl ₄ Zn	Zirconium tetrachloride	II-61	GeH ₄	Germane-d ₂	I-35
			GeH ₄	Germane-d ₃	I-36
			GeH ₄	Germane-d ₄	I-37
Cl ₅ P	Phosphorus pentachloride	II-116			
Cl ₅ Sb	Antimony pentachloride	II-119	Ge ₂ H ₆	Digermane	II-125
			Ge ₂ H ₆	Digermane-d ₆	II-126
FGeH ₃	Germyl fluoride	II-84			
FGeH ₃	Germyl fluoride-d ₃	II-85	HNO ₃	Nitric acid-H ¹⁴ NO ₃	II-102
FHO	Hypofluorous acid	II-26	HNO ₃	Nitric acid-d-D ¹⁴ NO ₃	II-103
FH ₃ Si	Silyl fluoride	I-42	HNO ₃	Nitric acid-H ¹⁵ NO ₃	II-104
FMnO ₃	Permanganyl fluoride	II-83	HNO ₃	Nitric acid-d-D ¹⁵ NO ₃	II-105
FNO	Nitrosyl fluoride- ¹⁶ O ¹⁴ NF	II-15	HN ₃	Hydrazoic acid	II-50
FNO	Nitrosyl fluoride- ¹⁶ O ¹⁵ NF	II-16	HN ₃	Hydrazoic acid-d	II-51
FNO	Nitrosyl fluoride- ¹⁸ O ¹⁴ NF	II-17			
FNO	Nitrosyl fluoride- ¹⁸ O ¹⁵ NF	II-18	H ₂ O	Water	I-4
FNO ₂	Nitryl fluoride-F ¹⁴ NO ₂	II-41	H ₂ O	Water-H ₂ ¹⁸ O	II-6
FNO ₂	Nitryl fluoride-F ¹⁵ NO ₂	II-42	H ₂ O	Water-d ₁	I-5
FNO ₃	Fluoronitrate-FO ¹⁴ NO ₂	II-106	H ₂ O	Water-d ₂	I-6
FNO ₃	Fluoronitrate-FO ¹⁵ NO ₂	II-107	H ₂ O ₂	Hydrogen peroxide	II-47
FNS	Thionitrosyl-S-fluoride	II-29	H ₂ S	Hydrogen sulfide	I-9
F ₂ HN	Difluoroamine	II-52	H ₂ S	Hydrogen sulfide-d ₂	I-10
F ₂ N ₂	trans-1,2-Difluorodiazine	II-40	H ₂ S ₂	Hydrogen persulfide	II-48
F ₂ O	Oxygen difluoride	I-7	H ₂ Se	Hydrogen selenide	I-12
F ₂ Kr	Krypton difluoride	II-1	H ₂ Se	Hydrogen selenide-d ₁	I-13
F ₂ OS	Thionyl fluoride	II-55			
F ₂ O ₂ S	Sulfuryl fluoride	II-96	H ₃ N	Ammonia	I-14
F ₂ O ₂ Se	Selenium dioxide difluoride	II-101	H ₃ N	Ammonia-d ₃	I-15
F ₂ Pb	Lead (II) fluoride	II-13	H ₃ N	Ammonia-t ₃	II-34
F ₂ S ₂	Difluoro disulfane	II-49	H ₃ P	Phosphine	I-17
F ₂ Sn	Tin (II) fluoride	II-10	H ₃ P	Phosphine-d ₃	I-18
			H ₃ Sb	Stibine	I-23
F ₃ HSi	Trifluorosilane	II-71	H ₃ Sb	Stibine-d ₃	I-24
F ₃ HSi	Trifluorosilane-d	II-72			
F ₃ N	Nitrogen trifluoride- ¹⁴ NF ₃	I-16	H ₄ Si	Silane	I-25
F ₃ N	Nitrogen trifluoride- ¹⁵ NF ₃	II-35	H ₄ Si	Silane-d ₂	I-26
F ₃ NS	Thionitrosyl trifluoride	II-70	H ₄ Si	Silane-d ₃	I-27
F ₃ OP	Phosphoryl fluoride	II-76	H ₄ Si	Silane-d ₄	I-28
F ₃ P	Phosphorus trifluoride	I-19			
F ₃ PS	Thiophosphoryl trifluoride	II-79	IF ₇	Iodine heptafluoride	II-124
			I ₃ P	Phosphorus triiodide	II-37
F ₄ Si	Silicon tetrafluoride	I-29			
			I ₄ Hf	Hafnium tetraiodide	II-67
F ₅ P	Phosphorus pentafluoride	II-114	I ₄ Si	Silicon tetraiodide	I-32
F ₅ V	Vanadium pentafluoride	II-117	I ₄ Ti	Titanium tetraiodide	II-60
			I ₄ Zr	Zirconium tetraiodide	II-63
F ₆ Ir	Iridium (VI) fluoride	II-121			
F ₆ Mo	Molybdenum (VI) fluoride	I-52	NO ₂	Nitrogen dioxide- ¹⁴ NO ₂	II-2
F ₆ S	Sulfur hexafluoride	I-50			
F ₆ Se	Selenium hexafluoride	I-51			

Empirical Formula	Name	Volume and Number	Empirical Formula	Name	Volume and Number
NO ₂	Nitrogen dioxide- ¹⁵ N ₂ O ₂	II-3	CHCl ₃	Trichloromethane	I-89
N ₂ O	Nitrous oxide	I-1	CHCl ₃	Trichloromethane-d ₁	I-90
N ₂ O	Nitrous oxide- ¹⁴ N ¹⁵ N ¹⁶ O	I-2	CHF ₃	Trifluoromethane	I-88
N ₂ O	Nitrous oxide- ¹⁵ N ₂ O	I-3	CHF ₃	Trifluoromethane-d	II-139
O ₂ S	Sulfur dioxide	I-11	CHN	Hydrogen cyanide	I-62
O ₂ S	Sulfur dioxide-S ¹⁸ O ₂	II-7	CHN	Hydrogen cyanide-d ₁	I-63
O ₃	Ozone- ¹⁶ O ₃	II-4	CHNO	Isocyanic acid	II-134
O ₃	Ozone- ¹⁸ O ₃	II-5	CHNO	Isocyanic acid-d	II-135
O ₃ S	Sulfur trioxide	II-33	CH ₂ BrCl	Bromochloromethane	I-102
O ₄ Os	Osmium tetroxide-Os ¹⁶ O ₄	II-68	CH ₂ BrCl	Bromochloromethane-d ₁	I-103
O ₄ Os	Osmium tetroxide-Os ¹⁸ O ₄	II-69	CH ₂ BrCl	Bromochloromethane-d ₂	I-104
O ₄ Ru	Ruthenium tetroxide	II-64	CH ₂ Br ₂	Dibromomethane	I-98
			CH ₂ Br ₂	Dibromomethane-d ₁	I-99
			CH ₂ Br ₂	Dibromomethane-d ₂	I-100
			CH ₂ Cl ₂	Dichloromethane	I-95
			CH ₂ Cl ₂	Dichloromethane-d ₁	I-96
			CH ₂ Cl ₂	Dichloromethane-d ₂	I-97
			CH ₂ O	Formaldehyde	I-68
			CH ₂ O	Formaldehyde-d ₁	I-69
			CH ₂ O	Formaldehyde-d ₂	I-70
			CH ₂ O ₂	Formic acid	I-105
			CH ₂ O ₂	Formic acid-d ₂	I-106
			CH ₃ BO	Borine carbonyl- ¹⁰ BH ₃ CO	II-150
			CH ₃ BO	Borine carbonyl-d ₃ ¹⁰ BD ₃ CO	II-151
			CH ₃ BO	Borine carbonyl- ¹¹ BH ₃ CO	II-152
			CH ₃ BO	Borine carbonyl-d ₃ ¹¹ BD ₃ CO	II-153
			CH ₃ Br	Methylbromide	I-84
			CH ₃ Br	Methylbromide-d ₃	I-85
			CH ₃ Cl	Methylchloride	I-82
			CH ₃ Cl	Methylchloride-d ₃	I-83
			CH ₃ F	Methylfluoride	I-80
			CH ₃ F	Methylfluoride-d ₃	I-81
			CH ₃ GeI ₃	Methyltriiodogermane	II-159
			CH ₃ HgI	Methylmercuric iodide	II-154
			CH ₃ I	Methyliodide	I-86
			CH ₃ I	Methyliodide-d ₃	I-87
			CH ₄	Methane	I-71
			CH ₄	Methane-d ₁	I-72
			CH ₄	Methane-d ₂	I-73
			CH ₄	Methane-d ₃	I-74
			CH ₄	Methane-d ₄	I-75
			CH ₄ O	Methanol (Gas)	I-107
			CH ₄ O	Methanol (Liquid)	I-108
			CH ₄ O	Methanol-d ₁ (Gas)	I-109
			CH ₄ O	Methanol-d ₁ (Liquid)	I-110
			CH ₄ O	Methanol-d ₃ (Gas)	I-111
			CH ₄ O	Methanol-d ₃ (Liquid)	I-112
			CH ₄ O	Methanol-d ₄ (Gas)	I-113
			CH ₅ N	Methylamine	I-114
			CH ₅ N	Methylamine-d ₂	I-115
			CH ₅ N	Methylamine-d ₃	I-116
			CH ₅ N	Methylamine-d ₅	I-117
			CH ₆ Ge	Methylgermane	II-157
			CH ₆ Ge	Methyl-d ₃ -germane	II-158
			CH ₄ Si	Methylsilane	II-155
			CH ₆ Si	Methylsilane-d ₃	II-156

Compounds Containing Carbon

Empirical Formula	Name	Volume and Number	Empirical Formula	Name	Volume and Number
C_3H_4	Methylacetylene	I-179	C_4H_6	1,3-Butadiene	I-205
C_3H_4	Methylacetylene- d_1	I-180	C_4H_6	1,3-Butadiene-1- d_1 , trans	I-206
C_3H_4	Methyl- d_3 -acetylene	I-181	C_4H_6	1,3-Butadiene-1,1,2- d_3	I-207
C_3H_4	Methylacetylene- d_4	I-182	C_4H_6	1,3-Butadiene-1,1,4,4- d_4	I-208
C_3H_4O	Propenal	I-185	C_4H_6	1,3-Butadiene- d_6	I-209
C_3H_5N	Ethylcyanide	I-188	C_4H_6	2-Butyne	I-210
C_3H_6	Cyclopropane	I-186	C_4H_8	Cyclobutane	I-211
C_3H_6	Cyclopropane- d_6	I-187	C_4H_8	Cyclobutane- d_8	I-212
C_3H_6O	Acetone	I-189	C_4H_8	2-Methylpropene	I-213
C_3H_6O	Acetone- d_3	I-190	C_4H_8	2-Methyl- d_3 -propene-3,3,3- d_3	I-214
C_3H_6O	Acetone- d_6	I-191	C_4H_8O	2-Butanone (trans form)	I-215
C_3H_6O	Methyl acetate	I-197	$C_4H_8O_2$	1,4-Dioxane	II-207
$C_3H_6O_2$	Methyl acetate- d_3	I-198	C_4H_{10}	n-Butane (trans form)	I-216
$C_3H_6O_2$	Methyl- d_3 -acetate	I-199	C_4H_{10}	n-Butane (gauche form)	I-217
$C_3H_6O_2$	Methyl acetate- d_6	I-200			
C_3H_8	Propane	I-192	C_5H_2O	Diethynyl ketone	II-208
C_3H_8	Propane- d_2	I-193			
C_3H_8	Propane- d_3	I-194	C_6CrO_6	Hexacarbonylchromium	II-210
C_3H_8	Propane- d_6	I-195	C_6MoO_6	Hexacarbonylmolybdenum	II-211
C_3H_8	Propane- d_8	I-196	C_6N_2	Dicyanodiacetylene	II-209
			C_6O_6W	Hexacarbonyltungsten	II-212
C_4HBr	Bromodiacetylene	II-205	C_6H_6	Benzene	I-218
C_4HCl	Chlorodiacetylene	II-204	C_6H_6	Benzene- d_6	I-219
C_4HI	Iododiacetylene	II-206	C_6H_{12}	Cyclohexane	I-220
C_4H_2	Butadiyne	I-201	C_6H_{12}	Cyclohexane- d_{12}	I-221
C_4H_4O	Furan	I-202	$-(CH_2)_n-$	Poly-(methylene) $_n$	I-222
C_4H_4S	Thiophene	I-203	$-(CH_2)_n-$	Poly-(methylene- d_2) $_n$	I-223
C_4H_4S	Thiophene- d_4	I-204			