

Vibrational and Electronic Energy Levels of Polyatomic Transient Molecules. Supplement A

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A summary is presented of recently published, critically evaluated experimental vibrational and electronic energy level data for neutral and ionic transient molecules and high temperature species possessing from three to sixteen atoms. Although the emphasis is on species with lifetimes too short for study using conventional sampling techniques, there has been selective extension of the compilation to include data for isolated molecules of inorganic species such as the heavy-metal oxides, which are important in a wide variety of industrial chemical systems. Radiative lifetimes and the principal rotational constants are included. Observations in the gas phase, in molecular beams, and in rare-gas and diatomic molecule matrices are evaluated. The types of measurement surveyed include conventional and laser-based absorption and emission techniques, laser absorption with mass analysis, and photoelectron spectroscopy. Not counting isotopic species, 904 molecules are surveyed, and 2696 distinct references are given. © 1998 American Institute of Physics and American Chemical Society. [S0047-2689(98)00102-0]

Key words: electronic energy levels; emission spectra; experimental data; free radicals; gas phase; high-temperature molecules; infrared spectra; laser-excited fluorescence; matrix isolation; molecular ions; photoelectron spectroscopy; polyatomic molecules; radiative lifetimes; Raman spectra; rotational constants; transient molecules; ultraviolet absorption; vibrational energy levels.

Contents

1. Introduction.....	115	8.15. Hydrocarbons with More Than Eight Atoms.....	342
2. Scope of Review.....	116	8.16. Non-Hydrocarbons with More Than Eight Atoms.....	357
3. Types of Measurement.....	116	8.17. Molecules Related to Benzene.....	368
4. Guide to the Compilation.....	119	9. Molecule Index.....	373
5. Abbreviations.....	121		
6. Acknowledgment.....	122		
7. General References.....	122		
8. Tables.....	123		
8.1. H ₃ ⁺ , H ₃ , and Triatomic Dihydrides.....	123		
8.2. Triatomic Monohydrides.....	136		
8.3. Triatomic Nonhydrides.....	159		
8.4. Four-Atomic Trihydrides.....	211		
8.5. Four-Atomic Dihydrides.....	217		
8.6. Four-Atomic Monohydrides.....	231		
8.7. Four-Atomic Nonhydrides.....	241		
8.8. Five-Atomic Tetra- and Trihydrides.....	265		
8.9. Five-Atomic Dihydrides.....	275		
8.10. Five-Atomic Monohydrides.....	286		
8.11. Five-Atomic Nonhydrides.....	292		
8.12. Six-Atomic Molecules.....	307		
8.13. Seven-Atomic Molecules.....	323		
8.14. Eight-Atomic Molecules.....	332		

1. Introduction

Most chemical processes—including not only laboratory and industrial chemical syntheses but also those which occur in flames, propellant systems, the initiation of energetic materials, atmospheric pollution, chemical vapor deposition, and plasma processing—consist of a complicated sequence of interrelated reactions in which neutral and charged molecular fragments play essential roles. Although these fragments are present in only very small concentration, they are highly chemically reactive. If a specific molecular fragment is removed from the system, as by introducing a scavenger molecule, the reactions in which that fragment participates stop. Other parts of the overall process continue, resulting in very significant changes in product distribution and yield.

In the early studies of complex chemical processes, it was necessary to postulate mechanisms involving such transient intermediates, present in concentrations too small for direct detection. Conventional end product analysis aids in the se-

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lection of suitable mechanisms, but generally does not yield a complete description of the system. Consequently, the improvement of industrial chemical processes often is achieved by semiempirical experimentation. The determination of the detailed chemical mechanism would, in turn, permit the development of rational strategies for removing undesired products and enhancing the yield of the desired species.

In recent years, there has been great progress in the development of techniques suitable for monitoring chemical reaction intermediates. Molecular spectroscopy is especially well suited to this task. Optical detection can be used not only for gas-phase measurements, but also for studies of processes which occur on surfaces or in the condensed phase. It also permits remote sensing, an important advantage. A wide variety of recently developed laser-based spectroscopic detection schemes are not only highly sensitive but also space and time specific. Although the development of spectroscopy-based diagnostics for chemical reaction systems is in its infancy, already the laboratory application of sophisticated sampling and observation techniques has yielded a wealth of vibrational and electronic spectral data for reaction intermediates.

For many years, the most important source of vibrational and electronic energy level data for small polyatomic reaction intermediates was the compilation of spectroscopic data for small polyatomic molecules (3–12 atoms) given by Herzberg.¹ To meet the need for an updated, critically evaluated compilation, a series of publications^{2–4} have appeared in this journal, culminating in the publication in 1994 of a monograph⁵ which presented evaluated spectral data for more than 1550 small polyatomic transient molecules. Vibrational fundamentals in the ground and excited electronic states and radiative lifetimes were included. To aid in spectral identification, the principal rotational constants were also given to three decimal places. These tables have provided the basis for a personal computer software database (Vibrational and Electronic Energy Levels of Small Polyatomic Transient Molecules, National Institute of Standards and Technology Standard Reference Database 26), designed to supplement the published compilation by providing a capability for rapid searches by molecule or wave number.

The rapid growth in the scientific literature concerned with the spectroscopic study of transient molecules and with their detection in chemical reaction systems continues. Since the October 1993 cutoff in the data evaluation for the 1994 monograph,⁵ substantive new spectroscopic data have been published for many species included in it, and the first data have become available for several hundred other molecules. There has been especially great progress in the spectroscopic characterization of transient species produced by the reaction of metal atoms with oxygen and other small molecules. This paper attempts to provide a comprehensive, critically evaluated summary of these new data on the vibrational and electronic energy levels of small polyatomic transient molecules, in order to support further research and new technologies

such as those of plasma processing and chemical vapor deposition.

2. Scope of Review

This review provides a critical evaluation of the vibrational and electronic spectral data available in mid-1997 for approximately 900 small polyatomic transient molecules. Of these, about 570 were not represented in the 1994 monograph; the remainder are molecules included in that monograph for which new data have become available. Despite this rapid progress, many gaps remain in our knowledge of the energy levels of the species represented, and many new and potentially important transient molecules are still to be discovered.

Data have been selectively included for some molecules which are important in environmental and industrial chemical reaction systems but which can be studied only with difficulty using conventional sampling techniques because of the ease with which they decompose, rearrange, or polymerize. Also included are data derived from spectra of many high-temperature species, such as metal oxides, studied in molecular beams and in rare-gas matrices.

3. Types of Measurement

Studies in the gas phase offer the potential for the most precise, detailed measurements. Because of the high chemical reactivity of transient molecules, it is difficult to obtain gas-phase infrared spectra of them. The well known advantages of Fourier transform infrared measurements, coupled with sophisticated digital data handling procedures, have permitted the acquisition of gas-phase survey spectra for a number of transient molecules. Diode lasers and other laser-based techniques with limited tunability have been used to obtain high resolution spectra of individual vibrational transitions of these species.

Although vibrational frequencies of ground-state molecular ions have frequently been estimated from structure in Rydberg transitions of the parent neutral species, such data are not included in this compilation, since many of these transitions have residual valence character, resulting in significant variations in vibrational frequencies from one Rydberg state to another.

As in the earlier compilations, spectral data obtained in rare gas and small covalent molecule matrices are included. The application of matrix isolation sampling for the stabilization and spectroscopic study of uncharged reaction intermediates has recently been reviewed.⁶ Because nitrogen and the rare gases are transparent through the entire infrared spectral region, matrix isolation measurements provide a potentially valuable survey tool. In these matrices, infrared absorptions are typically sharp, with half band widths between 0.1 and 1 cm⁻¹. Rotational structure is, with few exceptions, quenched. Multiple trapping sites occur, often resulting in

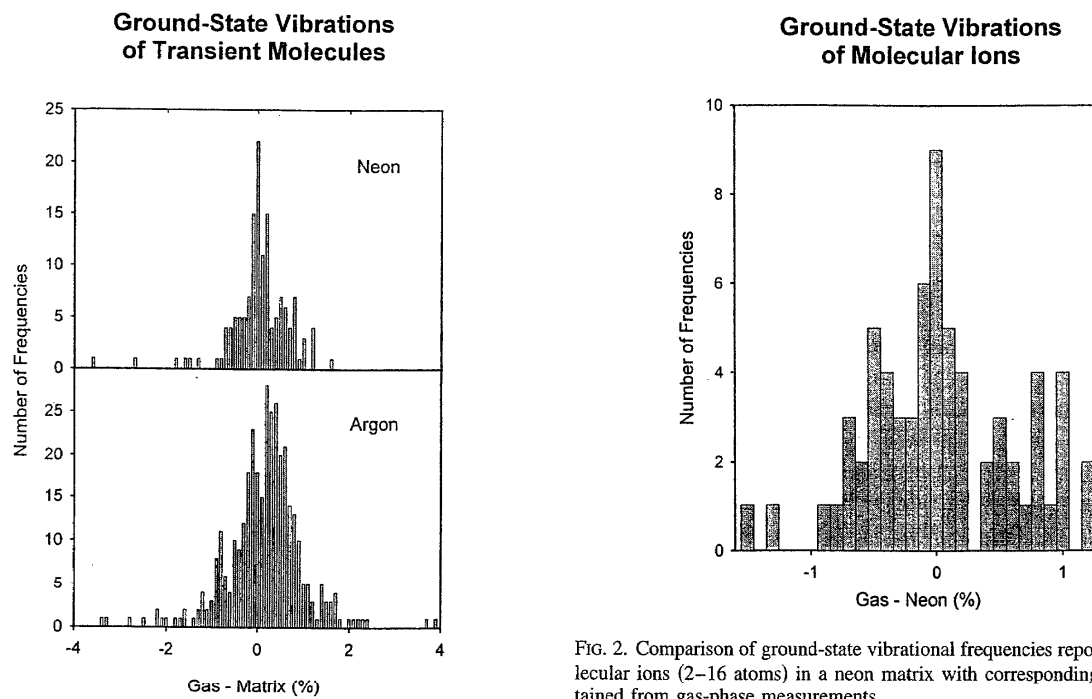


FIG. 1. Comparison of ground-state vibrational frequencies reported for transient molecules (2–16 atoms) in neon and argon matrices with corresponding values obtained from gas-phase measurements.

the appearance of several absorption maxima—usually one or two of which predominate—over a range of a few cm^{-1} .

Matrix shifts for covalently bonded molecules trapped in solid neon or argon often are quite small. A comparison⁷ of the positions of the ground-state vibrational fundamentals of over two hundred diatomic molecules observed in the gas phase and in nitrogen and rare-gas matrices has shown that, typically, the smallest matrix shift occurs for neon matrix observations, with successively greater matrix shifts for the heavier rare gases and for nitrogen. Except for very weakly bonded molecules and for the alkali metal and Group IIIa halides, matrix shifts of most diatomic molecules isolated in solid argon are smaller than 2%. Similar conclusions resulted from a comparison of neon- and argon-matrix shifts for the ground-state vibrational fundamentals of larger molecules.⁸ The generalization that matrix interactions are minimal for neon and that they increase as the mass of the rare gas is increased and become even more important for nitrogen and most other small molecule matrices is supported by experimental observations on larger molecules, as well. Figure 1 compares the observed matrix shifts for the ground-state fundamental vibrations of transient molecules trapped in solid neon and argon. For neon matrices, the maximum in the distribution lies near 0.0%, and for argon matrices, near 0.2%. For both neon and argon matrices, fewer than one-tenth of the matrix shifts are greater than 1%.

For molecular ions, neon is the matrix of choice. Polarization and charge-transfer interactions become successively

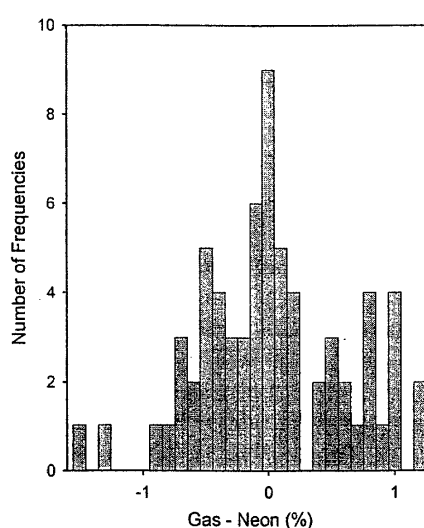


FIG. 2. Comparison of ground-state vibrational frequencies reported for molecular ions (2–16 atoms) in a neon matrix with corresponding values obtained from gas-phase measurements.

more important for molecules isolated in the heavier rare gases. Charge delocalization sometimes also occurs for ionic species trapped in the rare gases.^{9,10} The anomalously large matrix shift for ν_3 of ClHCl^- may be attributed to this phenomenon. As is shown in Fig. 2, data for small cation species trapped in solid neon are consistent with the matrix shift generalizations given above. Only twelve comparisons are available for molecular cations observed both in the gas phase and trapped in solid argon. The absolute values of five of the observed matrix shifts are greater than 1%. For several other vibrations of molecular ions which have been observed in both neon and argon matrices but not in the gas phase, there are deviations greater than 1% between the neon- and argon-matrix frequencies. Very few comparisons are possible for molecular anions. A number of these species have been generated in rare-gas (usually argon) matrices by charge transfer between a precursor molecule and an alkali metal atom. Recent studies of such species as CO_2^- and SO_2^- generated instead by photoionization and/or Penning ionization and trapped in solid neon indicate that shifts on the order of 50 cm^{-1} may be attributed to the relatively strong interaction of the anion with the nearby alkali metal cation. On the other hand, when the uncharged molecule has a relatively large electron affinity, as is true for C_2 and for NO_2 , charge transfer occurs at a relatively great separation, and a substantial fraction of the anion population may be trapped in sites in which interaction with the alkali metal cation is minimal.

Matrix shifts for vibrations associated with ionic bonds are often considerably larger than those associated with uncharged molecules or with intramolecular vibrations of mo-

lular ions. Criteria for inclusion of data for species which include an ionic bond are exemplified by the selection process for the heavy-metal oxides. Often the stable dioxide structures include an $M^+O_2^-$ species with significant covalent bond character for the attachment of M^+ , as evidenced by a substantial shift in the O_2^- stretching fundamental as M^+ is varied. Such species are included in the compilation. On the other hand, there is little evidence for substantial metal-atom participation in the vibrations characteristic of the O_3^- moiety of $M^+O_3^-$. Accordingly, spectral data are given for O_3^- , but not for $M^+O_3^-$.

Data are beginning to appear for molecules trapped in a hydrogen matrix. Insufficient information is available for generalization on the magnitude of matrix shifts in this medium. For the few species heretofore studied, including several transient molecules present in this compilation, the matrix shifts have been comparable to those in a neon matrix.

Many other matrix materials have also been employed for spectroscopic studies. However, complications due to reaction or to relatively strong interaction (e.g., hydrogen bonding) of the transient molecule with the matrix frequently occur. Therefore, observations in such media as solid hydrocarbons and aqueous solutions and studies of condensed reaction products without an inert carrier have been excluded.

Because of the prevalence of electronic emission spectra and the sensitivity, rapid time response, and cumulative detection capability of the photographic plate in the visible and ultraviolet spectral regions, the study of the electronic spectra of gas-phase reaction intermediates has a comparatively long history. Flash photolysis has permitted the preparation of relatively high concentrations of transient species. Many electronic band systems of gas-phase transient molecules have been discovered through flash photolysis studies. More recently, a wide variety of laser-based techniques have also been used for electronic spectral observations, often with exceptionally high detection sensitivity. The spatial configuration of the laser beam makes possible crossed molecular beam-laser beam studies, providing an extremely powerful tool for studies of the energy levels of molecules in molecular beams. Laser-based techniques show great promise for application in the development of diagnostics for chemical reaction intermediates in the environment, the laboratory, and the industrial plant. Laser studies may be broadly classified according to whether the interaction of the molecule with the laser beam(s) is followed by photon or mass detection. Photon-based observations are amenable to remote sensing applications. Because pulsed lasers offer an exceptionally wide range of time specificity, they are very useful for determining radiative lifetimes and rates of elementary chemical reactions.

Much valuable information on the energy levels of molecular cations has been obtained from photoelectron spectroscopy. These tables include selective coverage of the voluminous literature on photoelectron spectroscopic

measurements. The number of stable molecules which possess more than six atoms for which photoelectron spectra have been reported is too great to permit the inclusion of low-to-moderate resolution photoelectron spectral data for molecular cations with more than six atoms. Those who need such data for larger molecules may find the reviews by Turner *et al.*,¹¹ Rabalais,¹² and Kimura *et al.*¹³ helpful. Several criteria are important in determining whether a given reference should be included in the present work. The first of these is resolution. In the few instances in which high resolution photoelectron data are available, these data are heavily weighted. Where direct spectroscopic observation is possible, the measurements generally are of considerably higher precision than are the photoelectron data, which are then omitted from the tables. A second criterion is the availability of adiabatic ionization potentials. In order to obtain information on the positions of electronic transitions from photoelectron spectral data, it is necessary to subtract the first ionization potential from the energy of the photoelectron band. Where there is little change in the molecular geometry in the transition, the difference between the vertical ionization potentials gives a reasonable approximation to the position of the electronic transition. However, this is not the general case. Therefore, priority is given to papers which include adiabatic ionization potentials.

For most photoelectron spectroscopic transitions, structure has not been resolved. Many of these states are dissociative. Further information on the dissociation products can be obtained from values of the appearance potentials for various products in photoionization studies on the parent molecule. Such studies are beyond the scope of this review. The tables of ionization and appearance potentials by Lias and co-workers^{14,15} constitute a valuable source of information on the appearance potentials of photofragments.

The range of tunability of visible and ultraviolet lasers, like that of infrared lasers, is limited. Therefore, a preliminary survey using conventional gas-phase and/or matrix-isolation spectroscopic studies is often desirable. A comparison of the positions of the electronic band origins of diatomic molecules in the gas phase and in rare-gas and nitrogen matrices has been published.¹⁶ As in the determination of ground-state vibrational energy levels, neon is the matrix material of choice, with a sharp maximum at 0.0% in the distribution of matrix deviations for valence transitions of covalently bonded molecules. In argon-matrix observations, most such band origins are shifted by less than 2% from the gas-phase values. At the somewhat higher temperatures often used for electronic spectral observations in matrices of the heavier rare gases or of nitrogen, relatively broad phonon bands become prominent. The blue shift of the phonon maximum from the zero-phonon line in absorption measurements, and the red shift in emission measurements, typically amount to approximately 1 to 1.5%. Rydberg transitions of molecules in matrices often are greatly broadened and experience much larger shifts. Further details of the

behavior of electronic transitions of matrix-isolated molecules have previously been discussed.^{3,6,16}

The radiative lifetime of a relatively large molecule isolated in a rare-gas matrix is frequently related to the radiative lifetime in the gas phase by a simple refractive index correction.¹⁷ In a neon matrix, such a correction typically decreases the radiative lifetime by about 15%. For such molecules, often intramolecular mechanisms for nonradiative energy transfer are available both in the gas phase and in the matrix. On the other hand, the density of excited states is much lower for small molecules, and matrix shifts may alter perturbation interactions between strongly coupled electronic states, providing a path for nonradiative deactivation. In this circumstance, fluorescence which is prominent in the gas phase may even be completely quenched in the matrix.

4. Guide to the Compilation

The goal of this paper is to extend the previously published compilation⁵ of critically evaluated vibrational and electronic spectroscopic data for small polyatomic transient molecules. The literature has been surveyed through June 1997; only limited addition of more recent data has been possible. Unfortunately, with a few exceptions it is not possible to include data for stable molecules. However, the spectra of many of these species are relatively well established, and sources of data such as the tables of Herzberg¹ and Shimanouchi¹⁸ remain extremely useful. In obtaining spectral identifications with the help of the present compilation, it is crucial that the possible contribution of the absorptions or emissions by a stable molecule also be considered.

Considerable effort has been expended to provide a critical evaluation of the data. However, for many species the available data are meager. The identities of some species have been proposed on the basis of chemical evidence. While such evidence may be quite compelling, it is not definitive. Many examples could be cited in which a spectrum was later reassigned to characteristic impurities in the sample. Where chemical evidence has provided a reasonable basis for the assignment of vibrational or electronic bands to a transient molecule, data have been included in this compilation, in the hope that further testing of the assignment will be facilitated.

While every effort has been made to make these tables as complete as possible, for various reasons omissions do occur. There remains some selectivity in the coverage of electronic spectral data for larger molecules. It is planned to support this database, with further selective extension, by the preparation of additional supplements. Data from the earlier tables have sometimes been omitted from this paper because more recent data dictate a reassignment or because there has been a subsequent refinement. An important example of this latter situation is the replacement of low resolution photoelectron spectral data by spectroscopic studies with apprecia-

bly higher resolution and greater precision. Candidate molecules or energy levels may also have been inadvertently omitted. Suggestions of additions or needed revisions to the data to be included in subsequent extensions of this database are welcome, as are inquiries regarding new data added after the publication cutoff data for this compilation.

Molecular formulas are used in this compilation. In order to permit a compact index, an attempt has been made to provide as much structural information as possible in a minimal amount of space. This restriction is especially severe for larger molecules. The following formula abbreviations have been used:

<i>br</i>	bridged
<i>cyc</i>	cyclic. If parentheses follow, only the atoms enclosed in them are included in the ring.
<i>c</i>	<i>cis</i>
<i>t</i>	<i>trans</i>

Where heavy isotopic peaks are resolved, data are given for the most abundant isotopic species (e.g., ⁷Li, ¹¹B, ³⁵Cl, ⁷⁹Br).

As in the earlier compilations, the tables are grouped by the number of atoms in the molecule and, secondarily, by the number of hydrogen atoms present. Molecules within a given section of the tables are arranged in the order of increasing number of valence electrons. For species with the same number of valence electrons, molecules with a simple chain of three heavy atoms are listed in the order, first, of the number of valence electrons in the central atom of the chain and, second, of the row which this atom occupies in the Periodic Table. For larger molecules, the sequence is somewhat arbitrary, but criteria of increasing molecular size and grouping in the Periodic Table (e.g., the order N, P, O, S) are used. Halogen-substituted species are placed immediately after the related hydrides. Data for molecules related to benzene are presented in a separate section. As in the earlier tables, data are included for both the normal and the fully deuterium-substituted molecule. However, only the hydrogen-containing species is listed in the index.

The heading for each electronic state gives its symmetry, the point group to which the molecule belongs in that electronic state, and, where available, references to the determination of a quantitative molecular structure. For C_{2v} molecules, there is potential ambiguity in the definition of the molecular symmetry axes. The convention in which the *x* axis is chosen perpendicular to the plane of the molecule, recommended by the Joint Commission for Spectroscopy of IAU and IUPAP,¹⁹ has been adopted. Often this has required the interchange of published assignments of energy levels with B₁ and B₂ symmetry.

Most authors of papers on photoelectron spectroscopy have proposed assignments for the various photoelectron bands, using arguments based on molecular orbital theory and often on semiempirical or *ab initio* calculations. These assignments have been included in the present tables. Where

several conflicting assignments have been given in the literature, an attempt has been made to choose the most satisfactory one. Generally, the assignments of photoelectron spectra have been made with the presumption that the point group to which the molecular cation belongs is the same in all of its excited states. Structural data for these excited states are extremely rare. Therefore, the molecular point group which has been adopted in the analysis of the photoelectron spectrum is given in these tables. In practice, it is likely that there is some variation in excited-state molecular symmetries. Thus, a bent molecular ion may become linear in some of its excited states. For highly symmetric species, Jahn-Teller distortion may reduce the molecular symmetry.

The energy of the electronic transition follows the state designation and symmetry information. Where possible, T_0 , the energy separation between the electronic energy level of interest and the ground electronic, vibrational, and rotational states of the molecule, is given. However, where only low resolution data or photoelectron data are available, often only band maxima have been given in the literature. With photoelectron data, T_0 is derived by subtracting the value of the first ionization potential from that of the higher ionization potential which corresponds to the state of interest. When data for the first adiabatic ionization potential are available, the footnote phrase "from vertical ionization potential" implies that the first adiabatic ionization potential is known but that the higher ionization potential is measured to the peak maximum; the phrase "from vertical ionization potentials" implies that the energy difference between the higher and the first absorption maximum was used. If the first photoelectron transition has a gradual onset, a better value of the first ionization potential may have been obtained from photoionization data or from the extrapolation of Rydberg series in the spectrum of the parent molecule. Supplementary sources of data for the first ionization potential are cited in the tables. However, if the difference between the first adiabatic ionization potential obtained in the photoelectron spectrum and that obtained in other measurements amounts to only 10 or 20 meV, the photoelectron spectroscopic value is used, because of the advantage of a consistent set of measurements. Where threshold energies differ by one quantum in a vibrational progression, a best value for the ionization potential is chosen which coincides with the most probable position of the vibrationally unexcited transition. Because of inherent uncertainties in the determination of higher ionization potentials in many photoelectron spectral measurements, photoelectron peaks above about 18 eV are often omitted. Except where otherwise indicated, the units of all quantities in these tables are cm^{-1} . Error estimates are those of the authors of the original literature. The numbers in parentheses give these estimated errors in relation to the last digits of the electronic or vibrational frequency (e.g., 1234.567(89) = 1234.567 ± 0.089). Where the error includes a decimal point, the decimal point is retained inside the parentheses. When the uncertainty is not explicitly indicated, the value is given to the

estimated number of significant figures. As in the tables of Herzberg,¹ T_0 values are given to the center of multiplet structure. For doublet states, the two components differ by $\pm A$ (the spin-orbit splitting constant), and the energy difference is measured from the average of the two bands, whereas for triplet states the three components fall at 0, $\pm A$ with respect to the position from which the band energy is measured. This convention is also followed here unless specific states (e.g., $^2\Pi_{3/2}$) are given. In matrix isolation absorption and many laser excitation studies only the lowest component of the ground electronic state is accessible. Except for transitions with relatively small values of A , this is also likely to be true for studies using cooled molecular beams. Often these latter studies give T_0 values from the lowest energy component with a precision better than that to which A is known.

The wavelength range (nm) in which various electronic transitions have been observed is also tabulated. This range is a composite of the values typical of absorption and emission observations. Laser-excited fluorescence studies often include both excitation and resolved emission measurements. Since the position of the band origin is given, ambiguity should not arise. For information on the range in which the band system is observed for a given type of measurement, see the original literature cited for that measurement technique.

The format of the vibrational tables is similar to that used in the earlier compilations. The vibrational numbering convention is that used by Herzberg.¹ Within a given symmetry species, vibrations are numbered starting with the highest frequency. The same convention is followed for deuterated species. Therefore, a given type of vibration may be numbered differently for the deuterated than for the unsubstituted molecule. For triatomic molecules, the bending vibration is always designated as ν_2 . For aromatic molecules, an alternate vibrational numbering scheme developed by Wilson¹⁹ has often been used in the literature. Where both the Herzberg and the Wilson numbering schemes have been used for the published data, the Herzberg numbering is adopted, and the Wilson numbering is sometimes shown in parentheses. For a few species, only the Wilson numbering has been used. To avoid confusion, this is retained in the present tables, and the use of the Wilson numbering is indicated in a footnote. Where possible, the values of $\Delta G(\frac{1}{2})$, the separation between the $v=0$ and $v=1$ levels for the vibration of interest, have been used. The expression of uncertainties is similar to that described for electronic band origins. Where vibrational frequencies have been determined with a precision greater than two decimal places, the tabulated values have been rounded off. If a bending fundamental is split by Renner-Teller interaction, the position of the unperturbed fundamental is given. Where specific components of such a split fundamental have been studied, they may also be listed, with the transition designated in a footnote. For a more complete treatment of the Renner effect and definitions of the

parameters included in many of these footnotes, see the discussion by Herzberg¹ and the references cited for the molecule of interest. A few of the species in these tables possess out-of-plane vibrations which have resolved inversion splitting structure. For these, the specific component for which the vibrational frequency is reported is designated in a footnote. Relative intensities of vibrational bands are dependent on the technique used for the measurement. When possible, the relative intensities of *ground-state infrared absorptions* are included. It is not feasible to give the corresponding relative intensities for other types of observation. Relative intensity abbreviations include:

vw	very weak
w	weak
m	medium
s	strong
vs	very strong
sh	shoulder
br	broad

Where radiative lifetimes have been measured, they are cited following the vibrational energy level table for the appropriate electronic state. τ_0 , the radiative lifetime of the vibrationless transition, is given whenever possible. If the lifetime is accessible only for excited vibrational states, the subscripts give the vibrational quantum numbers of the observed band.

When spin-orbit splitting occurs and the splitting constant, A , is known, it is included in the compilation.

Finally, as an aid in the recognition of vibrational bands and electronic band systems observed with comparatively high resolution, the principal rotational constants are summarized. Where possible, the values associated with the vibrationless transition (A_0 , B_0 , C_0) are given. Occasionally these values have not been determined, and the subscript gives the vibrational quantum numbers appropriate to the band for which the rotational constants have been measured. These constants are truncated at three decimal places. Often a far more detailed set of molecular constants, with much greater precision, has been derived from the analysis of high resolution spectra. Microwave spectroscopy is an important source of detailed, highly precise rotational data for molecules in their ground states. The references to the experimental literature, which are included in the compilation, should facilitate the location of such high resolution data.

5. Abbreviations

Many sophisticated laser techniques—frequently employing two or more laser beams—have been used for studies of transient molecules. The laser is frequently used both in the preparation of the transient molecule and in the detection scheme. For example, ions may be generated by multiphoton ionization and detected by absorption of radiation from a probe laser. Often the developers of such techniques have

designated them by complicated acronyms. In these tables, an attempt has been made to avoid relatively lengthy and unfamiliar acronyms by designating only the generic type of detection, using the abbreviations defined below. (Velocity modulation, designated as a separate detection technique in the first of this series of data evaluations,² is widely used and is considered to be a measurement tool rather than a type of observation. The type of laser used for the absorption measurement in an infrared detection scheme employing velocity modulation is instead specified in these tables.)

AB	near infrared-visible-ultraviolet absorption
CC	color-center laser
CR	cavity ringdown
DL	diode laser absorption
DPI	depletion photoionization
DR	double resonance
ED	electron diffraction
EF	electron-excited fluorescence
EM	near infrared-visible-ultraviolet emission
ESR	electron spin resonance
FD	fluorescence depletion
HFD	high frequency deflection
IB	ion beam
ID	ion drift, ion depletion (see specific reference)
IR	infrared absorption (conventional or Fourier transform)
LD	laser difference frequency
LF	laser-excited fluorescence (excitation and resolved emission)
LMR	laser magnetic resonance
LS	laser Stark spectroscopy
MO	molecular orbital calculations
MPD	multiphoton dissociation
MPI	multiphoton ionization
MW	microwave and millimeter wave
ND	neutron diffraction
PD	electron photodetachment
PE	photoelectron spectroscopy
PEFCO	photoelectron-photon coincidence
T-PEFCO	threshold photoelectron-photon coincidence
PEPICO	photoelectron-photoion coincidence
PF	photofragment spectroscopy
PI	photoionization
PIFCO	photoion-photon coincidence
PIR	photoionization resonance
PRI	photoinduced Rydberg spectroscopy
Ra	Raman
SEP	stimulated emission pumping
TF	tunable far-infrared laser
TPE	threshold photoelectron spectroscopy, including ZEKE detection
UV	near infrared-visible-ultraviolet absorption and emission

6. Acknowledgment

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8. Tables

8.1. H_3^+ , H_3 , and Triatomic Dihydrides

H₃⁺

\tilde{X}	D _{3h}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1'	1	Ring breathing	3178.18 ^a	gas	IR,PI LD,EM	10,12,13 19,22
e'	2	Deformation	2521.42 ^a	gas	LD,IR	1,4,15,22

$$B_0=43.510; C_0=20.698 \quad \text{LD}^{1,4}\text{DL}^4\text{IR}^{10}\text{EM}^{22}$$

$$\text{H}_2\text{D}^+$$

\tilde{X}	C_{2v}					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Ring breathing	2992.51	gas	LD	3,6,11
	2	Deformation	2205.87	gas	LD,DL	7
b_2	3	Deformation	2335.45	gas	LD,DL	7

$$A_0=43.438(2); B_0=29.134; C_0=16.601 \quad \text{LD,MW}^{6,7,11}$$

D₂H⁺

\tilde{X}	C_{2v}					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Ring breathing	2736.98	gas	LD	5,11,14
	2	Deformation	1968.17	gas	DL	8,14
b_2	3	Deformation	2078.43	gas	DL	8,14

$\pi_0 = 36.199$, $D_0 = 21.389$, $S_0 = 15.573$ $L_0 = 12$

D₃

X	D_{3h}					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	Ring breathing	2300.84 ^a	gas	EM,DL	21
e'	2	Deformation	1834.67 ^a	gas	IB,DL	2,9

^aHot bands arising from ν_1 and ν_2 of H_3^+ have been observed,^{16,19} as have been the first¹⁷ and second^{18,20} overtones of ν_2 . Several of the corresponding hot bands and overtones of D_3^+ have also been reported.²¹

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H₃

Higher Rydberg states have been detected using photoionization and field ionization. The ionization limit observed for vibrationally and rotationally unexcited H₃ from its 2p ²A₂["] state is 29562.6(5).^{12,15,16} Ion depletion studies have also yielded frequencies for the ring breathing vibration of a number of these higher Rydberg states.²³ Near the lowest ionization threshold, predissociation has been found to be induced by very weak electric fields.²⁵ Rotational and vibrational interactions, autoionization, and predissociation in the np Rydberg manifold have been studied.²⁷

$$\begin{array}{lll}
 3d^2A'_1 & D_{3h} & \text{Structure: EM}^8 \\
 T_0^a = 18511 & \text{gas} & 3d - 2p^2 A''_2 \quad 568-615 \text{ nm} \\
 & & 3d - 3p^2 E' \quad 3891-4456 \text{ cm}^{-1} \\
 B_0 = 42.99; C_0 = 22.735 & \text{EM}^8 &
 \end{array}$$

$3d\ 2E''$	D_{3h}	Structure: EM^8
$T_o = 18409$	gas	$3d-2p^2 A_2''$ 568–615 nm
	$EM^8PF^{14,26}$	$3d-3p^2 E'$ 3891–4456 cm $^{-1}$

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	Ring breathing	3168 ^b	gas	PI	19,23
e'	2	Deformation	2518	gas	EM,PF	22

$$\begin{array}{lll}
 B_0 = 42.99; C_0 = 22.735 & \text{EM}^8 & \\
 \text{3d } ^2\text{E}' & \text{D}_{3h} & \text{Structure: EM}^8 \\
 \text{T}_0^a = 18037 & \text{gas} & 3d - 2p^2 A_2'' \text{ 568--615 nm} \\
 & \text{EM}^8 & 3d - 3p^2 E' \text{ 3891--4456 cm}^{-1} \\
 B_0 = 42.99; C_0 = 22.735 & \text{EM}^8 &
 \end{array}$$

$$\begin{array}{lll}
 3p\ ^2A_2'' & \text{D}_{3h} & \text{Structure: EM}^3 \\
 T_a^a = 17789 & \text{gas} & 3p^2 A_2'' - 2s^2 A_1' \ 556-574 \text{ nm} \\
 r=37(4) \text{ ns} & \text{gas} & \text{EM}^{10} \\
 B_o=47.45^\circ, C_o=23.495 & \text{EM}^8
 \end{array}$$

$2s\ ^2A_1'$	D_{3h}	Structure: EM ³
gas	EM ^{2,3} LF ⁷	$3p^2A_2'' - 2s^2A_1'$ 553–569 nm
	EM ^{4,21}	$3p^2E' - 2s^2A_1'$ 700–765 nm
Vib.	No.	Approximate type of mode
		cm ⁻¹
a'_1	1	Ring breathing
e'	2	Deformation
$B_0 = 23.09$; $C_0 = 11.544(6)$		EM ^{3,21}

^aMeasured with respect to lowest bound state, $2s\ ^2A_1'$. Structure observed¹ in the dissociation spectrum of H₂ has been reinterpreted⁵ as arising from the predissociation of H₃ ($2s\ ^2A_1'$) into H+H₂. Unstructured emission observed¹¹ between 190 and 280 nm, with a maximum near 230 nm, upon charge transfer between K and H₃⁺ or D₃⁺ has been attributed to transitions originating in bound Rydberg states of H₃ or D₃ and terminating in the dissociative ground-state continuum. When charge transfer with Cs was studied,²⁸ emission between 200 and 400 nm was detected for all four isotopic species of H₃. A double maximum for D₃ was interpreted as arising from emission to both sheets of the ground-state potential surface. Photo-fragment spectroscopy¹⁴ has placed the $2p\ ^2A_2''$ state 5.563(20) eV above the ground-state H+H₂ dissociation limit.

^bObserved for $N=1$ rotational level.

^cPredisassociated by vibronic interaction with the $2p\ ^2E'$ repulsive ground state; linewidth is approximately 15 cm⁻¹ for H₃ and 6 cm⁻¹ for D₃.²

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BeH₂

\tilde{X}	$D_{\infty h}$
Vib.	No.
	Approximate type of mode
	cm ⁻¹
Π_u	2
Σ_u^+	3

\tilde{X}	$D_{\infty h}$
Vib.	No.
	Approximate type of mode
	cm ⁻¹
Π_u	2
Σ_u^+	3

Reference

- ¹T. J. Tague, Jr. and L. Andrews, J. Am. Chem. Soc. **115**, 12111 (1993).

MgH₂

\tilde{X}	$D_{\infty h}$
Vib.	No.
	Approximate type of mode
	cm ⁻¹
Π_u	2
Σ_u^+	3

\tilde{X}	$D_{\infty h}$
Vib.	No.
	Approximate type of mode
	cm ⁻¹
Π_u	2
Σ_u^+	3

References

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TiH₂

\tilde{X}		C _{2v}	Structure: IR ¹			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	2	Bend	496.1T	Kr	IR	1
b ₂	3	TiH a-stretch	1435.5	Ar	IR	1,2
			1416.5vs	Kr	IR	1
			1412.1vs			

TiD₂

\tilde{X}		C _{2v}	Structure: LMR ⁵			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	2	Bend	376.5T	Kr	IR	1
b ₂	3	TiD a-stretch	1041.1	Ar	IR	1,2
			1028.1vs	Kr	IR	1
			1024.5vs			

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ZrH₂

\tilde{X}		C _{2v}	Structure: LMR ⁵			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b ₂	3	ZrH ₂ a-stretch	1518.6	Ar	IR	1

ZrD₂

\tilde{X}		C _{2v}	Structure: LMR ⁵			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b ₂	3	ZrD ₂ a-stretch	1092.5	Ar	IR	1

Reference

- ¹G. V. Chertihin and L. Andrews, J. Phys. Chem. **99**, 15004 (1995).

HfH₂

\tilde{X}		C _{2v}	Structure: LMR ⁵			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b ₂	3	HfH ₂ a-stretch	1629.1	Ar	IR	1
			1622.4			

HfD₂

\tilde{X}		C _{2v}	Structure: LMR ⁵			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b ₂	3	HfD ₂ a-stretch	1166.7	Ar	IR	1
			1161.0			

Reference

- ¹G. V. Chertihin and L. Andrews, J. Phys. Chem. **99**, 15004 (1995).

FeH₂

In an argon, krypton, or xenon matrix, three broad absorptions appear¹ between 400 and 450 nm. Irradiation at 440 nm results in photodecomposition, producing Fe+H₂.^{1,2}

 \tilde{X}

\tilde{X}		D _{>h}	Structure: LMR ⁵			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
3	2	Bend	335	Ar	IR	2
			322	Kr	IR	1
			323	Xe	IR	1
3	3	FeH a-stretch	1674.72	gas	LMR	5
			1660.8	Ar	IR	2,4
			1647	Kr	IR	1,2
			1636	Xe	IR	1,2

$$B_0 = 3.013 \text{ LMR}^5$$

FeD₂

In krypton and xenon matrices, three broad absorptions appear¹ between 400 and 450 nm, each slightly shifted from their FeH₂ counterparts. Irradiation at 440 nm results in the formation of Fe+D₂.^{1,2}

$$T_0 = 9530(180) \text{ gas PE}^3$$

 \tilde{X}

\tilde{X}		D _{>h}	Structure: LMR ⁵			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
3	2	Bend	235	Ar	IR	2
			232	Xe	IR	1
3	3	FeD a-stretch	1204.2	Ar	IR	2,4
			1195	Kr	IR	2
			1188	Xe	IR	1

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CoH₂Photodissociation into Co+H₂ was observed² on irradiation at 22000. $T_0 \approx 2600$ gas PE¹ $T_0 = 535(90)$ gas PE¹ \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	2	Bend	380	Ar	IR	2
	3	CoH a-stretch	1684.5	Ar	IR	2,3
			1647	Kr	IR	2

CoD₂ $T_0 \approx 2600$ gas PE¹ $T_0 = 440(60)$ gas PE¹ \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	3	CoD a-stretch	1223.4	Ar	IR	2,3
			1215	Kr	IR	2

References¹A. E. S. Miller, C. S. Feigerle, and W. C. Lineberger, J. Chem. Phys. **84**, 4127 (1986).²R. L. Rubinovitz, T. A. Cellucci, and E. R. Nixon, Spectrochim. Acta A **43**, 647 (1987).³W. E. Billups, S.-C. Chang, R. H. Hauge, and J. L. Margrave, J. Am. Chem. Soc. **117**, 1387 (1995).**NiH₂** $T_0 \approx 1600$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	2200T	gas	PE	1
			2007	Ar	IR	2
			1985	Kr	IR	2
<i>b</i> ₂	2	Bend	771	Ar	IR	2
	3	Asym. stretch	1969	Ar	IR	2
			1947	Kr	IR	2

NiD₂ $T_0 \approx 1600$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1445	Ar	IR	2
			1430	Kr	IR	2
<i>b</i> ₂	2	Bend	602	Ar	IR	2
	3	Asym. stretch	1426	Ar	IR	2
			1411	Kr	IR	2

References¹A. E. S. Miller, C. S. Feigerle, and W. C. Lineberger, J. Chem. Phys. **84**, 4127 (1986).²S. Li, R. J. Van Zee, W. Weltner, Jr., M. G. Cory, and M. C. Zerner, J. Chem. Phys. **106**, 2055 (1997).**PtH₂** \bar{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			2349T	Ar	IR	1
			2314T	Kr	IR	1
			2168T	Ar	IR	1
			2158T	Kr	IR	1

References¹S. Li, H. A. Weimer, R. J. Van Zee, and W. Weltner, Jr., J. Chem. Phys. **106**, 2583 (1997).**ZnH₂** \bar{X} D_{oh}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_u	2	Bend	630.9	Ar	IR	1,2
			625.3	Kr	IR	1
Σ_u^+	3	ZnH ₂ a-stretch	1870.8	Ar	IR	1,2
			1861.0	Kr	IR	1

ZnD₂ \bar{X} D_{oh}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_u	2	Bend	454.4	Ar	IR	2
Σ_u^+	3	ZnD ₂ a-stretch	1357.2	Ar	IR	1,2
			1350.9	Kr	IR	1

References¹Z. L. Xiao, R. H. Hauge, and J. L. Margrave, High Temp. Sci. **31**, 59 (1991).²T. M. Greene, W. Brown, L. Andrews, A. J. Downs, G. V. Chertihin, N. Runeberg, and P. Pyykkö, J. Phys. Chem. **99**, 7925 (1995).**CdH₂** \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_u	2	Bend	604.6	Ar	IR	1
			601.7			
Σ_u^+	3	CdH ₂ a-stretch	1753.5	Ar	IR	1

CdD₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_u	2	Bend	434.2	Ar	IR	1
			432.5			
Σ_u^+	3	CdD ₂ a-stretch	1264.4	Ar	IR	1

References

¹T. M. Greene, W. Brown, L. Andrews, A. J. Downs, G. V. Chertihin, N. Runeberg, and P. Pyykkö, *J. Phys. Chem.* **99**, 7925 (1995).

HgD₂ \tilde{X} D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_u	2	Bend	555.2	Ar	IR	1
			553.3			
	3	Asym. stretch	555.8	D ₂	IR	1
			553.3			
			561.3	N ₂	IR	2
			1365.4	Ar	IR	1
Σ_u^+	3	Asym. stretch	1363.6			
			1362.8			
			1368.5	D ₂	IR	1
			1395.6	N ₂	IR	2

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¹N. Legay-Sommaire and F. Legay, *Chem. Phys. Lett.* **207**, 123 (1993).

²N. Legay-Sommaire and F. Legay, *J. Phys. Chem.* **99**, 16945 (1995).

NbH₂ \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1611m	Ar	IR	1
	2	Bend	519ms	Ar	IR	1
b_2	3	Asym. stretch	1569s	Ar	IR	1

NbD₂ \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1154m	Ar	IR	1
	2	Bend	373ms	Ar	IR	1
b_2	3	Asym. stretch	1128s	Ar	IR	1

Reference

¹R. J. Van Zee, S. Li, and W. Weltner, Jr., *J. Chem. Phys.* **102**, 4367 (1995).

HgH₂ \tilde{X} D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_u	2	Bend	773.0	Ar	IR	1
			771.1			
	3	Asym. stretch	769.7			
			769.2	Kr	IR	1
			772.5	H ₂	IR	1
			781.0	N ₂	IR	2
			1895.8	Ar	IR	1
			1895.4			
			1893.1	Kr	IR	1
			1891.2			
			1889.2			
			1902.7	H ₂	IR	1
			1943.7	N ₂	IR	2

UH₂ \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1406.1	Ar	IR	1
			1399.0			
b_2	3	Asym. stretch	1392.2			
			1370.7	Ar	IR	1
			1365.3			
			1360.6			

Reference

¹P. F. Souter, G. P. Kushto, L. Andrews, and M. Neurock, *J. Phys. Chem. A* **101**, 1287 (1997).

UD₂

		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1003.5 998.3 992.5	Ar	IR	1
<i>b</i> ₂	3	Asym. stretch	978.7 975.7 972.5	Ar	IR	1

Reference

¹P. F. Souter, G. P. Kushto, L. Andrews, and M. Neurock, J. Am. Chem. Soc. 119, 1682 (1997).

AlH₂

$\tilde{A}^2B_1(\Pi)$ D_{∞h} Structure: AB¹
 $T_0 < 15200$ gas AB¹ $\tilde{A}-\tilde{X}$ 658.4 nm
 Other bands were also observed, but their analysis has not been reported.
 There is evidence for a predissociation limit at 15450.
 B_0 (cm⁻¹) = 3.57 AB¹

		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	AlH s-stretch	1770.0m 1766vw	Ar Kr	IR IR	3-5 2
	2	Bend	766.4m 760m	Ar Kr	IR IR	4,5 2
<i>b</i> ₂	3	AlH a-stretch	1806.6vs 1799w	Ar Kr	IR IR	3-5 2

$A_0 = 13.6$; $B_0 = 4.4$; $C_0 = 3.3$ AB¹

AID₂

		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	AID s-stretch	1278.9 1275vw	Ar Kr	IR IR	3,4 2
	2	Bend	554.1 560m	Ar Kr	IR IR	4 2
<i>b</i> ₂	3	AID a-stretch	1325.0 1320w	Ar Kr	IR IR	3,4 2

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⁴P. Pullumbi, C. Mijoule, L. Manceron, and Y. Bouteiller, Chem. Phys. 185, 13 (1994).

⁵P. Pullumbi, Y. Bouteiller, and L. Manceron, J. Chem. Phys. 101, 3610 (1994).

GaH₂

		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1727.6wm 1730.8	Ar Kr	IR IR	1,2 1
	2	Bend	740.1wm	Ar	IR	2
<i>b</i> ₂	3	Asym. stretch	1799.2vs 1796.4	Ar Kr	IR IR	1,2 1

GaD₂

		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1244.7 1244.2wm	Ar Kr	IR IR	1,2 1
	2	Bend	528	Ar	IR	2
<i>b</i> ₂	3	Asym. stretch	1303.3 1302.4s	Ar Kr	IR IR	1,2 1

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¹Z. L. Xiao, R. H. Hauge, and J. L. Margrave, Inorg. Chem. 32, 642 (1993).

²P. Pullumbi, C. Mijoule, L. Manceron, and Y. Bouteiller, Chem. Phys. 185, 13 (1994).

InH₂

		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1548.6wm	Ar	IR	1
	2	Bend	607.4wm	Ar	IR	1
<i>b</i> ₂	3	Asym. stretch	1615.6vs	Ar	IR	1

InD₂

		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1115.3	Ar	IR	1
	2	Bend	429.1	Ar	IR	1
<i>b</i> ₂	3	Asym. stretch	1165.8	Ar	IR	1

Reference

¹P. Pullumbi, C. Mijoule, L. Manceron, and Y. Bouteiller, Chem. Phys. 185, 13 (1994).

SiH_2^+

$\tilde{A} \ ^2B_1(\text{II})$	$\tilde{X} \ ^3B_1$	C_{2v}	Structure: ESR ^{4–6} AB ⁷ LMR ^{15,17} IR ^{17,26}
gas PF ^{1,2}			
$\tilde{A}-\tilde{X} \ 567-659 \text{ nm}$			
Predissociation into $\text{Si}^+ + \text{H}_2$ and into $\text{SiH}^+ + \text{H}$ was observed.			
$B=3.956(1) \text{ PF}^1$			
$\tilde{X} \ ^2A_1$	C_{2v}		
$B=5.094(2); C=3.772(4)$	PF ¹		

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 CH_2

$4p$	C_{2v}				
$T_0=74254$	gas	MPI ³⁷			
\tilde{D}					
$T_0=71592 \text{ gas AB}^1\text{MPI}^{36}$					
$\tilde{D}-\tilde{X} \ 139.7 \text{ nm}$					
\tilde{C}					
$T_0=70917 \text{ gas AB}^1\text{MPI}^{36}$					
$\tilde{C}-\tilde{X} \ 141.0 \text{ nm}$					
$3d \ ^3A_2$	C_{2v}	Structure: AB ⁷			
$T_0=70634 \text{ gas AB}^1\text{MPI}^{36}$		$3d^3A_2-\tilde{X} \ 141.5 \text{ nm}$			
Diffuse. First member of Rydberg series converging to 83851. Higher members observed (AB ²) at 76553, 79241, and 80688.					
$B_0=6.89^{\text{a}}$	AB ¹				
$3p$	C_{2v}				
$T_0=64126$	gas	MPI ³⁷			
$\tilde{c} \ ^1A_1$	C_{2v}				
gas AB ³			$\tilde{c}-\tilde{a} \ 330-362 \text{ nm}$		
$\tilde{b} \ ^1B_1^{\text{b}}$	C_{2v}	Structure: AB ^{3,29}			
$T_0=10255(20) \text{ gas AB}^{1,3,27,44}\text{LMR}^{21}\text{LF}^{33,38,40,42,43}$		$\tilde{b}-\tilde{a} \ 465-920 \text{ nm}$			

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	Bend	570T	gas	AB	3

$\tau(0,14,0)=4.6(1)\mu\text{s LF}^{39}$
 $\tau(0,16,0)=1.3(3)\mu\text{s LF}^{11}$
 $3.8(3)\mu\text{s LF}^{39}$
 $B_0=7.74 \text{ AB}^1$
 Barrier to linearity=1617²⁹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
$\tilde{a} \ ^1A_1^{\text{b}}$	1	CH ₂ s-stretch	2806.01(7)	gas	LF,LD,IR	10,20,31,40

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CH ₂ s-stretch	2806.01(7)	gas	LF,LD,IR	10,20,31,40
	2	Bend	1352.6	gas	AB,LF	3,27,28,40

$\tau \approx 18 \text{ s}^{\text{c}}$
 $A_0=20.118(2); B_0=11.205(2); C_0=7.069(2) \text{ AB}^{3,27,28}$
 Barrier to linearity=8600(400) LF⁴⁵

$\tilde{X} \ ^3B_1$	C_{2v}	Structure: ESR ^{4–6} AB ⁷ LMR ^{15,17} IR ^{17,26}
Vib.	No.	Approximate type of mode
a_1	2	Bend
		963.10
b_2	3	CH ₂ a-stretch
		3190(5) ^d

$A_0=73.811; B_0=8.450; C_0=7.184 \text{ IR}^{25}$
 Barrier to linearity=1931(30)²⁶

 CD_2

$4p$	C_{2v}
$T_0=74228$	gas MPI ³⁷

\tilde{D}	$T_0=70947 \text{ gas AB}^1\text{MPI}^{36}$	$\tilde{D}-\tilde{X} \ 140.95 \text{ nm}$
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\tilde{C}	$T_0=71510 \text{ gas AB}^1\text{MPI}^{36}$	$\tilde{C}-\tilde{X} \ 139.8 \text{ nm}$
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$3d \ ^3A_2$	C_{2v}	Structure: AB ⁷
$T_0=70591.7 \text{ gas AB}^1\text{MPI}^{36}$		$3d^3A_2-\tilde{X} \ 141.6 \text{ nm}$
$B_0=3.595 \text{ AB}^1$		

$3p$	C_{2v}
$T_0=64082$	gas MPI ³⁷
$\tilde{b} \ ^1B_1^{\text{b}}$	C_{2v}
gas LF ^{13,41}	

$\tilde{b} \ ^1B_1^{\text{b}}$	C_{2v}
gas LF ^{13,41}	
$\tau(0,16,0)=6.0(7)\mu\text{s LF}^{13}$	

$\tilde{a} \ ^1A_1^{\text{b}}$	C_{2v}
$T_0=3140(50) \text{ gas PE}^{23}\text{LF}^{41}\text{SEP}^{41}$	

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	Bend	1005(1)	gas	LF	13

$A_0=11.37(32); B_0=5.476(48); C_0=3.701(45) \text{ LF}^{41}\text{SEP}^{41}$

 $\tilde{X} \ ^3B_1$

$\tilde{X} \ ^3B_1$	C_{2v}
Vib.	No.

a_1	2	Bend	752.37	gas	DL	19
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$A_0=37.787; \frac{1}{2}(B+C)_0=3.962; \frac{1}{2}(B-C)_0=0.267 \text{ LMR}^{18,22}\text{MW}^{46}$

^aValue given for ¹³CH₂.

^bThe $\tilde{a} \ ^1A_1^{\text{b}}$ and $\tilde{b} \ ^1B_1^{\text{b}}$ states are perturbed by strong Renner-Teller interaction.^{13,14,29,44} They are also strongly perturbed by interaction with the $\tilde{X} \ ^3B_1$ state.^{27,28,35}

^cCalculated value.²¹

^dFrom analysis of perturbations involving combination bands.³¹

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SiH₂

$\tilde{A}^1B_1^a$ C_{2v} Structure: AB^{1,2}
 $T_0 = 15530.4(5)$ gas AB^{1,2}LF^{10,11,13} $\tilde{A}-\tilde{X}$ 480–650 nm
 Onset of predissociation into Si (¹D) + H₂ near 21450.¹¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1990	gas	LF	13
	2	Bend	854	gas	AB,LF	1,11,13

$\tau_0 = 1.10(17)\mu s$ gas LF^{6,7,11}
 $A_0 = 17.75$; $B_0 = 4.9$; $C_0 \approx 2.8$ AB²
 Barrier to linearity $\approx 8000^3$

$\tilde{\alpha}^3B_1$ C_{2v}
 $T_0 = 7340(240)^c$ gas PI⁸
 Barrier to predissociation into Si (³S) + H₂ between 17070 and 17690.¹¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1978.51 ^d	gas	LF	10,13,15
			1964 ^d	Ar	IR	5
	2	Bend	999.03	gas	AB,LF	2,4,9,13
<i>b</i> ₂	3	Asym. stretch	995	Ar	IR	5
			1954 ^e	gas	LF	10,13,15
			1973	Ar	IR	5

$A_0 = 8.099$; $B_0 = 7.024$; $C_0 = 3.703$ AB²DL⁹

SiD₂

$\tilde{A}^1B_1^a$ C_{2v}
 $T_0 = 15539.875(2)$ gas LF^{11,14} $\tilde{A}-\tilde{X}$ 463–652 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	Bend	618.55	gas	AB,LF	1,11,14

$A_0 = 9.629$; $B_0 = 2.456$; $C_0 = 1.926$ LF¹⁴

$\tau_0 = 0.93(38)\mu s$ gas LF¹¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1427 ^d	Ar	IR	5
	2	Bend	731 ^f	gas	LF	11
			720	Ar	IR	5
<i>b</i> ₂	3	Asym. stretch	1439	Ar	IR	5

$A_0 = 4.334$; $B_0 = 3.519$; $C_0 = 1.919$ LF¹⁴

^aThe $\tilde{A}^1B_1^a$ and \tilde{X}^1A_1 states are perturbed by strong Renner-Teller interaction.³ The combined effects of Renner-Teller and spin-orbit interaction have been considered in detail by Ref. 12.

^bExtrapolated values.²

^cPossibly 6290(240).⁸

^dIn Fermi resonance with $2\nu_2$, observed for SiH₂ at 2005.7 (gas) and 1993 (Ar) and for SiD₂ at 1445 (Ar).

^e $\frac{1}{2}(2\nu_3)$.

^f ω .

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GeH₂

\tilde{A}^1B_1		C _{2v}	Structure: LF ⁴			
			$\tilde{A}-\tilde{X}$ 489–750 nm			
Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
a_1	1	Sym. stretch	1798.4	gas	LF	4
	2	Bend	783.0	gas	LF	2–4
$\tau_0=2.29(7)\mu s$		gas	LF ^{3,4}			
$A_0=16.415(21)$			$B_0=4.518(3)$	$C_0=3.460(4)$	LF ⁴	
\tilde{X}^1A_1		C _{2v}				
Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
a_1	2	GeH stretch	1887wm	Ar	IR	1
		GeH stretch	1864wm	Ar	IR	1
		Bend	913	gas	LF	2
			920wm	Ar	IR	1
$A_0=6.998(15)$			$B_0=6.531(8)$	$C_0=3.332(3)$	LF ⁴	

GeD₂

\tilde{A}^1B_1		C _{2v}	$\tilde{A}-\tilde{X}$ 535–700 nm			
Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
a_1	2	Bend	563	gas	LF	2,3
$\tau_0=2.5(5)\mu s$		gas	LF ³			
\tilde{X}		C _{2v}				
Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
a_1	2	GeD stretch	1338ms			
			1329vs	Ar	IR	1
			1325vs			
		Bend	659	gas	LF	2
			658m	Ar	IR	1

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NH₂⁺

\tilde{b}^1B_1 C_{2v}
 $T_0=20490(160)^a$ gas PE¹

Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
a_1	2	Bend	920(150)	gas	PE	1

\tilde{a}^1A_1 C_{2v}
 $T_0=10530(80)$ gas PI²

Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
a_1	1	Sym. stretch	2900(50)	gas	PE	1
	2	Bend	1350(50)	gas	PE	1

\tilde{X}^3B_1 D_{∞h}^b Structure: LD⁴

Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
Π_u^-	2	Bend	439T	gas	LD	4
	3	Asym. stretch	3359.94	gas	LD	3,4
$B_0=7.959$ LD ⁴						

ND₂⁺

\tilde{a}^1A_1 C_{2v}

Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
a_1	1	Sym. stretch	2210(50)	gas	PE	1
	2	Bend	940(50)	gas	PE	1

\tilde{X}^3B_1 D_{∞h}^b

Vib.	No.	Approximate sym.	cm ⁻¹	Med.	Type	Refs.
Π_u^-	2	Bend	330(30)H	gas	PE	1

^aCorrected for revision² of first adiabatic ionization potential of NH₂.

^bQuasilinear; barrier to linearity < $\frac{1}{2}\nu_2$.⁴

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NH₂

Rydberg series with members at 93054, 95753, 97193, and 98049, converging to NH₂⁺ ($\tilde{A}^{-1}A_1$) at 100410. PI¹⁹

$\tilde{A}^{-2}A_1(\text{II}_u)$ ^a		C _{2v}	Structure: AB ^{1,4}
T ₀ = 11122.6	gas	AB ^{1,8,22} LF ^{6,21,25,26} EM ²⁵	$\tilde{A}-\tilde{X}$ 342–2700 nm
	Ne,Ar,Kr,Xe ^b	AB ^{2,3,5,27}	$\tilde{A}-\tilde{X}$ 344–880 nm
N ₂ ^b		AB ⁵	$\tilde{A}-\tilde{X}$ 480–620 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	3325	gas	AB	1
	2	Bend	633	gas	AB	1

$$\tau_{090\Sigma}=10.0(1.7)\mu\text{s} \quad \text{gas LF}^7$$

$$\tau_{080\Pi}=10(3)\mu\text{s} \quad \text{gas LF}^{17}$$

Approximate ν^2 dependence.^{7,17} In another LF study,¹² τ varied from 25 to 46 μs for relatively unperturbed rotational sublevels, and there was a weaker $\sim 100 \mu\text{s}$ component associated with levels which are substantially perturbed.

$$B_0=8.78 \quad \text{AB}^1$$

Barrier to linearity = 730¹⁴

$\tilde{X}^{-2}B_1$ ^a		C _{2v}	Structure: AB ¹ IR ²⁹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	3219.37	gas	LF,EM	6,15,16
				LD,IR		21,23
	2	Bend	3220w ^c	N ₂	IR	5
			1497.32	gas	UV,LF	1,6,8–10
					LMR,IR	13,20
			1499m	N ₂	IR	5
<i>b</i> ₂	3	Asym. stretch	3301.11	gas	LD,LF,IR	16,21,23

$$A_0=23.693; B_0=12.952; C_0=8.173 \quad \text{AB}^{1,8}\text{LMR}^{13} \quad \text{IR}^{20,29} \quad \text{MW}^{28}$$

Barrier to linearity = 12024¹⁴

ND₂

$\tilde{A}^{-2}A_1(\text{II}_u)$ ^a		C _{2v}	
gas	AB ¹		$\tilde{A}-\tilde{X}$ 500–680 nm
Ar	AB ²⁷		$\tilde{A}-\tilde{X}$ 380–825 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	2520T	gas	AB	1
	2	Bend	430	gas	AB	1

$$B_0=4.41 \quad \text{AB}^1$$

$\tilde{X}^{-2}B_1$ ^a		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	Bend	1108.75	gas	LMR	11,18

$$A_0=13.342; B_0=6.488; C_0=4.290 \quad \text{AB}^{1,18}\text{LMR}^{11}\text{MW}^{24}\text{IR}^{29}$$

^aThe $\tilde{A}^{-2}A_1$ and $\tilde{X}^{-2}B_1$ states are perturbed by strong Renner-Teller interaction.

^bA detailed comparison of the argon-matrix data with gas-phase data has been given in Refs. 22 and 27. Rotational structure is resolved in the rare-

gas matrices. In nitrogen,⁵ bands are very broad and red-shifted by approximately 400 cm⁻¹, with no evidence for rotational structure.

^cAssigned⁵ in matrix studies to ν_3 . Gas-phase observation of ν_1 at 3219.37 cm⁻¹ and demonstration¹⁶ that ν_1 is more intense than ν_3 dictate reassignment to ν_1 .

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PH₂

\tilde{A}	2A_1 ^a	C _{2v}	Structure: AB ⁴
$T_0 = 18276.569$	gas	AB ^{1,4,6} EM ^{2,3,5} LF ^{22,23}	$\tilde{A}-\tilde{X}$ 360–880 nm
18215(4)	Ar	AB ^{13,19}	$\tilde{A}-\tilde{X}$ 405–550 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	Bend	949.12	gas	UV	2,6
			949(7)	Ar	AB	13,19

$\tau = 4(1)\mu s$ gas LF^{11,20}EM¹²
 $A_0 = 20.414(5)$; $B_0 = 5.607$; $C_0 = 4.293$ AB^{4,6}EM⁵LF^{22,23}
 Barrier to linearity = 6840⁷

\tilde{X}	2B_1 ^a	C _{2v}	Structure: AB ⁴
Vib.	No.	Approximate type of mode	cm ⁻¹
a_1	1	Sym. stretch	2310(2)
	2	Bend	1101.91

AB^{4,16}LMR^{8,14,15}MW^{17,18}LF²²
 Barrier to linearity = 25100⁷

PD₂

\tilde{A}	2A_1 ^a	C _{2v}	
$T_0 = 18282.1$	gas	AB ¹ EM ^{2,3}	
		$\tilde{A}-\tilde{X}$ 360–880 nm	
Vib.	No.	Approximate type of mode	cm ⁻¹
a_1	2	Bend	689.5
			665(25)

\tilde{X}	2B_1 ^a	C _{2v}	
Vib.	No.	Approximate type of mode	cm ⁻¹
a_1	2	Bend	795.5
			797w
		Ar	IR

$A_0 = 4.857(2)$; $B_0 = 4.044(4)$; $C_0 = 2.180(2)$ AB⁹

^aThe \tilde{A} 2A_1 and \tilde{X} 2B_1 states are perturbed by strong Renner-Teller interaction.

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MARILYN E. JACOX

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H₂O⁺

\tilde{B}	2B_2	C _{2v}
$T_0 = 36757(12)$	gas	PE ⁹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	2968 ^a	gas	PE	9
	2	Bend	1596 ^a	gas	PE	9

\tilde{A}	$^2A_1(\Pi_u)$ ^b	D _{∞h}	Structure: PE ^{4,7} EM ⁷
$T_{070} = 13409.3$	gas	EM ^{1,5,17} AB ¹⁴	$\tilde{A}-\tilde{X}$ 400–750 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	3547(16)	gas	PE	9
	2	Bend	876.8 ^b	gas	EM,PE	5,9

$\tau = 10.5(1.0)\mu s$ gas EF⁶
 $B_{070} = 8.57$ FM⁵

\tilde{X}	2B_1 ^b	C _{2v}	Structure: EM ^{5,7} LMR ⁸ LD ¹¹ CC ¹⁵
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	3212.86	gas	PE,LD	9,11
			3182.7	Ne	PI,CC	13,15
b_2	2	Bend	1408.42	gas	EM,PE	1,5,9
			1401.7	Ne	DL	12
b_2	3	Asym. stretch	3259.04	gas	LD,CC	11,15
			3219.5	Ne	IR	16

$A_0 = 29.036(2)$; $B_0 = 12.423$; $C_0 = 8.469$ LMR⁸LD¹¹DL¹²CC¹⁵

Barrier to linearity $\leq 9187^{7,17}$

D₂O⁺

\tilde{B} 2B_2 C_{2v}
 $T_0 = 37430(50)$ gas PE^{2,4}
 $38498(12)$ gas PE⁹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	2282 ^a	gas	PE	2,9
	2	Bend	1099 ^a	gas	PE	9

\tilde{A} $^2A_1(\Pi_u)$ ^b D_{∞h}
 $T_{050} = 10456(30)$ gas PE^{2,4} EM¹⁰ $\tilde{A}-\tilde{X}$ 490–670 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	2531(8)	gas	PE	9
	2	Bend	640(9)	gas	PE	9

τ is ~12% greater than for H₂O⁺.³

\tilde{X} 2B_1 ^b C_{2v} Structure: EM¹⁰

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	2344(6)	gas	PE	2,4,9
			2326.7	Ne	IR	16
<i>a</i> ₂	2	Bend	1044.27(5)	gas	EM	10
			1040.5	Ne	IR	16
<i>b</i> ₂	3	Asym. stretch	2392.7	Ne	IR	16

$A_0 = 16.03$; $B_0 = 6.240(3)$; $C_0 = 4.407(3)$ EM¹⁰

^aBest fit of simulated photoelectron spectrum.

^bThe \tilde{A} $^2A_1(\Pi_u)$ and \tilde{X} 2B_1 states are perturbed by strong Renner-Teller interaction.

^cVibrational numbering uncertain.

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H₂S⁺

\tilde{B} 2B_2 C_{2v}
 $T_0 = 34770(160)$ gas PE²

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	2259T	gas	PE	5

\tilde{A} 2A_1 ^a C_{2v} Structure: EF³
 $T_0 = 18518$ gas EF^{1,3} PE^{2,5,6} $\tilde{A}-\tilde{X}$ 400–500 nm
 Predissociated above 23300 into H₂+S⁺.^{1,2}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	2600T	gas	PE	6
	2	Bend	950(5)	gas	PE	2,5,6

$\tau = 4.2(4)\mu s$ gas EF⁴
 $B_{020} = 5.03$ EF³
 Barrier to linearity $\cong 4600^1$

\tilde{X} 2B_1 ^a C_{2v} Structure: EF^{1,3}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	2490(5)	gas	PE	6
	2	Bend	1159.0	gas	EF,PE	3,6

$A_0 = 10.18(2)$; $B_0 = 8.63(1)$; $C_0 = 4.60(6)$ EF³

D₂S⁺

\tilde{A} 2A_1 C_{2v}
 $T_0 = 18574$ gas EF³ $\tilde{A}-\tilde{X}$ 400–500 nm
 $B_{030} = 2.46$ EF³

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	Bend	838.6	gas	EF	3

$A_0 = 5.37(2)$; $B_0 = 4.32(1)$; $C_0 = 2.34(2)$ EF³

^aThe \tilde{A} 2A_1 and \tilde{X} 2B_1 states are perturbed by strong Renner-Teller interaction.

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H₂XeH

\tilde{X} D _{∞h}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_u	2	Bend	700.8	Xe	IR	1
Σ_u^+	3	Asym. stretch	1180.6 1165.9	Xe	IR	1,2

D₂XeD

\tilde{X} D _{∞h}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_u	2	Bend	513.5wT	Xe	IR	1
Σ_u^+	3	Asym. stretch	856.2 845.8	Xe	IR	1

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8.2. Triatomic Monohydrides**Li₂H**

\tilde{B}'	gas	MPI ³	$\tilde{B}' - \tilde{X}$ 360–380 nm
\tilde{B}	C _{2v}	Structure: MPI ^{1,2}	
T ₀ =18940	gas	MPI ^{1,2}	$\tilde{B} - \tilde{X}$ 457–528 nm
Vib.	No.	Approximate type of mode	cm ⁻¹
a ₁	1		332
	2		195
b ₂	3		309

A₀=6.87; B₀=0.455; C₀=0.427 MPI^{1,2}

\tilde{X} C_{2v} Structure: MPI^{1,2}
A₀=13.250; B₀=0.750; C₀=0.710 MPI^{1,2}

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BeBeH

\tilde{X} C _{∞v}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	BeH stretch	2013.7	Ar	IR	1
Π	2	Bend	540.0	Ar	IR	1

BeBeD

\tilde{X} C _{∞v}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	BeD stretch	1504.2	Ar	IR	1

Reference

- ¹T. J. Tague, Jr. and L. Andrews, J. Am. Chem. Soc. **115**, 12111 (1993).

ZnZnH

\tilde{X}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	ZnH stretch	1657.6	Ar	IR	1

ZnZnD

\tilde{X}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		ZnD stretch	1198.6	Ar	IR	1

Reference

- ¹T. M. Greene, W. Brown, L. Andrews, A. J. Downs, G. V. Chertihin, N. Runeberg, and P. Pyykkö, J. Phys. Chem. **99**, 7925 (1995).

CdCdH

\tilde{X}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CdH stretch	1521.0	Ar	IR	1

CdCdD

\tilde{X}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CdD stretch	1098.0	Ar	IR	1

Reference

- ¹T. M. Greene, W. Brown, L. Andrews, A. J. Downs, G. V. Chertihin, N. Runeberg, and P. Pyykkö, J. Phys. Chem. **99**, 7925 (1995).

TiCH

\tilde{A} $^2\Pi$	C _{∞v}				
gas	LF ¹				
		$\tilde{A}-\tilde{X}$ 620–725 nm			

\tilde{X} $^2\Sigma^+$		C _{∞v}	Structure: LF ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	578(5)	gas	LF	1
Σ ⁺	3	TiC stretch	855(5)	gas	LF	1
$B_0=0.486$ LF ¹						

TiCD

\tilde{A} $^2\Pi$	C _{∞v}				
gas	LF ¹				
		$\tilde{A}-\tilde{X}$ 620–725 nm			
\tilde{X} $^2\Sigma^+$	C _{∞v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Π	2	Bend	442(5)	gas	LF
Σ ⁺	3	TiC stretch	815(5)	gas	LF
$B_0=0.416$ LF ¹					

Reference

¹M. Barnes, A. J. Merer, and G. F. Metha, J. Mol. Spectrosc. **181**, 168 (1997).

VCH

\tilde{A}	gas	LF ¹				
			$\tilde{A}-\tilde{X}$ 600–810 nm			
\tilde{X} $^3\Delta_1$	C _{∞v}		Structure: LF ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	564	gas	LF	1
Σ ⁺	3	VC stretch	838	gas	LF	1
Spin-orbit splitting=139 gas LF ¹						
$B_0=0.494$ LF ¹						

VCD

\tilde{A}	gas	LF ¹				
			$\tilde{A}-\tilde{X}$ 600–810 nm			
\tilde{X} $^3\Delta_1$	C _{∞v}		Structure: LF ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	442	gas	LF	1
Σ ⁺	3	VC stretch	813	gas	LF	1
$B_0=0.422$ LF ¹						

Reference

¹M. Barnes, P. G. Hajigeorgiou, R. Kasrai, A. J. Merer, and G. F. Metha, J. Am. Chem. Soc. **117**, 2096 (1995).

WCH

\tilde{C} $^2\Phi_{5/2}$	C _{∞v}				
T ₀ =14109.7	gas	LF ¹			
		$\tilde{C}-\tilde{X}$ 652–709 nm			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	
Π	2	Bend	554	gas	
Σ ⁺	3	WC stretch	862	gas	
$B_0=0.392$ LF ¹					

\tilde{B} $^2\Pi_{1/2}$	C _{∞v}				
T ₀ =13391.7	gas	LF ¹			
		$\tilde{B}-\tilde{X}$ 680–747 nm			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	
Π	2	Bend	505	gas	
Σ ⁺	3	WC stretch	819	gas	
$B_0=0.392$ LF ¹					

\tilde{A} $^2\Delta_{3/2}$	C _{∞v}				
T ₀ =12090.3	gas	LF ¹			
		$\tilde{A}-\tilde{X}$ 749–827 nm			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	
Π	2	Bend	489	gas	
Σ ⁺	3	WC stretch	936	gas	
$B_0=0.395$ LF ¹					

\tilde{a} $^4\Sigma^-$	C _{∞v}				
T ₀ =813	gas	LF ¹			
		Structure: LF ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	
Π	2	Bend	612(5)	gas	
Σ ⁺	3	WC stretch	971(5)	gas	
$B_0=0.41T$ LF ¹					

\tilde{X} $^2\Delta_{3/2}$	C _{∞v}				
		Structure: LF ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	
Π	2	Bend	660(5)	gas	
Σ ⁺	3	WC stretch	1006(5)	gas	
$B_0=0.408$ LF ¹					

WCD

$\tilde{C}^2\Phi_{5/2}$ C_{∞v}
 $T_0=14079.8$ gas LF¹

 $\tilde{C}-\tilde{X}$ 652–710 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	576	gas	LF	1
Σ^+	3	WC stretch	670T	gas	LF	1
$B_0=0.336$ LF ¹						

$\tilde{B}^2\Pi_{1/2}$ C_{∞v}
 $T_0=13416.4$ gas LF¹

 $\tilde{B}-\tilde{X}$ 680–746 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	450	gas	LF	1
Σ^+	3	WC stretch	809	gas	LF	1
$B_0=0.328(2)$ LF ¹						

$\tilde{A}^2\Delta_{3/2}$ C_{∞v}
 $T_0=12122.4$ gas LF¹

 $\tilde{A}-\tilde{X}$ 749–825 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	376	gas	LF	1
Σ^+	3	WC stretch	896	gas	LF	1
$B_0=0.337$ LF ¹						

$\tilde{a}^4\Sigma^-$ C_{∞v}
 $T_0=834$ gas LF¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	451(5)	gas	LF	1
Σ^+	3	WC stretch	930(5)	gas	LF	1
$B_0=0.347$ LF ¹						

Reference

¹M. Barnes, D. A. Gillett, A. J. Merer, and G. F. Metha, J. Chem. Phys. **105**, 6168 (1996).

ScNH

$\tilde{A}^2\Pi$ C_{∞v}
 $T_0=15023.033$ gas LF¹
 $A=102.23$ gas LF¹
 $B_0=0.424$ LF¹

 $\tilde{X}^2\Sigma^+$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	511(5)	gas	LF	1
Σ^+	3	ScN stretch	935(5)	gas	LF	1
$B_0=0.441$ LF ¹						

ScND

$\tilde{A}^2\Pi_{1/2}$ C_{∞v}
 $T_0=14952$ gas LF¹

 $\tilde{X}^2\Sigma^+$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	391(5)	gas	LF	1
Σ^+	3	ScN stretch	877(5)	gas	LF	1
$B_0=0.441$ LF ¹						

Reference

¹T. C. Steinle, J. Xin, A. J. Marr, and S. Beaton, J. Chem. Phys. **106**, 9084 (1997).

BeOH

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		BeO stretch	1245.5T	Ar	IR	1

Reference

¹C. A. Thompson and L. Andrews, J. Phys. Chem. **100**, 12214 (1996).

MgOH

$\tilde{X}^2\Sigma^+$ C_{∞v} Structure: MW^{1,3}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	188(2)H	gas	LF	2
Σ^+	3	MgO stretch	750(3)	gas	LF	2
$B_0=0.494$ MW ¹						

MgOD

$\tilde{X}^2\Sigma^+$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	130(2)H	gas	LF	2
$B_0=0.448$ MW ^{1,3}						

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³ B. P. Nuccio, A. J. Apponi, and L. M. Ziurys, J. Chem. Phys. **103**, 9193 (1995).

CaOH

\tilde{G} $^2\Pi$ $C_{\infty v}$
 $T_0=32630.2$ gas DR²²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	311	gas	DR	22
$B_0=0.347$ DR ²²						

\tilde{F} C_s
 $T_0=30215$ gas LF,MPI²¹

\tilde{E} $^2\Sigma^+$	$C_{\infty v}$	$E-\tilde{X}$ 326–335 nm
$T_0=29879$ gas LF,MPI ²¹		

\tilde{D} $^2\Sigma^+$	$C_{\infty v}$	Structure: LF ²¹	$D-\tilde{X}$ 335–355 nm
$T_0=28153$ gas LF,MPI ²¹			

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	412(2)	gas	LF,MPI	21
Σ^+	3	CaO stretch	674.0(1.5)	gas	LF,MPI	21
$B_0=0.364$ LF ²¹						

\tilde{C} $^2\Delta$	$C_{\infty v}$	$C-\tilde{X}$ 440–470 nm
$T_0=21896T$ gas LF ^{12,14}		

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	312	gas	LF	12
Σ^+	3	CaO stretch	528	gas	LF	12
$B_{010\Pi}=0.322$ LF ¹²						

\tilde{B} $^2\Sigma^+$	$C_{\infty v}$	Structure: LF ⁷	$\tilde{B}-\tilde{X}$ 555 nm
$T_0=18022.268(1)$ gas CL ² LF ^{4,6,10}			

Absorption maximum at 18236(15) in a krypton matrix.⁵

$B_0=0.339$ LF^{4,6}

\tilde{A} $^2\Pi$	$C_{\infty v}$	Structure: LF ^{3,20}	$\tilde{A}-\tilde{X}$ 600–650 nm
$T_0=15998.122$ gas CL ² LF ^{3,6,8,10,13,16,17,20}			

Absorption maximum at 16096(15) in a krypton matrix.⁵

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	2	Bend	361.36 ^a	gas	LF	13,16,18
Σ^+	3	CaO stretch	630.68 ^a	gas	LF	13

$A=66.818$ gas LF^{3,6,17}
 $\varepsilon\omega_2=-36.26$ gas LF^{13,16,18}
 $B_0=0.341$ LF^{3,6}

\tilde{X} $^2\Sigma^+$		$C_{\infty v}$	Structure: LF ^{3,7,20} MW ¹⁹
Vib. sym.	No.	Approximate type of mode	cm ⁻¹
Σ^+	1	OH stretch	3778
Π	2	Bend	352.93
Σ^+	3	CaO stretch	609.02
$B_0=0.334$		LF ^{3,4,6,12,16} MW ⁹ MODR ¹⁵	

CaOD

\tilde{G} $^2\Pi$	$C_{\infty v}$	$T_0=32628.3$	gas DR ²²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	267	gas	DR	22

$B_0=0.327$ DR²²

\tilde{C} $^2\Delta$	$C_{\infty v}$	$T_0=21907.128$	gas LF ^{12,14}	$\tilde{C}-\tilde{X}$ 440–470 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	235.38	gas	LF	12
Σ^+	3	CaO stretch	549	gas	LF	12

$A_0=11.494$ gas LF¹²

$B_0=0.294$ LF¹²

\tilde{B} $^2\Sigma^+$	$C_{\infty v}$	$T_0=18013.894(4)$	gas LF ⁷

$B_0=0.307$ LF⁷

\tilde{A} $^2\Pi$	$C_{\infty v}$	$T_0=15995.016$	gas LF ^{3,8,17,20}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	273.33	gas	LF	18
Σ^+	3	CaO stretch	623(2) ^b	gas	LF	3

$A_0=66.762$ LF¹⁷

$\varepsilon\omega_2=-26.96$ LF¹⁸

$B_0=0.309$ LF^{3,17}

\tilde{X} $^2\Sigma^+$	$C_{\infty v}$	$T_0=66.818$	gas LF ^{3,6,17} MW ¹⁹

^aThe Fermi resonance between ν_3 and $2\nu_2$ is analyzed by Ref. 13.

^b $\omega_1 + \frac{1}{2}x_{13}$.

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CaSH

\bar{C}^2A'		C_s		$\bar{C}-\bar{X}$ 597–647 nm			
$T_0=16082$	gas	LF ¹					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.	
a'	3	CaS stretch	312	gas	LF	1	
\bar{B}^2A''		C_s		$\bar{B}-\bar{X}$ 607–659 nm			
$T_0=15859.4638(15)$	gas	LF ¹ LS ³					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.	
a'	3	CaS stretch	320	gas	LF	1	
$A_0=10.434, B_0=0.143, C_0=0.141$ LS ³							
\bar{A}^2A'		C_s		Structure: LF ²			
$T_0=15380.2847(2)$	gas	LF ^{1,2}		$\bar{A}-\bar{X}$ 626–679 nm			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.	
a'	3	CaS stretch	318	gas	LF	1	
$A_0=9.091; B_0=0.147; C_0=0.145$ LF ²							
\bar{X}^2A'		C_s		Structure: LF ² MW ⁴			
$A_0=9.694; B_0=0.142; C_0=0.140$				LF ² MW ⁴			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.	
a'	3	CaS stretch	326	gas	LF	1	

$\tilde{X}^2\Sigma^+$		$C_{\infty v}$	Structure: LF ³			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	2	Bend	282(10)	gas	LF	3
	3	SrO stretch	510(10)	gas	LF	3
			470.6	Ar	IR	5

$B_0=0.225$ LF^{3,14}MW⁹

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ZnOH

\tilde{X}						
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		⁶⁴ ZnO stretch	649.6	Ar	IR	1,2

ZnOD

\tilde{X}						
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		⁶⁴ ZnO stretch	648.1	Ar	IR	1

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HNB

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	NH stretch	3693.7	Ar	IR	1,2
Π	2	Bend	432.5	Ar	IR	2
Σ^+	3	NB stretch	1829.6	Ar	IR	2

DNB

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	ND stretch	2777.4	Ar	IR	1,2
	3	NB stretch	1776.1	Ar	IR	2

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HBS⁺

\tilde{B} $2\Sigma^+$		$C_{\infty v}$
$T_0=38000(1000)$ gas PE ^{2,3}		
Vib.	No.	Approximate type of mode

\tilde{A} $2\Sigma^+$		$C_{\infty v}$	Structure: PE ^{2,3,6}			
$T_0=19827$ gas EM ¹ EF ⁴			$\tilde{A}-\tilde{X}$ 479–635 nm			
Vib.	No.	Approximate type of mode	cm^{-1}			
Σ^+	1	BH stretch	2214.8(4)	gas	EF	4
Σ^+	3	BS stretch	1050.9(4)	gas	EF	4

$\tau \geq 2300(200)$ ns gas EF⁵

$B_0=0.615^a$ EM¹

\tilde{X} $2\Pi_{3/2}$		$C_{\infty v}$	Structure: PE ^{2,3,6}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	BH stretch	2746.8(4)	gas	EF	4
Π	2	Bend	659(1)	gas	EF	4
Σ^+	3	BS stretch	984.1(4) ^b	gas	EF	4

$A_{010}=-321.4$, $\varepsilon\omega_2=-45(1)$.⁴

$B_0 = 0.576^a$ EM¹

DBS⁺

$\tilde{A}^2\Sigma^+$ C_{∞v}
 $T_0=19913$ gas EF⁴ $\tilde{A}-\tilde{X}$ 462–646 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	BD stretch	1706.6(4)	gas	EF	4
Σ^+	3	BS stretch	1011.1(4)	gas	EF	4

$\tilde{X}^2\Pi_{3/2}$ C_{∞v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	BD stretch	2071.1(4)	gas	EF	4
Σ^+	3	BS stretch	933.9(4) ^c	gas	EF	4

^aFrom analysis of bands originally attributed to BS and reassigned by Ref. 4 to HBS⁺.

^b75.9(4) in $\tilde{X}^2\Pi_{1/2}$ state.

^c937.4(4) in $\tilde{X}^2\Pi_{1/2}$ state.

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HCC

3pσ Rydberg state^a C_{∞v}
 $T_0=72100$ (1300) gas MPI³⁴

$T_0=51387(25)$ T Ar AB ⁸		195–160 nm				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CC stretch	2175(25)	Ar	AB	8
		Bend	630(25) ^b	Ar	AB	8

$\tilde{B}^2A'_1?$ C_s
 $T_0=51387(25)$ T Ar AB⁸ $\tilde{D}-\tilde{X}$ 250–313 nm

Approximately 20 emission bands of HCC have been assigned to transitions between a few common upper-state energy levels and HCC (\tilde{X}) with appreciable bending excitation. Analysis of these bands has yielded accurate vibrational and rotational constants for several highly excited ground-state vibrational levels. However, a definitive assignment of the upper-state energy levels has not been achieved.

$\tau \approx 120$ ns gas LF³⁶

$T_0 \leq 29360$ Ar AB^{2,8}

An absorption band system between 340 and 246 nm, with band spacings of approximately 2700, 1300 and 840 cm⁻¹, which has been observed on vacuum UV photolysis of C₂H₂ in an argon matrix, is tentatively attributed to HCC.

$\tilde{A}^2\Pi$ C_{∞v}
 $T_0^c=3692.61$ gas CC^{10,14}PE³²
3685.8 Ne AB³⁸
3732 Ar AB^{21,38}

In neon and argon matrices, a complicated absorption band system of HCC extends from approximately 3600 to 9000.^{21,38} This band system is extensively perturbed by high vibrational levels of the ground state. A few of the individual bands have been observed in the gas phase by color-center laser absorption,^{10,14,29} by high resolution emission spectroscopy,²⁴ or by time-resolved emission spectroscopy.³⁰ Because of the extensive perturbations and because of the high energy input in the gas-phase studies, high ground-state vibrational levels are prominent both in the gas phase^{10,14,19,26,27,32,37,40–42} and in neon and argon matrices.^{20,21,38}

Quasicontinuous 400–900 nm emission results on 136–110 nm photolysis of C₂H₂ or HCCBr in the gas phase.^{5,11,13,17} The fluorescence lifetimes vary from 6 to 20 μs.^{11,13,17} Unstructured emission from 1 to 5 μm has been detected¹⁸ upon 193 nm photolysis of gas-phase C₂H₂, with maximum intensity between 3600 and 5000. The HCC fluorescence resulting from the 193 nm photolysis of HCCBr extends from 500 nm to 5 μm,¹⁸ with lifetime increasing from ~5 μs near 500 nm to ~60 μs near 4000. Unstructured HCC emission between 400 and 500 nm has also been observed¹⁵ on vacuum UV irradiation of C₂H₂ isolated in the solid rare gases.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	3	CC stretch	1706.2T	Ne	AB	38
$\tilde{X}^2\Sigma^+$ C _{∞v}			Structure: MW ³¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CH stretch	3298.85T	gas	CC	29
Π	2	Bend	371.60 ^d	gas	DL	26,27
Σ^+	3	CC stretch	1840.57	gas	DL	23
			1835.5	Ne	IR	38
			1846.2m	Ar	IR	1,3,20,21

$A \approx 10$ IR¹⁴
 $B_{000}=1.457$ MW^{4,6,7,9} LMR¹²
 $B_{020}=1.451$ LMR²⁸

DCC

3pσ Rydberg state^a C_{∞v}
 $T_0=72100$ (1300) gas MPI³⁴

$T_0=51493(25)$ T Ar AB ⁸		194–170 nm				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CC stretch	2183(25)	Ar	AB	8
		Bend	520(25) ^b	Ar	AB	8

Fluorescence observed on laser excitation between 34500 and 40000 (250–290 nm) of gas-phase DCC from 2 to 10 μs after its formation by 193 nm photolysis of C₂D₂ has been attributed³⁵ to transitions of DCC between high vibrational levels of the ground state and an undetermined excited state, possibly 2²P.

$\tilde{A}^2\Pi$ C_{∞v}
 $T_0 < 3800$ Ar AB²¹

A complicated absorption band system extends to approximately 9700 in neon and argon-matrix studies of DCC.^{21,38} As for HCC, the band system is extensively perturbed by high vibrational levels of the ground state. A few

of the bands, some of which arise from excited ground-state vibrational energy levels, have been studied in the gas phase using infrared laser absorption.^{22,29,33}

$\tilde{X}^2\Sigma^+$	$C_{\infty v}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CD stretch	2537.1	Ne	IR	38
Σ^+	3	CC stretch	1743.18	gas	DL	25
			1739.6	Ne	IR	38
			1746.3m	Ar	IR	1,3,20,21

$B_0 = 1.203 \text{ MW}^{16}$

^aTentative assignment.

^bObserved band spacing; $2\nu_2$ if upper state is linear.

^cThe $\tilde{A}(000)$ band of HCC is strongly perturbed by high vibrational energy levels of the ground state which possess $^2\Pi$ vibronic symmetry. Accordingly, there is extensive mixing of the $^2\Pi$ vibronic levels in this spectral region, and designation of the band origin is approximate.^{10,14,38,41}

^dDerived from $(\nu_2 + \nu_3) - [(\nu_2 + \nu_3) - \nu_2]$. The detailed assignment of $(\nu_2 + \nu_3)$ is given in Ref. 27 and that of $(\nu_2 + \nu_3) - \nu_2$ in Ref. 26.

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AlOH

\tilde{B}		$\tilde{B}-\tilde{X}$ 230–240 nm			
$T_0 = 41747$	gas	MPI ⁴			
			760.3	gas	MPI 4
			581.0	gas	MPI 4

\tilde{A}		$\tilde{A}-\tilde{X}$ 227–255 nm			
$T_0 = 40073$	gas	MPI ⁴	Kr	AB ^{2,3}	$\tilde{A}-\tilde{X}$ 245–252 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		OH stretch	3258.4T	gas	MPI	4
			807.6	gas	MPI	4
			636.6	gas	MPI	4

$\tilde{X}^1\Sigma^+$ $C_{\infty v}$ Structure: MW⁵

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	OH stretch	3790	Ar	IR	1
	3	AlO stretch	895	gas	MPI	4
			810.3	Ar	IR	1

$B_0 = 0.525 \text{ MW}^5$

AlOD

\tilde{B}		$\tilde{B}-\tilde{X}$ 230–240 nm			
$T_0 = 41706$	gas	MPI ⁴			
			570.8 ^a	gas	MPI 4

\tilde{A}
 $T_0=40025$ gas MPI⁴

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			667.5	gas	MPI	4
			569.3	gas	MPI	4

$\tilde{X}^1\Sigma^+$ C_{∞v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	3	AlO stretch	795.2	Ar	IR	1

$B_0=0.473$ MW⁵

^aw.

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InOH

A broad absorption with maximum near 271 nm observed in a krypton matrix has been assigned² to InOH.

β^3A'' a C_s Structure: LF³
 $T_0=27188.00$ gas LF³ 345-377 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	829.6(4)	gas	LF	3
	3	InO stretch	496.9	gas	LF	3

$\tau > 2.3 \mu s$ gas LF³
 $A = 20.13; \frac{1}{2}(B+C) = 0.273$ LF³

α^3A' a C_s Structure: LF³
 $T_0=26914.34$ gas LF³ 345-377 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	820.41(3)	gas	LF	3

$\tau > 2.3 \mu s$ gas LF³
 $A = 26.70(3); \frac{1}{2}(B+C) = 0.271$ LF³

\tilde{X}^1A' C_s Structure: LF³MW⁵

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	378(2)	gas	LF	3
			421.8	Ar	IR	1
3		InO stretch	542(5)	gas	LF	3
			522.8	Ar	IR	1

$A_0=42.049; B_0+C_0=0.535; B_0-C_0=0.002$ LF³MW⁵

InOD

β^3A'' a C_s
 $T_0=27130.97$ gas LF³ 345-377 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	609.2(3)	gas	LF	3
	3	InO stretch	491.9(2)	gas	LF	3

$A=11.20; \frac{1}{2}(B+C)=0.255$ LF³

α^3A' a C_s
 $T_0=26862.42$ gas LF³ 345-377 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	604.1(2)	gas	LF	3
	3	InO stretch	489.5(2)	gas	LF	3

$A=13.73(7); \frac{1}{2}(B+C)=0.252$ LF³

\tilde{X}^1A' C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	301(5)	gas	LF	3
	3	InO stretch	524(9)	gas	LF	3
			595.7	Ar	IR	1

$A_0=20.637; B_0+C_0=0.490$ LF³MW⁵

^aComponents of the $\tilde{\alpha}^3\Pi$ state, split by strong Renner-Teller interaction.⁴

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HSiN

In an argon matrix, absorption maxima at 42000, 38800, 37600, and 28600 (238, 258, 266, and 350 nm) have been assigned¹ to HSiN.

\tilde{X}	$C_{\infty v}$					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	SiH stretch	2152.2m	Ar	IR	1
Σ^+	3	SiN stretch	1162.2s	Ar	IR	1

DSiN

\tilde{X}	$C_{\infty v}$					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	SiD stretch	1580.5wm	Ar	IR	1
Σ^+	3	SiN stretch	1145.4s	Ar	IR	1

Reference

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HCO⁺

\tilde{X}	$C_{\infty v}$	Structure: MW ^{2-4,15}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CH stretch	3088.74	gas	LD,CC	5,6,16,17
Π	2	Bend	829.72	gas	DL	10,11
Σ^+	3	CO stretch	2183.95	gas	DL	7,8,14

$$B_0 = 1.488 \text{ MW}^{2-4}$$

DCO⁺

\tilde{X}	$C_{\infty v}$					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CD stretch	2584.56	gas	DL	12
Π	2	Bend	647(25)	gas	PE	1,13
Σ^+	3	CO stretch	1904.06	gas	DL	9

$$B_0 = 1.201 \text{ MW}^{2-4}$$

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HCS⁺

\tilde{X}	$C_{\infty v}$	Structure: MW ⁶				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CH stretch	3141.68	gas	CC	4,5
Π	2	Bend	766.45	gas	DL	3

$$B_0 = 0.712 \text{ MW}^{1.2} \text{ DL}^3 \text{ CC}^{4.5}$$

DCS⁺

$$B_0 = 0.601 \text{ MW}^6$$

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HNC

\tilde{A} ^a	$T_0 = 32850$	gas	AB ⁷	$\tilde{A} - \tilde{X}$ 250–305 nm		
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CN stretch	1005	gas	UV	7

\tilde{X} C_{∞v} Structure: MW^{4,5,9,12}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	NH stretch	3652.66	gas	IR,EM	3,6,8,11
			3643.1	Ne	IR	10
			3620s	Ar	IR	2
			3583s N ₂	Ar	IR	1,2
			3567s	N ₂	IR	2
			464.24	gas	IR	8
Π	2	Bend	477s	Ar	IR	2
			535s N ₂	Ar	IR	1
			559s	N ₂	IR	2
			2023.86	gas	IR	8
Σ^+	3	NC stretch	2025.4	Ne	IR	10
			2029w	Ar	IR	2
			2032w N ₂	Ar	IR	1
			2035w	N ₂	IR	2

 $B_0 = 1.512 \text{ MW}^4 \text{ IR}^8$

DNC

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	ND stretch	2787.07	gas	IR	3,6
			2780.7	Ne	IR	10
			2769s	Ar	IR	2
			2733s N ₂	Ar	IR	1
			2728s	N ₂	IR	2
			374s	Ar	IR	2
Π	2	Bend	413s N ₂	Ar	IR	1
			432s	N ₂	IR	2
			1938.7	Ne	IR	10
			1940w	Ar	IR	2
Σ^+	3	NC stretch	1940w N ₂	Ar	IR	1
			1937w	N ₂	IR	2

 $B_0 = 1.273 \text{ MW}^4$ ^aTentative identification.

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HNSi

In an argon matrix, has a weak absorption maximum near 40000 (250 nm).⁵ \tilde{X} C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	NH stretch	3588.44	gas	EM	2,4
			3583	Ar	IR	1,5
Π	2	Bend	523	Ar	IR	1,5
Σ^+	3	NSi stretch	1198	Ar	IR	1,5

$B_0 = 0.634 \text{ gas } \text{EM}^{2,4} \text{MW}^3$

DNSi

 \tilde{X} C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	ND stretch	2669	Ar	IR	1
Π	2	Bend	395	Ar	IR	1
Σ^+	3	NSi stretch	1166	Ar	IR	1

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HCO

 $4p \text{ } ^2\Pi(A'')$ C_s
 $T_0 = 64073.5 \text{ gas DR}^{40,42}$

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	863.8(5)	gas	DR	40,42

 $3p \text{ } ^2\Pi$ C_{∞v}
 $T_0 = 45540.1(3.3) \text{ gas MPI}^{19,20,28}$

$3p^2\Pi - \tilde{X} \text{ 187--222 nm}$

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	799.9(1.4) ^a	gas	MPI	28
Σ^+	3	CO stretch	2177(3)	gas	MPI	20

$A = 0.9(3); \epsilon = 0.071 \text{ gas MPI}^{28}$

$B = 1.492(12) \text{ MPI}^{28}$

\bar{B}^2A'	C_s	Structure: MPI,LF ⁴⁰
$T_0=38695.48$ gas	EM ⁶ LF ^{25,30,33,35,36,39,43} MPI ²⁹ SEP ^{30,35}	$\bar{B}-\bar{X}$ 235–475 nm
38595(35) Ar	AB ^{5,11}	$\bar{B}-\bar{X}$ 210–260 nm
38567(35) CO	AB ⁵	$\bar{B}-\bar{X}$ 210–260 nm

Lifetime measurements^{32,35,37,38} give evidence for predissociation. Large decrease in fluorescence quantum yield above 41465.³³

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CH stretch	2596.4(2)	gas	EM,LF	6,25,30,33
			2570(29)	Ar	MPI AB	29,5,11
			2570(29)	CO	AB	5
	2	Mixed	1380.9(2)	gas	LF,MPI	25,29,30
						33
			1375(35)	Ar	AB	5,11
			1375(35)	CO	AB	5
	3	Mixed	1065.9(2)	gas	LF,MPI	25,29,30
						33
			1035(35)	Ar	AB	5,11
			1035(35)	CO	AB	5

$$A_0=15.976; \frac{1}{2}(B_0+C_0)=1.151; \frac{1}{4}(B_0-C_0)=0.021 \quad LF^{35,43}$$

$$\tau_0=89(2) \text{ ns} \quad LF^{27,32,35,37,38}$$

$\tilde{A}^2A''(\Pi)$ $C_{\infty v}$

$T_0=9297(3)$ gas	AB ^{1,3,8} LF ^{24,26} DM ³¹ CR ⁴¹	$\tilde{A}-\tilde{X}$ 460–860 nm
Bands with K'>0 are diffuse.		

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CH stretch	3319(3)	gas	AB	1,3,8
	2	Bend	805	gas	AB	1,3,8
	3	CO stretch	1812.2	gas	AB	1,3,8

$$\tau_{090}=46(4) \text{ ns} \quad LF^{15}$$

$$B_0=1.34 \quad UV^{1,3,8}$$

\tilde{X}^2A' C_s

Structure: MW⁷ UV⁸
Vibrational term energies up to 20777 have been reported by Ref. 36.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CH stretch	2434.48	gas	LF,PE	17,18,25
				DL,LD	22,26	
				IR	23	
			2483m	Ar	IR	5
			2488m	CO	IR	4
	2	Bend	1080.76	gas	UV,LS	1,3,8
				LMR	9,10	
				LF	25,26	
			1087s	Ar	IR	5
3			1090s	CO	IR	2,4
			1868.17	gas	IR	12,23
				LMR	13,25	
			1863vs	Ar	IR	5
			1861vs	CO	IR	2,4

$$A_0=24.329; B_0=1.494; C_0=1.399 \quad UV^{1,3,8}MW^{16}$$

DCO

$4p^2\Pi(A'')$	C_s
$T_0=64033.3$ gas	DR ^{40,42}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	677.3(5)	gas	DR	40,42

$3p^2\Pi$	$C_{\infty v}$
$T_0=45444.0(3.6)$ gas	MPI ^{20,28}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	645.5(1.6) ^a	gas	MPI	28
Σ^+	3	CO stretch	1900(5)	gas	MPI	20

$$A=0.9(3); \epsilon=0.069 \quad MPI^{28}$$

$$B=1.221(12) \quad MPI^{28}$$

\bar{B}^2A' C_s

$T_0=38628.4(2.0)$ gas	MPI ²⁹ LF ^{34,40}	$\bar{B}-\bar{X}$ 250–450 nm
38568(70) Ar	AB ⁵	$\bar{B}-\bar{X}$ 200–260 nm
38569(35) CO	AB ⁵	$\bar{B}-\bar{X}$ 204–260 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CD stretch	1944.2(2.0)	gas	MPI,LF	29,40,42
	2	Mixed	1212.9(2.5)	gas	MPI,LF	29,40,42
			1150(35)	Ar	AB	5
			1150(35)	CO	AB	5
	3	Mixed	922.1(1.6)	gas	MPI,LF	29,40,42
			925(35)	Ar	AB	5
			925(35)	CO	AB	5

$$A_0=9.172; B_0=1.093; C_0=0.974 \quad LF^{40}$$

$\tilde{A}^2A''(\Pi)$ $C_{\infty v}$

$T_0=9162(3)$ gas	UV ^{1,3,8} LF ^{24,26}	$\tilde{A}-\tilde{X}$ 460–860 nm
Bands with K'>0 are diffuse.		

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CD stretch	2547(2)	gas	UV	1,3,8
Π	2	Bend	641.7(7)	gas	UV	1,3,8

$$B_0=1.10 \quad UV^{1,3,8}$$

\tilde{X}^2A' C_s

Vibrational term energies up to 18186 have been reported by Ref. 42.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CD stretch	1909.77	gas	LMR,LF,SEP	14,26,40,42
			1926s	Ar	IR	5
			1937s	CO	IR	4
	2	Bend	846.5	gas	UV,LF,SEP	1,3,8,26,40,42
			850s	Ar	IR	5
			852s	CO	IR	2,4
	3	CO stretch	1794.59	gas	LMR,LF,SEP	14,26,40,42
			1803m	Ar	IR	5
			1800m	CO	IR	2,4

$$A_0=14.734; B_0=1.281; C_0=1.171 \quad UV^{1,3,8}MW^{21}$$

$$^a\omega_2$$

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HCF

$\tilde{E}^1A'(3p)$		C_s						
$T_0=62154(2)$		gas	MPI ¹⁵					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.		
a'	2	Bend	1128(4)	gas	MPI	15		
	3	CF stretch	1614(4)	gas	MPI	15		
\tilde{A}^1A''		C_s	Structure: $AB^1LF^{5,6}$					
$T_0=17277.47$		gas	$AB^1CL^3LF^{5,10}$	$\tilde{A}-\tilde{X}$ 429–635 nm				
17320(15)		Ar	AB^2	$\tilde{A}-\tilde{X}$ 469–546 nm				
Evidence has been obtained ^{8,9} for perturbation of the \tilde{A} state by high vibrational levels of the ground state and by the low-lying triplet state.								
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.		
a'	2	Bend	1021.26	gas	AB,LF	1,7		
			1000(20)	Ar	AB	2		
$A_0=25.69; B_0=1.162; C_0=1.107$ AB^1LF^5								
$\tau_0=2.45(10) \mu s$ gas LF ⁴								
$\tau_1=2.57(16) \mu s; \tau_2=12.5(8) \mu s$ gas EM ¹³								
\tilde{a}^3A''		C_s						
$T_0=5210(140)$		gas	PE ^{12,14}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.		
a'	2	Bend	1047(25)	gas	PE	14		
	3	CF stretch	1232(25)	gas	PE	14		
\tilde{X}^1A'		C_s	Structure: $AB^1LF^{5,6}SEP^{11}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.		
a'	1	CH stretch	2643.04	gas	SEP	11		
	2	Bend	1403.20	gas	AB,LF	1,7		
					SEP	11		
	3	CF stretch	1406vw	Ar	IR	2		
			1189(25)	gas	PE	12,14		
			1181.5m	Ar	IR	2		
$A_0=15.563; B_0=1.223; C_0=1.129$ $AB^1LF^5SEP^{11}$								
DCF								
$\tilde{E}^1A'(3p)$		C_s						
$T_0=62175(2)$		gas	MPI ¹⁵					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.		
a'	1	CD stretch	2095(5)	gas	MPI	15		
	2	Bend	872(3)	gas	MPI	15		
	3	CF stretch	1582(4)	gas	MPI	15		

\tilde{A}^1A''	C _s	T ₀ =17293.426(3)	gas	CL ³ LF ⁶	$\tilde{A}-\tilde{X}$ 460–585 nm
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	780(5)	gas	CL	3

$$A_0=15.10; B_0=1.014; C_0=0.945 \text{ LF}^6$$

\tilde{a}^3A'	C _s	T ₀ <5140(700)	gas	PE ¹²
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	834(25)	gas	PE	12,14
	3	CF stretch	1216(25)	gas	PE	12,14

\tilde{X}^1A'	C _s	T ₀	gas	AB ¹	$\tilde{A}-\tilde{X}$ 550–820 nm
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	2	Bend	1046m	Ar	IR
	3	CF stretch	1193(25)	gas	PE
			1183m	Ar	IR

$$A_0=8.828; B_0=1.120; C_0=0.990 \text{ LF}^6$$

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HCCI

\tilde{A}^1A''	C _s	Structure: AB ⁸	T ₀ =12280.411(2)	gas	AB ^{1,8,9} LF ^{4,7}	$\tilde{A}-\tilde{X}$ 550–820 nm
				Ar	AB ²	$\tilde{A}-\tilde{X}$ 570–750 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	873.0	gas	AB	1,9
			855(50)	Ar	AB	2
	3	CCl stretch	987.4	gas	AB	9

Barrier to linearity=2250¹
B₀=0.609 AB⁸

\tilde{a}^3A''	C _s	T ₀ =1470(880)	gas	PE ^{5,6}
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	3	CCl stretch	850(60)	gas	PE	5,6

\tilde{X}^1A' C_s Structure: AB¹LF³

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	1201wm	Ar	IR	2
	3	CCl stretch	811.60	gas	DL	10
			815s	Ar	IR	2

$$A_0=15.759; B_0=0.605; C_0=0.581 \text{ AB}^1\text{LF}^3$$

DCCI

\tilde{A}^1A''	C _s	T ₀ =12274	gas	AB ¹	$\tilde{A}-\tilde{X}$ 550–820 nm
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	657.2	gas	AB	1

\tilde{X}^1A' C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	3	CCl stretch	805s	Ar	IR	2

$$A_0=8.75; B_0=0.557; C_0=0.525 \text{ AB}^1$$

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HCBr

\tilde{A}^1A''	C _s	Structure: AB ⁴	T ₀₂₀ =12786.06	gas	LF ³ AB ⁴	$\tilde{A}-\tilde{X}$ 550–785 nm
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	840T	gas	LF	3
	3	CBr stretch	783T	gas	LF	3

Barrier to linearity≤1600.

$$B_0=0.437 \text{ AB}^4$$

\tilde{A}^1A'' C_s
 $T_0=910(770)$ gas PE^{1,2}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	3	CBr stretch	725(70)	gas	PE	2
\tilde{X}^1A'	C _s	Structure: AB ⁴				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	3	CBr stretch	683(25)	gas	PE	1,2
$B_0=0.422$		AB ⁴				

DCBr

\tilde{A}^1A''	C _s	gas LF ³	$\tilde{A}-\tilde{X}$ 550–600 nm			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	620T	gas	LF	3
	3	CBr stretch	769T	gas	LF	3

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HSiF

\tilde{A}^1A''	C _s	Structure: LF ^{3,6}	$T_0=23260.02$	gas LF ^{2,6}	$\tilde{A}-\tilde{X}$ 390–470 nm	
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	SiH stretch	1547	gas	LF	5,6
	2	Bend	558	gas	LF	2,5,6
	3	SiF stretch	857	gas	LF	5,6

$\tau_0=175(6)$ ns gas LF^{2,5}
 $A_0=9.319$; $B_0=0.549$; $C_0=0.516$ LF^{3,5}
 Barrier to linearity = 9130(20) LF⁵

\tilde{X}^1A'	C _s	Structure: LF ^{3,4,6}				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	SiH stretch	1913s	Ar	IR	1
	2	Bend	860T	gas	LF	2
			859m	Ar	IR	1
	3	SiF stretch	834s	Ar	IR	1

$A_0=7.58$; $B_0=0.564$; $C_0=0.524$ LF^{3,4}

DSiF

\tilde{A}^1A''	C _s	$T_0=23338.723(6)$	gas LF ⁶	$\tilde{A}-\tilde{X}$ 383–453 nm
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Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Ref:
a'	1	SiD stretch	1174.3	gas	LF	6
	2	Bend	424.8	gas	LF	6
	3	SiF stretch	854.4	gas	LF	6

$A_0=5.086$; $B_0=0.518$; $C_0=0.467$ LF⁶

 \tilde{X}^1A' C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Ref:
a'	1	SiD stretch	1387m	Ar	IR	1
	2	Bend	641.5	gas	LF	6
			638w	Ar	IR	1
	3	SiF stretch	833m	Ar	IR	1

$A_0=3.997$; $B_0=0.549$; $C_0=0.481$ LF⁶

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HSiCl

\tilde{A}^1A''	C _s	Structure: UV ¹ LF ²	$T_0=20717.77$	gas UV ¹ LF ²	$\tilde{A}-\tilde{X}$ 410–600 nm
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Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Ref
a'	1	SiH stretch	1747.1	gas	LF	2
	2	Bend	563.9	gas	UV,LF	1,2
	3	SiCl stretch	532.3	gas	UV,LF	1,2

$\tau_0=432(20)$ ns gas LF²
 $A_0=9.840(2)$; $B_0=0.247$; $C_0=0.240$ UV¹LF²

 \tilde{X}^1A' C_s Structure: UV¹LF²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Ref
a'	1	SiH stretch	1968.8(2)	gas	LF	2
	2	Bend	805.9(2)	gas	UV,LF	1,2
	3	SiCl stretch	522.8	gas	UV,LF	1,2

$A_0=7.587$; $B_0=0.246$; $C_0=0.238$ UV¹LF²

DSiCl

\tilde{A}^1A''		C_s		$\tilde{A}-\tilde{X}$ 410–600 nm		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SiD stretch	1300.8	gas	LF	2
	2	Bend	408.6	gas	UV,LF	1,2
	3	SiCl stretch	543.2	gas	LF	2

$\tau_0=437(20)$ ns gas LF²
 $A_0=5.269$; $B_0=0.235$; $C_0=0.224$ UV¹LF²

\tilde{X}^1A'		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SiD stretch	1434.4	gas	LF	2
	2	Bend	592.3(3)	gas	LF	2
	3	SiCl stretch	518.1(2)	gas	LF	2

$A_0=3.970$; $B_0=0.240$; $C_0=0.226$ UV¹LF²

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HSiBr

\tilde{A}^1A''		C_s		Structure: UV ¹ LF ²		
T_0	gas	UV ¹ LF ²	$\tilde{A}-\tilde{X}$ 429–620 nm			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SiH stretch	1787.0	gas	LF	2
	2	Bend	535.3	gas	UV,LF	1,2
	3	SiBr stretch	416.5	gas	UV,LF	1,2

$A_0=9.897(2)$; $B_0=0.159$; $C_0=0.155$ UV¹LF²
 $\tau_0=598(18)$ ns gas LF²

\tilde{X}^1A'		C_s		Structure: UV ¹ LF ²		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SiH stretch	1976.2(3)	gas	LF	2
	2	Bend	771.9(2)	gas	UV,LF	1,2
	3	SiBr stretch	412.4(2)	gas	UV,LF	1,2

$A_0=7.576(8)$; $B_0=0.158$; $C_0=0.155$ UV¹LF²

DSiBr

\tilde{A}^1A''		C_s		$\tilde{A}-\tilde{X}$ 464–502 nm		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SiD stretch	1325.6	gas	LF	2
	2	Bend	375.9	gas	LF	2
	3	SiBr stretch	434.3	gas	LF	2

$\tau_0=586(21)$ ns gas LF²
 $A_0=5.274(3)$; $B_0=0.151$; $C_0=0.146$ LF²

\tilde{X}^1A'		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SiD stretch	1439.5	gas	LF	2
	3	SiBr stretch	408.0	gas	LF	2

$A_0=3.956(3)$; $B_0=0.153$; $C_0=0.148$ LF²

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HNO

$T_0=48240$ gas AB⁵ 198–208 nm
Diffuse bands.

\tilde{A}^1A'' C_s Structure: AB^{1,4}
 $T_0=13154.37$ gas AB^{1,4,25}LF^{13,21}CL²⁴CR²⁶ $\tilde{A}-\tilde{X}$ 544–770 nm
13118(2) Ar AB^{2,3} $\tilde{A}-\tilde{X}$ 590–762 nm
Onset of predissociation at 16450(10). LF¹³

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	NH stretch	2854.17	gas	AB	4
	2	Bend	981.18	gas	AB	1
			982	Ar	AB	2,3
	3	NO stretch	1420.77	gas	AB	1
			1422	Ar	AB	2,3

$\tau=25(4)$ μs LF^{12,14,23}
 $A_0=22.161$; $B_0=1.327$; $C_0=1.241$ AB^{1,4,25}MODR^{11,17}

\tilde{a}^3A'' C_s
 $T_0=6280(160)$ gas PE¹⁵

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	Bend	992(150)	gas	PE	15
	3	NO stretch	1468(140)	gas	PE	15

\tilde{X}^1A' C_s Structure: AB¹

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	NH stretch	2683.95	gas	IR,EM	9,16,22
			2716.3w ^a	Ar	IR	8
			2756m	N ₂	IR	8
	2	Bend	1500.82	gas	LS	10
			1505w	Ar	IR	8
			1511w	N ₂	IR	8
	3	NO stretch	1565.34	gas	LS	10
			1563.2vs ^a	Ar	IR	8
			1568.5s	N ₂	IR	8

$A_0=18.476$; $B_0=1.411$; $C_0=1.306$ AB^{1,25}MW^{7,19}IR^{10,16,22}

DNO

$T_0=48400$ gas AB⁵
Diffuse bands.

\tilde{A}^1A'' C_s
 $T_0=13180.3$ gas AB¹
Onset of predissociation at 17010(10) LF^{18,20}

196–206 nm
 $\tilde{A}-\tilde{X}$ 550–770 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	ND stretch	2176.49	gas	AB	4
	2	Bend	755.31	gas	AB	1
	3	NO stretch	1401.28	gas	AB	1

$\tau_{011}=32.4(1.6)$ μ s gas LF²³
 $A_0=12.629$; $B_0=1.199$; $C_0=1.088$ AB^{1,4}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	750(140)	gas	PE	15
	3	NO stretch	1452(140)	gas	PE	15

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	ND stretch	2025.14	gas	LS,IR	10,16
		2043wm	Ar	IR		8
		2074m	N ₂	IR		8
2	Bend	1153s	Ar	IR		8
		1158.5m	N ₂	IR		8
3	NO stretch	1546.88	gas	LS		10
		1547vs	Ar	IR		8
		1548vs	N ₂	IR		8

$A_0=10.524$; $B_0=1.292$; $C_0=1.146$ AB¹MW^{6,19}IR^{10,16}

^aRefined value from unpublished Fourier transform spectra.

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HSO⁺ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Ref.
			1150(160)	gas	PI	1

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HNF

\tilde{A}^2A' C_s Structure: AB³LF^{5,8}
 $T_0=20141.26(1)$ gas AB^{1,3}CL⁴LF^{5,7} $\tilde{A}-\tilde{X}$ 380–650 n
20140(20) Ar AB² $\tilde{A}-\tilde{X}$ 395–497 n

Onset of predissociation into NH+F identified at 23800(500) gas LF⁵PF⁶

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	Bend	1074	gas	AB	3
			1033 ^a	Ar	AB	2
	3	NF stretch	1121(5)	gas	AB	3

$\tau_0=3.6$ μ s gas LF⁵
 $A_0=27.570(5)$; $B_0=1.033$; $C_0=0.992$ AB³

 \tilde{X}^2A'' C_s Structure: AB³LF^{5,8}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Ref.
a'	1	NH stretch	3167(6)	gas	LF	7
	2	Bend	1439(3)	gas	CL,LF	4,7
			1432m	Ar	IR	2
	3	NF stretch	1020(8)	gas	LF	7
			1000s	Ar	IR	2

$A_0=17.688(8)$; $B_0=1.039$; $C_0=0.978$ AB³

DNF

\tilde{A}^2A'		C_s
$T_0=20169.10$	gas	LF ^{5,7,8}
20220	Ar	AB ²

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	Bend	837.4(8) ^a	gas	LF	5
			798 ^b	Ar	AB	2
	3	NF stretch	1124 ^c	gas	LF	5

$\tau_0=3.5(2) \mu\text{s}$ gas LF⁵
 $A_0=15.665(6); \frac{1}{2}(B_0+C_0)=0.900; \frac{1}{4}(B_0-C_0)=0.015$ LF⁸

\tilde{X}^2A''		C_s
Vib.	No.	Approximate type of mode
a'	1	ND stretch
	2	Bend
	3	NF stretch

$A_0=9.49; \frac{1}{2}(B_0+C_0)=0.929; \frac{1}{4}(B_0-C_0)=0.023$ LF⁸

^a $\omega_2+0.5 x_{12}$.

^bAverage value.

^c $\omega_3+2x_{33}+0.5x_{13}$.

^dOverlapped by NF₂ absorption.

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HO₂

Broad, unstructured gas-phase absorption between 200 and 280 nm, with maximum near 205 nm.^{2,3,5,6,37}

\tilde{A}^2A'		C_s
$T_0=7029.684(2)$	gas	AB ^{9,15} EM ^{10,16,19,20,35}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OH stretch	3268.5	gas	EM	35
	2	Bend	1285	gas	EM	35
	3	OO stretch	929.068	gas	AB,EM	15,19,28 35

$A_0=20.486; B_0=1.021; C_0=0.968$ EM^{16,20,35}

 \tilde{X}^2A'' C_s Structure:MW¹⁴UV²¹LMR²³ESR²³IR^{31,33}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OH stretch	3436.20	gas	LD	26
			3415.1	Ne	IR	36
			3412.5s ^a	Ar	IR	1,4,7
			3400	O ₂	IR	32
	2	Bend	1391.75	gas	DL,IR	24,39
			1397.8	Ne	IR	36
			1388.5vs ^a	Ar	IR	1,4,7
	3	OO stretch	1392	O ₂	IR	32
			1097.63	gas	LMR	18,29
			1100.3	Ne	DL,IR	29,38,39
			1101.1s ^a	Ar	IR	1,4,7
			1109	O ₂	IR	32

$A_0=20.356; B_0=1.118; C_0=1.056$ LMR^{8,11,12,18} MW^{13,17,25} EM¹⁶ IR⁴⁰

DO₂

\tilde{A}^2A'		C_s
$T_0=7041.1(1)$	gas	AB ⁹ EM ^{10,19,21}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	3	OO stretch	940(28)	gas	AB,EM	8,19

$A_0=11.147(7); B_0=0.970; C_0=0.887$ EM²¹

 \tilde{X}^2A'' C_s

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OD stretch	2549.22	gas	LD,DL	31
			2529.2	Ne	IR	36
			2529.5m ^a	Ar	IR	1,4,7
			2521	O ₂	IR	32
			1020.16	gas	LMR,DL	22,33
	2	Bend	1027.3	Ne	IR	36
			1019.9s ^a	Ar	IR	1,4,7
			1024	O ₂	IR	32
			1121.47	gas	LMR, DL	22,33
			1124.7	Ne	IR	36
	3	OO stretch	1122.9vw ^a	Ar	IR	7

$A_0=11.194; B_0=1.056; C_0=0.961$ MW^{14,27,34} EM²¹ LMR^{22,23,34} ESR²³

^aRefined value from unpublished Fourier transform spectra.

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HSO

\tilde{A}^2A'		C_s	Structure: LF ^{2,3}		$\tilde{A}-\tilde{X}$ 520–960 nm		
$T_0=14367$	gas	CL ¹					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a'	3	SO stretch	702(5)	gas	CL	1	

 $\tau_0=81(10) \mu s$ gas LF^{6,9}Values decrease as v'_3 increases. $A_{003}=9.735; B_{003}=0.565; C_{003}=0.527$ LF²

\tilde{X}^2A''		C_s	Structure: LF ^{2,3} MW ⁴				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a'	2	Bend	1063(5)	gas	CL	1	
	3	SO stretch	1009.36	gas	LMR	5	

 $A_0=9.990; B_0=0.684; C_0=0.638$ LF²MW⁴**DSO**

\tilde{A}^2A'		C_s	$\tilde{A}-\tilde{X}$ 520–960 nm		
$T_0=14371$	gas	CL ¹ LF ⁸			

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	Bend	575(10)	gas	LF	7
	3	SO stretch	702(10)	gas	CL	1

 $\tau^a=76 \mu s$ gas LF⁶
 $A_0^a=4.969(7); B_0^a=0.566; C_0^a=0.507$ LF^{7,8}

\tilde{X}^2A''		C_s	$\tilde{A}-\tilde{X}$ 520–960 nm		
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
a'	2	Bend	770(10)	gas	CL
	3	SO stretch	1029(15)	gas	CL

 $A_0=5.295; B_0=0.662; C_0=0.586$ LF³MW⁴

aMeasured at 606.0 nm.

bExtrapolated from values for 021 and 022.

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HS₂

\tilde{A}^2A'		C_s	$\tilde{A}-\tilde{X}$ 297–380 nm		
$T_0=14367$	gas	CL ¹			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
a'	1	SH stretch	2500T	gas	AB
	2	Bend	900T	gas	AB
	3	SS stretch	600T	gas	AB

\tilde{A}^2A'		C_s	$\tilde{A}-\tilde{X}$ 950–2100 nm		
$T_0=7255(7)$	gas	CL ⁴			

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
a'	3	SS stretch	504(4)	gas	CL

 $A_0=9.7(5)$ CL⁴

\tilde{X}^2A''		C_s	Structure: MW ⁵			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	Bend	904(8)	gas	CL	4
	3	SS stretch	595(4)	gas	CL	4
$A_0=9.906; B_0=0.267; C_0=0.259$		MW ⁵ LMR ⁶				

DS₂

\tilde{A}^2A'		C_s	Structure: $\tilde{A}-\tilde{X}$ 950–2100 nm			
T ₀		gas	CL ⁴			

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	3	SS stretch	502(15)	gas	CL	4

\tilde{X}^2A''		C_s	Structure: MW ⁵			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	Bend	696(20)	gas	CL	4
	3	SS stretch	591(10)	gas	CL	4
$A_0=5.184; B_0=0.260; C_0=0.247$		MW ⁵				

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HOBr⁺

\tilde{A}^2A'		C_s	Structure: PI ¹ TPE ²			
T ₀		gas	PI ¹ TPE ²			

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	3	OBr stretch	670T	gas	PI,TPE	1,2

\tilde{X}^2A''		C_s	Structure: MW ^{4,12}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	3	OBr stretch	730(20)	gas	PI,TPE	1,2

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HOI⁺

\tilde{X}^2A''		C_s	Structure: MW ^{4,12}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	3	OI stretch	702(60)	gas	PI	1

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HOCl

\tilde{X}		C_s	Structure: IR ^{2,14} MW ^{4,12}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OH stretch	3609.48	gas	IR	2,6,7,11,15
			3581	Ar	IR	3
	2	Bend	1238.62	gas	IR	1,5,6,8,9 11,13
a'	3	OCl stretch	1239	Ar	IR	3
			724.36	gas	IR	5,6,8,11,13
			728	Ar	IR	3

$A_0=20.464; B_0=0.504; C_0=0.491$ IR^{2,11,15,16}MW^{4,10}

DOCl

\tilde{X}		C_s	Structure: MW ^{4,12}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OD stretch	2665.58	gas	IR	1,2,11,14
			2647	Ar	IR	3
	2	Bend	909.63	gas	IR	1,11,14
a'	3	OCl stretch	911	Ar	IR	3
			723.25	gas	IR	11,14
			728	Ar	IR	3

$A_0=11.052; B_0=0.477; C_0=0.456$ IR^{2,14}MW^{4,12}

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HOBr

In the gas phase, a prominent absorption maximum near 280 nm (35700) has been assigned^{6,8} to HOBr.

In the gas phase, an absorption maximum near 350 nm (28600), with onset near 400 nm, has been assigned^{6,8} to HOBr.

In the gas phase, a maximum in the production of OH near 440 nm (22700) is believed to arise from excitation of HOBr to a dissociative triplet state.⁹

\tilde{X}		C_s	Structure: MW ³ IR ⁷			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OH stretch	3614.90	gas	IR	2,7
			3590	Ar	IR	1
	2	Bend	1162.57	gas	IR	4
			1164	Ar	IR	1
	3	OBr stretch	620.2	gas	IR	2,4,5
			626.0	Ar	IR	1

$$A_0=20.470; B_0=0.353; C_0=0.346 \quad \text{MW}^3\text{IR}^7$$

DOBr

\tilde{X}		C_s	Structure: MW ³			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OD stretch	2652	Ar	IR	1
	2	Bend	854	Ar	IR	1
	3	OBr stretch	621.8	Ar	IR	1

$$A_0=11.027; B_0=0.331; C_0=0.321 \quad \text{MW}^3$$

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HOI

\tilde{X}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OH stretch	3625.84	gas	IR	2,3
			3597m	N ₂	IR	1
	2	Bend	1069.8	gas	IR	2
			1075	Ar	IR	1
			1103m	N ₂	IR	1
	3	OI stretch	577	Ar	IR	1
			575m	N ₂	IR	1

$$A_0=20.935; B_0=0.279; C_0=0.275 \quad \text{EM}^3$$

DOI

\tilde{X}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OD stretch	2653	N ₂	IR	1
	2	Bend	808	N ₂	IR	1
	3	OI stretch	571	N ₂	IR	1

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FHF⁻

\tilde{X} D_∞h Structure: DL⁵

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	583.05	gas	DL	5
Π_u	2	Bend	1286.03	gas	DL	5
			1217m Cs	Ar	IR	1,2
Σ_u^+	3	Asym. stretch	1331.15	gas	DL	5
			1379	Ne	IR	4
			1377.0s	Ar	IR	3,4
			1364vs Cs	Ar	IR	1,2

$$B_0=0.334 \quad \text{DL}^5$$

FDF⁻

\tilde{X} D_∞h

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	588T	gas	DL	6
Π_u	2	Bend	928.73	gas	DL	6
			880m Cs	Ar	IR	1,2
Σ_u^+	3	Asym. stretch	934.19	gas	DL	6
			965s	Ar	IR	3,4
			969vs Cs	Ar	IR	1,2

$$B_e=0.342 \quad \text{DL}^6$$

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ClHCl⁻

An absorption maximum which appeared at 287 nm in argon-matrix studies² of the 122-nm photolysis of Ar:HCl or Ar:H₂O:Cl₂ samples and in argon-matrix studies⁶ of the electron bombardment of Ar:HCl samples has been assigned to an electronic transition of ClHCl⁻.

\tilde{X}		D _{∞h}	Structure: DL ⁵			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	263.4 ^b	Ne	IR	8
			259.3 ^{a,b}	Ar	IR	1-3,7
			252.8 ^b	Kr	IR	7
			248.8 ^b	Xe	IR	7
			722.90	gas	DL	5
			737.9	Ne	IR	8
Σ_u^+	3	Asym. stretch	728.9			
			695.6 ^a	Ar	IR	1-3,7
			662.8	Kr	IR	7
			644.1	Xe	IR	7
$B_0 = 0.0974 \text{ DL}^5$						

CIDCI⁻

\tilde{X}		D _{∞h}	Structure: DL ⁵			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	267 ^{a,b}	Ar	IR	1-3
			255.3 ^b	Kr	IR	7
			496.2	Ne	IR	8
			489.3			
			463 ^a	Ar	IR	1-3
			437.7	Kr	IR	7

^aAttributed by Ref. 1 to the uncharged species. Reassigned to the anion by Ref. 2, and Ref. 3 demonstrated that the absorptions did not appear when the atoms were present but a supplementary high energy source suitable for inducing photoionization or electron transfer was not. The increase in the rate of isotopic exchange in the ³⁷Cl+H³⁵Cl reaction for vibrationally excited HCl^a indicates that there is a potential barrier, rather than a minimum, for the ClHCl neutral species.

^b($\nu_1 + \nu_3$) - ν_3 .

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BrHBr⁻

\tilde{X}		D _{∞h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	165.1 ^b	Ne	IR	4
			163.1 ^b			
			162.6 ^b			
			164 ^{a,b}	Ar	IR	1,2
			158.5 ^b	Kr	IR	3
			151.9 ^b	Xe	IR	3
Σ_u^+	3	Asym. stretch	752.9	Ne	IR	4
			745.4			
			741.2			
			728 ^a	Ar	IR	1,2
			686.6	Kr	IR	3
			645.6	Xe	IR	3

BrDBr⁻

\tilde{X}		D _{∞h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	163.9 ^b	Ne	IR	4
			160.8 ^b			
			170 ^{a,b}	Ar	IR	1,2
			163.8 ^b	Kr	IR	3
			157.9 ^b	Xe	IR	3
			516.5	Ne	IR	4
Σ_u^+	3	Asym. stretch	510.7			
			507.6			
			498 ^a	Ar	IR	1,2
			465.5	Kr	IR	3
			434.6	Xe	IR	3

^aAttributed in Ref. 1 to the uncharged species. Reassigned to the anion by Ref. 2. See ClHCl⁻.

^b($\nu_1 + \nu_3$) - ν_3 .

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IHI⁻

\tilde{X}		$D_{\infty h}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	115.2 ^b	Ne	IR	4
			114.8 ^b			
			121 ^{a,b}	Ar	IR	1,2
			116.7 ^b	Kr	IR	3
			109.0 ^b	Xe	IR	3
Σ_u^+	3	Asym. stretch	671.8	Ne	IR	4
			653.3			
			644.7			
			682m ^a	Ar	IR	1,2
			647.4	Kr	IR	3
			580.8	Xe	IR	3

IDI⁻

\tilde{X}		$D_{\infty h}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	124 ^{a,b}	Ar	IR	1,2
Σ_u^+	3	Asym. stretch	470m ^a	Ar	IR	1,2

^aAttributed in Ref. 1 to the uncharged species. Reassigned by Ref. 2 to the anion. See ClHCl⁻.

^b($\nu_1 + \nu_3$) - ν_3 .

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HKrCl

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	HKr stretch	1476vs	Kr	IR	1
II	2	Bend	544w	Kr	IR	1

DKrCl

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	DKr stretch	1106s	Kr	IR	1

Reference

- M. Pettersson, J. Lundell, and M. Räsänen, J. Chem. Phys. **102**, 6423 (1995).

MARYL E. JACOX**HXeCl**

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	HXe stretch	1664	Kr	IR	1
			1649	Xe	IR	1

DXeCl

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	DXe stretch	1198	Xe	IR	1

Reference

- M. Pettersson, J. Lundell, and M. Räsänen, J. Chem. Phys. **102**, 6423 (1995).

HXeBr

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	HXe stretch	1524	Kr	IR	1
			1504vs	Xe	IR	1
II	2	Bend	489w	Xe	IR	1

DXeBr

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	DXe stretch	1100s	Xe	IR	1

Reference

- M. Pettersson, J. Lundell, and M. Räsänen, J. Chem. Phys. **102**, 6423 (1995).

HxeI

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	HxeI stretch	1193vs	Xe	IR	1
II	2	Bend	450w	Xe	IR	1

D_{Xe}I

\tilde{X}		$C_{\infty v}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	DXe stretch	893vs	Xe	IR	1

Reference

¹M. Pettersson, J. Lundell, and M. Räsänen, *J. Chem. Phys.* **102**, 6423 (1995).

HAr₂⁺ a

\tilde{X}		$D_{\infty h}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	237 ^b	Ar	IR	5
Σ_u^+	3	Asym. stretch	905wm ^c	Ar	IR	1,2,4,5

DAr₂⁺ a

\tilde{X}		$D_{\infty h}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	644s ^c	Ar	IR	1-5

^aMay be complexed with two or four additional Ar atoms.²

^b($\nu_1 + \nu_3$) - ν_3 .

^cAssigned in Ref. 1 to vibration of uncharged H (or D) atoms trapped in the Ar lattice. Reassigned by Ref. 2 to the cation. Peak at 644 cm^{-1} was prominent in deuteron radiolysis experiments.³ Ref. 4 demonstrated that the absorptions did not appear when H or D atoms were present but a supplementary high energy source suitable for inducing photoionization or electron transfer was not.

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HKr₂⁺ a

\tilde{X}		$D_{\infty h}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	155 ^b	Kr	IR	4
Σ_u^+	3	Asym. stretch	853m ^c	Kr	IR	1-4

DKr₂⁺ a

\tilde{X}		$D_{\infty h}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	165 ^b	Kr	IR	4
Σ_u^+	3	Asym. stretch	606m ^c	Kr	IR	1,4

^aMay be complexed with two or four additional Kr atoms.²

^b($\nu_1 + \nu_3$) - ν_3 .

^cAssigned in Ref. 1 to vibration of uncharged H (or D) atoms trapped in the Kr lattice. Reassigned by Ref. 2 to the cation. Ref. 3 demonstrated that the 852 cm^{-1} absorption did not appear when H atoms were present but a supplementary high energy source suitable for inducing photoionization or electron transfer was not.

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8.3. Triatomic Nonhydrides**Li₃**

\tilde{C}^2E''	D_{3h}^a	$T_0=21541$	gas	MPI^3	$\tilde{C}-\tilde{X}$	450–472 nm
$B=0.57$	MPI^4					

\tilde{A}^2E'	D_{3h}^a	$T_0=14583$	gas	$\text{MPI}^{2,3,5}$	$\tilde{A}-\tilde{X}$	660–706 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	326	gas	MPI	2

Structure in the \tilde{A} state is characterized² by a radial frequency $\omega_0=191$ and linear and quadratic Jahn-Teller parameters $k=0.77$ and $g=0.15$.
 $B=0.57$ MPI^4

\tilde{X}^2E'	D_{3h}^a	Structure: MPI^4				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	302	gas	MPI	3
			303	Xe	Ra	1

Barrier to pseudorotation=26; pseudorotation frequency=34 MPI^3
 $B=0.584$ MPI^4

^aSubject to dynamic Jahn-Teller distortion.

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Na₃

\tilde{D} $^2E''(^2A_2)$ D_{3h}(C_{2v})^a
gas MPI^{4,5,14,19}DPI¹²TPE¹⁴
Fragments into Na₂ + Na in less than 1 ps.¹⁹

\tilde{C} $^2E''(^2A_2)$ D_{3h}(C_{2v})^a
 $T_0=20813$ gas MPI^{2,4,6}DPI^{8,12}
Higher vibrational bands are predissociated.⁷

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	135	gas	MPI,PF	8

Extensive vibronic structure has been tentatively assigned⁸ to energy levels derived from excitation of ν_2 (e'), perturbed by dynamic Jahn-Teller interaction.

$\tau_0=7(3)$ ns gas MPI⁷
 $\tau_{\text{frag}}=1.12$ ns for the lowest vibrational level. MPI²⁵

\tilde{B}'' 2B_2
 $T^b=19200$ gas DPI¹²

\tilde{B} $^2A'_1$ D_{3h}^a Structure: MPI²⁴
 $T_0=16124.63$ gas MPI^{1-6,17,20-24}DPI¹²TPE¹⁸ $\tilde{B}-\tilde{X}$ 550–625 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			128	gas	MPI,TPE	1,3,18
			105	gas	MPI,TPE	18
			74	gas	MPI,TPE	18

$\tau_{\tilde{B}}(16255)=14(5)$ ns gas MPI⁷

$\tau_{\tilde{B}'}(17418)=7(3)$ ns gas MPI⁷

$A_0=0.082(2)$; $B_0=0.138(4)$; $C_0=0.051$ MPI^{22,24}

Vibronic pseudorotation occurs,³ with a beat structure period of approximately 3 ps.^{17,20,21}

\tilde{A} $^2E''(^2A_2)$ D_{3h}(C_{2v})^a Structure: MPI^{15,16}
 $T_0=14894.769(4)$ gas MPI^{1,2,4-6,11,13,15,16,26}DPI¹² $\tilde{A}-\tilde{X}$ 658–675 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	Sym. stretch	150	gas	MPI	11
	2	Bend	127	gas	MPI	4,6,11,26
	3	Asym. stretch	47	gas	MPI	4,6,11

Vibronic structure assigned in Ref. 11. First three bands are partially localized; pseudorotation barrier=196.

$\tau_0=60(10)$ ns gas MPI⁷

$A_0=0.166$; $B_0=0.085$; $C_0=0.056$ MPI^{15,16}

A' 2A_1 D_{3h}
 $T^b=13500$ gas DPI¹²

\tilde{X} $^2E'(^2B_2)$ D_{3h}(C_{2v})^a Structure: MPI^{15,16}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	Sym. stretch	139	gas	MPI,SEP	4,6,9,10
	2	Bend	49.5	gas	MPI,SEP	4,6,9,10
	3	Asym. stretch	87	gas	MPI,SEP	6,9,10

$A_0=0.177$; $B_0=0.085$; $C_0=0.057$ MPI^{15,16,22}

^aDistorted by Jahn-Teller interaction.
^bBand maximum.

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Cu₃ \tilde{B}

In the ion depletion study of Ref. 10, a weak, broad band with maximum near 20600 (485 nm) has been attributed to the $\tilde{B}-\tilde{X}$ transition of Cu₃.

 \tilde{A}

$^2A'_1$ D_{3h}
 $T_0=18524$ gas DPI^{3,8,10}LF^{5,6}AB⁹ $\tilde{A}-\tilde{X}$ 522–567 nm

In an argon matrix, an absorption maximum at 18760 (533 nm) has been assigned¹ to Cu₃.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	146.4(5)	gas	DPI,LF	3,5,8
e'	2	Deformation	243	gas	DPI LF,AB	3,8 5,9

$\tau_0=35(5)$ ns gas MPI³LF⁵

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	269.5	gas	LF	5,7

^aDistorted by Jahn-Teller interaction.

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Ag₃

In an argon matrix study with mass selection,⁸ an absorption maximum at 31150 (321 nm) behaves appropriately for assignment to Ag₃, as does an emission maximum at 26700 (374 nm). In a similar krypton matrix study,⁶ the corresponding absorption maximum lies at 30200 (331 nm), and there is an emission maximum at 26200 (381 nm).

In the krypton-matrix mass selection study,⁶ there is also a less prominent absorption maximum near 27500 (364 nm).

$^2E''$ D_{3h}^a
 $T_0=26969.0$ gas MPI^{4,5,9}LF⁷ $^2E''-\tilde{X}$ 365–385 nm
In an argon matrix study with mass selection,⁸ an absorption maximum at 25900 (386 nm) behaves appropriately for assignment to Ag₃. In a similar krypton matrix study,⁶ the absorption maximum appears at 24900 (402 nm).

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	158.2	gas	MPI,LF	4,5,7,9
e'	2	Deformation	96	gas	MPI,LF	4,5,7,9

$\tau<80$ ns gas LF⁷

In argon, krypton, and xenon matrices, an absorption maximum near 23700 (422 nm) has been attributed^{2,3,6} to Ag₃.

$^2A'_1$? D_{3h}
 $T_0=19809(2)$ gas EM¹⁰ 495–505 nm
In an argon matrix study with mass selection,⁸ an absorption maximum at 20300 (492 nm) and an emission maximum at 16100 (622 nm) behave appropriately for assignment to Ag₃. In a krypton matrix study with mass selection,⁶ absorption maxima at 21830 (458 nm) and 19500 (514 nm) and emission maxima at 17900 (560 nm) and 16000 (626 nm) also behave appropriately for assignment to Ag₃.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	107 ^b	gas	EM	10
e'	2	Deformation	94 ^b	gas	EM	10

\tilde{X} $^2E'$ D_{3h}^a

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	121	gas	LF	7,9
e'	2	Deformation	99	gas	LF	7,9

^aDistorted by Jahn-Teller interaction.

^b107^aAg₂¹⁰⁹Ag.

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Au₃

In an argon matrix study with mass selection,⁴ absorption maxima at 43300 (231 nm), 38800 (258 nm), 35100 (285 nm), and 33100 (302 nm) behave appropriately for assignment to Au₃. An absorption maximum at 34200 (292 nm) reported in an earlier study¹ without mass selection may corre-

spond to the 285 nm maximum. In a krypton matrix study with mass selection,³ the four absorption maxima appear at 42900 (233 nm), 39200 (255 nm), 35500 (282 nm), and 32500 (308 nm).

In an argon matrix study with mass selection,⁴ an absorption maximum at 27000 (370 nm) behaves appropriately for assignment to Au_3 . This absorption maximum experiences a relatively large shift, to 28600 (349 nm), in the corresponding krypton matrix study.³

In an argon matrix study with mass selection,⁴ absorption maxima at 21900 (457 nm) and 19600 (511 nm) behave appropriately for assignment to Au_3 . The 21200 (471 nm) band reported in an earlier study¹ without mass selection may have been contributed in part by Au_3 . The counterparts of these two bands appear at 21900 (457 nm), and 19500 (513 nm) in a krypton matrix study with mass selection.³

In an argon matrix study with mass selection,⁴ emission maxima for Au_3 were observed at 18900 (529 nm), 17300 (579 nm), and 12400 (809 nm). In a similar krypton matrix study,³ the corresponding emission maxima were at 19600 (510 nm), 17300 (579 nm), and 12300 (816 nm).

$\tilde{\alpha}^4E'$		D _{3h} ^a	$T_0=13354.15$ gas MPI ²		$\tilde{\alpha}-\tilde{X}$ 710–750 nm	
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	181.7	gas	MPI	2
e'	2	Deformation	56T	gas	MPI	2
$\tau_0=28(2)$ μs		gas	MPI ²			

\tilde{X}^2E'	D _{3h}
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^aDistorted by Jahn-Teller interaction.

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Nb₃

In an argon matrix, absorption maxima have been observed¹ near 20300 (492 nm), 18800 (532 nm), and 17000 (587 nm).

\tilde{X}		D _{3h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	335(3)	Ar	Ra	1
e'	2	Deformation	227(3)	Ar	Ra	1

Reference

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cyc-Fe₂N

\tilde{X}	C _{2v}
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			811.2	Ar	IR	2
			779.0	N ₂	IR	1,2
			719.0	N ₂	IR	2

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FeFeN

\tilde{X}	C _s
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	FeN stretch	771.0T	Ar	IR	1

Reference

- G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. **100**, 14609 (1996).

B₃

$\tilde{X}^2A'_1$	D _{3h}	Structure: ESR ¹
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
e'	2		886.2	Ar	IR	2

References

- Y. M. Hamrick, R. J. Van Zee, and W. Weltner, Jr., J. Chem. Phys. **96**, 1767 (1992).
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Al₃

\tilde{X}	D _{3h}	516–602 nm
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Overlapping continuum with high-frequency edge at 19378(10). This continuum may be associated with unresolved high vibrational levels of another electronic state.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1		270.6	gas	MPI	1
	2		205	gas	MPI	1

Lifetimes (possibly radiative) vary¹ from 98 to 21 μs .

4A_2 **C_{2v}**

Ref. 3 has suggested that the gas-phase band spacing of 133, assigned by Ref. 1 to the ground state of Al_3 , may instead be contributed by a fundamental of the low-lying 4A_2 state.

 2B_1 **C_{2v}**
 $T_0=1770.4(3)$ Ne IR³
 $1756.9(3)$ Ar IR³

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	319(2) 317	Ne Ar	IR IR	3 3

 $\bar{X} \ ^2A'_1$ **D_{3h}** Structure: ESR²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
e'	2		245(2) 245	Ne Ar	IR IR	3 3

^a Possibly a 1_0^n band.

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BeOBe

\bar{X}	$D_{\infty h}$					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1412.4 1407.5 1403.0 1408.3	Ar Kr Xe N ₂	IR IR IR IR	1 1 1 2

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TiOTi

\bar{X}	C_{2v}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	736.4	Ar	IR	1

Reference

- G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).

FeOFe

\tilde{A}
 $T_0=3150(240)$ gas PE³

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Asym. stretch	868.6 878.1 848.6	Ar N ₂ O ₂	IR IR IR	1 2 1

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NiNiO **\tilde{X}**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	NiO stretch	844.6T 849.5T	Ar N ₂	IR IR	1 1

Reference

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CuOCu

In an argon matrix,¹ excitation at 20725 (482 nm) yields an emission appropriate to a resonance Raman or resonance fluorescence of CuOCu.

\tilde{D}
 $T_0=14930(400)$ gas PE²

\tilde{C}
 $T_0=14120(400)$ gas PE²

\tilde{B}
 $T_0=12590(400)$ gas PE²

\tilde{A}
 $T_0=11540(400)$ gas PE²

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	625T	Ar	LF	1
	2	Bend	200U	gas	PE	2

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²L.-S. Wang, H. Wu, S. R. Desai, and L. Lou, Phys. Rev. B **53**, 8028 (1996).

Al₂C

\tilde{X}	C_{2v}
Vib. sym.	No. type of mode
b_2	3 CAI a-stretch

cm^{-1} Med. Type Refs.
802.0 Ar IR 1

Reference

- ¹G. V. Chertihin, L. Andrews, and P. R. Taylor, J. Am. Chem. Soc. **116**, 3513 (1994).

FeCC

\tilde{A}
$T_0=8000(800)$ gas PE ¹

Reference

- ¹J. Fan and L.-S. Wang, J. Phys. Chem. **98**, 11814 (1994).

FeCC⁻

Threshold for electron detachment from ground-state FeCC⁻=15400(800)
gas PE¹

Reference

- ¹J. Fan and L.-S. Wang, J. Phys. Chem. **98**, 11814 (1994).

MgCN

$\tilde{X}^2\Sigma^+$	$C_{\infty v}$
$B_0=0.170$	MW ¹

Reference

- ¹M. A. Anderson, T. C. Steimle, and L. M. Ziurys, Astrophys. J. **429**, L41 (1994).

MgNC

$\tilde{X}^2\Sigma^+$	$C_{\infty v}$	Structure: MW ²				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Π	2	Bend	86T	gas	MW	3
$B_0=0.199$		MW ¹				

References

- ¹K. Kawaguchi, E. Kagi, T. Hirano, S. Takano, and S. Saito, Astrophys. J. **406**, L39 (1993).
²M. A. Anderson and L. M. Ziurys, Chem. Phys. Lett. **231**, 164 (1994).
³E. Kagi, K. Kawaguchi, S. Takano, and T. Hirano, J. Chem. Phys. **104**, 1263 (1996).

CaNC

$\tilde{C}^2\Pi$ $C_{\infty v}$
Unstructured absorption gas LF¹
 $\tau=165(38)$ ns gas LF¹ $\tilde{C}-\tilde{X}$ 385–418 nm

$\tilde{A}^2\Pi$ $C_{\infty v}$
 $T_0=16229.54$ gas LF^{1,3-6,9} CL² $\tilde{A}-\tilde{X}$ 572–670 nm
 $\tau(607 \text{ nm})=40.8(1.5)$ ns gas LF¹
 $A=77.64$ gas LF^{6,9}
 $B_0=0.150$ LF^{6,9}

$\tilde{X}^2\Sigma^+$ $C_{\infty v}$ Structure: LF⁹
 $B_0=0.135$ gas LF^{4,6,9} MW^{7,8}

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- ¹L. Pasternack and P. J. Dagdigian, J. Chem. Phys. **65**, 1320 (1976).
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⁶T. C. Steimle, D. A. Fletcher, K. Y. Jung, and C. T. Scurlock, J. Chem. Phys. **97**, 2909 (1992); **100**, 4025 (1994).
⁷T. C. Steimle, S. Saito, and S. Takano, Astrophys. J. **410**, L49 (1993).
⁸C. T. Scurlock, T. C. Steimle, R. D. Suenram, and F. J. Lovas, J. Chem. Phys. **100**, 3497 (1994).
⁹C. T. Scurlock, D. A. Fletcher, and T. C. Steimle, J. Chem. Phys. **101**, 7255 (1994).

cyc-BCC

\tilde{X}	C_{2v}
Vib. sym.	No. type of mode
a_1	2 CBC s-stretch

cm^{-1} Med. Type Refs.
1194.6 Ar IR 1,2

References

- ¹J. M. L. Martin, P. R. Taylor, J. T. Yustein, T. R. Burkholder, and L. Andrews, J. Chem. Phys. **99**, 12 (1993).
²J. D. Presilla-Márquez, C. W. Larson, P. G. Carrick, and C. M. L. Rittby, J. Chem. Phys. **105**, 3398 (1996).

YCC

$T_0=12889.5$ gas LF¹ 725–950 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	2	CY s-stretch	494.5(5.0)	gas	LF	1
	3	CY a-stretch	331.6(2.2)	gas	LF	1

 \tilde{X}^2A_1 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CY s-stretch	561.1(2.8)	gas	LF	1
b_2	3	CY a-stretch	369.6(1.7)	gas	LF	1

Reference

¹T. C. Steimle, A. J. Marr, J. Xin, A. J. Merer, K. Athanassenas, and D. Gillett, *J. Chem. Phys.* **106**, 2060 (1997).

cyc-BBN

In the argon-matrix study,² the $\nu_1 + \nu_3$ combination band at 1998.3 and other combination bands between 3250 and 6150 are moderately intense, suggesting strong vibronic coupling to a low-lying excited electronic state.

\tilde{X}		C_{2v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	BN s-stretch	1116 ^a	Ar	IR	2
b_2	3	BN a-stretch	882.3s	Ar	IR	2-4
			910.4	N ₂	IR	1
			890.3	N ₂	IR	1

^a($\nu_1 + \nu_3$) - ν_3 .

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- ¹P. Hassanzadeh and L. Andrews, *J. Phys. Chem.* **96**, 9177 (1992).
²L. Andrews, P. Hassanzadeh, T. R. Burkholder, and J. M. L. Martin, *J. Chem. Phys.* **98**, 922 (1993).
³C. A. Thompson and L. Andrews, *J. Am. Chem. Soc.* **117**, 10125 (1995).
⁴S. Li, R. J. Van Zee, and W. Weltner, Jr., *Chem. Phys. Lett.* **262**, 298 (1996).

BNB

\tilde{X}		$D_{\infty h}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1736.5	Ar	IR	1,2

References

- ¹L. Andrews, P. Hassanzadeh, T. R. Burkholder, and J. M. L. Martin, *J. Chem. Phys.* **98**, 922 (1993).
²C. A. Thompson and L. Andrews, *J. Am. Chem. Soc.* **117**, 10125 (1995).

Ga₂P

\tilde{X}		C_{2v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	281.2	Ar	IR	1

Reference

- ¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **97**, 11393 (1993).

Ga₂As

\tilde{X}		C_{2v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	205.4	Ar	IR	1

Reference

- ¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **97**, 11393 (1993).

Ga₂Sb

\tilde{X}		C_{2v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	167.4	Ar	IR	1

Reference

- ¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **97**, 11393 (1993).

In₂P

\tilde{X}		C_{2v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	249.3T	Ar	IR	1

Reference

- ¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **98**, 2275 (1994).

In₂As

\tilde{X}		C_{2v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	176.9T	Ar	IR	1

Reference

- ¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **98**, 2275 (1994).

In₂Sb

\tilde{X}		C_{2v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	143.9T	Ar	IR	1

Reference

¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **98**, 2275 (1994).

FeOFe⁻

Threshold for electron detachment from ground-state FeOFe⁻ = 13150(320) gas PE¹

Reference

¹H. Wu, S. R. Desai, and L.-S. Wang, *J. Am. Chem. Soc.* **118**, 5296 (1996).

CuOCu⁻

Threshold for electron detachment from ground-state CuOCu⁻ = 8880(240) gas PE¹

Reference

¹L.-S. Wang, H. Wu, S. R. Desai, and L. Lou, *Phys. Rev. B* **53**, 8028 (1996).

ScCO**X̄**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	CO stretch	1950brT	Ar	IR	1	

Reference

¹S. B. H. Bach, C. A. Taylor, R. J. Van Zee, M. T. Vala, and W. Weltner, Jr., *J. Am. Chem. Soc.* **108**, 7104 (1986).

TiCO**X̄**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	CO stretch	1854.4	Ar	IR	1	

Reference

¹G. V. Chertihin and L. Andrews, *J. Am. Chem. Soc.* **117**, 1595 (1995).

TiNN**X̄**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	NN stretch	1847.1	Ar	IR	1	
3	TiN stretch	766.2	Ar	IR	1	

¹L. Andrews, G. V. Chertihin, A. Citra, and M. Neurock, *J. Phys. Chem.* **100**, 11235 (1996).
²G. V. Chertihin, L. Andrews, and M. Neurock, *J. Phys. Chem.* **100**, 14609 (1996).

Reference

¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **98**, 5891 (1994).

VCO**X̄ 6Σ** **C_{∞v}** **Structure: ESR²**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ ⁺	1	CO stretch	1904	Ar	IR	1
			1890	Kr	IR	1
			1868	Xe	IR	1

References

- ¹L. Hanlan, H. Huber, and G. A. Ozin, *Inorg. Chem.* **15**, 2592 (1976).
²R. J. Van Zee, S. B. H. Bach, and W. Weltner, Jr., *J. Phys. Chem.* **90**, 583 (1986).

TaCO**X̄**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	CO stretch	1831T 1819T	Ar	IR	1

Reference

¹R. L. DeKock, *Inorg. Chem.* **10**, 1205 (1971).

CrCO**X̄**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	CO stretch	1977	Ar	IR	1

Reference

¹S. B. H. Bach, C. A. Taylor, R. J. Van Zee, M. T. Vala, and W. Weltner, Jr., *J. Am. Chem. Soc.* **108**, 7104 (1986).

NFeN**X̄** **C_{2v}**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b ₂	3	Asym. stretch	903.4 903.6	Ar N ₂	IR IR	2 1,2

References

FeNN**Reference**

\tilde{X}	C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
sym.					
a'	1	NN stretch	2017.8	Ar	IR 1
			2017.8	N ₂	IR 1
			2012.0	N ₂	IR 1

Reference

¹G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. **100**, 14609 (1996).

FeCO

$\tilde{\alpha}$	$5\Sigma^-$	$C_{\infty v}$			
$T_0=1135(25)$	gas	PE ³			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
sym.					
Σ^+	1	CO stretch	1990(15)	gas	PE 3
II	2	Bend	180(60)	gas	PE 3
Σ^+	3	FeC stretch	460(15)	gas	PE 3

\tilde{X}	$3\Sigma^-$	$C_{\infty v}$	Structure: MW ^{4,6} DL ⁵		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
sym.					
Σ^+	1	CO stretch	1946.47	gas	PE, DL 2,3,5
			1898T	Ar	IR 1
			1884T	Kr	IR 1
II	2	Bend	330(50)	gas	PE 3
Σ^+	3	FeC stretch	530(10)	gas	PE 3

$B_0=0.146$ MW^{4,6}DL⁵

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- ¹C. H. F. Peden, S. F. Parker, P. H. Barrett, and R. G. Pearson, J. Phys. Chem. **87**, 2329 (1983).
- ²R. J. Ryther and E. Weitz, J. Phys. Chem. **95**, 9841 (1991).
- ³P. W. Villalta and D. G. Leopold, J. Chem. Phys. **98**, 7730 (1993).
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- ⁵K. Tanaka, K. Sakaguchi, and T. Tanaka, J. Chem. Phys. **106**, 2118 (1997).
- ⁶K. Tanaka, M. Shirasaka, and T. Tanaka, J. Chem. Phys. **106**, 6821 (1997).

CoCO

\tilde{X}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
sym.					
1	CO stretch	1956	Ar	IR	1
		1949			
		1952	Kr	IR	1
		1944			
		1947	Xe	IR	1
		1941			

¹L. A. Hanlan, H. Huber, E. P. Kündig, B. R. McGarvey, and G. A. Ozin, J. Am. Chem. Soc. **97**, 7054 (1975).

NiCO

$T_0=11050(240)$ gas PE²

 \tilde{X}

\tilde{X}	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	CO stretch		1940(80)	gas	PE 2	
				1996	Ar	IR 1	

References

- ¹R. L. DeKock, Inorg. Chem. **10**, 1205 (1971).
- ²A. E. Stevens, C. S. Figgler, and W. C. Lineberger, J. Am. Chem. Soc. **104**, 5026 (1982).

PdCO \tilde{X}

\tilde{X}	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	CO stretch		2050	Ar	IR 1,2	
				2045	Kr	IR 2	
				316.8T	Kr	IR 2	

References

- ¹E. P. Kündig, M. Moskovits, and G. A. Ozin, Can. J. Chem. **50**, 3587 (1972).

- ²J. H. Darling and J. S. Ogden, J. Chem. Soc., Dalton Trans. **1973**, 1079.

PtCO \tilde{X}

\tilde{X}	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	CO stretch		2052	Ar	IR 1	

Reference

- ¹E. P. Kündig, D. McIntosh, M. Moskovits, and G. A. Ozin, J. Am. Chem. Soc. **95**, 7234 (1973).

CuCO \tilde{X}

\tilde{X}	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	CO stretch		2010.4	Ar	IR 1	

Reference

¹H. Huber, E. P. Kündig, M. Moskovits, and G. A. Ozin, J. Am. Chem. Soc. **97**, 2097 (1975).

CUO

\tilde{X} $C_{\infty v}$

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	UC stretch	852.6	Ar	IR	1
	3	UO stretch	804.4	Ar	IR	1

Reference

¹T. J. Tague, Jr., L. Andrews, and R. D. Hunt, J. Phys. Chem. **97**, 10920 (1993).

UCO

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	CO stretch	1832T 1817T	Ar	IR	1

Reference

¹J. L. Slater, R. K. Sheline, K. C. Lin, and W. Weltner, Jr., J. Chem. Phys. **55**, 5129 (1971).

NPuN

\tilde{X} $D_{\infty h}$

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1029.7	Ar	IR	1

Reference

¹D. W. Green and G. T. Reedy, J. Chem. Phys. **69**, 552 (1978).

C₃

$^1\Sigma_g^+$ $D_{\infty h}$
 $T_0=52826(30)$ Ar AB¹⁵

170–190 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1080(30)	Ar	AB	15
Π_u	2	Bend	300(30)H	Ar	AB	15
Σ_u^+	3	Asym. stretch	780(30)H ^a	Ar	AB	15

 \tilde{B}

In 2-photon ionization studies of jet-cooled C_3 , a complicated group of bands, all with rotational structure appropriate for a $\Sigma_u^+ - \Sigma_g^+$ vibronic transition arising from the \tilde{X} state, has been observed between 266 and 302 nm.²⁰ Lifetimes of these bands range from 0.4 to 2.5 μs , and the B' value for the first intense band, at 3153(5), is 0.396(3). These same bands, as well as bands at somewhat lower energies (possibly below the ionization threshold) and some bands arising from a $\Sigma_u^+ - \Pi_g$ vibronic transition, have also been studied using LF measurements on cooled beams.²³ In a neon matrix,³⁹ absorptions attributable to the $\tilde{B} - \tilde{X}$ transition of C_3 are observed between 281 and 302 nm, with the first intense band at 33149(20), and in an argon matrix,³⁰ absorptions arising from this transition of C_3 are observed between 280 and 284 nm.

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			444(20)	Ne	AB	39
			390	Ar	AB	30

In an argon matrix,³⁰ bands attributable to C_3 have been observed between 367 and 373 nm.

\tilde{A}	$^1\Pi_u$	$D_{\infty h}$	Structure: UV ⁶
$T_0=24675.5$	gas	$\text{EM}^{1,2,6}\text{AB}^{3,6,9}\text{LF}^{17,23,33,35,36,42,43}$	$\tilde{A} - \tilde{X}$ 335–640 nm
24640	Ne	$\text{AB}^{4,5,8}\text{EM}^2\text{LF}^{11}$	$\tilde{A} - \tilde{X}$ 347–488 nm
24370 ^b	Ar	$\text{AB}^{4,5,7}\text{LF}^{11}$	$\tilde{A} - \tilde{X}$ 352–411 nm
24350	Kr	AB ⁷	
23610	Xe	AB ^{4,7}	
24635	N ₂	AB ⁷	

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1085.9	gas	AB,LF	6,33
			1094(6)	Ne	AB	5
			1093(6)	Ar	AB	5,7
			1090	Kr	AB	7
			1120	Xe	AB	7
			1050	N ₂	AB	7
Π_u	2	Bend	307.9 ^c	gas	AB	6
Σ_u^+	3	Asym. stretch	541.7 ^d	gas	LF	36

$\tau_0=200(10)$ ns gas LF^{12,13}

In a neon or argon matrix,¹¹ efficient intersystem crossing into the $\tilde{a} - \tilde{3}\Pi_u$ state occurs, and $\tau \leq 10$ ns.
 $B_0=0.430$ UV⁶

\tilde{b}	$^3\Pi_g$	$D_{\infty h}$	Structure: DL,EM ²⁹
$T_0=23570(210)$	gas	$\text{DL}^{29}\text{EM}^{29,37}$	$\tilde{b} - \tilde{a}$ 1530–1640 nm
23584 ^e	Ne	IR ³²	

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Π_u	2	Bend	345(40) ^f	gas	EM	37
			$A=14.77(24)$ EM ³⁷			
			$B_0=0.424$ DL ²⁹ EM ²⁹			

$\tilde{a}^3\Pi_u$		D _{∞h}	Structure: DL,EM ²⁹
$T_0=17090(210)$	gas	DL ²⁹ EM ^{29,37} PE ³¹	$\tilde{b}-\tilde{a}$ 1530–1640 nm
$T_0=17080$	Ne	EM ⁵ LF ¹¹	$\tilde{a}-\tilde{X}$ 585–631 nm
	IR ³²		$\tilde{b}-\tilde{a}$ 1250–1538 nm
16930	Ar	EM ^{5,34}	
	IR ³²		$\tilde{b}-\tilde{a}$ 1253–1544 nm

Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.
sym.		type of mode			meas.	
Σ_g^+	1	Sym. stretch	1156.5 ^g	Ne	IR	32
			1154.2 ^g	Ar	IR	32
Π_u	2	Bend	505(40) ^h	gas	EM	37
Σ_u^+	3	Asym. stretch	1449.53	gas	DL	38
			1454.1	Ne	IR	32
			1455.3	Ar	IR	32

 $\tau \approx 0.02$ s Ne EM³ $A=15.16(4)$ EM³⁷ $B_0=0.417$ DL^{29,38}EM²⁹ $\tilde{X}^1\Sigma_g^+$ D_{∞h} Structure: UV⁶

This state of C₃ is highly anharmonic. The term values of many excited vibrational energy levels have been determined in SEP^{21,26,27} and LF²³ studies. Analysis of data from the SEP studies²⁸ using the semirigid bender model indicates that the molecule is linear in its ground state but that on excitation of ν_3 a potential barrier appears at the linear configuration.

Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.
sym.		type of mode			meas.	
Σ_g^+	1	Sym. stretch	1224.5	gas	AB,LF	9,17,23,33
			1226	Ne	EM	5
			1212(2)	Ar	IR,EM	19,34
			1215.8T	Kr	IR	41
			1234 ^g	N ₂	IR	19
Π_u	2	Bend	63.41 ⁱ	gas	UV,DL	6,22,23
					SEP,TF	24,25
			70T ^j	Ne	AB	5
			82H	Ar	EM	34
Σ_u^+	3	Asym. stretch	2040.02	gas	IR,DL	16,18
					LF	23
			2042	Ne	IR	4
			2038s	Ar	IR	4,10
			2033.3	Kr	IR	41
			2036.0	H ₂	IR	40
			2031	N ₂	IR	19

 $B_0=0.431$ UV⁶IR¹⁶DL¹⁸TF²⁵^aAlternate assignment gives 1320.^bIn the LF studies,¹¹ a second site was observed with $T_0=24408$.

^c ω . Large Renner splitting, with $\epsilon=0.537$.⁶ Detailed comparisons of gas-phase with neon- and argon-matrix band positions are given in Refs. 8 and 14. Ref. 14 also gives a more detailed analysis of electronic orbital angular momentum effects in the gas-phase molecule.

^dIn the gas phase,³⁵ the $\tilde{A}(002)-\tilde{X}(000)$ transition appears at 26348.0, giving $2\nu_3=1672.5$, and the corresponding transition observed in a neon matrix⁸ yields $2\nu_3=1680$. Ref. 36 suggests that this fundamental of C₃ $\tilde{A}^1\Pi_u$ has a large quartic anharmonicity.

^eSum of values for $\tilde{a}-\tilde{X}$ and $\tilde{b}-\tilde{a}$ transitions.^f ω . Large Renner splitting, with $\epsilon=0.447$.³⁷^g($\nu_1+\nu_3$) – ν_3 .^h ω . Large Renner splitting, with $\epsilon=0.566$.³⁷

ⁱMost precise value with tunable far infrared laser spectrometer (TF).²⁵ $\cong 45$ in $\tilde{X}(011)$.²²

^jGreatly broadened in a rare-gas matrix by interaction with lattice modes.¹¹

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SiC_C

\tilde{A}	1B_2	C_{2v}	Structure: Pi^5AB^{11}
$T_0=20065.505$	gas	$EM^1AB^{3,11}LF^{4,15}$	$\tilde{A}-\tilde{X}$ 402–507 nm
20142	Ne	AB^2LF^4	$\tilde{A}-\tilde{X}$ 409–611 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CC stretch	1462	gas	EM, AB	1,3,4,
			1462	Ne	AB, LF	11,15
			1457	Ar	AB	2,4
	2	CSi s-stretch	455.04	gas	LF, EM	4,11,15
			462	Ne	AB, LF	2,4
			448	Ar	AB	9
b_2	3	CSi a-stretch	487	gas	LF	15

$$\begin{aligned}\tau_0 &= 370 \text{ ns} & \text{gas} & LF^4 \\ & 310 \text{ ns} & \text{Ne} & LF^4 \\ A_0 &= 1.589; B_0 = 0.411; C_0 = 0.324 & AB^{11}\end{aligned}$$

\tilde{X}	1A_1	C_{2v}	Structure: $Pi^5MW^{6,7}AB^{11}$
The triangular configuration lies 1883(200) below the linear configuration. ¹⁷			

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CC stretch	1746.0(2.8)	gas	EM, LF	1,15
			1746s	Ne	IR, LF	2,4
			1741.3	Ar	IR	8,14
	2	CSi s-stretch	840.6(1.2)	gas	EM, LF	1,10,15
			836m	Ne	IR, LF	2,4
			824.3	Ar	IR	8,14
b_2	3	CSi a-stretch	196.37(4)	gas	MW, LF	13,15
			172H ^a	Ne	LF	4
			160.4	Ar	IR	14

$$A_0 = 1.729; B_0 = 0.436; C_0 = 0.348 \quad MW^{6,7,12,13,16}AB^{11}$$

^aIn neon-matrix emission measurements, Ref. 4 identified $2\nu_3$ at 344. The gas-phase term energy for $2\nu_3$, obtained in the SEP study of Ref. 17, is 352.85.

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¹⁶L. H. Couder, *J. Mol. Spectrosc.* **160**, 225 (1993).

¹⁷S. C. Ross, T. J. Butenhoff, E. A. Rohlfing, and C. M. Rohlfing, *J. Chem. Phys.* **100**, 4110 (1994).

Si₃

\tilde{G}	1B_2	C_{2v}
$T_0=25753(13)T$	Ne	AB^6

$\tilde{G}-\tilde{X}$ 330–390 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1			350T	Ne	AB	6

\tilde{F}	1B_1	C_{2v}
$T_0=19146(7)$	Ne	AB^6

$\tilde{F}-\tilde{X}$ 430–540 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	360T	Ne	AB	6

\tilde{E}	1A_1	C_{2v}
$T_0=17250(6)$	Ne	$AB^{1,6}$

$\tilde{E}-\tilde{X}$ 530–580 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	Bend	250T	Ne	AB	6

\tilde{D}	1B_1	C_{2v}
$T_0=13470(80)$	gas	PE^2

12839(3) Ne AB^6

$\tilde{D}-\tilde{X}$ 700–780 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	480(40)	gas	PE	2

\tilde{C}
$T_0=8880(80)$ gas PE ²

\tilde{B}
$T_0=7180(80)$ gas PE ²

\tilde{A}
$T_0=3630(80)$ gas PE ² EM ³

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	Sym. stretch	480(25)	gas	PE, EM	2,3

\tilde{A}'	$^3A'_2$	D_{3h}
$T_0 < 350$	gas	TPE ⁴

Vib.	No.	/	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1		Sym. stretch	501(10)	gas	TPE	4
e'_1	2		Deformation	337(10)	gas	EM, TPE	2–4

\tilde{X}^1A_1 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	550T	Ne	IR	5
			550.6	Ar	IR	5
			546.7	Kr	IR	5
<i>b</i> ₂	3	Asym. stretch	525.5	Ne	IR	5
			525.1	Ar	IR	5
			523.1	Kr	IR	5

References

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Ge₃ 1B_1 C_{2v}T₀=13640(100) gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			266T	gas	PE	1

 3B_1 C_{2v}T₀=8070(100) gas PE¹ 3A_1 C_{2v}T₀=6540(100) gas PE¹ 1B_2 C_{2v}T₀=1690(100) gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			355T	gas	PE	1

 $\tilde{X}\tilde{A}'^3A'_2, ^1A_1$ D_{3h}, C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>e'</i>	3		150T	gas	PE	1

Reference

- ¹G. R. Burton, C. Xu, C. C. Arnold, and D. M. Neumark, J. Chem. Phys. **104**, 2757 (1996).

OBeO⁺ \tilde{X} D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	988.6T	Ar	IR	1
			993.0T	Kr	IR	1
			993.6T	Xe	IR	1
			970.8T	N ₂	IR	2

References

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²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

NiCO⁻

Threshold for electron detachment from ground-state NiCO⁻ = 6490(100) gas PE¹

Reference

- ¹A. E. Stevens, C. S. Feigerle, and W. C. Lineberger, J. Am. Chem. Soc. **104**, 5026 (1982).

NFeO

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
FeN stretch		FeO stretch	971.1	N ₂	IR	1
			798.1	Ar	IR	2
			796.4	N ₂	IR	1

References

- ¹L. Andrews, G. V. Chertihin, A. Citra, and M. Neurock, J. Phys. Chem. **100**, 11235 (1996).
²G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. **100**, 14609 (1996).

FeNO

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NO stretch	1746.8 1731.0	N ₂	IR	1

Reference

- ¹L. Andrews, G. V. Chertihin, A. Citra, and M. Neurock, J. Phys. Chem. **100**, 11235 (1996).

CuOO⁺ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	OO stretch	1544.7T	Ar	IR	1

Reference

¹G. V. Chertikhin, L. Andrews, and C. W. Bauschlicher, Jr., *J. Phys. Chem. A* **101**, 4026 (1997).

NUO \tilde{X} $C_{\infty v}$

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	UN stretch	983.6	Ar	IR	1,2
	3	UO stretch	818.9	Ar	IR	2

References

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²G. P. Kushto, P. F. Souter, L. Andrews, and M. Neurock, *J. Chem. Phys.* **106**, 5894 (1997).

UNO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		NO stretch	1757.3	Ar	IR	1
			1749.2			

Reference

¹G. P. Kushto, P. F. Souter, L. Andrews, and M. Neurock, *J. Chem. Phys.* **106**, 5894 (1997).

AICO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
1		CO stretch	1867.7	Ar	IR	1-3
			1883.1	N ₂	IR	3
			1878.3			
3		AIC stretch	656.2wT	N ₂	IR	3

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GaCO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
1		CO stretch	1898.1	Ar	IR	1
			1886.5			
			1897.0	N ₂	IR	1
			1886.0			

Reference

¹A. Feltrin, M. Guido, and S. Nunziante Cesaro, *Vib. Spectrosc.* **8**, 175 (1995).

GaP₂ \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	322	Kr	IR	1
	2	Bend	220.9	Ar	IR	1

Reference

¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **97**, 11393 (1993).

GaAs₂ \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	231.0	Ar	IR	1
	2	Bend	171.4	Ar	IR	1

Reference

¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **97**, 11393 (1993).

GaSb₂ \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	183T	Ar	IR	1
	2	Bend	158T	Ar	IR	1

Reference

¹S. Li, R. J. Van Zee, and W. Weltner, Jr., *J. Phys. Chem.* **97**, 11393 (1993).

C₃⁻

Threshold for electron detachment from ground-state C₃⁻ = 15980(160) gas PE¹⁻³

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ _g ⁺	1	Sym. stretch	1075(100)	gas	PE	3
Σ _u ⁺	3	Asym. stretch	1721.8T	Ar	IR	4

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Si₃⁻

Threshold for electron detachment from ground-state Si₃⁻ = 18565(10) gas PE¹TPE²

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
2A ₁		C _{2v}				
T ₀ =9946(2)	Ne	AB ³				
			2A ₁ -X̄ 820-1040 nm			

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	Sym. stretch	495(5)	Ne	AB	3
	2	Bend	402(5)	Ne	AB	3

X̄ 2A₁**References**

- T. N. Kitsopoulos, C. J. Chick, A. Weaver, and D. M. Neumark, J. Chem. Phys. **93**, 6108 (1990).
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- J. Fulara, P. Freivogel, M. Grutter, and J. P. Maier, J. Phys. Chem. **100**, 18042 (1996).

Ge₃⁻

Threshold for electron detachment from ground-state Ge₃⁻ = 17990(80) gas PE¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
X̄ 2A ₁		C _{2v}				
			290T	gas	PE	1

Reference

¹G. R. Burton, C. Xu, C. C. Arnold, and D. M. Neumark, J. Chem. Phys. **104**, 2757 (1996).

CCN

C̄ 2Σ⁺ C_{∞v}
T₀=26661.80 gas AB¹LF¹⁴ **C̄-X̄ 310-375 nm**
Evidence for predissociation above 29100.¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ ⁺	1	Stretch	1859.20	gas	AB,LF	1,14
Π	2	Bend	469.9(4)	gas	AB,LF	1,14
Σ ⁺	3	Stretch	1257(4) ^a	gas	LF	14

B₀=0.413 AB¹LF¹⁴

B̄ 2Σ⁻ C_{∞v}
T₀=22413.25 gas AB¹ 22180 Ar AB³ **B̄-X̄ 442-446 nm**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π	2	Bend	445T	gas	AB	1

B₀=0.405 AB¹

Ā 2Δ C_{∞v}
T₀=21259.203 gas AB¹LF⁴EM⁹ **Ā-X̄ 376-471 nm**
21377 Ar LF²AB³ **Ā-X̄ 373-550 nm**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ ⁺	1	Stretch	1770.77	gas	AB	1
			1732(2)	Ar	LF	2
Π	2	Bend	457.1 ^a	gas	AB,LF	1,6,14
Σ ⁺	3	Stretch	1241.64	gas	AB	1
			1225(2)	Ar	LF	2

τ=170 ns Ar LF²
A_{eff}=-0.807 gas AB¹LF^{4,6}
B₀=0.414 AB¹LF^{4,6}MODR⁷

X̄ 2Π C_{∞v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ ⁺	1	Stretch	1923.25	gas	LF,EM	5,8,9
					LMR	12
					DL	10,11
Π	2	Bend	1717	Ar	LF	2
			324	gas	AB,LF	1,8
			179.27 ^b	gas	LF	14
Σ ⁺	3	Stretch	1050.76	gas	LF,EM	5,8,9
			1066	Ar	LF	2

A₀=42.328(2), εω₂=132.8 gas LF⁸DL^{10,11}LMR¹²
B₀=0.398 AB¹LF^{4,6}DL^{10,11}LMR¹²MW¹³

^aω.

^bμ²Σ component.

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SiNSi

Σ_u^+	$D_{\infty h}$	Structure: MPI ¹
$T_0 = 34314.29$ gas	Σ_u^+	$2\Sigma_u^+ - \tilde{X}$ 275–315 nm
$B_0 = 0.113$ MPI ¹		

\tilde{X} $2\Pi_{g,1/2}$	$D_{\infty h}$	Structure: MPI ¹
$B_0 = 0.112$ MPI ¹		

Reference

- ¹D. J. Brugh and M. D. Morse, Chem. Phys. Lett. **267**, 370 (1997).

OBeO

\tilde{X}	$D_{\infty h}$					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1431.4	Ne	IR	1
			1413.2	Ar	IR	1
			1408.2	Kr	IR	1
			1415.7	Xe	IR	1

Reference

- ¹C. A. Thompson and L. Andrews, J. Chem. Phys. **100**, 8689 (1994).

OMgO

\tilde{X}	$D_{\infty h}$					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	767.7	Ar	IR	1
			723.6	N ₂	IR	2

References

- ¹L. Andrews and J. T. Yustein, J. Phys. Chem. **97**, 12700 (1993).
- ²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

cyc-MgO₂

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	MgO ₂ s-stretch	621.1T	Ar	IR	1

Reference

- ¹L. Andrews and J. T. Yustein, J. Phys. Chem. **97**, 12700 (1993).

OCaO

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b_2	3	Asym. strctch	515.7	Ar	IR	1
			497.0	N ₂	IR	2

References

- ¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, J. Phys. Chem. **98**, 6514 (1994).
- ²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

cyc-CaO₂

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	O–O stretch	736.2	Ar	IR	1
			742.1	N ₂	IR	2
	2	OCaO s-stretch	555.7	N ₂	IR	2
b_2	3	OCaO a-stretch	505.5	N ₂	IR	2

References

- ¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, J. Phys. Chem. **98**, 6514 (1994).
- ²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

OSrO

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b_2	3	Asym. strctch	532.4	Ar	IR	1
			496.1	N ₂	IR	2

References

- ¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, J. Phys. Chem. **98**, 6514 (1994).
- ²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

cyc-SrO₂

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	O–O stretch	729.9	Ar	IR	1
			729.8	N ₂	IR	2
<i>a</i> ₂	2	SrO ₂ s-stretch	509.2	Ar	IR	1
			473.1	N ₂	IR	2

References

- ¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).
²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 10088 (1996).

OBaO

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> ₂	3	Asym. stretch	570.2	Ar	IR	1

Reference

- ¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).

cyc-BaO₂

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	O–O stretch	754.5	Ar	IR	1
	2	BaO ₂ s-stretch	468.3	Ar	IR	1

Reference

- ¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).

OCrO

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	895(20)	gas	PE	6
			914.4	Ar	IR	7
<i>a</i> ₂	2	Bend	220(20)	gas	PE	6
	3	Asym. stretch	965.4	Ar	IR	1–5,7

References

- ¹J. H. Darling, M. B. Garton-Sprenger, and J. S. Ogden, *Sympos. Faraday Soc.* **8**, 75 (1974).
²M. Poliakoff, K. P. Smith, J. J. Turner, and A. J. Wilkinson, *J. Chem. Soc., Dalton Trans.* 651 (1982).
³M. J. Almond, A. J. Downs, and R. N. Perutz, *Inorg. Chem.* **24**, 275 (1985).
⁴M. J. Almond and A. J. Downs, *J. Chem. Soc., Dalton Trans.* 809 (1988).
⁵M. J. Almond and M. Hahne, *J. Chem. Soc., Dalton Trans.* 2255 (1988).
⁶P. G. Wenthold, K.-L. Jonas, and W. C. Lineberger, *J. Chem. Phys.* **106**, 9961 (1997).
⁷G. V. Chertihin, W. D. Bare, and L. Andrews, *J. Chem. Phys.* **107**, 2798 (1997).

OMoO

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	948.5	Ne	IR	1
			938.7	Ar	IR	1,2
<i>b</i> ₂	3	Asym. stretch	899	Ne	IR	1
			885	Ar	IR	1,2

References

- ¹W. D. Hewett, Jr., J. H. Newton, and W. Weltner, Jr., *J. Phys. Chem.* **79**, 3640 (1975).
²M. J. Almond and A. J. Downs, *J. Chem. Soc., Dalton Trans.* 809 (1988).

OWO

$T_0=12672 \text{ Ne AB}^1 \quad 605\text{--}790 \text{ nm}$						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	972	Ne	AB	1
	2	Bend	300	Ne	AB	1

Structure: IR²

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	992	Ne	IR	1
			974T	Ar	IR	1
<i>b</i> ₂	3	Asym. stretch	975.5	Kr	IR	2
			928	Ne	IR	1
			921	Ar	IR	1
			937.2	Kr	IR	2

References

- ¹W. Weltner, Jr. and D. McLeod, Jr., *J. Mol. Spectrosc.* **17**, 276 (1965).
²D. W. Green and K. M. Ervin, *J. Mol. Spectrosc.* **89**, 145 (1981).

OTIO $T_0=18880$ Ne EM¹

529–621 nm

 \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	965T	gas	IR	2
			962.0	Ne	IR	1
			946.9	Ar	IR	3
<i>b</i> ₂	3	Asym. stretch	944	gas	IR	2
			934.8	Ne	IR	1
			917.1	Ar	IR	3

References¹N. S. McIntyre, K. R. Thompson, and W. Weltner, Jr., *J. Phys. Chem.* **75**, 3243 (1971).²T. C. DeVore and T. N. Gallaher, *High Temp. Sci.* **16**, 269 (1983).³G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).**OZrO** \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	884.3	Ar	IR	1
<i>b</i> ₂	3	Asym. stretch	818.0	Ar	IR	1

Reference¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).**OHfO** \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	883.4	Ar	IR	1
<i>b</i> ₂	3	Asym. stretch	814.0	Ar	IR	1

Reference¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).**OTaO** \tilde{B} Ne AB¹B– \tilde{X} 558–616 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	2	Bend	281	Ne	AB	1

³L. Andrews, G. V. Chertihin, A. Citra, and M. Neurock, J. Phys. Chem. **100**, 11235 (1996).

cyc-FeO₂

\tilde{X}	C _{2v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	OO stretch	956.0	Ar	IR	1-4
			954.5	O ₂	IR	4
<i>a</i> ₂	2	OFeO s-stretch	548.4	Ar	IR	3,4
			549.7	O ₂	IR	4

References

- ¹S. Chang, G. Blyholder, and J. Fernandez, Inorg. Chem. **20**, 2813 (1981).
- ²M. Fanfarillo, A. J. Downs, T. M. Greene, and M. J. Almond, Inorg. Chem. **31**, 2973 (1992).
- ³L. Andrews, G. V. Chertihin, A. Ricca, and C. W. Bauschlicher, Jr., J. Am. Chem. Soc. **118**, 467 (1996).
- ⁴G. V. Chertihin, W. Saffel, J. T. Yustein, L. Andrews, M. Neurock, A. Ricca, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 5261 (1996).

cyc-RhO₂

\tilde{X}	C _{2v}	Structure: IR ¹				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	OO stretch	908	Ar	IR	1
			900	Xe	IR	1
<i>a</i> ₂	2	RhO stretch	422	Xe	IR	1

Reference

- A. J. L. Hanlan and G. A. Ozin, Inorg. Chem. **16**, 2848 (1977).

ONiO

\tilde{b} 1Σ _g ⁺	D _{∞h}					
T ₀ =6210(240)	gas	PE ²				
\tilde{a} 1Δ _g	D _{∞h}					
T ₀ =3230(160)	gas	PE ²				
\tilde{X} 3Σ _g ⁻	D _{∞h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ _g ⁺	1	Sym. stretch	750(30)	gas	PE	2
Σ _u ⁺	3	Asym. stretch	954.9	Ar	IR	1
			959.5	N ₂	IR	1

References

- A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. A **101**, 3109 (1997).
- H. Wu and L.-S. Wang, J. Chem. Phys. **107**, 16 (1997).

cyc-NiO₂

\tilde{A}
T₀=8000T gas PE³

\tilde{X}	C _{2v}	Structure: IR ¹				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	OO stretch	967.1	Ar	IR	1,2
			970.1	N ₂	IR	2
<i>a</i> ₂	2	NiO s-stretch	538.3	Ar	IR	2
	3	NiO a-stretch	511.7	Ar	IR	2

References

- H. Huber, W. Klotzbücher, G. A. Ozin, and A. Vander Voet, Can. J. Chem. **51**, 2722 (1973).
- A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. A **101**, 3109 (1997).
- H. Wu and L.-S. Wang, J. Chem. Phys. **107**, 16 (1997).

NiOO

\tilde{X}	C _s					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OO stretch	1221.3	Ar	IR	1

Reference

- A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. A **101**, 3109 (1997).

cyc-PdO₂

\tilde{X}	C _{2v}	Structure: IR ¹				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	OO stretch	1023.5	Ar	IR	1
	2	PdO s-stretch	427w	Ar	IR	1

References

- H. Huber, W. Klotzbücher, G. A. Ozin, and A. Vander Voet, Can. J. Chem. **51**, 2722 (1973).

cyc-PtO₂

\tilde{X}	C _{2v}	Structure: IR ¹				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	OO stretch	926.6	Ar	IR	1

Reference

- H. Huber, W. Klotzbücher, G. A. Ozin, and A. Vander Voet, Can. J. Chem. **51**, 2722 (1973).

OCuO

$T_0=20699$	Ne	LF ³	432–540 nm
20290	Ar	LF ^{1,3}	
19680	Xe	UV ²	440–586 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	CuO s-stretch	610	Ne	LF	3	
		615(5)	Xe	AB	2	
2	Bend	133	Ne	LF	3	

\tilde{E}
 $T_0=13640(160)$ gas PE⁵

\tilde{D}
 $T_0=9680(160)$ gas PE⁵

\tilde{C}
 $T_0=6540(160)$ gas PE^{4,5}

\tilde{B}
 $T_0=5080(160)$ gas PE^{4,5}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			640(80)	gas	PE	4,5

\tilde{A}
 $T_0=2580(160)$ gas PE^{4,5}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			640(80)	gas	PE	4,5

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	CuO s-stretch	658	Ne	LF	3
			668	Ar	LF	1
			670(5)	Xe	EM	2
Π_u	2	Bend	193	Ne	LF	3
Σ_u^+	3	CuO a-stretch	823.0	Ar	IR	6
			826.7	N ₂	IR	6

References

- D. E. Tevault, J. Chem. Phys. **76**, 2859 (1982).
- G. A. Ozin, S. A. Mitchell, and J. García-Prieto, J. Am. Chem. Soc. **105**, 6399 (1983).
- V. E. Bondybey and J. H. English, J. Phys. Chem. **88**, 2247 (1984).
- H. Wu, S. R. Desai, and L.-S. Wang, J. Chem. Phys. **103**, 4363 (1995).
- H. Wu, S. R. Desai, and L.-S. Wang, J. Phys. Chem. A **101**, 2103 (1997).
- G. V. Chertihin, L. Andrews, and C. W. Bauschlicher, Jr., J. Phys. Chem. A **101**, 4026 (1997).

CuOO

In a xenon matrix, an absorption maximum at 44640 (224 nm) has been attributed¹ to CuOO.

\tilde{A}
 $T_0=7400(400)$ gas PE²

\tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	OO stretch	1089.6	Ar	IR	3
	3	CuO stretch	530(50)	gas	PE	2
			550.4	Ar	IR	3
			530	Xe	IR	1

References

- G. A. Ozin, S. A. Mitchell, and J. García-Prieto, J. Am. Chem. Soc. **105**, 6399 (1983).
- H. Wu, S. R. Desai, and L.-S. Wang, J. Chem. Phys. **103**, 4363 (1995).
- G. V. Chertihin, L. Andrews, and C. W. Bauschlicher, Jr., J. Phys. Chem. A **101**, 4026 (1997).

AgOO

\tilde{X} C_s Structure: IR¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	OO stretch	1084.4	Ar	IR	1
	3	AgO stretch	1078.9	Ar	IR	1

Reference

- D. E. Tevault, R. R. Smardzewski, M. W. Urban, and K. Nakamoto, J. Chem. Phys. **77**, 577 (1982).

cyc-AuO₂

In an argon matrix, relatively weak absorption maxima at 40980 (244 nm), 36230 (276 nm), and 31950 (313 nm) and a prominent, broad absorption maximum at 23640 (423 nm) have been attributed¹ to Au(OO).

\tilde{X} C_{2v} Structure: IR¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	OO stretch	1091.7	Ar	IR	1

Reference

- D. McIntosh and G. A. Ozin, Inorg. Chem. **15**, 2869 (1976).

OZnO

\tilde{X} D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	748.2	Ar	IR	1
			748.1	N ₂	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Chem. Phys. **106**, 3457 (1997).

OCdO

\tilde{X}	$D_{\infty h}$	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3			Asym. stretch	626.6	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Chem. Phys. **106**, 3457 (1997).

OCeO

$T_0=20067$	Ne	EM ¹	498–649 nm
19836	Ar	EM ¹	504–660 nm
19259	N ₂	EM ¹	519–633 nm
$\tau=200(50)$ ms	Ar	EM ¹	

\tilde{X}	C_{2v}	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1			Sym. stretch	777w	Ne	IR,EM	1
					757.0w	Ar	IR,EM	1,2
					714	N ₂	IR,EM	1
b_2	3			Asym. stretch	759s	Ne	IR	1
					736.9s	Ar	IR	1,2
					719	N ₂	IR	1

References

¹R. L. DeKock and W. Weltner, Jr., J. Phys. Chem. **75**, 514 (1971).

²S. D. Gabelnick, G. T. Reedy, and M. G. Chasanov, J. Chem. Phys. **60**, 1167 (1974).

OPrO

\tilde{X}	$D_{\infty h}$	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3			Asym. stretch	730.4	Ar	IR	1,2

References

¹R. L. DeKock and W. Weltner, Jr., J. Phys. Chem. **75**, 514 (1971).

²S. D. Gabelnick, G. T. Reedy, and M. G. Chasanov, J. Chem. Phys. **60**, 1167 (1974).

OTbO

\tilde{X}	C_{2v}	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1			Sym. stretch	758.8	Ar	IR	2
b_2	3			Asym. stretch	718.8	Ar	IR	1,2

References

¹R. L. DeKock and W. Weltner, Jr., J. Phys. Chem. **75**, 514 (1971).

²S. D. Gabelnick, G. T. Reedy, and M. G. Chasanov, J. Chem. Phys. **60**, 1167 (1974).

OTbO

\tilde{X}	C_{2v}	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1			Sym. stretch	787.4	Ar	IR	1
b_2	3			Asym. stretch	735.4	Ar	IR	1

Reference

¹S. D. Gabelnick, G. T. Reedy, and M. G. Chasanov, J. Chem. Phys. **60**, 1167 (1974).

OUO

\tilde{X}	$D_{\infty h}$	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3			Asym. stretch	776.1	Ar	IR	1,2
					768.0	Kr	IR	1

References

¹S. D. Gabelnick, G. T. Reedy, and M. G. Chasanov, J. Chem. Phys. **58**, 4468 (1973).

²R. D. Hunt and L. Andrews, J. Chem. Phys. **98**, 3690 (1993).

OPuO

\tilde{X}	$D_{\infty h}$	Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3			Asym. stretch	794.2	Ar	IR	1
					786.8	Kr	IR	1

Reference

¹D. W. Green and G. T. Reedy, J. Chem. Phys. **69**, 544 (1978).

CCO

\tilde{A}	$^3\Pi$	$C_{\infty v}$	Structure: AB ³
$T_0=11651.182$	gas	AB ^{2,3,14} LF ⁶ DL ^{10-12,14}	$\tilde{A}-\tilde{X}$ 500–860 nm
11860	Ar	AB ⁵	$\tilde{A}-\tilde{X}$ 600–850 nm
Vib.	No.	Approximate type of mode	cm^{-1}
Σ^+	1	CO stretch	2045.7
Π	2	Bend	594.75 ^a
Σ^+	3	CC stretch	1283.60

The fluorescence decay pattern⁷ of CCO $\tilde{A}(101)$ and of higher vibronic levels is complex. There is a short-lived ($\sim 15 \mu\text{s}$) component and a long-lived ($333 + 105/-64 \mu\text{s}$) component which is, in turn, nonexponential, suggesting perturbation by the heretofore unobserved $\tilde{B}^1\Sigma^+$ and $\tilde{a}^1\Delta$ states, as well as by high ground-state vibrational levels.

$A = -35.36$; $\varepsilon = -0.172$ gas $\text{AB}^3\text{DL}^{10}$

$B_0 = 0.407 \text{ AB}^3\text{DL}^{10}$

$\tilde{b}^1\Sigma^+$ $C_{\infty v}$
 $T_0 = 8190(145)$ gas PE¹⁵

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CO stretch	2010(170)	gas	PE	15

$\tilde{a}^1\Delta$ $C_{\infty v}$
 $T_0 = 5310(145)$ gas PE¹⁵

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CO stretch	1950(170)	gas	PE	15

$\tilde{X}^3\Sigma^-$ $C_{\infty v}$ Structure: AB^3

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CO stretch	1970.86	gas	LF,DL	6,9,16
			1969s	Ar	IR	1,4
			1978 N ₂	Ar	IR	1
			1987	N ₂	IR	1
II	2	Bend	379.53	gas	AB,DL	3,11
			381m	Ar	IR	1
Σ^+	3	CC stretch	1063	gas	LF	6
			1064w	Ar	IR	1
			1074 N ₂	Ar	IR	1
			1077	N ₂	IR	1

$B_0 = 0.385 \text{ AB}^3\text{MW}^{8,13}$

ω_a .

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GeCO

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CO stretch	1918.9	Ar	IR	2
			1908	Kr	IR	1
			1921.3	N ₂	IR	2
			1924.0	CO	IR	2

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SnCO

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	CO stretch	1921	Kr	IR	1

Reference

¹A. Bos, *J. Chem. Soc. Chem. Commun.* **1972**, 26 (1972).

NCN

\tilde{A}_u $D_{\infty h}$
gas AB^6 250–290 nm

$\tilde{B}^3\Sigma_u^-$ $D_{\infty h}$
 $T_0 = 33512$ gas AB^6 $\tilde{B}-\tilde{X}$ 258–300 nm
33100 Ar AB^2 $\tilde{B}-\tilde{X}$ 240–302 nm
33215 N₂ AB^2 $\tilde{B}-\tilde{X}$ 240–301 nm

In the gas phase, bands are diffuse. Threshold for photodecomposition into C+N₂ observed in argon and nitrogen matrices^{2,4} near 280 nm.

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1100(10)	gas	AB	6
			1050(10)	Ar,N ₂	AB	2

$\tilde{b}^1\Pi_u$ $D_{\infty h}$
 $T_0 = x + 30045.76$ gas AB^5 Structure: AB^5
 $\tilde{b}-\tilde{a}$ 330–334 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1160T	gas	AB	6

$\varepsilon\omega_2 = -84.2$ gas AB^5

$B_0 = 0.395 \text{ AB}^5$

\tilde{A}	$^3\Pi_u$	D _{∞h}	Structure: AB ¹ LF ¹⁰
$T_0=30383.967$	gas	AB ¹ LF ^{10,12}	$\tilde{A}-\tilde{X}$ 326–329 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1254(5)	gas	LF	8
Π_u	2	Bend	534T	gas	LF	12

$A=-37.57$; $\varepsilon \omega_2=-91.12$ gas AB¹LF¹²

$\tau_0=183(6)$ ns gas LF⁸

$B_0=0.397$ AB¹LF^{10,12}

$\tilde{\alpha}$	$^1\Lambda_g$	D _{∞h}	Structure: AB ⁵
$T_0=x$	gas	AB ⁵	$\tilde{b}-\tilde{\alpha}$ 330–334 nm
$B_0=0.399$	AB ⁵		

\tilde{X}	$^3\Sigma_g^-$	D _{∞h}	Structure: AB ¹ LF ¹⁰
Vib. sym.	No.	Approximate type of mode	cm ⁻¹
Σ_g^+	1	Sym. stretch	1197 ^a
Π_u	2	Bend	437T
			423m
Σ_u^+	3	Asym. stretch	1466.51
			LMR
			1475vs
			1478vs
			N ₂

$B_0=0.397$ AB¹LF^{9,10}IR⁹

^aFrequency deduced from weak combination with ν_3 which appears at 2672.

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N₃⁺

\tilde{A}	$^3\Pi_u$	D _{∞h}	Structure: PF ²
$T_0^a=35370$	gas	PF ²	$\tilde{A}-\tilde{X}$ 245–283 nm
Vib. sym.	No.	Approximate type of mode	cm ⁻¹
Σ_g^+	1	Sym. stretch	1300T
Π_u	2	Bend	440T
Σ_u^+	3	Asym. stretch	1700T

$\tau>50$ ps gas PF²

$A_{100}=-39.67$ gas PF²

$B_{100}=0.429$ gas PF²

\tilde{b}	$^1\Sigma^+$
$T_0=14520(160)$	gas PE ¹

$\tilde{\alpha}$	$^1\Delta$
$T_0=9120(160)$	gas PE ¹

\tilde{X}	$^3\Sigma_g^-$	D _{∞h}	Structure: PE ¹ PF ²
Vib. sym.	No.	Approximate type of mode	cm ⁻¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1170(30)	gas	PE	1

$B_0=0.424$ PF²

$^aT_{100}=36671.81$ gas PF².

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PbOPb

\tilde{X}	D _{∞h}
Vib. sym.	No.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	486.8	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Chem. Phys. **105**, 2561 (1996).

MgF₂⁺

\tilde{C}	$^2\Sigma_g^+$	D _{∞h}
$T^a=12500(560)$	gas	PE ¹

\tilde{B}	$^2\Sigma_u^+$	D _{∞h}
$T^a=7500(560)$	gas	PE ¹

\tilde{A}	$^2\Pi_u$	D _{∞h}
$T^a=5240(560)$	gas	PE ¹

\tilde{X}	$^2\Pi_g$	D _{∞h}

^aFrom vertical ionization potential.

Reference

¹J. M. Dyke, D. Haggerston, M. P. Hastings, and T. G. Wright, Chem. Phys. **181**, 355 (1994).

OTiO⁻

\tilde{X}	C _{2v}
Vib. sym.	No.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	878.4	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Phys. Chem. **99**, 6356 (1995).

OZrO⁻

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	761.4	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Phys. Chem. **99**, 6356 (1995).

OHfO⁻

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	747.9	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Phys. Chem. **99**, 6356 (1995).

OCrO⁻

Threshold for electron detachment from ground-state OCrO⁻ = 19470(65) gas EB¹PE²

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OFeO⁻

Threshold for electron detachment from ground-state OFeO⁻ = 19040(240) gas PE^{1,2}

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ONiO⁻

Threshold for electron detachment from ground-state ONiO⁻ = 24610(80) gas PE²

\tilde{X}	$D_{\infty h}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	886.8	Ar	IR	1
			894.4	N ₂	IR	1

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¹A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. A **101**, 3109 (1997).

²H. Wu and L.-S. Wang, J. Chem. Phys. **107**, 16 (1997).

cyc-NiO₂⁻

Threshold for electron detachment from ground-state cyc-NiO₂⁻ = 6620(240) gas PE¹

Reference

¹H. Wu and L.-S. Wang, J. Chem. Phys. **107**, 16 (1997).

OCuO⁻

Threshold for electron detachment from ground-state OCuO⁻ = 27920(320) gas PE¹

\tilde{X}	${}^1\Sigma_g^+$	$D_{\infty h}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	600(80)	gas	PE	2
Σ_u^+	3	Asym. stretch	839.6T	Ar	IR	3

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¹H. Wu, S. R. Desai, and L.-S. Wang, J. Chem. Phys. **103**, 4363 (1995).

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CuOO⁻

Threshold for electron detachment from ground-state CuOO⁻ = 12130(80) gas PE¹

\tilde{X}						
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	OO stretch	1009.5	Ar	IR	2

References

¹H. Wu, S. R. Desai, and L.-S. Wang, J. Chem. Phys. **103**, 4363 (1995).

²G. V. Chertihin, L. Andrews, and C. W. Bauschlicher, Jr., J. Phys. Chem. A **101**, 4026 (1997).

BS₂

$\tilde{B} \quad {}^2\Sigma_u^+ / Ne \quad D_{\infty h}$
 $T_0 = 24072(5) / Ne \quad AB^2$
 $gas \quad AB^{1,3}$
 $\tilde{B} - \tilde{X} \quad 395-412 \text{ nm}$
 $\tilde{B} - \tilde{X} \quad 409-418 \text{ nm}$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	509	Ne	AB	2

$\tilde{A}^2\Pi_u$		D _{∞h}	$\tilde{A}-\tilde{X}$ 514–721 nm	
$T_0=13766(2)$	Ne AB ²		$\tilde{A}-\tilde{X}$ 592–800 nm	
gas	AB ^{1,3}			

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1	Sym. stretch	504(2)	Ne	AB	2
Π_u	2	Bend	311H	Ne	AB	2
Σ_u^+	3	Asym. stretch	1535H	Ne	AB	2

$A = -263(2)$ Ne AB²

$\tilde{X}^2\Pi_g$		D _{∞h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1	Sym. stretch	510	Ne	AB	2
Π_u	2	Bend	120 ^{aT}	Ne	AB	2
Σ_u^+	3	Asym. stretch	1014.6(5)s	Ne	IR	2
			1015.7	Ar	IR	4

$A = -440$ gas AB^{1,2}

^aEstimated from isotope shift in origin of $\tilde{A}-\tilde{X}$ transition.

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CIBS⁺

$\tilde{C}^2\Sigma^+$		C _{∞v}
$T^a=50500(1000)$	gas	PE ¹

$\tilde{B}^2\Pi_{3/2}$		C _{∞v}
$T_0=26019$	gas	EF ²

$\tilde{B}-\tilde{X}$ 405–516 nm

$\tilde{A}^2\Sigma^+$		C _{∞v}
$T_0=24961.5(4)$	gas	EF ²

$\tilde{A}-\tilde{X}$ 392–440 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ^+	1	BS stretch	1390.6(8)	gas	EF	2
	3	BCl stretch	516.0(8)	gas	EF	2

$\tau=240(13)$ ns gas EF²

$\tilde{X}^2\Pi_{3/2}$		C _{∞v}	Structure: PE ^{1,3}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ^+	1	BS stretch	1347.8(8)	gas	EF	2
	3	BCl stretch	508.9(8)	gas	EF	2

$A = -383$ gas EF²
 $B_0=0.093$ EF²

^aFrom vertical ionization potential.

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OAO

$\tilde{A}^2\Pi_u$		D _{∞h}				
$T_0=17465$	Ar	LF ¹	495–620 nm			

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1	Sym. stretch	790	Ar	LF	1
Π_u	2	Bend	70	Ar	LF	1
Σ_u^+	3	Asym. stretch	1015	Ar	LF	1

$\tilde{B}^2\Sigma_g^+$		D _{∞h}
$T_0=6860(180)$	gas	PR ³

$\tilde{A}^2\Pi_u$		D _{∞h}
$T_0=5240(180)$	gas	PE ³

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1	Sym. stretch	810(60)	gas	PE	3

$\tilde{X}^2\Pi_g$		D _{∞h}
$T_0=26019$	gas	EF ²

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1	Sym. stretch	635	Ar	LF	1
Π_u	2	Bend	70	Ar	LF	1
Σ_u^+	3	Asym. stretch	1129.5	Ar	IR	2

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CCO⁻

Threshold for electron detachment from ground-state CCO⁻ is 18470(145).²

X

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
			1900T	gas	PE	1,2

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NCO

\tilde{B}	$^2\Pi$	$C_{\infty v}$	
$T_0 = 31751.1(5)$	gas	UV ² LF ^{18,30} PD ²⁷	$\tilde{B} - \tilde{X}$ 265–320 nm
31616(25)	Ne	UV ³	$\tilde{B} - \tilde{X}$ 260–320 nm
31437(25)	Ar	UV ³	$\tilde{B} - \tilde{X}$ 232–315 nm
31339(25)	N ₂	UV ³	$\tilde{B} - \tilde{X}$ 256–315 nm

All vibrational states are predissociated.²⁷

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	Stretch	2303	gas	UV	2
			2295(50)	Ne	UV	3
			2303(50)	Ar	UV	3
	3	Stretch	1047	gas	UV	2
			1033(50)	Ne	UV	3
			1053(50)	Ar	UV	3
			1025(50)	N ₂	UV	3

$\tau_0 = 63(3) - 152(5)$ ns for various rovibronic bands
 $A = -76.6$ gas LF¹⁸
 $B_0 = 0.356$ LF¹⁸

\tilde{A}	$^2\Sigma^+$	$C_{\infty v}$	Structure: UV ²⁰
$T_0 = 22754.020(2)$	gas	AB ¹ LF ^{23,30} SEP ²⁶ EM ²⁸	$\tilde{A} - \tilde{X}$ 304–512 nm
22800(10)	Ne	AB ³	$\tilde{A} - \tilde{X}$ 398–440 nm
22712(2)	Ar	LF ⁸	$\tilde{A} - \tilde{X}$ 390–530 nm
22956(10)	N ₂	AB ³	$\tilde{A} - \tilde{X}$ 395–440 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	Stretch	2338.0	gas	UV	1
			2325(20)	Ne	UV	3
			2332(4)	Ar	UV,LF	3,8
			2321(20)	N ₂	UV	3
Π	2	Bend	680.8	gas	UV	1
			673(20)	Ne	UV	3
Σ^+	3	Stretch	1289.3 ^a	gas	UV	1
			1270(20)	Ne	UV	3
			1291(4)	Ar	UV,LF	3,8

$\tau_0 = 435(10)$ ns gas LF^{9,13}
 $350(30)$ ns gas LF^{11,12}
 170 ns Ar LF⁸
 $B_0 = 0.402$ UV¹

\tilde{X}	$^2\Pi$	$C_{\infty v}$	Structure: UV ^{1,7,20} MW ^{4–6}
Vib.	No.	Approximate type of mode	cm ⁻¹
Σ^+	1	Sym. stretch	1266.63(8) ^b
			gas LF,LMR SEP 14,15,17 19,23,26,29
Π	2	Bend	1275vw
			Ar IR,LF 3,8
Σ^+	3	Asym. stretch	534.06(3) ^c
			gas UV,LF 1,7,15 SEP 23,29 LF 8
			529.5 ^d
			Ar LMR 10,14,15 LF,DL 21,23 EM 28 SEP 25
			1921.28 ^e gas Ar IR,LF 3,8
			1923m Ar IR 3
			1935 N ₂ IR 3

$A_0 = -95.589(3)$, $\epsilon \omega_2 = -78.37(3)$ gas LF²³SEP^{26,29}EM²⁸
 $B_0 = 0.390$ UV¹MW¹⁶LMR²²EM²⁸

^aIn Fermi resonance with $2\nu_2$, at 1385.3.

^bFor \tilde{X} $^2\Pi_{1/2}$, 1362.87 gas LF^{14,15,19,23,24}SEP²⁶

^c ω_2 .

^dLowest frequency component ($^2\Sigma^+$) contributes a strong infrared absorption at 487.³ Four components ($^2\Sigma^+$, $^2\Delta_{5/2}$, $^2\Delta_{3/2}$, $^2\Sigma^-$) observed at 484, 531, 626, and 672 in LF experiments.⁸ Components of (0n0), n ≥ 2, have also been assigned in gas-phase studies.^{19,23,26}

^eFor \tilde{X} $^2\Pi_{1/2}$, 2017.7(5) gas LF^{14,15,23,24}

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CO_2^+

$\tilde{C} \ ^2\Sigma_g^+$ D_{∞h} Structure: MP²⁴
 $T_0=45157(3)$ gas TPE²¹PE²³MP²⁴ $\tilde{C}-\tilde{A}$ 585–640 nm

The band origin is perturbed by another state of $2\Sigma_g^+$ symmetry at 45188 which has a rotational constant of 0.353.²⁴ This state may be an excited vibrational level of the \tilde{A} or \tilde{B} state.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
Σ_g^+	1	Sym. stretch	1352(4) ^a	gas	PE	23
Π_u	2	Bend	614(4)	gas	TPE, PE	21,23
Σ_u^+	3	Asym. stretch	1567(4)	gas	PE	23

$$B_0=0.395 \text{ MP}^{24}$$

$\tilde{B} \ ^2\Sigma_u^+$ D_{∞h} Structure: EM⁹
 $T_0=34591.6$ ^b gas EM^{1,9,29,30}EF²⁹ $\tilde{B}-\tilde{X}$ 287–291 nm

Perturbations by the \tilde{A} state are considered in Refs. 14–16 and Ref. 29.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
Σ_g^+	1	Sym. stretch	1275(1)	gas	TPE, PE	21,23
Π_u	2	Bend	557(4)	gas	EM,PE	9,23
Σ_u^+	3	Asym. stretch	1847(10)	gas	PE,EM	17,23,30

$$\tau_0=140(7) \text{ ns} \text{ gas T-PEFCO}^{10}\text{PEFCO}^{13}\text{LF}^{16}$$

$$B_0=0.380 \text{ EM}^{1,29}$$

$\tilde{A} \ ^2\Pi_u$ D_{∞h} Structure: EM¹¹
 $T_0=28500.35$ gas EM^{2,6,11,30}LF^{27,30}PE³¹ $\tilde{A}-\tilde{X}$ 253–490 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
Σ_g^+	1	Sym. stretch	1126	gas	EM PE	2,5,11 23,31
Π_u	2	Bend	461	gas	EM,PE	11,23,31
Σ_u^+	3	Asym. stretch	2685.3	gas	LF,EM	30

$$\tau_0=102(8) \text{ ns} \text{ gas EF}^7\text{T-PEFCO}^{10}$$

$$124(6) \text{ ns} \text{ gas PEFCO}^{13}\text{HF}^{18}$$

$$A=-95.51 \text{ gas EM}^{11}\text{LF}^{27}\text{PE}^{31}$$

$$B_0=0.350 \text{ EM}^{2,11}\text{LF}^{27}$$

$\tilde{X} \ ^2\Pi_g$	D _{∞h}	Structure: EM ^{2-5,9,11}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
Σ_g^+	1	Sym. stretch	1244.3(3)	gas	EM,DL PE	4,5,8,12 22,31
Π_u	2	Bend	511.4(3)	gas	EM,DL PE	11,19,22 28,31
				TPE		
Σ_u^+	3	Asym. stretch	462.6 ^c 1423.08 1421.7	Ne gas Ne	IR DL,PE IR	32 19,31 25,32

$$A=-161.02(6),^d \varepsilon \omega_2=-98.8(3)^d \text{ gas EM}^{1,9,11}\text{DL}^{20,22}\text{TPE}^{28}$$

$$B_0=0.380 \text{ EM}^{1,3,9,11}$$

^aCorrected for Fermi resonance.

^bMeasured from lowest rotational level of \tilde{X} state,²⁹ 34672.33.

^c $\mu \ ^2\Sigma_u^+$ level (gas-phase value²⁰)=467.26.

^dReanalysis by Ref. 26 gives $A=-161.48(5)$ and $\varepsilon \omega_2=-100.4$.

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OCS⁺

$\tilde{C} \ ^2\Sigma^+$ C_{∞v}
 $T_0=54640(30)$ gas PI⁴PE¹⁰

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
Σ^+	1	CO stretch	2202(2)	gas	PE	10
Π	2	Bend	454(5)	gas	PE	10
Σ^+	3	CS stretch	926(5)	gas	PI,PE	4,10

$\tilde{B} \ ^2\Sigma^+$ C_{∞v}
 $T_0=39180(20)$ gas PI⁴PF⁸PE¹⁰

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
Σ^+	1	CO stretch	1850(8)	gas	PE	10
Π	2	Bend	515(3)	gas	PF,PE	9,10
Σ^+	3	CS stretch	829 ^a	gas	PF	9

$\tilde{A}^2\Pi_{3/2}$		$C_{\infty v}$				
$T_0=31404.099(7)$ gas		$EF^1LF^8PF^{9,11}$		$\tilde{A}-\tilde{X}$ 282–432 nm		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CO stretch	2027	gas	PE,PF	10,11
Π	2	Bend	336(20)H	gas	PE	10
Σ^+	3	CS stretch	807 ^b	gas	PF	9,11
$\tau_0=93(9) \text{ ns}^c$ gas PEFCO ⁵						
$\tau_0(\omega=3/2)=105(3) \text{ ns}; \tau_0(\omega=1/2)=77(3) \text{ ns}$ gas HFD ⁶ EF ⁷						
$A=-111.8 \text{ gas } EF^1PF^{9,11}$						
$B_0=0.186 \text{ LF}^8PF^{9,11}$						

$\tilde{X}^2\Pi_{3/2}$		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CO stretch	2066	gas	EF,PF	1,11
Π	2	Bend	476(16)	gas	PE	10
Σ^+	3	CS stretch	697 ^d	gas	PF	9,11
$A=-367.2 \text{ gas } EF^1PF^9$						
$B_0=0.194 \text{ LF}^8PF^{9,11}$						

^aRef. 10 gives value of 742(7).^b817 for $\omega=1/2,9,11$

^cAbsence of emission from states above the $\tilde{A}^2\Pi$ band origin in photoionization experiments² suggested that the molecule is predissociated into CO + S⁺ (⁴S⁰), as was later confirmed.³ PEFCO studies⁵ have yielded the branching ratio for photexcitation vs predissociation for the transition origin, permitting an estimate of 550(50) ns for the radiative lifetime.

^d699.7 for $\omega=1/2$.^{9,11}

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CS₂⁺

$\tilde{C}^2\Sigma_g^+$		$D_{\infty h}$		Structure: MP ¹⁵		
$T_0=49064$ gas		$PI^2PF^{13}MP^{15}$		$\tilde{C}-\tilde{B}$ 658–724 nm		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	652(2)	gas	PI,PE PF,MP	5,9,11 13–15,19
Π_u	2	Bend	348(9)	gas	PF,PE	13,14,19
Σ_u^+	3	Asym. stretch	1024(6)T	gas	PE	14
$\tau_0=11(2) \text{ ps}$ gas MP ¹⁵						
$B_0=0.111 \text{ PI}^2PF^{13}MP^{15}$						

$\tilde{B}^2\Sigma_u^+$		$D_{\infty h}$		Structure: EM ¹		
$T_0=35238.01$ gas		EM^1		$\tilde{B}-\tilde{X}$ 277–307 nm		
		35270	Ne	LF ⁷		
		35226				

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	602	gas	EM,PE	3,19
Π_u	2	Bend	351(5)	gas	PE	14,19
Σ_u^+	3	Asym. stretch	1320(5)	gas	PE	14,19

$\tau_0=290(10) \text{ ns}$ gas EF²PIFCO⁴PEFCO⁸UV¹²
There is also a long-lifetime component, with $\tau=1.44(22) \mu\text{s}$.^{8,12}

 $B_0=0.108 \text{ EM}^1$

$\tilde{A}^2\Pi_u$		$D_{\infty h}$		Structure: EM ³		
$T_0=20975$ gas		EM^3		$\tilde{A}-\tilde{X}$ 426–512 nm		
		21017	Ne	LF ^{6,7,17}		$\tilde{A}-\tilde{X}$ 400–638 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	550 ^T	gas	EM	3
			557	Ne	LF	6,7,18
Π_u	2	Bend	305HT	gas	EM	3
			309H	Ne	LF	6,7,18
Σ_u^+	3	Asym. stretch	1644H	Ne	LF	7

$\tau=4.09(19) \mu\text{s}$ gas PIFCO⁴ID¹⁰UV¹²
 $2.3(1) \mu\text{s}$ Ne LF^{6,7}

 $A=-176 \text{ gas } EM^3PE^{14,19}$ $B_0=0.101 \text{ EM}^3$

$\tilde{X}^2\Pi_g$		$D_{\infty h}$		Structure: EM ¹		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.

Σ_g^+	1	Sym. stretch	617 ^a	gas	EM,TPE	3,16,21
			618 ^a	Ne	LF	6,7,17,18
Π_u	2	Bend	332 ^b	gas	TPE	16,21
			349H	Ne	LF	6,7,17,18
Σ_u^+	3	Asym. stretch	1188(8)	gas	EM,PE	3,14,19
			1207.1	Ne	LF,IR	6,7,17,18

 $A=-440.39(3) \text{ gas } EM^1TPE^{16,21}$ $B_0=0.109 \text{ EM}^1$

^aStrong Fermi resonance with $2\nu_2$; assignment of Refs. 7 and 18 adopted for the \tilde{A} state.

^bRenner-Teller components are observed for $\tilde{X}^2\Pi_{g,3/2}$ at 330 (² Σ^+) and 336 (² $\Delta_{3/2}$) in a gas-phase TPE study²¹. For $\tilde{X}^2\Pi_{g,1/2}$, components at 320 (² $\Delta_{u,3/2}$) and 341 (² Σ_u^-) are observed in gas-phase TPE studies.^{16,21}

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BrCN⁺

$\tilde{C}^{\infty}\Sigma^{+}$ C_{∞v}
 $T_0=50200(200)$ gas PE^{1,2,13}

$\tilde{B}^{\infty}\Pi_{3/2}$ C_{∞v}
 $T_0=18759.78(4)$ gas EF⁷LF^{10,11}PE¹³
 $18586(14)$ Ne AB⁵ $\tilde{B}-\tilde{X}$ 445–620 nm
 $\tilde{B}-\tilde{X}$ 418–538 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	C≡N stretch	1958(2) 1830(10)	gas Ne	EF,LF,PE AB	7,10,13 5
Π	2	Bend	395.7(2)H 377(10)	gas Ne	EF,LF AB	7,10 5
Σ^+	3	CBr stretch	473.1(2) 478(10)	gas Ne	EF,LF,PE AB	7,10,13 5

$\tau_1=197(10)$ ns gas EF³PIFCO⁶T-PEFCO⁸

$\tau_2=713(40)$ ns gas PIFCO⁶T-PEFCO⁸

$A=-774(20)$ gas PE¹³

$B_0=0.127$ gas LF^{10,11}

$\tilde{A}^{\infty}\Sigma^{+}$ C_{∞v}
 $T_0=13699(1)$ gas EF⁷PE¹³ $\tilde{A}-\tilde{X}$ 708–853 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CN stretch	1930(2)	gas	EF	7
Π	2	Bend	421(2)	gas	EF	7
Σ^+	3	CBr stretch	584(2)	gas	EF	7

$\tau=2750(100)$ ns gas T-PEFCO⁸

$\tilde{X}^{\infty}\Pi_{3/2}$ C_{∞v} Structure: LF¹¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	C≡N stretch	1905.94	gas	EF,LF DL,PE	7,10 12,13
Π	2	Bend	287.2(2)H	gas	EF,LF	7,10
Σ^+	3	CBr stretch	649.4(5)	gas	EF,LF PE	7,10,13

$A=-1477(2)$ gas EF³PIFCO⁴
 $B_0=0.141$ gas LF^{10,11}DL¹²

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PNP

$\tilde{A}^{\infty}\Sigma_u^+$ D_{∞h}
 $T_0=28300$ Ar AB^{1,2}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	560(20)	Ar	AB	1,2

$\tilde{X}^{\infty}\Pi_g$ D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1179.3	Ar	IR	2

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N₂O⁺

$\tilde{C}^{\infty}\Sigma^{+}$ C_{∞v}
 $T_0=58245(32)$ gas PE¹PI⁵

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	Sym. stretch	1280(50)	gas	PE	1
	3	Asym. stretch	2300(50)	gas	PE	1

$\tilde{B}^2\Pi$		$C_{\infty v}$	$\tilde{B}-\tilde{A}$ 538–866 nm			
$T_0=38440(100)^a$		gas PE ¹ PF ¹⁵				

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	Sym. stretch	900 ^b T	gas	PE	1

$\tilde{A}^2\Sigma^+$		$C_{\infty v}$	Structure: EM ³ PF ¹⁶			
$T_0=28162.33$		gas EM ^{3,14} PF ^{6,10,11,16,18}	$\tilde{A}-\tilde{X}$ 317–421 nm			

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	Sym. stretch	1345.52	gas	EM,PF	3,10
Π	2	Bend	614.45	gas	EM	3,14
Σ^+	3	Asym. stretch	2451.7	gas	EM	3

$\tau=230(10)$ ns gas $\text{BF}^2\Pi\text{FCO}^4\text{PE}\text{FCO}^7\text{ID}^8\text{EM}^{12}\text{HFD}^{13}$
 $B_0=0.433$ EM^{3,14}PF^{10,11}

$\tilde{X}^2\Pi$		$C_{\infty v}$	Structure: EM ³			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	Sym. stretch	1126.51	gas	EM	3
			1135.5	Ne	IR	17
Π	2	Bend	452.42	gas	EM,PF	3,11,14
Σ^+	3	Asym. stretch	1737.6	gas	EM	3
			1741.4wm	Ne	IR	17

$A=-132.434$, $\epsilon\omega_2=-90.2$ gas EM^{3,14}PF^{11,16,18}
 $B_0=0.412$ EM³PF^{10,11,16}

^aCalculated using first ionization potential of 12.886(2) eV, from Ref. 5.
^bSomewhat irregular band spacings.

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ArBeO

 \tilde{X}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BeO stretch	1526.1	Ar	IR	1

Reference

¹C. A. Thompson and L. Andrews, J. Am. Chem. Soc. **116**, 423 (1994).

KrBeO

 \tilde{X}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BeO stretch	1521.8	Ar	IR	1
			1511.8	Kr	IR	2

References

¹C. A. Thompson and L. Andrews, J. Am. Chem. Soc. **116**, 423 (1994).
²C. A. Thompson and L. Andrews, J. Chem. Phys. **100**, 8689 (1994).

XeBeO

 \tilde{X}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BeO stretch	1516.7	Ar	IR	1
			1498.3	Xe	IR	2

References

¹C. A. Thompson and L. Andrews, J. Am. Chem. Soc. **116**, 423 (1994).
²C. A. Thompson and L. Andrews, J. Chem. Phys. **100**, 8689 (1994).

FBO

 \tilde{X} $C_{\infty v}$ Structure: MW,DL²

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	B=O stretch	2078.87	gas	DL	2
			2081	Ne	IR	1,3
			2071	Ar	IR	1
Π	2	Bend	502	Ne	IR	1,3
			493	Ar	IR	1

$B_0=0.312$ MW,DL²

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²Y. Kawashima, K. Kawaguchi, Y. Endo, and E. Hirota, J. Chem. Phys. **87**, 2006 (1987).
³M. E. Jacox and W. E. Thompson, J. Chem. Phys. **102**, 4747 (1995).

CIBO

\tilde{X}		$C_{\infty v}$	Structure: MW,DL ²			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1	B=O stretch	1958s	Ar	IR	1
	2	Bend	404s	Ar	IR	1
	3	BCl stretch	676.04	gas	DL	2
			673wm	Ar	IR	1

 $B_0=0.174 \text{ MW,DL}^2$ **References**

- ¹A. Snelson, High Temp. Sci. **4**, 318 (1972).
²K. Kawaguchi, Y. Endo, and E. Hirota, J. Mol. Spectrosc. **93**, 381 (1982).

CIBS

\tilde{X}		$C_{\infty v}$	Structure: MW ¹			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	BS stretch	1408	gas	IR	3,4
Π	2	Bend	300(40)	gas	MW	2
Σ^+	3	BCl stretch	525	gas	IR	3,4

 $B_0=0.093 \text{ MW}^{1,2}$ **References**

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BF₂⁺

$\tilde{X} \ ^1\Sigma_g^+$		$D_{\infty h}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	2026.1	Ne	IR	1

Reference

- ¹M. E. Jacox and W. E. Thompson, J. Chem. Phys. **102**, 4747 (1995).

BCl₂⁺

Emission between 280 and 350 nm which results on excitation of BCl₃ by radiation with a threshold of 74 nm has been attributed¹ to BCl₂⁺. A band spacing of 650(30) observed in this emission has not been definitively assigned.

\tilde{X}		$D_{\infty h}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1439.7	Ne	IR	2
			1436.3			

References

- ¹H. Biehl, J. C. Creasey, D. M. Smith, R. P. Tuckett, K. R. Yoxall, H. Baumgartel, H. W. Jochims, and U. Rockland, J. Chem. Soc. Faraday Trans. **91**, 3073 (1995).

- ²M. E. Jacox, K. K. Irikura, and W. E. Thompson, J. Chem. Phys. **104**, 8871 (1996).

BBr₂⁺

Emission between 250 and 300 nm, with a maximum at 290 nm (34500), which appears when BBr₃ is excited by radiation with a threshold at 78 nm (15.8 eV), has been attributed¹ to transitions of BBr₂⁺.

Reference

- ¹H. Biehl, D. M. Smith, R. P. Tuckett, K. R. Yoxall, H. Baumgartel, H. W. Jochims, and U. Rockland, Mol. Phys. **87**, 1199 (1996).

OAIO⁻

Threshold for electron detachment from ground-state OAIO⁻=34130(160) gas PE¹

Reference

- ¹S. R. Desai, H. Wu, C. M. Rohlfing, and L.-S. Wang, J. Chem. Phys. **106**, 1309 (1997).

OSiSe

\tilde{X}		$C_{\infty v}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	Si=O stretch	1254.2	Ar	IR	1

Reference

- ¹L. Andrews, P. Hassanzadeh, D. V. Lanzisera, and G. D. Brabson, J. Phys. Chem. **100**, 16667 (1996).

SiSe₂

\tilde{X}		$D_{\infty h}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	SiSe a-stretch	750.8	Ar	IR	1

Reference

- ¹L. Andrews, P. Hassanzadeh, D. V. Lanzisera, and G. D. Brabson, J. Phys. Chem. **100**, 16667 (1996).

OSnO

\tilde{X}		$D_{\infty h}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	863.1	Kr	IR	1
			877.8	N ₂	IR	1

Reference

- ¹A. Bos and J. S. Ogden, J. Phys. Chem. **77**, 1513 (1973).

OPbO

\tilde{X}	$D_{\infty h}$					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+_g	1	Sym. stretch	658.67 ^a	Ar	IR	1
Σ_u^+	3	Asym. stretch	764.8 775.9	Ar N_2	IR	1

^aFrom combination band.

Reference

¹G. V. Chertihin and L. Andrews, *J. Chem. Phys.* **105**, 2561 (1996).

cyc-PbO₂

\tilde{X}	C_{2v}					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	OO stretch	728.7 730.6	Ar N_2	IR IR	1,2 2
	2	PbO stretch	437.3 449.6	Ar N_2	IR IR	1,2 2

References

¹S. A. Konnov, L. V. Serebrennikov, and A. A. Mal'tsev, *Zh. Neorg. Khim.* **27**, 323 (1982); *Russ. J. Inorg. Chem.* **27**, 184 (1982).

²G. V. Chertihin and L. Andrews, *J. Chem. Phys.* **105**, 2561 (1996).

FCP

\tilde{X}	$C_{\infty v}$		Structure: MW ¹		
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
a'	1	C≡P stretch	1670.84	gas	IR
	2	Bend	375.43	gas	IR
	3	CF stretch	780.1(2)	gas	IR

$B_0=0.175$ MW¹IR²

References

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CICP

\tilde{X}	$C_{\infty v}$		Structure: MW ²		
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.
a'	1	C≡P stretch	1477.34	gas	IR
	2	Bend	303.41	gas	IR
	3	CCl stretch	573.92(6)	gas	IR

$B_0=0.101$ MW^{1,2}IR³

References

¹T. J. Dennis, S. Firth, H. W. Kroto, D. R. M. Walton, and C.-Y. Mok, *J. Chem. Soc., Chem. Commun.* 1430 (1990).

²S. Firth, S. Khalaf, and H. W. Kroto, *J. Chem. Soc. Faraday Trans.* **88**, 3393 (1992).

³D. McNaughton and D. N. Bruget, *J. Mol. Spectrosc.* **161**, 336 (1993).

NO₂⁺

$\tilde{\mathcal{B}}_2$
 $T_0^a=74580(100)$ C_{2v}
gas PE^{1,2}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1113(20)	gas	PE	1,2
	2	Bend	686(20)	gas	PE	1,2

\tilde{d}^3A_1
 $T^a=67600$ C_{2v}
gas PE²

\tilde{C}^1B_1	$T_0^a=60670(100)$	C_{2v}	gas	PE ²		
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1017(20)	gas	PE	1,2

\tilde{c}^3B_1
 $T_0^a=60100(100)$ C_{2v}
gas PE^{1,2}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1041(20)	gas	PE	1,2
	2	Bend	573(20)	gas	PE	1,2

\tilde{B}^1B_2
 $T_0^a=38940(100)$ C_{2v}
gas PE^{1,2}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1025(20)	gas	PE	2
	2	Bend	573(20)	gas	PE	1,2

\tilde{A}^1A_2
 $T_0^a=35900(100)$ C_{2v}
gas PE^{1,2}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	984(20)	gas	PE	1,2
	2	Bend	694(20)	gas	PE	1,2

\tilde{b}^3A_2
 $T_0^a=32110(100)$ C_{2v}
gas PE^{1,2}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	Bend	662(20)	gas	PE	1,2

\tilde{a}^3B_2 C_{2v}
 $T_0^a = 26170(100)$ gas PE^{1,2}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	Bend	654(20)	gas	PE	1,2
$\tilde{X}^1\Sigma_g^+$	D _{>h}	Structure: TPE ^{4,5,7}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1396	gas	TPE	5,7
			1362.4 ^b	Ne	IR	6
II	2	Bend	639	gas	TPE	5,7
Σ_u^+	3	Asym. stretch	2362	gas	TPE	5,7
			2348.2	Ne	IR	6

$B_0 = 0.417$ TPE^{4,7}

^aThe band origins given here have been calculated using a first ionization potential of 9.62(1) eV for NO₂, as found in the photoionization study of Ref. 3.

^b($\nu_1 + \nu_3$) - ν_3 .

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SNO⁺

\tilde{X}	C _{>v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	NO stretch	1975.2	Ar	IR	1

Reference

- L. Andrews, P. Hassanzadeh, G. D. Brabson, A. Citra, and M. Neurock, J. Phys. Chem. **100**, 8273 (1996).

SNS⁺

\tilde{X}	D _{>h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1499.7	Ar	IR	1,2

References

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- L. Andrews and P. Hassanzadeh, J. Chem. Soc., Chem. Commun. 1523 (1994).

SeNSe⁺

\tilde{X}	D _{>h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1250.7	Ar	IR	1,2

References

¹L. Andrews and P. Hassanzadeh, J. Chem. Soc., Chem. Commun. 1523 (1994).

²L. Andrews, P. Hassanzadeh, D. V. Lanzisera, and G. D. Brabson, J. Phys. Chem. **100**, 16667 (1996).

P₂O

\tilde{X}	C _{>v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	P=O stretch	1278.677	gas	DL	2
			1270.4	Ar	IR	1

References

¹Z. Mielke, M. McCluskey, and L. Andrews, Chem. Phys. Lett. **165**, 146 (1990).

²H.-B. Qian, P. B. Davies, and P. A. Hamilton, J. Chem. Soc. Faraday Trans. **91**, 2993 (1995).

INC

\tilde{X}	C _{>v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	NC stretch	2057	Ar	IR	1,2
			2063	Kr	IR	1
			2055			
			2053	Xe	IR	1
			2058	N ₂	IR	1
II	2	Bend	200T	Ar	IR	2
Σ^+	3	IN stretch	494	Ar	IR	2

References

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²U. Samuni, S. Kahana, R. Fraenkel, Y. Haas, D. Danovich, and S. Shaik, Chem. Phys. Lett. **225**, 391 (1994); Chem. Phys. Lett. **231**, 124 (1994).

BF₂

\tilde{C}	D _{>h}					
T _{vert}	gas	MPI ⁶				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Π_g	2	Bend	600(50)	gas	MPI	6

\tilde{B} 2A_1 (3s)	C_{2v}		
$T_{\text{vert}} = 59100 \text{ T}$	gas	$\text{EF}^1\text{EM}^{2,3}\text{MPI}^6$	$\tilde{B}-\tilde{A}, \tilde{X} 190-650 \text{ nm}$
$\tau \approx 11 \text{ ns}$	EF^1EM^3		

\tilde{A} 2B_1	C_{2v}		
$T_{\text{vert}} = 35100$	gas	$\text{EF}^1\text{EM}^{2,3}\text{MPI}^6$	$\tilde{A}-\tilde{X} 220-650 \text{ nm}$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	Bend	310(20)	gas	MPI	6

$\tau \approx 49 \text{ ns}$ EM^3

\tilde{X}	2A_1	C_{2v}	
Vib.	No.	Approximate type of mode	cm^{-1}
a_1	1	Sym. stretch	1151.4vs
	2	Bend	525(30)
			523.7m
b_2	3	Asym. stretch	1393.5vs
			1389.9
			1384.8

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- M. E. Jacox and W. E. Thompson, *J. Chem. Phys.* **102**, 4747 (1995).
- D. B. Atkinson, K. K. Irikura, and J. W. Hudgens, *J. Phys. Chem. A* **101**, 2045 (1997).

BCl₂

Emission which is observed between 200 and 260 nm on excitation of BCl₃ by radiation of wavelength shorter than 91 nm has been attributed⁵ to BCl₂.

\tilde{B}, \tilde{C}

Emission which is observed between 200 and 500 nm on excitation of BCl₃ by radiation of wavelength between 100 and 124 nm (9.99–12.27 eV) has been attributed^{8,9} to the $\tilde{B}-\tilde{X}$ and $\tilde{C}-\tilde{X}$ transitions of BCl₂. Other studies are consistent with contributions from two different emission systems, as summarized below.

Emission which is observed between 240 and 380 nm on excitation of BCl₃ by radiation of wavelength shorter than 118 nm^{4,5} and on electron impact⁶ has been attributed to BCl₂, as has been a similar chemiluminescence emission observed^{1,2} on reaction of H atoms with BCl₃. The electron impact study⁶ yielded a radiative lifetime of 1.65(20) μs and suggested that the lower state is the ground state of BCl₂.

Emission which is observed between 280 and 380 nm on excitation of BCl₃ by radiation of wavelength shorter than 124 nm^{4,5} and on electron impact⁶ has also been attributed to BCl₂.

\tilde{A}

Emission which is observed between 380 and 650 nm on excitation of BCl₃ by radiation of wavelength shorter than 138 nm^{4,5,8} and on electron impact⁶ has been attributed to the $\tilde{A}-\tilde{X}$ transition of BCl₂. The electron impact study indicated that the lower state is the ground state of BCl₂.

\tilde{X}		C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a_1	1	Sym. stretch	731w	Ar	IR	3	
b_2	3	Asym. stretch	977.5	Ne	IR	10	
			976.4				
			965.6vs	Ar	IR	3,7	

References

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BBR₂

Emission between 240 and 390 nm, with a lifetime ranging from 18(2) to 26(2) ns, which occurs when BBr₃ is excited by radiation of wavelength between 116 and 140 nm, has been attributed⁴ to two or more band systems of BBr₂.

In an argon matrix, an absorption maximum at 15900 (630 nm) has been attributed² to BBr₂.

\tilde{X}		C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a_1	1	Sym. stretch	551.0w	Ar	IR	2	
b_2	3	Asym. stretch	833.4s	Ar	IR	1-3	

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AlF₂

\tilde{X}		C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a_1	1	Sym. stretch	755.3	Ar	IR	1	
b_2	3	Asym. stretch	887.5	Ar	IR	1	

Reference

- P. Hassanzadeh, A. Citra, L. Andrews, and M. Neurock, *J. Phys. Chem.* **100**, 7317 (1996).

AlCl₂

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	461.0	Ar	IR	2,3
b_2	3	Asym. stretch	563.6	Ar	IR	1-3

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³P. Hassanzadeh, A. Citra, L. Andrews, and M. Neurock, *J. Phys. Chem.* **100**, 7317 (1996).

AlBr₂

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	348.3T	Ar	IR	1
b_2	3	Asym. stretch	458.4	Ar	IR	1

Reference

- ¹P. Hassanzadeh, A. Citra, L. Andrews, and M. Neurock, *J. Phys. Chem.* **100**, 7317 (1996).

AlI₂

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_2	3	Asym. stretch	385.9	Ar	IR	1

Reference

- ¹P. Hassanzadeh, A. Citra, L. Andrews, and M. Neurock, *J. Phys. Chem.* **100**, 7317 (1996).

GaF₂

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	642.0	Ar	IR	1
			638.9			
	2	Bend	158T	Ar	IR	1
b_2	3	Asym. stretch	746.6	Ar	IR	1
			744.5			

Reference

- ¹S. B. Osin, E. D. Samsonova, and V. F. Shevel'kov, *Zh. Fiz. Khim.* **68**, 2009 (1994); *Russ. J. Phys. Chem.* **68**, 1823 (1994).

CO₂⁻

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1253.8w	Ne	IR	2
	2	Bend	714.1w	Ne	IR	2
b_2	3	Asym. stretch	1658.3vs	Ne	IR	1,2

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FCO

\tilde{C}^2A''	C_s
$T_0=28500(25)$	gas AB ^{2,7,8}
≤ 35587	Ar AB ³
≤ 35211	CO AB ^{1,3}

In the gas phase,² bands are diffuse, and the onset of predissociation is estimated⁶ to lie at a wavelength longer than 294 nm. In an argon matrix,³ the threshold for photodissociation into F + CO was observed near 280 nm.

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	Bend	1458(6) ^a	gas	AB	8
			651(5) ^a	gas	AB	2,7,8
			650T	Ar	AB	3
			650T	CO	AB	1,3

\tilde{B}^2A'	C_s
$T_0 \leq 27000$	gas CL ⁵ AB ^{7,8}
≥ 24000	gas AB ⁸
≤ 29586	Ar AB ³
≤ 29516	CO AB ^{1,3}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	Bend	1180(35) ^a	gas	AB	8
			770(30) ^a	gas	AB	8
			700T	Ar	AB	3
			700T	CO	AB	1,3

\tilde{X}^2A'	C_s					
Vib.	No.					
a'	1	CO stretch	1861.64	gas	DL	4
			1857vs	Ar	IR	3
			1855vs	CO	IR	1
	2	Bend	627.5m	Ar	IR	3
			626m	CO	IR	1
	3	CF stretch	1026.13	gas	DL	4
			1023vs	Ar	IR	3
			1018s	CO	IR	1

$A_0=6.38$; $B_0=0.382$; $C_0=0.360$ DL⁴

^a ω .

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CS₂ \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b ₂	3	Asym. stretch	1159.4	Ne	IR	1

Reference

- ¹T. M. Halasinski, J. T. Godbout, J. Allison, and G. E. Leroi, *J. Phys. Chem.* **100**, 14865 (1996).

SiO₂⁻

Threshold for electron detachment from ground-state SiO₂⁻ = 16940(800) gas PE¹

Reference

- ¹L.-S. Wang, H. Wu, S. R. Desai, J. Fan, and S. D. Colson, *J. Phys. Chem.* **100**, 8697 (1996).

GeO₂⁻

Threshold for electron detachment from ground-state GeO₂⁻ = 20170(800) gas PE¹

Reference

- ¹L.-S. Wang, H. Wu, S. R. Desai, J. Fan, and S. D. Colson, *J. Phys. Chem.* **100**, 8697 (1996).

FCS \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CF stretch	1297.4	Ar	IR	1
	2	Bend	457T	Ar	IR	1
	3	C=S stretch	917.9	Ar	IR	1

Reference

- ¹N. Caspary, B. E. Wurfel, A. Thoma, G. Schallmoser, and V. E. Bondybey, *Chem. Phys. Lett.* **212**, 329 (1993).

PO₂

²B₁? C_{2v}
 $T_0=30378(3)$ gas AB¹LF³
Ar AB⁶
²B₁- \tilde{X} 268–600 nm
²B₁- \tilde{X} 292–301 nm

In LF studies,³ there was an apparently continuous background signal, with a maximum between 400 and 500 nm. The similarity of the behavior of this band system to that of the visible bands of NO₂ suggests that the quasicontinuum may be contributed by high vibrational levels of the ground state.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	Sym. stretch	933	gas	AB	1
			942	Ar	AB	6
	2	Bend	396	gas	AB	1

$\tau \approx 500$ ns gas LF³
 $\tau_{\text{cont}} \approx 4.5$ μ s gas LF³

\tilde{X} 2A₁ C_{2v} Structure: AB¹MW, LMR²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	PO s-stretch	1117(20)	gas	MW	2,3
		Bend	387(20)	gas	MW, LFM	2,3
b ₂	3	PO a-stretch	386.4	Ar	IR	7
			1327.53	gas	DL	8,9
			1319.1	Ar	IR	4,5,7

$A_0=3.486$; $B_0=0.287$; $C_0=0.264$ MW, LMR²

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⁸H.-B. Qian, P. B. Davies, I. K. Ahmad, and P. A. Hamilton, *Chem. Phys. Lett.* **235**, 255 (1995).
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SNO \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	NO stretch	1522.8 ^a	Ar	IR	1–3
	3	NS stretch	790.2m	Ar	IR	1–3

^a In Fermi resonance with 2ν₃.

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SeNO

\tilde{X}	C _s
Vib. sym.	No. Approximate type of mode
a'	1 NO stretch

Reference

- ¹L. Andrews, P. Hassanzadeh, D. V. Lanzisera, and G. D. Brabson, J. Phys. Chem. **100**, 16667 (1996).

NS₂

\tilde{X}	C _{2v}
Vib. sym.	No. Approximate type of mode
a ₁	1 NS ₂ s-stretch
b ₂	3 NS ₂ a-stretch

References

- ¹P. Hassanzadeh and L. Andrews, J. Am. Chem. Soc. **114**, 83 (1992).
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NSE₂

\tilde{X}	C _{2v}
Vib. sym.	No. Approximate type of mode
a ₁	1 Sym. stretch
b ₂	3 Asym. stretch

References

- ¹L. Andrews and P. Hassanzadeh, J. Chem. Soc., Chem. Commun. 1523 (1994).
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SeSeN

\tilde{X}	
Vib. sym.	No. Approximate type of mode
	SeN stretch

Reference

- ¹L. Andrews, P. Hassanzadeh, D. V. Lanzisera, and G. D. Brabson, J. Phys. Chem. **100**, 16667 (1996).

SO₂⁺

\tilde{F}^2A_1 C_{2v}
 $T_0=62200(500)$ gas PE²

\tilde{E}^2B_1 C_{2v}
 $T_0=33090(20)$ gas PE^{2,6,7}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	960(10)	gas	PE	7
	2	Bend	444(10)	gas	PE	7

\tilde{D}^2A_1 C_{2v}
 $T_0=32190(50)$ gas PE^{1,2,6,7}PF⁵ $\tilde{D}-\tilde{X}$ 300–317 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	912(1)	gas	PE,PF	1,2,5–7
	2	Bend	411(60)	gas	PF	5

\tilde{C}^2B_2 C_{2v}
 $T_0=28670(50)$ gas PE^{1,2,6,7}PF⁵ $\tilde{C}-\tilde{B}$ 511–437 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	800(4)	gas	PE	6,7
	2	Bend	371(10)	gas	PF,PE	4–7

\tilde{B}^2B_2 C_{2v}
 $T_0=7034(80)$ gas PE^{1,6,7}PF^{4,5} $\tilde{C}-\tilde{B}$ 437–511 nm^a

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	Bend	465(9)	gas	PE,PF	1,4–7
	3	Asym. stretch	1320(8)HT	gas	PE	7

$\tau \approx 25 \mu s$ gas PE⁵

\tilde{A}^2A_2 C_{2v}
 $T_0=5156(65)$ gas PE^{1,6,7}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	981(60)	gas	PE	6,7
	2	Bend	353(7)	gas	PE	6,7
	3	Asym. stretch	202(13)	gas	PE	6,7

\tilde{X}^2A_1 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	Bend	404.2(5)	gas	PE	1,6,7

Barrier to linearity ca. 3200 PE^{6,7}

^aAttributed by Ref. 5 to the $\tilde{C}-\tilde{A}$ transition.

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CF₂ \tilde{B}^a

$T_0=72740$ gas AB¹⁰ $\tilde{B}-\tilde{X}$ 131–138 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	2	Bend	625	gas	AB	10

 \tilde{A}^1B_1 C_{2v} Structure: AB¹⁰

$T_0=37226$ gas EM¹AB^{2,3,5,10}LF^{20,42} $\tilde{A}-\tilde{X}$ 220–380 nm
 37219(2) Ne LF¹⁷
 36878(2) Ar AB^{4,6,16}EM¹⁶LF¹⁷ $\tilde{A}-\tilde{X}$ 210–346 nm
 37054(2) N₂ LF¹⁷

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1011(2)	gas	LF	42
	2	Bend	496	gas	UV	1–3,5,10
			496(2)	Ne	LF	17
			496(2)	Ar	AB,LF	4,6,16,17
<i>b</i> ₂			496(2)	N ₂	LF	17
	3	Asym. stretch	1180(2)	gas	LF	42

$\tau_0=50(5)$ ns gas LP^{20,21,28,30,33}EM^{35,39}EF³⁷

31 ns Ne LF¹⁷

27 ns Ar LF¹⁷

23 ns Kr LF¹⁷

$A_0=4.577$; $B_0=0.334$; $C_0=0.311$ AB¹⁰

 \tilde{a}^3B_1 C_{2v}

$T_0=19828$ gas CL^{18,19,22,24,29}PE³⁴ $\tilde{a}-\tilde{X}$ 430–800 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	Bend	517	gas	CL	18,22, 24,29

$\tau \approx 1$ s gas CL¹⁹

 \tilde{X}^1A_1 C_{2v} Structure: MW⁸AB^{9,10}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	1225.08	gas	DL,IR	23,31,32
			1221.9	Ne	PE	34,36
			1222vs	Ar	IR,LF	12,17,40
	2	Bend	666.25	gas	UV,PE	6,11,17
<i>b</i> ₂			668vw	Ar	DL	41
	3	Asym. stretch	1114.44	gas	IR,LF	6,11,17
			1105.8	Ne	IR	32,38
			1102vs	Ar	IR	12,40

$$A_0=2.947; B_0=0.417; C_0=0.365 \quad \text{MW}^{8,15,26}\text{AB}^{9,10}$$

^aTentative assignment. This band system was associated with the $\tilde{C}-\tilde{X}$ transition in Ref. 10. Subsequent studies^{14,25} have dictated the reassignment to CF₃ of almost all of the bands between 136 and 160 nm which had tentatively been attributed¹⁰ to CF₂.

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SiF₂

\tilde{B}^1B_2 C_{2v}
 $T_0=62280$ gas UV¹¹MPI^{15,16,19} $\tilde{B}-\tilde{X}$ 158–165 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
a_1	1	Sym. stretch	790T	gas	AB	11,19
	2	Bend	320T	gas	MPI AB	11

\tilde{A}^1B_1 C_{2v} Structure: AB⁹
 $T_0=44113.9$ gas EM^{1,2}AB^{5,9}LF^{14,17} $\tilde{A}-\tilde{X}$ 213–276 nm
 $\cong 43964$ Ne AB⁷ $\tilde{A}-\tilde{X}$ 216–225 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
a_1	2	Bend	250.1(3)	gas	AB,LF	5,9,17,18
			253T	Ne	AB	7

$\tau \leq 20$ ns gas LF¹⁷
 $A_0=1.446$; $B_0=0.241$; $C_0=0.206$ AB⁹

\tilde{a}^3B_1 C_{2v} Structure: LF¹⁸
 $T_0=26319.478(6)$ gas EM¹⁰MPI^{16,19}LF¹⁸ $\tilde{a}-\tilde{X}$ 364–420 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
a_1	2	Bend	278.2	gas	EM,MPI,LF	10,16,18

$A_0=1.367$; $B_0=0.253$; $C_0=0.213$ LF¹⁸

\tilde{X}^1A_1 C_{2v} Structure: MW^{3,4}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
a_1	1	Sym. stretch	855.01	gas	IR,LF	6,13,17
			851s	Ne	IR	8
			843s	Ar	IR	7,8
	2	Bend	345	gas	MW,UV, LF,MPI	4,5 17,19
b_2	3	Asym. stretch	343	Ar	IR	7
			870.40	gas	IR	6,13
			864.6s	Ne	IR	8
			855vs	Ar	IR	7,8

$A_0=1.021$; $B_0=0.294$; $C_0=0.228$ MW^{3,4}

SiCl₂

- \tilde{A}^1B_1 C_{2v} Structure: LF¹⁰
 $T_0=30013.5(2)$ gas AB⁴EM^{5-7,15}LF^{8,10,13} $\tilde{A}-\tilde{X}$ 295–430 nm
In an argon matrix, unstructured absorption attributable to SiCl₂ has been observed¹ between 310 and 320 nm, with a maximum at approximately 315 nm.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
a_1	1	Sym. stretch	428.9	gas	LF	8,13
	2	Bend	149.8	gas	UV,LF	3,7,8,13

$\tau_{070}=77(3)$ ns gas LF^{8,15}
 $A_{060}=0.909$; $B_{060}=0.076$; $C_{060}=0.069$ LF¹⁰

\tilde{a}^3B_1 C_{2v}
 $T_0=18943T$ gas EM^{11,12,15} $\tilde{a}-\tilde{X}$ 500–650 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.		
a_1	2	Bend	159(2)	gas	EM	11,12

$\tau=11(2)$ ms gas EM¹²

\tilde{X}^1A_1		C_{2v}	Structure: ED ⁴ MW ^{9,14}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	521.6	gas	LF	13
			518.7	Ne	IR	2
			512.5s	Ar	IR	1,2
	2	Bend	200.6	gas	EM,LF	7,8,11-13
b_2	3	Asym. stretch	202.2	Ar	IR	2
			509.4	Ne	IR	2
			502vs	Ar	IR	1,2

$A_0=0.493$; $B_0=0.094$; $C_0=0.079$ MW⁹LF¹⁰

\tilde{X}^1A_1		C_{2v}	Structure: IR ²			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	663	gas	IR	2
			655	Ne	IR	2
			648	Ar	IR,Ra	2,7
	2	Bend	263(2)	gas	AB,EM	1,6,8
b_2	3	Asym. stretch	692	gas	IR	2
			685	Ne	IR	2
			676	Ar	IR	2

$A_0=0.513$; $B_0=0.262$; $C_0=0.173$ MW^{3,4}

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GeF₂

The high-temperature vapor of GeF₂ shows unstructured absorption between 136 and 156 nm, with a maximum near 146.3 nm.⁵

\tilde{A}^1B_1		C_{2v}	$\tilde{A}-\tilde{X}$ 222-243 nm			
$T_0=43860.9$	gas	AB ¹ LF ¹⁰				

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	623.8(6)	gas	LF	10
	2	Bend	159.6	gas	AB,LF	1,10

\tilde{a}^3B_1		C_{2v}	$\tilde{a}-\tilde{X}$ 305-370 nm			
$T_0=30582.1$	gas	EM ^{6,8,9} LF ¹⁰				

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	673.1(5)	gas	LF	10
	2	Bend	192.2	gas	EM,LF	6,8,10

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GeCl₂

\tilde{A}^1B_1	C_{2v}	$T_0=30622(2)$ gas AB ² LF ¹¹	$\tilde{A}-\tilde{X}$ 300-330 nm
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Structured absorption² is superposed on a continuum with maximum near 32280, presumably due to predissociation of GeCl₂ into GeCl + Cl. In the LF excitation spectrum,¹¹ there is a marked increase in the line density beyond approximately 31630.

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	354	gas	LF	11
	2	Bend	104	gas	AB,LF	2,11

$\tau=89.7(6.8)$ ns gas EM¹⁰

\tilde{a}^3B_1	C_{2v}	$T_0=22315(2)$ gas CL ¹ LF ¹¹	$\tilde{a}-\tilde{X}$ 400-490 nm
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Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	393	gas	LF	11
	2	Bend	118	gas	CL,LF	1,11

$\tau=17.4(6)$ μ s gas EM¹⁰

\tilde{X}^1A_1		C_{2v}	Structure: ED ⁸ MW ¹²			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	399	gas	CL,Ra	1,4
			398.6	Ar	IR	3,5,7,9
			390	N ₂	Ra	6
b_2	2	Bend	160(4)	gas	CL,AB	1,2
				Ra		4
			163	N ₂	Ra	6
b_2	3	Asym. stretch	373.5	Ar	IR	3,5,7,9
			362	N ₂	Ra	6

$A_0=0.241$; $B_0=0.087$; $C_0=0.064$ MW¹²

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NF₂⁺

$\tilde{A}^1B_1?$ C_{2v}
 $T_0 \approx 38400^a$ gas PE^{1,2}

$\tilde{\alpha}^3B_1$ C_{2v}
 $T_0 = 19610(320)$ gas PE^{1,2}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			520(20)	gas	PE	1,2

\tilde{X}^1A_1 C_{2v}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1250(20)	gas	PE	1,2
b_2	3	Asym. stretch	1233.5T	Ne	IR	3

^aFrom vertical ionization potential.

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PO₂⁻

Threshold for electron detachment from ground-state PO₂⁻=27600(80) gas PE²

\tilde{X}		C_{2v}	Structure: PE ²			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	Bend	520T	gas	PE	2
b_2	3	PO a-stretch	1198.8	Ar	IR	1

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SOO

\tilde{X}		C_s	Structure: AB ¹¹			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'			1006.1	Ar	IR	1
			739.9	Ar	IR	1

Reference

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SSO

An absorption band system between 190 and 230 nm has been attributed⁷ to SSO. However, an alternate assignment to the $\tilde{C}-\tilde{X}$ band system of SO₂ has been proposed.¹⁵

\tilde{C}^1A' C_s Structure: AB¹¹
 $T_0 = 29687.72$ gas AB^{1,7,11}LF^{12,13,16,17} $\tilde{C}-\tilde{X}$ 250–395 nm
29285(20) Xe AB⁴ $\tilde{C}-\tilde{X}$ 280–342 nm

Predissociation limit between 31172 and 31307 AB¹¹

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SO stretch	1032	gas	AB,LF	11,13,16
	2	Bend	253	gas	AB,LF	11,13,16
	3	SS stretch	410.6	gas	AB,LF	7,11,13, 16,17
			415(20)	Xe	AB	4

$\tau=66(4)$ ns gas LF¹⁷
 $A_0=1.016$; $B_0=0.149$; $C_0=0.130$ AB¹¹LF¹⁷

$\tilde{\alpha}^3A'$ C_s
 $T_0 = 13943$ gas AB^{10,15}LF¹³ $\tilde{\alpha}-\tilde{X}$ 430–670 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SO stretch	1089	gas	AB	10,15
	2	Bend	332	gas	AB	10,15
	3	SS stretch	505	gas	AB	10,15

\tilde{X}^1A' C_s Structure: MW^{2,6}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	SO stretch	1166.45	gas	IR,DL	1,3,14
			1156.2	Ar	IR,Ra	8,9
	2	Bend	380	gas	LF	16
			382	Ar	IR,Ra	8,9
	3	SS stretch	679.1	gas	IR,LF	1,3,13
			672.2	Ar	IR,Ra	8,9

 $A_0=1.398; B_0=0.169; C_0=0.150$ MW^{2,5,6}DL¹⁴

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S₃
 $T_0=23465(15)$ gas AB^{1,2,5,7}
 23210 Ar AB^{4,6,8}
 Kr AB^{1,2}

350–510 nm
355–435 nm
310–420 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	420	gas	AB	2
			450T	Ar	AB	4,8
	2	Bend	420	Kr	AB	2
			340T	Ar	AB	8

 \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	590	gas	AB	2
			581	gas	Ra	9
	2	Bend	583	Ar	Ra,IR	3,6
			281	gas	Ra	9
<i>b</i> ₂	3	Asym. stretch	680.0	Ar	IR	6,8
			676.2			
			674.5			

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SeO₂
 1B_2 C_{2v} Structure: AB¹⁰
 $T_0=31957.4$ gas AB^{1,9,10}EM⁴
 31065(20) Xe AB⁶

$^1B_2-\tilde{X}$ 225–345 nm
 $^1B_2-\tilde{X}$ 242–322 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	648.8	gas	AB	1,9
	2	Bend	620T	Xe	AB	6
			258	gas	AB	9

 3B_2 C_{2v} Structure: AB¹¹
 $T_0=23840$ gas AB^{2,11}EM⁴
 Kr EM⁶
 Xe EM⁶

$^3B_2-\tilde{X}$ 370–500 nm
 $^3B_2-\tilde{X}$ 475–600 nm
 $^3B_2-\tilde{X}$ 475–615 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	780	gas	AB	11
	2	Bend	199	gas	AB,EM	2,4,11
<i>b</i> ₂	3	Asym. stretch	863	gas	AB	11

 $\tau<200(100)$ μ s Xe EM⁶ \tilde{X}^1A_1 C_{2v} Structure: MW⁷

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	922.6	gas	AB,PE	9,11,12
			923.4	Ne	IR	5
			922.0w	Ar	IR	5,8,13
	2	Bend	372	gas	AB	9,11
			366.0s	Ar	IR	8,13
<i>b</i> ₂	3	Asym. stretch	365T	Kr,Xe	FM	6
			963	gas	IR,AB	3,11
			971.2	Ne	IR	5
			970.2			
			965.3s	Ar	IR	5,8,13

 $A_0=0.962; B_0=0.289; C_0=0.222$ MW⁷

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SeOO

\tilde{X}		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OO stretch	1061.0T	Ar	IR	1

Reference

- ¹G. D. Brabson, L. Andrews, and C. J. Marsden, *J. Phys. Chem.* **100**, 16487 (1996).

SeSeO

\tilde{X}		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SeO stretch	883.2	Ar	IR	1

Reference

- ¹G. D. Brabson, L. Andrews, and C. J. Marsden, *J. Phys. Chem.* **100**, 16487 (1996).

NF₂

\tilde{A} 2A_1		C_{2v}				
gas	AB ^{1,4,5}	$\tilde{A}-\tilde{X}$	237–278 nm			
Ar	AB ⁹	$\tilde{A}-\tilde{X}$	247–265 nm			
In an argon matrix, ^{6,9} evidence has been obtained for predissociation into NF+F.						
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	Bend	390 ^a	gas	AB	4,5
			408 ^a	Ar	AB	9

\tilde{X} 2B_1		C_{2v}		Structure: IR ² MW ⁸		
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1074.99	gas	IR,DL	2,10,
			1071.6	Ne	IR	11,12
			1069m	Ar	IR	15
			1070	N ₂	IR	7,9
		Bend	573w	N ₂	IR	2,3
	3	Asym. stretch	942.48	gas	IR,DL	2,10,
			936.0	Ne	IR	13,14
			932vs	Ar	IR	15
			931	N ₂	IR	7,9
						2,3

$$A_0=2.351; B_0=0.396; C_0=0.338 \quad \text{MW}^8 \text{DL}^{12}$$

^aAverage value.

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NFCI

\tilde{A}		C_s				
gas	EM ¹	Ar	AB ³ EM ⁴	$\tilde{A}-\tilde{X}$	380–500 nm	$\tilde{A}-\tilde{X}$
In the gas phase, an emission maximum is observed ¹ near 23500 (425 nm). In an argon matrix, an absorption maximum is observed ³ at 27800 (360 nm), and an emission maximum appears ⁴ near 21200 (472 nm). Ref. 4 discusses reasons for the disparity between the gas-phase and the argon-matrix observations.						
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	Bend	340(20)	Ar	AB	3
$\tau < 20 \text{ ns}$ gas EM ¹ = 610(40) ns Ar EM ⁴						

\tilde{X}	C_s					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	NF stretch	917 ^a	Ar	IR	2
	2	Bend	390(40)	gas	EM	1
			390T	Ar	EM	4
	3	NCl stretch	720 ^a	Ar	IR	2

^aSimilar values, not explicitly given, were obtained in krypton and xenon matrix experiments.

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PF_2

\tilde{M}	C_{2v}					
$T_0=68505(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	972(7) ^a	gas	MPI	4
	2	Bend	345(3) ^a	gas	MPI	4

\tilde{L}	C_{2v}					
$T_0=67922(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	956(2)	gas	MPI	4
	2	Bend	358(3)	gas	MPI	4

\tilde{K}	C_{2v}					
$T_0=66763(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	956(2)	gas	MPI	4
	2	Bend	364(1)	gas	MPI	4

\tilde{J}	C_{2v}					
$T_0=66118(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	981(4) ^a	gas	MPI	4
	2	Bend	366(1) ^a	gas	MPI	4

\tilde{I}	C_{2v}					
$T_0=65958(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	980(4) ^a	gas	MPI	4
	2	Bend	365(3) ^a	gas	MPI	4

\tilde{H}	C_{2v}					
$T_0=60962(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1004(2)	gas	MPI	4

\tilde{G}	C_{2v}					
$T_0=58184(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	998(2)	gas	MPI	4

\tilde{F} 2A_1	C_{2v}					
$T_0=55126(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1008(2)	gas	MPI	4

\tilde{E} 2B_1	C_{2v}					
$T_0=51932(5)$	gas MPI ⁴					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1016(2) ^a	gas	MPI	4
	2	Bend	408(2) ^a	gas	MPI	4

\tilde{a} 4A_2	C_{2v}
$T_0=23998(6)$	gas EM ³

\tilde{V}	C_{2v}
$T_0=505(2)$	gas EM
$T_0=219$	gas EM

$\tau \geq 1.9 \text{ ms}$ gas EM³

\tilde{X} 2B_1	C_{2v}	Structure: MW ²
Vib.	No.	Approximate type of mode
a_1	1	Sym. stretch
		841(4)
		852.1ms
	2	Bend
		366
b_2	3	Asym. stretch
		848(24)
		831.4s
		Ar
		IR

$A_0=0.933$; $B_0=0.310$; $C_0=0.232$ MW²

^a ω .

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PCl₂

$\tilde{E}(4p)$ C_{2v}
 $T_0=51320(10)$ gas MPI³

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	600(15)	gas	MPI	3
	2	Bend	240(15)	gas	MPI	3

$\tilde{D}(2A_1(4s))$ C_{2v}
 $T_0=42760(15)$ gas MPI³

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	620(20)	gas	MPI	3
	2	Bend	230(20)	gas	MPI	3

$\tilde{A}(2A_1)$ C_{2v}

In the gas phase, unstructured emission at wavelengths longer than 370 nm, with a maximum at approximately 460 nm, has been attributed² to PCl₂.
 $\tau=29(6) \mu s$ gas EM²

$\tilde{X}(2B_1)$ C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	525(10)	gas	MPI	3
			525	Ar	IR	1
b_2	3	Asym. stretch	452	Ar	IR	1

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PBr₂

$\tilde{A}(2A_1)$ C_{2v}

In the gas phase, unstructured emission at wavelengths longer than 510 nm, with a maximum at 527 nm, has been attributed² to PBr₂.
 $\tau=21(9) \mu s$ gas EM²

$\tilde{X}(2B_1)$ C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	369	Ar	IR	1
	2	Bend	140T	gas	EM	3
b_2	3	Asym. stretch	410	Ar	IR	1

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AsF₂

\tilde{H} C_{2v}
 $T_0=57480(15)$ gas MPI³

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	810(10)	gas	MPI	3
	2	Bend	310(20)	gas	MPI	3

\tilde{F} C_{2v}
 $T_0=54355(15)$ gas MPI³

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	813(6)	gas	MPI	3
	2	Bend	308(5)	gas	MPI	3

When gas-phase AsF₃ is subjected to 124 nm radiation, an emission band system, tentatively assigned to AsF₂, is observed between 330 and 610 nm, with a maximum near 450 nm.¹ The radiative lifetime of the species which contributes this band system is 25.5(1.8) μs .²

$\tilde{X}(2B_1)$ C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	705(10)	gas	MPI	3
	2	Bend	285(10)	gas	MPI	3

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O₃⁻

$\tilde{C}(2A_1)$ C_{2v}
 $T_0=21420(40)$ gas PF¹⁰

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	760(20)	gas	PF	10
	2	Bend	190(20)	gas	PF	10

Threshold for electron detachment from ground-state O₃⁻ = 16970(20) gas PE⁹PD¹¹

$\tilde{A}(2A_2)$ C_{2v}
 $T_0=16508(16)$ gas PF^{8,10}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	815(10)	gas	PF	8,10
	2	Bend	275(10)	gas	PF	8,10

\tilde{X}^2B_1		C_{2v}	Structure: PD ¹¹ PE ¹³			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	975(50)	gas	PD,PF PE	8–10,13
			1016 Cs	Ar	Ra	3,5
			1011 Na	Ar	Ra	3,5
	2	Bend	550(50)	gas	PD,PF PE	9,10,13
			600w Cs	Ar	IR	4
			880(50)	gas	PE	13
b_2	3	Asym. stretch	796.3	Ne	IR	12,14
			804.3	Ar	IR	6,7
			789s Cs	Ar	IR	2,4
			802s Cs			
			802s Na	Ar	IR	1,2,4

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CIOO

An unstructured absorption between 225 and 270 nm, with a maximum near 246 nm in the gas phase^{3–6} and near 250 nm in a neon⁸ or an argon⁷ matrix, has been assigned to CIOO, which photodecomposes in that spectral region.

\tilde{X}		C_s	Structure: IR ⁸				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a'	1	OO stretch	1443 ^a	gas	IR	4	
			1438.6	Ne	IR	8	
			1442.8vs ^b	Ar	IR	2,7	
			1438	N ₂	IR	1	
			1428				
	2		1436	O ₂	IR	7	
			413.7	Ne	IR	8	
			408.3vw ^b	Ar	IR	2,7	
			201.4	Ne	IR	8	
			192.4w	Ar	IR	7	
	3	ClO stretch	203	O ₂	IR	7	

^aAbsorption maximum; spectral slit width 13 cm^{-1} .

^bPeaks at 1415, 435, and 227 cm^{-1} may be contributed by ClOO trapped in a metastable site in the argon lattice.^{7,8}

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BrOO

In an argon matrix, a prominent absorption between 240 and 300 nm, with maximum at 38200 (262 nm), has been assigned³ to BrOO. Irradiation at 280 nm leads to photoisomerization to OBrO.⁴

X

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		OO stretch	1485.1	Ar	IR	1–4

References

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IOO

In an argon matrix, an absorption maximum at 39370 (254 nm) has been attributed¹ to IOO. Irradiation of the sample in the spectral region of this absorption results in photoisomerization to OIO.

Reference

- G. Maier and A. Bothur, Chem. Ber. **130**, 179 (1997).

SeO₂⁻

Threshold for electron detachment from ground-state SeO₂⁻ = 14710(400) gas PE¹

\tilde{X}		C_{2v}	Structure: IR ⁸			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	810(80)	gas	PE	1

Reference

- J. T. Snodgrass, J. V. Coe, K. M. McHugh, C. B. Freidhoff, and K. H. Bowen, J. Phys. Chem. **93**, 1249 (1989).

SSF

\tilde{A}^2A'		C_s	Structure: LF ^{1,2}				
$T_0=14922$	gas	LF ^{1,2}	$\tilde{A}-\tilde{X}$ 485–700 nm				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a'	1	SF stretch	768	gas	LF	1	
	2	Bend	217	gas	LF	1	
	3	SS stretch	495	gas	LF	1	

 $A_0=0.662; B_0=0.157; C_0=0.126 \text{ LF}^{1,2}$

\tilde{X}^2A''		C_s	Structure: LF ¹				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a'	1	SF stretch	705	gas	LF	1	
	2	Bend	293	gas	LF	1	
	3	SS stretch	684	gas	LF	1	

 $A_0=0.886; B_0=0.162; C_0=0.137 \text{ LF}^1 \text{MW}^3$

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SSCI

$T_0=21650\text{T}$ gas ^a		AB ^{1,2,5}	378–481 nm
≤ 21925 Ar		AB ³	389–456 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	Stretch	480	gas	AB	5
			491(20)	Ar	AB	3
	3	Stretch	407	gas	AB	5

\tilde{A}		C_s					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a'	1	SS stretch	630	gas	LF	7,8	
	2	Bend	249	gas	LF	7,8	
	3	SCI stretch	554	gas	LF	7,8	

\tilde{X}^2A''		C_s	Structure: MW ⁶				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	SS stretch	662(3)	gas	AB,Ra,LF	5,7,8
			665	Ar	IR	4
	2	Bend	196	gas	Ra,LF	7,8
	3	SCI stretch	450	gas	Ra,LF	7,8
			404	Ar	IR	4

 $A_0=0.611; B_0=0.094; C_0=0.082 \text{ MW}^6$

^aDiffuse band system. Ref. 3 also reported two very weak bands approximately 600 cm^{-1} below the principal progression which they attributed to ground-state vibrational excitation.

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OCIO

\tilde{E}		$AB^{12,19,45}$	$\tilde{E}-\tilde{X}$ 148–157 nm
Vib.	No.	Approximate type of mode	cm^{-1}

1	Sym. stretch	1000T	gas	AB	12
2	Bend	508	gas	AB	12,19

\tilde{D}		$AB^{12,19,42,45}$	$\tilde{D}-\tilde{X}$ 155–163 nm
Vib.	No.	Approximate type of mode	cm^{-1}

1	Sym. stretch	1051(20)	gas	AB	12,19,42
2	Bend	521(20)	gas	AB	12,19,42

\tilde{C}^2A_1		C_{2v}	$\tilde{C}-\tilde{X}$ 176–183 nm
$T_0=54689(20)$	gas	$AB^{12,19,42,45}$	

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	1020(20)	gas	AB	12,19,42

 $^2A_1, ^2B_2$

These two as yet undetected states are expected to lie close to the \tilde{A} state and to interact strongly with it. One or both of them may contribute to the much-studied predissociation of the \tilde{A} state.^{21,24,29,35,40,46,47}

\tilde{A}^2A_2		C_{2v}^a	Structure: AB ²⁵
$T_0=21017.2$		gas	$AB^{1,2,5,14,25,31,32,34,35,45}LF^{15,16}MPI^{33,36}PF^{41,43,47}$
Vib.	No.	Approximate type of mode	cm^{-1}

a_1	1	Sym. stretch	711.5	gas	AB	5,14,34
	2	Bend	292.5	gas	AB	5,14,34
b_2	3	Asym. stretch	441.2	gas	AB	25,34

 $\tau_1^b=56(20) \text{ ps}$ gas $AB^{21,24}LF^{29}$ $\tau_2^c=336(27) \text{ ps}$ gas LF^{29} $A_0=1.057; B_0=0.311; C_0=0.240 \text{ AB}^{25}$

\tilde{X}^2B_1		C_{2v}	Structure: MW ^{6,30} IR ²⁶			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	945.59s	gas	IR,AB	3–5,14–16
					LF	20,23,26
					LMR	
					DL,LS	28,37,39
			944.8m	Ne	IR	44
			947.6m	Ar	IR,Ra	13,17,18
						44
			942	Kr	Ra	18
			940(2)	Xe	Ra	18
			950(2)	N ₂	Ra	18
a_2	2	Bend	447.70s	gas	IR,AB	4,5,14–16
					LF	22,25,38
						39
			448.7m	Ne	IR	44
			451s	Ar	IR,LF	13,17,18
			447s			44
			447	Kr	LF	18
b_2	3	Asym. stretch	1110.11vs	gas	IR	3,14,26
						38,39
			1107.6vs	Ne	IR	44
			1106.5vs	Ar	IR	13,17,44
			1100.8vs			

$A_0=1.737$; $B_0=0.332$; $C_0=0.278$ MW^{6–10,27}IR^{37,38}

^aDouble minimum in potential for antisymmetric stretch,^{11,25} with barrier height of 1153.³⁴

^bFor $F_1(J=N+\frac{1}{2})$ spin states.

^cFor $F_2(J=N-\frac{1}{2})$ spin states.

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OBrO

\tilde{A}^2A_2 C_{2v}
 $T_0 \leq 16509$ gas AB² $\tilde{A}-\tilde{X}$ 420–605 nm
 16785(20) Ar AB^{3,5} $\tilde{A}-\tilde{X}$ 400–650 nm

Irradiation of the argon-matrix deposit in this spectral region leads to photoisomerization to BrOO.⁵

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	615T	gas	AB	2
			631T	Ar	AB	5
a_2	2	Bend	200T	gas	AB	2
			221T	Ar	AB	5
b_2	3	Asym. stretch	365HT	gas	AB	2

\tilde{X}^2B_1 C_{2v} Structure: MW⁴

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	795.7	Ar	IR	3,5
			317.0	Ar	IR	5
			848.6	gas	IR	4
b_2	3	Asym. stretch	845.2	Ar	IR	1,3,5

$A_0=0.935$; $B_0=0.275$; $C_0=0.212$ MW⁴

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OIO

\tilde{A}		C_{2v}			
$T_0=14553(7)$ T		gas AB ²		$\tilde{A}-\tilde{X}$ 480–662 nm	
In an argon matrix, structured absorption between 467 and 634 nm, with maximum at 18180 (550 nm), has been assigned ³ to OIO.					

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	631(3)	gas	AB	2
	2	Bend	177(2)	gas	AB	2

\tilde{X}		C_{2v}				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	765(25)	gas	PE	1
			768.0w	Ar	IR	3
			769.5	O ₂	IR	3
b_2	2	Bend	192(35)	gas	PE	1
	3	Asym. stretch	800.3vs	Ar	IR	3
			801.3	O ₂	IR	3

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NF₂⁻

\tilde{X}		C_{2v}				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	994.7m	Ne	IR	1
	2	Bend	431H	Ne	IR	1
b_2	3	Asym. stretch	862.4s	Ne	IR	1

Reference

¹M. E. Jacox and W. E. Thompson, J. Chem. Phys. **102**, 6 (1995).

BrOCl

\tilde{X}		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	ClO stretch	675.9	Ar	IR	1
	3	BrO stretch	559.5	Ar	IR	1

Reference

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BrOBr

In the gas phase, a prominent absorption maximum near 50000 (200 nm) and a less prominent absorption maximum near 31800 (314 nm), with a broad shoulder which has its onset near 22700 (440 nm) and on which structure with separations of ~500 is partially resolved, have been assigned⁴ to BrOBr.

In a nitrogen matrix, a strong absorption maximum at approximately 47000, a weaker shoulder near 31200, and a much weaker shoulder near 22100 have been attributed³ to electronic transitions of BrOBr.

\tilde{X} C_{2v} Structure: EXAFS³MW^{5,6}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	526.1s	Ar	IR	1,2
			528	N ₂	IR	3
b_2	3	Asym. stretch	623.4w	Ar	IR	2
			626	N ₂	IR	3

$A_0=1.108$; $B_0=0.046$; $C_0=0.044$ MW^{5,6}

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SF₂

\tilde{E} 4p Rydberg state C_{2v} $T_0=62015(30)$ gas MPI¹⁰

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	931(59)	gas	MPI	10
	2	Bend	383(42)	gas	MPI	10

\tilde{C} gas MPI¹⁰ $\tilde{C}-\tilde{X}$ 165–175 nm

\tilde{B} 1B₁(4s) C_{2v} $T_0=54433(30)$ gas MPI¹⁰

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Sym. stretch	991(12)	gas	MPI	10
	2	Bend	361(24)	gas	MPI	10

\tilde{A}

$T_0 = 18100(1000)$ gas CL^{7,8,11} $\tilde{A}-\tilde{X}$ 550–850 nm
Chemiluminescence in the reaction of F₂ with CS₂, originally assigned^{7,8} to FCS, has been reassigned⁹ to SF₂.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	2	Bend	243(6)	gas	CL	11
\tilde{X} C _{2v} Structure: MW ^{1,2,4}						
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	838.53	gas	IR,CL	5,7–9,11
			834	Ne	IR	3
			832vs	Ar	IR	3,12
			825	N ₂	IR	3
2		Bend	355(2)	gas	MW,CL	2,7–9,11
			358	Ne	IR	3
			358m	Ar	IR	3
			358	N ₂	IR	3
<i>b</i> ₂	3	Asym. stretch	813.04	gas	IR,CL	5,11
			807.5	Ne	IR	3
			804vs	Ar	IR	3,12
			795	N ₂	IR	6

$A_0=0.898$; $B_0=0.307$; $C_0=0.228$ MW^{1,2,13}

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FSCI

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	SF stretch	781vs	Ne	IR	1
			778vs	Ar	IR	1
2		Bend	277vw	Ne	IR	1
			274vw	Ar	IR	1
3		SCl stretch	552s ^a	Ne	IR	1
			543s ^a	Ar	IR	1

$A_0=0.738$; $B_0=0.157$; $C_0=0.130$ MW²

^aIn Fermi resonance with $2\nu_2$.

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FSBr

 \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	SF stretch	765	Ar	IR	1
	3	SBr stretch	434	Ar	IR	1

Reference

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SeF₂ \tilde{X} C_{2v} Structure: IR¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	699	Ar	IR	1
<i>b</i> ₂	3	Asym. stretch	672	Ar	IR	1

Reference

- A. Haas and H. Willner, Z. Anorg. Allg. Chem. **454**, 17 (1979).

CICIO

In an argon matrix, an absorption maximum at 38500 (260 nm) has been attributed^{5,6} to CICIO.

 \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	CIO stretch	961.6s	Ar	IR,Ra	3–6
			963	N ₂	IR	1,2
2		Bend	240.2wm	Ar	IR,Ra	3,5,6
3		CICl stretch	374.2ms	Ar	IR,Ra	3,5,6
			377	N ₂	IR	1,2

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BrClO

\tilde{X}	C_s					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	ClO stretch	940.5	Ar	IR	1,2

References

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ClBrO

In an argon matrix, irradiation at 633 nm results in isomerization to BrClO, and irradiation at 870 nm to BrOCl.²

\tilde{X}	C_s					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	BrO stretch	819.6	Ar	IR	1,2

References

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BrBrO

In an argon matrix, irradiation at 870 nm results in photoisomerization to BrOBr.³

\tilde{X}	C_s					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	BrO stretch	805.4	Ar	IR	1-3
	3	BrBr stretch	236	Ar	IR	1

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KrF₂

Continuous absorption in the gas phase between 210 and 320 nm, most intense at 210 nm.⁵

\tilde{X}	$D_{\infty h}$		Structure: IR ^{3,7}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	449	gas	Ra	2
			452	Kr	Ra	4
Π_u	2	Bend	235.05	gas	IR	2,6
			236m	Ar	IR	1
Σ_u^+	3	Asym. stretch	592.52	gas	IR	2,3,7
			580s	Ar	IR	1

$$B_0 = 0.125 \text{ IR}^{3,6,7}$$

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XeF₂

$$6p \ ^1\Sigma_g \quad D_{\infty h} \quad T_0 = 87400 \text{ gas AB}^{3,4,11} \quad 6p \ ^1\Sigma_g - \tilde{X} \ 114 \text{ nm}$$

A higher member of this Rydberg series has also been reported.¹¹

$$5d \ ^1\Pi_{u,1/2} \quad D_{\infty h} \quad T_0 = 86000 \text{ gas AB}^{3,4,11} \quad 5d \ ^1\Pi_{u,1/2} - \tilde{X} \ 116 \text{ nm}$$

Higher members of this Rydberg series have also been reported.¹¹

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	484(24)	gas	AB	11
Π_u	2	Bend	200T	gas	AB	11

$$5d \ ^1\Pi_{u,3/2} \quad D_{\infty h} \quad T_0 = 80800 \text{ gas AB}^{3,4,11} \quad 5d \ ^1\Pi_{u,3/2} - \tilde{X} \ 124 \text{ nm}$$

Higher members of this Rydberg series have also been reported.¹¹

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	524(8)	gas	AB	11
Π_u	2	Bend	200T	gas	AB	11

$$6s \ ^1\Pi_{u,1/2} \quad D_{\infty h} \quad T_0 = 73870 \text{ gas AB}^{3,4,8,11} \quad 6s \ ^1\Pi_{u,1/2} - \tilde{X} \ 135 \text{ nm}$$

Higher members of this Rydberg series have also been reported.^{3,4,11}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	500(16)	gas	AB	11

$6s^1\Pi_{g,3/2}$ D_{∞h}
 $T_0=69300$ gas AB^{3,4,8,11}
 $6s^1\Pi_{u,3/2}-\tilde{X}$ 144 nm
Higher members of this Rydberg series have also been reported.^{3,4,11}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1	Sym. stretch	532(8)	gas	AB	11
Π_u	2	Bend	73(8)	gas	AB	11

$\tilde{B}^1\Sigma_u$ D_{∞h}
 $T^a=63300$ gas AB^{3,4,11}
 $\tilde{B}-\tilde{X}$ 158 nm

$\tilde{A}^1\Pi_g$ D_{∞h}
 $T^a=43500$ gas AB^{3,4,6,11}
 $\tilde{A}-\tilde{X}$ 230 nm

$\tilde{X}^1\Sigma_g$		D _{∞h}	Structure: IR ^{7,14,16}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1	Sym. stretch	516.5(5)	gas	Ra	9,13
			512	Ar	Ra	10
			512	Xe	Ra	10
Π_u	2	Bend	213.08s	gas	IR	5,14,15
			215	Ar	IR	12
Σ_u^+	3	Asym. stretch	561.94s	gas	IR	1,5,7,14
			547	Ar	IR	2

$B_0=0.113$ IR^{7,14,15}

^aAbsorption maximum.

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Ar₂F

$4^2\Gamma$ C_{2v}

Unstructured gas-phase emission¹⁻³ between 250 and 340 nm, with maximum near 34250 (292 nm). Predissociated at 248 nm to ArF + Ar.⁵ In a neon matrix, the emission maximum is shifted to 33240 (301 nm),⁷ and in an argon matrix to 28170 (355 nm).⁶

$\tau=185(46)$ ns gas EM⁴
 $180(10)$ ns Ar EM⁶

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ArKrF

Unstructured gas-phase emission^{1,2} between 240 and 370 nm, with maximum near 32790 (305 nm). In solid argon, the emission maximum appears near 29150 (343 nm) and has a radiative lifetime of 60(5) ns.³

References

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Kr₂F

$9^2\Gamma$ C_{2v}
gas AB^{6,8,9,12}

Broad absorption, with maximum near 31750 (315 nm). Predissociated into KrF (D) + Kr.¹²

$6^2\Gamma$ C_{2v}
gas AB¹²

Broad absorption, with maximum near 14080 (710 nm). Predissociated into KrF (C) + Kr.¹²

$4^2\Gamma$ C_{2v}

Unstructured gas-phase emission¹⁻⁴ between 340 and 480 nm, with maximum near 24390 (410 nm). In solid neon, the emission maximum is shifted to 23240 (430 nm).¹¹ In solid argon, the emission maximum appears near 22780 (439 nm), and in solid krypton the excitation maximum appears near 36360 (275 nm) and the emission maximum near 22080 (453 nm).¹⁰
 $\tau=200(28)$ ns gas LF⁵EF⁷
140(10) ns Kr LF¹⁰

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KrXeF

Unstructured gas-phase emission between 340 and 460 nm, with maximum near 400 nm, has been attributed¹⁻³ to KrXeF.

References

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Xe₂F

$4^2\Gamma$ C_{2v}
gas EM¹⁻³

$4^2\Gamma-1,2^2\Gamma$

Unstructured emission maximum at 16300 (614(5) nm), with bandwidth (FWHM) of 125(5) nm.

Ne LF⁵

$4^2\Gamma-1,2^2\Gamma$

Excitation maximum near 28600 (350 nm), and emission maximum at 14000 (714 nm).

Xe EM⁴

$4^2\Gamma-1,2^2\Gamma$

Unstructured emission maximum at 12900 (775 nm), with bandwidth (FWHM) of 2300 cm⁻¹.

$\tau=152(+19,-10)$ ns gas EM²
250 ns Ne LF⁵
190(10) ns Xe EM⁴

References

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8.4. Four-Atomic Trihydrides**TiH₃**

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		TiH stretch	1580.6	Ar	IR	1

TiD₃

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		TiD stretch	1147.2	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, *J. Am. Chem. Soc.* **116**, 8322 (1994).

ZrH₃

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		ZrH stretch	1545.3	Ar	IR	1

ZrD₃

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		ZrD stretch	1110.3	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 15004 (1995).

HfH₃

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		HfH stretch	1646	Ar	IR	1

HfD₃

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		HfD stretch	1180.2	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 15004 (1995).

FeH₃

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		FeH ₃ stretch	1646.1	Ar	IR	1

FeD₃ \tilde{X}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		FeD ₃ stretch	1193.2	Ar	IR	1

Reference¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 12131 (1995).**ThH₃** \tilde{X} C_{3v}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
e	3	ThH ₃ stretch	1435.4 1434.1	Ar	IR	1

ThD₃ \tilde{X} C_{3v}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
e	3	ThD ₃ stretch	1025.9 1024.8	Ar	IR	1

Reference¹P. F. Souter, G. P. Kushto, L. Andrews, and M. Neurock, *J. Phys. Chem. A* **101**, 1287 (1997).**UH₃** \tilde{X} C_{3v}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
e	3	UH ₃ stretch	1346.8	Ar	IR	1

UD₃ \tilde{X} C_{3v}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
e	3	UD ₃ stretch	962.5	Ar	IR	1

Reference¹P. F. Souter, G. P. Kushto, L. Andrews, and M. Neurock, *J. Am. Chem. Soc.* **119**, 1682 (1997).**BH₃** \tilde{X} D_{3h}Structure: IR³

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'' ₂	2	OPLA	1147.50 1129.2	gas Ar	DL,IR IR	2,5 1,4
e'	3	BH ₃ stretch	2601.57 2587.3	gas Ar	IR IR	3 4
	4	Deformation	1196.7	gas	IR	5

 $B_0 = 7.874; C_0 = 3.879 \text{ IR}^3$ **BD₃** \tilde{X} D_{3h}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'' ₂	2	OPLA	878.5	Ar	IR	4
e'	3	BD ₃ stretch	1953.4	Ar	IR	4

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AlH₃ \tilde{X} D_{3h}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'' ₂	2	OPLA	697.8m	Ar	IR	1-3
e'	3	AlH ₃ stretch	1882.8vs	Ar	IR	1-3
	4	Deformation	783.4m	Ar	IR	1-3

AlD₃ \tilde{X} D_{3h}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'' ₂	2	OPLA	512.5s	Ar	IR	1-3
e'	3	AlD ₃ stretch	1377.9vs	Ar	IR	1-3
	4	Deformation	568.5m	Ar	IR	1-3

References

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GaH₃

\tilde{X} D _{3h}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	717.4ms	Ar	IR	1
e'	3	GaH ₃ stretch	1923.2vs	Ar	IR	1
	4	Deformation	758.7s	Ar	IR	1

GaD₃

\tilde{X} D _{3h}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	517.5s	Ar	IR	1
e'	3	GaD ₃ stretch	1387.7vs	Ar	IR	1
	4	Deformation	544.0m	Ar	IR	1

Reference

¹P. Pullumbi, Y. Bouteiller, L. Manceron, and C. Mijoule, Chem. Phys. **185**, 25 (1994).

InH₃

\tilde{X} D _{3h}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	613.2m	Ar	IR	1
e'	3	InH ₃ stretch	1754.5vs	Ar	IR	1
	4	Deformation	607.8s	Ar	IR	1

InD₃

\tilde{X} D _{3h}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	439.7	Ar	IR	1
e'	3	InD ₃ stretch	1261.2	Ar	IR	1
	4	Deformation	433.3	Ar	IR	1

Reference

¹P. Pullumbi, Y. Bouteiller, L. Manceron, and C. Mijoule, Chem. Phys. **185**, 25 (1994).

CH₃⁺

\tilde{A}'^1E'	D _{3h}
$T_0=50510(280)$	gas PE ²
\tilde{a}'^3E'	D _{3h}
$T_0=39700(280)$	gas PE ²
$\tilde{X}'^1A'_1$	D _{3h}
	Structure: LD ^{3,4}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	1380(20)	gas	PE	1,2
e'	3	CH stretch	3108.38	gas	LD	3-5
$B_0=9.362$ LD ^{3,5}						

CD₃⁺

\tilde{X}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	1070(30)	gas	PE	2

References

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SiH₃⁺

\tilde{X} D _{3h} Structure: DL ¹						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	838.067	gas	DL	1,2
e'	4	SiH stretch	938.397	gas	DL	2
$B_0=5.214; C_0=2.585$ DL ^{1,2}						

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CH₃**4f** $^2E'$ ^aD_{3h} $T_0 = 72508$ gas MPI¹²Higher member of Rydberg series observed at 74961. MPI¹²**4p** $^2A''_2$ D_{3h} $T_0 = 69853.44(13)$ gas MPI¹⁵ $B_0 = 9.90$ gas MPI¹⁵**3d** $^2A'_1$ D_{3h}Structure: AB² $T_0 = 66805$ gas AB^{1,2}Ar AB³ $3d^2A'_1 - \bar{X}$ 147–150 nm $3d^2A'_1 - \bar{X}$ ~150.3 nmFirst member of Rydberg series converging to 79392(5). Higher members observed at 72326, 74851, 76256, 77090, and 77643. AB² $B_0 = 10.72(8)$ AB²**3d** $^2E''$ D_{3h}Structure: AB² $T_0 = 66536$ gas AB^{1,2} MPI¹⁰Ar AB³ $3d^2E'' - \bar{X}$ 144–150 nm $3d^2E'' - \bar{X}$ ~150.3 nmDiffuse. First member of Rydberg series converging to 79392(5). Higher members observed at 72165, 74851, 76256, 77090, and 77643. AB²

Vib.

No.

Approximate type of mode

cm⁻¹

Med.

Type

meas.

Refs.

a''₂

2

OPLA

Structure: MPI¹² $T_0 = 59972$ gas MPI^{12,28}Higher member of Rydberg series observed at 69837. MPI¹²

Vib.

No.

Approximate type of mode

cm⁻¹

Med.

Type

meas.

Refs.

a'₁

1

CH stretch

Structure: AB² $T_0 = 46205$ gas AB^{1,2,7} Ra²⁴ $3s^2A'_1 - \bar{X}$ 216 nmDiffuse. First member of Rydberg series converging to 79392(5). Next member observed at 71042. AB²

Vib.

No.

Approximate type of mode

cm⁻¹

Med.

Type

meas.

Refs.

a'₁

1

CH stretch

Structure: AB² $T_0 = 46205$ gas AB^{1,2,7} Ra²⁴ $3s^2A'_1 - \bar{X}$ 216 nmDiffuse. First member of Rydberg series converging to 79392(5). Next member observed at 71042. AB²

Vib.

No.

Approximate type of mode

cm⁻¹

Med.

Type

meas.

Refs.

a'₁

1

CH stretch

Structure: AB² $T_0 = 46205$ gas AB^{1,2,7} Ra²⁴ $3s^2A'_1 - \bar{X}$ 216 nmDiffuse. First member of Rydberg series converging to 79392(5). Next member observed at 71042. AB²

Vib.

No.

Approximate type of mode

cm⁻¹

Med.

Type

meas.

Refs.

a'₁

1

CH stretch

Structure: AB² $T_0 = 46205$ gas AB^{1,2,7} Ra²⁴ $3s^2A'_1 - \bar{X}$ 216 nmDiffuse. First member of Rydberg series converging to 79392(5). Next member observed at 71042. AB²

Vib.

No.

Approximate type of mode

cm⁻¹

Med.

Type

meas.

Refs.

a'₁

1

CH stretch

Structure: AB² $T_0 = 46205$ gas AB^{1,2,7} Ra²⁴ $3s^2A'_1 - \bar{X}$ 216 nmDiffuse. First member of Rydberg series converging to 79392(5). Next member observed at 71042. AB²**X** $^2A''_2$ D_{3h} Structure: AB²IR^{9,18}CAR^{23,26}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i> ₁	1	CH stretch	3004.43(2)	gas	CAR	13,16,23
<i>a''₂</i>	2	OPLA	606.453	gas	IR,DL	5,9,21
			617vs	Ne	IR	4
			603 ^b	Ar	IR	3,8
			611	N ₂	IR	3
<i>e'</i>	3	CH stretch	3160.821	gas	LD	11,25,27
			3162wm	Ne	IR	4
			3150	Ar	IR	6
			3171.4	H ₂	IR	29
			3170.6			
4		Deformation	1396w	Ne	IR	4
			1398 ^c	Ar	IR	8
			1402.7	H ₂	IR	29
			1401.6			

 $B_0 = 9.578$ AB²DL⁹; $C_0 = 4.742$ DL⁹**CD₃****4f** $^2E'$ ^aD_{3h} $T_0 = 72431$ gas MPI¹²Higher member of Rydberg series observed at 74885. MPI¹²**4p** $^2A''_2$ D_{3h} $T_0 = 69777.40(4)$ gas MPI¹⁵ $B_0 = 4.846(2)$ gas MPI¹⁵**3d** $^2A'_1$ D_{3h} $T_0 = 66715$ gas AB^{1,2}

Ar

AB³Structure: AB² $3d^2A'_1 - \bar{X}$ 145–150 nm $3d^2A'_1 - \bar{X}$ ~150.3 nmFirst member of Rydberg series converging to 79315(5). Higher members observed at 72296, 74781, 76181, 77023, 77562, and 77933. AB²**a''₂**

2

OPLA

1040H

gas

AB

2

 $B_0 = 5.14$ AB²**3d** $^2E''$ D_{3h} $T_0 = 66465$ gas AB^{1,2} MPI¹⁰

Ar

AB³Structure: AB² $3d^2E'' - \bar{X}$ 146–150 nm $3d^2E'' - \bar{X}$ ~150.3 nmDiffuse. First member of Rydberg series converging to 79315(5). Higher members observed at 72180, 74753, 76166, 77023, 77562, and 77933. AB²**a''₂**

2

OPLA

1031H

gas

AB,MPI

2,10

3p $^2A''_2$ D_{3h}Structure: MPI¹ $T_0 = 59886$ gas MPI^{12,17}Higher members of Rydberg series observed at 69789, 73645, and 75557. MPI¹²**a'₁**

1

CD stretch

Structure: MPI¹ $a''₂$

2

OPLA

1032

gas

MPI

12,17

$B_0 = 4.76(2)$; $C_0 = 2.38$ MPI^{12,17}

$3s^2 A'_1$ D_{3h} Structure: AB²
 $T_0 = 46629$ gas AB^{1,2,7}Ra²⁴ 3s²A'_1-X 204-225 nm
 First member of Rydberg series converging to 79315(5). Higher members observed at 70910, 74246, 75869, and 76830. AB²

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	CD stretch	1684 ^d	gas	AB	7
a''_2	2	OPLA	1094	gas	AB,Ra	7,24

$B_0 = 4.42$ AB²

$\bar{X}^2 A''_2$ D_{3h}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	CD stretch	2157.5(2)	gas	Ra, CAR	19,20
a''_2	2	OPLA	457.81	gas	DL	14,18
			463s	Ne	IR	4
			453 ^b	Ar	IR	3,8
			463	N ₂	IR	3
e'	3	CD stretch	2381.09	gas	IR,DL	22,31
			2381w	Ne	IR	4
			2369	Ar	IR	6
	4	Deformation	1026vw	Ne	IR	4
			1029	Ar	IR	6

$B_0 = 4.802$ AB²DL¹⁴

^aTentative assignment.

^bBand center. Rotational structure assigned.⁸

^cR(0₀) transition.

^dApproximate value; perturbed by Fermi resonance.

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SiH₃

$\tilde{M}^2 A''_2$ (6p) D_{3h}
 $T_0 = 60341$ gas MPI⁷

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	800(30)	gas	MPI	7

$\tilde{L}^2 (5d)$ D_{3h}
 $T_0 = 59615(30)^c$ gas MPI⁸

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	839(26)	gas	MPI	8

$\tilde{J}' (4d)$ D_{3h}
 $T_0 = 57726(30)$ gas MPI⁸

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	835(26)	gas	MPI	8

$\tilde{J}^2 A''_2$ (5p) D_{3h}
 $T_0 = 56929$ gas MPI⁷

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	821(4)	gas	MPI	7

$\tilde{I}' (4d)$ D_{3h}
 $T_0 = 56253(30)$ gas MPI⁸

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	814(25)	gas	MPI	8

\tilde{D}^{\prime}	$2A'_1$	(3d)	D _{3h}			
$T_0=49787(30)$	gas	MPI ⁸				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	810(31)	gas	MPI	8
<hr/>						
\tilde{E}^{\prime}	$2A''_2$	(4p)	D _{3h}			
$T_0=48438$	gas	MPI ^{5,7}				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	796(7)	gas	MPI	5,7
e'	4	Deformation	870(5)H	gas	MPI	7
<hr/>						
\tilde{A}^{\prime}	$2A_1$	C _{3v}				
		gas	AR ⁹			
				$\tilde{A}-\tilde{X}$	205-250 nm	
<hr/>						
\tilde{X}^{\prime}	$2A_1$	C _{3v}	Structure: ESR ^{1,2} DL ³			
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	Umbrella	727.94 ^a	gas	DL,MPI	3,6,7
			721.05 ^b	gas	DL,MPI	3,6,7
e	3	SiH ₃ stretch	2185.2	gas	DL	10
<hr/>						
Barrier to inversion = 1935	gas	PE ⁴ MPI ⁷				
B ₀ =4.763	DL ³					
<hr/>						
SiD₃						
\tilde{P}^{\prime}	$2A''_2$	(7p)	D _{3h}			
$T_0=62002$	gas	MPI ⁷				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	615(3)	gas	MPI	7
<hr/>						
\tilde{N}^{\prime}	$2E'$	(5f)	D _{3h}			
$T_0=61005$	gas	MPI ^{7,8}				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	619(7)	gas	MPI	7
<hr/>						
\tilde{M}^{\prime}	$2A''_2$	(6p)	D _{3h}			
$T_0=60267$	gas	MPI ⁷				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	615(4)	gas	MPI	7
<hr/>						
\tilde{K}^{\prime}	$2E'$	(4f)	D _{3h}			
$T_0=58417$	gas	MPI ^{7,8}				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	602(10)	gas	MPI	8

\tilde{J}'	(4d)	D _{3h}				
$T_0=57840(30)^c$	gas	MPI ⁸				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	603(20)	gas	MPI	8
<hr/>						
\tilde{J}^{\prime}	$2A''_2$	(5p)	D _{3h}			
$T_0=56874$	gas	MPI ⁷				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	608(3)	gas	MPI	7
<hr/>						
\tilde{I}'	(4d)	D _{3h}				
$T_0=56205(30)$	gas	MPI ⁸				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	600(17)	gas	MPI	8
<hr/>						
\tilde{H}^{\prime}	$2E'$	(4p)	D _{3h}			
$T_0 \approx 50000$	gas	MPI ^{7,8}				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	602(5)	gas	MPI	7
<hr/>						
\tilde{D}^{\prime}	$2A'_1$	(3d)	D _{3h}			
$T_0=49685(30)^c$	gas	MPI ⁸				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2	2	OPLA	600(28)	gas	MPI	8
<hr/>						
\tilde{E}^{\prime}	$2A''_2$	(4p)	D _{3h}			
$T_0=48391$	gas	MPI ⁷				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	SiD ₃ stretch	1576(3)	gas	MPI	7
a''_2	2	OPLA	589(3)	gas	MPI	7
e'	4	Deformation	635(6)H	gas	MPI	7
<hr/>						
\tilde{X}^{\prime}	$2A_1$	C _{3v}				
<hr/>						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	Umbrella	545 ^a	gas	MPI	7
			542 ^b	gas	MPI	7

Barrier to inversion = 1925 gas MPI⁷^a1⁻-0⁺ transition.^b1⁺-0⁻ transition.^cExtrapolated value.

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NH₃⁺

\tilde{A}^2E D_{3h}
 $T_0 = 36590(100)$ gas PE^{3,4}

Broad, partially resolved vibrational structure has been discussed³ in terms of the expected Jahn-Teller distortion. Continuous background absorption may be associated with the formation of NH₂⁺, for which the threshold is ~44700,^{1,3} or NH⁺.

$\tilde{X}^2A''_2$ D_{3h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	NH stretch	3150(100)	gas	PE	5,10
a''_2	2	OPLA	903.39	gas	PE,DL	3,7
			916.8	Ne	IR	11
e'	3	NH stretch	3388.65	gas	LD	6
			3404.6	Ne	IR	11
	4	Bend	1507.1	gas	PE,TPE	8-10
			1516.8	Ne	IR	11

$B_0 = 10.644$; $C_0 = 5.247$ LD⁶DL⁷

ND₃⁺

$\tilde{X}^2A''_2$ D_{3h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	ND stretch	2453(56)	gas	PE	10
a''_2	2	OPLA	725(25)	gas	PE	2
			683.3	Ne	IR	11
e'	3	ND stretch	2551.8	Ne	IR	11
	4	Bend	1138(56)	gas	PE	10
			1114.6	Ne	IR	11

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H₃S⁺

\tilde{X} C_{3v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	SH stretch	2521.05	gas	LD	3
	2	Umbrella	1033.31	gas	DL	2
e	3	SH stretch	2525.74	gas	LD	1,3

$B_0 = 4.895$; $C_0 = 4.228$ LD^{1.3}DL²MW⁴

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⁴S. K. Lee, H. Ozeki, S. Saito, and S. Yamamoto, Chem. Phys. Lett. **224**, 21 (1994).

8.5. Four-Atomic Dihydrides**HBeBeH**

\tilde{X} D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	BeH a-stretch	1900.6	Ar	IR	1

DBeBeD

\tilde{X} D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	BeD a-stretch	1533.0	Ar	IR	1

Reference

- ¹T. J. Tague, Jr. and L. Andrews, J. Am. Chem. Soc. **115**, 12111 (1993).

HMgMgH

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		MgH stretch	1491.8	Ar	IR	1

DMgMgD \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		MgD stretch	1084.3	Ar	IR	1

Reference¹T. J. Tague, Jr. and L. Andrews, *J. Phys. Chem.* **98**, 8611 (1994).**br-(MgH)₂** \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1022.8	Ar	IR	1
			605.4	Ar	IR	1

br-(MgD)₂ \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			744.6	Ar	IR	1

Reference¹T. J. Tague, Jr. and L. Andrews, *J. Phys. Chem.* **98**, 8611 (1994).**HZnZnH** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		ZnH stretch	1740.3	Ar	IR	1

DZnZnD \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		ZnD stretch	1254.9	Ar	IR	1

Reference¹T. M. Greene, W. Brown, L. Andrews, A. J. Downs, G. V. Chertihin, N. Runeberg, and P. Pyykkö, *J. Phys. Chem.* **99**, 7925 (1995).**HHgHgH** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		HgH stretch	1792.0T	N ₂	IR	1

DHgHgD \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		HgD stretch	1286.6T	N ₂	IR	1

Reference¹N. Legay-Sommaire and F. Legay, *J. Phys. Chem.* **99**, 16945 (1995).**cyc-U₂H₂** \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>b</i> _{2u}		UH stretch	1182.4	Ar	IR	1

cyc-U₂D₂ \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>b</i> _{2u}		UD stretch	845.6	Ar	IR	1

Reference¹P. F. Souter, G. P. Kushto, L. Andrews, and M. Neurock, *J. Am. Chem. Soc.* **119**, 1682 (1997).**CrCH₂** \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₂ s-stretch	2907.4	Ar	IR	1
	3	CrC stretch	567.0	Ar	IR	1
<i>b</i> ₁	4	OPLA	687.7	Ar	IR	1
	5	CH ₂ a-stretch	2966.7	Ar	IR	1
<i>b</i> ₂	6	CH ₂ rock	450.3	Ar	IR	1

CrCD₂

\tilde{X}	C _{2v}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CrC stretch	525.7	Ar	IR	1
<i>b</i> ₁	4	OPLA	538.8	Ar	IR	1
<i>b</i> ₂	6	CD ₂ rock	346.2	Ar	IR	1

Reference

¹W. E. Billups, S.-C. Chang, R. H. Hauge, and J. L. Margrave, Inorg. Chem. **32**, 1529 (1993).

CoCH₂

\tilde{X}	C _{2v}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₂ s-stretch	2918.0	Ar	IR	1,2
	2	CH ₂ scissors	1327.0	Ar	IR	1,2
	3	CoC stretch	655.4	Ar	IR	1,2
<i>b</i> ₁	4	OPLA	757.4	Ar	IR	1,2
<i>b</i> ₂	5	CH ₂ a-stretch	2979.7	Ar	IR	1,2
	6	CH ₂ rock	587.5	Ar	IR	1,2

CoCD₂

\tilde{X}	C _{2v}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CD ₂ scissors	993.2	Ar	IR	1,2
	3	CoC stretch	608.2	Ar	IR	1,2
<i>b</i> ₁	4	OPLA	599.5	Ar	IR	1,2
<i>b</i> ₂	6	CD ₂ rock	455.4	Ar	IR	1,2

References

¹W. E. Billups, S.-C. Chang, R. H. Hauge, and J. L. Margrave, J. Am. Chem. Soc. **115**, 2039 (1993).

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NiCH₂

\tilde{X}	C _{2v}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₂ s-stretch	2917.1	Ar	IR	1
	2	CH ₂ scissors	1328.5	Ar	IR	1
	3	NiC stretch	696.2	Ar	IR	1
<i>b</i> ₁	4	OPLA	791.9	Ar	IR	1
<i>b</i> ₂	5	CH ₂ a-stretch	2973.2	Ar	IR	1
	6	CH ₂ rock	586.7	Ar	IR	1

NiCD₂

\tilde{X}	C _{2v}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CD ₂ scissors	993.6	Ar	IR	1
	3	NiC stretch	641.7	Ar	IR	1
<i>b</i> ₁	4	OPLA	627.0	Ar	IR	1
<i>b</i> ₂	6	CD ₂ rock	455.1	Ar	IR	1

Reference

¹S.-C. Chang, R. H. Hauge, Z. H. Kafafi, J. L. Margrave, and W. E. Billups, Inorg. Chem. **29**, 4373 (1990).

HBBH

\tilde{X}	C _{2v}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		BH stretch	2679.9	Ar	IR	1

Reference

¹T. J. Tague, Jr. and L. Andrews, J. Am. Chem. Soc. **116**, 4970 (1994).

cyc-Al₂H₂

\tilde{X}	D _{2h}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> _{1u}			1160.9	Ar	IR	1,2
<i>b</i> _{2u}			844	Ar	IR	1,2

cyc-Al₂D₂

\tilde{X}	D _{2h}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> _{1u}			852.8	Ar	IR	1
<i>b</i> _{2u}			617.2	Ar	IR	1

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H(cyc-AlHAl)

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>			1668.7	Ar	IR	1,2
			1127	Ar	IR	1,2
			890	Ar	IR	1,2

D(cyc-AIDAI)

\tilde{X}	C_s					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'			1210	Ar	IR	1,2
			811	Ar	IR	1,2

References

- ¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **97**, 10295 (1993).
²J. C. Stephens, E. E. Bolton, H. F. Schaefer III, and L. Andrews, *J. Chem. Phys.* **107**, 119 (1997).

HAIAIH

\tilde{X}	$D_{\infty h}$					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1646.9T	Ar	IR	1

DAIAID

\tilde{X}	$D_{\infty h}$					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1194.3T	Ar	IR	1

Reference

- ¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **97**, 10295 (1993).

cyc-Ga₂H₂

In an argon or a krypton matrix, irradiation at 400–520 nm results in rearrangement to HGaGaH.¹

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		GaH stretch	1002.1	Ar	IR	1
			1022.3	Kr	IR	1

cyc-Ga₂D₂

\tilde{X}						
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		GaD stretch	728.2	Ar	IR	1
			732.0	Kr	IR	1

Reference

- ¹Z. L. Xiao, R. H. Hauge, and J. L. Margrave, *Inorg. Chem.* **32**, 642 (1993).

HGaGaH

In an argon or a krypton matrix, irradiation at 320–380 nm results in rearrangement to cyc-Ga₂H₂.

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		GaH stretch	1686.2	Ar	IR	1

DGaGaD \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		GaD stretch	1206.5	Ar	IR	1
			1218.2	Kr	IR	1

Reference

- ¹Z. L. Xiao, R. H. Hauge, and J. L. Margrave, *Inorg. Chem.* **32**, 642 (1993).

CaNH₂

\tilde{C}^2A_1 C_{2v} Structure: LF^{2,7}
 $T_0=17375.167$ gas CL¹LF^{2,7} $\tilde{C}-\tilde{X}$ 575 nm

$$A=12.95; \frac{1}{2}(B+C)=0.302; \frac{1}{4}(B-C)=0.0018 \quad \text{LF}^{2,7}$$

\tilde{B}^2B_1 C_{2v} Structure: LF⁷

$T_0=15885.28$ gas CL¹LF^{3-5,7} $\tilde{B}-\tilde{X}$ 620–650 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	3	CaN stretch	545.8(5)	gas	LF	4

$$A=14.37; \frac{1}{2}(B+C)=0.302; \frac{1}{4}(B-C)=0.005 \quad \text{LF}^{5,7}$$

\tilde{A}^2B_2 C_{2v} Structure: LF⁷

$T_0=15464.367$ gas CL¹LF³⁻⁶ $\tilde{A}-\tilde{X}$ 620–650 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	3	CaN stretch	549.5(1.0)	gas	LF	3,4

$$A=11.449; B_0=0.307; C_0=0.299 \quad \text{LF}^{5,6}$$

\tilde{X}^2A_1 C_{2v} Structure: LF^{2,6,7}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	3	CaN stretch	524(10)	gas	LF	3

$$A=13.057; B_0=0.300; C_0=0.293 \quad \text{LF}^{2,5,6}$$

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- ²R. F. Wormsbecher, R. E. Penn, and D. O. Harris, *J. Mol. Spectrosc.* **97**, 65 (1983).

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⁷Z. Morbi, C. Zhao, and P. F. Bernath, J. Chem. Phys. **106**, 4860 (1997); J. Chem. Phys. **107**, 1297 (1997).

HBCH

\tilde{X}		$C_{\infty v}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
Σ^+	1	CH stretch	3248.8	Ar	IR	1	
	2	BH stretch	2724.6	Ar	IR	1	
	3	B=C stretch	1475.3	Ar	IR	1	

DBCD

\tilde{X}		$C_{\infty v}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
Σ^+	1	CD stretch	2464.3	Ar	IR	1	
	2	BD stretch	2097.8	Ar	IR	1	

Reference

¹P. Hassanzadeh, Y. Hannachi, and L. Andrews, J. Phys. Chem. **97**, 6418 (1993).

HCCH⁺

$\tilde{C} \ 2\Sigma^+$		$D_{\infty h}$					
$T_0=92460(80)$		gas	PE ⁶				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
Σ_g^+	2	CC stretch	1370T	gas	PE	6	
$\tilde{B} \ 2\Sigma_u^+$		$D_{\infty h}$					
$T_0=56380(80)$		gas	PE ^{1,3}				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
Σ_g^+	1	CH s-stretch	2500(20)	gas	PE	1,3	
	2	CC stretch	1815(20)	gas	PE	1,3	
$\tau < 14 \text{ fs}$		gas	PE ³				

$\tilde{A} \ 2A_g^a$ C_{2h} Structure: MPD¹¹
 $T_0=39109.7(1.0)$ gas PE³MPD¹¹ $\tilde{A}-\tilde{X} \ 240-255 \text{ nm}$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_g	1	CH s-stretch	2530(20)	gas	PE	3
	2	CC stretch	1730(20)	gas	PE	3
	3	Bend	793 ^b	gas	MPD	11
a_u	4	Torsion	282H ^b	gas	MPD	11
b_u	6	Bend	436H ^b	gas	MPD	11

$\tau_0=150 \text{ ps}$ gas MPD¹¹
 $A_0=12.15$; $B_0=0.94(5)$ MPD¹¹

$\tilde{X} \ 2\Pi_u$ $D_{\infty h}$ Structure: LD⁹

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	2	CC stretch	1829(3)	gas	PE	1,3
Σ_u^+	3	CH a-stretch	3135.98	gas	LD	5,9
			3137.6	Ne	IR	8
			3105.5	Ar	IR	4,8
Π_g	4	Deform.	694 ^c	gas	PE,TPE	7,10
Π_u	5	Deform.	837(12) ^c	gas	PE	3

$A=-30.91(2)$ gas LD^{5,9}

$B_0=1.105$ LD^{5,9}

DCCD⁺

$\tilde{B} \ 2\Sigma_u^+$ $D_{\infty h}$ $T_0=56655(80)$ gas PE³

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	CC stretch	2275(20)	gas	PE	1,3
	2	CD s-stretch	1475(20)	gas	PE	1,3

$\tilde{A} \ 2A_g$ C_{2h} $T_0=39906(80)$ gas PE³

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_g	1	CD s-stretch	2280(20)	gas	PE	3
	2	CC stretch	1450(20)	gas	PE	3
	3	Bend	339(12)	gas	PE	3
b_u	6	Bend	516(12)	gas	PE	3

$\tilde{X} \ 2\Pi_u$ $D_{\infty h}$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	CD s-stretch	2572(16)	gas	PE	3
	2	CC stretch	1651(4)	gas	PE	1,3
Σ_u^+	3	CD a-stretch	2329.1	Ne	IR	8
			2311T	Ar	IR	4,8
Π_u	5	Bend	702(12) ^c	gas	PE	3

^aThreshold for formation of $\text{HCC}^+ \leq 48000 \text{ cm}^{-1}$ ^{2,3}

^bAll three bending fundamentals have exceptionally large anharmonicities.

^c ω_4 ; $\epsilon_4=0.30$.¹⁰ The identification^{7,10} of the Renner components of ν_4 (Π_g) of HCCH^+ in the photoelectron spectrum of HCCH ($\tilde{A} \ 1A_g$) necessitates the reassignment to ν_5 (Π_u) of the peaks previously assigned³ to ν_4 of HCCH^+ and DCCD^+ .

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HBeOH \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OH stretch	3841.8	Ar	IR	1
	2	BeH stretch	2117.7	Ar	IR	1
	3	BeO stretch	1206.9	Ar	IR	1
		Deformation	552.0	Ar	IR	1

DBeOD \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OD stretch	2834.2	Ar	IR	1
	2	BeD stretch	1663.6	Ar	IR	1
	3	BeO stretch	1098.3	Ar	IR	1

Reference

- ¹C. A. Thompson and L. Andrews, J. Phys. Chem. **100**, 12214 (1996).

HZnOH \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		HZn stretch	1955.0	Ar	IR	1
		ZnO stretch	660.0	Ar	IR	1
			481.4	Ar	IR	1

DZnOD \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		DZn stretch	1410.0	Ar	IR	1

Reference

- ¹T. M. Greene, W. Brown, L. Andrews, A. J. Downs, G. V. Chertihin, N. Runeberg, and P. Pyykkö, J. Phys. Chem. **99**, 7925 (1995).

HCdOH \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		HCd stretch	1836.9	Ar	IR	1

DCdOD \bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		DCd stretch	1319.8	Ar	IR	1

Reference

- ¹T. M. Greene, W. Brown, L. Andrews, A. J. Downs, G. V. Chertihin, N. Runeberg, and P. Pyykkö, J. Phys. Chem. **99**, 7925 (1995).

HBNH \bar{X} $C_{\infty v}$

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	NH stretch	3710.5m	Ar	IR	3,5,6
			3700.5m			
	2	BH stretch	2775.3	Ar	IR	6
Π	3	BN stretch	1786.19	gas	DL	4
			1788.7m	Ar	IR	3,5,6
			1782.8m			
Π	4	BH deform.	676.9	Ar	IR	6
			671.6			
Σ^+	5	NH deform.	463.3m	Ar	IR	3,5,6
			461.0m			

 $B_0 = 1.099 \text{ DL}^4$ **DBND** \bar{X} $C_{\infty v}$

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	ND stretch	2836	Ar	IR	3
	2	BD stretch	2/30	Ar	IR	3
	3	BN stretch	1734	Ar	IR	3
Π	4	Bend	360	Ar	IR	3

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H₂NB \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1		NH ₂ scissors	1533.2	Ar	IR	1
b_1	4	OPLA	596.0	Ar	IR	1

Reference

¹C. A. Thompson, L. Andrews, J. M. L. Martin, and J. El-Yazal, *J. Phys. Chem.* **99**, 13839 (1995).

H₂CSi

1B_2 C_{2v}
 $T_0=29312.883(4)$ gas AB¹

$^1B_2-\tilde{X}$ 310–340 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CH ₂ scissors	1101.96	gas	AB	1
	3	CSi stretch	702.00	gas	AB	1

$A_0=8.537$; $B_0=0.509$; $C_0=0.479$ AB¹

\tilde{X} 1A_1 C_{2v} Structure: MW²

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CH ₂ rock	331T	gas	MW	2

$A_0=10.167(18)$; $B_0=0.553$; $C_0=0.521$ AB¹MW²

D₂CSi

1B_2 C_{2v}
 $T_0=29272$ gas AB¹

$^1B_2-\tilde{X}$ 310–340 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CD ₂ scissors	829	gas	AB	1
	3	CSi stretch	691	gas	AB	1

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br-Si₂H₂

\tilde{X} C_{2v} Structure: MW^{1,2}
 $A_0=5.244$; $B_0=0.243$; $C_0=0.240$ MW^{1,2}

br-Si₂D₂

\tilde{X} C_{2v}
 $A_0=2.688$; $B_0=0.240$; $C_0=0.234$ MW²

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HGaOH

\tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		OH stretch	3675	Ar	IR	1
		HGa stretch	1669.8	Ar	IR	1,2
		HGaO bend	784.9	Ar	IR	1,2
		GaO stretch	646.4	Ar	IR	1,2
		GaOH bend	520.5	Ar	IR	1

DGaOD

\tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		OD stretch	2708	Ar	IR	1
		DGa stretch	1213.8	Ar	IR	1
		GaO stretch	644.6	Ar	IR	1
		DGaO bend	582.9	Ar	IR	1

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²P. Pullumbi, C. Mijoule, L. Manceron, and Y. Bouteiller, *Chem. Phys.* **185**, 13 (1994).

H₂CO⁺

\tilde{C} 2B_2 C_{2v}
 $T_0=43172(11)$ gas PE^{1,4,5}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1			1895T	gas	PE	4,5
			1412T	gas	PE	1,4,5

\tilde{B} 2A_1 C_{2v}
 $T_0=39928(6)$ gas PE^{1,4,5}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CO stretch	1304(4)	gas	PE	1,4,5

\tilde{A} 2B_1 C_{2v}
 $T_0=25929(5)$ gas PE^{1,4,5}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1			1488(4)	gas	PE	4,5
			1250(4)	gas	PE	1,4,5
b_1	4	OPLA	263(4)	gas	PE	4,5

\tilde{X}^2B_2 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH stretch	2580(4)	gas	PE	1,4,5
	2	CO stretch	1675(4)	gas	PE	4,5
	3	CH ₂ scissors	1210(4)	gas	PE	1,4,5
<i>b</i> ₁	4	OPLA	777(4)	gas	PE	4,5

 $A_0=8.875(3); B_0=1.31(1); C_0=1.14(1)$ TPE⁶ D_2CO^+ \tilde{C}^2B_2 C_{2v} $T_0=44600(160)$ gas PE^{4,5}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁		1604T	gas	PE	4,5	
		943T	gas	PE	1,4,5	
		819T	gas	PE	4,5	

 \tilde{B}^2A_1 C_{2v} $T_0=39816(5)$ gas PE^{1,4,5}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CO stretch	1311(4)	gas	PE	1,4,5
	3	CD ₂ scissors	957(4)	gas	PE	1,4,5

 \tilde{A}^2B_1 C_{2v} $T_0=25756(7)$ gas PE^{1,4,5}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2		1282(4)	gas	PE	1,4,5
	3		1064(4)	gas	PE	4,5
<i>b</i> ₁	4	OPLA	777(4)	gas	PE	5

 $\tau_0=64(22)\mu s$ gas PEPICO³ \tilde{X}^2D_2 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CD stretch	1948(4)	gas	PE	1,4,5
	2	CO stretch	1657(4)	gas	PE	4,5
	3	CD ₂ scissors	920(4)	gas	PE	1,4,5
<i>b</i> ₁	4	OPLA	648(4)	gas	PE	4,5

 H_2CS $\tilde{E}^3p_z^1B_2$ C_{2v} $T_0=55096$ gas AB^{8,19} $\tilde{E}-\tilde{X}$ 181.5 nm $\tilde{D}^3p_y^1A_1$ C_{2v} $T_0=53134$ gas AB^{8,19} $\tilde{D}-\tilde{X}$ 188.2 nm $\tilde{C}^3s^1B_2$ C_{2v} $T_0=47110.821(9)$ gas AB^{1,8,15,19} $\tilde{C}-\tilde{X}$ 212.1 nm $A_0=8.557; B_0=0.603; C_0=0.562$ AB¹⁹ $\tilde{B}^1A_1^a$ C_{2v} $T_0=45197$ gas AB^{8,18} $\tilde{B}-\tilde{X}$ 185–215 nmAll but the first absorption band show evidence for predissociation.¹⁸ \tilde{A}^1A_2 C_{2v} $T_0=16394.628(4)$ gas AB^{7,9,10}LF^{24,28,32,33} $\tilde{A}-\tilde{X}$ 440–610 nm $\tilde{Vib.}$

Vib.

No.

Approximate

type of mode

cm⁻¹

Med.

Type

meas.

Refs.

*a*₁ 1 CH stretch 3033.4^b2 CH₂ scissors 1334.5^b

3 CS stretch 819.7

4 OPLA 370.3

5 CH stretch 3054.9^b6 CH₂ rock 785.2

gas LF 28

AB,LF 9,28

AB,LF 7,9,28

AB,LF 9,28,32

LF 28

LF 28

LF 28

 $\tau_0=140(3)\mu s$ gas LF^{22,25} $A_0=9.434; B_0=0.538; C_0=0.509$ AB¹⁰LF^{24,28} \tilde{a}^3A_2 C_{2v} $T_0=14507.38$ gas AB^{7,11}LF^{20,24,29}CL^{26,30} $\tilde{a}-\tilde{X}$ 610–800 nm $\tilde{Vib.}$

Vib.

No.

Approximate

type of mode

cm⁻¹

Med.

Type

meas.

Refs.

*a*₁ 2 CH₂ scissors 1318.6

3 CS stretch 861.6

4 OPLA 326

6 CH₂ rock 762.3

gas AB,LF 11,29

AB,LF 11,23

LF,CL 20,21,26

30,32

LF 23

 $\tau>1.5\text{ ms}$ gas LF^{22,25} $A_0=9.383; B_0=0.552; C_0=0.521$ AB¹¹LF²⁴

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\tilde{X}^1A_1		C_{2v}	Structure: MW ^{2,4,5} IR ^{3,14}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CH stretch	2971.03	gas	IR	3,14
			2970w	Ar	IR	6,17
			2973w	N ₂	IR	6
	2	CH ₂ scissors	1455.50	gas	LF,IR	13,31
			1447	Ar	IR	17
	3	CS stretch	1059.20	gas	LS,IR	12,14
b_1			1063w	Ar	IR	6,17
			1062w	N ₂	IR	6
	4	OPLA	990.19	gas	LS,IR	12,14
			993s	Ar	IR	6,17
			995s	N ₂	IR	6
	5	CH stretch	3024.61	gas	IR	3,14
b_2	6	CH ₂ rock	991.01	gas	LS,IR	12,14
			988m	Ar	IR	6,17

$A_0=9.729$; $B_0=0.590$; $C_0=0.555$ MW^{2,4,5}AB¹⁰IR³¹

D₂CS

$\tilde{C}^3s^1B_2$		C_{2v}	Structure: AB ^{8,19}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CD stretch	1783	gas	AB	8,19
	2	CD ₂ scissors	746	gas	AB	19

$A_0=4.350$; $B_0=0.510$; $C_0=0.456$ AB¹⁹

$\tilde{B}^1A_1^a$		C_{2v}	Structure: AB ¹⁸			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	3	CS stretch	467	gas	AB	18
b_1	4	OPLA	263H	gas	AB	18

\tilde{A}^1A_2		C_{2v}	Structure: AB ^{7,9,10}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CD stretch	2139(2)	gas	AB	9
	2	CD ₂ scissors	1013(2)	gas	AB	9
	3	CS stretch	771.3(5)	gas	AB	7,9
b_1	4	OPLA	275.33	gas	AB	9
b_2	5	CD stretch	2324.85	gas	AB	9
	6	CD ₂ rock	599(2)	gas	AB	9

$\tau_0 = 182 \mu\text{s}$ gas LF²⁵
 $A_0=4.736$; $B_0=0.458$; $C_0=0.417$ AB¹⁰

\tilde{a}^3A_2		C_{2v}	Structure: AB ²¹			
$T_0=14613.54$	gas	AB ^{7,11} CL ^{26,30}	$\tilde{a}-\tilde{X}$ 610–815 nm			

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CD ₂ scissors	1012	gas	AB,CL	11,30
	3	CS stretch	798	gas	AB,CL	7,11,30
	4	OPLA	219.4(5.0)	gas	AB,CL	21,26,30
	6	CD ₂ rock	572.6(5.0)	gas	CL	30

$A_0=4.716$; $B_0=0.469$; $C_0=0.426$ AB¹¹

\tilde{X}^1A_1		C_{2v}	Structure: MW ^{2,4,5} AB ¹⁰ LF ²⁷			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CD stretch	2158.5	gas	IR	14
			2155m	Ar	IR	17
	2	CD ₂ scissors	1171.8	gas	IR	14
			1167m	N ₂	IR	17
b_1	3	CS stretch	936.13	gas	IR,LS	14,16
			941vw	Ar	IR	6,17
			939wm	N ₂	IR	6
	4	OPLA	781.2	gas	IR	14
b_2			783m	Ar	IR	6,17
	6	CD ₂ rock	784s	N ₂	IR	6
			757.4	gas	IR	14

$A_0=4.883$; $B_0=0.497$; $C_0=0.450$ MW²AB¹⁰LF²⁷

^aBarrier to inversion ≈ 50 .¹⁸

^bFrom fit to combination bands.

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H₂CSe

\tilde{A}^1A_2 C_{2v} Structure: LF⁷
 $T_0 \approx 13555$ gas LF^{5,6}CL⁸ $\tilde{A}-\tilde{X}$ 685–833 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	C=Se stretch	662	gas	LF	6
<i>b</i> ₁	4	OPLA	355T	gas	LF	6

$A^a = 9.015$; $B^a = 0.377$; $C^a = 0.360$ LF⁷

\tilde{a}^3A_2 C_{2v} Structure: LF⁹
 $T_0 = 12162.514$ gas AB¹CL^{3,8}LF^{5,6,9} $\tilde{a}-\tilde{X}$ 658–865 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CH ₂ scissors	1312	gas	LF	5,6
	3	C=Se stretch	704	gas	AB, CL	1,3,5,6,8
<i>b</i> ₁	4	OPLA	297 ^b	gas	AB, LF	1,5,6
	6	HCSe bend	812H	gas	LF	5,6

$A_0 = 9.320$; $B_0 = 0.391$; $C_0 = 0.375$ LF⁹

\tilde{X}^1A_1 C_{2v} Structure: MW^{2,4}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CH ₂ scissors	1460(30)	gas	CL	8
	3	C=Se stretch	860(10)	gas	CL	8
<i>b</i> ₁	4	OPLA	906(10)	gas	CL	8
	6	HCSe bend	914(20)	gas	CL	8

$A_0 = 9.690$; $B_0 = 0.414$; $C_0 = 0.396$ MW^{2,4}LF⁷

D₂CSe

\tilde{A}^1A_2 C_s
 $T_0 = 13631.4$ gas LF⁶ $\tilde{A}-\tilde{X}$ 671–734 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	C=Se stretch	619	gas	LF	6

$A_0 = 4.711$; $B_0 = 0.316$; $C_0 = 0.296$ LF⁷

\tilde{a}^3A_2 C_{2v}
 $T_0 = 12258.060$ gas LF^{6,9} $\tilde{a}-\tilde{X}$ 661–815 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CD ₂ scissors	996	gas	LF	6
	3	C=Se stretch	667	gas	LF	6
<i>b</i> ₁	4	OPLA	208 ^b	gas	LF	6
<i>b</i> ₂	6	DCSe bend	563H	gas	LF	6

$A_0 = 4.688$; $B_0 = 0.327$; $C_0 = 0.306$ LF⁹

\tilde{X}^1A_1 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	C=Se stretch	789T	gas	LF	6

$A_0 = 4.865$; $B_0 = 0.344$; $C_0 = 0.321$ MW⁴LF^{7,9}

^av₄ = 1.

^bFrom fit to double minimum potential. Barrier to inversion 13.1 for H₂CSe. 16.2 for D₂CSe.⁶

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H₂SiO

\tilde{X} C_{2v} Structure: MW⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Si=O stretch	1202	Ar	IR	1,2
		SiH ₂ deform.	697	Ar	IR	2

$A_0 = 5.559$; $B_0 = 0.623$; $C_0 = 0.558$ MW³

D₂SiO

\tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Si=O stretch	1189	Ar	IR	1,2
		SiD ₂ deform.	533	Ar	IR	2

$A_0 = 2.789$; $B_0 = 0.555$; $C_0 = 0.461$ MW⁴

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t-N₂H₂

$\tilde{C}^1B_u^a$ C_{2h}
 $T_0=67894$ gas AB⁹ $\tilde{C}-\tilde{X}$ 135–147 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	2	Bend	1180	gas	AB	9
	3	NN stretch	1849	gas	AB	9

\tilde{B}^1B_u C_{2h}
 $T_0=57926.5$ gas AB^{2,9} Structure: AB⁹ $\tilde{B}-\tilde{X}$ 150–175 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	1	NH stretch	3092	gas	AB	9
	2	Bend	1180	gas	AB	2,9
	3	NN stretch	1875	gas	AB	2,9

$A_0=15.63$; $B_0=1.32$; $C_0=1.22$ AB⁹

\tilde{A}^1B_g C_{2h}
 $T^e=23896$ gas AB^{4,5,8} $\tilde{A}-\tilde{X}$ 300–440 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	2	Bend	1215(15)	gas	AB	5
	3	NN stretch	1550(20)	gas	AB	5

\tilde{X}^1A_g C_{2h} Structure: IR^{2,6}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	1	NH stretch	3128	N ₂	Ra	3
	2	NH bend	1583	N ₂	Ra	3
	3	N=N stretch	1529	N ₂	Ra	3
a_u	4	Torsion	1288.65	gas	IR	10,11
			1283	Ar	IR	7
			1286	N ₂	IR,Ra	1,3,7
b_u	5	NH stretch	3120.29	gas	IR	2,6,10,11
			3118	Ar	IR	7
			3137	N ₂	IR	7
6	NH bend	1316.41	gas	IR	10,11	
		1313	Ar	IR	7	
		1321	N ₂	IR	3,7	

$A_0=10.001$; $B_0=1.304$; $C_0=1.150$ IR^{6,10,11}

t-N₂D₂

\tilde{B}^1B_u C_{2h}
 $T_0=58086^d$ gas AB² $\tilde{B}-\tilde{X}$ 159–172 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	2	Bend	950	gas	AB	2

\tilde{A}^1B_g C_{2h}
 $\tilde{A}-\tilde{X}$ 320–430 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	2	Bend	910(10)	gas	AB	5
	3	NN stretch	1440(20)	gas	AB	5

\tilde{X}^1A_g C_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	2	ND bend	1215	N ₂	Ra	3
	3	N=N stretch	1539	N ₂	Ra	3
	4	Torsion	948.30	gas	IR	12
b_u	5	ND stretch	946	N ₂	IR	1,3,7
			2315.05	gas	IR	6,12
	6	ND bend	2308	N ₂	IR	7
			968.70	gas	IR	12
			972	N ₂	IR	7

$A_0=6.025$; $B_0=1.089$; $C_0=0.920$ IR^{6,12}

^a4pπ Rydberg transition.

^b3pπ Rydberg transition.

^c5₀¹ vibronic band origin.⁸

^d1-0 subband origin.

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H₂CCI

\tilde{X}		C _{2v}	Structure: ESR ³ MW ⁴				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
a ₁	2	CH ₂ scissors	1391wm	Ar	IR	2	
	3	CCl stretch	827s	Ar	IR	1,2	
b ₁	4	OPLA	402s	Ar	IR	1,2	

 $A_0=9.152(3); B_0=0.532; C_0=0.502$ MW⁴LMR⁵**D₂CCI**

\tilde{X}		C _{2v}	Structure: ESR ³ MW ⁴				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
a ₁	2	CD ₂ scissors	1045m	Ar	IR	1,2	
	3	CCl stretch	788m	Ar	IR	1,2	
b ₁	4	OPLA	291m	Ar	IR	1,2	

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H₂CBr

\tilde{X}		C _{2v}	Structure: ESR ³ MW ⁴				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
		CH ₂ scissors	1355.7s	Ar	IR	1,2	
		CH ₂ rock	953w	Ar	IR	1	
		CBr stretch	693.4s	Ar	IR	1,2	
		Umbrella	368vs	Ar	IR	1,2	

 $B+C=0.744(8)$ LMR³**D₂CBr**

\tilde{X}		C _{2v}	Structure: ESR ³ MW ⁴				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
		CD ₂ scissors	1016.4s	Ar	IR	1,2	
		CD ₂ rock	708w	Ar	IR	1	
		CBr stretch	656.6m	Ar	IR	1,2	
		Umbrella	263vs	Ar	IR	1,2	

 $B+C=0.621(15)$ LMR³**References**

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H₂PO

\tilde{X}		C _s	Structure: MW ²				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
		PH stretch	2275.7	Ar	IR	1	
			2273.8				
		P=O stretch	1153.1	Ar	IR	1	
			1151.7				
			1147.7				
		PH ₂ rock	833.2	Ar	IR	1	
		PH ₂ wag	791.6	Ar	IR	1	

 $A_0=5.208; B_0=0.634; C_0=0.594$ MW²**D₂PO**

\tilde{X}		C _s	Structure: MW ²				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
		PD stretch	1648.2	Ar	IR	1	
		P=O stretch	1147.3	Ar	IR	1	
			1141.7				
			1137.7				
		PD ₂ rock	628.4	Ar	IR	1	
		PD ₂ wag	602.9	Ar	IR	1	

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HPSH

\tilde{X}		C _s	Structure: MW ²				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
		PH stretch	2187.0T	Ar	IR	1	
			461.2T	Ar	IR	1	

Reference

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H₂PF

\tilde{X}		C _s	Structure: MW ⁵				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
a'	1	PH ₂ s-stretch	2283.12	gas	IR	2-4	
			2304	Ar	IR	1	
	2	PH ₂ deform.	1109.24	gas	IR	2,3	
			1090	Ar	IR	1	
	3	PH ₂ deform.	976.50	gas	IR	3	
			934	Ar	IR	1	
	4	PF stretch	807.36	gas	IR	2,3	
			795vs	Ar	IR	1	
a''	5	PH ₂ a-stretch	2286.97	gas	IR	2-4	
			2310	Ar	IR	1	

 $A_0=4.364; B_0=0.505; C_0=0.499$ IR^{3,4}MW⁵

D₂PF

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	PD ₂ s-stretch	1660	gas	IR	2
			1673	Ar	IR	1
	3	PF stretch	808	gas	IR	2
			798	Ar	IR	1
<i>a''</i>	4	PD ₂ s-deform.	701	Ar	IR	1
	5	PD ₂ a-stretch	1667	gas	IR	2
			1680	Ar	IR	1

 $A_0=2.272; B_0=0.465; C_0=0.453 \text{ MW}^5$ **References**

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H₂PCI

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	PH ₂ s-stretch	2303.44	gas	IR	1-3
	2	PH ₂ deform.	1095.74	gas	IR	1,2
	3	PH ₂ deform.	861.05	gas	IR	1,2
	4	PCI stretch	511.23	gas	IR	1,2
<i>a''</i>	5	PH ₂ a-stretch	2309.68	gas	IR	1-3

 $A_0=4.353; B_0=0.225; C_0=0.224 \text{ IR}^2 \text{MW}^4$ **D₂PCI**

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	PD ₂ s-stretch	1665T	gas	IR	1
	3	PD ₂ deform.	633	gas	IR	1
	4	PCI stretch	510	gas	IR	1
	5	PD ₂ a-stretch	1670T	gas	IR	1

 $A_0=2.256; B_0=0.212; C_0=0.209 \text{ MW}^4$ **References**

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⁴P. Dréan, M. Paplewski, J. Demaison, J. Breidung, W. Thiel, H. Beckers, and H. Bürger, Inorg. Chem. **35**, 7671 (1996).

H₂PBr

\tilde{X}	C _s					
A ₀	B ₀	C ₀	MW ¹			

References

- ¹M. Paplewski, P. Dréan, J. Demaison, H. Beckers, and H. Bürger, Chem. Commun. 2015 (1996).

H₂PI

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.

<i>a'</i>	3	PH ₂ deform.	752.52	gas	IR	1
	4	PI stretch	337.60	gas	IR	1
<i>a''</i>	6	PH ₂ deform.	742.68	gas	IR	1

 $A_0=4.350(45); B_0=0.105; C_0=0.105 \text{ IR,MW}^1$ **Reference**

- ¹H. Beckers, H. Bürger, J. Demaison, P. Dréan, J.-M. Flaud, and M. Paplewski, J. Mol. Spectrosc. **172**, 78 (1995).

HOOHContinuous absorption, 120–300 nm.^{1,2,7,12}

\tilde{X}	C ₂					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i>	1	OH s-stretch	3617.95 ^a	gas	IR,Ra	3,4,10,
			3609.8			17
			3593	Ar	Ra	11
	2	OH s-bend	1393.5	gas	Ra	10
			1385	Ar	Ra	11
	3	OO stretch	877.93 ^a	gas	IR	20
			865.94	gas	Ra,IR	10,20
			869	Ar	Ra	11
	4	Torsion	370.89 ^a	gas	IR	6,17,18,
			254.55			19
<i>b</i>	5	OH a-stretch	372br	Ar	IR	9
			264			
			378vs,br	N ₂	IR	5,9
			3618.84 ^a	gas	IR	3,4,17
			3610.66			
			3597.0 ^b	Ar	IR	9,23
			3587.8 ^b			
			3583.6 ^b	Kr	IR	23
			3574.0 ^b			
			3568.0 ^b	Xe	IR	23
<i>c</i>	6	OH a-bend	3560.0 ^b			
			3587s	N ₂	IR	5,9
			3582s			
			1273.68 ^a	gas	IR,DL	3,16
			1264.58			
			1277.0ms ^b	Ar	IR	9,23
			1270.9vs ^b			
			1273.7 ^b	Kr	IR	23
			1268.7 ^b			
			1270.3 ^b	Xe	IR	23
<i>d</i>	7	OH a-bend	1265.7 ^b			
			1294vs	N ₂	IR	5,9

 $A_0=10.069; B_0=0.874; C_0=0.838 \text{ IR}^{4,17,20} \text{MW}^{8,13-15,22}$

DOOD

	\tilde{X}	C_2				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i>	1	OD s-stretch	2668	gas	Ra	10
			2653.5	Ar	Ra	11
	2	OD s-bend	1029	gas	Ra	10
			1021.5	Ar	Ra	11
	3	OO stretch	867	gas	Ra	10
			871	Ar	Ra	11
	4	Torsion	251	Ar	IR	9
			286br	N ₂	IR	9
	<i>b</i>	OD a-stretch	2661m	gas	IR	3
			2655.7	Ar	IR	9,23
			2651.3			
			2645.3			
			2645.2	Kr	IR	23
			2639	Xe	IR	23
			2633.2			
			2628.4			
			2646	N ₂	IR	9
			947s	gas	IR	3
	6	OD a-bend	951.3vs	Ar	IR	9,23
			949.9	Kr	IR	23
			947.8	Xe	IR	23
			966vs	N ₂	IR	9

^aTransitions to two lowest torsional levels associated with this fundamental are given.

^bRef. 23 attributes the doubling in the matrix absorptions to excitation from the two lowest torsional levels.

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HSSH

 \tilde{X} C₂ Structure: MW, IR^{13,14}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i>	1	SH s-stretch	2555.78	gas	IR	3,5
	2	Deformation	883	gas	IR	3
	3	SS stretch	515.92	gas	IR,MW	3,15
	4	Torsion	417.48	gas	IR	2,3,11,15
	5	SH a-stretch	2558.63	gas	IR	1,3,5,16
	6	Deformation	882.0	gas	IR	1,3

$$A_0 = 4.899; \quad B_0 = 0.233; \quad C_0 = 0.232 \quad \text{MW}^{4.8,9,12} \text{IR}^{10,12}$$

DSSD

 \tilde{X} C₂

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i>	4	Torsion	306	gas	IR	3
	5	SD a-stretch	1863	gas	IR	3
	6	Deformation	646.4	gas	IR	3

$$A_0 = 2.550; \quad B_0 = 0.218; \quad C_0 = 0.218 \quad \text{MW}^{4.6,7,13} \text{IR}^{13}$$

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8.6. Four-Atomic Monohydrides

BeCCH

\tilde{X}	$C_{\infty v}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	2	CC stretch	2018.9	Ar	IR	1
	3	BeC stretch	855.0	Ar	IR	1
II	4	HCC bend	675.7	Ar	IR	1

BeCCD

\tilde{X}	$C_{\infty v}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	2	CC stretch	1873.4	Ar	IR	1
	3	BeC stretch	848.9	Ar	IR	1
II	4	DCC bend	553.8	Ar	IR	1

Reference

¹C. A. Thompson and L. Andrews, J. Am. Chem. Soc. **118**, 10242 (1996).

MgCCH

\tilde{A}	$^2\Pi$	$C_{\infty v}$
$T_0=22807$	gas	LF ²
$\tilde{A}-\tilde{X}$ 415–440 nm		

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+		MgC stretch	552	gas	LF	2

$A=36.4$ gas LF²
 $B_0=0.175$ gas LF²

\tilde{X}	$^2\Sigma^+$	$C_{\infty v}$
Vib.	No.	Approximate type of mode
Σ^+	2	CC stretch
	3	MgC stretch
II	4	HCC bend

$B_0=0.167$ MW¹

MgCCD

\tilde{X}	$^2\Sigma^+$	$C_{\infty v}$
Vib.	No.	Approximate type of mode
Σ^+	2	CC stretch
	3	MgC stretch

Σ^+ 2 CC stretch 1856.4 Ar IR 3
 3 MgC stretch 485.1 Ar IR 3

References

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CaCCH

\tilde{B} $^2\Sigma^+$ $C_{\infty v}$
 $T_0=16600$ T gas LF⁵

\tilde{A}	$^2\Pi$	$C_{\infty v}$	$T_0=15525.393$	gas	LF ^{1-3,5,6}	$\tilde{A}-\tilde{X}$ 640–665 nm
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
II	5	CaCC bend	101.39	gas	LF	6

$A=70.821$ gas LF^{1-3,5}
 $\epsilon_5\omega_5=3.528(14)$ gas LF⁶
 $B_0=0.115$ LF^{2,3,5}

\tilde{X} $^2\Sigma^+$ $C_{\infty v}$ Structure: LF^{2,5}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	3	CaC stretch	399(10)	gas	LF	1
II	5	CaCC bend	102.94	gas	LF	6

$B_0=0.113$ LF^{2,3,5} MW⁴

References

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⁵M. Li and J. A. Coxon, J. Mol. Spectrosc. **176**, 206 (1996).
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SrCCH

\tilde{A} $^2\Pi$ $C_{\infty v}$
 $T_0=14176(10)$ gas LF¹ $\tilde{A}-\tilde{X}$ 685–725 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	3	SrC stretch	354(10)	gas	LF	1
		$A=275(10)$ gas LF ¹				

\tilde{X} $^2\Sigma^+$ $C_{\infty v}$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	3	SrC stretch	343(10)	gas	LF	1
II	5	SrCC bend	70(5)H	gas	LF	1

$B_0=0.083$ MW²

References

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FeCCD

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		FeC stretch	500(100)	gas	PE	1

Reference

- ¹J. Fan and L.-S. Wang, J. Phys. Chem. **98**, 11814 (1994).

FeCCH⁻

Threshold for electron detachment from ground-state FeCCH⁻=11400(400)
gas PE¹

Reference

- ¹J. Fan and L.-S. Wang, J. Phys. Chem. **98**, 11814 (1994).

HBNB

 \tilde{X} C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	BH stretch	2805.6	Ar	IR	1,2
	2	BNB a-stretch	1846.3	Ar	IR	1,2
	3	BNB s-stretch	1089.3	Ar	IR	1,2
II	4	HBN deform.	746.2	Ar	IR	1,2

DBNB

 \tilde{X} C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	BD stretch	2218.7	Ar	IR	1,2
	2	BNB a-stretch	1717.6	Ar	IR	1,2
	3	BNB s-stretch	1062.4	Ar	IR	1,2
II	4	DBN deform.	596.4	Ar	IR	1

References

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HBCC

 $\tilde{X}^1\Sigma^+$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C≡C stretch	1995.2	Ar	IR	1

DBCC

 $\tilde{X}^1\Sigma^+$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		BD stretch	2173.3	Ar	IR	1
		C≡C stretch	1907.7	Ar	IR	1

Reference

- ¹L. Andrews, P. Hassanzadeh, J. M. L. Martin, and P. R. Taylor, J. Phys. Chem. **97**, 5839 (1993).

HCCB

 $\tilde{X}^1\Sigma^+$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C≡C stretch	2039.3	Ar	IR	1

DCCB

 $\tilde{X}^1\Sigma^+$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C≡C stretch	1918.8	Ar	IR	1

Reference

- ¹L. Andrews, P. Hassanzadeh, J. M. L. Martin, and P. R. Taylor, J. Phys. Chem. **97**, 5839 (1993).

cyc-HBC₂ \tilde{X}^1A_1 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	3	BC ₂ s-stretch	1122.7	Ar	IR	1,2

cyc-DBC₂ \tilde{X}^1A_1 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	3	BC ₂ s-stretch	1075.7	Ar	IR	1,2

References

- ¹J. M. L. Martin, P. R. Taylor, P. Hassanzadeh, and L. Andrews, J. Am. Chem. Soc. **115**, 2510 (1993).
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HCCAI

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CC stretch	1976.4	Ar	IR	1,2
		HCC deform.	683.5	Ar	IR	1,2
		AIC stretch	512.8	Ar	IR	1,2

DCCAI

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CC stretch	1860.8	Ar	IR	1
		DCC deform.	505.2	Ar	IR	1
		AIC stretch	456.0	Ar	IR	1

References

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cyc-HC₃

\tilde{X}^2B_2 C_{2v} Structure: MW^{2,4}
 $A_0=1.486$; $B_0=1.135$; $C_0=0.640$ MW^{1,3,4}

cyc-DC₃

\tilde{X}^2B_2 C_{2v}
 $A_0=1.485$; $B_0=0.931$; $C_0=0.570$ MW^{2,3}

References

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⁴S. Yamamoto and S. Saito, J. Chem. Phys. **101**, 5484 (1994).

HC₃

$\tilde{X}^2\Pi_{1/2}$ C_{∞v} Structure: MW⁷

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CH stretch	3238.0w	Ar	IR	5,6
	2	C ₃ a-stretch	1832.6m	Ar	IR	1,5,6
			1824.7s			
Π	3	C ₃ s-stretch	1167br	Ar	IR	5,6
			1159.8w			
II	4	HCC bend	28 ^a	gas	MW	4

$$\begin{aligned}A &= 14.44 \text{ gas } \text{MW}^{2-4} \\B_0 &= 0.373 \text{ MW}^{2-4}\end{aligned}$$

DC₃

$\tilde{X}^2\Pi_{1/2}$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CD stretch	2424.0wmT	Ar	IR	5,6
	2	C ₃ a-stretch	1778.8sh	Ar	IR	1,5,6
			1771.5m			
Σ^+	3	C ₃ s-stretch	1148br	Ar	IR	5,6
			1140.4w			

$$\begin{aligned}A &= 12.53 \text{ gas } \text{MW}^4 \\B_0 &= 0.337 \text{ MW}^4\end{aligned}$$

^a 2Σ^μ component.

References

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HCCSi

$\tilde{X}^2\Pi$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	2	CC stretch	1989.8	Ar	IR	1
	3	SiC stretch	636.0T	Ar	IR	1

DCCSi

$\tilde{X}^2\Pi$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	2	CC stretch	1874.9	Ar	IR	1
	3	SiC stretch	622.4T	Ar	IR	1

Reference

- ¹D. Han, C. M. L. Rittby, and W. R. M. Graham, J. Chem. Phys. **106**, 6222 (1997).

HCCN

 $^3\Sigma^-$?

In the gas phase, a prominent absorption band system beginning at 340 nm has been attributed^{2,3} to HCCN, but has not been analyzed. An absorption band system assigned to HCCN was observed in an argon matrix between 240 and 340 nm, with band separations of approximately 1050.⁴

$\tilde{X} \ ^3\Sigma^-$		$C_{\infty v}$ ^a	Structure: ESR ¹ MW ⁵⁻⁷			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CH stretch	3246.66	gas	CC	8
			3229s	Ar	IR	4
	2	CCN a-stretch	1735s	Ar	IR	4
	3	CCN s-stretch	1178m	Ar	IR	4
	4	H deform.	383(20)	gas	CC,MW	8,9
			369.5 ^b	Ar	IR	4
	5	Skel. deform.	187(20)	gas	CC	8

$$B_0 = 0.366 \text{ MW}^{5,7,9}$$

DCCN

$\tilde{X} \ ^3\Sigma^-$		$C_{\infty v}$ ^a	Structure: MW ^{1,4}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1	CD stretch	2424ms	Ar	IR	4
	2	CCN a-stretch	1730s	Ar	IR	4
	3	CCN s-stretch	1127w	Ar	IR	4
	4	CCN bend	367(15)	gas	MW	9
			405wmT	Ar	IR	4
	5	D deform.	90(15)	gas	MW	9

$$B_0 = 0.330 \text{ MW}^{6,9}$$

^aQuasi-linear,^{6,7} with barrier to linearity⁹ of ~235.

^bAssignment proposed by Ref. 9. A prominent argon-matrix absorption of HCCN at 458 and of DCCN at 317.5, with counterparts at 435(20) and 311(20), respectively, in the millimeter-wave spectra, may result from $l=0$ transitions associated with overtones or combination bands.

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HCCO

$\tilde{B} \ ^2\Pi$ $C_{\infty v}$
 $T_0 = 33423.92(2)$ gas PF^{6,7}LF⁸ $\tilde{B}-\tilde{X}$ 263–300 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	3	CCO s-stretch	1037	gas	PF,LF	7,8
Π	4	Deformation	416T	gas	PF	7
	5	Deformation	365T	gas	PF	7

$$B_0 = 0.324 \text{ PF}^7\text{LF}^8$$

Analysis of the submillimeter-wave spectrum¹ indicates that HCCO possesses a low-lying excited electronic state which, together with the ground state, is derived from a Π state by Renner-Teller interaction.

$\tilde{X} \ ^2A''$ C_s Structure: MW^{1,4}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	CCO a-stretch	2022.64	gas	DL	3
			2023.7	Ne	IR	5
			2019.5	Ar	IR	2
a''	6	CCH bend	494	gas	LF	8

$$A_0 = 41.5(1.5); B_0 = 0.363; C_0 = 0.359 \text{ MW}^{1,4}$$

DCCO

$\tilde{X} \ ^2A''$ C_s

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2	CCO a-stretch	1995.2	Ne	IR	5
			1989.9	Ar	IR	2

$$A_0 = 21.75(12); B_0 = 0.331; C_0 = 0.325 \text{ MW}^1$$

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HCCS

$A \ ^2\Pi$ $C_{\infty v}$
 $T_0 = 24299.690(6)$ gas AB^{1,2}EM³LF^{4,6} $\tilde{A}-\tilde{X}$ 377–452 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	2	CC stretch	1843T	gas	AB	2
	3	CS stretch	740	gas	AB	1,2
	5	CCS bend	328H	gas	AB,EM	2,3

$$B_0 = 0.174 \text{ AB}^2$$

$\tilde{X}^2\Pi$		$C_{\infty v}$	Structure: MW ⁵			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	2	CC stretch	2189T	gas	EM	3
	3	CS stretch	782	gas	EM	3
Π	5	CCS bend	411H ^a	gas	EM	3

$A \approx -185^a$ gas MW⁵
 $B_0 = 0.196$ AB²MW⁵

DCCS

$\tilde{A}^2\Pi$		$C_{\infty v}$	$\tilde{A}-\tilde{X}$ 376–420 nm			
$T_0 = 24359$	gas	AB ² LF ⁴	cm^{-1}	Med.	Type meas.	Refs.
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	2	CC stretch	1718T	gas	AB	2
	3	CS stretch	725	gas	AB	2

^aRef. 7 estimated that $A = -270$, $\omega_4 = 565$, and $\omega_5 = 380$ by considering the relative intensities of microwave lines arising from vibrationally excited HCCS.

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HNCN

\tilde{B}^2A'		C_s	Structure: AB ¹			
$T_0 = 28994.1$	gas	AB ¹ LF ²	$\tilde{B}-\tilde{X}$ 344–365 nm			
$\tau_0 = 20(5)$	ns	gas LF ²	cm^{-1}	Med.	Type meas.	Refs.
$A_0 = 22.30$	$B_0 = 0.376$	$C_0 = 0.369$	AB ¹ LF ²			

\tilde{X}^2A''		C_s	Structure: AB ¹			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	NH stretch	3297.5m	Ar	IR	4
	2	NCN a-stretch	1843s	Ar	IR	4
	3		1146vs	Ar	IR	4
a''	6	NCN deform.	440T	gas	LF	2

$A_0 = 21.18(11)$; $B_0 = 0.370$; $C_0 = 0.363$ AB¹LF²MW³

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HCNN

\tilde{A}		$T_0 = 30500$	gas	AB ^{1,2,4,5}	$\tilde{A}-\tilde{X}$ 289–328 nm	
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1048	gas	AB	4

\tilde{X}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CH stretch	3233.7wm	Ar	IR	3,6
			3229m	N ₂	IR	3
		CNN a-stretch	1786.6s	Ar	IR	3,6
			1784s	Kr	IR	3
			1800s	N ₂	IR	3
		H deform.	862.2vs	Ar	IR	3,6
			860vs	Kr	IR	3
			871m	N ₂	IR	3
			863m			

DCNN

\tilde{X}		T_0	gas	AB ^{1,2,4,5}	$\tilde{B}-\tilde{X}$ 344–365 nm	
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CNN a-stretch	1771vs	Ar	IR	3
		D deform.	725vs	Ar	IR	3

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HNCO

Between 132.5 and 120 nm, continuous absorption, with some diffuse bands.⁵

Relatively sharp absorption bands at 72940, 73910, and 74680 (137.1, 135.3, and 133.9 nm), which may be contributed by a Rydberg transition.⁵

Between 147 and 163 nm, diffuse bands are superposed on a continuum, with maximum near 157 nm.⁵

Continuous absorption between 163 and 185 nm, with a maximum near 166 nm.⁵

Continuous absorption has its onset near 41000 (244 nm), and extends beyond 200 nm.⁴

\tilde{A}^1A'' C_s
gas AB⁴ $\tilde{A}-\tilde{X}$ 228–282 nm

Complicated group of progressions, possibly resulting from the occurrence of *cis*- and *trans*- rotamers in the excited state.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>		NCO bend	555T	gas	AB	4

$A \approx 4.37; B \approx 0.388; C \approx 0.357$ AB⁴

\tilde{X}^1A' C_s Structure: MW⁷

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	NH stretch	3538.25s	gas	IR,Ra	1,2,12 14,15
			3516.8wm	Ar	IR	12
			3505.7wm			
2		NCO a-stretch	2268.89vs	gas	IR	1,2,9,10
			2259.0vs	Ar	IR	12,13
3		NCO s-stretch	1327vw	gas	IR	1,2
4		HNC, NCO bend	776.62wm	gas	IR	6
			769.8wm	Ar	IR	12,13
5		HNC, NCO bend	577.35w	gas	IR	1,2,6
			573.7wm	Ar	IR	12,13
<i>a''</i>	6	Torsion	656.29	gas	IR	6,11

$A_0=30.638; B_0=0.369; C_0=0.364$ MW⁷IR¹⁴

DNCO

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	ND stretch	2637.20	gas	IR	8
			2606.9m	Ar	IR	12
2		NCO a-stretch	2235vs	gas	IR	3
			2231.0vs	Ar	IR	12
3		NCO s-stretch	1310	gas	IR	3
4		DNC, NCO bend	578.6w	Ar	IR	12
5		DNC, NCO bend	475.4w	Ar	IR	12
<i>a''</i>	6	Torsion	602.9	gas	IR	3

$A_0=17.09; B_0=0.344; C_0=0.336$ MW⁷

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HOCS+

\tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1		OH stretch	3435.16	gas	LD	1

$A_0=26.11; B_0=0.192; C_0=0.190$ LD¹MW²

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t-HOCO

\tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OH stretch	3635.70	gas	LD	7
			3631.7wm	Ne	IR	9
			3628.1wm			
			3602.9	Ar	IR	2
			3456	CO	IR	1
2		C=O stretch	1852.57	gas	DI	4
			1848.0s	Ne	IR	9
			1843.6	Ar	IR	2
			1833	CO	IR	1
3		HOC bend	1210.4m	Ne	IR	9
			1211.2	Ar	IR	2
			1261	CO	IR	1
4		C-O stretch	1050.4wm	Ne	IR	9
			1064.6	Ar	IR	2
			1077	CO	IR	1
5		OCO bend	615	CO	IR	1
<i>a''</i>	6	Torsion	508.1wm	Ne	IR	9
			515	Ar	IR	2

$A_0=5.596; B_0=0.381; C_0=0.356$ MW³LMR⁵

t-DOCO

\tilde{X}		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OD stretch	2684.10	gas	LD	6
			2680.7wm	Ne	IR	9
			2678.1wm			
	2	C=O stretch	2558	CO	IR	1
			1851.65	gas	DL	4
			1848.0m	Ne	IR	9
			1846.2s			
	3	Mixed	1841.7	Ar	IR	2
			1825	CO	IR	1
			1082.5m	Ne	IR	9
			1092.6	Ar	IR	2
	5	OCO bend	1117	CO	IR	1
			610m	CO	IR	1
a''	6	Torsion	472wm	CO	IR	1

$A_0 = 5.160$; $R_0 = 0.356$; $C_0 = 0.333$ MW³LD⁶LMR⁸

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HCO₂

\tilde{B}		C_s				
$T_0 = 4320$	gas	PE ¹	Structure: PE ¹			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	3	CO ₂ bend	700(50)	gas	PE	1
\tilde{A}					C_s	
$T_0 = 220$	gas	PE ¹	Structure: PE ¹			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	3	CO ₂ bend	550T	gas	PE	1
\tilde{X}					C_s	
$T_0 = 570T^b$	gas	PE ¹	Structure: PE ¹			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	3	CO ₂ bend	570T ^b	gas	PE	1

DCO₂

\tilde{B}		C_{2v}	
$T_0 = 4110$	gas	PE ¹	
Vib.	No.	Approximate type of mode	cm^{-1}

\tilde{A} $^2A_1^a$ C_{2v}
 $T_0 = 65T$ gas PE¹

\tilde{X} $^2B_2^a$ C_{2v}

^aStrong vibronic interaction between the \tilde{X} and \tilde{A} states. Differences in the zero-point vibrational energies result in a reversal of the symmetries of these two states for DCO₂.

^bLarge anharmonicity.

Reference

- ¹E. H. Kim, S. E. Bradforth, D. W. Arnold, R. B. Metz, and D. M. Neumark, J. Chem. Phys. **103**, 7801 (1995).

HFCS⁺

\tilde{B}		C_s				
$T_0 = 28720(120)$	gas	PE ¹				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'			1210(80)	gas	PE	1
			980(80)	gas	PE	1

\tilde{A} $^2A''$ C_s
 $T_0 = 10810(180)$ gas PE¹

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'			1450(80)	gas	PE	1
			750(80)	gas	PE	1

\tilde{X} $^2A'$ C_s

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'			1360(60)	gas	PE	1

Reference

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HBF₂

\tilde{X}	C _{2v}	Structure: IR ^{2,3} MW ⁵				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	BH stretch	2620.8(5)	gas	IR	1-4
			2644	N ₂	IR	6
	2	BF ₂ s-stretch	1164.2(5)	gas	IR	1-4
			1156	N ₂	IR	6
	3	BF ₂ scissors	541.5(5)	gas	IR	1-4
			542	N ₂	IR	6
<i>b</i> ₁	4	OPLA	923.5(5)	gas	IR	1-4
			918	N ₂	IR	6
<i>b</i> ₂	5	BF ₂ a-stretch	1402vs	gas	IR	1,2,4
			1385.1	N ₂	IR	6

 $A_0=2.485; B_0=0.350; C_0=0.306$ MW^{5,7}**DBF₂**

\tilde{X}	C _{2v}	Structure: IR ^{2,3} MW ⁵				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	BD stretch	1960.0(5)	gas	IR	2-4
			1980	N ₂	IR	6
	2	BF ₂ s-stretch	1126.4(5)	gas	IR	2-4
			1121.3	N ₂	IR	6
	3	BF ₂ scissors	540	gas	IR	2,4
	4	OPLA	785	gas	IR	2,4
<i>b</i> ₁			772.7	N ₂	IR	6
	5	BF ₂ a-stretch	1392	gas	IR	2,4
			1374.5	N ₂	IR	6

 $A_0=1.764; B_0=0.350; C_0=0.292$ MW⁵**References**

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HBCl₂

\tilde{X}	C _{2v}	Structure: IR ⁴ MW ⁸				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	BH stretch	2616.9w	gas	IR	1-4
			2625	Ar	IR	6
	2	BCl ₂ s-stretch	740T	gas	IR	5
	4	OPLA	784T	gas	IR	5
			788	Ar	IR	6
	5	HBCl bend	1089s	gas	IR	1-3,5
<i>b</i> ₂			1082.5	Ar	IR	6,7
	6	BCl ₂ a-stretch	892s	gas	IR	1-3,5
			886.3	Ar	IR	6,7

 $A_0=1.565; B_0=0.106; C_0=0.099$ IR⁴MW⁸**DBCi₂**

\tilde{X}	C _{2v}	Structure: IR ⁴ MW ⁸				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	BD stretch	1969	gas	IR	4
	4	OPLA	645T	gas	IR	5
	5	BCl ₂ a-stretch	1005T	gas	IR	5
			970	Ar	IR	6

 $A_0=1.173; B_0=0.106; C_0=0.097$ IR⁴MW⁸**References**

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HBBr₂

\tilde{X}	C _{2v}	Structure: IR ¹				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	BH stretch	2622.4	gas	IR	1-3
	2	BBR ₂ s-stretch	595	gas	IR	2
	4	OPLA	731	gas	IR	2
	5	BBR ₂ a-stretch	1036	gas	IR	2,3
			1032	Ar	IR	4
	6	HBBr deform.	772	gas	IR	2,3
			768	Ar	IR	4

 $A_0=1.256; B_0=0.041; C_0\approx 0.039$ gas IR¹**DBBBr₂**

\tilde{X}	C _{2v}	Structure: IR ¹				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	BD stretch	1960.1	gas	IR	1-3
	2	BBR ₂ s-stretch	585	gas	IR	2
	4	OPLA	625vw	gas	IR	2
	5	BBR ₂ a-stretch	870	gas	IR	2,3
	6	DBBBr deform.	656	gas	IR	2,3

 $A_0=0.822; B_0=0.041; C_0\approx 0.039$ gas IR¹**References**

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HCO₂⁻

Threshold for electron detachment from ground-state HCO₂⁻ = 28220(120)
gas PE¹

\bar{X}		C _{2v}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a ₁	3	CO ₂ bend	730	gas	PE 1

DCO₂⁻

Threshold for electron detachment from ground-state DCO₂⁻ = 28320(120)
gas PE¹

\bar{X}		C _{2v}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a ₁	3	CO ₂ bend	730	gas	PE 1

Reference

¹E. H. Kim, S. E. Bradford, D. W. Arnold, R. B. Metz, and D. M. Neumark, J. Chem. Phys. **103**, 7801 (1995).

HCOCl

\bar{A} 1A''		C _s			
$T_0=32754.7$		gas AB ^{5,6}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	2	CO stretch	1153.8	gas	AB 5
	4	CCl stretch	633.6	gas	AB 5
	5	CICO deform.	306.3	gas	AB 5
a''	6	Umbrella (OPLA)	779.5	gas	AB 5

Barrier to planarity = 1608.8 gas AB⁵

\bar{X} 1A'		C _s			
		Structure: MW ^{2,4}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	1	CH stretch	2929.18w	gas	IR 1,3,6
	2	CO stretch	1784.14vs	gas	IR 1,3,6,8
	3	CH bend	1307.21m	gas	IR 1,3,6,7
	4	CCl stretch	738.80vs	gas	IR 1,3,6,9
	5	CCl bend	457.02vw	gas	IR 1,6,9
a''	6	OPLA	932.15vw	gas	IR 1,3,6,11

$A_0=2.601$; $B_0=0.205$; $C_0=0.190$ MW^{2,4}IR⁷

DCOCl

\bar{A} 1A''		C _s	
$T_0=32775.3$		gas	AB ⁵
$\bar{A}-\bar{X}$ 283–313 nm			

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	CO stretch	1092.0	gas	AB 5	
	4	CCl stretch	633.4	gas	AB 5	
	5	CICO deform.	303.1	gas	AB 5	
a''	6	Umbrella (OPLA)	566.5	gas	AB 5	

\bar{X} 1A'		C _s			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	1	CD stretch	2204.3	gas	IR 4,10
	2	CO stretch	1749.1	gas	IR 4,10
	3	CD bend	986.11	gas	IR 4,10
	4	CCl stretch	700.77	gas	IR 4,10,11
	5	CCl bend	455.78	gas	IR 10
a''	6	OPLA	781.50	gas	IR 10,11

$A_0=1.904$; $B_0=0.203$; $C_0=0.184$ MW^{2,4}DL¹⁰

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- D.-L. Joo, D. J. Clouthier, and A. J. Merer, J. Mol. Spectrosc. **174**, 353 (1995).

HCOBr

\bar{X}		C _s			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	1	CH stretch	2911.9	gas	IR 1
	2	CO stretch	1797.9 ^a	gas	IR 1
			1799.5	Ar	IR 2
	3	H deform.	1270.9 ^b	gas	IR 1
	4	CBr stretch	646.3	gas	IR 1
a''	6	OPLA	893.4	gas	IR 1

DCOBr

\tilde{X}	C_s					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CD stretch	2205.4	gas	IR	1
	2	CO stretch	1748.3	gas	IR	1
			1746.2	Ar	IR	2
	3	D deform.	975.5 ^c	gas	IR	1
	4	CBr stretch	605.2	gas	IR	1
	5	CBr deform.	358T	gas	IR	1
a''	6	OPLA	746.8	gas	IR	1

^aUncorrected for Fermi resonance with $2\nu_6$.^bUncorrected for Fermi resonance with $2\nu_4$.^cUncorrected for Fermi resonance with ($\nu_4 + \nu_5$).**References**¹G. Yarwood, H. Niki, and P. D. Maker, *J. Phys. Chem.* **95**, 4773 (1991).²C. Lugez, A. Schriver, L. Schriver-Mazzuoli, E. Lasson, and C. J. Nielsen, *J. Phys. Chem.* **97**, 11617 (1993).**HCOI**

\tilde{X}	C_s					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CH stretch	2930Tw	gas	IR	1
	2	CO stretch	1776.3vs	gas	IR	1
	3	HCO deform.	1247.9wm	gas	IR	1
	4	CI stretch	561m	gas	IR	1

Reference¹I. Barnes, K. H. Becker, and J. Starcke, *Chem. Phys. Lett.* **246**, 594 (1995).**c-HNSO^a**gas AB³ 238–269 nmDiffuse absorption merges into continuum with maximum near 217 nm.³Photolysis in an argon matrix by 254 nm radiation leads to rapid formation of *c*-HOSN.⁵

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	5	NSO bend	285T	gas	AB	3

 $\tilde{\alpha}^3\text{A}'^b$ C_s Weak, unstructured absorption 325–350 nm.³In an argon matrix, converted to *t*-HNSO by irradiation at wavelengths longer than 300 nm.⁶

$\tilde{X}^1\text{A}'$		C_s	Structure: MW ²			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	NH stretch	3308.5	gas	IR	7,9
			3308wm	Ar	IR	3,4
			3303	N_2	IR	4
	2	SO stretch	1257.58s	gas	IR	1,9,10
			1249vs	Ar	IR	3,4
	3	NS stretch	1086.83w	gas	IR	1,9,11
a''			1083s	Ar	IR	3,4
			1094	N_2	IR	4
	4	HNS bend	906.3m	gas	IR	1,9
			900s	Ar	IR	3,4
			923	N_2	IR	4
	5	NSO bend	447.85m	gas	IR	1,9,11
a''			447s	Ar	IR	3,4
			455	N_2	IR	4
	6	Torsion	757.0s	gas	IR	1,9
			755vs	Ar	IR	3,4
			774	N_2	IR	4

 $A_0 = 1.645$; $B_0 = 0.329$; $C_0 = 0.274$ MW²**c-DNSO**gas AB³ 242–264 nm
Diffuse, merges into continuum.

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	5	NSO bend	285T	gas	AB	3
\tilde{X}						C_s
a'	1	ND stretch	2454.8	gas	IR	8,9
			2453	Ar	IR	3,4
	2	SO stretch	1254.4s	gas	IR	1,9
			1245	Ar	IR	3,4
	3	NS stretch	1052.3w	gas	IR	1,9
			1048	Ar	IR	3,4
a''	4	DNS bend	755.0m	gas	IR	1,9
			752	Ar	IR	3,4
	5	NSO bend	400.2m	gas	IR	1,9
			400	Ar	IR	3,4
	6	Torsion	595.7s	gas	IR	1,9
			594	Ar	IR	3,4

 $A_0 = 1.644(3)$; $B_0 = 0.318$; $C_0 = 0.259$ MW²^aStable rotamer.^bTentative assignment.**References**

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¹⁰D.-L. Joo and D. J. Clouthier, J. Chem. Phys. **104**, 8852 (1996).

¹¹D.-L. Joo, H. H. Harjanto, and D. J. Clouthier, J. Mol. Spectrosc. **178**, 78 (1996).

HPSS

\bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		PH stretch	2172.6	Ar	IR	1
		P=S stretch	690.6	Ar	IR	1
		S=S stretch	682.4	Ar	IR	1

DPSS

\bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		P=S stretch	690.3	Ar	IR	1
		S=S stretch	682.3	Ar	IR	1
		DPS deform.	588.1	Ar	IR	1

Reference

¹Z. Mielke and L. Andrews, J. Phys. Chem. **97**, 4313 (1993).

HOClO

In an argon matrix, a dissociative excited state has its maximum near 400 nm (25000).¹

\bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	OH stretch	3527.1vs	Ar	IR	1
	2	HOCl deform.	1176.9wm	Ar	IR	1
	3	ClO stretch	973.9s	Ar	IR	1
	4	OCl stretch	591.5wm	Ar	IR	1

DOCIO

\bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	OD stretch	2605.4vs	Ar	IR	1
	2	ClO stretch	974.7vs	Ar	IR	1
	3	DOCl deform.	864.5wm	Ar	IR	1
	4	OCl stretch	594.5wm	Ar	IR	1

Reference

¹K. Johnsson, A. Engdahl, and B. Nelander, J. Phys. Chem. **100**, 3923 (1996).

HClO₂

In an argon matrix, the onset of photodecomposition is near 290 nm (34500).¹

\bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	HCl stretch	2116wm	Ar	IR	1
	2	ClO ₂ a-stretch	1132.0vs	Ar	IR	1
	5	ClO ₂ s-stretch	969.4m	Ar	IR	1

DClO₂

\bar{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	DCl stretch	1550.4wm	Ar	IR	1
	2	ClO ₂ a-stretch	1115.9vs	Ar	IR	1
	3	ClO ₂ s-stretch	969.3m	Ar	IR	1

Reference

¹K. Johnsson, A. Engdahl, and B. Nelander, J. Phys. Chem. **100**, 3923 (1996).

8.7. Four-Atomic Nonhydrides

Ta₄

In an argon matrix,¹ there is a broad absorption maximum centered at 18800 nm (532 nm).

In an argon matrix,¹ a broad absorption maximum at 13020 (768 nm) has partially resolved high frequency shoulders separated by intervals of ~240.

\bar{X} T_d

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			270.2	Ar	Ra	1
			185.1	Ar	Ra	1
			130.6(2)	Ar	Ra	1

Reference

¹H. Wang, R. Craig, H. Haouari, J.-G. Dong, Z. Hu, A. Vivoni, J. R. Lombardi, and D. M. Lindsay, J. Chem. Phys. **103**, 3289 (1995).

BeBeCO

\bar{X} C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CO stretch	1943.6	Ar	IR	1
			483.3	Ar	IR	1

Reference

¹L. Andrews, T. J. Tague, Jr., G. P. Kushto, and R. D. Davy, Inorg. Chem. **34**, 2952 (1995).

BeBeNN

\tilde{X}		$C_{\infty v}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
1		NN stretch	1876.1	Ar	IR	1

Reference

¹C. A. Thompson, L. Andrews, and R. D. Davy, J. Phys. Chem. **99**, 7913 (1995).

BeNNBe

\tilde{X}						
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BeN stretch	1111.7	Ar	IR	1

Reference

¹C. A. Thompson, L. Andrews, and R. D. Davy, J. Phys. Chem. **99**, 7913 (1995).

(BeN)₂

\tilde{X}		C_{2v} ?				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BeN stretch	787.7	Ar	IR	1

Reference

¹C. A. Thompson, L. Andrews, and R. D. Davy, J. Phys. Chem. **99**, 7913 (1995).

FeCCC

\tilde{A}		$C_{\infty v}$				
$T_0=4500(700)$		gas	PE ¹			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		Fe=C stretch	700(150)	gas	PE	1

Reference

¹J. Fan, L. Lou, and L.-S. Wang, J. Chem. Phys. **102**, 2701 (1995).

AICCAI

\tilde{X}		$D_{\infty h}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_u^+	3	AIC a-stretch	605.1	Ar	IR	1

Reference

¹G. V. Chertihin, L. Andrews, and P. R. Taylor, J. Am. Chem. Soc. **116**, 3513 (1994).

FeCCC⁻

Threshold for electron detachment from ground-state FeCCC⁻ = 13600(600) gas PE¹

Reference

¹J. Fan, L. Lou, and L.-S. Wang, J. Chem. Phys. **102**, 2701 (1995).

cyc-(BeO)₂

\tilde{X}		D_{2h}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1131.2	Ar	IR	1
			1120.8	Kr	IR	1
			1104.4	Xe	IR	1
			1118.0	N ₂	IR	2
			866.3	Ar	IR	1
			857.2	Kr	IR	1
			845.4	Xe	IR	1
			892.4	N ₂	IR	2
			522.4	Ar	IR	1
			516.9	Kr	IR	1
			514.4	Xe	IR	1
			539.1	N ₂	IR	2

References

¹C. A. Thompson and L. Andrews, J. Chem. Phys. **100**, 8689 (1994).

²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

BeOBeO

\tilde{X}		$D_{\infty h}$				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1572.9T	Ar	IR	1,2
			1552Tbr	N ₂	IR	2

References

¹C. A. Thompson and L. Andrews, J. Chem. Phys. **100**, 8689 (1994).

²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

MgOMgO

\tilde{X}	$C_{\infty v}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			971.7A ^a	Ar	IR	1
			943.1B	Ar	IR	1
			591.7A	Ar	IR	1
			567.2B	Ar	IR	1

^aIt is uncertain whether the absorptions labelled A and B are contributed by two different MgOMgO structures (A and B) or whether they correspond to the same molecule trapped in two different matrix sites.

Reference

¹L. Andrews and J. T. Yustein, *J. Phys. Chem.* **97**, 12700 (1993).

cyc-(CaO)₂

\tilde{X}	D_{2h}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_{2u}	5		584.6	Ar	IR	1
			559.5	N ₂	IR	2
b_{3u}	6		516.3	Ar	IR	1
			497.0	N ₂	IR	2

References

¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).

²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 10088 (1996).

CaOCaO

\tilde{X}	$C_{\infty v}$					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			610.7	Ar	IR	1

Reference

¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).

cyc-(SrO)₂

\tilde{X}	D_{2h}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_{2u}	5		530.1	Ar	IR	1
			502.7	N ₂	IR	2
b_{3u}	6		441.3	Ar	IR	1
			423T	N ₂	IR	2

References

¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).

²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 10088 (1996).

cyc-(BaO)₂

\tilde{X}	D_{2h}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_{2u}	5		501.2	Ar	IR	1,2
b_{3u}	6		401.4	Ar	IR	1,2

References

¹D. M. Thomas and L. Andrews, *J. Mol. Spectrosc.* **50**, 220 (1974).

²L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).

BaOBaO

\tilde{X}	D_{2h}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			460.0	Ar	IR	1,2

References

¹D. M. Thomas and L. Andrews, *J. Mol. Spectrosc.* **50**, 220 (1974).

²L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).

cyc-(TiO)₂

\tilde{X}	D_{2h}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			704.2	Ar	IR	1
			680.1	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).

cyc-(ZrO)₂

\tilde{X}	D_{2h}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			652.4	Ar	IR	1
			593.1	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).

cyc-(HfO)₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			651.3	Ar	IR	1
			583.3	Ar	IR	1

Reference¹G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).**(FeO)₂** \tilde{A} $T_0=3550(400)$ gas PE⁶ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	5		670(70)	gas	PE	6
			517.4	Ar	IR	1-4
			535.5	N ₂	IR	5

References

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cyc-(NiO)₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			650.2	Ar	IR	1
			481.6	Ar	IR	1

Reference¹A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, *J. Phys. Chem. A* **101**, 3109 (1997).**NiONiO** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			973.9	Ar	IR	1

Reference¹A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, *J. Phys. Chem. A* **101**, 3109 (1997).**Cu₂O₂** \tilde{C} $T_0=6780(400)$ gas PE¹ \tilde{B} $T_0=5320(400)$ gas PE¹ \tilde{A} $T_0=3630(400)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			650(30)	gas	PE	1

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			630(30)	gas	PE	1

Reference¹L.-S. Wang, H. Wu, S. R. Desai, and L. Lou, *Phys. Rev. B* **53**, 8028 (1996).**ZnOZnO** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			882br	Ar	IR	1
			877br			
			877br	N ₂	IR	1
			848.5	Ar	IR	1

Reference¹G. V. Chertihin and L. Andrews, *J. Chem. Phys.* **106**, 3457 (1997).**CdOCdO** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			682.0	Ar	IR	1
			675.5			

Reference¹G. V. Chertihin and L. Andrews, *J. Chem. Phys.* **106**, 3457 (1997).

C₄

$^3\Sigma_u^-$ D_{∞h}
 $T_0=26323(15)$ Ne AB¹¹ $^3\Sigma_u^- - \tilde{X}$ 325–380 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1822(20)	Ne	AB	11
	2	C–C stretch	903(20)	Ne	AB	11
Π_g	4	Bend	370(20)H	Ne	AB	11
Π_u	5	Bend	175(20)H	Ne	AB	11

$^1\Delta_g$ D_{∞h}
 $T_0=2640(50)$ gas PE⁶

\tilde{X} $^3\Sigma_g^-$		D _{∞h} ^a	Structure: ESR ^{2,5} DL ^{3,7}			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	2032(50)	gas	PE	6
Σ_u^+	3	Asym. stretch	1548.61	gas	DL	3
			1547.0	Ne	IR	8,10
			1543.4	Ar	IR	1
			1539.5	Kr	IR	12
Π_g	4	Bend	352(15)	gas	PE,DL	6,9
Π_u	5	Bend	160(4) ^b	gas	DL	7
			172.4	Ar	IR	4
			170.4			

$B_0=0.166$ DL^{4,8}

^aESR measurements^{2,5} suggest that C₄ may deviate by a few degrees from the linear structure.

^bEstimated from *I*-doubling parameter for ν₅.

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Si₂C₂

\tilde{X}		D _{2h}		
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.
b_{1u}	3		982.9vs	Ar
b_{2u}	4		382.2wm	Ar

Reference

¹J. D. Presilla-Márquez, S. C. Gay, C. M. L. Rittby, and W. R. M. Graham, J. Chem. Phys. **102**, 6354 (1995).

Si₄

$^1B_{1u}$ D_{2h}
 $T_0=23635T^a$ gas EM³
 $21432(9)$ Ne AB^{1,7} $^1B_{1u} - \tilde{X}$ 418–448 nm
 $^1B_{1u} - \tilde{X}$ 410–470 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	1		431(14)	Ne	AB	7
	2		310(25)	gas	EM	3
			317(14)	Ne	AB	1,7

$^3B_{1u}$ D_{2h}
 $T_0=16220(160)$ gas PE²TPE⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	1		450(65)	gas	PE,TPE	2,4

$^1B_{3u}$ D_{2h}
 $T_0=11050(80)$ gas PE²TPE⁴
 $10807(2)$ Ne AB⁷ $^1B_{3u} - \tilde{X}$ 790–930 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g			300(8)	gas	TPE	4
			300(6)	Ne	AB	7

$^3B_{3u}$ D_{2h}
 $T_0=6580(80)$ gas PE²TPE⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	2		312	gas	PE,TPE	2,4

\tilde{X} 1A_g D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	1		470	N ₂	Ra	5
	2		360(25)	gas	PE	2
			345	N ₂	Ra	5
b_{1u}	4		501.9	Ne	IR	6
			500.9	Ar	IR	6
			498.5	Kr	IR	6

^aAttributed by Refs. 1 and 3 to Si₃.

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Ge₄

\tilde{a} $^3B_{3u}$ D_{2h}
 $T_0=6000T$ gas PE¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1		Breathing	173T	gas	TPE	1

Reference

- ¹ G. R. Burton, C. Xu, C. C. Arnold, and D. M. Neumark, J. Chem. Phys. **104**, 2757 (1996).

Fe₂O₂⁻

Threshold for electron detachment from ground-state Fe₂O₂⁻ = 19040(400)
gas PE¹

Reference

- ¹ H. Wu, S. R. Desai, and L.-S. Wang, J. Am. Chem. Soc. **118**, 5296 (1996).

Cu₂O₂⁻

Threshold for electron detachment from ground-state Cu₂O₂⁻ = 19850(240)
gas PE¹

Reference

- ¹ L.-S. Wang, H. Wu, S. R. Desai, and L. Lou, Phys. Rev. B **53**, 8028 (1996).

C₄⁻

Threshold for electron detachment from ground-state C₄⁻
 $= 31320(80)$ gas PE^{1,2}

(3) $^2\Pi_u$ D_{∞h}
 Ne AB⁶ 335–347 nm

(2) $^2\Pi_g$ D_{∞h}
 $T_0=26069(14)$ Ne AB⁶ 372–384 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	2		755(20)	Ne	AB	6
Π_g	4		349(10)H	Ne	AB	6
Π_u	5		271(10)H	Ne	AB	6

\tilde{C} $^2\Pi_u$ D_{∞h}
 $T_0=21871.53$ gas PD⁴
 21896(5) Ne AB³LF⁵

$\tilde{C}-\tilde{X}$ 410–460 nm

$\tilde{C}-\tilde{X}$ 410–527 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1		2241(5)T	Ne	AB	3
	2		752.4	gas	PD	4
			759(5)	Ne	AB	3
Π_u	5	Deformation	223H	gas	PD	4

$A_0=-37(9)$ gas PD⁴

$B_0=0.156$ PD⁴

\tilde{B} $^2\Sigma_u^+$ D_{∞h}
 $T_0=10789(2)$ Ne AB⁶

$\tilde{B}-\tilde{X}$ 672–927 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1		2056(5)	Ne	AB	6
	2		930(4)	Ne	AB	6
Π_u	5		259(2)H	Ne	AB	6

\tilde{A} $^2\Sigma_g^+$ D_{∞h}
 Ne AB⁶

$\tilde{A}-\tilde{X}$ 907–1206 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	2		817(3)	Ne	AB	6
Π_u	5		275(2)	Ne	AB	6

\tilde{X} $^2\Pi_g$ D_{∞h}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
Σ_g^+	1		2047(20)	Ne	LF	5
	2		936(20)	Ne	LF	5
Π_g	4		396(20)H	Ne	LF	5

$A_0=-39(9)$ gas PD⁴

$B_0=0.167$ PD⁴

References

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Si₄⁻

Threshold for electron detachment from ground-state Si₄⁻
 $= 17510(80)$ gas PE¹TPE²

²B_{1u} D_{2h}
 $T_0 = 11460(3)$ Ne AB³ ²B_{1u}– \tilde{X} 780–875 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g	1		352(4)	Ne	AB	3
	2		278(4)	Ne	AB	3

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g	2		365	gas	TPE	2

References

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CCCN

\tilde{X}	² S ⁺	C _{∞v}	Structure: MW ⁴			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
II	5	Bend	179T	gas	MW	3

$B_0 = 0.165$ MW^{1,2}

References

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cyc-NaNO₂

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	NO ₂ s-stretch	1293.0	Ar	IR	2
	2	NO ₂ scissors	825.6	Ar	IR	2
	3	ONaO s-stretch	306	Ar	IR	1
<i>b</i> ₂	5	NO ₂ a-stretch	1222.8	Ar	IR	1,2
	6	ONaO a-stretch	207	Ar	IR	1

References

¹M. Barbeschi, L. Bencivenni, and F. Ramondo, Chem. Phys. **112**, 387 (1987).

²W.-J. Lo, M.-Y. Shen, C.-H. Yu, and Y.-P. Lee, J. Chem. Phys. **104**, 935 (1996).

t-NaONO

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NO stretch	1446.2	Ar	IR	1
		ON stretch	1159.1	Ar	IR	1
		ONO deform.	787.1	Ar	IR	1

Reference

¹W.-J. Lo, M.-Y. Shen, C.-H. Yu, and Y.-P. Lee, J. Chem. Phys. **104**, 935 (1996).

cyc-KNO₂

\tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	NO ₂ s-stretch	1313.4	Ar	IR	2
	2	NO ₂ scissors	807.3	Ar	IR	1,2
	3	OKO s-stretch	227	Ar	IR	1
<i>b</i> ₂	5	NO ₂ a-stretch	1217.3	Ar	IR	1,2

References

¹M. Barbeschi, L. Bencivenni, and F. Ramondo, Chem. Phys. **112**, 387 (1987).

²W.-J. Lo, M.-Y. Shen, C.-H. Yu, and Y.-P. Lee, J. Chem. Phys. **104**, 935 (1996).

t-KONO

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NO stretch	1412.4	Ar	IR	1
		ON stretch	1171.0	Ar	IR	1
		ONO deform.	783.1	Ar	IR	1

Reference

¹W.-J. Lo, M.-Y. Shen, C.-H. Yu, and Y.-P. Lee, J. Chem. Phys. **104**, 935 (1996).

OCBeO

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CO stretch	2189.5	Ar	IR	1
		BeO stretch	1498.2	Ar	IR	1

Reference

¹L. Andrews and T. J. Tague, Jr., *J. Am. Chem. Soc.* **116**, 6856 (1994).

COBeO

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CO stretch	2056.5	Ar	IR	1
		BeO stretch	1533.9	Ar	IR	1

Reference

¹L. Andrews and T. J. Tague, Jr., *J. Am. Chem. Soc.* **116**, 6856 (1994).

OTiCO

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CO stretch	1866.7	Ar	IR	1
		TiO stretch	952.8	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, *J. Am. Chem. Soc.* **117**, 1595 (1995).

N₂FeO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NN stretch	2262.6	Ar	IR	1
			953.2	N ₂	IR	2
		FeO stretch	887.3	Ar	IR	1,3
			853.9	N ₂	IR	2

References

¹G. V. Chertihin, W. Saffel, J. T. Yustein, L. Andrews, M. Neurock, A. Ricca, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 5261 (1996).

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³G. V. Chertihin, L. Andrews, and M. Neurock, *J. Phys. Chem.* **100**, 14609 (1996).

OUCO

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CO stretch	1799.6	Ar	IR	1
		UO stretch	804.4	Ar	IR	1

Reference

¹T. J. Tague, Jr., L. Andrews, and R. D. Hunt, *J. Phys. Chem.* **97**, 10920 (1993).

(AlO)₂

\tilde{B}
 $T_0=25980(640)$ gas PE³

\tilde{A}
 $T_0=3950(160)$ gas PE³

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Sym. stretch	730(80)	gas	PE	3

 \tilde{X}^a

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1176.3T	Ar	IR	2
			945.4	Ar	IR	1,2
		Sym. stretch	660(80)	gas	PE	3

^aAb initio calculations cited in Ref. 2 suggest a linear structure, whereas those considered in Ref. 3 favor a rhombic, D_{2h} configuration. Isotopic substitution studies would be consistent with either structure. For the rhombic configuration, the absorption at 1176.3 may be contributed by a combination band.

References

¹I. L. Rozhanski, G. V. Chertikhin, L. V. Serebrennikov, and V. F. Shevel'kov, *Zh. Fiz. Khim.* **62**, 2351 (1988); *Russ. J. Phys. Chem.* **62**, 1215 (1988).

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³S. R. Desai, H. Wu, C. M. Rohlfing, and L.-S. Wang, *J. Chem. Phys.* **106**, 1309 (1997).

CNCN

\tilde{X} C_{∞v} Structure: MW⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	C≡N s-stretch	2302.00	gas	IR	1,7
			2294.4	Ar	IR	2
	2	C≡N a-stretch	2059.73	gas	IR	1,7,8
			2053.7	Ar	IR	2
	3	N-C stretch	934.73 ^a	gas	IR	6
			979.2 ^a	Ar	IR	2
II	4	Bend	463.69	gas	IR	2,5,6
			468.5	Ar	IR	2
			467.2			
	5	Bend	194.93	gas	IR	1,6
			200T ^b	Ar	IR	1

$B_0=0.173$ IR^{1,3,8} MW^{1,3,5}

^aIn Fermi resonance with 2ν₄.

^bFrom combination bands.

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CCCS

In an argon matrix, a weak absorption maximum at 26460 (378 nm) has been assigned² to CCCS.

$\tilde{X}^1\Sigma$		$C_{\infty v}$	Structure: MW ³			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	1		2058.22	gas	DL	5
			2046.2vs	Ar	IR	2
	2		1533.2wm	Ar	IR	2
	3		725.6w	Ar	IR	2

$$B_0=0.096 \text{ MW}^{1,4}$$

References

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NFeO₂

\tilde{X}		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	FeN stretch	980.5	N ₂	IR	1
	2	FeO s-stretch	837.2	N ₂	IR	1
a''	5	FeO a-stretch	928.0	N ₂	IR	1

Reference

- ¹L. Andrews, G. V. Chertihin, A. Citra, and M. Neurock, J. Phys. Chem. **100**, 11235 (1996).

NUO₂

\tilde{X}	C_1					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		UO stretch	845.5	Ar	IR	1
		UO stretch	718.3	Ar	IR	1

Reference

- ¹G. P. Kushto, P. F. Souter, L. Andrews, and M. Neurock, J. Chem. Phys. **106**, 5894 (1997).

SBCS

\tilde{X}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.
			1201.9	Ar

Reference

- ¹P. Hassanzadeh, L. Andrews, and R. D. Davy, J. Phys. Chem. **97**, 7412 (1993).

(AlO)₂⁻

Threshold for electron detachment from ground-state (AlO)₂⁻ = 15170(160) gas PE¹

Reference

- ¹S. R. Desai, H. Wu, C. M. Rohlfing, and L.-S. Wang, J. Chem. Phys. **106**, 1309 (1997).

NSCN⁺

\tilde{C}^2A''	C_s
$T^a=29370(120)$	gas PE ¹
\tilde{B}^2A'	C_s
$T^a=21700(120)$	gas PE ¹
\tilde{A}^2A''	C_s
$T^a=17670(120)$	gas PE ¹
\tilde{X}^2A'	C_s

^aFrom vertical ionization potentials.

Reference

- ¹A. W. Allaf, R. L. Johnston, and R. J. Suffolk, Chem. Phys. Lett. **233**, 33 (1995).

N₄⁺ **\tilde{A}, \tilde{B}**

A broad, unstructured absorption¹⁻³ between 270 and 650 nm, with a maximum near 330 nm, leads to the formation of N₂⁺+N₂. Detection of the fluorescence of N₂⁺ (B) at the higher energies in this range suggests that the potential energy surface for a bound excited state of N₄⁺ which correlates with N₂⁺ (B) has an avoided crossing with the surface for the dissociative \tilde{A} state.³

$\tilde{X}^2\Sigma_u^-$		D _{∞h}	Structure: ESR, MO ⁴ DL ⁷			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	N≡N s-stretch	2283T ^a	Ne	IR	6
Σ_u^+	3	N≡N a-stretch	2234.51 2237.6	gas Ne	DL IR	7 5,6

$$B_0 = 0.112 \text{ DL}^7$$

^aCalculated using observed values for asymmetrically ¹⁵N-substituted species.

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- L. B. Knight, Jr., K. D. Johannessen, D. C. Cobranchi, E. A. Earl, D. Feller, and E. R. Davidson, *J. Chem. Phys.* **87**, 885 (1987).
- M. E. Jacox and W. E. Thompson, *Res. Chem. Intermed.* **12**, 33 (1989).
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- T. Ruchti, T. Speck, J. P. Connelly, E. J. Bieske, H. Linnartz, and J. P. Maier, *J. Chem. Phys.* **105**, 2591 (1996).

BeO₃ **\tilde{X}**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		O ₃ a-stretch	869.8	Ne	IR	1
			871.8	Ar	IR	1
			436.1	Ar	IR	1

Reference

- C. A. Thompson and L. Andrews, *J. Chem. Phys.* **100**, 8689 (1994).

OTiOO **\tilde{X}**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			972.1	Ar	IR	1

Reference

- G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).

OZrOO **\tilde{X}**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			941.1	Ar	IR	1

Reference

- G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).

OHfOO **\tilde{X}**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			945.8T	Ar	IR	1

Reference

- G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 6356 (1995).

MoO₃ **\tilde{X}** **C_{3v}**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	MoO stretch	976	Ne	IR	1
<i>e</i>	3	MoO stretch	922.2	Ne	IR	1

Reference

- W. D. Hewett, Jr., J. H. Newton, and W. Weltner, Jr., *J. Phys. Chem.* **79**, 2640 (1975).

WO₃ **\tilde{X}** **D_{3h}**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>e'</i>	3	WO stretch	918.3	Ar	IR	1,2
			916	Kr	IR	1

References

- D. W. Green and K. M. Ervin, *J. Mol. Spectrosc.* **89**, 145 (1981).
- M. J. Almond and A. J. Downs, *J. Chem. Soc. Dalton Trans.* 809 (1988).

FeO₃ **\tilde{X}** **D_{3h}**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			850(50)	gas	PE	2
			975.8T	Ar	IR	1

References

- ¹G. V. Chertihin, W. Saffel, J. T. Yustein, L. Andrews, M. Neurock, A. Ricca, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 5261 (1996).
²H. Wu, S. R. Desai, and L.-S. Wang, J. Am. Chem. Soc. **118**, 5296 (1996).

(cyc-O₂Fe)O \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	OO stretch	1147.5	Ar	IR	1
		FeO stretch	928.1	Ar	IR	1

Reference

- ¹G. V. Chertihin, W. Saffel, J. T. Yustein, L. Andrews, M. Neurock, A. Ricca, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 5261 (1996).

ONiOO

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OO stretch	1393.7	Ar	IR	1

Reference

- ¹A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. A **101**, 3109 (1997).

cyc-(O₂Ni)O \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	OO stretch	1095.5T	Ar	IR	1

Reference

- ¹A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, J. Phys. Chem. A **101**, 3109 (1997).

CuO₃ \tilde{E} T₀=21060T gas PE¹ \tilde{D} T₀=9280(160) gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			660(90)	gas	PE	1

 \tilde{C} T₀=6700(160) gas PE¹ \tilde{B} T₀=2580(160) gas PE¹ \tilde{A} T₀=1610(160) gas PE¹

Reference

- ¹H. Wu, S. R. Desai, and L.-S. Wang, J. Phys. Chem. A **101**, 2103 (1997).

UO₃ \tilde{X} C_{2v}Structure: IR²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	UO stretch	843.5	Ar	IR	1,2
	2	UO ₂ s-stretch	745.6	Ar	IR	1-3
			740.7	Kr	IR	1
<i>b</i> ₂	5	UO ₂ a-stretch	852.6	Ar	IR	1-3
			848.1	Kr	IR	1

References

- ¹S. D. Gabelnick, G. T. Reedy, and M. G. Chasanov, J. Chem. Phys. **58**, 4468 (1973).

- ²S. D. Gabelnick, G. T. Reedy, and M. G. Chasanov, J. Chem. Phys. **59**, 6397 (1973).

- ³R. D. Hunt and L. Andrews, J. Chem. Phys. **98**, 3690 (1993).

Ge₂O₂ \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g	2	Ge-Ge breathing	400(60)	gas	PE	3
<i>b</i> _{2u}	5	GeO stretch	667.7	Ar	IR	1,2
			667	N ₂	IR	1
<i>b</i> _{3u}	6	GeO stretch	600.0	Ar	IR	1,2
			599	N ₂	IR	1

References

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- ²L. Andrews and M. McCluskey, J. Mol. Spectrosc. **154**, 223 (1992).

- ³J. B. Nicholas, J. Fan, H. Wu, S. D. Colson, and L.-S. Wang, J. Chem. Phys. **102**, 8277 (1995).

Sn₂O₂ \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> _{2u}	5		611.7	Kr	IR	1
			612.4	N ₂	IR	1
<i>b</i> _{3u}	6		524.4	Kr	IR	1
			523	N ₂	IR	1

Reference

¹A. Bos and J. S. Ogden, J. Phys. Chem. **77**, 1513 (1973).

Pb₂O₂

\tilde{X}	D _{2h}	Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a _g	1				518T ^a	Ar	IR	3
					522T ^a	N ₂	IR	3
b _{1g}	3				390T ^a	Ar	IR	3
					395T ^a	N ₂	IR	3
b _{2u}	5				555.6	Ar	IR	1-3
					557.6	N ₂	IR	1,3
b _{3u}	6				466.4	Ar	IR	1,3
					468.0	N ₂	IR	1,3

^aFrom combination band.

References

¹J. S. Ogden and M. J. Ricks, J. Chem. Phys. **56**, 1658 (1972).

²S. A. Konnov, L. V. Serebrennikov, and A. A. Mal'tsev, Zh. Neorg. Khim. **27**, 323 (1982); Russ. J. Inorg. Chem. **27**, 184 (1982).

³G. V. Chertihin and L. Andrews, J. Chem. Phys. **105**, 2561 (1996).

PbPb(O₂)

\tilde{X}	Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
				648.7	Ar	IR	1
				645.1			
				644.0	N ₂	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Chem. Phys. **105**, 2561 (1996).

NSCN

\tilde{X}	Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1		C≡N stretch	2225m	gas	IR	1
	2		N=S stretch	1374vs	gas	IR	1

Reference

¹A. W. Allaf and R. J. Suffolk, J. Chem. Res., Synop. 186 (1994).

FeO₃⁻

Threshold for electron detachment from ground-state FeO₃⁻=26300(320) gas PE¹

Reference

¹H. Wu, S. R. Desai, and L.-S. Wang, J. Am. Chem. Soc. **118**, 5296 (1996).

CuO₃⁻

Threshold for electron detachment from ground-state CuO₃⁻=25740(320) gas PE¹

Reference

¹H. Wu, S. R. Desai, and L.-S. Wang, J. Phys. Chem. A **101**, 2103 (1997).

Ge₂O₂⁻

Threshold for electron detachment from ground-state Ge₂O₂⁻=5040(400) gas PE¹

Reference

¹J. B. Nicholas, J. Fan, H. Wu, S. D. Colson, and L.-S. Wang, J. Chem. Phys. **102**, 8277 (1995).

BrCNO⁺

\tilde{D}

T^a=65110(200) gas PE¹

\tilde{C}

T^a=57200(200) gas PE¹

\tilde{B}

T^a=46400(200) gas PE¹

\tilde{A}

T^a=23100(200) gas PE¹

^aFrom vertical ionization potentials.

Reference

¹T. Pasinszki and N. P. C. Westwood, J. Phys. Chem. **99**, 6401 (1995).

t-(NO)₂⁺

\tilde{A}

Dissociates into NO+NO⁺, with onset at 10700(1000) and maximum at 16400(1000). AB^{2-5,7}PE⁸

 \tilde{X} C_{2h}

\tilde{X}	Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	a _g	1	NO s-stretch	2090	gas	PI,PE	6,8
				2064.7 ^a	Ne	TPE	11,12
		2	NNO s-bend	323	gas	TPE	10
	a _u	4	Torsion	118T	gas	TPE	11,12
	b _u	5	NO a-stretch	1618	gas	TPE	12
				1619.2	Ne	IR	10,12
				1593.3	Ar	IR	1,9

^a($\nu_1 + \nu_5$) - ν_5 .

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⁷M. F. Jarrold, A. J. Illies, and M. T. Bowers, *J. Chem. Phys.* **79**, 6086 (1983).
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⁹J. Hacaloglu, S. Suzer, and L. Andrews, *J. Phys. Chem.* **94**, 1759 (1990).
¹⁰M. E. Jacox and W. E. Thompson, *J. Chem. Phys.* **93**, 7609 (1990).
¹¹I. Fischer, A. Strobel, J. Staeker, G. Niedner-Schatteburg, K. Müller-Dethlefs, and V. E. Bondybey, *J. Chem. Phys.* **96**, 7171 (1992).
¹²A. Strobel, N. Knoblauch, J. Agreiter, A. M. Smith, G. Niedner-Schatteburg, and V. E. Bondybey, *J. Phys. Chem.* **99**, 872 (1995).

ArBeO₂

\tilde{X}		C_{2v}				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1288.9	Ar	IR	1,2
			1264.1	Ar	IR	1,2

References

- ¹C. A. Thompson and L. Andrews, *J. Chem. Phys.* **100**, 8689 (1994).
²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 10088 (1996).

FNCO

\tilde{X}		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a'</i>	1	NCO a-stretch	2174.8vs	Ne	IR	2
			2174.4s	Ar	IR	1,2
			2167s			
	2	NCO s-stretch	1245T	Ne	IR	2
	3	NF stretch	860.8wm	Ne	IR	2
			862.0m	Ar	IR	1,2
<i>a''</i>	4	NCO deform.	701.5wm	Ne	IR	2
			695wm	Ar	IR	1
	5	FNC deform.	203.5w	Ne	IR	2
<i>a''</i>	6	NCO deform.	533.8wm	Ne	IR	2
			529m	Ar	IR	1

References

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²J. Jacobs, B. Jülicher, G. Schatte, H. Willner, and H.-G. Mack, *Chem. Ber.* **126**, 2167 (1993).

BrCNO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CNO a-stretch	2211	gas	IR	2
			2271.3	Ar	IR	1
		CNO s-stretch	2252.5			
			1321	gas	IR	2
			1305.6	Ar	IR	1

References

- ¹G. Maier and J. H. Teles, *Angew. Chem.* **99**, 152 (1987); *Angew. Chem. Int. Ed. Engl.* **26**, 155 (1987).
²T. Pasinszki and N. P. C. Westwood, *J. Phys. Chem.* **99**, 6401 (1995).

SiO₃ \tilde{X}

\tilde{X}		C_{2v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Si=O stretch	1363.5vs	Ar	IR	1
	2	OSiO s-stretch	877.1wm	Ar	IR	1
<i>b</i> ₁	4	OPLA	287.8vs	Ar	IR	1
<i>b</i> ₂	5	OSiO a-stretch	855.3ms	Ar	IR	1
	6	Deformation	292.0s	Ar	IR	1

Reference

- ¹B. Tremblay, P. Roy, L. Manceron, M. E. Alikhani, and D. Roy, *J. Chem. Phys.* **104**, 2773 (1996).

OPb(O₂) \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			702.6	Ar	IR	1
			531.3	Ar	IR	1

Reference

- ¹G. V. Chertihin and L. Andrews, *J. Chem. Phys.* **105**, 2561 (1996).

c-(NO)₂

A diffuse gas-phase absorption with onset below 38500 (>260 nm) and maximum at 48800 (205 nm) has been assigned^{4,7} to c-(NO)₂.

\tilde{X}		C _{2v}	Structure: MW ⁹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	NO s-stretch.	1868.25	gas	IR,DL	3,13,15
			1867.2	Ne	IR	14,16,17
			1866m	Ar	IR	1
			1867.0	N ₂	IR	2,17
			1862m	CO ₂	IR,Ra	1,8
	2	N-N stretch	1866	NO	IR,Ra	5,6,10
			262	CO ₂	Ra	8
	3	NNO s-bend	266	NO	IR,Ra	5,6,10
			161	CO ₂	Ra	8,10
			187	NO	Ra	5,6,10
<i>a</i> ₂	4	Torsion	88.2 ^a	gas	IR	11
			97	NO	IR,Ra	5,6,10
<i>b</i> ₂	5	NO a-stretch	1789.09	gas	IR,DL	3,11,12
						13,18
			1780.6	Ne	IR	14,16,17
			1778.7			
			1776s	Ar	IR	1
	6	NNO a-bend	1781.2	N ₂	IR	2,17
			1768s	CO ₂	IR	1
			1762	NO	IR,Ra	6,10
			202	CO ₂	Ra	8,10
			214	NO	Ra	5,6,10

$$A_0 = 0.862; B_0 = 0.187; C_0 = 0.154 \text{ MW}^9 \text{ IR}^{15}$$

^aFrom observation of $\nu_5 - \nu_4 = 1700.8 \text{ cm}^{-1}$.

References

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- ¹⁷F. Legay and N. Legay-Sommaire, *J. Chem. Phys.* **102**, 7798 (1995).
- ¹⁸A. Dkhissi, P. Soulard, A. Perrin, and N. Lacome, *J. Mol. Spectrosc.* **183**, 12 (1997).

t-(NO)₂ \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g	1	Sym. stretch	1861.1vw	N ₂	IR	4
<i>b</i> _u	5	Asym. stretch	1762.5	Ne	IR	3
			1760.6			
			1759.6	N ₂	IR	2,4
			1740	CO ₂	IR	1

References

- ¹W. G. Fateley, H. A. Bent, and B. Crawford, Jr., *J. Chem. Phys.* **31**, 204 (1959).
- ²W. A. Guillory and C. E. Hunter, *J. Chem. Phys.* **50**, 3516 (1969).
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NNO₂

³A₁ C_{2v}
 $T_0 = x + 8110$ gas PE¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	NO ₂ scissors	540(40)	gas	PE	1

³A₂ C_{2v}
 $T_0 = x^a$ gas PE¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	NO ₂ s-stretch	1240T	gas	PE	1
	2	NN stretch	880T	gas	PE	1
	3	NO ₂ scissors	680T	gas	PE	1

^aIt is uncertain whether this or a ¹A' state which is calculated¹ to be nearly isoenergetic is the ground state of NNO₂.

Reference

- ¹D. W. Arnold and D. M. Neumark, *J. Chem. Phys.* **102**, 7035 (1995).

SNNO \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NO stretch	1815.2	Ar	IR	1

Reference

- ¹L. Andrews, P. Hassanzadeh, G. D. Brabson, A. Citra, and M. Neurock, *J. Phys. Chem.* **100**, 8273 (1996).

$A = 1600(80)$ gas TPE⁶
 $\tau = 5.5(5)$ ns gas T-PEFCO⁶

\tilde{C}^2A_2'' D_{3h}
 $T_0 = 21780(120)$ gas PE¹TPE⁶

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	BCL stretch	403(16)	gas	PE,TPE	1,6

\tilde{B}^2E' D_{3h}
 $T^a = 9280(120)$ gas PE¹TPE⁶
 $A = 800T$ gas TPE⁶

\tilde{A}^2E'' D_{3h}
 $T_0 = 5080(120)$ gas PE¹TPE⁶

$\tilde{X}^2A'_2$ D_{3h}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
e'	3	BCL stretch	1103.8	Ne	IR	7
			1090	Ar	IR	2

^aFrom vertical ionization potential.

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FCO₂

\tilde{B}^2A_1 C_{2v}
 $T_0 = 13150$ gas AB^{1,2}
 $13103(5)$ Ne AB⁴

$\tilde{B}-\tilde{X}$ 500–795 nm
 $\tilde{B}-\tilde{X}$ 508–763 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.		
a'_1	1	OCO s-stretch	1607(10)	Ne	AB	4		
	2	CF stretch	1110	gas	AB	1		
			1119(5)	Ne	AB	4		
b'_1	4	OCPLA	610	gas	AB	1		
			614(5)	Ne	AB	4		
			840	gas	AB	1		
			838(5)	Ne	AB	4		

\tilde{A}^2A_2 C_{2v}
 $T_0 = 4670(40)$ gas PE³

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	OCO s-stretch	1274(40)	gas	PE	3
	2	CF stretch	903(40)	gas	PE	3
	3	OCO bend	532(40)	gas	PE	3

\tilde{X}^2B_2 C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'_1	1	OCO s-stretch	1465(70)	gas	PE	3
			1475Tvs	Ne	IR	4
	2	CF stretch	950(70)	gas	PE	3
			968.9m	Ne	IR	4
			975.1	Ar	IR	4
			975.5	N ₂	IR	4
	3	OCO bend	520	gas	AB,PE	1,3
			518.5wm	Ne	IR	4
			521.6	Ar	IR	4
b'_1	4	OPLA	522.8	N ₂	IR	4
			735.2wm	Ne	IR	4
b'_2	5	OCO a-stretch	730.6	Ar	IR	4
			728.5	N ₂	IR	4
			1097.7ms	Ne	IR	4
			1113.6	Ar	IR	4
			1116.5	N ₂	IR	4
b'_6	6	FCO bend	474.3wm	Ne	IR	4
			482.5	Ar	IR	4
			481.3	N ₂	IR	4

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NNO₂⁻

\tilde{B}^2A_2 C_{2v}

Photoelectron spectra² of NNO₂⁻ indicate that at both 532 and 355 nm dissociation into N₂O+O⁻ occurs. Fast ion beam translational spectroscopy studies⁶ indicate that a structured absorption with onset near 580 nm arises from excitation to the \tilde{B} state. Two different dissociation channels occur in this energy range; dissociation on the ground-state potential surface leads to the formation of N₂O+O⁻, and excited-state dissociation results in the formation of NO+NO⁻. When 266 and 213 nm photons are used, the photoelectron spectra⁵ suggest that electron detachment to form excited states of uncharged NNO₂ occurs.

NSeNSe \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			902.9	Ar	IR	1

Reference

¹L. Andrews, P. Hassanzadeh, D. V. Lanzisera, and G. D. Brabson, *J. Phys. Chem.* **100**, 16667 (1996).

NO₃⁺

\tilde{B}^1E' D_{3h}
 $T^{\circ}=24120(120)$ gas PE¹

\tilde{A}^1E'' D_{3h}
 $T^{\circ}=12100(120)$ gas PE¹

\tilde{b}^3E' D_{3h}
 $T^{\circ}=8630(120)$ gas PE¹

\tilde{a}^3E'' D_{3h}
 $T^{\circ}=5080(120)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	1050(60)	gas	PE	1

$\tilde{X}^1A'_1$ D_{3h}

^aFrom vertical ionization potentials.

Reference

¹D. Wang, P. Jiang, X. Zian, and G. Hong, *J. Chem. Phys.* **106**, 3003 (1997).

BF₃⁺

$\tilde{E}^2A'_1$ D_{3h}
 $T_0=47800$ gas PE³

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	BF stretch	800T	gas	PE	3

$\tau=10$ ns gas EM⁵

\tilde{D}^2E' D_{3h}
 $T_0=34860(160)$ gas PE^{1,4}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	BF stretch	742(20)	gas	PE	1,4

Additional structure reported by Ref. 4 has been attributed to vibronic coupling.

$\tilde{C}^2A''_2$ D_{3h}
 $T_0=27110(80)$ gas PE^{1,2,4}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	BF stretch	830(20)	gas	PE	2,4

\tilde{B}^2E' D_{3h}
 $T_0=10890(240)$ gas PE^{1,2,4}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	BF stretch	770(60)	gas	PE	1,4

\tilde{A}^2E'' D_{3h}
 $T_0=5890(240)$ gas PE^{1,4}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
e'	3	BF ₃ stretch	1661.6	Ne	IR	6

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¹P. J. Bassett and D. R. Lloyd, *J. Chem. Soc. A* 1551 (1971).

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⁶M. E. Jacox and W. E. Thompson, *J. Chem. Phys.* **102**, 4747 (1995).

BCl₃⁺

$\tilde{E}^2A'_1$ D_{3h}
 $T_0=49200(120)$ gas PE¹TPE⁶

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	BCl stretch	403(16)	gas	PE,TPE	1,6

\tilde{D}^2E' D_{3h}
 $T_0=29700(120)$ gas PE¹TPE⁶

Emission which is observed between about 300 and 650 nm on excitation of BCl₃⁺ by radiation of wavelength shorter than 81 nm (15.36 eV) arises from the \tilde{D} state of BCl₃⁺.³⁻⁵ The emission between 330 and 420 nm has been attributed to the $\tilde{D}-\tilde{X}$ transition of BCl₃⁺, and that between 420 and 580 nm, with vibrational spacings of ~445, to the $\tilde{D}-\tilde{A}, \tilde{B}$ transitions.³ The T-PEPICO measurements of Ref. 6 indicate that the quantum yield for these emissions is ≤ 0.04 and that fragmentation into BCl₂⁺+Cl predominates.

A broad absorption with maximum at 320 nm (31200) which appears on argon-resonance photolysis of BCl₃ isolated in an argon matrix and which can be destroyed by prolonged exposure of the sample to 340–600 nm radiation has been assigned² to the $\tilde{D}-\tilde{X}$ transition of BCl₃⁺.

\tilde{X}^2B_2 C_{2v}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	NN stretch	1359.6wm	Ne	IR	4
			1355w	Ar	IR	1,3,4
	2	NO ₂ s-stretch	950T	gas	PE	5
			1004.5wm	Ne	IR	4
b_2			1008w	Ar	IR	1,3,4
	5	NO ₂ a-stretch	1199.3m	Ne	IR	4
			1205.5m	Ar	IR	1,3

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 $t(\text{NO})_2^-$

In gas-phase photoelectron spectra¹ taken at 532 nm, evidence was obtained for both electron detachment and photodissociation into NO+NO⁻.

 \tilde{X} C_{2h}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
b_u	5	Asym. stretch	1424.1T	Ne	IR	2

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- ¹L. A. Posey and M. A. Johnson, J. Chem. Phys. **88**, 5383 (1988).
²M. E. Jacox and W. E. Thompson, J. Chem. Phys. **93**, 7609 (1990).

 NO_3 \tilde{B}^2E' D_{3h}

$T_0=15089$ gas AB^{1,2,4-7}LF^{8,9,17,21}

$\tilde{B}-\tilde{X}$ 450–795 nm

All bands are diffuse.^{2,7,21} The threshold for the production of NO+O₂≤16780, and that for the production of O+NO₂ is 17090(20).²⁰ Above this second threshold, the predominant photodissociation products are O(³P) and NO₂.^{19,20}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	930	gas	AB	2,6
			1450	gas	AB	6
			850	gas	AB	6

$\tau_0=340(20)\mu\text{s}$ gas LF¹⁰

 \tilde{A}^2E'' D_{3h}

$T_0=7000(110)$ gas PE¹⁴

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	804(4)T	gas	PE	14
e'	4	Deformation	541(8)T	gas	PE	14

 $\tilde{X}^2A'_2$ D_{3h}

Structure: DL¹¹

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'_1	1	Sym. stretch	1050	gas	LF,PE	8,9,14,17
a''_2	2	OPLA	762.33	gas	IR,LF	12,17
e'	3	NO stretch ^a	1492.39	gas	LF,DL	8,9,11,17
	4	Deformation ^b	360	gas	IR	12
					LF,PE	8,9,14,17

$B_0=0.459$ DL^{11,13}IR¹²

^aAnomalous structure appears in the high resolution spectrum of this band because of vibronic coupling with the \tilde{B}^2E' state.¹⁶ Coupling with this state and with the \tilde{A}^2E'' state leads to the appearance of infrared absorptions of several prominent overtone and combination bands between 1900 and 2600.^{15,18} Among these bands, also observed in fluorescence,^{8,17} is one near 2020, attributed by Ref. 17 to the occurrence of pseudorotation on the ground-state potential surface but by Ref. 15 to intensification of the fourth overtone of ν_4 , which has appreciable “negative” anharmonicity, as a result of vibronic interaction with the \tilde{A} and \tilde{B} states.

^bCoupled to the \tilde{B}^2E' state through this mode.¹⁴

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SSNO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NO stretch	1772.5	Ar	IR	1
		SS stretch	670.9	Ar	IR	1

Reference

- ¹L. Andrews, P. Hassanzadeh, G. D. Brabson, A. Citra, and M. Neurock, *J. Phys. Chem.* **100**, 8273 (1996).

SeSeNSe \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			735.1	Ar	IR	1

Reference

- ¹L. Andrews, P. Hassanzadeh, D. V. Lanzisera, and G. D. Brabson, *J. Phys. Chem.* **100**, 16667 (1996).

SeSeSeN \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			527	Ar	IR	1

Reference

- ¹L. Andrews, P. Hassanzadeh, D. V. Lanzisera, and G. D. Brabson, *J. Phys. Chem.* **100**, 16667 (1996).

t-O₄⁺

In the gas phase, the high frequency tail of a photodissociation continuum, resulting in the formation of O₂⁺+O₂, has been observed¹⁻³ between 450 and 680 nm.

 \tilde{X} C_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a _g	1	O=O s-stretch	1644.1 ^a	Ne	IR	4
b _u	5	O=O a-stretch	1164.4	Ne	IR	4
			1118	Ar	IR	5

^a($\nu_1 + \nu_5$) - ν_5 .

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cyc-O₄⁺ \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		O=O s-stretch	1628.3 ^a	Ne	IR	1,2
		O=O a-stretch	1320.3	Ne	IR	1,2

^a($\nu_1 + \nu_5$) - ν_5 .

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- ¹W. E. Thompson and M. E. Jacox, *J. Chem. Phys.* **91**, 3826 (1989).
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CF₃⁺ \tilde{X} D_{3h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₂ '	2	OPLA	798.1	Ne	IR	5
e'	3	CF stretch	1662.4 ^a	Ne	IR	3,5
			1667	Ar	IR	1,2,4

^aAbsorptions at 1650.6 and 1669.5 which appear in some neon-matrix experiments³⁻⁵ are attributed to CF₃⁺ in other trapping sites.

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SiF₃⁺ \tilde{X} D_{3h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
e'	3	SiF ₃ stretch	1159.7	Ne	IR	1

Reference

¹M. E. Jacox, K. K. Irikura, and W. E. Thompson, J. Chem. Phys. **103**, 5308 (1995).

FCO₂⁻

Threshold for electron detachment from ground-state FCO₂⁻=34500(240) gas PE²

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CO stretch	1399 ^a	gas	PE	2
			1316 Cs	Ar	IR	1
	2	OCO bend	899 ^a	gas	PE	2
			883 Cs	Ar	IR	1
<i>b</i> ₂	3	CF stretch	596 ^a	gas	PE	2
	5	CO stretch	1749 Cs	Ar	IR	1

^aFrom Franck-Condon simulation of observed photoelectron spectrum.

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NO₃⁻

\tilde{X}		D _{3h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>e'</i>	3	NO ₃ stretch	1356.2	Ne	IR	1

Reference

- ¹D. Forney, W. E. Thompson, and M. E. Jacox, J. Chem. Phys. **99**, 7393 (1993).

CINO₂

The onset of continuous absorption by gas-phase CINO₂ occurs near 25000 (400 nm).³ The maximum in this absorption is beyond the 185 nm limit of the observations.

\tilde{X} 1A ₁		C _{2v}					Structure: MW ⁵
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
<i>a</i> ₁	1	NO ₂ s-stretch	1267.3s	gas	IR	1,7	
			1264.3	Ar	IR	4	
	2	NO ₂ scissors	792.76s	gas	IR	1,7,8	
			787.0	Ar	IR	4	
<i>b</i> ₁	3	Cl-N stretch	363.8vs	gas	IR	1,7	
			365.0	Ar	IR	4	
	4	OPLA	651.7w	gas	IR	1,7	
<i>b</i> ₂	5	NO ₂ a-stretch	1683.89s	gas	IR	1,6,7	
			1674.8	Ar	IR	4	
	6	NO ₂ rock	408.1vw	gas	IR	1	

$$A_0=0.443; B_0=0.173; C_0=0.124 \text{ MW}^{2.5} \text{ IR}^6$$

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c-CIONO

\tilde{X}		C _s		Structure: MW ⁴			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
<i>a'</i>	1	N=O stretch	1715vs	gas	IR	2,3	
			1714s	Ar	IR	1	
			1717	N ₂	IR	5	
	2	ONO bend ^a	858w	gas	IR	2	
			856m	Ar	IR	1	
	3	ClO stretch ^a	644m	gas	IR	2	
<i>a''</i>	4	O-N stretch ^a	406vs	gas	IR	2	
			398m	Ar	IR	1	
	5	ClON bend	270T	gas	IR	2	
<i>a''</i>	6	Torsion ^a	344vw	gas	IR	2	

$$A_0=0.608; B_0=0.152; C_0=0.122 \text{ MW}^4$$

^aSee Ref. 6.

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t-CIONO^a

\tilde{X}		Structure: MW ⁵				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		N=O stretch	1752ms	Ar	IR	1

^aOriginally believed to be OCINO. Reassigned by Ref. 2 to t-CIONO.

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BrNO₂

In the gas phase, unstructured absorption maxima have been observed⁵ at 50250 (199 nm), 40490 (247 nm), and 26880 (372 nm).

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	NO ₂ s-stretch	1292	gas	IR	3-5
			1290.7	Ar	IR	1,2,5
			1291	O ₂	IR	4
	2	NO ₂ deform.	787	gas	IR	3-5
			782.9s	Ar	IR	2,5
			784	O ₂	IR	4
<i>b</i> ₁	3	NBr stretch	281.8w	Ar	IR	5
	4	OPLA	605	gas	IR	5
			605.6	Ar	IR	5
<i>b</i> ₂	5	NO ₂ a-stretch	1667	gas	IR	3-5
			1659.2vs	Ar	IR	2,5
			1655	O ₂	IR	4
	6	NO ₂ wag	290Tsh	Ar	IR	5

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- D. Scheffler, H. Grothe, H. Willner, A. Frenzel, and C. Zetsch, *Inorg. Chem.* **36**, 335 (1997).

t-BrONO

\tilde{X}		C _s				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	N=O stretch	1727.9vs	Ar	IR	1-3
	2	ONO deform.	833.2ms	Ar	IR	2,3
	3	BrO stretch	584.8vs	Ar	IR	1-3
	4	O-N stretch	391.1wm	Ar	IR	2,3
	5	BrON deform.	197T	Ar	IR	3
	6	Torsion	150HT	Ar	IR	3
<i>a''</i>						

References

- M. Feuerhahn, R. Minkwitz, and U. Engelhardt, *J. Mol. Spectrosc.* **77**, 429 (1979).
- D. E. Tevault, *J. Phys. Chem.* **83**, 2217 (1979).
- D. Scheffler, H. Grothe, H. Willner, A. Frenzel, and C. Zetsch, *Inorg. Chem.* **36**, 335 (1997).

OSOO

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OS stretch	1229.6	Ar	IR	1
		OO stretch	1041.3	Ar	IR	1
			597.6	Ar	IR	1

Reference

- S.-H. Jou, M.-Y. Shen, C.-H. Yu, and Y.-P. Lee, *J. Chem. Phys.* **104**, 5745 (1996).

S₄ **\tilde{A}**

gas AB^{1,2,4} $\tilde{A}-\tilde{X}$ 425-575 nm
Unstructured absorption maximum at 19300 (518 nm) in an Ar matrix.⁶
Unstructured absorption maximum at 18870 (530 nm) in a Kr matrix.^{1,2}

 \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			680	gas	Ra	7
			661.6	Ar	IR	3,5,6
			660s	Kr	IR	3
			660	Xe	IR	3
			575	gas	Ra	7
			483wmT	Kr	IR	3
			303	gas	Ra	7
			320mT	Kr	IR	3
			270wmT	Kr	IR	3

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SeO₃

\tilde{X}		D _{3h}		Structure: Ra ¹		
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a''</i>	2	Deform.	356	Ar	IR	2
			358	N ₂	IR	2
<i>e'</i>	3	SeO stretch	995	Ar	IR	2,3
			1005.5	N ₂	IR	2

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OSeOO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		OO stretch	1141.5	Ar	IR	1
			952.0	Ar	IR	1

Reference

¹G. D. Brabson, L. Andrews, and C. J. Marsden, *J. Phys. Chem.* **100**, 16487 (1996).

cyc-(SeOSe)=O \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		Se=O stretch	966.2	Ar	IR	1
			636.5	Ar	IR	1
			457	Ar	IR	1

Reference

¹G. D. Brabson, L. Andrews, and C. J. Marsden, *J. Phys. Chem.* **100**, 16487 (1996).

BF₃⁻ \tilde{X} C_{3v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁	1	BF ₃ stretch	847.2wm	Ne	IR	1
	2	Umbrella	595.3wm	Ne	IR	1
<i>e</i>	3	BF ₃ stretch	1026.4m	Ne	IR	1
	4	Deformation	409T	Ne	IR	1

Reference

¹M. E. Jacox and W. E. Thompson, *J. Chem. Phys.* **102**, 4747 (1995).

BCl₃⁻ \tilde{X} C_{3v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>e</i>	3	BCl ₃ stretch	685.8	Ne	IR	1

Reference

¹M. E. Jacox, K. K. Irikura, and W. E. Thompson, *J. Chem. Phys.* **104**, 8871 (1996).

CF₃

Rydberg state D_{3h}
gas AB⁶MPI¹⁰ 139–165 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₂ ^{''}	2	OPLA	820	gas	AB,MPI	6,10

4s ²A₁' D_{3h}
 $T_0 \approx 51665$ gas AB⁷EM^{11–14,16,20} 4s²A₁'– \tilde{X} 180–300 nm

Calculations¹⁴ suggest that this state is of mixed valence-Rydberg character, with increasing Rydberg contribution at large C–F distances.

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁ '	1	Sym. stretch	804T	gas	EM	12

$\tau = 12(3)$ ns gas EM^{17,18}
Ref. 22 obtained a lifetime of 17.3(4) ns for this state, suggesting that it is identical to the upper state for the 450–750 nm emission bands of CF₃.

3p ²A₂'', ²E' D_{3h}
 $T_0 \approx 51600$ gas EM^{8,11,13,14} 3p–3s²A₁' 450–750 nm
 $\tau = 17.5(5)$ ns gas EM^{17,18,22}EF²¹

The lower state of the visible emission of CF₃, calculated¹⁴ to be the 3s ²A₁' state, which assumes increasing valence character at large C–F distances, is both observed and calculated to be dissociative.

\tilde{X} ²A₁ C_{3v} Structure: ESR¹MW⁹DL¹⁵

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
<i>a</i> ₁	1	CF stretch	1089	gas	IR,EM	2,12	
			1086	Ne	CARS	19	
			1087s	Ar	IR	5,23	
	2	Umbrella	701(3)	gas	IR,EM	2,12	
			701.4	Ne	IR	5,23	
	3	CF stretch	703m	Ar	IR	3,4	
			1260.16	gas	IR,DL	2,15	
			1253.8	Ne	IR	5,23	
			1251vs	Ar	IR	3,4	
	4		508.7	Ne	IR	5,23	
			512w	Ar	IR	4	

$B_0 = 0.364$ MW⁹; $C_0 = 0.189$ DL¹⁵

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²²J. C. Creasey, I. R. Lambert, R. P. Tuckett, and A. Hopkirk, *Mol. Phys.* **71**, 1355 (1990).
²³D. Forney, M. E. Jacox, and K. K. Irikura, *J. Chem. Phys.* **101**, 8290 (1994).

SiF₃

Unstructured emission bands between 290 and 340 nm and between 350 and 800 nm which appear with varying relative intensities on photoexcitation of SiF₄ by radiation at 99.1, 95.5, or 92.2 nm have been attributed⁴ to SiF₃. The 350–800 nm band may correspond to the unstructured emission centered at 632 nm which has been observed^{3,5} in the flowing afterglow of an SiF₄ discharge and in the reaction of F atoms with Si and which has been tentatively assigned to SiF₃. A third emission band, between 240 and 280 nm, becomes more prominent at higher excitation energies. This latter band occurs in the same spectral region as the 210–260 nm emission band system observed in a discharge through SiF₄. Although that band system was initially assigned² to SiF₃, subsequent studies⁶ have demonstrated that it is entirely contributed by SiF₂.

\tilde{X}		C_{3v}				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	SiF stretch	834.9	Ne	IR	7
			832s	Ar	IR	1
e	2	Umbrella	406s	Ar	IR	1
	3	SiF stretch	958.6	Ne	IR	7
			954vs	Ar	IR	1
	4	Deformation	290wm	Ar	IR	1

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- ¹D. E. Milligan, M. E. Jacox, and W. A. Guillory, *J. Chem. Phys.* **49**, 5330 (1968).
²J. L.-F. Wang, C. N. Krishnan, and J. L. Margrave, *J. Mol. Spectrosc.* **48**, 346 (1973).
³V. M. Donnelly and D. L. Flamm, *J. Appl. Phys.* **51**, 5273 (1980).
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⁵S. Vanhaelemeersch, J. Van Hoeymissen, D. Vermeylen, and J. Peeters, *J. Appl. Phys.* **70**, 3892 (1991).
⁶W. B. Griffith, Jr. and C. W. Mathews, private communication.
⁷M. E. Jacox, K. K. Irikura, and W. E. Thompson, *J. Chem. Phys.* **103**, 5308 (1995).

NF₃⁺

\tilde{E} 2E C_{3v}
 $T^a=65920(900)$ gas PE¹

\tilde{D} 2A_1 C_{3v}
 $T_0=50600(560)$ gas PE¹

\tilde{C} 2E C_{3v}
 $T_0=33800(560)$ gas PE¹

\tilde{B} 2A_2 C_{3v}
 $T^a=28900(720)$ gas PE¹

\tilde{A} 2E C_{3v}
 $T_0=20330(650)$ gas PE¹

\tilde{X} 2A_1 C_{3v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	NF ₃ stretch	967.0wm	Ne	IR	3
	2	Umbrella	565(40)	gas	PE	2
e	3	NF ₃ stretch	658.6m	Ne	IR	3
	4	Deformation	1233.5vs	Ne	IR	3
			522.1wm	Ne	IR	3

Barrier to inversion ≈ 6000 .²

^aFrom vertical ionization potential.

References

- ¹P. J. Bassett and D. R. Lloyd, *J. Chem. Soc. Dalton Trans.* 248 (1972).
²J. Berkowitz and J. P. Greene, *J. Chem. Phys.* **81**, 3383 (1984).
³M. E. Jacox and W. E. Thompson, *J. Chem. Phys.* **102**, 6 (1995).

ClO₃

In a neon matrix,¹ bands between 300 and 440 nm have been attributed to ClO₃, and photodissociation of ClO₃ to produce OCLO+O has been observed at wavelengths longer than 420 nm.

\tilde{X} C_{3v} Structure: IR¹

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	ClO stretch	905.0w	Ne	IR	1
	2	Deform.	566.6wm	Ne	IR	1
e	3	ClO stretch	1081.3vs	Ne	IR	1
	4	Deform.	475.8m	Ne	IR	1

Reference

- ¹H. Grothe and H. Willner, *Angew. Chem.* **106**, 1581 (1994); *Angew. Chem. Int. Ed. Engl.* **33**, 1483 (1994).

CF₃⁻

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1		1050.1wT	Ne	IR	1
e	3	CF ₃ stretch	778.4mT	Ne	IR	1

Reference

¹D. Forney, M. E. Jacox, and K. K. Irikura, J. Chem. Phys. **101**, 8290 (1994).

SiF₃⁻

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	SiF ₃ stretch	770.3	Ne	IR	1
	2	Umbrella	401H	Ne	IR	1
e	3	SiF ₃ stretch	757.0	Ne	IR	1
			750.9			

Reference

¹M. E. Jacox, K. K. Irikura, and W. E. Thompson, J. Chem. Phys. **103**, 5308 (1995).

CIOOCI

An unstructured gas-phase absorption with maximum at 40800 (245 nm) has been assigned^{1,4-6,8} to CIOOCl. Cl atoms have been detected⁶ on irradiation of CIOOCl in this band.

In an argon matrix, photodecomposition occurs at wavelengths shorter than 360 nm (27800).⁷



Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a	1	OO stretch	750	gas	IR	4
			754.0m	Ar	IR	2,7
b	2	CIO s-stretch	560	gas	IR	4
			543.0m	Ar	IR	7
b	5	CIO a-stretch	653	gas	IR	4
			647.7vs	Ar	IR	2,7
	6	Deformation	418.5w	Ar	IR	7

A₀=0.437; B₀=0.080; C₀=0.071 MW³

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⁷J. Jacobs, M. Kronberg, H. S. P. Müller, and H. Willner, J. Am. Chem. Soc. **116**, 1106 (1994).

⁸K. J. Huder and W. B. DeMore, J. Phys. Chem. **99**, 3905 (1995).

CICIO₂

In the gas phase, prominent, unstructured absorption maxima appear²⁻⁴ at 44200 and 33800 (226 and 296 nm). In a neon matrix, unstructured absorption maxima are observed^{1,3,4} at 42700 and 34100 (234 and 293 nm), with half band widths of 40 and 45 nm, respectively.

In an argon matrix, the threshold⁴ for photodecomposition into Cl+OCIO, followed by cage recombination to form CIOClO, lies near 16400 (610 nm). The maximum for this process⁵ lies near 33300 (300 nm).



Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CIO ₂ s-stretch	1041.5(5)s	gas	IR	2,3
			1041.2s	Ne	IR	1,3
			1040.7	Ar	IR	5
	2	CIO ₂ scissors	522.7(5)wm	gas	IR	2,3
			522.5wm	Ne	IR	1,3
	3	ClCl stretch	440.5(5)s	gas	IR	2,3
			440.4s	Ne	IR	1,3
	4	Umbrella	271.4wm	Ne	IR	1,3
	5	ClO ₂ a-stretch	1218.2(5)vs	gas	IR	2,3
			1216.4vs	Ne	IR	1,3
			1213.2	Ar	IR	5
	6	ClO ₂ rock	251.4vw	Ne	IR	1,3

A₀=0.315; B₀=0.120; C₀=0.093 MW⁶

References

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BrClO₂

In an argon matrix, an absorption maximum near 30800 (325 nm) is associated¹ with isomerization to BrOClO.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	ClO ₂ s-stretch	1019.2	Ar	IR	1
a''	5	ClO ₂ a-stretch	1193.5	Ar	IR	1

Reference

- K. Johnsson, A. Engdahl, J. Kölm, J. Nieminen, and B. Nelander, J. Phys. Chem. **99**, 3902 (1995).

IClO₂

In an argon matrix, an absorption maximum near 28600 (350 nm) is associated¹ with photodissociation into ICl + O₂.

\tilde{X}		C _s				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	ClO ₂ s-stretch	987.4	Ar	IR	1
a''	5	ClO ₂ a-stretch	1161.2	Ar	IR	1

Reference

¹K. Johnsson, A. Engdahl, J. Kölöm, J. Nieminen, and B. Nelander, *J. Phys. Chem.* **99**, 3902 (1995).

ClOCIO

In an argon matrix, an absorption maximum at 22200 (450 nm) is associated² with isomerization to ClClO₂.

In an argon matrix, photoisomerization into ClOOCl occurs between 488 and 665 nm.¹

 \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1		ClO _t stretch	994.5vs	Ar	IR	1,2
2		ClO stretch	695.9w	Ar	IR	1
3		OCl stretch	440.3w	Ar	IR	1
4		ClOCl deform.	338.7vw	Ar	IR	1

References

¹J. Jacobs, M. Kronberg, H. S. P. Müller, and H. Willner, *J. Am. Chem. Soc.* **116**, 1106 (1994).

²K. Johnsson, A. Engdahl, J. Kölöm, J. Nieminen, and B. Nelander, *J. Phys. Chem.* **99**, 3902 (1995).

BrOCIO

In an argon matrix, an absorption maximum at 19000 (525 nm) is associated¹ with isomerization to BrClO₂.

 \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1		ClO _t stretch	991.1	Ar	IR	1

Reference

¹K. Johnsson, A. Engdahl, J. Kölöm, J. Nieminen, and B. Nelander, *J. Phys. Chem.* **99**, 3902 (1995).

ClOBrO \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		BrO _t stretch	834.4T	Ar	IR	1

Reference

¹K. Johnsson, A. Engdahl, J. Kölöm, J. Nieminen, and B. Nelander, *J. Phys. Chem.* **99**, 3902 (1995).

IOClO

In an argon matrix, an absorption maximum at 18200 (550 nm) is associated¹ with isomerization to IOClO₂.

 \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1		ClO _t stretch	974.9	Ar	IR	1

Reference

¹K. Johnsson, A. Engdahl, J. Kölöm, J. Nieminen, and B. Nelander, *J. Phys. Chem.* **99**, 3902 (1995).

FSeSeF \tilde{X} C₂

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a	1	SeF s-stretch	610.1	Ar	IR	1
b	5	SeF a-stretch	588.2	Ar	IR	1
	6	Deformation	202.9	Ar	IR	1

Reference

¹A. Haas and H. Willner, *Z. Anorg. Allg. Chem.* **454**, 17 (1979).

SeSeF₂ \tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	SeF ₂ s-stretch	599	Ar	IR	1
	2	Se=Se stretch	404	Ar	IR	1
	4	Deformation	220.5	Ar	IR	1
a''	5	SeF ₂ a-stretch	570	Ar	IR	1

Reference

¹A. Haas and H. Willner, *Z. Anorg. Allg. Chem.* **454**, 17 (1979).

8.8. H₅⁺ Five-Atomic Tetra- and Trihydrides**TiH₄** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		TiH stretch	1663.8	Ar	IR	1,2
			1656.7			
			1657.8T	Kr	IR	1

TiD₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		TiD stretch	1205.6	Ar	IR	1,2
			1200.0			
			1201.1	Kr	IR	1

References

¹Z. L. Xiao, R. H. Hauge, and J. L. Margrave, *J. Phys. Chem.* **95**, 2696 (1991).

²G. V. Chertihin and L. Andrews, *J. Am. Chem. Soc.* **116**, 8322 (1994).

ZrH₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		ZrH stretch	1623.6	Ar	IR	1,2

ZrD₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		ZrD stretch	1166.6	Ar	IR	1,2

References

¹G. V. Chertihin and L. Andrews, *J. Am. Chem. Soc.* **117**, 6402 (1995).

²G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 15004 (1995).

HfH₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		HfH stretch	1678.4	Ar	IR	1,2

HfD₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		HfD stretch	1201.7	Ar	IR	1,2

References

¹G. V. Chertihin and L. Andrews, *J. Am. Chem. Soc.* **117**, 6402 (1995).

²G. V. Chertihin and L. Andrews, *J. Phys. Chem.* **99**, 15004 (1995).

ThH₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		ThH stretch	1444.8	Ar	IR	1
			1443.3			

ThD₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		ThD stretch	1032.1	Ar	IR	1
			1031.1			

Reference

¹P. F. Souter, G. P. Kushto, L. Andrews, and M. Neurock, *J. Phys. Chem. A* **101**, 1287 (1997).

UH₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		UH stretch	1483.6	Ar	IR	1
			1481.7			

UD₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		UD stretch	1060.7T	Ar	IR	1
			1059.8			

Reference

¹P. F. Souter, G. P. Kushto, L. Andrews, and M. Neurock, *J. Am. Chem. Soc.* **119**, 1682 (1997).

BH₄⁻

\bar{X}	T _d					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>t</i> ₂	3	BH stretch	2212.8	Ar	IR	1

Reference

¹T. J. Tague, Jr. and L. Andrews, J. Am. Chem. Soc. **116**, 4970 (1994).

AIH₄

\bar{X}	T _d					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>e</i>	2	Deformation	750T	Ar	IR	1
<i>f</i> ₂	3	AIH stretch	1609.3	Ar	IR	1
	4	Deformation	766.6	Ar	IR	1

AID₄⁻

\bar{X}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>e</i>	2	Deformation	535T	Ar	IR	1
<i>f</i> ₂	3	AID stretch	1173.9	Ar	IR	1
	4	Deformation	566.6	Ar	IR	1

Reference

¹P. Pullumbi, Y. Bouteiller, and L. Manceron, J. Chem. Phys. **101**, 3610 (1994).

NH₄⁺

\bar{X}	T _d	Structure: LD ^{1,4} CC ² TPE ⁸				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>f</i> ₂	3	NH stretch	3343.14	gas	LD,CC	1-3,6
	4	Deformation	1447.22	gas	DL	5,7

$B_0=5.929$ LD^{1,4}CC²DL⁷

ND₄⁺

\bar{X}	T _d					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>f</i> ₂	3	ND stretch	2495.0	gas	LD	4

$B_0=2.986(4)$ LD⁴TPE⁸

References

- ¹M. Crofton and T. Oka, J. Chem. Phys. **79**, 3157 (1983).
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- ⁵M. Polak, M. Gruebele, B. W. DeKock, and R. J. Saykally, Mol. Phys. **66**, 1193 (1989).
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- ⁸R. Signorell, H. Palm, and F. Merkt, J. Chem. Phys. **106**, 6523 (1997).

NH₄

$3p\ ^2F_2$	T _d			
$T_0 \approx 15078^{ab}$	gas	EM ^{1,2,4}		
	Diffuse.			
$3s\ ^2A_1$	T _d	Structure: TPE ⁸		
	gas		$3p^2F_2-3s^2A_1$	663.5 nm
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.
<i>a</i> ₁	1	NH stretch	2552T	gas
<i>e</i>	2	Deformation	1581T	gas
			EM	1,4
			EM	1,4

ND₄

$3p\ ^2F_2$	T _d			
$T_0^a=14828.285(4)$	gas	EM ^{1,2,4} AB ^{3,5}		
			$3p^2F_2-3s^2A_1$	675 nm
		Three weak bands have been observed ⁴ 775, 1138, and 1722 cm ⁻¹ above the band origin. However, the assignment of these bands has not yet been established.		
		$B_0=3.122$ gas AB ⁵		
$3s\ ^2A_1$	T _d			
	gas		$3p^2F_2-3s^2A_1$	675 nm
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.
<i>a</i> ₁	1	ND stretch	1960T	gas
<i>e</i>	2	Deformation	1080.25(7)	gas
			EM	1,4
			EM	7

$\tau \approx 30 \mu s$ gas AB⁵
 $B_0=2.856(4)$ gas TPE⁸

^aMeasured with respect to the lowest Rydberg state, $3s\ ^2A_1$. The ground state is dissociative.

^bEstimated⁵ by scaling of data for ND₄.

References

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HBeHBeH \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1493.4	Ar	IR	1
			1402.1	Ar	IR	1

Reference¹T. J. Tague, Jr. and L. Andrews, J. Am. Chem. Soc. **115**, 12111 (1993).**MgCH₃**

\tilde{A} ² E	C_{3v}	Structure: LF ¹
$T_0=20030.296(2)$	gas	LF ¹

\tilde{X} ² A ₁	C_{3v}	Structure: LF ¹ MW ²
$B_0=0.367$	LF ¹ MW ²	

References¹R. Rubino, J. M. Williamson, and T. A. Miller, J. Chem. Phys. **103**, 5964 (1995).²M. A. Anderson and L. M. Ziurys, Astrophys. J. **452**, L157 (1995).**CaCH₃**

\tilde{B} ² A ₁	C_{3v}	Structure: LF ¹
$T_0=16003(10)$	gas	LF ¹

 $\tilde{B}-\tilde{X}$ 620–630 nm

\tilde{A} ² E	C_{3v}	Structure: LF ^{1,2,4}
$T_0=14743.382^a$	gas	LF ^{1,2,4}

 $\tilde{A}-\tilde{X}$ 630–730 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CH ₃ umbrella	1048(10)	gas	LF	1
	3	CaC stretch	413(10)	gas	LF	1
e	6	CaCH deform.	391(5)H	gas	LF	1

$$A=72.71 \text{ gas } \text{LF}^{1,2,4}$$

$$A_0=5.386; B_0=0.254 \text{ LF}^{2,4}$$

\tilde{X} ² A ₁	C_{3v}	Structure: LF ²

$$A_0=5.448; B_0=0.252 \text{ LF}^{2,4}\text{MW}^3$$

^aPredisassociated above ~ 16200 .¹**References**¹C. R. Brazier and P. F. Bernath, J. Chem. Phys. **86**, 5918 (1987).²C. R. Brazier and P. F. Bernath, J. Chem. Phys. **91**, 4548 (1989).³M. A. Anderson and L. M. Ziurys, Astrophys. J. **460**, L77 (1996).⁴A. J. Marr, F. Grieman, and T. C. Steinle, J. Chem. Phys. **105**, 3930 (1996).**SrCH₃**

\tilde{B} ² A ₁	C_{3v}	gas	LF ¹	$\tilde{B}-\tilde{X}$ 670–680 nm

\tilde{A} ² E	C_{3v}	gas	LF ¹	$\tilde{A}-\tilde{X}$ 670–740 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CH ₃ umbrella	1054(10)	gas	LF	1
	3	SrC stretch	373(10)	gas	LF	1
e	6	SrCH deform.	342(5)H	gas	LF	1

$$A=273(20) \text{ gas } \text{LF}^1$$

\tilde{X} ² A ₁	C_{3v}	Structure: MW ²

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CH ₃ umbrella	1072(10)	gas	LF	1
	3	SrC stretch	362(10)	gas	LF	1
e	6	SrCH deform.	279(5)H	gas	LF	1

$$B_0=0.194 \text{ MW}^2$$

^aPredisassociated above ~ 15000 .¹**References**¹C. R. Brazier and P. F. Bernath, J. Chem. Phys. **86**, 5918 (1987).²M. A. Anderson, J. S. Robinson, and L. M. Ziurys, Chem. Phys. Lett. **257**, 471 (1996).**ZnCH₃**

\tilde{C} ² A ₁ ^a	C_{3v}	AB ¹	$\tilde{C}-\tilde{X}$ 260–274 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CH ₃ deform.	950T	gas	AB	1

\tilde{A} ² E	C_{3v}	Structure: LF ⁶
$T_0=24082.82$	gas	AB ¹ EM ² LF ^{5,6}

 $\tilde{A}-\tilde{X}$ 379–437 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	CH ₃ deform.	1048	gas	AB,LF	1,4,5
	3	ZnC stretch	460	gas	LF	5

$$\tau=40(3) \text{ ns } \text{gas } \text{EM}^3\text{LF}^4$$

$$A=253 \text{ gas } \text{AB}^1\text{EM}^2\text{LF}^4$$

$$A_0=4.949; B_0=0.317 \text{ LF}^6$$

References¹C. R. Brazier and P. F. Bernath, J. Chem. Phys. **86**, 5918 (1987).²C. R. Brazier and P. F. Bernath, J. Chem. Phys. **91**, 4548 (1989).³M. A. Anderson and L. M. Ziurys, Astrophys. J. **460**, L77 (1996).⁴A. J. Marr, F. Grieman, and T. C. Steinle, J. Chem. Phys. **105**, 3930 (1996).

\tilde{X}^2A_1		C_{3v}		Structure: LF ⁶			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a_1	2	CH ₃ deform.	1064	gas	EM,LF	3-5	
	3	ZnC stretch	445	gas	LF	4,5	
e	6	ZnCH deform.	315T	gas	LF	5	

$A_0=5.15$; $B_0=0.310$ LF⁶

^aAssignment to \tilde{C} state suggested⁴ by analogy with ZnH.

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CdCH₃

\tilde{C}^2A_1		C_{3v}		Structure: LF ⁸			
$T_0=34916$ gas AB ¹						$\tilde{C}-\tilde{X}$ 264–287 nm	
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a_1	2	CH ₃ deform.	960T	gas	AB	1	

\tilde{A}^2E		C_{3v}		Structure: LF ⁸			
$T_0=22835.22$ gas AB ¹ EM ²⁻⁵ LF ⁶⁻⁹						$\tilde{A}-\tilde{X}$ 400–458 nm	
Perturbations in the $^2E_{3/2}$ band origin suggest another nearby state of 2E symmetry. ⁸							
a_1	2	CH ₃ deform.	1019(3)	gas	LF	7,8	
	3	CdC stretch	400(3)	gas	LF	6-8	

$\tau=70(4)$ ns gas EM²⁻⁴LF⁷

$A=650$ gas LF⁸

$A_0=4.969$; $B_0=0.244$ LF⁸

\tilde{X}^2A_1		C_{3v}		Structure: LF ⁸			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a_1	2	CH ₃ deform.	1000(3)	gas	EM,LF	5-7	
	3	CdC stretch	355(3)	gas	EM,LF	5-7	

$A_0=5.157$; $B_0=0.234$ LF⁸

References

- P. J. Young, R. K. Gosavi, J. Connor, O. P. Strausz, and H. E. Gunning, *J. Chem. Phys.* **58**, 5280 (1973).
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H₂C=BH

\tilde{X}		C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a_1		BH stretch	2724.6	Ar	IR	1	
		C=B stretch	1469.7	Ar	IR	1	
b_1		H ₂ CB OPLA	705.7	Ar	IR	1	
		Torsion	611.8	Ar	IR	1	
b_2		HBC deform.	902.6	Ar	IR	1	

D₂C=BD

\tilde{X}		C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
a_1		BD stretch	2070.2	Ar	IR	1	
b_1		D ₂ CB OPLA	537.1	Ar	IR	1	
b_2		DBC deform.	739.1	Ar	IR	1	

Reference

- P. Hassanzadeh, Y. Hannachi, and L. Andrews, *J. Phys. Chem.* **97**, 6418 (1993).

AlCH₃

\tilde{X}^1A_1
 $B_0=0.393$ MW¹

Reference

- J. B. Robinson and I. M. Ziurys, *Astrophys. J.* **472**, L131 (1996).

C₂H₃⁺

\tilde{X} C_{2v} (bridged) Structure: LD^{1,2}

\tilde{X}		C _{2v} (bridged)		Structure: LD ^{1,2}			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
A_{1u}^+	6	CH a-stretch	3142.2	gas	LD	2,5	

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⁶M. Cordonnier and L. H. Coudert, *J. Mol. Spectrosc.* **178**, 59 (1996).

HBNH₂

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NH stretch	3598.4	Ar	IR	1
			3588.4			
		BH stretch	2418.2	Ar	IR	1
		BN stretch	1556.4	Ar	IR	1
		OPLA	608.5	Ar	IR	1

DBND₂

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		ND stretch	2685.9	Ar	IR	1
			2678.7			

Reference

¹C. A. Thompson, L. Andrews, J. M. L. Martin, and J. El-Yazal, *J. Phys. Chem.* **99**, 13839 (1995).

C₂H₃

Rydberg state

T₀=59410 gas AB³

164–169 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1306	gas	AB	3

\tilde{A}^2A'' C_s
T₀≤20020 gas AB¹

500–400 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'		CC stretch	1200	gas	AB	1
		CCH bend	920	gas	AB	1

\tilde{X} C^a

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''	7	Mixed OPLA	895.16	gas	DL	4
			895.4	Ne	IR	5
			900	Ar	IR	2

A₀=7.913; B₀=1.083; C₀=0.949 DL⁴

C₂D₃

\tilde{X}		C _s					
Vib. sym.		Approximate type of mode		cm ⁻¹	Med.	Type meas.	Refs.
a''	7	Mixed OPLA		704	Ar	IR	2

^aRapid tunneling, giving effective C_{2v} symmetry.⁴

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CH₃N

\tilde{A}^3E C_{3v} Structure: EM⁴
T₀=31830.913(12) gas AB¹, EM^{1,2,4,5,7} LF^{8,10} $\tilde{A}-\tilde{X}$ 288–356 nm
31576(20) N₂ AB^{3,6} $\tilde{A}-\tilde{X}$ 281–317 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	CH ₃ stretch	2943	gas	LF	8,10
	2	CH ₃ deform.	1239	gas	LF	8
			1166T	N ₂	AB	6
e	3	CN stretch	758(4)	gas	UV,LF	1,2,8
			755(22)	N ₂	AB	3,6
e	5	CH ₃ deform.	1500T ^a	gas	LF	8
	6	CH ₃ rock	728(4)	gas	EM	5

$\tau_0=415(8)$ ns gas LF^{9,10}
 $A=-22.52$ gas EM^{4,7}
 $A_0=5.423$; $B_0=0.845$ EM^{4,7}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	CH ₃ s-stretch	2943(4)	gas	EM	2,5
	2	CH ₃ deform.	1349(4)	gas	EM	2,5
	3	CN stretch	1040(4)	gas	EM,LF	2,5,10
e			1029	N ₂	AB	3
	4	CH ₃ a-stretch	2989(4)	gas	EM	5
	5	CH ₃ deform.	1490(4)	gas	EM	5
e	6	CH ₃ rock	903(8)	gas	EM	2,5

$A_0=5.61$; $B_0=0.929$ EM^{4,7}

CD₃N

\tilde{A} 3E		C_{3v}			$\tilde{A}-\tilde{X}$ 294–365 nm		
$T_0=31774.158(2)^b$		gas	$AB^1, EM^{2,4,5}$		$\tilde{A}-\tilde{X}$ 277–318 nm		
31516(30) N ₂		AB ^{3,6}					

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CD ₃ deform.	1044T	N ₂	AB	6
	3	CN stretch	759(4)	gas	UV	1,2
			695T	N ₂	AB	6
<i>e</i>		CD ₃ rock	579(4) ^c	gas	EM	2

 $B_0=0.691$ EM⁴ \tilde{X} 3A_2

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CD ₃ deform.	941(4)	gas	EM	5
	3	CN stretch	1110(4)	gas	EM	2,5
	6	CD ₃ rock	749(8)	gas	EM	2,5

 $B_0=0.744$ EM⁴^aFrom combination bands.^bCalculated assuming A(CD₃N)=A(CH₃N).^cObserved as sequence band.**References**

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SiH₂NH

In an argon matrix, an absorption maximum at 41700 (240 nm) has been assigned¹ to SiH₂NH.

\tilde{X}						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		SiH ₂ stretch	2253	Ar	IR	1
		SiH ₂ stretch	2175	Ar	IR	1
		Si=N stretch	1097	Ar	IR	1

SiD₂ND \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		SiD ₂ stretch	1638	Ar	IR	1
		SiD ₂ stretch	1585	Ar	IR	1
		Si=N stretch	1063	Ar	IR	1

Reference

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CH₂OH⁺

\tilde{X}		C_s	Structure: MW ⁵			
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OH stretch	3422.802	gas	LD	2
		CO stretch	1650(30)	gas	PE	1
		CH ₂ deform.	13'0(30)	gas	PE	1

 $A_0=6.591$; $B_0=1.146$; $C_0=0.973$ LD²MW^{3,4}**CD₂OD⁺**

\tilde{X} C_s
 $A_0=3.407$; $B_0=0.903$; $C_0=0.711$ MW⁵

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CH₃S⁺

\tilde{X} 3A_2		C_{3v}				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CS stretch	733(5)	gas	PI,TPE	1,2

CD₃S⁺

\tilde{X}		C_{3v}				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CS stretch	730(60)	gas	PI	1

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²C.-W. Hsu and C. Y. Ng, J. Chem. Phys. **101**, 5596 (1994).

CH₃O

\tilde{A}^2A_1 C_{3v} Structure: LF^{22,23}
 $T_0=31644.2$ gas EM^{1,2,11,12}AB⁵LF^{6,8,18,19,22-24,27-29,31} $\tilde{A}-\tilde{X}$ 271-421 nm
 $31291(3)$ Ar LF²¹ $\tilde{A}-\tilde{X}$ 270-420 nm

Evidence for predissociation above 35437 (3793 above the origin of the \tilde{A} state),^{13,29,32} with CH₃+O the principal products.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₃ stretch	2947.8	gas	LF	31
	2	Umbrella	1289.3	gas	LF	19,27,31
	3	CO stretch	1308(4)	Ar	LF	21
<i>e</i>	4	CH ₃ stretch	662.4	gas	AB,EM,LF	5,12,19,31
	5	CH ₂ scissors	657(2)	Ar	LF	21
<i>e</i>	4	CH ₃ stretch	3077.8T	gas	LF	31
	5	CH ₂ scissors	1403.0	gas	LF	19,31
	6	HCO deform.	1410(3)	Ar	LF	21
			929.5	gas	LF	31

$\tau=2.57(13)$ μ s gas EM^{2,9}LF^{4,14,15,16,18,20,32}
 $A_0=4.981(3)$; $B_0=0.743$ LF^{22-24,28}

\tilde{X}^2E C_{3v}^a Structure: LMR^{3,7}MW^{10,17}LF²²
High vibrational levels have been studied using SEP.^{25,28,30}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₃ stretch	2840T ^b	gas	LF	19,27
	2	CH ₃ umbrella	1412(4)T ^c	gas	LF	27
	3	CO stretch	1406(2)T	Ar	LF	21
<i>e</i>	4	CH ₃ stretch	1047	gas	LF,EM	6,9,12, 19,27,30
	5	CH ₂ scissors	1044(2)	Ar	LF	21
	6	HCO deform.	2778	gas	SEP	30
			2758(3)	Ar	LF	21
<i>e</i>	4	CH ₃ stretch	914	gas	SEP	30
	5	CH ₂ scissors	651.5	gas	LF,SEP	19,26,27,30
	6	HCO deform.				

$A=-61.97(7)$ gas LMR⁷MW¹⁰EM^{11,12}LF^{19,22,23,27}
 $A_0=5.206(4)$; $B_0=0.932$ LMR⁷MW^{10,17}LF^{22,23}

CD₃O

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CD ₃ stretch	2015	gas	LF	19
	2	CD ₃ umbrella	971	gas	LF	19
	3	CO stretch	663	gas	EM,LF	12,19
<i>e</i>	5	CD ₂ scissors	1047	gas	LF	19

\tilde{X}^2E		C _{3v} ^a				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	
<i>a</i> ₁	2	CO stretch	1000T	gas	LF	19
	3	CD ₃ umbrella	893T	gas	LF	19
<i>e</i>	5	CD ₂ scissors	1174T	gas	LF,EM	6,12,19
	6	DCO deform.	496	gas	LF	19

$A=-56(2)$ gas EM¹²
 $B_0=0.740$ MW¹⁷

^aSomewhat distorted by Jahn-Teller coupling.

^bRef. 30 indicates that reassignment is necessary.

^cRef. 30 favors assignment at 1359.

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CH₂OH

$\tilde{B}^2A'(3p)$		C_s	$\tilde{B}-\tilde{X}$ 217–244 nm		
$T_0=41065(3)$	gas	MPI ^{3,4,7} AB ⁵			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	4	CO stretch	1621(7)	gas	MPI 3,4,7
	5	CH ₂ scissors	1465(7)	gas	MPI 3,7
	6	HCOH deform.	1357(7)	gas	MPI 3,7
	7	HCOH deform.	1107(7)	gas	MPI 3,7
a''	9	Torsion	993(8)	gas	MPI 7

$\tilde{A}^2A'(3s)$		C_s	$\tilde{A}-\tilde{X}$ 243–285 nm		
$T_0=35050$	gas	AB ⁵			
Threshold for photodecomposition into H ₂ CO+H near 280 nm. ^{1,2}					

\tilde{X}^2A''		C_s^a			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	1	OH stretch	3650m	Ar	IR 1,2
			3637m	N ₂	IR 1
	4	CH ₂ scissors	1459w	Ar	IR 2
	5	OH deform.	1334m	Ar	IR 1,2
	6	CO stretch	1176(7)	gas	MPI 7
			1183vs	Ar	IR 1,2
			1183s	N ₂	IR 1
	7	HCOH deform.	1048s	Ar	IR 1,2
			1056m	N ₂	IR 1
a''	8	Torsion	420m	Ar	IR 1,2
			482m	N ₂	IR 1
	9	H ₂ CO OPLA	234(5)	gas	MPI 7

CD₂OD

$\tilde{B}^2A'(3p)$		C_s	$\tilde{B}-\tilde{X}$ 216–244 nm		
$T_0=40913$	gas	MPI ^{3,7} AB ⁶			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	4	CO stretch	1565	gas	MPI,AB 3,6,7
	5	CD ₂ scissors	1109	gas	MPI,AB 3,6,7
a''	8	D ₂ CO OPLA	952(23)	gas	MPI 7
	9	Torsion	723(23)	gas	MPI 7

$\tilde{A}^2A'(3s)$		C_s	$\tilde{A}-\tilde{X}$ 350–530 nm		
$T_0=35124$	gas	AB ⁶			
Threshold for photodecomposition into D ₂ CO+D near 280 nm. ^{1,2}					

\tilde{X}^2A''		C_s^a			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
a'	1	OD stretch	2694wm	Ar	IR 2
			2682m	N ₂	IR 1
	4	CO stretch	1221(4)	gas	MPI 7
			1223m	Ar	IR 2
			1222m	N ₂	IR 1
	5	CD ₂ scissors	1020(5)	gas	MPI 7
			1041m	Ar	IR 2
a''	7		765wm	Ar	IR 2
	8	Torsion	329(23)	gas	MPI 7
	9	D ₂ CO OPLA	177(23)	gas	MPI 7

^a Rapid interconversion of nonplanar forms.⁷

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CH₃S

A diffuse, unstructured absorption at 218.5 nm, observed on flash photolysis¹ of a number of sulfur-containing compounds or on pulse radiolysis¹² of CH₃SH in the presence of an F-atom source, has been attributed to CH₃S.

\tilde{A}^2A_1		C_{3v}	Structure: LF ⁹	$\tilde{A}-\tilde{X}$ 350–530 nm		
$T_0=26396.8$	gas	EM ² LF ^{6,9,11}	Predissociation threshold ≥ 27885 . ¹¹ In an argon matrix, CH ₂ SH is formed. ⁵			

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	CH ₃ umbrella	1098(2)	gas	LF	11
	3	CS stretch	401(2)	gas	EM,LF	2,6,9,11
e	6	HCS deform.	635(10)	gas	LF	11

$T_0=760(60)$ ns gas LF⁸; 1130(70) ns gas LF^{10,11}
 $A_0=5.343(47)$; $B_0=0.346$ LF⁹

\tilde{X}^2E		C_{3v}	Structure: MW ⁷ LF ⁹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CH ₃ stretch	2776T	gas	LF	11
	2	CH ₃ umbrella	1313(5)	gas	PD,LF	4,6,11
	3	CS stretch	727(3)	gas	EM,PE	2–4,6
					PD,LF	11
e	4	CH ₃ stretch	2706T	gas	LF	11
	5	CH ₃ deform.	1496(6)	gas	LF	11
	6	HCS deform.	586T	gas	LF	11

$A=-259.1$ gas LF^{9,11}TPE¹³
 $A_0=5.68(4)$; $B_0=0.450$ MW⁷LF⁹

CD₃S

\tilde{A}^2A_1 C_{3v}
 $T_0=26574$ gas LF⁶
 $\tilde{A}-\tilde{X}$ 352–378 nm
 Predissociation threshold ≤ 27728 .

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	CD ₃ umbrella	837(1)	gas	LF	6
	3	CS stretch	395(1)	gas	LF	6

$\tau_0=0.45(11)$ μ s gas LF⁶

 \tilde{X}^2E C_{3v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	CD ₃ umbrella	1100(50)	gas	PD	4
	3	CS stretch	667(1)	gas	PD,LF	4,6
e	5	DCS deform.	780(30)H	gas	PD	4

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H₂CFH⁺ \tilde{X} C₁

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	FH stretch	3308.9	Ne	IR	1
			3291.0			
7		HF wag	921.2	Ne	IR	1
8		CF stretch	714.9	Ne	IR	1

D₂CFD⁺ \tilde{X} C₁

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1		FD stretch	2442.2	Ne	IR	1
			2429.4			

Reference

¹C. L. Lugez, D. Forney, M. E. Jacox, and K. K. Irikura, J. Chem. Phys. **106**, 489 (1997).

CH₃Cl⁺

\tilde{C}^2A_1 C_{3v}
 $T^a=82400(900)$ gas PE²

\tilde{B}^2E C_{3v}

$T^a=33170(900)$ gas PE^{1,4,7}

Position of first maximum is given. A Jahn-Teller splitting of ~ 5000 is observed.^{4,7}

A weak, broad absorption with onset near 400 nm (25000) and maximum at 335 nm (29800) which appears on argon-resonance photolysis of CH₃Cl isolated in an argon matrix and which is destroyed by exposure of the sample to 290–1000 nm radiation has been attributed⁶ to the $\tilde{B}-\tilde{X}$ and $\tilde{A}-\tilde{X}$ transitions of CH₃Cl⁺.

\tilde{A}^2A_1 C_{3v}
 $T_0=20260(900)$ gas PE^{1,4,7}

 \tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH ₃ deform.	1550(50)	gas	PE	3,4
			1503.3w	Ne	IR	8
		CH ₃ deform.	1239.5wm	Ne	IR	8
		CH ₃ umbrella	1073(50)	gas	PE	2,4
		CH ₃ rock	870(50)	gas	PE	3,4
		CH ₃ deform.	649.4wm	Ne	IR	8
			626.9wm			
		CCl stretch	654(50)	gas	PE	3,4

CD₃Cl⁺ \tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CD ₃ stretch	2013.4m	Ne	IR	8
		CD ₃ deform.	975.2w	Ne	IR	8
		CD ₃ deform.	927.9wm	Ne	IR	8
		CCl stretch	628.9wm	Ne	IR	8

^aFrom vertical ionization potential.

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H₂CCIH⁺

\tilde{X}	C ₁					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	CH ₂	a-stretch	3225.5wm	Ne	IR	1
2	CH ₂	s-stretch	3053.3m	Ne	IR	1
3	CIH	stretch	2626.1m	Ne	IR	1
7	CCl	stretch	798.8vw	Ne	IR	1
8		Wag	655.0wm	Ne	IR	1

D₂CCID⁺

\tilde{X}	C ₁					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	CD ₂	a-stretch	2444.2m	Ne	IR	1
2	CD ₂	s-stretch	2219.4w	Ne	IR	1
3	CID	stretch	1909.9m	Ne	IR	1
7	CCl	stretch	575.6wm	Ne	IR	1

Reference

¹C. L. Lugez, D. Forney, M. E. Jacox, and K. K. Irikura, J. Chem. Phys., **106**, 489 (1997).

CH₃Br⁺

\tilde{C} ²A₁ C_{3v}
 $T^a=75500(900)$ gas PE¹

\tilde{B} ²E C_{3v}
 $T_0=31930(900)$ gas PE^{1,3}

A Jahn-Teller splitting of ~5600 is observed.¹⁻³ (Onset of the transition is given.)

A broad absorption with maximum at 267 nm (37400) which appears on argon-resonance photolysis of CH₃Br isolated in an argon matrix and which has a photodecomposition threshold at a wavelength longer than 500 nm has been assigned⁵ to the $\tilde{B}-\tilde{X}$ transition of CH₃Br⁺.

\tilde{A} ²A₁ C_{3v}
 $T_0=19820(900)$ gas PE¹⁻³

A broad absorption with maximum at 318 nm (28700) which appears on argon-resonance photolysis of CH₃Br isolated in an argon matrix and which has a photodecomposition threshold at a wavelength longer than 500 nm has been assigned⁵ to the $\tilde{A}-\tilde{X}$ transition of CH₃Br⁺.

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CH stretch	3130(100)	gas	PE	2,3
			3121.9wm	Ne	IR	6
	2	CH stretch	2930(120)	gas	PE	3
			2909.5s	Ne	IR	6
	3	Deformation	1417.3wm	Ne	IR	6
			1412.5wm			
	4	Deformation	1290(80)	gas	PE	1-3
			1296.7m	Ne	IR	6
	5	Deformation	850(80)	gas	PE	1-3
a''			815.0w	Ne	IR	6
	6	CBr stretch	468(80)	gas	PE	3
			549.7wm	Ne	IR	6
			537.7wm			
a''	7	CH ₂ a-stretch	3055.3m	Ne	IR	6
	8	Deformation	1276.8wm	Ne	IR	6
	9	Deformation	650.8m	Ne	IR	6

CD₃Br⁺

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CD stretch	2165(80)	gas	PE	2
	2	CD stretch	2091.6ms	Ne	IR	6
	4	Deformation	947(80)	gas	PE	2
			948.2wm	Ne	IR	6
a''	7	CD ₂ a-stretch	2128.1m	Ne	IR	6
	8	Deformation	974.7ms	Ne	IR	6
	9	Deformation	491.8m	Ne	IR	6

^aFrom vertical ionization potential.

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H₂CBrH⁺

\tilde{X}	C ₁					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	CH ₂ a-stretch	3221.4w	Ne	IR	1	
2	CH ₂ s-stretch	2979.3wm	Ne	IR	1	
3	BrH stretch	2395.6wm	Ne	IR	1	

D₂CBrD⁺

\tilde{X}	C ₁					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	CD ₂	a-stretch	2440.3wm	Ne	IR	1
3		BrD stretch	1727.1wm	Ne	IR	1

Reference

¹C. L. Lugez, D. Forney, M. E. Jacox, and K. K. Irikura, J. Chem. Phys. **106**, 489 (1997).

H₂PSH

\tilde{X}	C _{∞v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		SH stretch	2563.9	Ar	IR	1
		PH ₂ wag	967.5	Ar	IR	1
		P-S stretch	484.1	Ar	IR	1

Reference

¹Z. Mielke and L. Andrews, J. Phys. Chem. **97**, 4313 (1993).

8.9. Five-Atomic Dihydrides**HGa₃H**

\tilde{X}	D _{∞h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		GaH stretch	1765.8	Ar	IR	1
			1753.2	Kr	IR	1
		GaH stretch	1752.8	Ar	IR	1
			1740.4	Kr	IR	1

DGa₃D

\tilde{X}	D _{∞h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		GaD stretch	1276.2	Ar	IR	1
			1267.2	Kr	IR	1
		GaD stretch	1262.9	Ar	IR	1
			1253.3	Kr	IR	1

Reference

¹Z. L. Xiao, R. H. Hauge, and J. L. Margrave, Inorg. Chem. **32**, 642 (1993).

HBeCCH

\tilde{X}	C _{∞v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	2	BeH stretch	2119.4	Ar	IR	1
	4	BeC stretch	838.9	Ar	IR	1
Π	5	HCC bend	689.1	Ar	IR	1
	6	HIDeC bend	550.0	Ar	IR	1

DBeCCD

\tilde{X}	C _{∞v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	2	BeD stretch	1535.0	Ar	IR	1
	4	BeC stretch	784.9	Ar	IR	1
Π	6	DBeC bend	450.0	Ar	IR	1

Reference

¹C. A. Thompson and L. Andrews, J. Am. Chem. Soc. **118**, 10242 (1996).

HBCBH

\tilde{X}	D _{∞h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	4	BCB a-stretch	1872.0	Ar	IR	1,2

DBCBD

\tilde{X}	D _{∞h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	BD a-stretch	2190.9	Ar	IR	1,2
	4	BCB a-stretch	1727.4	Ar	IR	1,2

References

¹P. Hassanzadeh and L. Andrews, J. Am. Chem. Soc. **114**, 9239 (1992).

²P. Hassanzadeh, Y. Hannachi, and L. Andrews, J. Phys. Chem. **97**, 6418 (1993).

HBeOBeH

\tilde{X}	D _{∞h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		BeH stretch	2092.4	Ar	IR	1
		BeOBe a-stretch	1403.1	Ar	IR	1
		Deformation	578.7	Ar	IR	1

DBeOBeD \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BeD stretch	1642.7	Ar	IR	1
		BeOBe a-stretch	1279	Ar	IR	1
		Deformation	481	Ar	IR	1

Reference

¹C. A. Thompson and L. Andrews, *J. Phys. Chem.* **100**, 12214 (1996).

cyc-HC=CHB $\tilde{X}^2\text{A}_1$ C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BC ₂ s-stretch	1170.6	Ar	IR	1,2
		HCC deform.	896.7	Ar	IR	2

cyc-DC=CDB $\tilde{X}^2\text{A}_1$ C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BC ₂ s-stretch	1169.4	Ar	IR	1,2
		DCC deform.	771.4	Ar	IR	2

References

¹J. M. L. Martin, P. R. Taylor, P. Hassanzadeh, and L. Andrews, *J. Am. Chem. Soc.* **115**, 2510 (1993).

²L. Andrews, P. Hassanzadeh, J. M. L. Martin, and P. R. Taylor, *J. Phys. Chem.* **97**, 5839 (1993).

cyc-HC=CBH $\tilde{X}^2\text{A}'$ C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BH stretch	2651.4	Ar	IR	1
		Ring stretch	1119.7	Ar	IR	1

cyc-DC=CBD $\tilde{X}^2\text{A}'$ C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BD stretch	2004.0	Ar	IR	1

Reference

¹L. Andrews, P. Hassanzadeh, J. M. L. Martin, and P. R. Taylor, *J. Phys. Chem.* **97**, 5839 (1993).

HCCBH $\tilde{X}^2\text{A}'$ C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a'</i>	3	C≡C stretch	2080.4	Ar	IR	1

DCCBD $\tilde{X}^2\text{A}'$ C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a'</i>		C≡C stretch	1940.3	Ar	IR	1

Reference

¹L. Andrews, P. Hassanzadeh, J. M. L. Martin, and P. R. Taylor, *J. Phys. Chem.* **97**, 5839 (1993).

cyc-HC=CHAI $\tilde{X}^2\text{A}_1$ C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		HCC deform.	609	Ar	IR	1,2

References

¹T. R. Burkholder and L. Andrews, *Inorg. Chem.* **32**, 2491 (1993).

²G. V. Chertihin, L. Andrews, and P. R. Taylor, *J. Am. Chem. Soc.* **116**, 3513 (1994).

HCCAIH \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		C≡C stretch	2000.8	Ar	IR	1,2
		AlH stretch	1804.1	Ar	IR	1,2

DCCAID \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		C≡C stretch	1882.7	Ar	IR	1
		AlD stretch	1314.8	Ar	IR	1

References

¹T. R. Burkholder and L. Andrews, *Inorg. Chem.* **32**, 2491 (1993).

²G. V. Chertihin, L. Andrews, and P. R. Taylor, *J. Am. Chem. Soc.* **116**, 3513 (1994).

(C₂H₂)Al

\tilde{X} ² B ₂		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		HCC deform.	569.0	Ar	IR	1,2
		AlC stretch	424.8	Ar	IR	1,2

(C₂D₂)Al

\tilde{X} ² B ₂		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		DCC deform.	429.7	Ar	IR	1
		AlC stretch	405.8	Ar	IR	1

References

- ¹T. R. Burkholder and L. Andrews, Inorg. Chem. **32**, 2491 (1993).
²G. V. Chertihin, L. Andrews, and P. R. Taylor, J. Am. Chem. Soc. **116**, 3513 (1994).

(C₂H₂)Ga

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH stretch	2992	Ar	IR	1
		CC stretch	1682	Ar	IR	1
		HCC deform.	595	Ar	IR	1
		HCC deform.	566	Ar	IR	1
		GaC stretch	408	Ar	IR	1

(C₂D₂)Ga

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CD stretch	2242	Ar	IR	1
		CC stretch	1563	Ar	IR	1
		DCC deform.	486	Ar	IR	1
		DCC deform.	426	Ar	IR	1
		GaC stretch	375	Ar	IR	1

Reference

- ¹T. R. Burkholder and L. Andrews, Inorg. Chem. **32**, 2491 (1993).

(C₂H₂)In

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH stretch	3062	Ar	IR	1
		CC stretch	1668	Ar	IR	1
		HCC deform.	595	Ar	IR	1
		HCC deform.	544	Ar	IR	1
		InC stretch	418	Ar	IR	1

(C₂D₂)In

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CD stretch	2312	Ar	IR	1
		CC stretch	1547	Ar	IR	1
		DCC deform.	436	Ar	IR	1
		DCC deform.	409	Ar	IR	1
		InC stretch	387	Ar	IR	1

Reference

- ¹T. R. Burkholder and L. Andrews, Inorg. Chem. **32**, 241 (1993).

H₂C=C=C:

Photoisomerization to HCCCH occurs at 254 nm.^{1,2}

\tilde{C} ¹ A ₁	$T_0=38650(160)$	\tilde{C} ¹ A ₁	C_{2v}	Ar	AB^8	$\tilde{C}-\tilde{X}$ 213–259 nm
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1		C ₃ s-stretch	900T	Ar	AB	8

\tilde{B} ¹ B ₁	$T_0=18720(80)$	\tilde{B} ¹ B ₁	C_{2v}	Ar	AB^8	$\tilde{B}-\tilde{X}$ 382–535 nm
In an argon matrix, randomization of carbon-13 labelling occurs on irradiation at wavelengths longer than 22500 (444 nm). ⁷						

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1		C ₃ s-stretch	1000T	Ar	AB	8

$\tilde{A}^1 A_2$ C_{2v}
 $T_0 \leq 13960$ Ar AB⁸

$\tilde{a}^3 B_1$ C_{2v}
 $T_0 = 10390(70)$ gas PE⁶

$\tilde{X}^1 A_1$ C_{2v} Structure: MW⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₂ s-stretch	3059.6	Ar	IR	1,2
			3049.5			
	2	C ₃ a-stretch	1963.2	Ar	IR	1,2
<i>b</i> ₁	3	CH ₂ scissors	1449.3	Ar	IR	1,2
			1446.9			
	5	H ₂ CC OPLA	1003.0	Ar	IR	1,2
<i>b</i> ₂			999.2			
	8	CH ₂ rock	1025.0	Ar	IR	1

A₀=9.633; B₀=0.353; C₀=0.340 MW^{3,5}

D₂C=C=C:

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CD ₂ s-stretch	2212.5	Ar	IR	1
			2200.5			
	2	C ₃ a-stretch	1944.4	Ar	IR	1
<i>b</i> ₁			1933.4			
	3	CD ₂ scissors + C ₃ s-stretch	1208.7	Ar	IR	1
	4	C ₃ s-stretch + CD ₂ scissors	950.8	Ar	IR	1
	5	D ₂ CC OPLA	803.2	Ar	IR	1
<i>b</i> ₂			800.3			
	8	CD ₂ rock	832.6	Ar	IR	1
			829.2			

A₀=4.842; B₀=0.314; C₀=0.294 MW⁴

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HCCCH

In an argon matrix, a broad absorption near 31750 (315 nm) which shows some fine structure and bands between 38460 and 47600 (210–260 nm) have been attributed⁸ to triplet HCCCH. Prolonged 313-nm irradiation of

HCCCH isolated in solid argon leads both to its isomerization to H₂C=C=C, which has a singlet ground state,^{5,9} and to the randomization of carbon-13 substitution in HCCCH.⁹

An absorption band system of singlet HCCCH has also been identified² in the 310–370-nm spectral region, but a detailed analysis has not been reported. Beyond 28900 the bands are diffuse, and a continuous absorption is superposed on the short wavelength end of the band system. The molecule is linear in at least one of the states of the transition. A progression involving an upper-state vibrational frequency of 1094 has been tentatively identified. The preliminary analysis found B'=0.310 and B''=0.324.

$\tilde{X}^1 A$ C₂ Structure: ESR, IR⁸

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
7		CH stretch	3266.0s	Ar	IR	5–8
			3285	Kr	IR	3
		C ₃ a-stretch	1619.4w	Ar	IR	6–8
		CCH s-bend	550.4m	Ar	IR	6,8
			402.6	Ar	IR	4–8
			401.5wm			
			408	Kr	IR	3
		HCC wag	248.5	Ar	IR	4–8
			245.9s			
			258	Kr	IR	3

DCCCCD

$\tilde{X}^3 A$ C₂

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
7		CD stretch	2457.9s	Ar	IR	4,6,7
			2482	Kr	IR	3
		C ₃ a-stretch	1529.0w	Ar	IR	6,7
		CCD, C ₃ bend	384.0m	Ar	IR	6,7
			392	Kr	IR	3
		DCC wag	171m	Ar	IR	4

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cyc-HC=CHSi

In an argon matrix, cyc-HC=CHSi shows a weak, broad absorption between 260 and 320 nm, with maximum at 286 nm.^{1,3} Irradiation in this spectral region results in rearrangement to form HCCSiH.

\tilde{X}	C_{2v}		Structure: MW ²			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CH stretch	3048.7wm	Ar	IR	1,3
	3	CH deform.	875.3m	Ar	IR	1,3
	4	SiC stretch	761.9ms	Ar	IR	1,3
b_1	6	CH deform.	677.2vs	Ar	IR	1,3
b_2	7	CH stretch	3026.5wm	Ar	IR	1,3
	8	CH deform.	1085.8s	Ar	IR	1,3
	9	SiC stretch	672.1s	Ar	IR	1,3

$A_0=1.118$; $B_0=0.397$; $C_0=0.292$ MW²

cyc-DC=CDSi

\tilde{X}	C_{2v}		Structure: MW ²			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CD stretch	2313.6w	Ar	IR	3
	4	SiC stretch	769.1vs	Ar	IR	3
b_1	6	CD deform.	518.4s	Ar	IR	3
b_2	7	CD stretch	2248.7w	Ar	IR	3
	8	CD deform.	938.6wm	Ar	IR	3
	9	SiC stretch	583.9s	Ar	IR	3

$A_0=0.876$; $B_0=0.355$; $C_0=0.252$ MW²

References

- G. Maier, H. P. Reisenauer, and H. Pacl, Angew. Chem. **106**, 1347 (1994); Angew. Chem. Int. Ed. Engl. **33**, 1248 (1994).
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HCCSiH

In an argon matrix, a broad absorption maximum at 500 nm is associated with isomerization to cyc-HC=CHSi and to H₂C=C=Si:^{1,2}

\tilde{X}	C_s		Structure: MW ²			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CH stretch	3304.2m	Ar	IR	1,2
	2	C=C stretch	1995.7m	Ar	IR	1,2
	3	SiH stretch	1969.9vs	Ar	IR	1,2
	4	SiH deform.	814.8ms	Ar	IR	1,2
	5	CH deform.	613.9m	Ar	IR	1,2
	6	SiC stretch	605.0m	Ar	IR	1,2
	8	CH deform.	722.8wm	Ar	IR	1,2

DCCSiD

\tilde{X}	C_s		Structure: MW ²			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CD stretch	2570.1w	Ar	IR	2
	2	C≡C stretch	1879.7ms	Ar	IR	2
	3	SiD stretch	1435.3vs	Ar	IR	2
	4	SiD deform.	641.6s	Ar	IR	2
	5	SiC stretch	575.5wm	Ar	IR	2
a''	8	CD deform.	479.9wm	Ar	IR	2

References

- G. Maier, H. P. Reisenauer, and H. Pacl, Angew. Chem. **106**, 1347 (1994); Angew. Chem. Int. Ed. Engl. **33**, 1248 (1994).
- G. Maier, H. Pacl, H. P. Reisenauer, A. Meudt, and R. Janoschek, J. Am. Chem. Soc. **117**, 12712 (1995).

$H_2C=C=Si$:

In an argon matrix, absorption maxima at 340, 325, and 310 nm have been assigned^{1,2} to $H_2C=C=Si$: Irradiation at 340 nm leads to isomerization to HCCSiH.

C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	C=C stretch	1667.9s	Ar	IR	1,2
b_1	5	CH deform.	957.7s	Ar	IR	1,2

$D_2C=C=Si$:

C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	C=C stretch	1648.3	Ar	IR	2

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cyc-H₂SiCC

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	SiH ₂ stretch	2214.4ms	Ar	IR	1,2
	2	CC stretch	1769.8w	Ar	IR	1,2
	3	SiH ₂ scissors	1023.1vs	Ar	IR	1,2
	4	SiC ₂ s-stretch	836.5ms	Ar	IR	1,2
b_1	6	SiH ₂ stretch	2228.9ms	Ar	IR	1,2
	7	SiH deform.	676.4wm	Ar	IR	2
b_2	8	SiH deform.	757.4m	Ar	IR	2

cyc-D₂SiCC

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	SiD ₂ stretch	1601.9m	Ar	IR	2
	2	CC stretch	1771.1w	Ar	IR	2
	3	SiC ₂ s-stretch	842.3vs	Ar	IR	2
	4	SiD ₂ scissors	722.2wm	Ar	IR	2
<i>b</i> ₁	6	SiD ₂ stretch	1626.3m	Ar	IR	2
	7	SiD deform.	500.5wm	Ar	IR	2
<i>b</i> ₂	8	SiD deform.	604.2wm	Ar	IR	2

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- ¹G. Maier, H. P. Reisenauer, and H. Pacl, Angew. Chem. **106**, 1347 (1994); Angew. Chem. Int. Ed. Engl. **33**, 1248 (1994).
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H₂C=C=C:-

Threshold for electron detachment from ground-state H₂C-C-C:- = 14470(65) gas PE^{1,2}

² A ₁		C _{2v}	
(Dipole-Bound State)		T ₀ =14284.420(5) gas PD ³	

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	C ₃ a-stretch	1965	gas	PD	4
	4	C ₃ s-stretch	1117	gas	PD	4
<i>b</i> ₁	6	H ₂ CC OPLA	221.45	gas	PD	4
	9		276.69	gas	PD	4

$$A_0=9.651; \frac{1}{2}(B_0+C_0)=0.346 \text{ PD}^3$$

 \tilde{X} ²B₁

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> ₁	6	H ₂ CC OPLA	385.43	gas	PD	4
	9		309.24	gas	PD	4

$$A_0=9.731; B_0=0.344; C_0=0.332 \text{ PD}^3$$

References

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H₂CCN

\tilde{X} ² B ₁		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> ₁	5	H ₂ CC OPLA	663.79	gas	PE, DL	1,3

$$A_0=9.53; B_0=0.342; C_0=0.329 \text{ MW}^2$$

D₂CCN

\tilde{X} ² B ₁		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> ₁	5	D ₂ CC OPLA	538 ^a	gas	PE	1

^a From computer fit.

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H₂CCO⁺

\tilde{E} ² A ₁		C _{2v}				
T ₀	68343(11)	gas	PE ^{2,4}			
<i>a</i> ₁	3	CH ₂ scissors	1177(4) ^a	gas	PE	2,4
			671(4) ^a	gas	PE	4
			397(4) ^a	gas	PE	4

 \tilde{D} ²A₁

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			1488(4) ^a	gas	PE	4
			1164(4) ^a	gas	PE	4
			709(4) ^a	gas	PE	4

 \tilde{C} ²B₂

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2		1697(4) ^a	gas	PE	4
	3	CH ₂ scissors	1087(4) ^a	gas	PE	1,2,4

\tilde{B}^2B_1 C_{2v}
 $T_0=40259(10)$ gas PE¹⁻⁴

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CCO a-stretch	2001(4) ^a	gas	PE	4
	4	CCO s-stretch	1001(4) ^a	gas	PE	1,2,4
			347(4)	gas	PE	4

\tilde{A}^2B_2 C_{2v}
 $T_0=33706(6)$ gas PE¹⁻⁴

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			1528(4) ^a	gas	PE	4
			1164(4) ^a	gas	PE	4
			690(4)	gas	PE	4
<i>b</i> ₁		OPLA	461(4)	gas	PE	4

\tilde{X}^2B_1 C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CCO a-stretch	2263(4) ^a	gas	PE	2-4
	3		1379(4)	gas	PE	4
	4		1227(4) ^a	gas	PE	4

D₂CCO⁺

\tilde{E}^2A_1 C_{2v}
 $T_0=68261(11)$ gas PE^{2,4}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			1133(4) ^a	gas	PE	4
			834(4)	gas	PE	2,4

\tilde{D}^2A_1 C_{2v}
 $T_0=54117T$ gas PE^{2,4}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			1267(4)	gas	PE	4
			1204(4) ^a	gas	PE	4
			1169(4) ^a	gas	PE	4

\tilde{C}^2B_2 C_{2v}
 $T_0=52222(3)$ gas PE¹⁻⁴

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			1102(4)	gas	PE	4
			800(4) ^a	gas	PE	2,4

\tilde{B}^2B_1 C_{2v}
 $T_0=39383(8)$ gas PE²⁻⁴

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			1732(4)	gas	PE	4
			1056(4) ^a	gas	PE	4
			848(4)	gas	PE	4

\tilde{A}^2B_2 C_{2v}
 $T_0=34077(9)$ gas PE²⁻⁴

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			2140(4) ^a	gas	PE	4
			1007(4) ^a	gas	PE	4
			106(4)	gas	PE	4
<i>b</i> ₁		OPLA	299(4)	gas	PE	4

\tilde{X}^2B_1 C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			2286(4) ^a	gas	PE	2-4
			1091(4) ^a	gas	PE	2-4
			925(4) ^a	gas	PE	4

^a ω .

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Be(OH)₂

\tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OH stretch	3805.9	Ar	IR	1
		OBeO a-stretch	1493.7	Ar	IR	1
		Deformation	456	Ar	IR	1

Be(OD)₂

\tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OD stretch	2805.4	Ar	IR	1
		OBeO a-stretch	1477.3	Ar	IR	1

Reference

- C. A. Thompson and L. Andrews, J. Phys. Chem. 100, 12214 (1996).

Mg(OH)₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		MgO a-stretch	866.9	Ar	IR	1

Reference¹T. J. Tague, Jr. and L. Andrews, *J. Phys. Chem.* **98**, 8611 (1994).**HN=C=NH** \tilde{X} C₂

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i>	2	NCN s-stretch	1285(20) ^a	gas	IR	2
			1275 ^a	Ar	IR	1
<i>b</i>	5	NCN deform.	537m	Ar	IR	1
	7	NCN a-stretch	2104.7	gas	IR	2
			2097s	Ar	IR	1
<i>a</i>	8	NH deform.	890(10)	gas	IR	2
			886vs	Ar	IR	1

 $A_0=12.650$; $B_0=0.346$; $C_0=0.346$ IR,MW³⁻⁵**DN=C=ND** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i>	1	ND stretch	2545s ^b	Ar	IR	1
	5	NCN deform.	471m	Ar	IR	1
<i>b</i>	6	ND stretch	2545s ^b	Ar	IR	1
	7	NCN a-stretch	2107vs	Ar	IR	1
	8	ND deform.	752s	Ar	IR	1

^aCalculated from ($\nu_2 + \nu_8$) combination band.^bBoth ND-stretching frequencies presumed equal.**References**

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HCNNH

In an argon matrix, an absorption maximum at 41700 (240 nm) has been assigned¹ to HCNNH.

 \tilde{X} C₁

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
1		NH stretch	3250.1w	Ar	IR	1
2		CH stretch	3142.5w	Ar	IR	1
3		CNN a-stretch	2032.7vs	Ar	IR	1
4		HNN bend	1278.1wm	Ar	IR	1
5		CNN s-stretch	1187.5wm	Ar	IR	1
6		Torsion	792.1w	Ar	IR	1
7		HCN bend	606.5m	Ar	IR	1
8		CNN bend	461.4w	Ar	IR	1

DCNND \tilde{X} C₁

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i>	2	CD stretch	2398.1wm	Ar	IR	1
	3	CNN a-stretch	1958.4vs	Ar	IR	1
<i>b</i>			1953.1vs			
	5	CNN s-stretch	984.5wm	Ar	IR	1
<i>c</i>	7	DCN bend	569.4wm	Ar	IR	1
	8	CNN bend	425.7m	Ar	IR	1
			418.0m			

Reference¹G. Maier, J. Eckwert, A. Bothur, H. P. Reisenauer, and C. Schmidt, *Liebigs Ann.* **1996**, 1041.**H₂CCS** $\tilde{A}'^1\text{A}''$ C_s $T_0 \approx 17995$ gas AB¹⁰ $\tilde{A}-\tilde{X}$ 450–550 nm

Diffuse bands.

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a'</i>	2	C=C stretch	1402	gas	AB	10
	8	CCS bend	284T	gas	AB	10

\tilde{X}^1A_1		C_{2v}	Structure: MW ²⁻⁵ IR ^{9,11}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CH ₂ s-stretch	3020	gas	IR	9
			3012w	Ar	IR	1,6,7
	2	C=C stretch	1757	gas	IR	9
			1755vs	Ar	IR	1,6-8
b_1	3	CH ₂ scissors	1331.67	gas	IR	9,11
			1323m	Ar	IR	1,6,7
	4	C=S stretch	850	gas	IR	9
			858	Ar	IR	7
b_1	5	H ₂ C=C OPLA	698.09	gas	IR	11
			692s	Ar	IR	1,6-8
b_2	6	CCS bend	413.85	gas	IR	12
			404vw	Ar	IR	1,7
b_2	7	CH ₂ a-stretch	3107.33	gas	IR	9,11
			3097	Ar	IR	7
			3068			
	8	CH ₂ rock	921.60	gas	IR	9,11
			918	Ar	IR	7
b_2	9	CCS bend	357.45	gas	IR	12
			356	Ar	IR	7

$A_0=9.554$; $B_0=0.189$; $C_0=0.185$ MW²⁻⁵IR^{9,11,12}

D₂CCS

\tilde{A}^1A''		C_s	$\tilde{A}-\tilde{X}$ 450-550 nm			
$T_0 \approx 1800$	2	gas AB ¹⁰	Diffuse bands.			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	C=C stretch	1400	gas	AB	10
	8	CCS bend	256T	gas	AB	10

\tilde{X}^1A_1		C_{2v}	Structure: MW ²⁻⁵ IR ¹¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CD ₂ s-stretch	2237.48	gas	IR	11
			2229	Ar	IR	1,7
	2	C=C stretch	1740	gas	IR	11
			1738vs	Ar	IR	1,7
b_1	3	CD ₂ scissors	1030	Ar	IR	7
	4	C=S stretch	755	Ar	IR	7
	5	D ₂ C=C OPLA	558.04	gas	IR	11
			555s	Ar	IR	1,7
b_2	6	CCS bend	375	Ar	IR	7
	7	CD ₂ a-stretch	2330T	Ar	IR	7
	8	CD ₂ rock	775	Ar	IR	7
b_2	9	CCS bend	308	Ar	IR	7

$A_0=4.784$; $B_0=0.168$; $C_0=0.162$ IR¹¹

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H₂CCSe

\tilde{X}		C_{2v}	Structure: MW ²			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CH ₂ s-stretch	3032.3w	Ar	IR	3
	2	C=C stretch	1699.7vs	Ar	IR	3
	3	CH ₂ scissors	1334.8w	Ar	IR	3
	4	C=Se stretch	662.1vw	Ar	IR	3
b_1	5	CH ₂ wag	736.8m	Ar	IR	3
b_2	7	CH ₂ a-stretch	3063.3vw	Ar	IR	3

$A_0=9.35(25)$; $B_0=0.129$; $C_0=0.127$ MW^{1,2}

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H₂CNO

\tilde{X}		Structure: MW ²				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'		C=N stretch	1664.0vs	Ar	IR	1
			1197.0s	Ar	IR	1
			1106.4m	Ar	IR	1
		CH ₂ OPLA	792.9wm	Ar	IR	1

Reference

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cyc-H₂CO₂

\tilde{X}		C_{2v}	Structure: MW ²			
A_0	B_0	C_0	MW ^{1,2}			

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H₂CSO

\tilde{X}		C _s	Structure: MW ²			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	CH ₂ a-stretch	3117vw	Ar	IR	5
	2	CH ₂ s-stretch	2985w	gas	IR	6
			3013vw	Ar	IR	3,5
	3	CH ₂ scissors	1355m	gas	IR	6
			1357m	Ar	IR	3,5
	4	SO stretch	1175.75s	gas	IR	1,4,6
			1165s	Ar	IR	3,5
	5	CS stretch	972w	Ar	IR	3,5
<i>a''</i>	6	CH ₂ rock	866T ^a	gas	IR	7
			860w	Ar	IR	3,5
	7	CSO bend	397w	Ar	IR	3,5
	8	H ₂ CS OPLA	762.07s	gas	IR	1,4,7
			765s	Ar	IR	3,5

$A_0=1.348$; $B_0=0.313$; $C_0=0.254$ MW^{1,2}IR⁷

D₂CSO

\tilde{X}		C _s	Structure: MW ²			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	CD ₂ a-stretch	2339w	Ar	IR	5
	2	CD ₂ s-stretch	2137wT	gas	IR	6
			2180vw	Ar	IR	5
	3	SO stretch	1189s	gas	IR	6
			1181s	Ar	IR	5
	4	CD ₂ scissors	1117wT	gas	IR	6
			1100m	Ar	IR	5
	5	CS stretch	835wT	gas	IR	6
<i>a''</i>			845w	Ar	IR	5
	6	CD ₂ rock	719w	Ar	IR	5
	7	CSO bend	345m	Ar	IR	5
	8	D ₂ CS OPLA	602m	gas	IR	6
			607m	Ar	IR	5

$A_0=1.148$; $B_0=0.277$; $C_0=0.223$ MW²

^aEstimated from analysis of perturbations.⁷

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CH₂F₂⁺

$\tilde{G}^{\circ} 2A_1$ C_{2v}
 $T_0=83900(1000)$ gas PE^{1,3}

$\tilde{F}^{\circ} 2B_2$ C_{2v}
 $T_0=44706(35)$ gas PE^{1-3,5}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CF ₂ stretch	708	gas	PE	5
<i>a</i> ₁	3	CF ₂ stretch	668	gas	PE	5

$\tilde{D}^{\circ} 2B_1$ C_{2v}
 $T_0=44206(35)$ gas PE^{1-3,5}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CF ₂ stretch	711(5)	gas	PE	1-3,5

$\tilde{C}^{\circ} 2A_2$ C_{2v}
 $T_0=23358(35)$ gas PE^{1-3,5}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CF ₂ stretch	1050(5)	gas	PE	1-3,5
	4	CF ₂ scissors	500(80)	gas	PE	1,3

$\tilde{B}^{\circ} 2A_1$ C_{2v}
 $T_0=22938(35)$ gas PE⁵

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CF ₂ stretch	1054(5)	gas	PE	5

$\tilde{A}^{\circ} 2B_1$ C_{2v}
 $T_0=15185(35)$ gas PE^{1-3,5}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₂ stretch	2744m	Ar	IR	4
	2	CH ₂ scissors	1150(5)	gas	PE	1-3,5
	3	CF ₂ stretch	1162(5)	gas	PE	5
	4	CF ₂ scissors	527H	gas	PE	5
	6	CH ₂ stretch	2854wm	Ar	IR	4
	8	CH ₂ wag	1408s	Ar	IR	4
	9	CF ₂ stretch	1255vs	Ar	IR	4

CD₂F₂⁺

$\tilde{C}^{\circ} A_2$		C _{2v}	
Vib.	No.	Approximate type of mode	cm ⁻¹
a ₁	3	CF ₂ stretch	970(80)
	4	CF ₂ scissors	500(80)
$\tilde{X}^{\circ} B_2$		C _{2v}	
Vib.	No.	Approximate type of mode	cm ⁻¹
a ₁	1	CD ₂ stretch	2062m
b ₁	7		980w
h ₂	8	CD ₂ wag	1063s
	9	CF ₂ stretch	1262vs

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⁵T. Pradeep and D. A. Shirley, J. Electron Spectrosc. Relat. Phenom. **66**, 125 (1993).

CH₂Cl₂⁺

$\tilde{G}^{\circ} A_1$		C _{2v}	
T ^a =	72500(1000)	gas	PE ¹

$\tilde{F}^{\circ} B_1$		C _{2v}	
T ^a =	43970(240)	gas	PE ¹

$\tilde{E}^{\circ} A_1$		C _{2v}	
T ^a =	37280(240)	gas	PE ¹

$\tilde{D}^{\circ} B_2$		C _{2v}	
T ₀ =	28530(35)	gas	PE ^{1,2}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	3	CCl ₂ stretch	541(5)	gas	PE	2

$\tilde{C}^{\circ} A_2$		C _{2v}	
T ₀ =	7673(35)	gas	PE ^{1,2}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	4	CCl ₂ scissors	297(5)	gas	PE	2

$\tilde{B}^{\circ} A_1$		C _{2v}	
T ₀ =	6713(35)	gas	PE ^{1,2}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	4	CCl ₂ scissors	276(5)	gas	PE	2

 $\tilde{A}^{\circ} B_1$ C_{2v}
T₀=300(35) gas PE²

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	3	CCl ₂ stretch	677(5)	gas	PE	2
	4	CCl ₂ scissors	400(5)	gas	PE	2

 $\tilde{X}^{\circ} B_2$ C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	3	CCl ₂ stretch	713(5)	gas	PE	1,2

^aFrom vertical ionization potential.**References**

- ¹A. W. Potts, H. J. Lempka, D. G. Streets, and W. C. Price, Philos. Trans. R. Soc. London, Ser. A **268**, 59 (1970).

- ²T. Pradeep and D. A. Shirley, J. Electron Spectrosc. Relat. Phenom. **66**, 125 (1993).

t-HSSSH**X**C₂

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		SH a-stretch	2542	gas	IR	1
		SSH a-bend	860	gas	IR	1
		S ₃ a-stretch	480	gas	IR	1

A=0.470; B=0.092; C=0.079 MW¹**Reference**

- ¹M. Liedtke, A. H. Saleck, K. M. T. Yamada, G. Winnewisser, D. Kremer, E. Kraka, A. Dolgner, J. Hahn, and S. Dobos, J. Phys. Chem. **97**, 11204 (1993).

c-HSSSH**X**C_sEstimated⁴ to be 87 cm⁻¹ (0.25 kcal/mol) higher in energy than t-HSSSH.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		SH s-stretch	2548	gas	IR	4
		SSH a-bend	865	gas	IR	4
		S ₃ a-stretch	480	gas	IR	4
		S ₃ bend	240(30)	gas	MW	3

A=0.470; B=0.092; C=0.079 MW¹⁻⁴**References**

- ¹D. Mauer, G. Winnewisser, K. M. T. Yamada, J. Hahn, and K. Reinartz, Z. Naturforsch. A **43**, 617 (1988).

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8.10. Five-Atomic Monohydrides

FeC=C=CH

\tilde{A}
 $T_0=7260(630)$ gas PE¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Fe=C stretch	670(150)	gas	PE	1

Reference

¹J. Fan, L. Lou, and L.-S. Wang, J. Chem. Phys. **102**, 2701 (1995).

FeCCCH^a

\tilde{A}
 $T_0=6860(760)$ gas PE¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Fe≡C stretch	880(150)	gas	PE	1

^aAcetylenic bonding.

Reference

¹J. Fan, L. Lou, and L.-S. Wang, J. Chem. Phys. **102**, 2701 (1995).

FeC=C=CH⁻

Threshold for electron detachment from ground-state FeC=C=CH⁻ = 12700(500) gas PE¹

Reference

¹J. Fan, L. Lou, and L.-S. Wang, J. Chem. Phys. **102**, 2701 (1995).

FeCCCH^{-a}

Threshold for electron detachment from ground-state FeCCCH⁻ = 10250(650) gas PE¹

^aAcetylenic bonding.

Reference

¹J. Fan, L. Lou, and L.-S. Wang, J. Chem. Phys. **102**, 2701 (1995).

C₄H

Absorptions between 250 and 300 nm which were attributed by Ref. 1 to C₄H have since been reassigned (D. Forney and J. P. Maier, private communication) to C₆H₂. Groups of absorptions observed in an argon matrix¹ between 13408 and 13906, 17629 and 17939, 21972 and 25667, and 26867 and 32104 are tentatively attributed to C₄H.

$\tilde{X}^2\Sigma$		C _{∞v}	Structure: MW ⁹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CH stretch	3307.4w	Ar	IR	7
	2	C≡C stretch	2083.9wm	Ar	IR	7
	3	C≡C stretch	2063.7	Ne	IR	8
Π	7	Skel. bend	2060.6ms	Ar	IR	1,7
			131T	gas	MW	6

$B_0=0.165$ MW²⁻⁵

C₄D

An absorption at 17685 and a group of bands between 26925 and 30883 in the argon-matrix observations¹ are tentatively attributed to C₄D.

$\tilde{X}^2\Sigma$		C _{∞v}	Structure: MW ⁹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CD stretch	2579.3w	Ar	IR	7
	2	C≡C stretch	2056.5wm	Ar	IR	7
	3	C≡C stretch	2052.9	Ne	IR	8
			2049.6ms	Ar	IR	1,7

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- D. Forney, M. E. Jacox, and W. E. Thompson, J. Mol. Spectrosc. **170**, 178 (1995).
- M. C. McCarthy, C. A. Gottlieb, P. Thaddeus, M. Horn, and P. Botschwina, J. Chem. Phys. **103**, 7820 (1995).

HCCCO

\tilde{X}^2A'		C _s	Structure: MW ⁴			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2		2308.6	Ar	IR	3

$A_0=8.709(2)$; $B_0=0.153$; $C_0=0.150$ MW^{1,2}

DCCCO

\tilde{X}^2A'		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	2		2287.4	Ar	IR	3
$A_0=8.488(3)$; $B_0=0.142$; $C_0=0.139$	MW ²					

References

- ¹A. L. Cooksy, J. K. G. Watson, C. A. Gottlieb, and P. Thaddeus, *Astrophys. J.* **386**, L27 (1992).
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⁴A. L. Cooksy, J. K. G. Watson, C. A. Gottlieb, and P. Thaddeus, *J. Chem. Phys.* **101**, 178 (1994).

HCCCS

$\tilde{X}^2\Pi$		$C_{\infty v}$
$B_0=0.090$		MW ¹

DCCCS

$\tilde{X}^2\Pi$		$C_{\infty v}$
$B_0=0.084$		MW ¹

Reference

- ¹Y. Hirahara, Y. Ohshima, and Y. Endo, *J. Chem. Phys.* **101**, 7342 (1994).

HCSCN⁺

\tilde{E}^2A'		C_s					
T ^a	41100(1200)	gas	PE ¹				
\tilde{D}^2A''		C_s					
T ^a	32300(1200)	gas	PE ¹				
\tilde{C}^2A'		C_s					
T ₀	29000(1200)	gas	PE ¹				
\tilde{B}^2A'		C_s					
T ₀	21800(1200)	gas	PE ¹				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.	
			1900T	gas	PE	1	

\tilde{A}^2A'' C_s
 $T^a=13700(1200)$ gas PE¹

 \tilde{X}^2A'

^aFrom vertical ionization potential.

Reference

- ¹G. Pfister-Guillouzo, F. Gracian, A. Senio, F. Bourdon, Y. Vallée, and J.-L. Ripoll, *J. Am. Chem. Soc.* **115**, 324 (1993).

HCOCN

\tilde{A}^1A''		C_s				
T ₀	26283.396	gas	LF ^{1,4,6} AB ⁴	$\tilde{A}-\tilde{X}$ 345–385 nm		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	3	CO stretch	1308	gas	LF	1,6
	4	HCO bend	1121	gas	LF	1,6
	5	CC stretch	943	gas	LF	1,6
	6	CCO bend	530	gas	LF	1,6
	7	CCN bend	217	gas	LF	1,6
a''	8	Bend	418	gas	LF	1,6
	9	Bend	376	gas	LF	1,6

$A_0=1.957$; $B_0=0.169$; $C_0=0.154$ AB,LF⁴

X^1A'		C_s				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CH stretch	2892m	gas	IR	5
	2	C≡N stretch	2229m	gas	IR,EM	2,5,6
	3	C=O stretch	1716s	gas	IR,EM	2,5,6
	4	CH rock	1383	gas	IR	5
	5	C-C stretch	914ms	gas	IR,EM	2,5,6
	6	CCO bend	626vw	gas	IR	5
	7	CCN bend	230	gas	LF,IR	1,2,6
a''	8	CH wag	960	gas	EM	6
	9	CCN bend	278	gas	LF,EM	1,6

$A_0=2.251$; $B_0=0.167$; $C_0=0.155$ MW^{3,7}

DCOCN

\tilde{A}^1A''		C_s				
T ₀	26335.6	gas	LF ⁶	$\tilde{A}-\tilde{X}$ 345–380 nm		
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CD stretch	2232	gas	LF	6
	3	CO stretch	1279	gas	LF	6
	4	CC stretch	931	gas	LF	6
	5	CD rock	732	gas	LF	6
	6	CCO bend	527	gas	LF	6
	7	CCN bend	214	gas	LF	6
a''	8	Bend	406	gas	LF	6
	9	Bend	315	gas	LF	6

$A^a=1.561$; $B^a=0.169$; $C^a=0.152$ LF⁶

\tilde{X}^1A'		C_s
$A_0=1.737$; $B_0=0.168$; $C_0=0.152$		LF ⁶

^aFrom analysis of g_0^1 band.

References

- ¹R. H. Judge, D. C. Moule, A. Biernacki, M. Benkel, J. M. Ross, and J. Rustenburg, *J. Mol. Spectrosc.* **116**, 364 (1986).
²D. J. Clouthier and D. C. Moule, *J. Am. Chem. Soc.* **109**, 6259 (1987).
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⁵ W. Lewis-Bevan, R. D. Gaston, J. Tyrrell, W. D. Stork, and G. L. Salmon, *J. Am. Chem. Soc.* **114**, 1933 (1992).

⁶ D. J. Clouthier, J. Karolczak, J. Rae, W.-T. Chan, J. D. Goddard, and R. H. Judge, *J. Chem. Phys.* **97**, 1638 (1992).

⁷ M. Bogey, C. Demuyunc, J. L. Destombes, Yo. Vallee, and J. L. Ripoll, *J. Mol. Spectrosc.* **172**, 344 (1995).

HCSCN

\tilde{X}		C_s	
Vib. sym.	No.	Approximate type of mode	cm^{-1}
		Med.	Type meas.
a'	2	CN stretch	2221
	3	CH rock	1320
	4	CS stretch	1103
	5	CC stretch	889
a''	8	CH wag	824

$A_0=1.445$; $B_0=0.107$; $C_0=0.099$ MW¹

References

¹ M. Bogey, C. Demuyunc, J.-L. Destombes, A. Gaumont, J.-M. Denis, Y. Vallée, and J.-L. Ripoll, *J. Am. Chem. Soc.* **111**, 7399 (1989).

² C. O. Kappe, M. W. Wong, and C. Wentrup, *Tetrahedron Lett.* **34**, 6623 (1993).

cyc-C₃HCl

In an argon matrix, weak absorption maxima at 37000 (270 nm) and 38500 (260 nm) have been attributed¹ to cyc-C₃HCl. Exposure of the molecule to 254 nm radiation results in its isomerization to HCCCCl and HCIC=C=C:

\tilde{X}		C_s	
Vib. sym.	No.	Approximate type of mode	cm^{-1}
		Med.	Type meas.
a'	1	CH stretch	3139.2w
	2	Ring	1687.0m
	3	Ring	1283.4wm
	4	Ring	1059.7vs
	5	CH deform.	897.2w
	6	CCl stretch	580.5wm
a''	8	CH deform.	883.7wm

Reference

¹ G. Maier, T. Preiss, H. P. Reisenauer, B. A. Hess, Jr., and L. J. Schaad, *J. Am. Chem. Soc.* **116**, 2014 (1994).

HCCCCl

In an argon matrix, HCCCCl possesses broad, overlapping absorption maxima in the 270–300 nm spectral region.¹ Irradiation in that spectral region results in isomerization to cyc-C₃HCl.

\tilde{X}	C_s					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CH stretch	3305.5s	Ar	IR	1
	2	C=C stretch	1983.0s	Ar	IR	1
	3	C-C stretch	1054.4vs	Ar	IR	1
	4	CCl stretch	725.1vs	Ar	IR	1
	5	CH deform.	475.4m	Ar	IR	1
a''	8	CH deform.	511.5wm	Ar	IR	1

Reference

¹ G. Maier, T. Preiss, H. P. Reisenauer, B. A. Hess, Jr., and L. J. Schaad, *J. Am. Chem. Soc.* **116**, 2014 (1994).

HCIC=C=C:

In an argon matrix, HCIC=C=C: possesses broad, overlapping absorption maxima in the 270–300 nm spectral region.¹ Irradiation in that spectral region results in isomerization to cyc-C₃HCl.

\tilde{X}	C_s					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CH stretch	3103.5vw	Ar	IR	1
	2	C_3 a-stretch	1967.6vs	Ar	IR	1
	3	CH deform.	1293.0w	Ar	IR	1
	4	C_3 s-stretch	1146.5w	Ar	IR	1
	5	CCl stretch	794.1ms	Ar	IR	1
a''	8	CH deform.	839.9vw	Ar	IR	1

Reference

¹ G. Maier, T. Preiss, H. P. Reisenauer, B. A. Hess, Jr., and L. J. Schaad, *J. Am. Chem. Soc.* **116**, 2014 (1994).

HFCCO

\tilde{X}	C_s		Structure: MW ²			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	CH stretch	3025.9	Xe	IR	3
	2	CO stretch	2142.2vs	Xe	IR	3
	3		1383.0m	Xe	IR	3
	4		1168.0m	Xe	IR	3
	5		1031.5w	Xe	IR	3
	7	Deformation	150(10)T	gas	MW	2
a''	8	HFCC OPLA	500(50)T	gas	MW	2
			755	Xe	IR	3
	9	CCO bend	380(40)T	gas	MW	2
			525.5w	Xe	IR	3

$A_0=1.795$; $B_0=0.154$; $C_0=0.142$ MW¹

References

- ¹R. D. Brown, P. D. Godfrey, and B. Kleibömer, Chem. Phys. **105**, 301 (1986).
²R. D. Brown, P. D. Godfrey, and K. H. Wiedenmann, J. Mol. Spectrosc. **136**, 241 (1989).
³G. Davidovics, M. Monnier, and A. Allouche, Chem. Phys. **150**, 395 (1991).

 $\text{CF}_2=\text{NH}^+$ $\tilde{X} \ ^2\text{A}''$ C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'		C=N stretch	1480(40)	gas	PE	1

Reference

- ¹G. Pawelke, R. Dammel, and W. Poll, Z. Naturforsch. B **47**, 351 (1992).

 $\text{CF}_2=\text{NH}$ \tilde{X} C_s Structure: MW⁷

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	NH stretch	3402.3m	gas	IR	1,5
			3388.4m	Ar	IR	5,6
	2	C=N stretch	1785.2vs	gas	IR	1,5
			1777.2vs	Ar	IR	5,6
	3	CF ₂ a-stretch	1307vs	gas	IR	1,5
			1294.9vs	Ar	IR	5,6
	4	CNH deform.	1031.2s	gas	IR	1,5
			1027.0s	Ar	IR	5,6
	5	CF ₂ s-stretch	947.2s	gas	IR	1,5
			941.9s	Ar	IR	5,6
a''	6	CF ₂ scissors	572.7vw	Ar	IR	5,6
	7	CF ₂ rock	542.0m	gas	IR	1,5
			541.4m	Ar	IR	5,6
	8	Torsion	831.65s	gas	IR	1,4,5
			827.8s	Ar	IR	5,6
	9	F ₂ CN OPLA	695.2w	gas	IR	1,5
			694.3w	Ar	IR	5,6

$A_0=0.382$; $B_0=0.372$; $C_0=0.188$ MW^{2,3}

 $\text{CF}_2=\text{ND}$

\tilde{X}	C_s					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	ND stretch	2519.3m	gas	IR	1,5
			2508.2m	Ar	IR	5,6
	2	CN stretch	1780.0vs	gas	IR	1,5
			1771.9vs	Ar	IR	5,6
	3	CF ₂ a-stretch	1280vs	gas	IR	1,5
			1266.8vs	Ar	IR	5,6
	4	CF ₂ s-stretch	954.0m	gas	IR	1,5
			951.5m	Ar	IR	5,6
	5	CND deform.	836.0s	gas	IR	1,5
			832.7s	Ar	IR	5,6
a''	6	CF ₂ scissors	563.7vw	Ar	IR	5,6
	7	CF ₂ rock	491.8m	gas	IR	1,5
			491.5m	Ar	IR	5,6
8		F ₂ CN OPLA	705.1m	gas	IR	1,5
			702.4m	Ar	IR	5,6
9		Torsion	613.9w	gas	IR	1,5
			612.1w	Ar	IR	5,6

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 HONO_2

In the gas phase, continuous absorption begins near 330 nm, with a maximum near 38500 (260 nm), then increasing absorption out to the 190 nm observation limit.^{4,5,31} Between 330 and 220 nm, photodissociation to OH+NO₂ predominates.^{6,16,33,39} However, at 193 nm photodissociation to HONO+O is the major process.³³ Vacuum ultraviolet spectral observations^{7,12} show an absorption maximum near 54900 (182 nm), a more prominent maximum near 73500 (136 nm), and overlapping absorption bands at higher energies. The onset of the photoproduction of OH ($A \ ^2\Sigma$) + NO₂ has been observed¹² at 68000 (147.5 nm).

\bar{X}	C _s	Structure: MW ^{1,3,43}				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OH stretch	3550.0m	gas	IR	2
			3522.3	Ar	IR	28
			3519.3			
			3491.8	N ₂	IR	8,36
<i>a'</i>	2	NO ₂ a-stretch	1709.5vs	gas	IR,DL LS	2,9,11,20, 21,23,35
			1699.4	Ar	IR	28
			1696.2			
			1698.3	N ₂	IR	8,36
<i>a'</i>	3	Mixed	1325.74s	gas	IR,DL	2,19,26, 27,40,41
			1321.4	Ar	IR	28
			1318.7			
			1346.1	N ₂	IR	8,36
<i>a'</i>	4	Mixed	1303.52vs	gas	IR,DL	2,26,27, 40,41
			1304.4	Ar	IR	28
			1311.9	N ₂	IR	8,36
			879.11s	gas	IR,DL	2,18,32, 34,37,40, 41
<i>a'</i>	5	ON stretch	896.9	Ar	IR	2
			889.5			
			903.1	N ₂	IR	8,36
			646.83w	gas	IR	2,25,46
<i>a'</i>	6	NO ₂ scissors	656.6	Ar	IR	36
			664.1	N ₂	IR	8,36
			580.30w	gas	IR	2,25,44
			588.0	Ar	IR	36
<i>a'</i>	7	NO ₂ rock	597.5	N ₂	IR	8,36
			763.15s	gas	IR	2,25,44, 45
			763.6	Ar	IR	28
			767.7	N ₂	IR	8,36
<i>a''</i>	8	ONO ₂ OPLA	458.23m	gas	IR	2,15,24
			479	N ₂	IR	8

$A_0=0.434$; $B_0=0.404$; $C_0=0.209$ MW^{1,10,13,14,17,22}IR^{11,24,25}

DONO₂

\bar{X}	C _s	Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OD stretch	2621.5m	gas	IR	2		
			2601.5	Ar	IR	36		
			2599.1					
			2576.5	N ₂	IR	8,36		
<i>a'</i>	2	NO ₂ a-stretch	1688.36vs	gas	IR	2,42		
			1678.6	Ar	IR	36		
			1674.9					
			1669.2	N ₂	IR	8,36		
<i>a'</i>	3	NO ₂ s-stretch	1308.4vs	gas	IR	2		
			1310.4	Ar	IR	36		
			1311.1	N ₂	IR	8,36		
			1013.6m	gas	IR	2		
<i>a'</i>	4	DON bend	1013.4	Ar	IR	36		
			1012.2					
			1032.4	N ₂	IR	8,36		
			888.0s	gas	IR	2		
<i>a'</i>	5	ON stretch	894.2	Ar	IR	36		
			884.5					
			906.0	N ₂	IR	8,36		
			642.14s	gas	IR	2,38		
<i>a'</i>	6	NO ₂ scissors	660	N ₂	IR	8		
			541.58w	gas	IR	2,38		
			559.9	N ₂	IR	8,36		
			762.87s	gas	IR	2,29		
<i>a''</i>	7	NO ₂ rock	763.7	Ar	IR	36		
			767.7	N ₂	IR	8,36		
			343.85m	gas	IR	2,15,30		
			361	N ₂	IR	8		

$A_0=0.433$; $B_0=0.377$; $C_0=0.201$ MW^{1,43}IR³⁸

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c,c-HOONO

In an argon matrix, an absorption maximum near 36400 (275 nm) has been assigned² to c,c-HOONO.



Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	OH stretch	3285.4	Ar	IR	1,2	
2	N=O stretch	1600.3	Ar	IR	1,2	
3	HOO bend	1395.0	Ar	IR	1,2	
4	OO stretch	927.2	Ar	IR	1,2	
5	ONO bend	794.3	Ar	IR	1,2	
6	N-O stretch	629.1	Ar	IR	1,2	

c,c-DOONO



Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	OD stretch	2434.4	Ar	IR	1
	2	N=O stretch	1594.1	Ar	IR	1
	3	DOO bend	1091.9	Ar	IR	1
	4	OO stretch	927.2	Ar	IR	1
	5	ONO bend	792.5	Ar	IR	1
	6	N-O stretch	630.8	Ar	IR	1

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t,perp-HOONO

In an argon matrix, an absorption maximum near 45500 (220 nm) has been assigned³ to t,perp-HOONO.



Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	OH stretch	3545.5w	Ar	IR	1-3
			3563.5 ^a	Ar	IR	1-3
			3541.7	N ₂	IR	1,2
	2	N=O stretch	1703.6s	Ar	IR	1-3
			1708.3 ^a	Ar	IR	1-3
			1701.4	N ₂	IR	1,2
	3	HOO bend	1364.4m	Ar	IR	1-3
			1372.7 ^a	Ar	IR	1-3
			1394.9	N ₂	IR	1,2
	4	OO stretch	952.0m	Ar	IR	1-3
			957.4 ^a	Ar	IR	1-3
			960.5	N ₂	IR	1,2
	5	ONO bend	772.8m	Ar	IR	1-3
			782.9 ^a	Ar	IR	1-3
			793.6	N ₂	IR	1,2

t,perp-DOONO



Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
	1	OD stretch	2615.4	Ar	IR	1,2
			2633.5 ^a	Ar	IR	1,2
			2622.3	N ₂	IR	2
	2	N=O stretch	1703.7	Ar	IR	1,2
			1'08.3 ^a	Ar	IR	1,2
			1701.8	N ₂	IR	2
	3	DOO bend	1089.7	Ar	IR	1,2
			1091.7 ^a	Ar	IR	1,2
			1090.7	N ₂	IR	2
	4	OO stretch	950.3	Ar	IR	1,2
			955.7 ^a	Ar	IR	1,2
			957.3	N ₂	IR	2
	5	ONO bend	772.1	Ar	IR	1,2
			782.0 ^a	Ar	IR	1,2
			790.3	N ₂	IR	2

^aLess stable matrix site.

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HSPS₂

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		SH stretch	2571.3	Ar	IR	1
		Mixed	861.2	Ar	IR	1
		Mixed	765.7	Ar	IR	1
		Mixed	647.2	Ar	IR	1

DSPS₂

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		SD stretch	1860.3	Ar	IR	1
		Mixed	813.6	Ar	IR	1
		Mixed	687.4	Ar	IR	1

Reference

- ¹Z. Mielke and L. Andrews, *J. Phys. Chem.* **97**, 4313 (1993).

HCFCIBr⁺

\tilde{F}, \tilde{G}

$T^a=51600(500)$ gas PE¹

\tilde{E}

$T^a=26700(400)$ gas PE¹

\tilde{D}

$T^a=23100(400)$ gas PE¹

\tilde{C}

$T^a=11600(400)$ gas PE¹

\tilde{B}

$T^a=7180(300)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1120(50)	gas	PE	1

\tilde{A}

$T^a=2660(400)$ gas PE¹

^aFrom vertical ionization potentials.

Reference

- ¹I. Novak, S. C. Ng, and A. W. Potts, *Chem. Phys. Lett.* **215**, 561 (1993).

8.11. Five-Atomic Nonhydrides

FeC₄⁻

Threshold for electron detachment from ground-state FeC₄⁻ $\leq 17800(1600)$ gas PE¹

Reference

- ¹J. Fan, L. Lou and L.-S. Wang, *J. Chem. Phys.* **102**, 2701 (1995).

C₅

$T_0=41967(35)T$ Ne AB¹³ 220–240 nm
44228T Ar AB⁸ 211–226 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1813	Ar	AB	8
			695(40)	Ne	AB	13
			654	Ar	AB	8

¹ Π_u
 $T_0=19599(8)$ Ne AB¹³

¹ $\Pi_u-\tilde{X}$ 490–510 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			697(10)	Ne	AB	13
			542(10)	Ne	AB	13
			286(10)	Ne	AB	13

\tilde{X} ¹ Σ_g^- D_{∞h} Structure: ESR²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	2	Sym. stretch	779(10)	gas	TPE	9,10
			776T	Ar	IR	15
Σ_u^+	3	Asym. stretch	2169,441	gas	IR,DL	4–7
			2166.4	Ne	IR	2,12,13
			2164	Ar	IR	1–3
			2157.0	Kr	IR	15
			2164.5	H ₂	IR	14
	4	Asym. stretch	1444.3	Ne	IR	12
			1446.6	Ar	IR	11
			1443.2	Kr	IR	15
Π_g	5	Bend	218(13)T	gas	DL,TPE	7,9,10
Π_u	6	Bend	535(10)	gas	TPE	9,10
Π_u	7	Bend	118(3)T	gas	DL,TPE	7,9,10

$B_0=0.0853$ IR⁴DL⁵⁻⁷

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C₄Si

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CC stretch	2095.46	gas	DL	3
			2080.1	Ar	IR	2

$$B_0 = 0.051 \text{ MW}^1 \text{DL}^3$$

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SiCCCSi

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	C ₃ stretch	1968.19	gas	DL	2
			1955.2 vs	Ar	IR	1
	4	SiC stretch	898.9 w	Ar	IR	1

$$B_0 = 0.0316 \text{ DL}^2$$

References

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Si₃C₂

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	"Breathing"	681.1	Ar	IR	1
b_2	7	SiC a-str.	956.7	Ar	IR	1
	8	CSiC a-stretch	597.8	Ar	IR	1

Reference

- ¹J. D. Presilla-Márquez, C. M. L. Rittby, and W. R. M. Graham, J. Chem. Phys. **104**, 2818 (1996).

C₅⁻

Threshold for electron detachment from ground-state
 $C_5^- = 23020(10)$ gas TPE¹PE²

$\tilde{B} \quad 2\Pi_g \quad D_{\infty h}$
 $T_0 = 27847(15) \quad \text{Ne} \quad AB^6$ $\tilde{B} - \tilde{X} \quad 311 - 360 \text{ nm}$

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		1756(17)	Ne	AB	6
	2		724(16)	Ne	AB	6

$\tilde{A} \quad 2\Pi_g \quad D_{\infty h}$
 $T_0 = 20200T \quad \text{gas} \quad MPD^3$ $\tilde{A} - \tilde{X} \quad 470 - 500 \text{ nm}$
 $20200(8) \quad \text{Ne} \quad AB^6$ $\tilde{A} - \tilde{X} \quad 457 - 495 \text{ nm}$

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		1679(9)	Ne	AB	6
	2		718(43)	gas	MPD	3
			737(8)	Ne	AB	6



Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	3	Asym. stretch	1822.3	Ne	IR	5
			1831.8	Ar	IR	4
Π_u	7	Bend	200T	gas	TPE	1

$$A = 22T \quad \text{gas} \quad TPE^1$$

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Be(CO)₂

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CO stretch	1975.3	Ar	IR	1
			609.6	Ar	IR	1

Reference

¹L. Andrews, T. J. Tague, Jr., G. P. Kushto, and R. D. Davy, Inorg. Chem. **34**, 2952 (1995).

NNBeNN

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NN stretch	1823.4	Ar	IR	1

Reference

¹C. A. Thompson, L. Andrews, and R. D. Davy, J. Phys. Chem. **99**, 7913 (1995).

NN(cyc-BeN₂) \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1691.8	Ar	IR	1
			1682.9	N ₂	IR	1
			928.4	N ₂	IR	1

Reference

¹C. A. Thompson, L. Andrews, and R. D. Davy, J. Phys. Chem. **99**, 7913 (1995).

OBeOBeO

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1482.5T	Ne	IR	1,2
			1465.1T	Ar	IR	1,2

References

¹C. A. Thompson and L. Andrews, J. Chem. Phys. **100**, 8689 (1994).

²L. Andrews, G. V. Chertihiin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

Fe₂O₃ \tilde{A} T₀=2020(400) gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			650(80)	gas	PE	1

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			810(100)	gas	PE	1

Reference

¹H. Wu, S. R. Desai, and L.-S. Wang, J. Am. Chem. Soc. **118**, 5296 (1996).

Cu₂O₃ \tilde{B} T₀=6290(400) gas PE¹ \tilde{A} T₀=3870(400) gas PE¹ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			640T	gas	PE	1

Reference

¹L.-S. Wang, H. Wu, S. R. Desai, and L. Lou, Phys. Rev. B **53**, 8028 (1996).

C₄O \tilde{X} 3Σ⁻ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			2221.7vs	Ar	IR	1
			1922.7s	Ar	IR	1
			1431.5wm	Ar	IR	1
			774.8w	Ar	IR	1
			484.0wm	Ar	IR	1

B₀=0.078 MW²

References

¹G. Maier, H. P. Reisenauer, U. Schäfer, and H. Balli, Angew. Chem. **100**, 590 (1988); Angew. Chem. Int. Ed. Engl. **27**, 566 (1988).

²Y. Ohshima, Y. Endo, and T. Ogata, J. Chem. Phys. **102**, 1493 (1995).

C₄S \tilde{B} T₀≈41700 Ar AB¹ \tilde{A} T₀=22220 Ar AB¹ \tilde{A} – \tilde{X} 380–450 nm \tilde{X} 3Σ⁻ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1757	Ar	IR	1

B₀=0.051 MW²

References

¹G. Maier, J. Schrot, and H. P. Reisenauer, Chem. Ber. **124**, 2613 (1991).

²Y. Hirahara, Y. Ohshima, and Y. Endo, Astrophys. J. **408**, L113 (1993).

C(CN)₂

In an argon matrix, two overlapping series of relatively sharp absorptions between 280 and 360 nm, with spacings of 1650–1870 cm⁻¹, as well as weak, broad absorptions at 21300 (470 nm) and 26500 (377 nm), have been attributed² to C(CN)₂.

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1916w	Ar	IR	2
			1826w	Ar	IR	2
		CN a-stretch	1756m	Ar	IR	1,2
		C ₃ a-stretch	1158wT	Ar	IR	1
		CCN deform.	392wT	Ar	IR	1

References

- ¹
- W. H. Smith and G. E. Leroi, Spectrochim. Acta A
- 25**
- , 1917 (1969).
-
- ²
- I. R. Dunkin and A. McCluskey, Spectrochim. Acta A
- 50**
- , 209 (1994).

Fe₂O₃⁻

Threshold for electron detachment from ground-state Fe₂O₃⁻=24690(320)
gas PE¹

Reference

- ¹
- H. Wu, S. R. Desai, and L.-S. Wang, J. Am. Chem. Soc.
- 118**
- , 5296 (1996).

Cu₂O₃⁻

Threshold for electron detachment from ground-state Cu₂O₃⁻=28560(240)
gas PE¹

Reference

- ¹
- L.-S. Wang, H. Wu, S. R. Desai, and L. Lou, Phys. Rev. B
- 53**
- , 8028 (1996).

NCCNO⁺

\tilde{D} $^2\Sigma^+$ C_{∞ v}
 $T^a=60270(430)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+			1030(150)	gas	PE	1

\tilde{C} $^2\Pi$ C_{∞ v}
 $T^a=46800(430)$ gas PE¹

\tilde{B} $^2\Sigma^+$ C_{∞ v}
 $T^a=22750(280)$ gas PE¹

\tilde{A} $^2\Pi$ C_{∞ v}
 $T^a=20740(280)$ gas PE¹

 \tilde{X} $^2\Pi$ C_{∞ v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+			2200(150)	gas	PE	1
			1630(150)	gas	PE	1

^aFrom vertical ionization potentials.

Reference

- ¹
- T. Pasinszki and N. P. C. Westwood, J. Phys. Chem.
- 100**
- , 16856 (1996).

c-LiOONO \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	N=O stretch	1423.4	Ar	IR	1
			1422.0			
	2	Mixed	966.2	Ar	IR	1
	3	NO stretch	874.2	Ar	IR	1
	4	OO stretch	792.3	Ar	IR	1
	5	LiO stretch	642.5	Ar	IR	1

Reference

- ¹
- W.-J. Lo, Y.-P. Lee, J.-H. M. Tsai, H.-H. Tsai, T. P. Hamilton, J. G. Harrison, and J. S. Beckman, J. Chem. Phys.
- 103**
- , 4026 (1995).

t-LiOONO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1		N=O stretch	1581.6	Ar	IR	1
			1580.4			
		Mixed	998.3	Ar	IR	1
			995.6			
		LiO stretch	647.7	Ar	IR	1
			645.0			
		OO stretch	600.4	Ar	IR	1

Reference

- ¹
- W.-J. Lo, Y.-P. Lee, J.-H. M. Tsai, H.-H. Tsai, T. P. Hamilton, J. G. Harrison, and J. S. Beckman, J. Chem. Phys.
- 103**
- , 4026 (1995).

c-NaOONO \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	N=O stretch	1437.4	Ar	IR	1
			1434.6			
	2	Mixed	961.4	Ar	IR	1
	3	NO stretch	840.7	Ar	IR	1
	4	OO stretch	770.9	Ar	IR	1
			768.7			

Reference

¹W.-J. Lo, Y.-P. Lee, J.-H. M. Tsai, H.-H. Tsai, T. P. Hamilton, J. G. Harrison, and J. S. Beckman, J. Chem. Phys. **103**, 4026 (1995).

t-NaOONO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	N=O stretch	1549.3	Ar	IR	1	
		1540.6				
	Mixed	996.3	Ar	IR	1	
		994.1				
OO stretch	609.4	Ar	IR	1		
	607.4					

Reference

¹W.-J. Lo, Y.-P. Lee, J.-H. M. Tsai, H.-H. Tsai, T. P. Hamilton, J. G. Harrison, and J. S. Beckman, J. Chem. Phys. **103**, 4026 (1995).

c-KOONO

In an argon matrix,² a broad absorption (~70 nm, fwhm) with maximum near 30800 (325 nm) has been assigned to c-KOONO. Irradiation in the spectral region of this band leads to the formation of KNO₃.

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	N=O stretch	1444.5	Ar	IR	1
	2	Mixed	952.3	Ar	IR	1
	3	NO stretch	831.6	Ar	IR	1
	4	OO stretch	749.1	Ar	IR	1

References

¹W.-J. Lo, Y.-P. Lee, J.-H. M. Tsai, H.-H. Tsai, T. P. Hamilton, J. G. Harrison, and J. S. Beckman, J. Chem. Phys. **103**, 4026 (1995).

²W.-J. Lo, Y.-P. Lee, J.-H. M. Tsai, and J. S. Beckman, Chem. Phys. Lett. **242**, 147 (1995).

t-KOONO

In an argon matrix,² a broad absorption (~70 nm, fwhm) with maximum near 26700 (375 nm) has been assigned to t-KOONO. Irradiation in the spectral region of this band leads to the formation of KNO₃.

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	N=O stretch	1528.4	Ar	IR	1	
		987.4	Ar	IR	1	
	OO stretch	602.2	Ar	IR	1	

References

¹W.-J. Lo, Y.-P. Lee, J.-H. M. Tsai, H.-H. Tsai, T. P. Hamilton, J. G. Harrison, and J. S. Beckman, J. Chem. Phys. **103**, 4026 (1995).

²W.-J. Lo, Y.-P. Lee, J.-H. M. Tsai, and J. S. Beckman, Chem. Phys. Lett. **242**, 147 (1995).

NNBeO₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			928.9	N ₂	IR	1

Reference

¹L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 10088 (1996).

O₂TiCO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CO stretch	1887.7	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Am. Chem. Soc. **117**, 1595 (1995).

O₂TiN₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NN stretch	1901.3	Ar	IR	1
			909.4	Ar	IR	1

Reference

¹G. V. Chertihin and L. Andrews, J. Phys. Chem. **98**, 5891 (1994).

N₂FeO₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NN stretch	2271.3	Ar	IR	1
			1002.0	Ar	IR	1

Reference

¹G. V. Chertihin, W. Saffel, J. T. Yustein, L. Andrews, M. Neurock, A. Ricca, and C. W. Bauschlicher, Jr., J. Phys. Chem. **100**, 5261 (1996).

O₂UCO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CO stretch	1893.4	Ar	IR	1
		UO ₂ a-stretch	870.9	Ar	IR	1

Reference

¹T. J. Tague, Jr., L. Andrews, and R. D. Hunt, *J. Phys. Chem.* **97**, 10920 (1993).

Al₂O₃ \tilde{B}

$T_0=9600(640)$ gas PE²

 \tilde{A}

$T_0=4920(160)$ gas PE²

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1211.2	Ar	IR	1
		Sym. stretch	850(80)	gas	PE	2

References

¹L. Andrews, T. R. Burkholder, and J. T. Yustein, *J. Phys. Chem.* **96**, 10182 (1992).

²S. R. Desai, H. Wu, C. M. Rohlfing, and L.-S. Wang, *J. Chem. Phys.* **106**, 1309 (1997).

Al₂S₃ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			687	Ar	IR	1
			672	Ar	IR	1
			445	Ar	IR	1
			347	Ar	IR	1

Reference

¹B. S. Ault, *J. Phys. Chem.* **98**, 77 (1994).

NCCNO

In the gas phase, absorption has been observed³ between 210 and 290 nm, with a maximum at 251.7 nm (39730).

 \tilde{X} $C_{\infty v}$ Structure: MW⁴

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ	1	CNO a-stretch	2328vs	gas	IR	3
			2356vs	Ar	IR	1
	2	CN stretch	2192m	gas	IR	3
			2192w	Ar	IR	1
	3	CNO s-stretch	1442s	gas	IR	3
			1445wm	Ar	IR	1
	4	CC stretch	714.75vw	gas	IR	2,3
			717vw	Ar	IR	1
Π	5	CNO bend	422T	gas	IR	3
	6	NCC bend	403.93w	gas	IR	2,3
			407vw	Ar	IR	1
	7	CCN bend	86T	gas	IR	3

$B_0=0.077 \text{ IR}^2 \text{MW}^4$

References

¹G. Maier and J. H. Teles, *Angew. Chem.* **99**, 152 (1987); *Angew. Chem. Int. Ed. Engl.* **26**, 155 (1987).

²B. Guo, T. Pasinszki, N. P. C. Westwood, K. Zhang, and P. F. Bernath, *J. Chem. Phys.* **105**, 4457 (1996).

³T. Pasinszki and N. P. C. Westwood, *J. Phys. Chem.* **100**, 16856 (1996).

⁴Th. Brupbacher, R. K. Bohn, W. Jäger, M. C. L. Gerry, T. Pasinszki, and N. P. C. Westwood, *J. Mol. Spectrosc.* **181**, 316 (1997).

Al₂O₃⁻

Threshold for electron detachment from ground-state Al₂O₃⁻=29930(240) gas PE¹

Reference

¹S. R. Desai, H. Wu, C. M. Rohlfing, and L.-S. Wang, *J. Chem. Phys.* **106**, 1309 (1997).

O₂BeO₂ \tilde{X} D_{2d}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1307.4	Ar	IR	1,2

References

¹C. A. Thompson and L. Andrews, *J. Chem. Phys.* **100**, 8689 (1994).

²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 10088 (1996).

O₂MgO₂

$\tilde{\chi}$	D _{2d}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			669.9	Ar	IR	1,2

References

- ¹L. Andrews and J. T. Yustein, *J. Phys. Chem.* **97**, 12700 (1993).
²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 10088 (1996).

O₂CaO₂

$\tilde{\chi}$	D _{2d}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			494.4	Ar	IR	1,2
			449.5	N ₂	IR	2

References

- ¹L. Andrews, J. T. Yustein, C. A. Thompson, and R. D. Hunt, *J. Phys. Chem.* **98**, 6514 (1994).
²L. Andrews, G. V. Chertihin, C. A. Thompson, J. Dillon, S. Byrne, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 10088 (1996).

(cyc-O₂Fe)O₂

$\tilde{\chi}$				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.
			1095.4	Ar
			1097.6	O ₂
			968.9	Ar
			970.2	O ₂
				IR
				1
				IR
				1
				IR
				1

Reference

- ¹G. V. Chertihin, W. Saffel, J. T. Yustein, L. Andrews, M. Neurock, A. Ricca, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 5261 (1996).

(cyc-O₂Fe)OO

$\tilde{\chi}$				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.
			1421	Ar
			1413.4	O ₂
			1390	Ar
			1368.4	O ₂
			1174.7	Ar
			1166.6	O ₂
				IR
				1
				IR
				1
				IR
				1
				IR
				1

Reference

- ¹G. V. Chertihin, W. Saffel, J. T. Yustein, L. Andrews, M. Neurock, A. Ricca, and C. W. Bauschlicher, Jr., *J. Phys. Chem.* **100**, 5261 (1996).

O₂RhO₂

$\tilde{\chi}$	D _{2d}	Structure: IR ¹				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OO stretch	1048	Ar	IR	1
			1045	Xe	IR	1
			1038	O ₂	IR	1

Reference

- ¹A. J. L. Hanlan and G. A. Ozin, *Inorg. Chem.* **16**, 2848 (1977).

O₂NiO₂

$\tilde{\chi}$	C _{2h}	Structure: IR ¹				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OO stretch	1063.9	Ar	IR	1,2

References

- ¹H. Huber, W. Klotzbücher, G. A. Ozin, and A. Vander Voet, *Can. J. Chem.* **51**, 2722 (1973).
²A. Citra, G. V. Chertihin, L. Andrews, and M. Neurock, *J. Phys. Chem. A* **101**, 3109 (1997).

O₂PdO₂

$\tilde{\chi}$	D _{2d}	Structure: IR ¹				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OO stretch	1110.8	Ar	IR	1
		PdO stretch	504w	Ar	IR	1

Reference

- ¹H. Huber, W. Klotzbücher, G. A. Ozin, and A. Vander Voet, *Can. J. Chem.* **51**, 2722 (1973).

O₂PtO₂

$\tilde{\chi}$	D _{2d}	Structure: IR ¹				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OO stretch	1050.2	Ar	IR	1
			1047.2			

Reference

- ¹H. Huber, W. Klotzbücher, G. A. Ozin, and A. Vander Voet, *Can. J. Chem.* **51**, 2722 (1973).

O₂CuO₂ $T_0=15330(320)$ gas PE¹ \bar{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		OO stretch	1010(90)	gas	PE	1

Reference¹H. Wu, S. R. Desai, and L.-S. Wang, J. Phys. Chem. A **101**, 2103 (1997).**cyc-C₃F₂** \bar{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1		1930.5wm	Ar	IR	1
	3		797.4w	Ar	IR	1
b_2	7		1353.3vs	Ar	IR	1
	8		876.0w	Ar	IR	1
	9		543.8w	Ar	IR	1

Reference¹G. Maier, T. Preiss, and H. P. Reisenauer, Chem. Ber. **127**, 779 (1994).**FCCCCF** \bar{X} C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	C≡C stretch	2226.5vs	Ar	IR	1
	3		1115.7s	Ar	IR	1
	4		832.2wm	Ar	IR	1

Reference¹G. Maier, T. Preiss, and H. P. Reisenauer, Chem. Ber. **127**, 779 (1994).**F₂C=C=C:** \bar{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	C ₃ a-stretch	2025.4vs	Ar	IR	1
	2		1436.3s	Ar	IR	1
	3		841.6wm	Ar	IR	1
	4		544.8w	Ar	IR	1
b_1	5		684.1w	Ar	IR	1
b_2	7	CF ₂ a-stretch	1293.1m	Ar	IR	1
	8		568.4vw	Ar	IR	1

Reference¹G. Maier, T. Preiss, and H. P. Reisenauer, Chem. Ber. **127**, 779 (1994).**cyc-C₃Cl₂**

In an argon matrix, weak absorption maxima at 37300 (268 nm) and 38800 (258 nm) have been attributed¹ to cyc-C₃Cl₂. Isomerization to ClCCCCl and Cl₂C=C=C occurs when the molecule is exposed to radiation in the region of these absorptions.

 \bar{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	Ring	1736.6wm	Ar	IR	1
	2	Ring	1263.2w	Ar	IR	1
	3	CCl stretch	527.5vw	Ar	IR	1
b_2	7	Ring	1095.9vs	Ar	IR	1
	8	CCl stretch	681.7w	Ar	IR	1

Reference¹G. Maier, T. Preiss, H. P. Reisenauer, B. A. Hess, Jr., and L. J. Schaad, J. Am. Chem. Soc. **116**, 2014 (1994).**CICCCCCI**

In an argon matrix, ClCCCCl possesses strongly overlapping absorption bands between 275 and 320 nm.¹ Irradiation in that spectral region results in isomerization to cyc-C₃Cl₂.

 \bar{X} C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	C=C stretch	2079.9vs	Ar	IR	1
	2	C-C stretch	1192.8wm	Ar	IR	1
	3	CCl stretch	743.0s	Ar	IR	1

Reference¹G. Maier, T. Preiss, H. P. Reisenauer, B. A. Hess, Jr., and L. J. Schaad, J. Am. Chem. Soc. **116**, 2014 (1994).**Cl₂C=C=C:**

In an argon matrix, Cl₂C=C=C: possesses strongly overlapping absorption bands between 275 and 320 nm.¹ Irradiation in that spectral region results in isomerization to cyc-C₃Cl₂.

 \bar{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	C ₃ a-stretch	1976.7vs	Ar	IR	1
	2	C ₃ s-stretch	1238.1wm	Ar	IR	1
	3	ClCC deform.	542.9w	Ar	IR	1
b_2	7	CCl ₂ a-stretch	890.5m	Ar	IR	1

Reference¹G. Maier, T. Preiss, H. P. Reisenauer, B. A. Hess, Jr., and L. J. Schaad, J. Am. Chem. Soc. **116**, 2014 (1994).

Si₂O₃⁻

Threshold for electron detachment from ground-state Si₂O₃⁻ = 15330(800) gas PE¹

There is evidence for the presence of a presumably less stable isomer of Si₂O₃⁻ which has a threshold for electron detachment = 7260(800) gas PE¹

Reference

¹L.-S. Wang, H. Wu, S. R. Desai, J. Fan, and S. D. Colson, *J. Phys. Chem.* **100**, 8697 (1996).

Ge₂O₃⁻

Threshold for electron detachment from ground-state Ge₂O₃⁻ = 19610(800) gas PE¹

Reference

¹L.-S. Wang, H. Wu, S. R. Desai, J. Fan, and S. D. Colson, *J. Phys. Chem.* **100**, 8697 (1996).

O₂N-NO

In nitrogen- and neon-matrix studies,^{4,8} excitation in the region of a weak absorption between 700 and 900 nm, with maximum near 720 nm, leads to isomerization forming O=N-O-N=O.

		C _s	Structure: MW ³				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.	
<i>a'</i>	1	N=O stretch	1829.54	gas	IR	1,5,7,10	
			1830.2s	Ne	IR	8	
			1830.0(2.5)	Ar	IR	11	
			1840m	N ₂	IR	4,11	
			1867vs	O ₂	IR	2	
	2		1861	NO	IR,Ra	6	
			1652	gas	IR	5	
			1643.3s	Ne	IR	8	
			1630.0(2.5)	Ar	IR	11	
			1630vs	N ₂	IR	4,11	
<i>b'</i>	3	NO ₂ s-stretch	1596s	O ₂	IR	2	
			1593	NO	IR,Ra	6	
			1304.25	gas	IR	1,5,9	
			1302.5vs	Ne	IR	8	
			1298.0(2.5)	Ar	IR	11	
	4		1302s	N ₂	IR	4,11	
			1303	O ₂	IR	2	
			1298	NO	IR,Ra	6	
			773	gas	IR	1,5	
			773.1wm	Ne	IR	8	
<i>c'</i>	5	NO ₂ deform.	777.1(2.5)	Ar	IR	11	
			776wm	N ₂	IR	4,11	
			788	O ₂	IR	2	
			787	NO	IR,Ra	6	
			414	gas	IR	5	
<i>d'</i>	6	N-N stretch	420w	N ₂	IR	4	
			405	NO	Ra	6	
			241	gas	IR	5	
<i>e'</i>	7	NO ₂ wag	266	NO	Ra	6	
			205	NO	Ra	6	

<i>a''</i>	8	NNO ₂ OPLA	627	NO	Ra	6
	9	Torsion	63	gas	IR	5
			70	NO	Ra	6

A₀=0.415; B₀=0.141; C₀=0.105 MW³IR^{9,10}

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O=N-O-N=O

In nitrogen-matrix studies,¹ absorption maxima were observed at 398, 381, and 363 nm. Excitation in the 370–480 nm spectral region resulted in photoisomerization into O₂N-NO.^{1,3}

		C _{2v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.	
<i>a'</i>	1	N=O s-stretch	1740	NO	Ra	2	
			973.6m	Ne	IR	3	
			968.4(2.5)	Ar	IR	4	
			969w	N ₂	IR	1	
			973m	NO	IR,Ra	2	
	2		387m	N ₂	IR	1	
			395	NO	Ra	2	
			366s	N ₂	IR	1	
			140	NO	Ra	2	
			105H	NO	Ra	2	
<i>b'</i>	3	NON bend	1697.2vs	Ne	IR	3	
			1685.9(2.5)	Ar	IR	4	
			1690s	N ₂	IR	1	
			1687vs	NO	IR	2	
			1661w	N ₂	IR	1	
	4		877vw	N ₂	IR	1	
			865vw	NO	IR	2	
			704vv	N ₂	IR	1	
			705vv	NO	IR	2	

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FONO₂⁺ $T^a=48090(120)$ gas PE¹ $\tilde{E}^{\prime\prime} 2A'$ C_s
 $T^a=40180(120)$ gas PE¹ $\tilde{D}^{\prime\prime} 2A'$ C_s
 $T^a=30980(120)$ gas PE¹ $\tilde{C}^{\prime\prime} 2A''$ C_s
 $T^a=24450(120)$ gas PE¹ $\tilde{B}^{\prime\prime} 2A'$ C_s
 $T^a=3550(120)$ gas PE¹ $\tilde{A}^{\prime\prime} 2A'$ C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			1090(40)	gas	PE	1

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			660(40)	gas	PE	1

^aFrom vertical ionization potentials.**Reference**¹D. Wang, P. Jiang, and Q. Zhang, Chem. Phys. Lett. **262**, 771 (1996).**CIONO₂⁺** $T^a=58250(120)$ gas PE¹ $T^a=36630(120)$ gas PE¹ $\tilde{E}^{\prime\prime} 2A''$ C_s
 $T^a=25170(120)$ gas PE¹ $\tilde{D}^{\prime\prime} 2A'$ C_s
 $T^a=18640(120)$ gas PE¹ $\tilde{B}, \tilde{C}^{\prime\prime} 2A'', 2A'$ C_s
 $T^a=11130(120)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			545(50)	gas	PE	1
			415(50)	gas	PE	1

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			4680(120)	gas	PE	1

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			770(50)	gas	PE	1

$\tilde{X}^{\prime\prime} 2A''$		C _s				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			1240(60)	gas	PE	1

^aFrom vertical ionization potentials.**Reference**¹D. Wang, Y. Li, P. Jiang, X. Wang, and B. Chen, Chem. Phys. Lett. **260**, 99 (1996).**BrONO₂⁺** $\tilde{E}^{\prime\prime} 2A'$ C_e
 $T^a=24700(200)$ gas PE¹ $\tilde{D}^{\prime\prime} 2A''$ C_s
 $T^a=20330(200)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'		BrO stretch	670(60)	gas	PE	1

 $\tilde{C}^{\prime\prime} 2A'$ C_s
 $T^a=15330(200)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'		NO ₂ a-stretch	1130(60)	gas	PE	1
		NO ₂ s-stretch	1050(60)	gas	PE	1

 $\tilde{B}^{\prime\prime} 2A'$ C_s
 $T^a=9520(200)$ gas PE¹ $\tilde{A}^{\prime\prime} 2A'$ C_s
 $T^a=5240(200)$ gas PE¹ $\tilde{X}^{\prime\prime} 2A''$ C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			400(60)	gas	PE	1

 $A=2500(200)$ gas PE¹^aFrom vertical ionization potential.**Reference**¹D. Wang and P. Jiang, J. Phys. Chem. **100**, 4382 (1996).

cyc-F₂CO₂

$\tilde{\chi}$	C_{2v}	Structure: IR ^{2,3}				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CO ₂ s-stretch	1466.59vs 1456 ^a	gas Ar	IR IR	1,2 2
	2	CF ₂ s-stretch	917.90m 916 ^a	gas Ar	IR IR	1,2 2
	3	CF ₂ scissors	772m	gas	IR	1
	4	OCO deform.	511vw	gas	IR	1
	6	CF ₂ a-stretch	1259vs 1249 ^a	gas Ar	IR IR	1 2
	7	CF ₂ rock	557w	gas	IR	1
b_2	8	CO ₂ a-stretch	1062.10s 1055 ^a	gas Ar	IR IR	1,2 2
	9	CF ₂ wag	617w	gas	IR	1

$A_0=0.266$; $B_0=0.211$; $C_0=0.163$ IR²

^aUnpublished work cited in Ref. 2.

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FONO₂

$\tilde{\chi}$	C_s	Structure: ED ⁴				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	NO ₂ a-stretch	1760.9vs 1758.4	gas Ne	IR,Ra IR	1-4 4
	2	NO ₂ s-stretch	1301.2vs 1300.9	gas Ne	IR,Ra IR	1-4 4
	3	OF stretch	927m 929.8	gas Ne	IR,Ra IR	1-4 4
	4	NO ₂ scissors	803.4s 802.7	gas Ne	IR,Ra IR	1-4 4
	5	O-N stretch	632.9vw 635.0	gas Ne	IR,Ra IR	1-4 4
	6	NO ₂ rock	458.5ms 459.9	gas Ne	IR,Ra IR	1-4 4
	7	OF rock	302.6vw 300.0	gas Ne	IR,Ra IR	2,3 4
	8	NO ₂ wag	707.3m 710.0	gas Ne	IR IR	2,4 4
	9	Torsion	151.6vvw 161.0	gas Ne	IR,Ra IR	2,3 4

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CIONO₂

The onset of continuous absorption by gas-phase CIONO₂ occurs near 25000 (400 nm).³ Except for two possible shallow maxima, the absorption continues to rise out to the 185 nm cutoff of the observations.

$\tilde{\chi}$	C_s	Structure: MW ^{5,10} IR ⁸ ED ¹⁷				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	NO ₂ a-stretch	1736.9vs 1727.6s 1734.0s 1730.0vs	gas Ar N ₂ O ₂	IR,Ra IR IR IR	1,2,4,13 21 9,11,16 11,16 7,16
	2	NO ₂ s-stretch	1292.7vs 1285.8vs 1291.4vs 1289.7s	gas Ar N ₂ O ₂	IR,Ra IR IR IR	1,2,4,13 18,21 9,11,16 11,16 7,16
	3	ClO stretch	809.4s 801.6m 809.7wm 808.1m	gas Ar N ₂ O ₂	IR,Ra IR IR IR	1,2,4,13 21 9,11,16 11,16 7,16
	4	Mixed	780.22ms 775.6m 778.6m 776.3m 563.1s 560.9ms 563.2wm 559.0m	gas Ar N ₂ O ₂ Ar N ₂ O ₂ O ₂	IR,Ra IR IR IR IR,Ra IR IR IR	1,2,4,13 20,21 9,11,16 11,16 7,16 1,2,21 9,11,16 11,16 7,16
	5	Mixed	434.0m 426 436.5w 435.0	gas Ar N ₂ O ₂	IR,Ra IR IR IR	1,2,21 9,11 11 7
	6	NO ₂ rock	273.3vvw 711.0w 710.8w 708.4w 704.6wm	gas gas Ar N ₂ O ₂	IR,Ra IR IR IR IR	1,2,21 1,2,21 1,2,21 9,11,16 11,16 7,16
	7	ClO rock	120.16	gas	IR,Ra	1,2,6,8
	8	NO ₂ wag	273.3vvw 711.0w 710.8w 708.4w 704.6wm	gas gas Ar N ₂ O ₂	IR,Ra IR IR IR IR	1,2,21 1,2,21 9,11,16 11,16 7,16
	9	Torsion	120.16	gas	IR,Ra	1,2,6,8

$A_0=0.404$; $B_0=0.093$; $C_0=0.075$ MW^{5,10,22}
Torsional barrier = 1900(100) gas IR^{6,8}

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BrONO₂

\tilde{X}		C _s	Structure: ED ²			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	NO ₂ a-stretch	1714vs	gas	IR	1
			1709vs	Ne	IR	1
			1702vs	N ₂	IR	1
	2	NO ₂ s-stretch	1288vs	gas	IR	1
			1285vs	Ne	IR	1
			1285vs	N ₂	IR	1
	3	NO ₂ scissors	806vs	gas	IR	1
			802vs	Ne	IR	1
			805vs	N ₂	IR	1
	4	OBr stretch	750w	Ne	IR	1
			750w	N ₂	IR	1
<i>a''</i>	5	NO ₂ rock	564s	gas	IR	1
			563s	Ne	IR	1
			574s	N ₂	IR	1
			569s			
<i>a''</i>	8	ONO ₂ OPLA	728wm	gas	IR	1
			723vw	Ne	IR	1
			725wm	N ₂	IR	1

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CF₃O

\tilde{A}^2A_1		C _{3v}	$\tilde{A}-\tilde{X}$ 330–400 nm			
T ₀	gas	LF ^{3–5}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CO stretch	634(2)	gas	LF	5
<i>e</i>	6	FCO deform.	425(2)	gas	LF	3–5

$\tau_0=30.1$ ns gas LF³
 $B_0=0.195$ LF⁵

\tilde{X}^2E		C _{3v}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
<i>a</i> ₁	1	CO stretch	1225T	gas	LF
	2	CF ₃ s-stretch	907T	gas	LF
<i>e</i>	4	CF ₃ a-stretch	1221.6	Ar	IR
	6	FCO deform.	483T	gas	LF

$A=-41(5)$ LF⁵
 $B_0=0.198$ LF⁵

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CF₃S

\tilde{A}^2A_1		C _{3v}	$\tilde{A}-\tilde{X}$ 350–380 nm			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CS stretch	313.8(4) ^a	gas	LF,FD	1
			$\tau_0=2.7(4)\mu\text{s}$	gas	LF ¹	

^a ω .

Reference

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CF₄⁺

\tilde{D}^2A_1		T _d	Structure: PE,EF ⁷ EM ¹⁴			
T ₀ ^a	gas	PE ^{2,3,5} TPEPICO ¹³				
	gas	EF ⁶	$\tilde{D}-\tilde{C}$ 350–420 nm			

Broad, unstructured emission maxima at 189 and 160 nm (52900 and 62500) which appear on He⁺ or electron impact on CF₄ have been interpreted as arising from the $\tilde{D}-\tilde{B}$ and $\tilde{D}-\tilde{A}$ transitions of CF₄⁺, respectively.⁶

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CF stretch	800(1)	gas	PE,EF	3,5,6

$\tau=2.1(2)$ ns gas EF¹
 $B_0=0.180(3)^b$ EF⁷

\tilde{C} 2F_2 T_d Structure: PE, EF⁷EM¹⁴
 $T_0^a=51230(160)$ gas PE^{2,3,5}T-PEPICO¹⁵

$T_0(\tilde{D}-\tilde{C})=27662.2$ gas EF⁶EM¹⁴ $\tilde{D}-\tilde{C}$ 350–420 nm
 Broad, unstructured emission maxima at 290 and 230 nm (34500 and 43500) which result from He⁺ or electron impact⁶ on CF₄ and from exposure of CF₄ to synchrotron radiation¹⁰ of wavelength shorter than 57.4 nm have been interpreted as arising from the $\tilde{C}-\tilde{A}$ and $\tilde{C}-\tilde{X}$ transitions of CF₄⁺, respectively.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CF stretch	727.4	gas	PE, EF EM	2,3,5,6 14

$A=+16(1)$ EF⁶⁻⁸

$\tau=9.0(9)$ ns gas EF¹EM^{9,11}T-PEFCO¹³

$B_0=0.169$ EM¹⁴

\tilde{B} 2E

$T_0^a=23800(1000)$ gas PE²⁻⁵

Dissociates into CF₃⁺ + F, probably by internal conversion to the \tilde{A} 2F_2 state. T-PEPICO¹²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CF stretch	810(80)	gas	PE	3-5
e		Deformation	500(100)	gas	PE	3-5

\tilde{A} 2F_2

$T_0^a=14100(1000)$ gas PE²⁻⁵

Direct dissociation into CF₃⁺ + F. T-PEPICO¹²

\tilde{X} 2F_1

Direct dissociation into CF₃⁺ + F. T-PEPICO¹²

^aMeasured with respect to onset of first photoelectron band, estimated by Ref. 3 at 15.35 eV.

^bFrom computer simulation of emission bands.

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CF₂Cl₂⁺

\tilde{H} C_{2v}
 $T^a=68200(1000)$ gas PE²⁻⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	CCl ₂ stretch	565T	gas	PE	4

\tilde{G} C_{2v}
 $T_0=56160(160)$ gas PE¹⁻⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	CCl ₂ stretch	550(80)	gas	PE	4

\tilde{E} , \tilde{F} 2A_2 , 2A_1 C_{2v}
 $T^a=36500(160)$ gas PE¹⁻⁴

A broad absorption with maximum near 310 nm (32300) which appears on argon-resonance photolysis of CF₂Cl₂ isolated in solid argon and which has a photodecomposition threshold between 420 and 340 nm has been assigned⁶ to the $\tilde{E}, \tilde{F}-\tilde{X}$ transitions of CF₂Cl₂⁺.

\tilde{D} 2B_2 C_{2v}
 $T_0=19300(35)$ gas PE^{1-4,8}

It has been proposed⁷ that an unstructured emission in the 200–400 nm region, with its principal contribution at wavelengths longer than 300 nm, which appears on electron bombardment of CF₂Cl₂ at electron energies greater than that for the onset of the \tilde{D} state of CF₂Cl₂⁺ may be contributed by the $\tilde{D}-\tilde{X}$ transition of CF₂Cl₂⁺.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	3	CF ₂ scissors	374(5)	gas	PE	1,3,4,8

\tilde{C} 2A_1 C_{2v}
 $T^a=13880(160)$ gas PE¹⁻⁴

\tilde{B} 2B_1 C_{2v}
 $T_0=10844(35)$ gas PE^{1-4,8}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CF ₂ stretch	1097(80)	gas	PE	3,4
	2	CCl ₂ stretch	565(80)	gas	PE	4
	4	CCl ₂ scissors	301(5)	gas	PE	4,8

\tilde{A} 2A_2 C_{2v}
 $T^a=6370(160)$ gas PE¹⁻⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	1	CF ₂ stretch	1056(5)	gas	PE	8

\tilde{X}^2B_2 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	3	CF ₂ scissors	609wm	Ar	IR	5
	4	CCl ₂ scissors	457(5)	gas	PE	8
		CF ₂ a-stretch	1234vs	Ar	IR	5
		CCl ₂ a-stretch	1067m	Ar	IR	5
		FCCl deform.	424m	Ar	IR	5
		FCCl deform.	406m	Ar	IR	5

^aFrom vertical ionization potential.

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 CCl_4^+ \tilde{D}^2A_1 T_d $T_0^a=68800(900)$ gas PE²Fragments, producing CCl_4^+ , in $<10^{-10}$ s. T-PEPICO⁶ \tilde{C}^2F_2 T_d $T_0^a=39290(900)$ gas PE²⁻⁴Fragments, producing CCl_2^+ , in $<10^{-10}$ s. T-PEPICO⁶ \tilde{B}^2E $T_0^a=15330(240)$ gas PE²⁻⁴Dissociates into $\text{CCl}_3^+ + \text{Cl}$, probably by internal conversion to the \tilde{A}^2T_2 state. PEPICO⁵ A^2F_2 T_d $T_0^a=6450(320)$ gas PE²⁻⁴Direct dissociation into $\text{CCl}_3^+ + \text{Cl}$. PEPICO⁵ \tilde{X}^2F_1 T_d Direct dissociation into $\text{CCl}_3^+ + \text{Cl}$. PEPICO⁵^aThe first ionization potential is taken as 11.47(1) eV, as in the photoionization study of Ref. 1.^bFrom vertical ionization potential.

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 SiF_4^+ \tilde{D}^2A_1 T_d Structure: PE, EF⁹ $T_0^a,b=50800(200)$ gas PE²TPE¹³EF⁷EM^{8,10} $\tilde{D}-\tilde{C}$ 530–590 nm $\tilde{D}-\tilde{C}$ band origin measured at 18146.8 in emission studies on a cooled beam.⁷Continuous emission between 570 and 730 nm, with a maximum near 610 nm (16400), has been assigned to the $\tilde{D}-\tilde{C}$ transition.^{8,10,12}Broad, unstructured emission maxima at 370 and 304 nm (27000 and 32900) which appear on ion, electron, or photon impact on SiF_4 have been interpreted as arising from the $\tilde{D}-\tilde{B}$ and $\tilde{D}-\tilde{A}$ transitions of SiF_4^+ , respectively.^{6,8}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	SiF stretch	743.4(5)	gas	EF,TPE	7,13
	2	Deformation	258(40)	gas	TPE	13
	3	SiF stretch	1178(40)	gas	TPE	13

 $\tau=9.30(4)$ ns gas EF¹EM¹²TPEFCO¹³ $B_0=0.136(1)^c$ EF⁹ \tilde{C}^2T_2 T_d Structure: PE, EF⁹ $T_0^a=33130(100)$ gas PE^{2,3,5}TPE¹³EF⁷EM^{8,10} $\tilde{D}-\tilde{C}$ 530–590 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	SiF stretch	706.6(5)	gas	PE,EF	3,5,7,13
	2	Deformation	159.0(5)	gas	EF	7
	4	Deformation	431.0(5)	gas	PE,EF	3,5,7,13

 $A=+6.9(2)$ EF^{7,9,11} $B_0=0.132^c$ PE,EF⁹ \tilde{B}^2E $T_0^a=22580(100)$ gasPE^{2,3,5}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	SiF stretch	685(50)	gas	PE	5

 \tilde{A}^2F_2 T_d $T_0^a=17000(1000)$ gasPE^{2,3,5} \tilde{X}^2F_1 ^aMeasured with respect to a first ionization potential of 15.19 eV, estimated^b by extrapolation of the photoionization efficiency curve for SiF_4 .^bFrom vertical ionization potential.^cFrom Franck-Condon analysis of the photoelectron spectrum and computer simulation of the $\tilde{D}-\tilde{C}$ emission.^dDynamic Jahn-Teller distortion, probably to C_{3v} .^{7,9}

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SiCl₄⁺

\tilde{D} 2A_1 T_d
 $T_0=48900(400)$ gas PE^{1,2}TPE⁸
Fragments to SiCl₂⁺. T-PEPICO⁷

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	SiCl stretch	379(16)	gas	TPE	8

\tilde{C} 2F_2
 $T_0=26620(160)$ gas PE^{1,2}TPE⁸

A broad, unstructured emission with maxima at 410 and 570 nm (24400 and 17500) has been assigned^{5,6,9} to the $\tilde{C}-\tilde{X}$ and $\tilde{C}-\tilde{A}$ transitions, respectively, of gas-phase SiCl₄⁺. Using pulsed synchrotron excitation⁶ of SiCl₄, as well as TPEFCO measurements,⁸ a radiative lifetime of 38.4(1) ns has been determined for the \tilde{C} state of SiCl₄⁺.

A broad absorption with maximum at 475 nm (21000) which appears on argon-resonance photolysis of SiCl₄ isolated in an argon matrix has been assigned⁴ to the $\tilde{C}-\tilde{X}$ transition of SiCl₄⁺. The energy difference is attributed to structural relaxation in the argon matrix. The absorption can be destroyed by exposure of the sample to 290–1000 nm radiation.

\tilde{B} 2E
 $T_0=13880(400)$ gas PE^{1,2}

\tilde{A} 2F_2
 $T_0=7750(160)$ gas PE^{1,2}

X 2F_1

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		SiCl ₂ a-stretch	717T	Ar	IR	4

⁹From vertical ionization potential.

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GeCl₄⁺

\tilde{D} 2A_1 T_d
 $T_0=51070(400)$ gas PE^{1,2}TPE⁵

\tilde{C} 2F_2 T_d
 $T_0=21620(240)$ gas PE^{1,2}TPE⁵

A broad, unstructured emission with maxima at 495 and 615 nm (20200 and 16300) has been assigned^{3,4} to the $\tilde{C}-\tilde{X}$ and $\tilde{C}-\tilde{A}$ transitions, respectively, of gas-phase GeCl₄⁺. Using pulsed synchrotron excitation of GeCl₄, a radiative lifetime of 65.4(4) ns has been determined⁴ for the \tilde{C} state of GeCl₄⁺.

\tilde{B} 2E
 $T_0=8880(1100)$ gas PE^{1,2}TPE⁵

\tilde{A} 2F_2
 $T_0=5650(1100)$ gas PE^{1,2}TPE⁵

X 2F_1 **References**

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OCIO₃

In a neon matrix, absorption between 380 and 510 nm, with a maximum at 23530 (425 nm) and with partially resolved structure, can be assigned¹ to OCIO₃.

X C_{3v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	ClO ₃ s-stretch	1161vs	Ne	IR	1
	2	ClO stretch	874w	Ne	IR	1
	3	Deform.	576m	Ne	IR	1
e	4	ClO ₃ a-stretch	1234s	Ne	IR	1
	5	Deform.	646m	Ne	IR	1

Reference

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Cl₂O₃

An unstructured gas-phase absorption between 220 and 320 nm, with maximum at 37450 (267 nm), has been assigned^{2,4,5,8} to Cl₂O₃.



Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		1225vs	gas	IR	1,4	
		1216.0	Ne	IR	6	
		1226.1	Ar	IR	3,6	
		1213.5	Ar	IR	6	
		1203.1	Ne	IR	6	
		1200.7	Ar	IR	6	
		1057s	gas	IR	4	
		1060	Ne	IR	6	
		1058.4	Ar	IR	3,6	
		740w	gas	IR	4	
		560wm	gas	IR	4	

$$A_0=0.288; B_0=0.070; C_0=0.059 \text{ MW}^7$$

References

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SF₄⁺

$$T^a=51600(800) \text{ gas PE}^{1-3}$$

$$T^a=46800(400) \text{ gas PE}^{1-3}$$

$$T^a=43000(400) \text{ gas PE}^{1-3}$$

$$\tilde{C}, \tilde{D}^2B_2, ^2A_1 \quad C_{2v}$$

$$T^a=31460(400) \text{ gas PE}^{1-3}$$

$$\tilde{A}, \tilde{B}^2A_2, ^2B_1 \quad C_{2v}$$

$$T^a=25580(400) \text{ gas PE}^{1-3}$$

$$\tilde{X} \quad ^2A_1 \quad C_{2v}$$

^aFrom vertical ionization potential. The adiabatic first ionization potential of SF₄ is estimated to be 11.90 eV.³

References

- ¹A. H. Cowley, M. Lattman, and M. L. Walker, *J. Am. Chem. Soc.* **101**, 4074 (1979).
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8.12. Six-Atomic Molecules**SiH₅⁺**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1		HH stretch	3693.97(6)	gas	PF	1

$$A_0=2.519^a; B_0=1.689(3); C_0=1.639(3) \text{ PF}^1$$

^aFrom *ab initio* calculations; fixed in the fit.

Reference

$$^1\text{D. W. Boo and Y. T. Lee, } J. \text{ Chem. Phys.} \text{ } \mathbf{103}, 514 \text{ (1995).}$$

LiBH₄

\tilde{X}	C _{3v}	Structure: MW ^{1,2}				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			700(150)	gas	MW	1
<i>e</i>			400(150)	gas	MW	1

$$B_0=0.769 \text{ MW}^{1,2}$$

LiBD₄

$$\tilde{X} \quad C_{2v}$$

$$B_0=0.625 \text{ MW}^2$$

References

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NaBH₄

\tilde{X}	C _{3v}	Structure: MW ^{1,2}				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			435T	gas	MW	1

$$B_0=0.322 \text{ MW}^1$$

NaBD₄

$$\tilde{X} \quad C_{3v}$$

$$B_0=0.268 \text{ MW}^2$$

References

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KBH₄

\tilde{X}	C _{3v}	Structure: MW ¹				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			360T	gas	MW	1

 $B_0=0.210$ MW¹**Reference**¹Y. Kawashima and E. Hirota, *J. Chem. Phys.* **102**, 6961 (1995).**HBeH₂BeH**

\tilde{X}						
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		BeH ₂ stretch	1982	Ar	IR	1
		BeHBe stretch	1536	Ar	IR	1
		BeHBe stretch	1420	Ar	IR	1
			872	Ar	IR	1
			558	Ar	IR	1

DBeD₂BeD

\tilde{X}						
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		BeD ₂ stretch	1567	Ar	IR	1
		BeDBe stretch	1067	Ar	IR	1
			651	Ar	IR	1

Reference¹T. J. Tague, Jr. and L. Andrews, *J. Am. Chem. Soc.* **115**, 12111 (1993).**HMgH₂MgH**

\tilde{X}						
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1531.0T	Ar	IR	1
			1164.2T	Ar	IR	1
			1013.7T	Ar	IR	1
			613.9T	Ar	IR	1

DMgD₂MgD

\tilde{X}						
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1127.7T	Ar	IR	1
			844.5T	Ar	IR	1
			736.9T	Ar	IR	1

Reference¹T. J. Tague, Jr. and L. Andrews, *J. Phys. Chem.* **98**, 8611 (1994).**U₂H₄** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		UH stretch	1547T	Ar	IR	1
		UH stretch	1495.0	Ar	IR	1

U₂D₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		UD stretch	1102.8	Ar	IR	1
		UD stretch	1067.9	Ar	IR	1

Reference¹P. F. Souter, G. P. Kushto, L. Andrews, and M. Neurock, *J. Am. Chem. Soc.* **119**, 1682 (1997).**CH₃CoH** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH ₃ stretch	2982.0	Ar	IR	2
		CH ₃ stretch	2942.1	Ar	IR	2
		CoH stretch	1722.6	Ar	IR	1,2
		Deformation	1168.5	Ar	IR	2
		Deformation	573.2	Ar	IR	2
			568.9			
		CoC stretch	530.8	Ar	IR	1,2

CD₃CoD \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CoD stretch	1241.0	Ar	IR	1

References¹W. E. Billups, S.-C. Chang, R. H. Hauge, and J. L. Margrave, *J. Am. Chem. Soc.* **115**, 2039 (1993).²W. E. Billups, S.-C. Chang, R. H. Hauge, and J. L. Margrave, *J. Am. Chem. Soc.* **117**, 1387 (1995).

CH₃ZnH

\tilde{X}		C _{3v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₃ stretch	2919.8	Ar	IR	1
	2	ZnH stretch	1866.1	Ar	IR	1
	3	CH ₃ deform.	1179.3	Ar	IR	1
	4	ZnC stretch	566.5	Ar	IR	1
<i>e</i>	7	CH ₃ rock	689.4s	Ar	IR	1
			686.6			
	8	CZnH deform.	442.6	Ar	IR	1

CD₃ZnD

\tilde{X}		C _{3v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	ZnD stretch	1344.7	Ar	IR	1
	3	CD ₃ deform.	921.1	Ar	IR	1
	4	ZnC stretch	516.1	Ar	IR	1
	7	CD ₃ rock	530.5	Ar	IR	1
			527.9			
	8	CD ₃ deform.	317.2	Ar	IR	1

Reference

¹T. M. Greene, L. Andrews, and A. J. Downs, J. Am. Chem. Soc. **117**, 8180 (1995).

CH₃CdH

\tilde{X}		C _{3v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₃ stretch	2928.5	Ar	IR	1
	2	CdH stretch	1760.5	Ar	IR	1
	4	CdC stretch	508.6	Ar	IR	1
	7	CH ₃ rock	687.3	Ar	IR	1
			685.4			
	8	CCdH deform.	433.2	Ar	IR	1
			430.1			

CD₃CdD

\tilde{X}		C _{3v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CdD stretch	1264.4	Ar	IR	1
	4	CdC stretch	465.3	Ar	IR	1
	7	CD ₃ rock	523.6	Ar	IR	1
			521.9			
	8	CCdD deform.	310.6	Ar	IR	1

Reference

¹T. M. Greene, L. Andrews, and A. J. Downs, J. Am. Chem. Soc. **117**, 8180 (1995).

CH₃HgH

\tilde{X}		C _{3v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CH ₃ stretch	2921.2w	Ar	IR	1,2
	2	HgH stretch	1954.9vs	Ar	IR	1,2
	3	CH ₃ deform.	1191.7vw	Ar	IR	1,2
	4	HgC stretch	534.0w	Ar	IR	1,2
<i>e</i>	5	CH ₃ stretch	2990.6mT	Ar	IR	1
	6	CH ₃ deform.	1425.0w	Ar	IR	1,2
	7	CH ₃ rock	779.8	Ar	IR	1,2
			777.9			
	8	CHgH deform.	526.5	Ar	IR	1,2

CD₃HgD

\tilde{X}		C _{3v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	CD ₃ stretch	2127.7w	Ar	IR	1,2
	2	HgD stretch	1400.7vs	Ar	IR	1,2
	3	CD ₃ deform.	1024.5vwT	Ar	IR	1
	4	HgC stretch	487.0	Ar	IR	2
<i>e</i>	5	CD ₃ stretch	2233.9m	Ar	IR	1
	6	CD ₃ deform.	1041.1wT	Ar	IR	1,2
	7	CD ₃ rock	593.5m	Ar	IR	1,2
			592.1			
	8	CHgD deform.	374.5	Ar	IR	2

References

¹N. Legay-Sommaire and F. Legay, Chem. Phys. Lett. **217**, 97 (1994).
²T. M. Greene, L. Andrews, and A. J. Downs, J. Am. Chem. Soc. **117**, 8180 (1995).

H₂CBH₂

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁		CH ₂ scissors	1414.7	Ar	IR	1
		BH ₂ scissors	1239.8	Ar	IR	1
<i>b</i> ₁		BH ₂ OPLA	972.4	Ar	IR	1
		BH ₂ a-stretch	2560.6	Ar	IR	1

D₂CBD₂

\tilde{X}		C _{2v}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁		BD ₂ scissors	1177	Ar	IR	1
		BD ₂ OPLA	794.8	Ar	IR	1
<i>b</i> ₁		BD ₂ a-stretch	1923	Ar	IR	1

Reference

¹P. Hassanzadeh, Y. Hannachi, and L. Andrews, J. Phys. Chem. **97**, 6418 (1993).

CH₃BH \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CH stretch	2979.2T	Ar	IR	1
		BH stretch	2560.6T	Ar	IR	1

CD₃BD \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		BD stretch	1935T	Ar	IR	1

Reference

¹P. Hassanzadeh, Y. Hannachi, and L. Andrews, *J. Phys. Chem.* **97**, 6418 (1993).

CH₃GaH

In an argon matrix, an absorption with maximum at 45900 (218 nm) behaves appropriately for assignment to CH₃GaH.¹

In an argon matrix, a weak, broad (100 nm FWHM) absorption at 16700 (600 nm) has also been attributed¹ to CH₃GaH. Irradiation of the sample in the region of this absorption leads to photodestruction of CH₃GaH.

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CH stretch	2970	Ar	IR	1
		CH stretch	2914	Ar	IR	1
		GaH stretch	1719.7	Ar	IR	1,2
			1712.5	Kr	IR	2
		CH ₃ deform.	1432	Ar	IR	1
		CH ₃ deform.	1176.4	Ar	IR	1,2
			1153.7	Kr	IR	2
		CH ₃ rock	753.1	Ar	IR	1,2
			747.8	Kr	IR	2
		CH ₃ rock	642	Ar	IR	1
		GaC stretch	528.6	Ar	IR	1,2
			520.0	Kr	IR	2

CD₃GaD \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CD stretch	2222	Ar	IR	1
		CD stretch	2137	Ar	IR	1
		GaD stretch	1243.6	Ar	IR	1,2
			1239.0	Kr	IR	2
		CD ₃ deform.	1026	Ar	IR	1
		CD ₃ deform.	747.0T ^a	Kr	IR	2
		CD ₃ rock	578.2	Ar	IR	1,2
		GaC stretch	525.0T ^b	Ar	IR	2

^aRef. 1 gives 899 for an argon matrix.

^bRef. 1 gives 484.

References

- ¹R. D. Lafleur and J. M. Parnis, *J. Phys. Chem.* **96**, 2429 (1992).
²Z. L. Xiao, R. H. Hauge, and J. L. Margrave, *Inorg. Chem.* **32**, 642 (1993).

CH₂=SiH₂

In an Ar or N₂ matrix, absorption maximum at 258 nm.¹⁻⁴ On irradiation at 254 nm, photoisomerizes to CH₃SiH.^{2,4}

 \tilde{X} C_{2v} Structure: MW⁵

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁		SiH s-stretch	2219m	Ar	IR	1-3
			2214m	N ₂	IR	2,4
		CH ₂ scissors	1350w	Ar	IR	2,3
			1350w	N ₂	IR	2,4
		Si=C stretch	985w	Ar	IR	1-3
			985w	N ₂	IR	1,2,4
		SiH ₂ scissors	927w	Ar	IR	1-3
			927w	N ₂	IR	2,4
<i>b</i> ₁		CH ₂ wag	741s	Ar	IR	1-3
			747s	N ₂	IR	2,4
<i>b</i> ₂		SiH a-stretch	2239m	Ar	IR	1-3
			2235m	N ₂	IR	2,4
		CH ₂ rock	817s	Ar	IR	1-3
			817s	N ₂	IR	2,4

A₀=3.493; B₀=0.493; C₀=0.432 MW⁵

CH₂=SiD₂

In an Ar matrix, absorption maximum at 259 nm.^{1,4}

 \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁		SiD s-stretch	1600m	Ar	IR	1,3
			1600	N ₂	IR	4
		CH ₂ scissors	1335	Ar	IR	3
			1352	N ₂	IR	4
		Si=C stretch	952w	Ar	IR	1,3
			952	N ₂	IR	4
<i>b</i> ₁		CH ₂ wag	719s	Ar	IR	1,3
			725	N ₂	IR	4
<i>b</i> ₂		SiD a-stretch	1635m	Ar	IR	1,3
			1635	N ₂	IR	4
		CH ₂ rock	759s	Ar	IR	1,3
			760	N ₂	IR	4
		SiD ₂ rock	396w	Ar	IR	1,3
			396	N ₂	IR	4

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- ¹G. Maier, G. Mihm, and H. P. Reisenauer, *Angew. Chem.* **93**, 615 (1981); *Angew. Chem. Int. Ed. Engl.* **20**, 597 (1981).
²H. P. Reisenauer, G. Mihm, and G. Maier, *Angew. Chem.* **94**, 864 (1982); *Angew. Chem. Int. Ed. Engl.* **21**, 854 (1982).
³G. Maier, G. Mihm, and H. P. Reisenauer, *Chem. Ber.* **117**, 2351 (1984).

⁴G. Maier, G. Mihm, H. P. Reisenauer, and D. Littmann, Chem. Ber. **117**, 2369 (1984).

⁵S. Bailleux, M. Bogey, J. Demaison, H. Bürger, M. Senzlober, J. Breidung, W. Thiel, R. Fajgar, and J. Pola, J. Chem. Phys. **106**, 10016 (1997).

H₂BNBH

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		BH stretch	2788.3	Ar	IR	1
		BH stretch	2426.5	Ar	IR	1
		BH stretch	2393.2	Ar	IR	1
		BNB a-stretch	1864.2	Ar	IR	1

Reference

¹C. A. Thompson, L. Andrews, J. M. L. Martin, and J. El-Yazal, J. Phys. Chem. **99**, 13839 (1995).

CH₂CCH

In the gas phase, absorption between 230 and 280 nm, with a maximum at 41320 (242 nm), has been assigned⁹ to CH₂CCH.

T₀=30109^a gas AB¹
Ar AB² 290–345 nm
288–359 nm

All bands in the gas-phase spectrum are diffuse.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C-C stretch	961(10)	gas	AB	1
			965(10)	Ar	AB	2

\tilde{X}^2B_1

C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH stretch	3322.29	gas	CC	5
			3308.8m	Ar	IR	2,4
		CH ₂ wag	687.18	gas	DL	6,8
			686.5m	Ar	IR	2
		C ₃ deformation	490T	gas	PE	3,7
			483.5m	Ar	IR	2,4

A₀=9.608; B₀=0.318; C₀=0.307 CC⁵DL⁸

CD₂CCD

\tilde{X}^2B_1

C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CD stretch	2546.8m	Ar	IR	2,4
			552.9m	Ar	IR	2
			479.8	Ar	IR	4
			350T	gas	PE	7

^aAssignment of gas-phase band origin is tentative. The extension of the progression to 27886 in the argon-matrix study suggests that as many as two quanta of the C-C stretching vibration may be excited in the gas phase band at 30109.

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cyc-(HC=NCH₂)⁺

\tilde{D}^2A' C_s
T^a=46200(800) gas PE¹⁻³

\tilde{C}^2A'' C_s
T^a=41200(800) gas PE¹⁻³

\tilde{B}^2A' C_s
T^a=23240(160) gas PE¹⁻³

\tilde{A}^2A'' C_s
T^a=7900(160) gas PE¹⁻³

\tilde{X}^2A' C_s

^aFrom vertical ionization potentials.

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CH₂CCH⁻

Threshold for electron detachment from ground-state CH₂CCH⁻=7410(65)
gas PE¹⁻²

CD₂CCD⁻

Threshold for electron detachment from ground-state CD₂CCD⁻=7380(65)
gas PE²

References

- J. M. Oakes and G. B. Ellison, J. Am. Chem. Soc. **105**, 2969 (1983).
- M. S. Robinson, M. L. Polak, V. M. Bierbaum, C. H. DePuy, and W. C. Lineberger, J. Am. Chem. Soc. **117**, 6766 (1995).

CH₃CC⁻

Threshold for electron detachment from ground-state CH₃CC⁻ = 21930(65) gas PE¹

\tilde{X}^2E C_{∞v}

Reference

- ¹M. S. Robinson, M. L. Polak, V. M. Bierbaum, C. H. DePuy, and W. C. Lineberger, J. Am. Chem. Soc. **117**, 6766 (1995).

CH₂=C=NH

\tilde{B} C_s
 T_0^a = 38900 gas AB^{2,8}
 38790 Ar AB⁴

\tilde{A} C_s
 T_0^a = 32840 gas AB^{2,8}
 32680 Ar AB⁴

$\tilde{A}-\tilde{X}$ 273–317 nm
 $\tilde{A}-\tilde{X}$ 273–306 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>			990	Ar	AB	4
\tilde{X} C _s Structure: MO ³ MW ⁶						
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	3	CCN a-stretch	2043.58	gas	IR	7
			2040vs	Ar	IR	1,4
	5	CCN s-stretch; NH deform.	1124wm	Ar	IR	1,4
	6	NH deform.	1000s	Ar	IR	1,4
	7	H ₂ CC OPLA	690m	Ar	IR	1,4
<i>a''</i>	11	Torsion	872m	Ar	IR	1,4

A₀ = 6.719; B₀ = 0.322; C₀ = 0.316 MW^{5,6}IR⁷

CD₂=C=ND

\tilde{B} C_s
 T_0 = 38790 Ar AB⁴

\tilde{A} C_s
 T_0 = 32680 Ar AB⁴

$\tilde{A}-\tilde{X}$ 272–306 nm

\tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	3	CCN a-stretch	1998vs	Ar	IR	1,4
	5	CD ₂ scissors	921m	Ar	IR	1,4
	6	ND deform.	800s	Ar	IR	1,4
<i>a''</i>	11	Torsion	648m	Ar	IR	1,4

^aBands originally tentatively assigned² to cyanoallyl.

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HCNCH₂

In an argon matrix, an intense absorption with maximum at 38800 (258 nm) and a weak, broad absorption with maximum near 31200 (320 nm) have been assigned¹ to HCNCH₂.

\tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	3	CNC a-stretch	1914.5s	Ar	IR	1
	5	CNC s-stretch	1183.6w	Ar	IR	1
	6	HCN deform.	866.3vs	Ar	IR	1
	8	CNC deform.	473.5wm	Ar	IR	1
<i>a''</i>	10	CH ₂ deform.	1127.3w	Ar	IR	1

DCNCD₂

\tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CNC a-stretch	1870.2vs	Ar	IR	1
			982.6w	Ar	IR	1
			900.1w	Ar	IR	1
			708.3ms	Ar	IR	1
			414.6w	Ar	IR	1

Reference

- ¹G. Maier, C. Schmidt, H. P. Reisenauer, E. Endlein, D. Becker, J. Eckwert, B. A. Hess, Jr., and L. J. Schaad, Chem. Ber. **126**, 2337 (1993).

cyclo-(HC=NCH₂)

(2H-Azirine)

 \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	CH stretch	3078.4w	Ar	IR	3
	2	CH stretch	3005.6w	Ar	IR	3
	3	C=N stretch	1669.1wm	Ar	IR	3
			1667.2wm			
	5		1240.5wm	Ar	IR	3
			1237.6wm			
	7		978.7vs	Ar	IR	3
			977.3			
			974.4			
			971.5			
			969.1			
a''	9	CH ₂ a-stretch	3061.5vw	Ar	IR	3
	12		772.9wm	Ar	IR	3
			767.1wm			

$$A_0=1.188; B_0=0.741; C_0=0.503 \text{ MW}^{1,2}$$

References¹R. G. Ford, J. Am. Chem. Soc. **99**, 2389 (1977).²M. Bogey, J.-L. Destombes, J.-M. Denis, and J.-C. Guillemin, J. Mol. Spectrosc. **115**, 1 (1986).³G. Maier, C. Schmidt, H. P. Reisenauer, E. Endlein, D. Becker, J. Eckwert, B. A. Hess, Jr., and L. J. Schaad, Chem. Ber. **126**, 2337 (1993).**HCCAsH₂** \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	CH stretch	3318	gas	IR	1	
2	CC stretch	2180	gas	IR	1	
	AsH ₂ stretch	2118	gas	IR	1	

Reference¹J.-C. Guillemin, L. Lassalle, P. Dréan, G. Włodarczak, and J. Demaison, J. Am. Chem. Soc. **116**, 8930 (1994).**CH₃CaS** \tilde{X} C_{3v}
 $B_0=0.117$ MW¹Structure: MW¹**CD₃CaS** \tilde{X} C_{3v}
 $B_0=0.100$ MW¹**Reference**¹J.-C. Guillemin, L. Lassalle, P. Dréan, G. Włodarczak, and J. Demaison, J. Am. Chem. Soc. **116**, 8930 (1994).**CH₂CHO** $\tilde{B}' 2A''$ C_s

$$T_0=28784.09(1) \text{ gas } AB^1LF^{2,4,6,12}FD^{10}PF^{13}$$

 $\tilde{B}-\tilde{X}$ 300–405 nm

The failure to detect fluorescence on excitation above 30000 suggests² the onset of predissociation near 330 nm. In the gas phase,¹³ both CH₃ and H atoms are produced above 28700. In the argon matrix experiments,⁵ the threshold for the photodecomposition of CH₂CHO to produce CH₃+CO was observed between 280 and 300 nm.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	4	CO stretch	1621	gas	FD,PF	10,12,13
	5	CH ₂ scissors	1405	gas	FD,PF	10,12,13
	6	OCH bend	1274	gas	PF,LF,DR	13,14
	7	CH ₂ rock	1122	gas	LF,FD	2,6,10,12
	8	CC stretch	917	gas	LF,FD	2,6,10,12
	9	CCO bend	449	gas	LF,FD	2,6,10,12
	10	HCO wag	595H	gas	LF,DR	14
	11	CH ₂ wag	436H	gas	PF,LF,DR	13,14
	12	Torsion	274H	gas	PF,LF,DR	13,14

$$\tau_0=190(2) \text{ ns gas LF}^{11,14}$$

$$A_0=2.103(4); B_0=0.344(1); C_0=0.296(1) \text{ LF}^6$$

 $\tilde{A}' 2A'$ C_s

$$T_0=8006 \text{ gas AB}^3$$

 $\tilde{A}-\tilde{X}$ 1000–1250 nm $\tilde{X} 2A''$ C_s Structure: MW^{8,9}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	4	CCO stretch	1543	gas	LF	2,6,12,14
			1542m HF	Ar	IR	5
			1525m			
	5	CH ₂ scissors	1486	gas	LF	14
	6	OCH deform.	1366	gas	LF	12,14
			1375m HF	Ar	IR	5
	7	CH ₂ rock	1143	gas	LF,PD	2,6,7,14
	8	CC stretch	957	gas	LF	14
	9	CCO bend	500	gas	LF,PD	2,6,7,14
	10	CHO wag	703H	gas	LF	14
a''	11	CH ₂ wag	557H	gas	LF	14
	12	Torsion	404H	gas	LF	14

$$A_0=2.224; B_0=0.382; C_0=0.326 \text{ LF}^6 \text{MW}^8$$

CD₂CDO $\tilde{B}' 2A''$ C_s

$$T_0=28840 \text{ gas LF}^2PF^{13}$$

 $\tilde{B}-\tilde{X}$ 300–411 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			980	gas	LF	2
			768	gas	LF	2

\tilde{X}^2A'' C_s

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'		C=O stretch	1540	gas	LF	2
			1513ms HF	Ar	IR	5
			1223w HF	Ar	IR	5
		CC stretch	1050	gas	LF	2
		CCO bend	445	gas	LF	2

 $A_0=1.442; B_0=0.336; C_0=0.272 \text{ MW}^9$

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 CH_2CHO^- Dipole-Bound State C_s
 $T_0=14712.747(5) \text{ gas } \text{PD}^{1-3}$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'		C-C stretch	1143	gas	PD	2
		CCO bend	499	gas	PD	2
a''		Torsion	102H	gas	PD	2

 $A_0=2.221(2); B_0=0.376; C_0=0.320 \text{ PD}^{2,3}$ \tilde{X} C_s Structure: PD²

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'		CCO bend	525.82	gas	PD	2,3
a''		Torsion	375T	gas	PD	2

 $A_0=2.494; B_0=0.362; C_0=0.316 \text{ PD}^{2,3}$ CD_2CDO^- Dipole-Bound State C_s
 $T_0=14665.97(5) \text{ gas } \text{PD}^{1,2,4}$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'			1382(10)	gas	PD	1
			981(10)	gas	PD	1
		CCO bend	437(10)	gas	PD	1
a''		Torsion	80HT	gas	PD	1

 $A_0=1.419(3); B_0=0.330; C_0=0.268 \text{ PD}^{2,4}$ \tilde{X}
 $A_0=1.554(1); B_0=0.319; C_0=0.264 \text{ PD}^2$

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- ⁴A. S. Mullin, K. K. Murray, C. P. Schulz, and W. C. Lineberger, *J. Phys. Chem.* **97**, 10281 (1993).

 CH_2NOH \tilde{X} C_s Structure: MW³

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	OH stretch	3650.29vs	gas	IR	1,4,7
	2	CH_2 stretch	3109.72w	gas	IR	1,4,7
	3	CH_2 stretch	2973.17wm	gas	IR	1,4,7
	4	=C=N stretch	1639.48wm	gas	IR	1,4,9
	5	CH_2 scissors	1410.48ms	gas	IR	1,4,9
	6	OH bend	1318.98vs	gas	IR	1,4,9
	7	CH_2 rock	1166m	gas	IR	1,4
	8	NO stretch	892.58s	gas	IR	1,4,6,9
	9	CNO deform.	530w	gas	IR	1,4
	10	CH_2 OPLA	952.61s	gas	IR	1,4,6
	11	CH_2 torsion	774.1m	gas	IR	1,4,6
	12	OH torsion	400T	gas	IR	1,4

 $A_0=2.258; B_0=0.396; C_0=0.336 \text{ MW}^{2,3,5,8}$

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- ⁸A. Klesing and D. H. Sutter, *Z. Naturforsch. A* **45**, 817 (1990).
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PH₃F₂

\tilde{X}		D _{3h}	Structure: IR ³			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a''</i>	3	PH ₃ deform.	1266.15m	gas	IR	1,3,5
	4	PF ₂ a-stretch	755.99vs	gas	IR	1,3,5
<i>e'</i>			755	Ar	IR	2
	5	PH ₃ a-stretch	2501.7s ^a	gas	IR	1,3,5
<i>e'</i>			2480w,br	Ar	IR	2
	6	PH ₃ deform.	965.45s ^b	gas	IR	1,3,5
			974	Ar	IR	2
	7	PF ₂ deform.	341.55m	gas	IR	4,5

 $A_0=2.868(15); B_0=0.159$ gas IR^{3,4}**PD₃F₂**

\tilde{X}		D _{3h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a''</i>	3	PD ₃ deform.	936m	gas	IR	5
	4	PF ₂ a-stretch	745vs	gas	IR	5
<i>e'</i>			749	Ar	IR	2
	5	PD ₃ a-stretch	1828s	gas	IR	5
<i>e'</i>	6	PD ₃ deform.	719	Ar	IR	2
	7	PF ₂ deform.	319m	gas	IR	5

^aEffective value; deperturbed value = 2488.48.^bDeperturbed value.**References**

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HNBBNH

\tilde{X}		D _{∞h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NH stretch	3682.8	Ar	IR	1
		BN stretch	1827.4	Ar	IR	1
		Deformation	413.3	Ar	IR	1

Reference

- ¹C. A. Thompson, L. Andrews, J. M. L. Martin, and J. El-Yazai, J. Phys. Chem. **99**, 13839 (1995).

H₂CCCC:

\tilde{X}		C _{2v}	Structure: MW ²			
			$A_0=9.489; B_0=0.150; C_0=0.148$ MW ^{1,2}			

D₂CCCC:

\tilde{X}		C _{2v}
$A_0=4.783; B_0=0.137; C_0=0.133$		MW ²

Reference

- ¹T. C. Killian, J. M. Vrtilek, C. A. Gottlieb, E. W. Gottlieb, and P. Thaddeus, Astrophys. J. **365**, L89 (1990).

HCCCNH⁺

\tilde{X}		C _{∞v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	NH stretch	3513.96	gas	LD	1
	3	CN stretch	2315.14	gas	DL	2

 $B_0=0.144$ LD¹MW³**References**

- ¹S. K. Lee and T. Amano, Astrophys. J. **323**, L145 (1987).
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HOC≡COH

\tilde{X}		C ₂

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i>	8	OH stretch	3586.2wm	Ar	IR	1
	10	Deformation	1212.4vw	Ar	IR	1

DOC≡COD

\tilde{X}		C ₂

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i>	8	OD stretch	2591.1s	Ar	IR	1
	9	Deformation	1302.9vs	Ar	IR	1
	10	Deformation	950.9wm	Ar	IR	1

Reference

- ¹G. Maier and C. Rohr, Liebigs Ann. **1996**, 307 (1996).

CH₂NO₂²B₁ C_{2v}

In the gas phase, photodissociation into CH₂NO+O and into H₂CO+NO has been observed between 240 and 270 nm.⁴

In an Ar matrix, threshold for photodecomposition into H₂CO + NO near 290 nm.¹

²A₂ C_{2v}T₀=12840(80) gas PE³

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	3	NO ₂ s-stretch	1444(25)	gas	PE	3
	4	CN stretch ?	936(25)	gas	PE	3
	5	NO ₂ scissors	605(25)	gas	PE	3

 \tilde{X} ²B₁ C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1	CH ₂ s-stretch	3055 HF	Ar	IR	1,2
	2	CH ₂ scissors	1419 HF	Ar	IR	1,2
	3	NO ₂ s-stretch	1300(25)	gas	PE	3
			1297 HF	Ar	IR	1,2
	4	CN stretch	954(25)	gas	PE	3
			986 HF	Ar	IR	1,2
	5	NO ₂ scissors	693 HF	Ar	IR	1,2
a ₂	6	Torsion	205 H	gas	PE	3
b ₁	7	CNO ₂ OPLA ^a	719 HF	Ar	IR	1,2
	8	H ₂ CN OPLA ^a	606 HF	Ar	IR	1,2
b ₂	9	CH ₂ a-stretch	3200 HF	Ar	IR	1,2
	10	NO ₂ a-stretch	1484 ^b HF	Ar	IR	1,2
			1461			
	11	CH ₂ rock	1095 HF	Ar	IR	1,2
	12	NO ₂ rock	484T ^c HF	Ar	IR	1,2

CD₂NO₂ \tilde{X} ²B₁ C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	2	NO ₂ s-stretch	1292(25)	gas	PE	3
			1296 DF	Ar	IR	1,2
	4	CN stretch + CD ₂ scissors	906 DF	Ar	IR	1,2
	5	NO ₂ scissors	668 DF	Ar	IR	1,2
a ₂	6	Torsion	140 H	gas	PE	3
b ₁	7	CNO ₂ OPLA	694 DF	Ar	IR	1,2
b ₂	10	NO ₂ a-stretch	1460 DF	Ar	IR	1,2

^aThe two out-of-plane modes are strongly mixed.^bIn Fermi resonance with ($\nu_4 + \nu_{12}$).^cEstimated from ($\nu_4 + \nu_{12}$).**References**¹M. E. Jacox, J. Phys. Chem. **87**, 3126 (1983).²M. E. Jacox, J. Phys. Chem. **91**, 5038 (1987).³R. B. Metz, D. R. Cyr, and D. M. Neumark, J. Phys. Chem. **95**, 2900 (1991).⁴D. R. Cyr, D. J. Leahy, D. L. Osborn, R. E. Continetti, and D. M. Neumark, J. Chem. Phys. **99**, 8751 (1993).**CH₂CFO** \tilde{B} T₀=29874 gas LF¹ $\tilde{B}-\tilde{X}$ 307–335 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	3	CO stretch	1790	gas	LF	1
	5	CF stretch	1253	gas	LF	1
	6	CH ₂ rock	911	gas	LF	1
	7	CC stretch	874	gas	LF	1
	8	FCO bend	537	gas	LF	1
	9	CCO bend	421	gas	LF	1

τ₀=81.0(5) nsgas LF¹ \tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	3	CO stretch	1724	gas	LF	1
	5	CF stretch	1211	gas	LF	1
	6	CH ₂ rock	906T	gas	LF	1
	7	CC stretch	847	gas	LF	1
	8	FCO bend	584	gas	LF	1
	9	CCO bend	416T	gas	LF	1

Reference

¹M. Furubayashi, I. Bridier, S. Inomata, N. Washida, and K. Yamashita, J. Chem. Phys. **106**, 6302 (1997).

FeCCCCH⁻

Threshold for electron detachment from ground state FeCCCCH⁻ = 13470(500) gas PE¹

Reference

¹J. Fan, L. Lou, and L.-S. Wang, J. Chem. Phys. **102**, 2701 (1995).

C₅H \tilde{X} ²Π A_{eff}=23.68; B₀=0.080 MW¹**C₅D** \tilde{X} ²Π A_{eff}=24.2; B₀=0.075 MW²**References**

¹C. A. Gottlieb, E. W. Gottlieb, and P. Thaddeus, Astron. Astrophys. **164**, L5 (1986).

²T. Hirota, II. Ozawa, Y. Sckimoto, and S. Yamamoto, J. Mol. Spectrosc. **174**, 196 (1995).

HCCCCO \tilde{X} C_s B_{eff}=0.076 MW¹

DCCCCO
 \tilde{X} C_s
 $B_{\text{eff}}=0.072$ MW¹
Reference
¹H. Kohguchi, Y. Ohshima, and Y. Endo, J. Chem. Phys. **101**, 6463 (1994).
HCCCCS
 \tilde{X} 2II C_{∞v}
 $B_0=0.048$ MW¹
DCCCCS
 \tilde{X} 2II C_{∞v}
 $B_0=0.046$ MW¹
Reference
¹Y. Hirahara, Y. Ohshima, and Y. Endo, J. Chem. Phys. **101**, 7342 (1994).
O=C=C=NOH
 \tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OH stretch	3583.2w	Ar	IR	1
	2	CCO stretch	2074.1vs	Ar	IR	1
	4	HON bend	1357.7wm	Ar	IR	1
	5	CNO stretch	889.1wT	Ar	IR	1
	6	CNO stretch	882.8m	Ar	IR	1
	10	HON deform.	512.5w	Ar	IR	1
<i>a''</i>	11	HON deform.	500.9vwT	Ar	IR	1

Reference
¹G. Maier, H. P. Reisenauer, B. Röther, and J. Eckwert, Liebigs Ann. **1996**, 303.
HOONO₂

In the gas phase, the threshold for unstructured absorption lies near 330 nm. This absorption increases steadily to a maximum at or beyond 190 nm.^{4-7,10,12}

 \tilde{X} C_s Structure: MW⁸

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OH stretch	3540.1wm	gas	IR	1,4,12
	2	NO ₂ a-stretch	1728.3vs	gas	IR	1,3,7,12
	3	OH bend	1396.9wm	gas	IR	1,3,4,7,12,14
	4	NO ₂ s-stretch	1304.2s	gas	IR	1-4,7,12
	5	O-O stretch	941.0w	gas	IR	1,3,11,12
			922.1w	gas	IR	1,3
			919.2w	gas	IR	1,3,11
	6	NO ₂ scissors	802.54m	gas	IR,DL	1-4,7,9,12,13
	7	N-O stretch	648	gas	IR	13
	8	NO ₂ rock	466	gas	IR	13
	10	NO ₂ wag	722	gas	IR	11,13
	12	NO ₂ torsion	145(6)	gas	MW	8

 $A_0=0.400$; $B_0=0.156$; $C_0=0.113$ gas MW⁸
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NNBeBeNN
 \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		NN stretch	1804.1	Ar	IR	1
			1890.0	N ₂	IR	1
			820.4	N ₂	IR	1

Reference
¹C. A. Thompson, L. Andrews, and R. D. Davy, J. Phys. Chem. **99**, 7913 (1995).
C₆

In an argon matrix, the growth behavior of the 1952.5 cm⁻¹ infrared absorption on sample warmup has been tentatively correlated with that of an absorption maximum near 246 nm.⁴

 $\tilde{A}^3\Sigma_u^-$
 $T_0=19558(5)$ Ne AB¹⁰
 19220 Ar AB¹²
 $\tilde{A}-\tilde{X}$ 430-512 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2096(5)	Ne	AB	10
	2		1566(5)	Ne	AB	10
	3		640(5)	Ne	AB	10

$\bar{X}^3\Sigma_g^-$ D_{∞h} Structure: ESR²DL⁸

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	2061(10)	gas	TPE	6
			2050T	Ar	IR	11
	2	Sym. stretch	1322(10)	gas	TPE	5,6
	3	Stretch	489(10)	gas	TPE	6
	4	Asym. stretch	1959.86	gas	DL	8
			1958.7	Ne	IR	9,10
			1952.0	Ar	IR	3,4,7
			1951.2	Kr	IR	11
			1199.4	Ne	IR	9
Π_g	5	Asym. stretch	1197.3	Ar	IR	1,3,7
			1197.0	Kr	IR	11
			240HT	gas	PE	5

 $B_0=0.048 \text{ DL}^8$

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SiC₄Si

\bar{X}	D _{∞h}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	4	C=C stretch	1807.4	Ar	IR	1
	5	Si-C stretch	719.1	Ar	IR	1

Reference

- J. D. Presilla-Márquez, C. M. L. Rittby, and W. R. M. Graham, *J. Chem. Phys.* **106**, 8367 (1997).

Si₆

\bar{X}	D _{4h}					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_{1g}			458	N ₂	Ra	1
			300	N ₂	Ra	1
b_{1g}			386	N ₂	Ra	1
b_{2g}			252	N ₂	Ra	1
e_g			404	N ₂	Ra	1
e_u			462.9	Ne	IR	2
			460.9	Ar	IR	2
			458.5	Kr	IR	2

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C₆⁻

Threshold for electron detachment from ground-state C₆⁻=33725(10) gas PE^{1,2}TPE³

 \bar{F} D_{∞h}
 $T_0=33680\text{T}$ gas PE³

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2058(5)	gas	PE	3
	2		1307(5)	gas	PE	3
	3		480(2)	gas	PE	3
Σ_u^+	5		837HT	gas	PE	3
Π_g	7		195HT	gas	PE	3
Π_u	8		313HT	gas	PE	3
	9		93HT	gas	PE	3

 $(3)^2\Pi_g$ D_{∞h}
 $T_0=22517(10)$ Ne AB⁸

401–445 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	2		1861(17)	Ne	AB	8
	3		626(15)	Ne	AB	8
	9		157(7)H	Ne	AB	8

 $(2)^2\Pi_g$ D_{∞h}
 $T_0=20064(8)$ Ne AB⁸

470–499 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	3		597(12)	Ne	AB	8
	7		237(6)H	Ne	AB	8
	9		152(6)H	Ne	AB	8

\tilde{C}	$^2\Pi_g$	D _{∞h}
$T_0=16476$	gas	PD ⁵
16458(5)	Ne	AB ⁴
16239	Ar	AB ⁶

$\tilde{C}-\tilde{X}$ 445–629 nm
 $\tilde{C}-\tilde{X}$ 539–608 nm
 $\tilde{C}-\tilde{X}$ 573–616 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	C≡C stretch	2052	gas	PD	5
			2064(5)	Ne	AB	4
	2	C≡C stretch	1767	gas	PD	5
			1817(5)	Ne	AB	4
	3	C-C stretch	602	gas	PD	5
			607(5)	Ne	AB	4
			605T	Ar	AB	6
Π_g	8		245H	gas	PD	5
Π_u	9		110H	gas	PD	5

\tilde{A}	$^2\Sigma_g^+$	D _{∞h}
$T_0=9352(2)$	Ne	AB ⁸

$\tilde{A}-\tilde{X}$ 737–1070 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2111(4)	Ne	AB	8
	2		1912(4)	Ne	AB	8
	3		651(3)	Ne	AB	8
Π_g	7		238(2)H	Ne	AB	8
	9		128(2)H	Ne	AB	8

\tilde{X}	$^2\Pi_u$	D _{∞h}
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Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	3	C-C stretch	564	gas	PD	5
Σ_u^+	4	Asym. stretch	1938.5	Ne	IR	7
			1936.7	Ar	IR	6
Π_g	7		220T	gas	PE	3
	8		201H	gas	PD	5
Π_u	9		111T	gas	PE	3

$A=-29(2)$ gas PE³

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C₅N

\tilde{X}	$^2\Sigma$	C _{∞v}
$B_0=0.0468$	MW ¹	

Reference

- Y. Kasai, Y. Sumiyoshi, Y. Endo, and K. Kawaguchi, Astrophys. J. **477**, L65 (1997).

NCC≡CCN⁺

\tilde{D}	$^2\Pi_u$	D _{∞h}
$T_0=25500(160)$	gas	PE ^{1,3}

\tilde{C}	$^2\Sigma_u^+$	D _{∞h}
$T_0=18720(160)$	gas	PE ^{1,3}

\tilde{B}	$^2\Sigma_g^+$	D _{∞h}
$T_0=17430(160)$	gas	PE ^{1,3}

\tilde{A}	$^2\Pi_g$	D _{∞h}
$T_0=16781(1)$	gas	EF ² LF ³

16679 Ne LF⁴
 16709 Ne LF⁴AB⁵

$\tilde{A}-\tilde{X}$ 528–720 nm
 $\tilde{A}-\tilde{X}$ 530–816 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	C≡N stretch	2151(3)	gas	LF	3
			2154(2)	Ne	LF	4
	2	C≡C stretch	2099(3)	gas	LF	3
Σ_u^+			2094(2)	Ne	LF	4
	3	C-C stretch	696(3)	gas	LF	3
Π			511(2)	Ne	LF	4
	1	Bend	440H	Ne	AB	5

$\tau=13(2)$ ns gas EF²PEFCO³

\tilde{X} $^2\Pi_u$ D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	C≡N stretch	2210(10)	gas	EF	2,3
			2220	Ne	LF	4
	2	C≡C stretch	1930(10)	gas	EF	2,3
Σ_u^+			1942	Ne	LF	4
	3	C-C stretch	570(10)	gas	EF	2,3
Π	4		613	Ne	LF	4
	5		2010 ^a	Ne	LF	4
			1208 ^a	Ne	LF	4

^a From overtones and combination bands.

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(NUO)₂

\tilde{X}	D _{2h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> _{2u}			823.3	Ar	IR	1
			674.1	Ar	IR	1
<i>b</i> _{3u}		UN stretch	479.5	Ar	IR	1

Reference

¹G. P. Kushto, P. F. Souter, L. Andrews, and M. Neurock, J. Chem. Phys. **106**, 5894 (1997).

C₅O

\tilde{X}	1Σ ⁺	C _{∞v}	Structure: MW ¹
	$B_0=0.046$	MW ¹	

Reference

¹T. Ogata, Y. Ohshima, and Y. Endo, J. Am. Chem. Soc. **117**, 3593 (1995).

NCCCCN

\tilde{X}	C _{∞v}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ ⁺	1	C≡C stretch	2287.1s	Ar	IR	1,2
	2	-C≡N stretch	2203.6s	Ar	IR	1,2
	3	-N≡C stretch	2044.8vs	Ar	IR	1,2
	4	C-C,C-N a-stretch	1202.3vw	Ar	IR	1,2
	5	C-C,C-N s-stretch	610.1vwT	Ar	IR	1,2

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¹A. M. Smith, G. Schallmoser, A. Thoma, and V. E. Bondybey, J. Chem. Phys. **98**, 1776 (1993).

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Fe₂O₄

\tilde{A}	D _{∞h}					
T ^a	=1450(160)	gas	PE ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			720(60)	gas	PE	1
\tilde{X}	D _{∞h}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			710(60)	gas	PE	1

Reference

¹H. Wu, S. R. Desai, and L.-S. Wang, J. Am. Chem. Soc. **118**, 5296 (1996).

Cu₂O₄

\tilde{C}	D _{∞h}					
T ^a	=3630(400)	gas	PE ¹			

\tilde{B}	D _{∞h}					
T ^a	=2420(400)	gas	PE ¹			

\tilde{A}	D _{∞h}					
T ^a	=1290(400)	gas	PE ¹			

Reference

¹L.-S. Wang, H. Wu, S. R. Desai, and L. Lou, Phys. Rev. B **53**, 8028 (1996).

Fe₂O₄⁻

Threshold for electron detachment from ground-state Fe₂O₄⁻=28720(320) gas PE¹

Reference

¹H. Wu, S. R. Desai, and L.-S. Wang, J. Am. Chem. Soc. **118**, 5296 (1996).

Cu₂O₄⁻

Threshold for electron detachment from ground-state Cu₂O₄⁻=28240(240) gas PE¹

Reference

¹L.-S. Wang, H. Wu, S. R. Desai, and L. Lou, Phys. Rev. B **53**, 8028 (1996).

ONCCNO⁺

\tilde{C}	2Π _u	D _{∞h}				
T ^a	=54200(200)	gas	PE ¹			

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ _g ⁺			1613(40)	gas	PE	1

\tilde{B}	2Π _g	D _{∞h}				
T ^a	=47200(1000)	gas	PE ¹			

\tilde{A}	2Π _u	D _{∞h}				
T ^a	=16700(200)	gas	PE ¹			

\tilde{X}	2Π _g	D _{∞h}				

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ _g ⁺			2097(40)	gas	PE	1

^aFrom vertical ionization potentials.

Reference

¹T. Pasinszki and N. P. C. Westwood, J. Am. Chem. Soc. **117**, 8425 (1995).

Al₂O₄

\tilde{A}
 $T_0=9040(640)$ gas PE¹

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Sym. stretch	930(60)	gas	PE	1

Reference

¹S. R. Desai, H. Wu, C. M. Rohlfsing, and L.-S. Wang, J. Chem. Phys. **106**, 1309 (1997).

ONCCNO

\tilde{X} D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u	4		2246.04	gas	IR	1,2
	5		1258.48	gas	IR	1,2

$B_0=0.042$ IR²

References

¹T. Pasinszki and N. P. C. Westwood, J. Am. Chem. Soc. **117**, 8425 (1995).

²B. Guo, T. Pasinszki, N. P. C. Westwood, and P. F. Bernath, J. Chem. Phys. **103**, 3335 (1995).

Al₂O₄⁻

Threshold for electron detachment from ground-state Al₂O₄⁻=32110(240)
gas PE¹

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Sym. stretch	1090(60)	gas	PE	1

Reference

¹S. R. Desai, H. Wu, C. M. Rohlfsing, and L.-S. Wang, J. Chem. Phys. **106**, 1309 (1997).

Si₂O₄

\tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b_{1u}	7	Si=O stretch	1293.3	Ar	IR	1
	8	Si-O stretch	786.4	Ar	IR	1
b_{2u}	9		442w	Ar	IR	1
b_{3u}	11	Si-O stretch	889.2	Ar	IR	1
			285w	Ar	IR	1

Reference

¹T. Mehner, H. J. Göcke, S. Schunck, and H. Schnöckel, Z. Anorg. Allg. Chem. **580**, 121 (1990).

F₂BNCO

\tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	1	NCO a-stretch	2333vs	gas	IR	1
	2	Mixed	1569Ts	gas	IR	1
			1553Ts			
	3	BF ₂ a-stretch	1413s	gas	IR	1
a''	4	BF ₂ s-stretch	1213s	gas	IR	1
	10		659	gas	IR	1

Reference

¹M. Alaei, E. G. Livingstone, and N. P. C. Westwood, J. Am. Chem. Soc. **115**, 2871 (1993).

Si₂O₄⁻

Threshold for electron detachment from ground-state Si₂O₄⁻=20980(800)
gas PE¹

Reference

¹L.-S. Wang, H. Wu, S. R. Desai, J. Fan, and S. D. Colson, J. Phys. Chem. **100**, 8697 (1996).

N₂O₄

In the gas phase, absorption increases between 300 and 180 nm, with some indication of maxima near 265 and 190 nm.⁷

A broad gas-phase absorption with onset near 380 nm has its maximum near 340 nm.⁷

\bar{X} D_{2h} Structure: ED¹IR^{10,11}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_g	1		1383(3)	Ne	Ra	8
			1383	Ar	Ra	5
			1387(3)	Xe	Ra	8
	2		807(3)	Ne	Ra	8
			813	Ar	Ra	5
			815(3)	Xe	Ra	8
	3		265(3)	Ne	Ra	8
			262	Ar	Ra	5
			257(3)	Xe	Ra	8
	4		82T	gas	IR ^a	6,12
b_{1g}	5		1718(3)	Xe	Ra	8
	6		480T	gas	IR ^a	6,12
b_{1u}	7		498(3)	Ne	Ra	8
	8		485(3)	Xe	Ra	8
b_{2g}	9		425	gas	IR	6
			657(3)	Xe	Ra	8
b_{3u}	9		1756.76	gas	IR,DL	6,10,13,14
			1749.2s	Ar	IR	2,3,9
			1735s			
			1761	N ₂	IR	4
			1737			
			1750	O ₂	IR	3
			1735			
	10		265T	gas	IR	11
	11		1261.08	gas	IR,DL	2,6,11,13,14
			1257.0s	Ar	IR	2,3,9
			1261	N ₂	IR	4
			1261	O ₂	IR	3
12			755.37 ^b	gas	IR	6,13
			747.85 ^b	gas	IR	6,13
			755sh	Ar	IR	3
			745.8			
			751	N ₂	IR	4
			755	O ₂	IR	3
			746			

$A_0=0.218$; $B_0=0.122$; $C_0=0.078$ IR^{10,11}DL¹⁴
 Barrier to internal rotation = 1900(200) gas IR¹²

^aFrom analysis of sequence bands near 540 cm^{-1} .

^bFermi resonance between ν_{12} and $\nu_6 + \nu_{10}$.

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CF₃CO

\tilde{A}
 $T_0=26023$ gas LF² $\tilde{A}-\bar{X}$ 353–455 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			781(30)	gas	LF	2
		CCO bend	357(30)	gas	LF	2
			$\tau=0.34 \mu\text{s}$	gas	LF ²	

 \bar{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CO stretch	1550(30)	gas	IR,LF	1,2
		CF ₃ stretch	1338	gas	IR	1
		CF ₃ stretch	1115	gas	IR	1
		CC stretch	1024	gas	IR	1
		CCO bend	633(30)	gas	LF	2

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ClOClO₃

An unstructured gas-phase absorption with maximum at 42740 (234 nm) has been assigned⁴ to ClOClO₃.

 \bar{X} C_s

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	1	ClO ₂ s-stretch	1283vs	gas	IR	1–3,5
			1293.2	Ne	IR	6
			1287vs	Ar	IR	2
	2	Cl=O stretch	1040s	gas	IR	1,2
			1040.2	Ne	IR	6
			1039s	Ar	IR	2
	3	O-Cl stretch	749wm	gas	IR	1,2
			747.5	Ne	IR	6
			746m	Ar	IR	2
	4	Cl-O stretch	646vs	gas	IR	1,2
a''			652.1	Ne	IR	6
			647vs	Ar	IR	2
	5	O=Cl=O bend	580sh	gas	IR	2
			586.9	Ne	IR	6
			582m	Ar	IR	2
	6	ClO ₃ deform.	511wm	gas	IR	2
			514.6	Ne	IR	6
			513m	Ar	IR	2
	7	O=Cl-O bend	355vw	Ar	IR	2
	9	ClO ₂ a-stretch	1283vs	gas	IR	1–3,5
			1277.6	Ne	IR	6
			1271vs	Ar	IR	2
	10	O=Cl=O bend	561m	gas	IR	2
			565.3	Ne	IR	6
			561ms	Ar	IR	2
	11	O=Cl-O bend	382w	Ar	IR	2

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- ⁶H. S. P. Müller and H. Willner, J. Phys. Chem. **97**, 10589 (1993).

8.13. Seven-Atomic Molecules

C₂H₅**3p Rydberg state**

In the gas phase, an absorption with maximum at 205 nm has been assigned^{6,7} to the 3p- \tilde{X} transition of C₂H₅. The detection of this band in MPI studies⁹ is consistent with the assignment to a Rydberg state.

3s Rydberg state

In the gas phase, a broad, unstructured absorption with maximum at 246 nm has been assigned^{1,3,6,7} to the 3s- \tilde{X} transition of C₂H₅.

\tilde{X}	C _s	Structure: MO ⁴				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	CH ₂	s-stretch	3033m	Ar	IR	2,4,5,8
			3032.6	H ₂	IR	11
	CH ₃	s-stretch	2920m	Ar	IR	4,5
	2-CH	stretch	2842s	Ar	IR	2,4,5,8
	CH ₂	deform.	1440m	Ar	IR	2,4,5,8
			1383	Ar	IR	8
	CH ₃	deform.	1366m	Ar	IR	2,4,5,8
	CC	stretch	1138w	Ar	IR	4,5,8
			1025	Ar	IR	8
	CCH ₂	umbrella	528.1	gas	DL	10
<i>a''</i>			540vs	Ar	IR	2,4,5,8
	CH ₂	a-stretch	3112s	Ar	IR	2,4,5,8
			3122.8	H ₂	IR	11
	CH ₃	a-stretch	2987s	Ar	IR	2,4,5
			2984.3	H ₂	IR	11
	CH ₃	deform.	1440m	Ar	IR	2,4,5,8
	H	deform.	1175m	Ar	IR	4,5,8

C₂D₅

\tilde{X}	C _s	Structure: PE ^{2,3}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	CD ₂	s-stretch	2199m	Ar	IR	4,5
	CD ₃	s-stretch	2094m	Ar	IR	4,5
	2-CD	stretch	2048m	Ar	IR	4,5
	CD ₃	deform.	1070m	Ar	IR	4,5
	CD ₃	deform.	1035m	Ar	IR	4,5
	CCD ₂	umbrella	398vs	Ar	IR	4,5
	CD ₂	a-stretch	2249m	Ar	IR	4,5
	CD ₃	a-stretch	2170s	Ar	IR	4,5
	CD ₃	deform.	1041m	Ar	IR	4,5
<i>a''</i>						

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- ¹¹N. Sogoshi, T. Wakabayashi, T. Momose, and T. Shida, J. Phys. Chem. A **101**, 522 (1997).

H₅O⁺

\tilde{X}	C ₂					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		H ₂ O a-stretch	3684.4	gas	PF	1,2
		H ₂ O s-stretch	3608.8	gas	PF	1

References

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H₂CCCH₂⁺

\tilde{B}
 $T^a=61400(800)$ gas PE¹

\tilde{A}^2E
 $T_0=35638(32)$ gas PE¹⁻⁴

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	2	CH ₂ scissors	1320(6)	gas	PE	3,4
<i>b</i> ₂	7	CH ₂ deform.	1030(6)	gas	PE	3,4

 \tilde{X}^2E
D₂
Structure: PE^{2,3}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> ₁	4	Torsion	745(5)	gas	PE	3,4
<i>b</i> ₂	6	C ₃ a-stretch	1802(10)	gas	PE	4

D₂CCCD₂⁺

\tilde{X}^2E
D₂

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> ₁	4	Torsion	565(50)	gas	PE	1

^aFrom vertical ionization potential.

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³Z. Z. Yang, L. S. Wang, Y. T. Lee, D. A. Shirley, S. Y. Huang, and W. A. Lester, Jr., Chem. Phys. Lett. **171**, 9 (1990).
⁴P. Baltzer, B. Wannberg, M. Lundqvist, L. Karlsson, D. M. P. Holland, M. A. MacDonald, and W. von Niessen, Chem. Phys. **196**, 551 (1995).

cyc-C₂SiH₄

\bar{X}	C_{2v}					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CH stretch	3061.2w	Ar	IR	1
	2	SiH stretch	2166.7vs	Ar	IR	1
	3	CC stretch	1467.2m	Ar	IR	1
	4	SiH ₂ deform.	975.9vs	Ar	IR	1
	5	CH deform.	900.6w	Ar	IR	1
	6	SiC stretch	778.5wm	Ar	IR	1
b_1	9	SiH stretch	2176.3s	Ar	IR	1
	10	Deformation	695.1ms	Ar	IR	1
	11	Deformation	589.2wm	Ar	IR	1
b_2	12	CH stretch	3041.6wm	Ar	IR	1
	13	CH deform.	1105.1m	Ar	IR	1
	14	SiC stretch	707.4m	Ar	IR	1
	15	SiH ₂ deform.	655.4m	Ar	IR	1

Reference

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CH₂=CHOH⁺

\tilde{D}^2A'	C_s					
$T^a=82300(400)$	gas	PE ^{1,2}				
\tilde{C}^2A'	C_s					
$T^a=61200(400)$	gas	PE ^{1,2}				
\tilde{B}^2A''	C_s					
$T^a=41300(400)$	gas	PE ^{1,2}				
\tilde{A}^2A'	C_s					
$T^a=35400(400)$	gas	PE ^{1,2}				
\tilde{X}^2A''	C_s					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'			1400(50)	gas	PE,PI	2,3

^aValues for syn- rotamer. From vertical ionization potential.

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CH₃CHS

\tilde{F}^1A''	C_s					
$T_0=51560(200)$	gas	AB ²				
\tilde{D} or \tilde{E}^1A'	C_s					
$T_0=52770(100)$	gas	AB ²				
\tilde{C}^1A'	C_s					
$T_0=45260(100)$	gas	AB ²				
\tilde{B}^1A'	C_s					
$T^a=47600(100)$	gas	AB ²				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'		C=S stretch	1154	gas	AB	2
\tilde{A}^1A''	C_s					
$T_0=17900(100)$	gas	AB ²				
$\tau_0=0.40(5) \mu\text{s}$	gas	LF ⁴				
\tilde{a}^3A''	C_s					
$T_0=16294.9$	gas	AB ^{2,3} LF ⁷				
						$\tilde{a}-\tilde{X} 571-630 \text{ nm}$
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'	9	CS stretch	747.2	gas	AB,LF	3,7
	10	CCS deform.	283.9	gas	LF	7
a''	14	Wag	249.4	gas	LF	7
	15	Torsion	52.0	gas	LF	7
						$\tau \sim 8.2 \mu\text{s}$ gas LF ⁴
\tilde{X}^1A'	C_s	Structure: MW ¹				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CH stretch	3010.4w	Ar	IR	6,8
		CH stretch	2982.0w	Ar	IR	6,8
		CH stretch	2944.4m	Ar	IR	6,8
		CII stretch	2908.7w	Ar	IR	6
			1435.3m	Ar	IR	6,8
			1433.4wm	Ar	IR	6
			1357.7s	Ar	IR	6,8
			1345.6vs	Ar	IR	6,8
		C=S stretch	1240	gas	AB	3
			1140.7s	Ar	IR	6,8
			1065.0wm	Ar	IR	6,8
			1020.2wm	Ar	IR	6,8
			820.1w	Ar	IR	6,8
			747.8s	Ar	IR	6,8
	15	Torsion	163.0	gas	LF	7
						$A_0=1.620; B_0=0.192; C_0=0.177 \text{ MW}^1$

CD₃CDS

$\tilde{\alpha}^3A''$ C_s
 $T_0=16367.2$ gas AB³LF⁷

$\tilde{\alpha}-\tilde{X}$ 580–630 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	8	DCS deform.	751.8	gas	LF	7
	9	CS stretch	677.2	gas	AB,LF	3,7
a''	15	Torsion	29.6	gas	LF	7

\tilde{X}^1A'	C _s					
Vib. sym.	No.					
a''	15	Torsion	122.6	gas	LF	7

^aMaximum of diffuse absorption with partially resolved structure.

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- E. Suzuki, O. Watanabe, A. Happoya, and E. Watari, Vib. Spectrosc. **5**, 353 (1993).

FCH₂CH₂

\tilde{X}	C _s	Structure: MO ¹				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH ₂ stretch	2885wm	Ar	IR	3
		CH ₂ stretch	2856m	Ar	IR	2,3
		Deformation	1449w	Ar	IR	3
		Deformation	1426wm	Ar	IR	3
		CH ₂ wag	1242vw	Ar	IR	3
		Mixed	1087s	Ar	IR	3
		CF stretch	1047s ^a	Ar	IR	2
		CCF deform.	427wm	Ar	IR	2

FCD₂CD₂

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		2-CD ₂ stretch	2090wm	Ar	IR	2
		CF stretch + CD ₂ scissors	1211s	Ar	IR	2
			1091m	Ar	IR	2
			1059wm	Ar	IR	2
		CF stretch + CD ₂ scissors	969s	Ar	IR	2
		CCF deform.	424w	Ar	IR	2

^aRef. 3 proposes the assignment of this absorption to t-CH₂FCH₂F. Although that product is known to absorb near 1047 cm⁻¹, such an assignment is inconsistent with the infrared absorption pattern² obtained in a study of the reaction of F atoms with C₂H₄-¹³C, as well as with the low yield of the more stable *gauche*-CH₂FCH₂F in the earlier work.²

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CH₂(OH)₂

\tilde{X}						
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OH stretch	3637.6m ^a T	Ar	IR	1
		OH stretch	3564.4m ^b T	Ar	IR	1
		COH deform.	1424.4wmT	Ar	IR	1
		COH deform.	1358.7wT	Ar	IR	1
		CH ₂ wag	1334.6wT	Ar	IR	1
		CO stretch	1056.5sT	Ar	IR	1
		CO torsion	547.7mT	Ar	IR	1

^aRotamer 1.

^bRotamer 2.

Reference

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C₂H₃CN⁺

\tilde{E}^2A' C_s
 $T^a=42430(20)$ gas PE¹

\tilde{D}^2A' C_s
 $T^a=27980(20)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		H deformation	1494T	gas	PE	1

$\tilde{C}^{\prime\prime} 2A''$ C_s
 $T_0=20340(60)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		H deformation	1363(32)	gas	PE	1
			678(32)	gas	PE	1

$\tilde{B}^{\prime\prime} 2A'$ C_s
 $T_0=16890(20)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		H deformation	1137(24)	gas	PE	1
			660(24)	gas	PE	1

$\tilde{A}^{\prime\prime} 2A'$ C_s
 $T_0=11530(20)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C≡N stretch	1903(16)	gas	PE	1
		H deformation	1194(24)	gas	PE	1
		C ₃ deformation	492(32)	gas	PE	1

$\tilde{X}^{\prime\prime} 2A''$ C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C≡N stretch	2121(16)	gas	PE	1
		C=C stretch	1335(24)	gas	PE	1
			928(30)	gas	PE	1
			839(30)	gas	PE	1
			347(30)	gas	PE	1

^aFrom vertical ionization potential.

Reference

¹J. Delwiche, M. Gochel-Dupuis, J. E. Collin, and J. Heinesch, J. Electron Spectrosc. Relat. Phenom. **66**, 65 (1993).

H₂C=(cyc-CN=CH)

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1818	Kr	IR	1

Reference

¹K. Banert, M. Hagedorn, E. Knözinger, A. Becker, and E.-U. Würthwein, J. Am. Chem. Soc. **116**, 60 (1994).

CH₃OCN

\tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	CH ₃ a-stretch	3042wT	gas	IR	1
	2	CH ₃ s-stretch	2968	gas	IR	1
	3	OCN a-stretch	2263vs	gas	IR	1
	6	CH ₃ rock	1213	gas	IR	1
	7	Skel. stretch	1112	gas	IR	1
	8	Skel. stretch	883T	gas	IR	1
	11	CH ₃ a-stretch	3017	gas	IR	1
	12	CH ₃ a-deform.	1462	gas	IR	1

Reference

¹T. Pasinszki and N. P. C. Westwood, J. Phys. Chem. **99**, 1649 (1995).

B(OH)₃

\tilde{X} C_{3h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	OH stretch	3705	gas	Ra	2
	2	BOH deform.	1020	gas	Ra	2
	3	BO stretch	866	gas	Ra	2
	4	BO ₃ OPLA	666.4	Ar	IR	3
<i>e'</i>			675.0	N ₂	IR	1
	5	Torsion	436.0	Ar	IR	3
			513.8	N ₂	IR	1
	6	OH stretch	3706	gas	IR	2
<i>e''</i>			3688.6	Ar	IR	3
			3668.5	N ₂	IR	1
	7	BO stretch	1429	gas	IR	2
			1414.9	Ar	IR	3
<i>8</i>			1426.2	N ₂	IR	1
			1017	gas	IR	2
			992.4	Ar	IR	3
			1009.9	N ₂	IR	1
<i>9</i>			432.1	Ar	IR	3
			448.9	N ₂	IR	1
			520(5) ^a	Ar	IR	3

B(OD)₃

\tilde{X}		C _{3h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	2	BO stretch	887	gas	Ra	2
	3	BOD deform.	706	gas	Ra	2
a''	4	BO ₃ OPLA	665.5	Ar	IR	3
	5	Torsion	317.6	Ar	IR	3
			376.0	N ₂	IR	1
e'	6	OD stretch	2726	gas	IR	2
			2719.8	Ar	IR	3
			2704.6	N ₂	IR	1
	7	BO stretch	1405	gas	IR	2
			1396.6	Ar	IR	3
	8	DOB deform.	810	gas	IR	2
			807.9	Ar	IR	3
			825.5	N ₂	IR	1
	9	BO ₂ deform.	385.9	Ar	IR	3
			404.7	N ₂	IR	1
e''	10	DOB deform.	390(10) ^a	Ar	IR	3

^aFrom combination bands.**References**

- ¹J. S. Ogden and N. A. Young, J. Chem. Soc. Dalton Trans. 1645 (1988).
²T. R. Gilson, J. Chem. Soc. Dalton Trans. 2463 (1991).
³L. Andrews and T. R. Burkholder, J. Chem. Phys. **97**, 7203 (1992).

CH₃CF₂

3p Rydberg state		C _s				
T ₀ =43275(40)U	gas	MPI ¹				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	9	CF ₂ wag	530	gas	MPI	1

Reference

- ¹J. L. Brum, R. D. Johnson III, and J. W. Hudgens, J. Phys. Chem. **98**, 3645 (1994).

C₅H₂⁺

\tilde{A}	² Π_g	D _{∞h}
T ₀ =20026(5)T	Ne	AB ¹

Reference

- ¹P. Freivogel, J. Fulara, D. Lessen, D. Forney, and J. P. Maier, Chem. Phys. **189**, 335 (1994).

HC₅H

³ Σ_u^-		D _{∞h}				
T ₀ =23033(5)	Ne	AB ¹				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	2	C≡C s-stretch	1817(5)	Ne	AB	1
	3		751(5)	Ne	AB	1

$$\tilde{X} \quad \text{D}_{\infty h}$$

$$A_0=9.26; B_0=0.077; C_0=0.076 \quad \text{MW}^2$$

References

- ¹J. Fulara, P. Freivogel, D. Forney, and J. P. Maier, J. Chem. Phys. **103**, 8805 (1995).
²M. C. McCarthy, M. J. Travers, A. Kovács, W. Chen, S. E. Novick, C. A. Gottlieb, and P. Thaddeus, Science **275**, 518 (1997).

t-HCCCHO

Irradiation at wavelengths longer than 590 nm leads to photodestruction and, in an argon matrix, to stabilization of HCC(cyc-CHOO).¹

 \tilde{X} **C_s**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'		CH stretch	3313.5s	Ar	IR	1
		C=C stretch	2093.8s	Ar	IR	1
		HCO bend	1401.5m	Ar	IR	1
		CO stretch	1221.2m	Ar	IR	1
		Mixed	1026.4s	Ar	IR	1
		OO stretch	933.3s	Ar	IR	1
		CCO bend	739.0m	Ar	IR	1
		HCC bend	655.6m	Ar	IR	1
a''		HCC bend	646.5m	Ar	IR	1
		Skel. OPLA	497.1w	Ar	IR	1

Reference

- ¹S. Wierlacher, W. Sander, C. Marquardt, E. Kraka, and D. Cremer, Chem. Phys. Lett. **222**, 319 (1994).

c-HCCCHO^a **\tilde{X}** **C_s**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1412	Ar	IR	1
		CO stretch	1234	Ar	IR	1
		OO stretch	983	Ar	IR	1
			914	Ar	IR	1

^aTentative identification.**Reference**

- ¹S. Wierlacher, W. Sander, C. Marquardt, E. Kraka, and D. Cremer, Chem. Phys. Lett. **222**, 319 (1994).

HCC(cyc-CHOO)^a

Photodecomposes into CO₂, C₂H₂, and propionic acid on irradiation near 25000 (403(20) nm).¹



Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH stretch	3319	Ar	IR	1
			1397	Ar	IR	1
			1375	Ar	IR	1
			1247	Ar	IR	1
			950	Ar	IR	1
			873	Ar	IR	1
			676	Ar	IR	1

^aTentative identification.

Reference

¹S. Wierlacher, W. Sander, C. Marquardt, E. Kraka, and D. Cremer, Chem. Phys. Lett. **222**, 319 (1994).

C₆H

$\tilde{A}^2\Pi$		$C_{\infty v}$		$\tilde{A}-\tilde{X}$ 484–528 nm		
$T_0=18992.4$ gas CR ⁸				$\tilde{A}-\tilde{X}$ 444–530 nm		
18854(5) Ne AB ⁷						
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	3	C≡C stretch	2076(5)	Ne	AB	7
	5	C≡C stretch	1630.8	gas	CR	8
			1638(5)	Ne	AB	7
	6	C-C stretch	659.5	gas	CR	8
			658(5)	Ne	AB	7

$A_0=-7.9$ gas CR⁸

$B_0=0.045$ CR⁸

 $\tilde{X}^2\Pi$ $C_{\infty v}$

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+		C≡C stretch	1962.2T	Ne	IR	7
			1953.4	Ar	IR	6

$A_{\text{eff}}=-15.1$ gas MW^{3–5}

$B_0=0.046$ MW^{1–5}

C₆D

$\tilde{A}^2\Pi$		$C_{\infty v}$		$\tilde{A}-\tilde{X}$ 524–526 nm		
$T_0=19043.9$ gas CR ⁸						
18900(10) Ne AB ⁸						
$A_0=-8.2$ gas CR ⁸						

 $\tilde{X}^2\Pi$ $C_{\infty v}$

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+		C≡C stretch	1862.4	Ar	IR	6

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H(C≡C)₂CN⁺

$\tilde{C}^2\Pi$ $C_{\infty v}$
 $T_0=27350(160)$ gas PE¹

$\tilde{B}^2\Sigma^+$ $C_{\infty v}$
 $T_0=21860(160)$ gas PE¹

$\tilde{A}^2\Pi$ $C_{\infty v}$
 $T_0=17190(5)$ gas EF¹
 $17098(20)$ Ne AB²LF³ $\tilde{A}-\tilde{X}$ 580–670 nm
 $\tilde{A}-\tilde{X}$ 494–820 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+				1870(160)	gas	EF 1
	3	C≡C stretch	2006(5)	Ne	AB	2
	4	CN stretch	1849(5)	Ne	AB	2
	5	C-C stretch	1187(5)	Ne	AB	2
	6	C-C stretch	600(160)	gas	EF	1
			584(5)	Ne	AB	2

$\tau_0=15(2)$ ns gas EF¹
 $A=17T$ Ne LF³

 $\tilde{X}^2\Pi$ $C_{\infty v}$

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	1	CH stretch	3403.4	Ne	LF	3
	2	C≡N stretch	2190(10)	gas	EF	1
			2194.6	Ne	LF	3
	3	C≡C stretch	2070(10)	gas	EF	1
			2070.2	Ne	LF	3
	4	C≡C stretch	1912.7	Ne	LF	3
	5	C-C stretch	1220(10)	gas	EF	1
			1213	Ne	LF	3
	6	C-C stretch	630(10)	gas	EF	1
			625	Ne	LF	3

$A=91T$ Ne LF³

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C₇

$^1\Sigma_u^+$	D _{∞h}
$T_0 = 39556(30)$ T	Ne AB ⁷
40470	Ar AB ⁴

 $^1\Sigma_u^+ - \tilde{X}$ 246–253 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			979(30)	Ne	AB	7
			428(30)	Ne	AB	7

$T_0 = 32765(22)$ T	Ne AB ⁷	246–306 nm
34360T	Ar AB ⁴	

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1256(25)	Ne	AB	7
			695(25)	Ne	AB	7

$^1\Pi_u$	D _{∞h}
$T_0 = 18440(7)$	Ne AB ⁷

 $^1\Pi_u - \tilde{X}$ 485–543 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			2170(10)	Ne	AB	7
			1873(10)	Ne	AB	7
			568(10)	Ne	AB	7
			478(10)	Ne	AB	7

\tilde{X}	D _{∞h}	Structure: DL ¹
Vib.	No.	Approximate type of mode
		cm ⁻¹
		Med.
		Type meas.
		Refs.
Σ_g^+	3	Sym. stretch
		548(90) gas
		582T Ar
Σ_u^+	4	Asym. stretch
		2138.315 gas
		2134.6 Ne
		2127.8 Ar
		2120.4 Kr
	5	Asym. stretch
		1898.376 gas
		1897.5 Ne
		1894.3 Ar
		1889.3 Kr
Π_u	7	Bend
		496(110)T gas
		PE
		5

 $B_0 = 0.031$ DL¹⁻³**References**

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Si₇

\tilde{X}	D _{5h}
Vib.	No.
	Approximate type of mode
	cm ⁻¹
	Med.
	Type meas.
	Refs.
a'_1	
	435 N ₂
	358 N ₂
e'_1	
	422.4 Ar
	420.4 Kr
e'_2	
	340 N ₂
	289 N ₂

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C₇⁻

Threshold for electron detachment from ground-state C₇⁻=27090(115) gas PE^{1,2}

\tilde{C}^-	$^2\Pi_u$	D _{∞h}
$T_0 = 35231(25)$	Ne AB ⁶	$\tilde{C}^- - \tilde{X}$ 278–284 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	3		491(25)	Ne	AB	6
\tilde{B}^-	$^2\Pi_u$	D _{∞h}				
$T_0 = 20330(10)$	gas	MPD ⁵				
20314(8)	Ne	AB ⁶				
						380–493 nm
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1990(40)	gas	MPD	5
	2		2058(10)	Ne	AB	6
	3		1569(9)	Ne	AB	6
			575(8)	Ne	AB	6

\tilde{A}^-	$^2\Pi_u$	D _{∞h}
$T_0 = 15950(10)$	gas	MPD ⁵
15954(5)	Ne	AB ⁶
		$\tilde{A}^- - \tilde{X}$ 495–627 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	Sym. stretch	1810(40)	gas	MPD	5
	2		1877(6)	Ne	AB	6
	3		1457(5)	Ne	AB	6
			500(40)	gas	MPD	5
			562(5)	Ne	AB	6

\tilde{X}	$^2\Pi_g$	D _{∞h}				
Vib.	No.	Approximate type of mode				
		cm ⁻¹				
		Med.				
		Type meas.				
Σ_u^+	5	Asym. stretch	1736.4	Ne	IR	4
			1734.8	Ar	IR	3

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C₆O $\tilde{X}^3\Sigma^-$ C_{∞v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+			2163.8	Ar	IR	1
			1447.8	Ar	IR	1

 $B_0=0.028$ MW²

References

- ¹G. Maier, H. P. Reisenauer, and A. Ulrich, Tetrahed. Lett. **32**, 4469 (1991).
²Y. Ohshima, Y. Endo, and T. Ogata, J. Chem. Phys. **102**, 1493 (1995).

C₅N₂

\tilde{A}		$\tilde{A}-\tilde{X}$ 375–610 nm		
$T_0=22762.9(2)T$	Ne	AB ¹ LF ¹		
22737.3(2)T	Ar	AB ¹ LF ¹		
22731.8(2)T	N ₂	AB ¹ LF ¹		

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1			1961.3(2)T	Ne	AB	1
			1960.5(2)T	Ar	AB	1
8			601.2(2)T	Ar	AB	1
9			508.5(2)T	Ar	AB	1

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
1	1		2070.7(2)T	Ne	LF	1
			2071.4(2)T	Ar	LF	1
			2071.6(2)T	N ₂	LF	1
	2		2030.7(2)T	Ne	LF	1
			2030.9(2)T	Ar	LF	1
			2032.0(2)T	N ₂	LF	1
			1639.2(2)T	Ne	LF	1
			1639.5(2)T	Ar	LF	1
			1639.7(2)T	N ₂	LF	1
	4		871.7(2)T	Ne	LF	1
	5		870.9(2)T	Ar	LF	1
	6		860.8(2)T	Ne	LF	1
			859.3(2)T	Ar	LF	1
	7		804.7(2)T	Ne	LF	1
			801.5(2)T	Ar	LF	1
	8		753.3(2)T	Ne	LF	1
			748.5(2)T	Ar	LF	1
	9		603.2(2)T ^a	Ne	LF	1
			609.6(2)T	Ar	LF	1
			601.9(2)T	Ar	LF	1
			527.7(2)T ^a	Ne	LF	1
			530.5(2)T	Ar	LF	1
			525.2(2)T	Ar	LF	1
			521.9(2)T	Ar	LF	1

^aCenter of split transition.

Reference

- ¹A. M. Smith, C. Engel, A. Thoma, G. Schallmoser, B. E. Wurfel, and V. E. Bondybey, Chem. Phys. **184**, 233 (1994).

(CN)₂CCO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CCO a-stretch	2183	Ar	IR	1
			2185s	CO	IR	1
			1202w	CO	IR	1
			953w	CO	IR	1
			566w	CO	IR	1

Reference

- ¹I. R. Dunkin and A. McCluskey, Spectrochim. Acta A **50**, 209 (1994).

(CN)₂COO

In an argon matrix, a weak, broad absorption at 26700 (375 nm) has been attributed¹ to (CN)₂COO.

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1290w	Ar	IR	1
			1275vw	Ar	IR	1
			1264vw	Ar	IR	1
			1245vw	Ar	IR	1

Reference

¹I. R. Dunkin and A. McCluskey, Spectrochim. Acta A **50**, 209 (1994).

Al₂O₅

\tilde{A}
 $T_0=10890(640)$ gas PE¹

 \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		Sym. stretch	900(60)	gas	PE	1

Reference

¹S. R. Desai, H. Wu, C. M. Rohlfing, and L.-S. Wang, J. Chem. Phys. **106**, 1309 (1997).

O₂N-O-NO₂

In the gas phase, continuous absorption begins near 420 nm, and has its maximum beyond 200 nm.¹¹

\tilde{X}	C ₂ ^a	Structure: ED ^b MW ^c				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a	1	NO ₂ a-stretch	1720vs	gas	IR	1,2,5,8
			1742.4vs	Ar	IR	12
			1745	N ₂	IR	3
			1752	CO ₂	IR	1
	2	NO ₂ s-stretch	1338m	gas	IR	2,8
			1339.8wm	Ar	IR	12
			1305	N ₂	IR	3
			1300	O ₂	IR	4
			1316	CO ₂	IR	1
			860m	gas	IR	2,8
	3		863.1w	Ar	IR	12
			1720vs	gas	IR	1,2,5,8
			1702.7s	Ar	IR	7,12
			1704	N ₂	IR	3
			1704	O ₂	IR	4
b	9	NO ₂ a-stretch	1700	CO ₂	IR	1
			1245.9s	gas	IR	1,2,5,8,11
			1243.0m	Ar	IR	7,12
			1247	N ₂	IR	3
			1241	O ₂	IR	4
	10	NO ₂ s-stretch	1248	CO ₂	IR	1
			743.4s	gas	IR	1,2,5,8,11
			737s	Ar	IR	7,12
			739	N ₂	IR	3
			736	O ₂	IR	4
			719	CO ₂	IR	1
			614mT	gas	IR	2
			639.7w	Ar	IR	12
			557s	gas	IR	2,5
			569.4m	Ar	IR	12
14	15		353vsT	gas	IR	2,9
			343.9vs	Ar	IR	12
			50w,brT	gas	IR	9

$A_0=0.221$; $B_0=0.063$; $C_0=0.060$ MW¹⁰

^aRef. 2 analyzed the spectrum in terms of a C_{2v} structure. However, the electron diffraction measurements of Ref. 6, analyzed using a dynamical model, and the low rotational temperature microwave spectral observations of Ref. 13 suggest instead a C₂ structure. The vibrational assignment has been revised to correspond with the results of *ab initio* calculations given in Ref. 13.

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SF₆⁺

\tilde{F} 2A_{1g}
 $T_0^a=94450(100)$ gas PE^{2,3,6}TPE⁷
Fragments into SF₂⁺, T-PEPICO⁵

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	1	Sym. stretch	690(40)	gas	PE	6,7
<i>e</i> _g	2		460T	gas	PE,TPE	6,7

\tilde{E} 2F_{1u}
 $T_0^a=57620(100)$ gas PE^{2,3,6}TPE⁷
Fragments into SF₃⁺, T-PEPICO⁵

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g	1	Sym. stretch	540(40)	gas	PE,TPE	6,7

\tilde{D} 2F_{2g}
 $T_0^a=33250(150)$ gas PE^{1-3,6}TPE⁷
Fragments into SF₅⁺, SF₄⁺, and SF₃⁺, T-PEPICO⁵

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g	1	Sym. stretch	590(40)	gas	PE,TPE	3,6,7

\tilde{C} 2E_g
 $T_0^a\leq 24360(100)$ gas PE^{1-3,6}TPE⁷
Low vibrational levels fragment into SF₅⁺+F, higher levels yield SF₄⁺, T-PEPICO⁵

$\tilde{A}, \tilde{B}^+ 2F_{1u}, 2F_{2u}$
 $T_0^a = 12780(240)$ gas PE^{1-3,6}TPE⁷
 Direct dissociation into SF₅⁺ + F. T-PEPICO^{4,5}

$\tilde{X}^+ 2F_{1g}$
 Direct dissociation into SF₅⁺ + F. T-PEPICO^{4,5}

^aMeasured with respect to photoionization threshold, 15.116 eV, determined by Ref. 7.

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8.14. Eight-Atomic Molecules

B₂H₆⁺

$\tilde{D}^+ 2B_{1u}$ D_{2h}
 $T_0^a = 38650(100)$ gas PE¹⁻⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g	1	BH _t stretch	2178(100) 1000(140)	gas gas	PE PE	2,4 2

$\tilde{C}^+ 2B_{2u}$ D_{2h}
 $T^a = 28000(100)$ gas PE²⁻⁴

$\tilde{B}^+ 2B_{3u}$ D_{2h}
 $T^a = 21300(180)$ gas PE²⁻⁴

$\tilde{A}^+ 2A_g$ D_{2h}
 $T^a = 11860(250)$ gas PE²⁻⁴

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g			1700(300)	gas	PE	2

$\tilde{X}^+ 2B_{2g}$ D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g			900(60)	gas	PE	2

B₂D₆⁺

$\tilde{D}^+ 2B_{1u}$ D_{2h}
 $T_0^a = 38650(100)$ gas PE²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g	1	BD _t stretch	1500(80) 650(160)	gas gas	PE PE	2 2

$\tilde{C}^+ 2B_{2u}$ D_{2h}
 $T^a = 28000(100)$ gas PE²

$\tilde{B}^+ 2B_{3u}$ D_{2h}
 $T^a = 21300(180)$ gas PE²

$\tilde{A}^+ 2A_g$ D_{2h}
 $T^a = 11860(250)$ gas PE²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g			1500(300)	gas	PE	2

$\tilde{X}^+ 2B_{2g}$ D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g			800(120)	gas	PE	2

^aFrom vertical ionization potential.

References

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- ⁴J. M. Dyke, D. Haggerston, O. Warschkow, L. Andrews, A. J. Downs, and P. F. Souter, *J. Phys. Chem.* **100**, 2998 (1996).

GaBH₆⁺

$\tilde{D}^+ 2A_1$ C_{2v}
 $T_0^a = 34130(120)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁		BH _t stretch	1950(100)	gas	PE	1
		GaH _t stretch	1480(100)	gas	PE	1

\tilde{C}^2B_2 C_{2v}
 $T^a=20250(240)$ gas PE¹

\tilde{B}^2B_1 C_{2v}
 $T^a=14760(120)$ gas PE¹

\tilde{A}^2A_1 C_{2v}
 $T^a=10890(120)$ gas PE¹

\tilde{X}^2B_1 C_{2v}

^aFrom vertical ionization potential.

Reference

¹J. M. Dyke, D. Haggerston, O. Warschkow, L. Andrews, A. J. Downs, and P. F. Souter, J. Phys. Chem. **100**, 2998 (1996).

Ga₂H₆⁺

\tilde{D}^2B_{1u} D_{2h}
 $T_0=31550(120)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g		GaH _t stretch	1540(40)	gas	PE	1

\tilde{C}^2B_{2u} D_{2h}
 $T^a=14040(240)$ gas PE¹

\tilde{B}^2B_{3u} D_{2h}
 $T^a=10970(240)$ gas PE¹

\tilde{A}^2A_g D_{2h}
 $T^a=8630(240)$ gas PE¹

\tilde{X}^2B_{2g} D_{2h}

Ga₂D₆⁺

\tilde{D}^2B_{1u} D_{2h}
 $T_0=31470(120)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g		GaH _t stretch	1135(40)	gas	PE	1

\tilde{C}^2B_{2u} D_{2h}
 $T^a=13470(240)$ gas PE¹

\tilde{B}^2B_{3u} D_{2h}
 $T^a=11050(240)$ gas PE¹

\tilde{A}^2A_g D_{2h}
 $T^a=8550(180)$ gas PE¹

\tilde{X}^2B_{2g} D_{2h}

^aFrom vertical ionization potential.

Reference

¹J. M. Dyke, D. Haggerston, O. Warschkow, L. Andrews, A. J. Downs, and P. F. Souter, J. Phys. Chem. **100**, 2998 (1996).

Ga₂H₆

\tilde{X} D_{2h}^a Structure: ED²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_g	1	GaH _t stretch	2000	Ar	Ra	3
			1986	N ₂	Ra	3
	2	GaH _b stretch	1474	Ar	Ra	3
			1471	N ₂	Ra	3
	3	GaH ₂ deform.	736	Ar	Ra	3
			733	N ₂	Ra	3
	b _{2u}	GaH _t stretch	1993m	gas	IR	1,2
			2015m	Ar	IR	1,2
			1996m			
	9	GaH ₂ rock	2000m	N ₂	IR	1,2
			760w	gas	IR	1,2
			773m	Ar	IR	1,2
			761m			
b_{1g}	11	GaH _t stretch	2011	Ar	Ra	3
			1997	N ₂	Ra	3
	b _{1u}	GaH _b stretch	1202s	gas	IR	1,2
			1213m	Ar	IR	1,2
			1208m			
	14	GaH ₂ rock?	1195m			
			1220s	N ₂	IR	1,2
	b _{3g}	GaH ₂ deform.	659m	Ar	IR	1,2
			655m			
b_{3u}	15	GaH ₂ deform.	653m			
			648m			
	16	GaH _t stretch	655s	N ₂	IR	1,2
			647w			
	17	GaH _b stretch	763	Ar	Ra	3
			761	N ₂	Ra	3
	18	GaH ₂ deform.	1976m	gas	IR	1,2
			1985s	Ar	IR	1,2
			1968m			
	17	GaH _t stretch	1985m	N ₂	IR	1,2
			1273s	gas	IR	1,2
	18	GaH _b stretch	1283s	Ar	IR	1,2
			1278m			
a_g	18	GaH ₂ deform.	1253m			
			1282m	N ₂	IR	1,2
			671vsT	gas	IR	1,2
			676vs	Ar	IR	1,2
			666s			
			673vs	N ₂	IR	1,2

Ga₂D₆

\tilde{X}		D _{2h} ^a				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a_g</i>	1	GaD _t stretch	1433	Ar	Ra	3
			1424	N ₂	Ra	3
	2	GaD _b stretch	1060	Ar	Ra	3
			1059	N ₂	Ra	3
	3	GaD ₂ deform.	527	Ar	Ra	3
			525	N ₂	Ra	3
<i>b_{2u}</i>	8	GaD _t stretch	1439m	gas	IR	1,2
	9	GaD ₂ rock	555w	gas	IR	1,2
	11	GaD _b stretch	1451	Ar	Ra	3
<i>b_{1u}</i>			1442	N ₂	Ra	3
	13	GaD _b stretch	860s	gas	IR	1,2
<i>b_{3g}</i>	14	GaD ₂ rock	439wm	gas	IR	1,2
	15	GaD ₂ deform.	543	Ar	Ra	3
<i>b_{3u}</i>			543	N ₂	Ra	3
	16	GaD _t stretch	1416mT	gas	IR	1,2
	17	GaD _b stretch	923s	gas	IR	1,2
	18	GaD ₂ deform.	484vsT	gas	IR	1,2

^aAs in Ref. 2, the axis definitions and vibrational numbering commonly used for diborane (J. L. Duncan, D. C. McKean, I. Torto, and G. D. Nivellini, J. Mol. Spectrosc. **85**, 16 (1981)) have been adopted.

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CH₂CHCH₂

\tilde{D} 2B ₂		C _{2v}				
T ₀ =41557.8(5)	gas	MPI ¹³	Ra ¹⁴			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a₂</i>	9	CH ₂ a-twist	596	gas	Ra	14
<i>b₁</i>	12	CH ₂ s-twist	564	gas	Ra	14

\tilde{C} 2B₁ C_{2v} Structure: MPI¹⁸
T₀=40305.5(5) gas AB²MPI^{11,13,18} $\tilde{C}-\tilde{X}$ 220–250 nm
In an argon matrix, a very prominent absorption maximum at 213 nm, with a shoulder at 220 nm, has been assigned⁵ to this transition of CH₂CHCH₂.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a₁</i>	7	C ₃ bend	385(2)	gas	MPI	11,18

A₀≈1.619; B₀≈0.351; C₀≈0.288 MPI¹¹

 \tilde{B}' 2A₁(3s) C_{2v}^a Structure: MPI¹⁸

T₀=40056.5(5) gas MPI^{7,8,13,18}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a₁</i>	7	C ₃ bend	379(2)	gas	MPI	7,8,18
<i>a₂</i>	9	CH ₂ a-twist	596(2)	gas	MPI	18
<i>b₁</i>	12	CH ₂ s-twist	491(7)T	gas	MPI	8

\tilde{A}' 2B₁ C_{2v}
T₀=24485 gas AB¹
24480 Ar AB⁵
 $\tilde{A}-\tilde{X}$ 370–410 nm
 $\tilde{A}-\tilde{X}$ 360–410 nm

Diffuse bands.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a₁</i>			1241	gas	AB	1
			1005	gas	AB	1
			908	gas	AB	1
			359	gas	AB	1

 \tilde{X} 2A₂ C_{2v} Structure: DL¹²

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a₁</i>	2	CH stretch	3048vw	Ar	IR	4–6
	4	CH ₂ scissors	1488(4)	gas	Ra	10,15
			1477w	Ar	IR	4–6
	5	CH ₂ rock	1245(3)	gas	Ra	10,15
	6	C ₃ stretch	1066(4)	gas	Ra	10,15
			990(20)	gas	PE	19
	7	C ₃ bend	427(4)	gas	MPI,Ra	7,8,10,15
				PE		18,19
<i>a₂</i>	9	CH ₂ a-twist	549(4)	gas	MPI,Ra	8,15,16
<i>b₁</i>	10	CH OPLA	968H	gas	Ra	10
			983.6s	Ar	IR	4–6,9
	11	CH ₂ s-wag	801.72	gas	DL,PE	12,19
			801.1vs	Ar	IR	3–6,9
	12	CH ₂ s-twist	518(4)	gas	MPI,Ra	7,8,10,15,16,18
				510.1s	Ar	IR
<i>b₂</i>	13	CH ₂ a-stretch	3105w	Ar	IR	4–6
	14	CII stretch	3016w	Ar	IR	4–6
	15	CH ₂ scissors	1463vw	Ar	IR	4–6
	16	CH bend	1389m	Ar	IR	4–6
	17	C ₃ stretch	1182	Ar	IR	6

A₀=1.802; B₀=0.346; C₀=0.290 DL¹²

CD₂CD₂ \tilde{D} 2B₂ C_{2v}
T₀=41532.1(5) gas MPI¹³ \tilde{C}' 2B₁ C_{2v}
T₀=40286.7(5) gas MPI^{13,18}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a₁</i>	7	C ₃ bend	321(2)	gas	MPI	18

$\tilde{B}^2A_1(3s)$ C_{2v}^a
 $T_0=40096(2)$ gas MPI^{7,13,18}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	7	C ₃ bend	316(2)	gas	MPI	7,18
a_2	9	CD ₂ a-twist	429(2)	gas	MPI	18

\tilde{A}^2B_1 C_{2v}
 $T_0=24745$ gas AB¹
Diffuse bands.

 $\tilde{A}-\tilde{X}$ 360–405 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1			1155	gas	AB	1
			981	gas	AB	1
			823	gas	AB	1

\tilde{X}^2A_2 C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	2	CD stretch	2285	Ar	IR	6
3	CD ₂ stretch	2214	Ar	IR	6	
4	C ₃ stretch	1272(8)	gas	Ra	17	
		1263	Ar	IR	6	
5	CD ₂ rock	1020(8)	gas	Ra	17	
		1018	Ar	IR	6	
		1007				
6	Deform.	835(20)	gas	MPI, PE	7,19	
7	C ₃ deform.	350(8)	gas	MPI, Ra	7,17–19	
				PE		
a_2	9	CD ₂ torsion	372H	gas	Ra	17
b_1	10	Deform.	762H	gas	Ra	17
11	Deform.	678(10)H	gas	PR	19	
		646.5vs	Ar	IR	6,9	
12	Deform.	403(8)	gas	Ra, MPI	17,18	
b_2	14	CD ₂ stretch	2209	Ar	IR	6
16	CD bend	1062	Ar	IR	6	

^aEvidence presented in Refs. 14 and 18 suggests that this state may be slightly nonplanar.

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CH₂CHCH₂[−]

Threshold for electron detachment from ground-state CH₂CHCH₂[−] = 3880(65) gas PE²

\tilde{X} C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1		C ₃ bend	425(20)	gas	PE	1,2

CD₂CD₂CD₂[−]

Threshold for electron detachment from ground-state CD₂CD₂CD₂[−] = 3740(50) gas PE²

\tilde{X} C_{2v}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1		C ₃ bend	350(20)	gas	PE	1,2

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H₂BSCH₃

\tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'	3	BH ₂ a-stretch	2603	Ar	IR	1
	4	BH ₂ s-stretch	2526	Ar	IR	1
	5	CH ₃ deform.	1496	Ar	IR	1
	6	CH ₃ deform.	1320	Ar	IR	1
	7	BH ₂ s-bend	1183	Ar	IR	1
	8	CH ₃ rock	1161	Ar	IR	1
	9	B-S stretch	830	Ar	IR	1
	10	BH ₂ rock	787	Ar	IR	1
	11	C-S stretch	693	Ar	IR	1
	14	CH ₃ stretch	2945	Ar	IR	1
	15	CH ₃ deform.	1448	Ar	IR	1
a''	16	CH ₃ rock	1138	Ar	IR	1
	17	BH ₂ OPLA	913	Ar	IR	1

Reference

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c-CH₃CH=NH

\tilde{X}		C _s	Structure: MW ²			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	NH stretch	3247vvw	Ar	IR	1
	2	CH ₃ a-stretch	3018m	Ar	IR	1
	3	CH stretch	2916	gas	IR	3
			2925ms	Ar	IR	1
			2920m			
	4	CH ₃ s-stretch	2886	gas	IR	3
			2885m	Ar	IR	1
	5	C=N stretch	1655	gas	IR	3
			1652vs	Ar	IR	1
	6	CH ₃ a-deform.	1438ms	Ar	IR	1
	7	CH ₃ s-deform.	1412m	Ar	IR	1
	8	Mixed	1250	gas	IR	3
			1252vs	Ar	IR	1
	9	CH deform.	1107	gas	IR	3
			1114ms	Ar	IR	1
	10	CNH deform.	1052wm	Ar	IR	1
	11	CH ₃ rock	950w	Ar	IR	1
	12	CCN deform.	485s	Ar	IR	1
<i>a''</i>	13	CH ₃ a-stretch	2988	gas	IR	3
			2990m	Ar	IR	1
	14	CH ₃ a-deform.	1454	gas	IR	3
			1435s	Ar	IR	1
			1433s			
	15	C=N torsion	1132vw	Ar	IR	1
	16	CH ₃ wag	1045	gas	IR	3
			1045vs	Ar	IR	1
<i>a'</i>	17	CH wag	678	gas	IR	3
			674w	Ar	IR	1

 $A_0 = 1.772; B_0 = 0.326; C_0 = 0.290 \text{ MW}^2\text{IR}^4$ **c-CH₃CD=ND**

\tilde{X}		C _s	Structure: MW ²			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	ND stretch	2425vvw	Ar	IR	1
	2	CD ₃ a-stretch	2256wm	Ar	IR	1
	3	CD stretch	2191m	Ar	IR	1
	4	CD ₃ s-stretch	2062w	Ar	IR	1
	5	C=N stretch	1631s	Ar	IR	1
			1628vs			
	6	Mixed	1159ms	Ar	IR	1
	7	Mixed	1072m	Ar	IR	1
	8	CD ₃ a-deform.	1032wm	Ar	IR	1
	9	Mixed	850w	Ar	IR	1
	10	CD rock	796s	Ar	IR	1
	11	CD ₃ rock	736m	Ar	IR	1
	12	CCN bend	405m	Ar	IR	1
	13	CD ₃ a-stretch	2216wm	Ar	IR	1
	14	CD ₃ a-deform.	1039s	Ar	IR	1
	15	CD ₃ wag	848ms	Ar	IR	1
	16	C=N torsion	811s	Ar	IR	1
	17	CD wag	527m	Ar	IR	1

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C₂H₃NH₂

\tilde{X}		C ₁ ^a	Structure: MW ³⁻⁵ IR ⁵			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	5	CH stretch	2987	gas	IR	2
			2976			
	6	C=C stretch	1672	gas	IR	2
			1668			
	7	NH ₂ scissors	1625T	gas	IR	2
	8	CH ₂ scissors	1454	gas	IR	2
	10	NH ₂ twist	1260	gas	IR	2
			1248			
	11	C-N stretch	1084	gas	IR	2
			1078			
	12	CH wag	1046	gas	IR	2
			1039			
	14	CH ₂ wag	812s	gas	IR	2
			805s			
	15	NH ₂ wag	615 ^b	gas	IR	2,4,5
			570 ^c	gas	IR	2,4,5
			470 ^d	gas	IR	4,5
			425 ^e	gas	IR	4,5
<i>a''</i>	16	C=C torsion	379 ^b	gas	IR	4,5
			333 ^d	gas	IR	4,5
			287 ^e	gas	IR	4,5

 $A_0 = 1.879; B_0 = 0.335; C_0 = 0.286 \text{ MW}^{1,3,5}$

^aThe barrier to inversion about the N atom is 356(3) cm⁻¹³⁻⁵ leading to splittings in a number of the vibrational band centers. The 0⁻-0⁺ transition has been observed^{5,6} in the far IR at 45.2 cm⁻¹. Ref. 5 has offered a tentative reassignment of the peaks associated with ν_{15} and ν_{16} .

^b1⁻-0⁺.^c1⁻-0⁻.^d1⁺-0⁺.^e1⁺-0⁻.**References**

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CH₃SCH₂⁺

\tilde{X}			Structure: MW ³⁻⁵ IR ⁵			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			870(70)	gas	PE	1
			390(100)	gas	PE	1

Reference

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C₂H₅O

\tilde{B} $^2A'$	C _s	$T_0=29210.207(6)$	gas	EM ^{1,2,4} LF ^{3,5,6}	$\tilde{B}-\tilde{X}$ 310–500 nm
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Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>		CO stretch	585H	gas	LF	5,6
		$\tau_0=1.7(2) \mu s$	gas	LF ³ EM ⁴		
		$A_0=1.117(2)$		$B_0=0.311(2)$		
		$C_0=0.268(2)$		LF ⁶		

\tilde{X} $^2A''$	C _s	$T_0=23519.6$	gas	LF ^{1,2}	$\tilde{A}-\tilde{X}$ 390–600 nm	
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH ₂ wag	1370	gas	LF	3
		C-O stretch	1067	gas	LF,EM	3–5
		C-C stretch	875 ^a	gas	LF	3
		CCO bend	442	gas	LF	3
		$A_0=1.318(2)$		$B_0=0.321(2)$		
		$C_0=0.284(3)$		LF ⁶		

^aAssignment to overtone of CCO bend cannot be excluded.

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HOCH₂CH₂

In the gas phase, the onset of continuous absorption has been reported¹ near 37700 (265 nm), with increasing absorption out to the 210 nm cutoff of the observations.

\tilde{X}	$T^a=46500(1000)$	D _{2h}	gas	PE ¹
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.
		OH stretch	3625.8T	Ar
		CH stretch	2991.0	Ar
		CH stretch	2922.4T	Ar
		CH stretch	2842.7	Ar
			1355.4	IR
			1172.5	IR
		CO stretch	1040.1	Ar
			873.9T	IR

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C₂H₅S

\tilde{B}	C _s	$T_0=23519.6$	gas	LF ^{1,2}	$\tilde{A}-\tilde{X}$ 390–600 nm
Predissociation occurs.					

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>		CH ₂ wag	1158.9	gas	LF	2
		CC stretch	1054.6	gas	LF	2
		CH ₃ wag	862.8	gas	LF	2
		CS stretch	420.5	gas	LF	1,2
		CCS bend	256.0	gas	LF	2
<i>a''</i>		CH ₂ twist	1203.3	gas	LF	2
		CH ₂ rock	718.4	gas	LF	2

$\tau=75$ ns gas LF¹

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>		CH ₃ s-stretch	2950T	gas	LF	2
		CH ₂ bend	1470(10)	gas	LF	2
		CH ₂ wag	1257(10)	gas	LF	2
		CC stretch	1075(10)	gas	LF	2
		CH ₃ wag	957(15)	gas	LF	2
		CS stretch	672.4	gas	LF	1,2
		CCS bend	296.0	gas	LF	2
<i>a''</i>		CH ₃ a-stretch	3050T	gas	LF	2
		CH ₂ twist	1290(10)	gas	LF	2
		CH ₃ wag	890(10)	gas	LF	2
		CH ₂ rock	478.3	gas	LF	2
		Torsion	271.9	gas	LF	2

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cyc-C₄H₄⁺

\tilde{F} 2A_g

$T^a=46500(1000)$ gas PE¹

\tilde{E} $^2B_{3u}$

$T^a=41600(1000)$ gas PE¹

\tilde{C} $^2B_{1u}$

$T^a=32000(1000)$ gas PE¹

\tilde{B} $^2B_{2u}$

$T^a=26300(1000)$ gas PE¹

$\tilde{X} \ ^2B_{2g}$		D _{2h}		Structure: PE ²		
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g			1129(40)	gas	PE	1

^aFrom vertical ionization potentials.

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cyc-C₄H₄

In an argon matrix,⁸ absorption is weak between 420 and 250 nm and is accompanied by dissociation to form C₂H₂. Below 250 nm, the absorption rises abruptly.

\tilde{X}		D _{2h}		Structure: IR ³⁻⁵ Ra ⁷		
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _g		CH stretch	3140(5)	Ne, Ar	Ra	7
		C=C stretch	1678(5)	Ne, Ar	Ra	7
		CH deform.	1059(5)	Ne, Ar	Ra	7
		CH deform.	989(5)	Ne, Ar	Ra	7
<i>b</i> _{1u}		CH stretch	3124w	Ar	IR	5,6,8
		C=C stretch	1527	Ne	IR	6
			1526w	Ar	IR	4-6,8
		CH deform.	1028vw	Ar	IR	5
<i>b</i> _{2g}		CH OPLA deform.	531(5)	Ne, Ar	Ra	7
<i>b</i> _{2u}		CH stretch	3105wm	Ar	IR	5,6,8
		CH deform.	1244	Ne	IR	6
			1242s	Ar	IR	1-6,8
		Ring deform.	719wm	Ar	IR	4-6,8
<i>b</i> _{3g}		CH stretch	3093(5)	Ne, Ar	Ra	7
		Ring deform.	723(5)	Ne, Ar	Ra	7
<i>b</i> _{3u}		CH OPLA	576	Ne	IR	6
			569vs	Ar	IR	1-6,8

cyc-C₄D₄

\tilde{X}		D _{2h}				
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> _{1u}		C=C stretch	1456w	Ar	IR	4
<i>b</i> _{2u}		CD deform.	1043wm	Ar	IR	4
		Ring deform.	609wm	Ar	IR	4
<i>b</i> _{3u}		CD OPLA	421vs	Ar	IR	4

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C₂H₃CH=C:

\tilde{b} ³ A''	C _s
$T_0=16420(240)$	gas PE ¹

\tilde{a} ³ A'	C _s
$T_0=15520(120)$	gas PE ¹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> '			1250(50)	gas	PE	1
			500(50)	gas	PE	1
			360(50)	gas	PE	1

$\tilde{X} \ ^1A'$

Broadening in the first band of the photoelectron spectrum of the anion permits estimation of a lifetime between 20 and 200 fs for rearrangement to form vinylacetylene.¹

Reference

- ¹R. F. Gunion, H. Köppel, G. W. Leach, and W. C. Lineberger, J. Chem. Phys. **103**, 1250 (1995).

C₂H₃CH=C:⁻

Threshold for electron detachment to form ground-state C₂H₃CH=C: =7370(120) gas PE¹

C₂D₃CD=C:⁻

Threshold for electron detachment to form ground-state C₂D₃CD=D: =7330(120) gas PE¹

Reference

- ¹R. F. Gunion, H. Köppel, G. W. Leach, and W. C. Lineberger, J. Chem. Phys. **103**, 1250 (1995).

CH₃CCOH

\tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OH stretch	3430.0 ^a	Ar	IR	1
		C=C stretch	2192.7	Ar	IR	1

CD₃CCOH \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OD stretch	2528.1 ^a	Ar	IR	1

^aProbably perturbed by interaction with NO trapped in an adjacent site.**Reference**¹J. A. Harrison and H. Frei, J. Phys. Chem. **98**, 12142 (1994).**H₂GaCl₂GaH₂** \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> _{2u}	8	GaH a-stretch	2020s	Ar	IR	1
	9	GaH ₂ rock	480w	Ar	IR	1
<i>b</i> _{1u}	13	GaH ₂ wag	578m	Ar	IR	1
	14	GaCl stretch	290w,sh	Ar	IR	1
<i>b</i> _{3u}	16	GaH s-stretch	1985m	Ar	IR	1
	17	GaH ₂ scissors	690s	Ar	IR	1
	18	GaCl s-stretch	265s	Ar	IR	1

D₂GaCl₂GaD₂ \tilde{X} D_{2h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>b</i> _{2u}	8	GaD a-stretch	1450s	Ar	IR	1
	9	GaD ₂ rock	355m	Ar	IR	1
<i>b</i> _{1u}	13	GaD ₂ wag	410m	Ar	IR	1
	14	GaCl stretch	305w,sh	Ar	IR	1
<i>b</i> _{3u}	16	GaD s-stretch	1420m	Ar	IR	1
	17	GaD ₂ scissors	495s	Ar	IR	1
	18	GaCl s-stretch	265s	Ar	IR	1

Reference¹C. R. Pulham, A. J. Downe, M. J. Goode, and I. M. Mills, Spectrochim. Acta A **51**, 769 (1995).**CH₃COO₂**A prominent gas-phase absorption with maximum near 207 nm has been attributed⁴⁻⁶ to CH₃COO₂.A weaker, broad gas-phase absorption with maximum near 245 nm has been attributed^{2,4-6} to CH₃COO₂.In an argon matrix, photodissociates in the 300–400 nm spectral region.³

\tilde{A}			$\tilde{A}-\tilde{X}$ 1348–1798 nm		
$T_0=5562(3)$	gas	AB ¹			

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		OO stretch	932(5) 530	gas gas	AB AB	1 1

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1843s	Ar	IR	3
			1425w	Ar	IR	3
			1420w	Ar	IR	3
			1367m	Ar	IR	3
			1278vw	Ar	IR	3
			1169vw	Ar	IR	3
			1152m	Ar	IR	3
			1099m	Ar	IR	3
			1082vw	Ar	IR	3
			1030w	Ar	IR	3
			971m	Ar	IR	3
			736m	Ar	IR	3

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C₆H₂⁺

\tilde{B}	$^2\Pi_u$	$D_{\infty h}$
$T_0=27350(160)$	gas	PE ¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			650(80)	gas	PE	1

\tilde{A}	$^2\Pi_g$	$D_{\infty h}$	$\tilde{A}-\tilde{X}$ 485–725 nm
$T_0=16658$	gas	EP ^{2,3} LF ³	$\tilde{A}-\tilde{X}$ 524–843 nm
		16570 Ne LF ⁴ AB ⁵	

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	CH stretch	3243(2)T	gas	LF	3
	2	C≡C stretch	2053(2)	gas	LF	3
			2065.7	Ne	LF,AB	4,5
	3	C≡C stretch	1880(2)	gas	LF	3
			1890.2	Ne	LF,AB	4,5
	4	C-C stretch	617(2)	gas	LF,EF	2,3
			620	Ne	LF,AB	4,5
Π_g	10	Skel. bend	244T	gas	LF	3
			248H	Ne	LF	4

 $\tau_0=17(2)$ ns gas EF²

$\tilde{X}^2\Pi_u$		$D_{\infty h}$				
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	2	C=C stretch	2182(2)	gas	EF	2,3
			2181.4	Ne	LF	4
	3	C=C stretch	1903(2)	gas	EF	2,3
			1903.4	Ne	LF	4
Σ_u^+	4	C-C stretch	632(2)	gas	EF	2,3
			635H	Ne	LF	4
	6	C=C stretch	2014T	Ne	LF	4
Π_g	10	Skel. bend	228(2)T	gas	LF	3
			230T	Ne	LF	4

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 HC_6H

\tilde{X}
 $A_0=8.95$; $B_0=0.0450$; $C_0=0.0447$ MW¹

Reference

- ¹M. C. McCarthy, M. J. Travers, A. Kovács, W. Chen, S. E. Novick, C. A. Gottlieb, and P. Thaddeus, *Science* **275**, 518 (1997).

 cyc-(HOC=COHC)=O \tilde{X}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		OH stretch	3506.3ms	Ar	IR	1
			1901.5m	Ar	IR	1
			1738.6s	Ar	IR	1
			1316.5vs	Ar	IR	1
			1279.4wm	Ar	IR	1
			1270.7wm	Ar	IR	1
			847.6wm	Ar	IR	1

Reference

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 C_7H

$\tilde{X}^2\Pi_{1/2}$
 $B_0=0.029$ MW¹

Reference

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 HC_5CN^+

\tilde{A}
 $T_0=17548(5)$ Ne AB¹

Reference

- ¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, *J. Chem. Phys.* **102**, 1510 (1995).

 C_8

$^3\Sigma_u^-$
 $T_0=15630(5)$ Ne AB³

$^3\Sigma_u^- - \tilde{X}$ 550–640 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1		2075(5)	Ne	AB	3
	4		471(5)	Ne	AB	3

$\tilde{X}^3\Sigma_g^-$ $D_{\infty h}$ Structure: ESR¹

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	4	Sym. stretch	565T	gas	PE	2
	5		2067.8	Ne	IR	3,6
			2063.9	Ar	IR	4
			2065.3	Kr	IR	5
	6		1707.8	Ne	IR	4–6
			1705.6	Ar	IR	4
			1706.0T	Kr	IR	5

Reference

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 C_8^-

Threshold for electron detachment from ground-state $\text{C}_8^- = 35330(50)$ gas PE^{1,2}

(2) $^2\Pi_u$
 $T_0=16305$ gas PD⁴
 $16295(5)$ Ne AB⁶

446–614 nm
438–614 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1		2125(10)	Ne	AB	6
	2		1821(10)	Ne	AB	6
	4		469(8)	Ne	AB	6
Π_g	8		487(5)H	Ne	AB	6

\tilde{C}	$^2\Pi_u$	D _{∞h}
$T_0=12963$	gas	PD ⁴
12933(5)	Ne	AB ³

$\tilde{C}-\tilde{X}$ 639–774 nm
 $\tilde{C}-\tilde{X}$ 665–775 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2087	gas	PD	4
	2		2055(5)	Ne	AB	3
	3		959(5)	Ne	AB	3
	4		453	gas	PD	4
			475(5)	Ne	AB	3

$^2\Sigma_u^+$	D _{∞h}
$T_0=9545(2)$	Ne AB ⁶

817–1048 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2010(4)	Ne	AB	6
	2		1956(4)	Ne	AB	6

\tilde{X}	$^2\Pi_g$	D _{∞h}
Vib.	No.	Approximate type of mode

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	6		1796.0	Ne	IR	5

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NC(C≡C)₂CN⁺

\tilde{E}	$^2\Pi_u$	D _{∞h}
$T_0=30420(160)$	gas	PE ¹

\tilde{D}	$^2\Pi_g$	D _{∞h}
$T_0=25580(160)$	gas	PE ¹

\tilde{B}, \tilde{C}	$^2\Sigma_g^+, ^2\Sigma_u^+$	D _{∞h}
$T_0=22190(160)$	gas	PE ¹

\tilde{A}	$^2\Pi_u$	D _{∞h}
$T_0=15260(10)$	gas	EF ¹

$\tilde{A}-\tilde{X}$ 630–770 nm
 $\tilde{A}-\tilde{X}$ 565–1510 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1	C≡C s-stretch	2475(5)T	Ne	AB	2
	2	CN s-stretch	1940(80)	gas	PE	1
			2029	Ne	AB,LF	2,3
	3	C-C s-stretch	1726(5)	Ne	AB	2
	4	C-C s-stretch	449	Ne	AB,LF	2,3

\tilde{X}		$^2\Pi_g$	D _{∞h}
Vib.	No.	Approximate type of mode	cm ⁻¹
Σ_g^+	1	C≡N s-stretch	2180(10)
			2185
	2	C≡C s-stretch	2100(10)
			2094
Σ_g^+	3	C-C s-stretch	1360(10)
			1354
	4	C-C s-stretch	460(10)
			458
			Ne
			LF
			1
			3
			1
			3

^aRef. 3 reported a second trapping site with $T_0=15160$.

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C₇O

$$\tilde{X} = \frac{C_{\infty v}}{B_0 = 0.019} \text{ MW}^{\frac{1}{2}}$$

Reference

- T. Ogata, Y. Ohshima, and Y. Endo, J. Am. Chem. Soc. **117**, 3593 (1995).

FB(NCO)₂

\tilde{X}	C _s		
Vib.	No.		
a'	1	NCO a-stretch	2296s
	2	NCO a-stretch	2233vs
		Mixed	1569Ts
			1560Ts
	5	BF stretch	1553Ts
			1276s
			gas
			IR
			1
			1
			1
			1

Reference

- M. Alaee, E. G. Livingstone, and N. P. C. Westwood, J. Am. Chem. Soc. **115**, 2871 (1993).

FC(O)OONO₂

In the gas phase, the onset of unstructured absorption with continuously increasing intensity out to the 190-nm observation limit occurs near 34500 (290 nm).¹

\tilde{X}	C_1	Structure: ED ¹				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
1		C=O stretch	1910	gas	IR	1
			1902.5wm	Ne	IR	1
2			1763	gas	IR	1
			1762.6m	Ne	IR	1
3			1302	gas	IR	1
			1302.0wm	Ne	IR	1
4			1194	gas	IR	1
			1191.7vs	Ne	IR	1
5			981	gas	IR	1
			983.3w	Ne	IR	1
6			925	gas	IR	1
			926.0w	Ne	IR	1
7			795	gas	IR	1
			792.0wm	Ne	IR	1
8			712	gas	IR	1
			703.2vw	Ne	IR	1
9			635	gas	IR	1
			631.3w	Ne	IR	1
10			530	gas	IR	1
			530.0w	Ne	IR	1
11			475T	gas	IR	1
12			420	gas	IR	1
13			395	gas	IR	1
14			343	gas	IR	1
15			331	gas	IR	1
17			75T	gas	IR	1

Reference

¹D. Scheffler, I. Schaper, H. Willner, H.-G. Mack, and H. Oberhammer, Inorg. Chem. **36**, 339 (1997).

CF₃ONO₂

\tilde{X}						
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
NO ₂		a-stretch	1744	gas	IR	2
			1735.5	Ar	IR	1
			1345.2	Ar	IR	1
NO ₂		s-stretch	1334	gas	IR	2
			1327.1	Ar	IR	1
CF ₃		stretch	1264	gas	IR	2
			1260.8s	Ar	IR	1
CF ₃		stretch	1244	gas	IR	2
			1249.5vs	Ar	IR	1
			1151	gas	IR	2
			1142.5vs	Ar	IR	1
C-O		stretch	924	gas	IR	2
			924.1w	Ar	IR	1
NO ₂		scissors	791	gas	IR	2
			785.2	Ar	IR	1
ONO ₂		OPLA	752.9	Ar	IR	1
			519.0	Ar	IR	1

References

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8.15. Hydrocarbons with More Than Eight Atoms**t-H₂C=CH—CH=CH₂⁺**

$T^a=50180(120)$ gas PE¹



$T^a=31950(120)$ gas PE¹

An absorption maximum which appears at 33890 (295 nm) in an argon matrix has been attributed² to t-butadiene cation.



$T^a=26300(120)$ gas PE¹



$T^a=19850(120)$ gas PE¹

A relatively weak absorption maximum which appears at 18720 (534 nm) in an argon matrix has been attributed² to t-butadiene cation.

 \tilde{X} C_{2h}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_u	11		1000.9	Ar	IR	3,4
b_u	17		3125.3	Ar	IR	3,4
	18		3073.7	Ar	IR	3,4
	19		3026.3	Ar	IR	3,4
	20		1477.5	Ar	IR	3,4
	21		1330T	Ar	IR	3,4
	22		1251.1	Ar	IR	3,4
	23		1006.1	Ar	IR	3,4

t-D₂C=CD—CD=CD₂⁺ **\tilde{X} C_{2h}**

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_u	11		739.6	Ar	IR	3,4
b_u	17		2358.1	Ar	IR	3,4
	18		2276.1	Ar	IR	3,4
	19		2215.4	Ar	IR	3,4
	20		1319.1	Ar	IR	3,4
	21		1047.2	Ar	IR	3,4
	22		996.1	Ar	IR	3,4

^aFrom vertical ionization potential.

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⁴T. Keszthelyi, R. Wilbrandt, T. Bally, and J.-L. Roulin, J. Phys. Chem. **100**, 16850 (1996).

C(CH₂)₃

\tilde{b}^1A_1
 $T_0=5640(50)$ gas PE³

 $\tilde{X}^3A'_2$ D_{3h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2		758	Ar	IR	1	
		755.5vs	Xe	IR	1,2	
		500	Ar	IR	1	
		499.0m	Xe	IR	1,2	
		3115.1w	Xe	IR	1,2	
		3100.1w				
e'		3031.2	Xe	IR	1,2	
		3019.1w				
		1418.4wm	Xe	IR	1,2	

C(CD₂)₃ $\tilde{X}^3A'_2$ D_{3h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a''_2		671.1s	Xe	IR	2	
		415.6vs	Xe	IR	2	
e'		2204.9w	Xe	IR	2	
		2192.5vw				
		1050.6wm	Xe	IR	2	

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³P. G. Wentholt, J. Hu, R. R. Squires, and W. C. Lineberger, J. Am. Chem. Soc. **118**, 475 (1996).

C(CH₂)₃⁻

Threshold for electron detachment from ground-state C(CH₂)₃⁻=3480(50)
gas PE¹

Reference

¹P. G. Wentholt, J. Hu, R. R. Squires, and W. C. Lineberger, J. Am. Chem. Soc. **118**, 475 (1996).

H₂CC(CH₃)CH₂⁺ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			410(30)	gas	PE	1

Reference

¹J. C. Schultz, F. A. Houle, and J. L. Beauchamp, J. Am. Chem. Soc. **106**, 7336 (1984).

cyc-C₄H₇⁺ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			990(100)	gas	PE	1

Reference

¹J. C. Schultz, F. A. Houle, and J. L. Beauchamp, J. Am. Chem. Soc. **106**, 7336 (1984).

CH₃CHCH=CH₂ $T_0=42039$ gas AB¹

227–238 nm

 \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'		H deformation	777.7	Xe	IR	2
	11	CH s-bend	1492	gas	Ra	3
	16	C—CH ₃ s-stretch	1265	gas	Ra	3
	17	C—C—C s-bend	872	gas	Ra	3
	18	Skel. s-bend	499	gas	Ra	3
	22	CH OPLA	289	gas	Ra	3
a''	22	CH ₃ torsion	952H	gas	Ra	3
	27		103H	gas	Ra	3

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¹A. B. Callear and H. K. Lee, Trans. Faraday Soc. **64**, 308 (1968).

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H₂CC(CH₃)CH₂gas AB^{1,5}

223–258 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1		CH stretch	2830T	gas	AB	1
		C ₃ deformation	350T	gas	AB	1

²A₁ (3s) C_{2v}
 $T_0=38369(30)$ gas MPI²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1		C ₃ deformation	432(30)	gas	MPI	2

\tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	5	CH ₂ stretch	3109	N ₂	IR	3,4
		CH ₂ stretch	3025	N ₂	IR	3,4
			1000	N ₂	IR	3,4
			475	N ₂	IR	3,4
		CH ₂ scissors	1492(4)	gas	Ra	6
	7		1498	N ₂	IR	3,4
		CH ₃ s-deform.	1392(4)	gas	Ra	6
	8		1394	N ₂	IR	3,4
		CH ₂ rock	1353(4)	gas	Ra	6
		C ₃ stretch	1037(4)	gas	Ra,PE	6,7
		C-CH ₃ stretch	865(4)	gas	Ra	6
		CCH ₂ OPLA	780(10)H	gas	PE	7
<i>a''</i>	14		795	N ₂	IR	3,4
		CH ₂ twist	538HT	gas	Ra	6
	15	C ₃ deformation	432(4)	gas	Ra,PE	6,7
		CH ₂ scissors	1471	N ₂	IR	3,4
		CH ₃ deformation	1400	N ₂	IR	3,4

Reference

¹P. G. Wenthold, M. L. Polak, and W. C. Lineberger, J. Phys. Chem. **100**, 6920 (1996).

 $n\text{-C}_4\text{H}_9$ \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	1-CH ₂ a-stretch	3105	Ar	IR	1-3
		1-CH ₂ s-stretch	3024	Ar	IR	1-3
			2886	Ar	IR	2
		2-CH ₂ a-stretch	2835	Ar	IR	2,3
		2-CH ₂ s-stretch	2809	Ar	IR	1-3
	2		1471	Ar	IR	2
		-CH ₂ - deform.	1463	Ar	IR	2,3
		2-CH ₂ deform.	1425	Ar	IR	2,3
		2-CH ₂ deform.	1183	Ar	IR	2,3
		(C ₃ H ₇)CH ₂ OPLA	1098	Ar	IR	2
	3		520s	Ar	IR	1,3

 $\text{D}_2\text{CC}(\text{CD}_3)\text{CD}_2$ \tilde{X} C_s

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1		845(20)	gas	PE	7
		C ₃ deformation	350(20)	gas	PE	7

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 $\text{H}_2\text{CC}(\text{CH}_3)\text{CH}_2^-$

Threshold for electron detachment from ground-state $\text{H}_2\text{CC}(\text{CH}_3)\text{CH}_2^-$
 $=4070(50)$ gas PE¹

 \tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C ₃ deform.	425(20)	gas	PE	1

 $\text{D}_2\text{CC}(\text{CD}_3)\text{CD}_2^-$

Threshold for electron detachment from ground-state $\text{D}_2\text{CC}(\text{CD}_3)\text{CD}_2^-$
 $=3980(65)$ gas PE¹

cyc-C₅D₅

\tilde{A}^2A_2'' D_{5h}
 $T_0=29819.434(2)$ gas AB⁴LF⁹

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
e_2''		CD bend	233	gas	AB	4

$B_0=0.233$; $C_0=0.117$ LF⁹

\tilde{X}^2E_1' D_{5h}
 $B_0=0.240$; $C_0=0.120$ LF⁹

^aAlternatively, may be contributed by $2\nu_{12}$.

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cyc-C₅H₆⁺

The gas-phase photoelectron spectrum¹ of cyc-C₅H₆ shows several overlapping bands with unresolved structure, corresponding to excited states of cyc-C₅H₆⁺ between 25300 and 50300.

\tilde{B}^2B_1 C_{2v}
 $T_0=27430$ Ar AB³

$\tilde{B}-\tilde{X}$ 300–370 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1			1062(15)	Ar	AB	3

\tilde{A}^2B_1 C_{2v}
 $T_0=16570(20)$ gas PE^{1–3}
 16500 T Ar AB³

$\tilde{A}-\tilde{X}$ 450–650 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1			807(15)	gas	PE	2,3
			835(15)	Ar	AB	3

		\tilde{X}^2A_2		C_{2v}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
a_1	3		2885	Ar	IR	3	
	4		1452(15)	gas	PE	1–3	
	6		1456	Ar	IR	3	
	7		1325	Ar	IR	3	
	9		1113(15)	gas	PE	1–3	
	10		944	Ar	IR	3	
	15		807(15)	gas	PE	1–3	
	17		2901	Ar	IR	3	
	18		866	Ar	IR	3	
	22		612	Ar	IR	3	
b_1	23		1423	Ar	IR	3	
	24		1313	Ar	IR	3	
	25		1266	Ar	IR	3	
	26		1071	Ar	IR	3	
			920	Ar	IR	3	

cyc-C₅D₆⁺

\tilde{B}^2B_1 C_{2v}
 $T_0=27600$ T Ar AB³

$\tilde{B}-\tilde{X}$ 300–370 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1			892(15)	Ar	AB	3

\tilde{A}^2B_1 C_{2v}
 $T_0=16500$ T Ar AB³

$\tilde{A}-\tilde{X}$ 450–650 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1			760(15)	gas	PE	3
			775(15)	Ar	AB	3

 \tilde{X}^2A_2 **C_{2v}**

$T_0=27600$ T Ar AB³

$\tilde{X}-\tilde{A}$ 300–370 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	4		1412(15)	gas	PE	3
	5		1418	Ar	IR	3
	6		1312	Ar	IR	3
	7		1035	Ar	IR	3
	8		954	Ar	IR	3
	9		823(15)	gas	PE	3
	17		760	Ar	IR	3
	18		686	Ar	IR	3
	22		458	Ar	IR	3
	23		1345	Ar	IR	3
b_1	27		1204	Ar	IR	3
			707	Ar	IR	3

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n-C₅H₁₁

1

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs
		1-CH ₂ a-stretch	3103	Ar	IR	1,2
		1-CH ₂ s-stretch	3025	Ar	IR	1,2
		2-CH ₂ a-stretch	2838	Ar	IR	1,2
		2-CH ₂ s-stretch	2802	Ar	IR	1,2
		2-CH ₂ deform.	1425	Ar	IR	1,2
		2-CH ₂ deform.	1181	Ar	IR	1,2
			1096	Ar	IR	1
		(C ₄ H ₉)CCH ₂ OPLA	519s	Ar	IR	1,2

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m-C₆H₄

(m-Benzyne)

\widetilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1		CH stretch	3037w	Ar	IR	1
		Mixed	1402wm	Ar	IR	1
b_1		CH OPLA	824wm	Ar	IR	1
		CH OPLA	751ms	Ar	IR	1
b_2		Mixed	1486wm	Ar	IR	1
		HCC deform.	936m	Ar	IR	1
		Ring deform.	547vs	Ar	IR	1

Reference

- ¹ R. Marquardt, W. Sander, and E. Kraka, *Angew. Chem.* **108**, 825 (1996); *Angew. Chem. Int. Ed. Engl.* **35**, 746 (1996).

C₆H₅

A broad gas-phase absorption with maximum near 40820 (245 nm) has been attributed⁷ to C₆H₅. In an argon matrix, a broad, very prominent absorption maximum at 41060, with a partially resolved shoulder at 38360, has also been attributed¹² to C₆H₅.

A prominent absorption near 34720 (288 nm) in argon-matrix studies has been assigned^{4,9,12} to C₆H₅.

$$\tilde{A}^{-2}B_1 \quad C_{2v}$$

$$T_0 = 18908 \text{ gas } AB^1$$

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			896	gas	AB	1
			722	gas	AB	1
			571	gas	AB	1

Gas-phase photoelectron spectra¹⁰ suggest that there is an excited electronic state of C₆H₅ at or below 13700.

$$\widetilde{X} \quad ^2A_1 \qquad \qquad C_{2v}$$

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	CH s-stretch	3085w	Ar	IR	6,8,11
	2	CH s-stretch	3071vw	Ar	IR	11
	3	CH s-stretch	3052vw	Ar	IR	11
	4	Ring stretch	1499w	Ar	IR	11
	5	Ring stretch	1441wm	Ar	IR	8,11
	6	CH deform.	1080w	Ar	IR	11
	7	Ring deform.	1027wm	Ar	IR	6,8,11
	8	Ring breathing	1011vw	Ar	IR	11
	9	Ring deform.	968(15)	gas	PE	10
10			976w	Ar	IR	8,11
			600(10)	gas	PE	10
			605w	Ar	IR	8,11
b_1	14	CH deform.	971vw	Ar	IR	11
	15	CH deform.	878w	Ar	IR	11
	16	CH deform.	707s	Ar	IR	2-6,8,11
	17	CH deform.	656ms ^a	Ar	IR	8
b_2	18	Deformation	416w	Ar	IR	11
	19	CH a-stretch	3073wm	Ar	IR	8,11
	20	CH a-stretch	3060vw	Ar	IR	8,11
b_3	21	Ring stretch	1593vw	Ar	IR	11
	22	Ring stretch	1433wm	Ar	IR	6,8,11
	23	CH deform.	1344vw	Ar	IR	11
	24	Ring deform.	1226wm	Ar	IR	6,11
	25	CH deform.	1086vw	Ar	IR	11
	26	Ring deform.	1067w	Ar	IR	8,11
	27	Ring stretch	586vw	Ar	IR	8,11

C₆D₅

A prominent absorption at 286 nm in argon-matrix studies of photolyzed benzene samples has been attributed⁴ to C₆D₆.

$$\begin{array}{ccc} \widetilde{A} & {}^2B_1 & C_{2v} \\ T_0 = 18840 & \text{Ar} & \text{AB}^4 \end{array}$$

$\tilde{A} = \tilde{X}$ 530–535 nm

$$\widetilde{X}^2A_1 \quad C_{2v}$$

Vib. sym.	No.	Approximate type of mode	cm^{-1}	McD.	Type meas.	Refs.
a_1			960(20) 588(10)	gas gas	PE PE	10 10
b_1		CD deform.	519s	Ar	IR	2,5

^aThis absorption is assigned to C₆H₅ formed by the photolysis of C₆H₅NO, but would be obscured by a very prominent CO₂ absorption when C₆H₅ is formed by the photolysis of a benzoyl peroxide. Its position and intensity agree more satisfactorily with the *ab initio* calculations of Ref. 11 than do those for the assignment proposed in that paper.

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C₆H₆⁺

\tilde{G} $^2A_{1g}$ D_{6h}
T^a=61290(100) gas PE^{1,5}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _{1g}	1 ^b (2)	CH stretch	2790(100)	gas	PE	1
	2 (1)	Ring stretch	930(100)	gas	PE	1

\tilde{F} $^2B_{1u}$ D_{6h}
T^a=50160(100) gas PE^{1,5}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _{1g}	1 ^b (2)	CH stretch	2340(100)	gas	PE	1

\tilde{E} $^2B_{2u}$ D_{6h}
T^a=45320(100) gas PE⁵

\tilde{D} $^2E_{1u}$ D_{6h}
T^a=38220(100) gas PE⁵

\tilde{C} $^2A_{2u}$ D_{6h}
T^a=25310(100) gas PE⁵

In the gas phase, the mass-selected ion-dip spectrum of C₆H₆⁺ shows a broad, unstructured absorption with onset near 19000 and increasing in intensity up to the detection limit of 24000.¹⁰

A broad, unstructured absorption near 24000 may have been contributed by the $\tilde{C}-\tilde{X}$ transition of C₆H₆⁺ produced by vacuum-ultraviolet photolysis of benzene isolated in a neon matrix.²

T₀≈19840 Ar LF³AB⁴ C-X 420-547 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>e</i> _{2g}	18 ^b (6)	Ring deform.	600(30)T	Ar	AB	4

\tilde{B} $^2E_{2g}$ D_{6h}
T₀=18113 gas PF⁹PRI¹³

$\tilde{B}-\tilde{X}$ 510-550 nm
In an argon matrix, a weak, sharp absorption at 18100 has been attributed³ to a vibronically allowed transition in the excitation of C₆H₆⁺ to the \tilde{B} state.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _{1g}	1 ^b (2)	CH stretch	2850(100)	gas	PE	1
	2 (1)	Ring stretch	990 ^c	gas	PF	9
	16 (8)	Ring stretch	1520(100)T	gas	PE	1
	17 (9)	CH bend	1140 ^c	gas	PF	9
	18 (6)	Ring deform.	645(100) ^d	gas	PE	1
	19 (17)	CH bend	574	gas	PF	9
	20 (16)	Ring deform.	224	gas	PF	9

\tilde{X} $^2E_{1g}$ D_{6h} Structure: TPE¹¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _{1g}	1 ^b (2)	CH stretch	2960(100)T	gas	PE	1
	2 (1)	Ring stretch	975(15)	gas	PE, MPI	1,6,8
	8 (4)	Ring deform.	415(20)	gas	PE	6
	11 (10)	CH bend	835(15)	gas	MPI	8
	16 (8)	Ring stretch	1561(20) ^d	gas	PE	1,6
			1480(10)T	Ar	LF	3
	17 (9)	CH bend	1230(15) ^d	gas	PE, MPI	1,6,8
	18 (6)	Ring deform.	676.4 ^d	gas	TPE, MPI	1,6,8,12
			630(10)T	Ar	LF	3
			365.4 ^e	gas	TPE	12
			345.4 ^f	gas	PE, TPE	6,12
<i>e</i> _{2u}	20 (16)	Ring deform.	295(5)	gas	PE, MPI	6,8

B₀=0.189; C₀=0.095 PE⁷TPE¹¹TPI¹⁴

C₆D₆⁺

\tilde{G} $^2A_{1g}$ D_{6h}
T₀≈62000 gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> _{1g}	1 ^b (2)	CD stretch	2240(100)	gas	PE	1
	2 (1)	Ring stretch	920(100)	gas	PE	1

\tilde{F} $^2B_{1u}$ D_{6h}
 $T^a \approx 50000$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_{1g}	1 ^b (2)	CD stretch	1610T	gas	PE	1

\tilde{C} $^2A_{2u}$ D_{6h}
 $T_0 \approx 19930^c$ Ar LF³AB⁴ $\tilde{C}-\tilde{X}$ 470–545 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
e_{2g}	18 ^b (6)	Ring deform.	590(60)T	Ar	AB	4

\tilde{B} $^2E_{2g}$ D_{6h}
 $T^a \approx 18600$ gas PE¹

In an argon matrix, a weak, sharp absorption at 18215 has been attributed³ to a vibronically allowed transition in the excitation of C₆D₆⁺ to the \tilde{B} state.

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_{1g}	1 ^b (2)	CD stretch	2140(100)	gas	PE	1
e_{2g}	16 (8)	Ring stretch	1450(100)	gas	PE	1
	17 (9)	CD bend	870(100)	gas	PE	1
	18 (6)	Ring deform.	600(100)	gas	PE	1

\tilde{X} $^2E_{1g}$ D_{6h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_{1g}	1 ^b (2)	CD stretch	2330(100)	gas	PE	1
	2 (1)	Ring stretch	928(20)	gas	PE	1,6
b_{2g}	8 (4)	Ring deform.	351(20)	gas	PE	6
e_{2g}	16 (8)	Ring stretch	1565(100)	gas	PE	1
	18 (6)	Ring deform.	1460(10)T 637(20) ^d	Ar gas	LF PE	3 1,6
			590(10)T 343(20) ^g	Ar gas	LF PE	3 6
e_{2u}	20 (12)	Ring deform.	278(20)	gas	PE	6

B = 0.155; C = 0.076 TPI¹⁴

^aFrom vertical ionization potentials. The first ionization potential of benzene is taken to equal 74555.0(4), or 9.2405 eV, from Ref. 7.

^bIn order to be consistent with other vibrational numberings in these tables, the vibrational numbering of Herzberg is used here. However, many authors use instead the vibrational numbering of Wilson (E. B. Wilson, Jr., Phys. Rev. **45**, 706 (1934)). This latter numbering is indicated in parentheses.

^cFrom analysis of combination bands.

^dj = ± 1/2.

^gj = +3/2.

^fj = -3/2.
^gj = 3/2.

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(CH₃)₃CCH=C:

\tilde{a}' $^3A'$ C_s
 $T_0 = 15930(120)$ gas PE¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			1230(50) 800(50) 520(50) 340(50)	gas gas gas gas	PE PE PE PE	1 1 1 1

\tilde{X} $^1A'$ C_s

Reference

- R. F. Gunion and W. C. Lineberger, J. Phys. Chem. **100**, 4395 (1996).

(CH₃)₃CCH=C⁻

Threshold for electron detachment from ground-state (CH₃)₃CCH=C⁻
= 5200(120) gas PE¹

\tilde{X} $^2A'$ C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			370(50)	gas	PE	1

Reference

- R. F. Gunion and W. C. Lineberger, J. Phys. Chem. **100**, 4395 (1996).

C₇H₂⁺

$\tilde{A}^2\Pi_u$ D_{∞h}
 $T_0=16676(5)T$ Ne AB¹

Reference

¹P. Freivogel, J. Fulara, D. Lessen, D. Forney, and J. P. Maier, Chem. Phys. **189**, 335 (1994).

HC₇H

$^3\Sigma_u^-$ D_{∞h}
 $T_0=19812(5)$ Ne AB¹
 19770(5)

$^3\Sigma_u^- - \tilde{X}$ 410–506 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	2	C≡C s-stretch	1961(5)	Ne	AB	1
	4		647(5)	Ne	AB	1
Π		CH bend	286HT	Ne	AB	1

DC₇D

$^3\Sigma_u^-$ D_{∞h}
 $T_0=19932(5)$ Ne AB¹
 19891(5)

$^3\Sigma_u^- - \tilde{X}$ 410–503 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	2	C≡C s-stretch	1935(5)	Ne	AB	1
	4		551(5)	Ne	AB	1
Π		CD bend	202HT	Ne	AB	1

Reference

¹J. Fulara, P. Freivogel, D. Forney, and J. P. Maier, J. Chem. Phys. **103**, 8805 (1995).

C₆H₅CH₂⁺

\tilde{X}^1A_1

C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	11		993(2)	gas	TPE	3
	12		820(2)	gas	TPE	3
	13	Ring deform.	525(2)	gas	PE, TPE	1–3
a_2	16		627T	gas	TPE	3
	17		316H	gas	TPE	3
b_1	23		400H	gas	TPE	3
	24		163H	gas	TPE	3
b_2	28		1547(2)	gas	TPE	3
	32		1183T	gas	TPE	3
	33		1115T	gas	TPE	3
	34		973(2)	gas	TPE	3
	35	Ring deform.	598(2)	gas	TPE	2,3
	36		353(2)	gas	TPE	3

C₆H₅CD₂⁺

\tilde{X}^1A_1 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	11		951(2)	gas	TPE	3
	12		737(2)	gas	TPE	3
	13	Ring deform.	500T	gas	PE, TPE	1–3
a_2	17		278H	gas	TPE	3
b_1	23		349T	gas	TPE	3
	24		145T	gas	TPE	3
b_2	32		880T	gas	TPE	3
	33		862T	gas	TPE	3
	34		779T	gas	TPE	3
	35	Ring deform.	575(2)	gas	TPE	2,3
	36		302(2)	gas	TPE	3

^aIf, as in Refs. 2 and 3, the x axis is chosen in the molecular plane, interchange of the b_1 and b_2 representations results. However, the international convention established in 1955 is followed in the symmetry designations given here. This necessitates a renumbering of the vibrational fundamentals, so that modes 18–24 (numbered 30–36 in Refs. 2 and 3) are of b_1 symmetry and 25–36 (numbered 18–29 in Refs. 2 and 3) of b_2 symmetry.

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C₆H₅CH₂**E**

A gas-phase absorption maximum near 230 nm has been attributed⁹ to the first Rydberg transition of C₆H₅CH₂.

D

A gas-phase absorption between 260 and 245 nm, with maximum near 255 nm, has been attributed⁹ to the $\tilde{D}-\tilde{X}$ transition of C₆H₅CH₂. This band may have been excited in the MPI studies of Ref. 16. In a neon matrix, the counterpart of this absorption maximum has been observed near 245 nm.⁶

\tilde{C}^2A_2 C_{2v}
 $T_0=32760$ gas AB^{2,5,7}
 32730 Ne AB⁶ $\tilde{C}-\tilde{X}$ 291–309 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a_1	7a	C-CH ₂ stretch	1145	gas	AB	7
	18a	CH deform.	968	gas	AB	7
	1	Ring breathing	931	gas	AB	7
	12a	Ring deform.	804	gas	AB	7
b_2	6a	Ring deform.	432	gas	AB	7
	6b	Ring deform.	525	gas	AB	7
	18b	CII deform.	276	gas	AB	7

$\tilde{B}^2B_1^a$ C_{2v}
 $T_0=22850T$ gas AB¹¹EM¹¹LF^{22,24}

Vibronically coupled to modes of b_2 symmetry in the \tilde{A} state.^{11,22,24,27}
 $A^b=0.179$; $B^b=0.088$; $C^b=0.059$ gas LF²⁴

\tilde{A}^2A_2		C_{2v}
$T_0=22001.5$	gas	$EM^{1,3,5,10}AB^4LF^{12,20-22,26}$
22003	Ne	AB^6
21862	Ar	LF^{13}

$\tilde{A}-\tilde{X}$ 429–455 nm
 $\tilde{A}-\tilde{X}$ 429–455 nm
 $\tilde{A}-\tilde{X}$ 430–510 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
6a	Ring deform.	910	Ar	LF	13	
			770T	gas	LF	20
		798	Ar	LF	13	
		437	gas	AB,LF	11,18,20,22	
		433	Ne	AB	6	
		456	Ar	LF	13	
6b	Ring deform.	388 ^c	gas	AB,LF	11,17,18,22	
		402	Ne	AB	6	
6b	Ring deform.	328 ^c	gas	AB,LF	11,17,18,22	
		344	Ne	AB	6	

$\tau_{0(1)}=0.4 \mu s$; $\tau_{0(2)}=1.85 \mu s$ gas $LF^{15,20,21}$
 $A^d=0.180$; $B^d=0.088$; $C^d=0.059$ $EM^{10,17,19}LF^{22,24}$

$\tilde{X}^2B_1^a$		C_{2v}			
Vib.	No.	Approximate type of mode			
		cm^{-1}	Med.	Type meas.	Refs.
a_1	CH stretch	3069w	Ar	IR	25
	8a Ring stretch	1603	gas	EM	5,8,18
	C-CH ₂ stretch	1510(25)T	gas	PE	23
		1469m	Ar	IR	25
19a	Ring stretch	1431	gas	EM	5,8,18
		1423	Ar	LF	13
	CH ₂ deform.	1409m	Ar	IR	25
7a		1258	gas	EM,LF	5,7,18,20
		1264m	Ar	IR	25
9a	CH deform.	1181	gas	EM	5,8
18a	CH deform.	1046	gas	EM	18
1	Ring breathing	987.4	gas	EM,LF	5,8,10,18,20
		982	Ar	LF	13
12a	Ring deform.	830	gas	EM,LF	5,8,18,20
6a	Ring deform.	524	gas	EM,LF,PE	5,8,10,20,23
		520	Ar	LF	13
a_2	17a CH deform.	963	gas	EM	18
	10a CH deform.	860	gas	EM	8,18
b_1	16a Ring deform.	393	gas	EM	8
		882.0w	Ar	IR	25
		762.0vs	Ar	IR	25
b_2	CH ₂ wag	710.9w	Ar	IR	25
		667.0s	Ar	IR	25
		465.0s	Ar	IR	25
16b	Ring deform.	430	gas	EM	8
	CH ₂ a-stretch	3111w	Ar	IR	25
8b	Ring stretch	1549	gas	EM	5,8,18
		1530	Ar	LF	13
		1446m	Ar	IR	25
		1305w	Ar	IR	25
9b	CH deform.	1152	gas	EM	5,8,18
15	CH deform.	1089	gas	EM	5,8
		1015w	Ar	IR	25
		948.1vw	Ar	IR	25
6b	Ring deform.	615	gas	EM,LF	5,8,10,18,20
		612	Ar	LF	13
18b	CH deform.	356	gas	EM,LF	5,8,18,20
		357	Ar	LF	13

$A_0=0.184$; $B_0=0.090$; $C_0=0.060$ $EM^{10,17,19}LF^{24}$

$C_6D_5CD_2$

$\tilde{B}^2B_1^a$	C_{2v}
$T_0=22455(10)$	gas $AB^{11}EM^{11}$

\tilde{A}	C_{2v}
$T_0=22093.7$	gas $EM^{5,10}$

21962 Ar LF¹³ $\tilde{A}-\tilde{X}$ 434–502 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			844	Ar	LF	13
			731	Ar	LF	13

$\tau \approx 1340$ ns gas LF¹⁴

$\tilde{X}^2B_1^a$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	8a	Ring stretch	1593	gas	EM	5,8
	19a	Ring stretch	1327	gas	EM	5,8
			1323	Ar	LF	13
	7a	C-CD ₂ stretch	1204	gas	EM	5,8
	1	Ring breathing	945.7	gas	EM	5,8,10
			945	Ar	LF	13
	9a	CD deform.	895	gas	EM	5,8
	18a	CD deform.	848	gas	EM	8
	12a	Ring stretch	791	gas	EM	5,8
	6a	Ring deform.	497.5	gas	EM	5,8,10
			495	Ar	LF	13
a_2	10a	CD deform.	750	gas	EM	8
	16a	Ring deform.	305	gas	EM	8
b_1	16b	Ring deform.	376	gas	EM	8
	8b	Ring stretch	1495	gas	EM	5,8
			1490	Ar	LF	13
	15	CD deform.	848	gas	EM	5,8
	6b	Ring deform.	589.1	gas	EM	5,8,10
			588	Ar	LF	13
b_2	18b	CD deform.	305	gas	EM	5,8
			303	Ar	LF	13

^aIn many of the references concerned with this molecule, the x axis is chosen in the molecular plane, resulting in an interchange of the B_1 and B_2 representations. However, the international convention established in 1955 is followed in the symmetry designations given here. In all of the references, the vibrational numbering follows that introduced by Wilson (E. B. Wilson, Jr., Phys. Rev. **45**, 706 (1934)). In view of this unanimity, the common vibrational numbering is retained.

^bFrom analysis of band at 22326.368(2).

^cData presented in Ref. 22 suggest that this level is mixed with a nearby a_1 level of the \tilde{B}^2B_1 state.

^dFrom analysis of band at 22432.395(2).

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C₈H

2Π		C_{∞v}		2Π-X 515-631 nm			
<i>T₀=15848(5)</i>		Ne	AB ¹				
Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.	sym.
Σ ⁺	3		2080(5)	Ne	AB	1	
	5		1607(5)	Ne	AB	1	
	6		783(5)	Ne	AB	1	

X̄ 2Π **C_{∞v}**
A_{eff}=-19.33 MW²
B=0.0196 MW²

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C₈H₂⁺

Č 2Π_u		D_{∞h}		Č-T₀-31300(200) 400T			
Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.	sym.
			400T	gas	PE	1	

Č̄ 2Π_g		D_{∞h}		Č̄-T₀-24600(200)			
Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.	sym.
Σ ⁺ _g		1960(80)	gas	PE	1		
		1380(80)	gas	PE	1		

Ā̄ 2Π_u **D_{∞h}**
T₀=14160(10) gas PE¹EF²
13975 Ne LF³AB⁴

Ā̄-X̄ 625-845 nm

Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.	sym.
Σ ⁺ _g	3	C≡C stretch	1950(80)	gas	PE	1	
			2029.5	Ne	LF,AB	3,4	
	5	C-C stretch	463	Ne	LF,AB	3,4	
τ ₀ ≤6 ns		gas	EF ²				
X̄ 2Π _g		D _{∞h}					
Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.	sym.
Σ ⁺ _g	2	C≡C stretch	2100(80)	gas	PE	1	
			2141.0	Ne	LF	3	
	3	C≡C stretch	2040.5	Ne	LF	3	
Π	5	C-C stretch	472.2	Ne	LF	3	
	17	Deform.	88.5HT	Ne	LF	3	

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m-(CH₂)₂C₆H₄

Ā̄ 1B₂ **C_{2v}**
T₀=7520U gas PE¹

Ā̄ 1A₁ **C_{2v}**
T₀=3350(65) gas PE¹

Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.	sym.
a ₁			1260(40)	gas	PE	1	
			1000(40)	gas	PE	1	
			265(20)	gas	PE	1	

X̄ 3B₂ **C_{2v}**

Vib.	No.	Approximate	cm ⁻¹	Med.	Type	Refs.	sym.
a ₁	CH ₂ scissors	1500(40)	gas	PE	1		
	Ring deform.	540(20)	gas	PE	1		
	Ring=C bend	290(20)	gas	PE	1		

Reference

- ¹P. G. Wenthold, J. B. Kim, and W. C. Lineberger, J. Am. Chem. Soc. **119**, 1354 (1997).

p-(CH₂)₂C₆H₄ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			3102m	Ar	IR	1,2
			3059m	Ar	IR	1,2
			3047m	Ar	IR	1
			3038m	Ar	IR	1
			3010w	Ar	IR	1
			1603m	Ar	IR	1,2
			1449m	Ar	IR	1,2
			1342m	Ar	IR	1,2
			1136w	Ar	IR	1,2
			966w	Ar	IR	1
			940w	Ar	IR	1,2
			868s	Ar	IR	1,2
			821m	Ar	IR	1
			780m	Ar	IR	1,2
			468s	Ar	IR	1,2

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m-(CH₂)₂C₆H₄⁻

Threshold for electron detachment from ground-state *m*-(CH₂)₂C₆H₄⁻ = 7410(65) gas PE¹

 $\tilde{X} \ ^2B_1$ C_{ν}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			275(20)	gas	PE	1

Reference

¹P. G. Wentholt, J. B. Kim, and W. C. Lineberger, *J. Am. Chem. Soc.* **119**, 1354 (1997).

C₉

$^1\Sigma_u^+$ D_{∞h}
 $T_0=33895(25)$ Ne AB⁹ $^1\Sigma_u^+ - \tilde{X}$ 292–295 nm
32480^a Ar AB²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			306(30)	Ne	AB	9
$T_0=26777(15)$ T	Ne	AB ⁹			324–374 nm	
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2535(20)	Ne	AB	9
	3		1041(20)	Ne	AB	9
	4		443(20)	Ne	AB	9

 $\tilde{X} \ ^1\Sigma_g^+$ D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	3	Sym. stretch	1258(50)	gas	PE	6
	4	Sym. stretch	484(48)	gas	PE	6
			448T	Ar	IR	12
Σ_u^+	5	Asym. stretch	2079.67	gas	DL	11
			2081.1	Ne	IR	12,13
			2078.1	Ar	IR	12
			2079.0T	Kr	IR	12
6		Asym. stretch	2014.28	gas	DL	3,7
			2010.0	Ne	IR	12,13
			1998.0 ^a	Ar	IR	1,2,5,12
			1994.2	Kr	IR	12
			2007.3	H ₂	IR	10
7		Asym. stretch	1602.8	Ne	IR	12
			1601.0	Ar	IR	8,12
			1600.8	Kr	IR	12

$B_0=0.0143$ DL^{3,7,11}

^aPreviously assigned to C₈. Reassignment to C₉ dictated by arguments presented in Ref. 4.

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- P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, *Chem. Phys.* **216**, 401 (1997).

C₉⁻

Threshold for electron detachment from ground-state C₉⁻ = 29720(80) gas PE^{1,2}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
$\tilde{D} \ ^2\Pi_g$				D _{∞h}		
$T_0=34536(24)$	Ne	AB ⁶				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	3		983(25)	Ne	AB	6
	4		474(24)	Ne	AB	6

\tilde{C} $^2\Pi_g$
 $T_0=29446(17)$ Ne AB⁶

$\tilde{C}-\tilde{X}$ 295–340 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2177(19)	Ne	AB	6
	2		1666(19)	Ne	AB	6
	4		435(17)	Ne	AB	6

\tilde{B} $^2\Pi_g$
 $T_0=16470(10)$ gas MPD⁵
 $16468(5)$ Ne AB⁶

$\tilde{B}-\tilde{X}$ 530–608 nm
 $\tilde{B}-\tilde{X}$ 528–608 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2001(6)	Ne	AB	6
\tilde{A} $^2\Pi_g$		D _{∞h}				
$T_0=13020(10)$ gas MPD ⁵ $13082(3)$ Ne AB ⁶						

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+			1200(40)	gas	MPD	5

\tilde{X} $^2\Pi_u$
D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+	6	Asym. stretch	1692.6	Ne	IR	4
			1686.7	Ar	IR	3
	7	Asym. stretch	1583.3T	Ar	IR	3

References

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- D. Forney, M. Grutter, P. Freivogel, and J. P. Maier, J. Phys. Chem. A **101**, 5292 (1997).

C₉H

\tilde{X} $^2\Pi_{1/2}$
 $B_0=0.0138$ MW¹

Reference

- M. C. McCarthy, M. J. Travers, P. Kalmus, C. A. Gottlieb, and P. Thaddeus, Astrophys. J. **467**, L125 (1996).

C₉H₂⁺

\tilde{A} $^2\Pi_g$
 $T_0=14387(5)$ T Ne AB¹

$\tilde{A}-\tilde{X}$ 609–695 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+		CC stretch	2005(5) T	Ne	AB	1

Reference

- P. Freivogel, J. Fulara, D. Lessen, D. Forney, and J. P. Maier, Chem. Phys. **189**, 335 (1994).

HC₉H

$^3\Sigma_u^-$
 $T_0=17218(5)$ Ne AB¹
17160(5)

$^3\Sigma_u^- - \tilde{X}$ 433–583 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	3	C≡C s-stretch	1969(5) 770(5)	Ne Ne	AB AB	1 1

DC₉D

$^3\Sigma_u^-$
 $T_0=17292(5)$ Ne AB¹
17236(5)

$^3\Sigma_u^- - \tilde{X}$ 432–581 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	3	C≡C s-stretch	1966(5)	Ne	AB	1

Reference

- J. Fulara, P. Freivogel, D. Forney, and J. P. Maier, J. Chem. Phys. **103**, 8805 (1995).

C₁₀

\tilde{X} $^3\Sigma_g^-$
D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+			2074.5 1915.4	Ne Ne	IR IR	1 1

Reference

- P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, Chem. Phys. **216**, 401 (1997).

C₁₀⁻

(2) $^2\Pi_g$		D _{∞h}	447–736 nm		
$T_0=13596(5)$		Ne AB ^{1,3}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Σ_g^+	1		2022(7)	Ne	AB 1,3
	2		1913(7)	Ne	AB 3
	5		400(6)	Ne	AB 1,3

$\tilde{C} \ ^2\Pi_g$		D _{∞h}	$\tilde{C}-\tilde{X}$ 804–968 nm		
$T_0=10338(5)$		Ne AB ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Σ_g^+	2		2092(5)	Ne	AB 1
	3		1992(5)	Ne	AB 1
	4		488(5)	Ne	AB 1
	5		372(5)	Ne	AB 1

$\tilde{B} \ ^2\Sigma_g^+$		D _{∞h}	$\tilde{B}-\tilde{X}$ 8964(2) Ne AB ³		
$\tilde{X} \ ^2\Pi_u$		D _{∞h}			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Σ_u^+			2094.5	Ne	IR 2

References

- ¹P. Freivogel, J. Fulara, M. Jakobi, D. Forney, and J. P. Maier, *J. Chem. Phys.* **103**, 54 (1995).
²P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, *Chem. Phys.* **216**, 401 (1997).
³P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, *J. Chem. Phys.* **107**, 4468 (1997).

C₁₀H

$^2\Pi$		C _{∞v}	$^2\Pi-\tilde{X}$ 627–722 nm		
$T_0=13852(5)$		Ne AB ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Σ^+	3		2097(5)	Ne	AB 1
	5		1520(5)T	Ne	AB 1

Reference

- ¹P. Freivogel, J. Fulara, M. Jakobi, D. Forney, and J. P. Maier, *J. Chem. Phys.* **103**, 54 (1995).

C₁₀H₂⁺

$\tilde{A} \ ^2\Pi_g$		D _{∞h}	$\tilde{A}-\tilde{X}$ 703–824 nm		
$T_0=12143(5)$		Ne AB ¹			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Σ_g^+		C=C stretch	2072(5)	Ne	AB 1
		C-C stretch	396(5)	Ne	AB 1

Reference

- ¹P. Freivogel, J. Fulara, D. Lessen, D. Forney, and J. P. Maier, *Chem. Phys.* **189**, 335 (1994).

C₁₁

$^1\Sigma_u^+$		D _{∞h}	$^1\Sigma_u^+-\tilde{X}$ 323–337 nm		
$T_0=29732(20)$		Ne AB ²			
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Σ_g^+			1166(30)	Ne	AB 2
			872(30)	Ne	AB 2
			528(30)	Ne	AB 2
			240(30)	Ne	AB 2

$T_0=22245(10)T$ Ne AB² 399–450 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2440(15)	Ne	AB 2	
	2		2298(15)	Ne	AB 2	
	5		357(15)	Ne	AB 2	

 $\tilde{X} \ ^1\Sigma_g^+$

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	7	Sym. stretch	440T	gas	PE 1	
Σ_u^+	7	Asym. stretch	1938.6	Ne	IR 3	
	8	Asym. stretch	1853.4	Ne	IR 3	

References

- ¹D. W. Arnold, S. E. Bradforth, T. N. Kitsopoulos, and D. M. Neumark, *J. Chem. Phys.* **95**, 8753 (1991).
²D. Forney, P. Freivogel, M. Grutter, and J. P. Maier, *J. Chem. Phys.* **104**, 4954 (1996).
³P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, *Chem. Phys.* **216**, 401 (1997).

C₁₁H

$\tilde{X} \ ^2\Pi$ D_{∞h}
 $A_0=25T; B_0=0.0076$ MW¹

Reference

- ¹M. C. McCarthy, M. J. Travers, P. Kalmus, C. A. Gottlieb, and P. Thaddeus, *Chem. Phys. Lett.* **264**, 252 (1997).

C₁₁H₂⁺

\tilde{A} $^2\Pi_u$ D_{∞h}
 $T_0=12674(5)T$ Ne AB¹

$\tilde{A}-\tilde{X}$ 681–789 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+		CC stretch	1993(5)T	Ne	AB	1

Reference

¹P. Freivogel, J. Fulara, D. Lessen, D. Forney, and J. P. Maier, Chem. Phys. **189**, 335 (1994).

HC₁₁H

$^3\Sigma_u^-$ D_{∞h}
 $T_0=15302(5)$ Ne AB¹
 15268(5)
 15251(5)

$^3\Sigma_u^- - \tilde{X}$ 519–656 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+		C≡C s-stretch	1973(5)	Ne	AB	1
		C≡C s-stretch	1885(5)	Ne	AB	1

DC₁₁D

$^3\Sigma_u^-$ D_{∞h}
 $T_0=15294(5)$ Ne AB¹

Reference

¹J. Fulara, P. Freivogel, D. Forney, and J. P. Maier, J. Chem. Phys. **103**, 8805 (1995).

C₁₂

\tilde{X} $^3\Sigma_g^-$ D_{∞h}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+			2003.9T	Ne	IR	1

Reference

¹P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, Chem. Phys. **216**, 401 (1997).

C₁₂⁻

(2) $^2\Pi_u$ D_{∞h}
 $T_0=11636(5)$ Ne AB^{1,3}

642–860 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2014(6)	Ne	AB	3
	2		1923(6)	Ne	AB	1,3
	4		1463(5)	Ne	AB	3
	6		320(4)	Ne	AB	1,3

\tilde{C} $^2\Pi_u$
 $T_0=8006(5)$ Ne AB¹

$\tilde{C}-\tilde{X}$ 1200–1250 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	6		323(5)	Ne	AB	1
\tilde{X} $^2\Pi_g$		D _{∞h}				

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+			2012.6	Ne	IR	2
			1819.3	Ne	IR	2

References

¹P. Freivogel, J. Fulara, M. Jakobi, D. Forney, and J. P. Maier, J. Chem. Phys. **103**, 54 (1995).

²P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, Chem. Phys. **216**, 401 (1997).

³P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, J. Chem. Phys. **107**, 4468 (1997).

C₁₂H

$^2\Pi$ C_{∞v}
 $T_0=12492(5)$ Ne AB¹

$^2\Pi-\tilde{X}$ 685–801 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ^+	3		2089(5)	Ne	AB	1
	5		1395(5)T	Ne	AB	1

Reference

¹P. Freivogel, J. Fulara, M. Jakobi, D. Forney, and J. P. Maier, J. Chem. Phys. **103**, 54 (1995).

C₁₂H₂⁺

\tilde{A} $^2\Pi_u$ D_{∞h}
 $T_0=10703(5)$ Ne AB¹

$\tilde{A}-\tilde{X}$ 777–935 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+		C≡C stretch	2154(5)	Ne	AB	1
		C-C stretch	322(5)	Ne	AB	1

Reference

¹P. Freivogel, J. Fulara, D. Lessen, D. Forney, and J. P. Maier, Chem. Phys. **189**, 335 (1994).

C₁₂⁻

(2) $^2\Pi_u$ D_{∞h}
 $T_0=11636(5)$ Ne AB^{1,3}

642–860 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2014(6)	Ne	AB	3
	2		1923(6)	Ne	AB	1,3
	4		1463(5)	Ne	AB	3
	6		320(4)	Ne	AB	1,3

C₁₃

$^1\Sigma_u^+$ D_{∞h}
 $T_0=26341(15)$ Ne AB² $^1\Sigma_u^+ - \tilde{X}$ 370–380 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			669(20)	Ne	AB	2
			424(20)	Ne	AB	2
			207(20)	Ne	AB	2

$T_0=18761(7)$ T Ne AB²

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1		2409(10)	Ne	AB	2
	2		2258(10)	Ne	AB	2
	6		307(10)	Ne	AB	2

\tilde{X} $^1\Sigma_g^+$ D_{∞h} Structure: DL¹

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_u^+			1808.96	gas	DL	1

$B_0=0.0047$ DL¹

References

- ¹T. F. Giesen, A. Van Orden, H. J. Hwang, R. S. Fellers, R. A. Provençal, and R. J. Saykally, *Science* **265**, 756 (1994).
²D. Forney, P. Freivogel, M. Grutter, and J. P. Maier, *J. Chem. Phys.* **104**, 4954 (1996).

C₁₃H₂⁺

\tilde{A}	$^2\Pi_g$	D _{∞h}	$T_0=11418(5)$ T	Ne	AB ¹	$\tilde{A}-\tilde{X}$ 746–874 nm
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+		CC stretch	1924(5)T	Ne	AB	1

Reference

- ¹P. Freivogel, J. Fulara, D. Lessen, D. Forney, and J. P. Maier, *Chem. Phys.* **189**, 335 (1994).

HC₁₃H

$^3\Sigma_u^-$	D _{∞h}	$T_0=13901(5)$	Ne	AB ¹	$^3\Sigma_u^- - \tilde{X}$ 562–722 nm
					13866(5)
					13852(5)
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Σ_g^+		C≡C s-stretch	1967(5)	Ne	AB
		C≡C s-stretch	1904(5)	Ne	AB
			619(5)	Ne	AB

Reference

- ¹J. Fulara, P. Freivogel, D. Forney, and J. P. Maier, *J. Chem. Phys.* **103**, 8805 (1995).

C₁₄⁻

(2) $^2\Pi_g$ D_{∞h}
 $T_0=10202(2)$ Ne AB^{1,2} 723–981 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	1			1810(4)	Ne	AB

\tilde{C} $^2\Pi_g$ D_{∞h}
 $T_0=6849(5)$ Ne AB¹ $\tilde{C}-\tilde{X}$ 1402–1460 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	7		283(2)	Ne	AB	1

References

- ¹P. Freivogel, J. Fulara, M. Jakobi, D. Forney, and J. P. Maier, *J. Chem. Phys.* **103**, 54 (1995).
²P. Freivogel, M. Grutter, D. Forney, and J. P. Maier, *J. Chem. Phys.* **107**, 4468 (1997).

C₁₄H

$^2\Pi$ C_{∞v}
 $T_0=11554(5)$ T Ne AB¹

Reference

- ¹P. Freivogel, J. Fulara, M. Jakobi, D. Forney, and J. P. Maier, *J. Chem. Phys.* **103**, 54 (1995).

C₁₄H₂⁺

\tilde{A} $^2\Pi_g$ D_{∞h}
 $T_0=9548(5)$ Ne AB¹ $\tilde{A}-\tilde{X}$ 858–1048 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+		C≡C stretch	2097(5)	Ne	AB	1
		C-C stretch	258(5)	Ne	AB	1

Reference

- ¹P. Freivogel, J. Fulara, D. Lessen, D. Forney, and J. P. Maier, *Chem. Phys.* **189**, 335 (1994).

HC₁₃H

$^3\Sigma_u^-$	D _{∞h}	$T_0=13901(5)$	Ne	AB ¹	$^3\Sigma_u^- - \tilde{X}$ 562–722 nm
					13866(5)
					13852(5)
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
Σ_g^+		C≡C s-stretch	1967(5)	Ne	AB
		C≡C s-stretch	1904(5)	Ne	AB
			619(5)	Ne	AB

C₁₅

$^1\Sigma_u^+$ D_{∞h}
 $T_0=23832(12)$ Ne AB¹

$^1\Sigma_u^+ - \tilde{X}$ 410–420 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			277(20)	Ne	AB	1
$T_0=16090(5)$ T	Ne	AB ¹			611–622 nm	
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	7		266(7)	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, M. Grutter, and J. P. Maier, *J. Chem. Phys.* **104**, 4954 (1996).

C₁₆⁻

$^2\Pi_g$ D_{∞h}
 $T_0=5784(5)$ Ne AB¹

$^2\Pi_g - \tilde{X}$ 1658–1730 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
Σ_g^+	8		245(5)	Ne	AB	1

Reference

¹P. Freivogel, J. Fulara, M. Jakobi, D. Forney, and J. P. Maier, *J. Chem. Phys.* **103**, 54 (1995).

8.16. Non-Hydrocarbons with More Than Eight Atoms

(CH₃)₂Zn⁺

$T_0=32220$ T gas EF²

633–635 nm

A

$T_0=16440$ gas PE¹EF²

$\tilde{A} - \tilde{X}$ 594–669 nm

X

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH ₃ s-deform.	1086	gas	EF	2

References

¹G. M. Bancroft, D. K. Creber, M. A. Ratner, J. W. Moskowitz, and S. Topiol, *Chem. Phys. Lett.* **50**, 233 (1977).

²F. J. Grieman, E. S. Fujikawa, and J. C. Phillips, *J. Phys. Chem.* **100**, 9276 (1996).

C₂H₅HgH**X**

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		HgH stretch	1954.8	Ar	IR	1
			1946.8			
			1926.8			
		CH ₂ rock	698.4	Ar	IR	1

Reference

¹T. M. Greene, L. Andrews, and A. J. Downs, *J. Am. Chem. Soc.* **117**, 8180 (1995).

cyc-C₅Cl₄

In a nitrogen matrix, structured absorption between 304 and 345 nm has been assigned¹ to cyc-C₅Cl₄.

X

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1512w	N ₂	IR	1
			1508w	N ₂	IR	1
			1356w	N ₂	IR	1
			1138w	N ₂	IR	1
			1127m	N ₂	IR	1
			892w	N ₂	IR	1
			882w	N ₂	IR	1
			703w	N ₂	IR	1
			550w	N ₂	IR	1
			530w	N ₂	IR	1

Reference

¹G. A. Bell and I. R. Dunkin, *J. Chem. Soc. Faraday Trans. 2* **81**, 725 (1985).

cyc-C₅Br₄

In an argon matrix, absorption maxima at 21790 (459 nm), 24330 (411 nm), and 25710 (389 nm) have been attributed¹ to cyc-C₅Br₄.

X

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1182	Ar	IR	1
			1170	Ar	IR	1

Reference

¹I. R. Dunkin and A. McCluskey, *Spectrochim. Acta A* **49**, 1179 (1993).

cyc-C₂H₂Si(CH₃)₂

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH stretch	3023.1wm	Ar	IR	1
		CH stretch	2992.5wm	Ar	IR	1
		CH stretch	2959.1wm	Ar	IR	1
		C=C stretch	1447.1wm	Ar	IR	1
		CH ₃ deform.	1444.2wm	Ar	IR	1
		CH ₃ deform.	1438.1wm	Ar	IR	1
		CH ₃ deform.	1400.2w	Ar	IR	1
			1250.2m	Ar	IR	1
			1216.1w	Ar	IR	1
			1132.0wm	Ar	IR	1
			1129.8wm	Ar	IR	1
			1127.7wm	Ar	IR	1
			991.6wm	Ar	IR	1
			922.1wm	Ar	IR	1
			884.1s	Ar	IR	1
			788.4vs	Ar	IR	1
			784.9m	Ar	IR	1
		SiC s-stretch	694.1wm	Ar	IR	1
		SiC a-stretch	691.1wm	Ar	IR	1
			616.1m	Ar	IR	1
			613.7m	Ar	IR	1

Reference

¹M. Trommer, W. Sander, and C. Marquard, Angew. Chem. **106**, 816 (1994); Angew. Chem. Int. Ed. Engl. **33**, 766 (1994).

cyc-C₄H₄SiH₂

In an argon matrix, an absorption maximum has been observed¹ at 36000 (278 nm).

\tilde{X}	C _{2v}				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.
<i>a</i> ₁		SiH ₂ s-stretch	2144m	Ar	IR
		C=C s-stretch	1478w	Ar	IR
		CH deform.	1346m	Ar	IR
		CH deform.	1088w	Ar	IR
		SiH ₂ scissors	955s	Ar	IR
		Ring deform.	490m	Ar	IR
<i>b</i> ₁		OPLA CH bend	713vs	Ar	IR
		CH stretch	3038w	Ar	IR
<i>b</i> ₂		CH stretch	3001vw	Ar	IR
		SiH ₂ a-stretch	2174s	Ar	IR
		C=C a-stretch	1608m	Ar	IR
		CH deform.	1285w	Ar	IR
		SiH ₂ wag	880vs	Ar	IR
		SiC ₂ a-stretch	677w	Ar	IR

Reference

¹V. N. Khabashesku, V. Balaji, S. E. Boganov, O. M. Nefedov, and J. Michl, J. Am. Chem. Soc. **116**, 7863 (1994).

cyc-C₄H₆Si:

In an argon matrix, absorption maxima have been observed¹ at 20800 (480 nm) and at 40000 (250 nm).

\tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁		=CH stretch	3028vw	Ar	IR	1
		CH ₂ stretch	2876vw	Ar	IR	1
		C=C stretch	1624s	Ar	IR	1
		CH ₂ scissors	1406m	Ar	IR	1
		=CH s-rock	1208w	Ar	IR	1
		CH ₂ rock	1108s	Ar	IR	1
		Ring deform.	881m	Ar	IR	1
		SiC ₂ s-stretch	616m	Ar	IR	1
		Ring deform.	463w	Ar	IR	1
<i>b</i> ₁		CH ₂ stretch	2962vw	Ar	IR	1
		=CH wag	770m	Ar	IR	1
<i>b</i> ₂		=CH stretch	3002vw	Ar	IR	1
		CH ₂ stretch	2854vw	Ar	IR	1
		=CH a-rock	1397s	Ar	IR	1
		CH ₂ wag	1307w	Ar	IR	1
		CH ₂ twist	1174w	Ar	IR	1
		Ring deform.	944w	Ar	IR	1
		SiC ₂ a-stretch	741s	Ar	IR	1
		Ring deform.	528w	Ar	IR	1

Reference

¹V. N. Khabashesku, V. Balaji, S. E. Boganov, O. M. Nefedov, and J. Michl, J. Am. Chem. Soc. **116**, 7863 (1994).

cyc-C₄H₅SiH

(1-Silacyclopenta-1,3-diene)

In an argon matrix, an absorption maximum has been observed¹ at 33800 (296 nm).

\tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>		=CH stretch	3066m	Ar	IR	1
		=CH stretch	3038s	Ar	IR	1
		=CH stretch	2994w	Ar	IR	1
		CH ₂ stretch	2939m	Ar	IR	1
		CH ₂ stretch	2865m	Ar	IR	1
		SiH stretch	2210m	Ar	IR	1
		C=C stretch	1551m	Ar	IR	1
		CH ₂ scissors	1406m	Ar	IR	1
		=CH bend	1354m	Ar	IR	1
		Mixed	1282w	Ar	IR	1
		=CH bend	1181w	Ar	IR	1
		CH ₂ wag	1092s	Ar	IR	1
		=CH bend	961m	Ar	IR	1
		Si=C stretch	929m	Ar	IR	1
		C-C ring	852w	Ar	IR	1
		C-C ring	819w	Ar	IR	1
		SiH deform.	680w	Ar	IR	1
		Ring deform.	488m	Ar	IR	1
<i>a''</i>		=CH bend	974m	Ar	IR	1
		=CH bend	790w	Ar	IR	1
		CH ₂ rock	736m	Ar	IR	1
		=SiH OPLA	637m	Ar	IR	1
		Ring deform.	466w	Ar	IR	1

Reference

¹V. N. Khabashesku, V. Balaji, S. E. Boganov, O. M. Nefedov, and J. Michl, J. Am. Chem. Soc. **116**, 7863 (1994).

cyc-C₄H₅SiH

(1-Silacyclopenta-1,4-diene)

In an argon matrix, an absorption maximum has been observed¹ at 37000 (270 nm).

\tilde{X}	C_s					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a'		=CH stretch	3095w	Ar	IR	1
		=CH stretch	3061w	Ar	IR	1
		=CH stretch	3025w	Ar	IR	1
		SiH stretch	2216m	Ar	IR	1
		C=C stretch	1543m	Ar	IR	1
		CH ₂ scissors	1426w	Ar	IR	1
		=CH bend	1278w	Ar	IR	1
		=CH bend	1149w	Ar	IR	1
		=CH bend	1103m	Ar	IR	1
		=CH bend	985w	Ar	IR	1
		Si=C stretch	936s	Ar	IR	1
		CH ₂ rock	859s	Ar	IR	1
		Ring deform.	470w	Ar	IR	1
a''		CH ₂ stretch	2904w	Ar	IR	1
		CH ₂ wag	1135w	Ar	IR	1
		=CH bend	913m	Ar	IR	1
		=CH OPLA	742s	Ar	IR	1
		=SiH OPLA	597m	Ar	IR	1

Reference

¹V. N. Khabashesku, V. Balaji, S. E. Boganov, O. M. Nefedov, and J. Michl, J. Am. Chem. Soc. **116**, 7863 (1994).

cyc-C₄H₄GeH₂

In an argon matrix, unstructured absorption has been observed¹ between 260 and 320 nm, with maximum near 36760 (272 nm).

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	2	=CH s-stretch	3015	Ar	IR	1
	3	GeH ₂ s-stretch	2099	Ar	IR	1
	4	C=C s-stretch	1532	Ar	IR	1
	5	CH bend	1321	Ar	IR	1
	6	CH scissor	1064	Ar	IR	1
	8	GeH ₂ scissor	906	Ar	IR	1
b_1	15	GeH ₂ a-stretch	2096	Ar	IR	1
	16	=CH twist	1031	Ar	IR	1
	17	=CH wag	722	Ar	IR	1
	18	Mixed	505	Ar	IR	1
b_2	22	C=C a-stretch	1593	Ar	IR	1
	23	=CH bend	1285	Ar	IR	1
	24	=CH scissor	1168	Ar	IR	1
	25	GeCH rock	783	Ar	IR	1
	26	GeH ₂ wag	682	Ar	IR	1

cyc-C₄D₄GeD₂ \tilde{X} C_{2v}

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	=CD s-stretch	2262	Ar	IR	1
	2	=CD a-stretch	2229	Ar	IR	1
	4	GeD ₂ s-stretch	1522	Ar	IR	1
	5	Mixed	1168	Ar	IR	1
	9	GeD ₂ scissor	641	Ar	IR	1
b_1	15	GeD ₂ a-stretch	1507	Ar	IR	1
	17	=CD wag	594	Ar	IR	1
b_2	24	DC-CD rock	878	Ar	IR	1
	25	DCGeCD scissor	707	Ar	IR	1
	27	GeD ₂ wag	486	Ar	IR	1

Reference

¹V. N. Khabashesku, S. E. Boganov, D. Antic, O. M. Nefedov, and J. Michl, Organometallics **15**, 4714 (1996).

cyc-C₄H₆Ge:

In an argon matrix, an absorption maximum at 40320 (248 nm) and absorption between 370 and 460 nm have been assigned¹ to this species.

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	=CH s-stretch	3017	Ar	IR	1
	3	C=C stretch	1615	Ar	IR	1
	5	=CH s-rock	1186	Ar	IR	1
	6	CH ₂ rock	1078	Ar	IR	1
	8	CGeC s-stretch	508	Ar	IR	1
b_1	15	CH ₂ a-stretch	2908	Ar	IR	1
	17	=CH wag	732	Ar	IR	1
b_2	21	CH ₂ s-stretch	2865	Ar	IR	1
	22	CH ₂ bend	1389	Ar	IR	1
	23	=CH rock	1346	Ar	IR	1
	25	C—C a-stretch	949	Ar	IR	1
	27	CGeC a-stretch	478	Ar	IR	1

cyc-C₄D₆Ge:

\tilde{X}	C_{2v}					
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1	1	=CD s-stretch	2256	Ar	IR	1
	3	C=C stretch	1579	Ar	IR	1
	4	C—C s-stretch	1072	Ar	IR	1
	6	=CD rock	804	Ar	IR	1
	7	CD ₂ wag	748	Ar	IR	1
b_1	15	CD ₂ a-stretch	2171	Ar	IR	1
	16	CD ₂ twist	837	Ar	IR	1
	17	DC=CD wag	551	Ar	IR	1
b_2	21	CD ₂ s-stretch	2077	Ar	IR	1
	22	C—C a-stretch	1164	Ar	IR	1
	23	CD ₂ bend	1024	Ar	IR	1
	24	Mixed	976	Ar	IR	1
	25	=CD a-rock	772	Ar	IR	1
	27	CGeC a-stretch	438	Ar	IR	1

Reference

¹V. N. Khabashesku, S. E. Boganov, D. Antic, O. M. Nefedov, and J. Michl, Organometallics **15**, 4714 (1996).

cyc-C₄H₅GeH

(1-Germacyclopenta-1,3-diene)

In an argon matrix, an absorption maximum at 33220 (301 nm) has been tentatively assigned¹ to this product.

\tilde{X}		C_s				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a'</i>	2	=CH s-stretch	3048	Ar	IR	1
	4	CH ₂ s-stretch	2894	Ar	IR	1
	5	GeH stretch	2115	Ar	IR	1
	6	C=C stretch	1574	Ar	IR	1
	8	=CH rock	1352	Ar	IR	1
	9	Mixed	1262	Ar	IR	1
	12	Mixed	959	Ar	IR	1
	13	Mixed	922	Ar	IR	1
	14	Ge=CC s-stretch	838	Ar	IR	1
	16	GeH rock	631	Ar	IR	1
<i>a''</i>	19	CH ₂ a-stretch	2972	Ar	IR	1
	22	GeCH rock	778	Ar	IR	1
	24	Mixed	698	Ar	IR	1

cyc-C₄D₅GeD

\tilde{X}		C_s				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a'</i>	3	=CD a-stretch	2232	Ar	IR	1
	4	CD ₂ s-stretch	2090	Ar	IR	1
	6	GeD stretch	1536	Ar	IR	1
	12	DC=CD rock	820	Ar	IR	1
	14	Ge=CC s-stretch	691	Ar	IR	1
	15	C—C=C bend	663	Ar	IR	1
	16	Ge—C stretch	529	Ar	IR	1
	17	GeD rock	454	Ar	IR	1
	22	C—Ge=C twist	621	Ar	IR	1
	24	DC=C—CD wag	465	Ar	IR	1
<i>a''</i>	25	DC=CD wag	416	Ar	IR	1

Reference

¹V. N. Khabashesku, S. E. Boganov, D. Antic, O. M. Nefedov, and J. Michl, Organometallics **15**, 4714 (1996).

cyc-C₄H₅GeH

(1-Germacyclopenta-1,4-diene)

In an argon matrix, an absorption maximum at 38170 (262 nm) has been tentatively assigned¹ to this product.

\tilde{X}		C_s				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a'</i>	5	GeH stretch	2127	Ar	IR	1
	6	C=C stretch	1553	Ar	IR	1
	13	Ge=C s-stretch	834	Ar	IR	1
	14	Mixed	787	Ar	IR	1
<i>a''</i>	23	Ge=CH wag	682	Ar	IR	1
	24	CH ₂ rock	635	Ar	IR	1

cyc-C₄D₅GeD

\tilde{X}		C_s				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a'</i>	4	CD ₂ s-stretch	2068	Ar	IR	1
	6	GeD stretch	1547	Ar	IR	1
	10	CD ₂ bend	1072	Ar	IR	1
	14	Skel. stretch	686	Ar	IR	1
<i>a''</i>	24	Wag	509	Ar	IR	1

Reference

¹V. N. Khabashesku, S. E. Boganov, D. Antic, O. M. Nefedov, and J. Michl, Organometallics **15**, 4714 (1996).

CaC₅H₅

\tilde{B} 2A_1		C_{5v}				
$T_0=16772(5)T$	gas	$LF^{1,3}$	$\tilde{B}-\tilde{X}$ 585–607 nm			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁		Ca-C ₅ stretch	300(5)T	gas	LF	1
\tilde{A} 2E_1						Structure: LF ⁴
$T_0=14535.44$	gas	LF^{1-3}	$\tilde{A}-\tilde{X}$ 645–745 nm			
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CH wag	738	gas	LF	3
	4	Ca-C ₅ stretch	330.19	gas	LF	1–4
<i>e</i> ₁	10	Ca-C ₅ bend	222H	gas	LF	3
<i>e</i> ₂	13	CH deform.	1039T	gas	LF	3
	15	CH wag	869T	gas	LF	3

$A=57.66$ gas LF^{1-4}
 $B_0=0.091$ LF^4

 \tilde{X} 2A_1 C_{5v} Structure: LF⁴

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
<i>a</i> ₁	4	Ca-C ₅ stretch	311	gas	LF	1,3
			$B_0=0.088$ LF^4			

CaC₅D₅

\tilde{A}^2E_1 C_{5v}
 $T_0=14536$ gas LF³ $\tilde{A}-\tilde{X}$ 645–690 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	3	CD wag	512	gas	LF	3
	4	Ca-C ₅ stretch	316	gas	LF	3
<i>e</i> ₁	10	Ca-C ₅ bend	221H	gas	LF	3
<i>e</i> ₂	13	CD deform.	727T	gas	LF	3
	15	CD wag	614T	gas	LF	3

A=56 gas LF³

References

- L. C. O'Brien and P. F. Bernath, J. Am. Chem. Soc. **108**, 5017 (1986).
- A. M. Ellis, E. S. J. Robles, and T. A. Miller, J. Chem. Phys. **94**, 1752 (1991).
- E. S. J. Robles, A. M. Ellis, and T. A. Miller, J. Am. Chem. Soc. **114**, 7171 (1992).
- T. M. Cerny, J. M. Williamson, and T. A. Miller, J. Chem. Phys. **102**, 2372 (1995).

H₂BSC₂H₅ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		CH stretch	2984	Ar	IR	1
		CH stretch	2980	Ar	IR	1
		CH stretch	2951	Ar	IR	1
		CH stretch	2941	Ar	IR	1
		CH stretch	2892	Ar	IR	1
		BH ₂ a-stretch	2599	Ar	IR	1
		BH ₂ s-stretch	2522	Ar	IR	1
			1456	Ar	IR	1
			1444	Ar	IR	1
BH ₂ scissors	1184		1167	Ar	IR	1
			1145	Ar	IR	1
			1085	Ar	IR	1
			1040	Ar	IR	1
			971	Ar	IR	1
		BH ₂ OPLA	923	Ar	IR	1
		B-S stretch	829	Ar	IR	1
		BH ₂ rock	798	Ar	IR	1

Reference

- J. D. Carpenter and B. S. Ault, J. Phys. Chem. **97**, 3697 (1993).

H₂C=(cyc-CN=CCH₃) \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1865	Kr	IR	1
			1554	Kr	IR	1
			1228	Kr	IR	1
			696	Kr	IR	1

Reference

- ¹K. Banert, M. Hagedorn, E. Knözinger, A. Becker, and E.-U. Würthwein, J. Am. Chem. Soc. **116**, 60 (1994).

bicyc-C₆H₃F₂N

(7-Aza-1,5-difluoro-2,4,6-bicyclo[4.1.0]heptatriene)

In an argon matrix, an absorption maximum at 39700 (252 nm) has been assigned¹ to bicyc-C₆H₃F₂N.

In an argon matrix,¹ irradiation at 27300 (366 nm) results in partial rearrangement to the phenylnitrene, 2,6-C₆H₃F₂N::.

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C=N stretch	1679.3m	Ar	IR	1
		C=C stretch	1609.8vs	Ar	IR	1
		C=C stretch	1519.7m	Ar	IR	1
			1412.2ms	Ar	IR	1
		C-C stretch	1318.0m	Ar	IR	1
		CF stretch	1258.9s	Ar	IR	1
		CF stretch	1179.7s	Ar	IR	1
		6-ring deform.	1138.3m	Ar	IR	1
		6-ring deform.	1006.7wm	Ar	IR	1
		6-ring OPLA	939.2m	Ar	IR	1
		CH OPLA	896.1w	Ar	IR	1
		3-ring deform.	856.9wm	Ar	IR	1
			796.0m	Ar	IR	1
			737.4wm	Ar	IR	1
			688.2wm	Ar	IR	1
			608.5vw	Ar	IR	1
			557.1vw	Ar	IR	1
			535.7vw	Ar	IR	1
			500.3w	Ar	IR	1
			430.0w	Ar	IR	1

Reference

- ¹J. Morawietz and W. Sander, J. Org. Chem. **61**, 4351 (1996).

bicyc-C₆F₅N

(Perfluoro-7-aza-2,4,6-bicyclo[4.1.0]heptatriene)

In an argon matrix,¹ irradiation at 27300 (366 nm) results in reversible isomerization to C₆F₅N::.

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		1664.2wm	Ar	IR	1	
		1589.3wm	Ar	IR	1	
		1395.3m	Ar	IR	1	
		1374.5m	Ar	IR	1	
		1308.7wm	Ar	IR	1	
		1222.6wm	Ar	IR	1	
		1187.8wm	Ar	IR	1	
		1095.3wm	Ar	IR	1	
		1007.6vs	Ar	IR	1	
		947.5wm	Ar	IR	1	
		937.3wm	Ar	IR	1	
		816.0w	Ar	IR	1	
		660.6w	Ar	IR	1	
		524.8vw	Ar	IR	1	

Reference

¹J. Morawietz and W. Sander, J. Org. Chem. **61**, 4351 (1996).

 $\mathbf{H(C\equiv C)_3CN^+}$

$\tilde{A} \ ^2\Pi$		$C_{\infty v}$	$T_0=14836$	Ne	AB^1LF^2	$\tilde{A}-\tilde{X} \ 594-820 \text{ nm}$
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	5	CN stretch	1969(5)	Ne	AB	1
	6	C-C stretch	1578(5)	Ne	AB	1
	7	C-C stretch	1017(5)	Ne	AB	1
	8	C-C stretch	459(5)	Ne	AB	1

$A=19T$ Ne LF²

 $\tilde{X} \ ^2\Pi$

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	2	C≡N stretch	2180.2	Ne	LF	2
	3	C≡C stretch	2159.6	Ne	LF	2
	4	C≡C stretch	2128.5	Ne	LF	2
	5	C=C stretch	1938.3	Ne	LF	2
	6	C-C stretch	957.2	Ne	LF	2
	7	C-C stretch	881.0	Ne	LF	2
	8	C-C stretch	464.5	Ne	LF	2

$A=92T$ Ne LF²

References

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

²A. M. Smith, J. Agreiter, and V. E. Bondybey, Chem. Phys. Lett. **244**, 379 (1995).

 $\mathbf{HC_7CN^+}$

\tilde{A}
 $T_0=15217(5)$ Ne AB¹

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

 $\mathbf{NCC_5CN^+}$

\tilde{A}		$C_{\infty v}$	$T_0=15897(5)$	Ne	AB^1	$\tilde{A}-\tilde{X} \ 558-630 \text{ nm}$
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+		C≡C stretch	2012(5)	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

 $\mathbf{H(C\equiv C)_4CN^+}$

$\tilde{A} \ ^2\Pi$		$C_{\infty v}$	$T_0=12978(5)$	Ne	AB^1	$\tilde{A}-\tilde{X} \ 663-771 \text{ nm}$
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	6	C≡C stretch	2106(5)	Ne	AB	1
	8	C-C stretch	1223(5)	Ne	AB	1
	10	C-C stretch	363(5)	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

 $\mathbf{NC(C\equiv C)_3CN^+}$

$\tilde{A} \ ^2\Pi_g$		$D_{\infty h}$	$T_0=13480$	Ne	AB^1LF^2	$\tilde{A}-\tilde{X} \ 638-1090 \text{ nm}$
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	2	C≡C s-stretch	2163(5)	Ne	AB	1
	3	C≡C s-stretch	1952(5)	Ne	AB	1
	4	C-C s-stretch	1105(5)	Ne	AB	1
	5	C-C s-stretch	395(5)	Ne	AB	1
Π		Bend	423(5)H	Ne	AB	1

 $\tilde{X} \ ^2\Pi_u$

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	1	CN s-stretch	2136.8(5)	Ne	LF	2
	2	C≡C s-stretch	2127.9(5)	Ne	LF	2
	3	C≡C s-stretch	1974.4(5)	Ne	LF	2
	5	C-C s-stretch	362T	Ne	LF	2

References

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

²J. Agreiter, A. M. Smith, and V. E. Bondybey, Chem. Phys. Lett. **241**, 317 (1995).

HC_9CN^+

\tilde{A}
 $T_0=13395(5)$ Ne AB¹

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

NCC_7CN^+

\tilde{A}		$C_{\infty v}$	$\tilde{A}-\tilde{X}$ 623–714 nm			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+		C≡C stretch	2022(5)T	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

$\text{H}(\text{C}\equiv\text{C})_5\text{CN}^+$

\tilde{A} $^2\Pi$		$C_{\infty v}$	$\tilde{A}-\tilde{X}$ 734–872 nm			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ^+	5	C≡N stretch	2154(5)T	Ne	AB	1
	6	C≡C stretch	2007(5)T	Ne	AB	1
	7	C≡C stretch	1849(5)T	Ne	AB	1
	8	C-C stretch	1611(5)T	Ne	AB	1
	12	C-C stretch	323(5)	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

$\text{NC}(\text{C}\equiv\text{C})_4\text{CN}^+$

\tilde{A} $^2\Pi_u$		$D_{\infty h}$	$\tilde{A}-\tilde{X}$ 704–832 nm			
Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
Σ_g^+	3	CN sym. stretch	2168(5)T	Ne	AB	1
	4	C-C sym. stretch	1852(5)T	Ne	AB	1
	6	C-C sym. stretch	310(5)	Ne	AB	1
II		Bend	350(5)H	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

$\text{HC}_{11}\text{CN}^+$

\tilde{A}
 $T_0=12026(5)$ Ne AB¹

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

NCC_9CN^+

\tilde{A}
 $T_0=12587(5)$ Ne AB¹

$\tilde{A}-\tilde{X}$ 682–795 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.
		C≡C stretch	2058(5)T	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

$\text{H}(\text{C}\equiv\text{C})_6\text{CN}^+$

\tilde{A} $^2\Pi$
 $T_0=10276(5)$ Ne AB¹

$\tilde{A}-\tilde{X}$ 929–974 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.
Σ^+	9	C-C stretch	1795(5)T	Ne	AB	1
	10	C-C stretch	1353(5)T	Ne	AB	1
	14	C-C stretch	276(5)	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

$\text{NC}(\text{C}\equiv\text{C})_5\text{CN}^+$

\tilde{A} $^2\Pi_g$
 $T_0=10831(5)$ Ne AB¹

$\tilde{A}-\tilde{X}$ 806–924 nm

Vib.	No.	Approximate type of mode	cm^{-1}	Med.	Type	Refs.
Σ_g^+	5	C-C sym. stretch	1572(5)T	Ne	AB	1
	7	C-C sym. stretch	272(5)	Ne	AB	1

Reference

¹D. Forney, P. Freivogel, J. Fulara, and J. P. Maier, J. Chem. Phys. **102**, 1510 (1995).

H₉O₄⁺ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		H ₂ O in-phase a-stretch	3730.4	gas	PF	1,2
		H ₂ O out-of- phase s-stretch	3644.9	gas	PF	1

References

¹L. I. Yeh, M. Okumura, J. D. Myers, J. M. Price, and Y. T. Lee, *J. Chem. Phys.* **91**, 7319 (1989).

²L. I. Yeh, Y. T. Lee, and J. T. Hougen, *J. Mol. Spectrosc.* **164**, 473 (1994).

C₂H₅IO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		IO stretch	1195.4wm 717.0ms	Ar Ar	IR IR	1 1

C₂D₅IO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		IO stretch	958.1wm 717.0ms	Ar Ar	IR IR	1 1

Reference

¹R. J. H. Clark and J. R. Dann, *J. Phys. Chem.* **100**, 532 (1996).

C₂H₅OI \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		C-O stretch	1106.0wm 1076.7w 1071.9w	Ar Ar	IR IR	1 1
		CH ₂ deform.	961.9m	Ar	IR	1
		OI stretch	570.5wm	Ar	IR	1

C₂D₅OI \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Ref.
		C-O stretch	1112.9w 1075.3w	Ar Ar	IR IR	1
		CD ₃ deform.	735.8m	Ar	IR	1
		OI stretch	565.6wm	Ar	IR	1

Reference

¹R. J. H. Clark and J. R. Dann, *J. Phys. Chem.* **100**, 532 (1996).

C₈O

$\tilde{X}^{\text{3}\Sigma^-}$
 $B_0=0.013$ MW¹ C_{∞v}

Reference

¹Y. Ohshima, Y. Endo, and T. Ogata, *J. Chem. Phys.* **102**, 1493 (1995).

C₉O

\tilde{X}
 $B_0=0.010$ MW¹ C_{∞v}

Reference

¹T. Ogata, Y. Ohshima, and Y. Endo, *J. Am. Chem. Soc.* **117**, 3593 (1995).

C₂H₅IO₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		IO ₂ a-stretch	837.2wm 829.3m	Ar	IR	1
		IO ₂ s-stretch	803.1wm 795.4m	Ar	IR	1

C₂D₅IO₂ \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
		IO ₂ a-stretch	837.1wm 829.3m	Ar	IR	1
		IO ₂ s-stretch	802.8wm 794.1m	Ar	IR	1

Reference

¹R. J. H. Clark and J. R. Dann, *J. Phys. Chem.* **100**, 532 (1996).

CH₃SiHOHCHO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
OH stretch		3656.4m	Ar	IR	1	
		3645.7wm	Ar	IR	1	
		2768.4w	Ar	IR	1	
		2760.7w	Ar	IR	1	
		2664.1wm	Ar	IR	1	
		2654.3w	Ar	IR	1	
		2133.4ms	Ar	IR	1	
		1656.9m	Ar	IR	1	
		1258.2ms	Ar	IR	1	
		940.3ms	Ar	IR	1	
C=O stretch		909.4vs	Ar	IR	1	
		867.8s	Ar	IR	1	
		748.2m	Ar	IR	1	

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			3158w	N ₂	IR	1
			1478w	Ar	IR	3
			1481w	N ₂	IR	1
			1397w	Ar	IR	3
			1398	N ₂	IR	1
			1320m	Ar	IR	3
			1321	N ₂	IR	1
			1303w	N ₂	IR	1
			1221w	Ar	IR	3
			1075w	Ar	IR	3
			1063w	Ar	IR	3
			1066w	N ₂	IR	1
OO stretch			1018m	Ar	IR	3
			1014	N ₂	IR	1
			1007w	Ar	IR	3
			949m	Ar	IR	3
			947	N ₂	IR	1
			940	N ₂	IR	1
			888m	Ar	IR	3
			888	N ₂	IR	1
			754m	Ar	IR	3
			758	N ₂	IR	1
			693w	Ar	IR	3
			694w	N ₂	IR	1
			603w	Ar	IR	3
			606w	N ₂	IR	1
			498w	N ₂	IR	1

Reference

¹M. Trommer, W. Sander, and A. Patyk, J. Am. Chem. Soc. **115**, 11775 (1993).

(CH₃)₂SiOHCHO \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
OH stretch		3660.2ms	Ar	IR	1	
		3651.2m	Ar	IR	1	
		3645.9m	Ar	IR	1	
		2764.2wm	Ar	IR	1	
		2658.9wm	Ar	IR	1	
		2653.7m	Ar	IR	1	
		1655.0m	Ar	IR	1	
		1258.8vs	Ar	IR	1	
		1051.3s	Ar	IR	1	
		952.2vs	Ar	IR	1	
C=O stretch		941.6vs	Ar	IR	1	
		868.8s	Ar	IR	1	
		805.7s	Ar	IR	1	
		547.8wm	Ar	IR	1	

References

¹G. A. Bell and I. R. Dunkin, J. Chem. Soc., Chem. Commun. 1213 (1983).

²G. A. Bell, I. R. Dunkin, and C. J. Shields, Spectrochim. Acta A **41**, 1221 (1985).

³I. R. Dunkin and C. J. Shields, J. Chem. Soc., Chem. Commun. 154 (1986).

cyc-C₅Cl₄O-1-O

In argon and nitrogen matrices,^{1,2} an absorption maximum near 25300 (395 nm) is associated with photoisomerization of and O-atom photodissociation from cyc-C₅Cl₄O-1-O.

Reference

¹M. Trommer, W. Sander, and A. Patyk, J. Am. Chem. Soc. **115**, 11775 (1993).

cyc-C₅H₄O-1-O

(Cyclopentadienone O-Oxide)

In a nitrogen matrix,² an absorption maximum at 23800 (420 nm) can be assigned to cyclopentadienone O-oxide.

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
CO stretch			1555w	Ar	IR	2
			1558w	N ₂	IR	2
			1371w	Ar	IR	2
			1380w	N ₂	IR	2
			1231m	Ar	IR	2
			1230m	N ₂	IR	2
			1135w	Ar	IR	2
			1137w	N ₂	IR	2
			1016w	Ar	IR	2
			1007w			
OO stretch			1017w	N ₂	IR	2
			1005w			
			974vw	Ar	IR	2
			976w	N ₂	IR	2
			835w	Ar	IR	2
			833w	N ₂	IR	2
			748w	Ar	IR	2
			750w	N ₂	IR	2
			668w	Ar	IR	2
			669w	N ₂	IR	2
Reference			506w	Ar	IR	2
			508w	N ₂	IR	2

1,2,3-cyc-C₂H₄O₃

\tilde{X}	C _s	Structure: MW ⁴				
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CH ₂ deform.	1214m	Xe	IR	2
		CO stretch + CH ₂ deform.	983s	Xe	IR	2
			982	CO ₂	IR	1,2
		CO stretch	927s	Xe	IR	2
			926	CO ₂	IR	1,2
		O ₃ s-stretch	846wm	Xe	IR	2
		COO bend + CO stretch	727vs	Xe	IR	2
			726	CO ₂	IR	1,2
		O ₃ a-stretch	647vs	Xe	IR	2
			648	CO ₂	IR	1,2
		O ₃ bend	409m	Xe	IR	2
			406	CO ₂	IR	1,2

 $A_0=0.273; B_0=0.252; C_0=0.147 \text{ MW}^3$ **1,2,3-C₂D₄O₃**

\tilde{X}	C _s					
Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		CD ₂ deform.	1089wm	Xe	IR	2
		CO stretch	892s	Xe	IR	2
		O ₃ a-stretch	636vs	Xe	IR	2
		O ₃ bend	387m	Xe	IR	2

 $A_0=0.231; B_0=0.222; C_0=0.131 \text{ MW}^4$ **cyc-C₅Br₄O-1-O**

In argon and nitrogen matrices,¹ an absorption between 380 and 520 nm is associated with photoisomerization of and O-atom photoelimination from cyc-C₅Br₄O-1-O.

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
CO stretch			1522w	Ar	IR	1
			1525w	N ₂	IR	1
			1363w	Ar	IR	1
			1355w			
			1371w	N ₂	IR	1
			1187m	Ar	IR	1
			1185m	N ₂	IR	1
			1107w	Ar	IR	1
			1107w	N ₂	IR	1
			1000w	Ar	IR	1
OO stretch			1001w	N ₂	IR	1
			997w			
			987w	N ₂	IR	1

Reference¹I. R. Dunkin and A. McCluskey, Spectrochim. Acta A **50**, 151 (1994).¹B. Nelander and L. Nord, Tetrahedron Lett. 2821 (1977).²C. K. Kohlmiller and L. Andrews, J. Am. Chem. Soc. **103**, 2578 (1981).³J. Zozom, C. W. Gillies, R. D. Suenram, and F. J. Lovas, Chem. Phys. Lett. **140**, 64 (1987).⁴J. Z. Gillies, C. W. Gillies, R. D. Suenram, and F. J. Lovas, J. Am. Chem. Soc. **110**, 7991 (1988).

1,2,4-cyc-C₂H₄O₃

\bar{X}	C ₂	Structure: MW ^{1,2,4}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
<i>a</i>	1	CH stretch	2973m	Ar	IR	3,7	
	2	CH stretch	2894vs	Ar	IR	3,7	
	4	CH ₂ wag	1387m	Ar	IR	3,7	
	5	CH ₂ twist	1196m	Ar	IR	3,7	
	6	CH ₂ rock	1129s	Ar	IR	3,6,7	
			1130	Xe	IR	6	
	7	C-O _e stretch	955wm	gas	IR	5	
			952vs	Ar	IR	3,6,7	
			945	Xe	IR	6	
	8	C-O _p stretch	926w	Ar	IR	3,7	
	9	O-O stretch	810w	gas	IR	5	
			808s	Ar	IR	3,6,7	
			802	Xe	IR	6	
	10	Skeletal	737vw	Ar	IR	3,7	
	11	Ring pucker	352vw	Ar	IR	3	
	b	12	CH stretch	2967s	Ar	IR	3,7
	13	CH stretch	2900m	Ar	IR	3,7	
	14	CH ₂ bend	1483vw	Ar	IR	3	
	15	CH ₂ wag	1346m	Ar	IR	3,7	
	16	CH ₂ twist	1202m	Ar	IR	3,7	
	17	CH ₂ rock	1143vw	Ar	IR	3	
	18	C-O _e stretch	1081.8s	gas	IR	5	
			1078vs	Ar	IR	3,6,7	
			1072	Xe	IR	6	
	19	C-O _p stretch	1029s	Ar	IR	3,6,7	
			1021	Xe	IR	6	
	20	Skeletal	698m	Ar	IR	3,7	
	21	Ring bend	193m	Ar	IR	3	

 $A_0=0.275$; $B_0=0.270$; $C_0=0.153$ MW¹**1,2,4-cyc-C₂D₄O₃**

\bar{X}	C _{2v}	Structure: MW ^{1,2,4}					
Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.	
<i>a</i>	1	CD stretch	2249m	Ar	IR	3,7	
	2	CD stretch	2118w	Ar	IR	3,7	
	3	CD ₂ wag+skel.	1160s	Ar	IR	3,7	
	4	CD ₂ def.+skel.	1135s	Ar	IR	3,7	
	5	Skeletal	1021s	Ar	IR	3,7	
	6	Skeletal	972m	Ar	IR	3,7	
	7	CD ₂ rock+skel.	911w	Ar	IR	3,7	
	8	CD ₂ twist	851m	Ar	IR	3,7	
	9	OO stretch	759s	Ar	IR	3,7	
	10	Skeletal	672m	Ar	IR	3,7	
	11	Ring pucker	338vw	Ar	IR	3	
	b	12	CD stretch	2246w	Ar	IR	3,7
	13	CD stretch	2092w	Ar	IR	3,7	
	14	CD ₂ wag	1143s	Ar	IR	3,7	
	15	CD ₂ bend	1059vs	Ar	IR	3,7	
	16	Skeletal	980w	Ar	IR	3,7	
	17	CD ₂ twist	930m	Ar	IR	3,7	
	18	CD ₂ rock	904m	Ar	IR	3,7	
	19	CD ₂ rock	830w	Ar	IR	3,7	
	20	Skeletal	707w	Ar	IR	3,7	

^aAssignments of Ref. 3 adopted, except for C-O and O-O stretching modes of sec-C₂H₄O₃, for which assignments of Ref. 7, suggested by data for ¹⁸O-substituted species, are used.
O_p=peroxy O-atom; O_e=ether O-atom.

References

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- C. K. Kohlmiller and L. Andrews, J. Am. Chem. Soc. **103**, 2578 (1981).
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CH₃C(=NO)CHO

\bar{X}^a	Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			C=O stretch	1717.0	Ar	IR	1
				1710.3			
			C=N stretch	1587.8	Ar	IR	1
			CH ₃ deform.	1394.5	Ar	IR	1
			CH ₃ deform.	1374.0	Ar	IR	1
				1241.1	Ar	IR	1
				1020.5	Ar	IR	1
				994.3	Ar	IR	1
				807.9	Ar	IR	1
				546.1	Ar	IR	1

CD₃C(=NO)CDO

\bar{X}^a	Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			C=N stretch	1586.6	Ar	IR	1

^aFour different conformers of this species were stabilized upon photoinduced reaction of NO₂ with CH₃CCH in an argon matrix.¹ Their relative energies could not be determined. The vibrational frequencies for conformer B1, which grew in on 514 nm irradiation of the initial sample deposit, are chosen as typical.

Reference

- J. A. Harrison and H. Frei, J. Phys. Chem. **98**, 12142 (1994).

CF₃OONO₂ \tilde{X} C_s

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a'</i>	1	NO ₂ a-stretch	1764s	gas	IR	1,2
	3	CF ₃ s-stretch	1296s	gas	IR	1,2
	4	CF ₃ a-stretch	1243vs	gas	IR	1,2
	5	CO stretch	956ms	gas	IR	1,2
	7	NO stretch	787s	gas	IR	1,2
	8	CF ₃ s-deform.	669wm	gas	IR	1
	9	NO ₂ scissors	600wm	gas	IR	1
	10	CF ₃ a-deform.	485wm	gas	IR	1
	11	NO ₂ rock	436vw	gas	IR	1
	12	CF ₃ rock	371wm	gas	IR	1
	13	COO deform.	284vw	gas	IR	1
	14	OON deform.	253wm	gas	IR	1
	15	CF ₃ a-stretch	1190vs	gas	IR	1,2
	16	ONO ₂ OPLA	702wm	gas	IR	1
	17	CF ₃ a-deform.	562wm	gas	IR	1

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²J. Chen, V. Young, T. Zhu, and H. Niki, J. Phys. Chem. **97**, 11696 (1993).

8.17. Molecules Related to Benzene**C₆H₅Cl⁺**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			710(10)	Ar	AB	6

\tilde{E}	2B_1	C _{2v}				
$T^a=33430(160)$	gas	PE ³				
$T_0=33410(10)$	Ar	AB ⁶				
		$\tilde{F}-\tilde{X}$ 287–300 nm				
Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			710(10)	Ar	AB	6

\tilde{E}	2B_2	C _{2v}
$T^a=31570(160)$	gas	PE ³

\tilde{D}	2A_1	C _{2v}
$T^a=25920(160)$	gas	PE ^{1,3}

\tilde{C}	2B_1	C _{2v}
$T_0=21250(80)$	gas	PE ^{1,3}

20750(10) Ar AB^{4,6} $\tilde{C}-\tilde{X}$ 434–482 nm

Photodissociation into C₆H₅⁺ + Cl occurs.^{2,7–9}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			1370(10)	Ar	AB	6
			900(80)	gas	PE	3
			900(10)	Ar	AB	6
			370(10)	Ar	AB	6

\tilde{B}	2B_2	C _{2v}
$T_0=18300(80)$	gas	PE ^{1,3} PF ^{7–9}

Ar AB⁶ $\tilde{B}-\tilde{X}$ 485–580 nm
 $\tilde{B}-\tilde{X}$ 498–534 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			940T	gas	PE,PF	3,7
			930(10)	Ar	AB	6
			385T	gas	PE,PF	3,7
			627T	gas	PF	7
<i>a</i> ₂			179T	gas	PF	7
			430T	Ar	AB	6

\tilde{A}	2A_2	C _{2v}
$T^a=5160(160)$	gas	PE ^{1,3}

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁			1020(80)	gas	PE	3

 \tilde{X} 2B_1

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	5		1429(2)	gas	PE,TPE	3,12
	6		1200(2)	gas	TPE	11,12
	7		1116(2)	gas	TPE	11,12
	8		995(2)	gas	TPE	11,12
	9		975(2)	gas	TPE	11,12
	10		716(2)	gas	TPE	11,12
	11		422(2)	gas	TPE, PF,MPI	3,5,7 10–12
	14		348(2)	gas	MPI,PE	10–12
	15		909(20)	gas	TPE	11
	16		812(20)	gas	TPE	11
	17		603(20)	gas	TPE	11
<i>b</i> ₁	18		558(20)	gas	TPE	11
	19		394(2)	gas	TPE	11,12
	20		154(10)	gas	MPI,PE	10,11
<i>b</i> ₂	29		531(2)	gas	MPI,PE	10–12
	30		311(2)	gas	MPI,PE	10–12

C₆D₅Cl⁺ **\tilde{X} 2B_1**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
<i>a</i> ₁	11		413(20)	gas	TPE	11

^aFrom vertical ionization potential.

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1,4-C₆H₄F₂⁺

\tilde{E} 2A_g D_{2h}
 $T^a = 41600(1000)$ gas PE²

\tilde{D} $^2B_{2u}$ D_{2h}
 $T^a = 36800(1000)$ gas PE²

\tilde{C} $^2B_{3g}$ D_{2h}
 $T^a = 27100(1000)$ gas PE²

\tilde{B} $^2B_{3u}$ D_{2h}
 $T_0 = 23754$ gas PE²ID^{5,6}
23200 Ne AB¹
Ar AB³
 $\tilde{B} - \tilde{X}$ 390–425 nm
 $\tilde{B} - \tilde{X}$ 395–425 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.	meas.	
a_g			1450(40)	gas	PE	2
			404(10)	gas	ID	6
			400T	Ne	AB	1
			400	Ar	AB	3
a_u	8		200H	gas	ID	6
b_{3u}	30		144H	gas	ID	5,6

\tilde{A} $^2B_{1g}$ D_{2h}
 $T_0 = 7180(160)$ gas PE²

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.	meas.	
a_g			1355(40)	gas	PE	2

\tilde{X} $^2B_{2g}$ D_{2h}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.	meas.	
a_g	1		3098	gas	TPE	8
	2		1640	gas	TPE	7,8
	3		1375	gas	TPE	7,8
	4		1149	gas	TPE	7,8
a_u	5		836	gas	TPE	7,8
	6		440	gas	ID,TPE	6–8
	8		358	gas	ID,TPE	6–8
	9		726	gas	TPE	8
b_{1g}	17		302	gas	TPE	7,8
b_{2g}	27		430	gas	TPE	8
b_{3g}	28		859T	gas	TPE	8
b_{3u}	29		508(10)H	gas	PE	4
	30		126	gas	ID,TPE	5,7,8

1,4-C₆D₄F₂⁺

\tilde{B} $^2B_{3g}$ D_{2h}
 $T_0 = 23340$ Ne AB¹

^aFrom vertical ionization potential.

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(2-FC₆H₄)CH₂

\tilde{A} T₀=21924 gas EM¹LF² $\tilde{A} - \tilde{X}$ 435–475 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.	meas.	
			1022	gas	LF	2
			704	gas	LF	2
			501	gas	LF	2
			386	gas	LF	2

$\tau_0 = 1220(40)$ ns gas LF²

\tilde{X}

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.	meas.	
			882	gas	LF	2

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(3-FC₆H₄)CH₂

\tilde{A} T₀=21691 gas EM¹LF² $\tilde{A} - \tilde{X}$ 445–477 nm

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type	Refs.
sym.				meas.	meas.	
			734	gas	LF	2
			466	gas	LF	2

$\tau_0 = 597(15)$ ns gas LF²

\tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			741	gas	LF	2
			506	gas	EM,LF ¹	1,2

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(4-FC₆H₄)CH₂

\tilde{A}	2A_2	C_{2v}	$T_0=21524.0(2)$	gas	EM ^{1,3} LF ²	$\tilde{A}-\tilde{X}$ 429–475 nm
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Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1760	gas	LF	2
	7a		1228	gas	LF	4
	13		1194	gas	LF	2,4
	18a		846	gas	LF	4
	1		813	gas	LF	2,4
	6b		524	gas	LF	4
	6a		411	gas	LF	4
	9b		398	gas	LF	4
	15		222	gas	LF	4

$\tau_0=532(5)$ ns gas LF^{2,4}

 \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
	1		835	gas	EM,LF	3,4
	6a		453	gas	EM,LF	1–4

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(4-CIC₆H₄)CH₂

\tilde{B}	2B_1	C_{2v}	$T_0=21732$	gas	LF ¹	$\tilde{A},\tilde{B}-\tilde{X}$ 436–462 nm
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Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
			1194	gas	LF	1
			1076	gas	LF	1
			960	gas	LF	1
			779	gas	LF	1
			340	gas	LF	1
			242	gas	LF	1
b_2	6b		623	gas	LF	1
			122	gas	LF	1

 \tilde{A} 2A_2 C_{2v} $T_0=21638$

gas

LF¹ $\tilde{A},\tilde{B}-\tilde{X}$ 436–462 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
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a_1 12

726

gas

LF

1

 \tilde{X} 2B_1 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
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a_1 8a

1544

gas

LF

2

9a

1079

gas

LF

2

1

811

gas

LF

2

12

726

gas

LF

1

6a

559

gas

LF

2

6b

634

gas

LF

1

209

gas

LF

1

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(4-NC-C₆H₄)CH₂ \tilde{A} 2B_1 C_{2v} $T_0=20738T$

gas

LF¹ $\tilde{A}-\tilde{X}$ 445–580 nm

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
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b_2 6b

516

gas

LF

1

$\tau_0=400T$ ns

gas

LF¹

$A_0=0.172$

$B_0=0.032$

$C_0=0.028$

LF¹

 \tilde{X} 2B_1 C_{2v} T_0

gas

LF¹

a_1

CN stretch

2216

gas

LF

1

8a

1558

gas

LF

1

9a

1143

gas

LF

1

1

818

gas

LF

1

6a

513

gas

LF

1

6b

637

gas

LF

1

$A_0 = 0.182$

$B_0 = 0.032$

$C_0 = 0.028$

LF¹

Reference

¹M. Fukushima, K. Saito, and K. Obi, J. Mol. Spectrosc. **180**, 389 (1996).

C₆H₅N

In an argon or a nitrogen matrix, irradiation at wavelengths longer than 450 nm leads to isomerization to cyclo-C₆H₅N.¹

 \tilde{a} 1A_1 C_{2v} $T_0=6300(700)$

gas

PE²PD³

\tilde{X}^3A_2 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		1552s	Ar	IR	1	
		1524m	Ar	IR	1	
		1497w	Ar	IR	1	
		1426wm	Ar	IR	1	
		1408w	Ar	IR	1	
		1309w	Ar	IR	1	
		1300T	gas	PE	2	
		1299w	Ar	IR	1	
		1286ms	Ar	IR	1	
		1250w	Ar	IR	1	
		1148m	Ar	IR	1	
		1079wm	Ar	IR	1	
		1008wm	Ar	IR	1	
		964w	Ar	IR	1	
		885w	Ar	IR	1	
		820wm	Ar	IR	1	
OPLA CH-bend		746vs	Ar	IR	1	
Ring torsion		654s	Ar	IR	1	
Ring breathing		515T	gas	PE,PD	2,3	

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³R. N. McDonald and S. J. Davidson, J. Am. Chem. Soc. **115**, 10857 (1993).

 $2,6\text{-C}_6\text{H}_3\text{F}_2\text{N}:$

In an argon matrix, a prominent absorption at 42400 (236 nm) and weaker absorptions at 35200 (284 nm) and 32000 (312 nm) have been assigned¹ to $2,6\text{-C}_6\text{H}_3\text{F}_2\text{N}$:

In an argon matrix, prolonged irradiation at 22500 (444 nm) results in reversible rearrangement to the azirine (cf. *bicyc-C₆H₃F₂N*).¹

 \tilde{X}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		1585.1m	Ar	IR	1	
		1577.7s	Ar	IR	1	
		1459.4m	Ar	IR	1	
		1452.2vs	Ar	IR	1	
		1411.2wm	Ar	IR	1	
		1312.4m	Ar	IR	1	
		1281.1wm	Ar	IR	1	
		1242.0m	Ar	IR	1	
		1229.3m	Ar	IR	1	
		1002.8w	Ar	IR	1	
		991.6s	Ar	IR	1	
		850.4wm	Ar	IR	1	
		836.6w	Ar	IR	1	
		772.4wm	Ar	IR	1	
		767.7m	Ar	IR	1	
		708.3wm	Ar	IR	1	
		699.6w	Ar	IR	1	
		553.2w	Ar	IR	1	
		501.2w	Ar	IR	1	

Reference

- ¹J. Morawietz and W. Sander, J. Org. Chem. **61**, 4351 (1996).

 $\text{C}_6\text{F}_5\text{N}:$

In an argon matrix,² irradiation at 22500 (444 nm) results in isomerization to the azirene (cf. *bicyc-C₆F₅N*).

 \tilde{X} C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
		1572.4wm	Ar	IR	2	
		1575m	N_2	IR	1	
		1564.4wm	Ar	IR	2	
		1565m	N_2	IR	1	
		1502.7vs	Ar	IR	2	
		1502vs	N_2	IR	1	
		1464.0m	Ar	IR	2	
		1462vs	N_2	IR	1	
		1455.6ms	Ar	IR	2	
		1454vs	N_2	IR	1	
		1359.4wm	Ar	IR	2	
		1359s	N_2	IR	1	
		1334.6w	Ar	IR	2	
		1281.4wm	Ar	IR	2	
		1284m	N_2	IR	1	
		1205.5w	Ar	IR	2	
		1204w	N_2	IR	1	
		1151.7w	Ar	IR	2	
		1149w	N_2	IR	1	
		1025.5s	Ar	IR	2	
		1029vs	N_2	IR	1	
		1008.5wm	Ar	IR	2	
		1007s	N_2	IR	1	
		1000.6wm	Ar	IR	2	
		990m	N_2	IR	1	
		982.3vs	Ar	IR	2	
		981vs	N_2	IR	1	
		916.1w	Ar	IR	2	
		610.7vw	Ar	IR	2	

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 $\text{C}_6\text{H}_5\text{N}^-$

Threshold for electron detachment from ground-state $\text{C}_6\text{H}_5\text{N}^- = 11700(160)$ gas PE^{1,2}

 \tilde{X}^2B_2 C_{2v}

Vib. sym.	No.	Approximate type of mode	cm^{-1}	Med.	Type meas.	Refs.
a_1		Ring breathing	1350T	gas	PE	1
			500T	gas	PE	1

C₆D₅N⁻

Threshold for electron detachment from ground-state C₆D₅N⁻ = 11620(160) gas PE¹PD²

 \tilde{X}^2B_2 **C_{2v}**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			434HT	gas	PD	1

References

- ¹M. J. Travers, D. C. Cowles, E. P. Clifford, and G. B. Ellison, *J. Am. Chem. Soc.* **114**, 8699 (1992).
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C₆H₅NH₂⁺ **\tilde{A}^2A_2** **C_{2v}**

T₀=10570(80) gas PE⁷

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	1		840(40)	gas	PE	7
	6a		524(10)	gas	PE	7

 \tilde{X}^2B_1 **C_{2v}**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁	8a ^a		1594	gas	TPE	2,5,6
	19a		1436(10)T	gas	PE	2
	13		1385	gas	TPE	2,5,6
	9a		1188	gas	TPE	2,4-6
	18a		996	gas	TPE	5,6
	12	Ring s-stretch	982	gas	PI,TPE	1,2,4-6
	1		814	gas	PI,TPE	1,2,4,6
	6a	N-ring s-str.	522	gas	PI,TPE	1-6
		C-N torsion	577H	gas	TPE	6
	16a		356	gas	PI,TPE	2-6
b ₁	17b		963T ^b	gas	TPE	6
	11		790 ^b	gas	TPE	6
b ₂		Inversion	658	gas	PI,TPE	2,3,5,6
	16b		629	gas	TPE	6
	4	Ring deform.	445	gas	TPE	6
	10b		179	gas	PI,TPE	3,5,6
b ₂	6b		582	gas	TPE	6
	15		550H	gas	TPE	5,6

C₆D₅ND₂⁺ **\tilde{X}^2B_1** **C_{2v}**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
b ₁		Inversion	465(10)	gas	PI	3

^aVibrational numbering and assignments parallel those given by Chernoff and Rice [J. Chem. Phys. **70**, 2511 (1979)] for aniline.

^bFrom combination band.

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C₆H₅O **\tilde{C}** **C_{2v}**

gas AB^{1,2}

T₀=25175(10) Ar AB^{4,5}

$\tilde{C}-\tilde{X}$ 380-395 nm
 $\tilde{C}-\tilde{X}$ 351-397 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1462(20)	Ar	AB	4
			1140(20)	Ar	AB	4
			920(20)	Ar	AB	4

 \tilde{B} **C_{2v}**

gas AB^{2,3}

T₀=16360 Ar AB^{4,5}

$\tilde{B}-\tilde{X}$ 559-612 nm
 $\tilde{B}-\tilde{X}$ 573-629 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			500T	gas	AB	3
			504(10)	Ar	AB	4

 \tilde{A} **C_{2v}**

gas PE⁶

 \tilde{X} **C_{2v}**

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a ₁			1490(25)T	gas	PE	6
			515(15)	gas	PE	6

C₆D₅O **\tilde{C}** **C_{2v}**

Ar AB⁴

$\tilde{C}-\tilde{X}$ 349-397 nm

Vib. sym.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1349(20)	Ar	AB	4
			809(20)	Ar	AB	4

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C₆H₅OH⁺

In an argon matrix, an absorption maximum at 36310 (275 nm) has been attributed³ to C₆H₅OH⁺.

B	C _s	T ₀ =21129T	gas	PE ^{1,3} PRI ⁴	405–465 nm
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In an argon matrix, an absorption maximum at 23640 (423 nm) has been attributed³ to C₆H₅OH⁺.

Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			392T	gas	PRI	4
			363T	gas	PRI	4
			347T	gas	PRI	4

A	C _s	T ^a =7180(230)	gas	PE ^{1,3}
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
a'			1100(80)	gas	PE	1

X	C _s
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			1669(33)	gas	PE	2
			1500(30)	gas	PE	2
			1395(28)	gas	PE	2
			1210(24)	gas	PE	2
			1040(21)	gas	PE	2
			1027(21)	gas	PE	2
			976(20)	gas	PE	2
			815(16)	gas	PE	2
			556(12)	gas	PE	2
			516(12)	gas	PE	2
		Ring torsion ?	169(12)	gas	PE	2

C₆D₅OD⁺

B	C _s	gas	PRI ⁴	425–442 nm
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Vib.	No.	Approximate type of mode	cm ⁻¹	Med.	Type meas.	Refs.
			234T	gas	PRI	4

^aFrom vertical ionization potential.

References

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9. Molecule Index

In the References column, unprefixed numbers give the page on which the table appears in the monograph.⁵ Numbers prefixed by "A" give the page number in the present paper.

Formula	Structure/Name	References
AgAuCu	CuAgAu	60
AgCu ₂	Cu ₂ Ag	59
AgO ₂	AgOO	A178
Ag ₃	Ag ₃	60,A161
AlBr ₂	AlBr ₂	A193
AlBr ₃ ⁺	AlBr ₃ ⁺	203
AlClO	OAlCl	85
AlCl ₂	AlCl ₂	90,A193
AlCl ₂ H	HAICl ₂	169
AlCl ₃ ⁺	AlCl ₃ ⁺	203
AlFO	FAIO	85
AlF ₂	AlF ₂	A192
AlF ₃ ⁺	AlF ₃ ⁺	202
AlHO	AlOH	36,A143
AlH ₂	AlH ₂	17,A129
AlH ₂ O	HAIOH	139
AlH ₃	AlH ₃	A212
AlH ₄ ⁻	AlH ₄ ⁻	A266
AlH ₄ Si	SiH ₃ AlH	307
AlI ₂	AlI ₂	A193
AlO ₂	cyc-AlO ₂	75
AlO ₂	OAlO	75,A183
AlO ₂ ⁻	OAlO ⁻	A189
AlO ₃	OAlOO	190
Al ₂ H ₂	cyc-Al ₂ H ₂	A219
Al ₂ H ₂	H(cyc-AlHAl)	A219
Al ₂ H ₂	HAIAIH	A220
Al ₂ O	Al ₂ O	63
Al ₂ O ₂	(AlO) ₂	184,A248
Al ₂ O ₂ ⁻	(AlO) ₂ ⁻	A249
Al ₂ O ₃	Al ₂ O ₃	283,A297
Al ₂ O ₃ ⁻	Al ₂ O ₃ ⁻	A297
Al ₂ O ₄	Al ₂ O ₄	A321
Al ₂ O ₄ ⁻	Al ₂ O ₄ ⁻	A321
Al ₂ O ₅	Al ₂ O ₅	A331
Al ₂ S ₃	Al ₂ S ₃	A297
Al ₃	Al ₃	62,A162
ArBeO	ArBeO	A188
ArBeO ₂	ArBeO ₂	A253
ArBrKr	ArKrBr	123
ArBrXe	ArXeBr	123
ArClKr	ArKrCl	122
ArDKr	ArKrD	57
ArFKr	ArKrF	121,A210
ArFXe	ArXeF	121
ArHXe	ArXeH	57
Ar ₂ Cl	Ar ₂ Cl	122
Ar ₂ H ⁺	HAr ₂ ⁺	56,A159
Ar ₂ H	Ar ₂ H	56

Formula	Structure/Name	References	Formula	Structure/Name	References
Ar ₂ F	Ar ₂ F	121,A210	BF ₂ H ₂ N ⁺	NH ₂ BF ₂ ⁺	332
AsBr ₃ ⁺	AsBr ₃ ⁺	221	BF ₂ O	F ₂ BO	200
AsClO	ClAsO	104	BF ₃ ⁺	BF ₃ ⁺	201,A255
AsCl ₃ ⁺	AsCl ₃ ⁺	221	BF ₃ ⁻	BF ₃ ⁻	A261
AsFH ₂	AsH ₂ F	152	BF ₄ ⁻	BF ₄ ⁻	302
AsF ₂	AsF ₂	109,A203	BGaH ₆ ⁺	GaBH ₆ ⁺	A332
AsF ₂ H	HAsF ₂	179	BHN	HNB	31,A141
AsF ₂ H ₃	AsH ₃ F ₂	325	BHO	HBO	35
AsF ₃ ⁺	AsF ₃ ⁺	221	BHO ₂	HOBO	162
AsGa ₂	Ga ₂ As	A165	BHS ⁺	HBS ⁺	31,A141
AsHO ₃	HOAsO ₂	273	BHS	HBS	35
AsH ₂	AsH ₂	23	BH ₂	BH ₂	17
AsH ₃ ⁺	AsH ₃ ⁺	129	BH ₂ N	HB ₂ NH	137,A222
AsH ₃ O	H ₃ AsO	243	BH ₂ N	H ₂ NB	A223
AsH ₃ O	H ₂ AsOH	243	BH ₃	BH ₃	124,A212
AsIn ₂	In ₂ As	A165	BH ₃ ⁻	BH ₃ ⁻	125
As ₂ Ga	GaAs ₂	A172	BH ₃ N	HB ₂ NH ₂	A269
As ₄ ⁺	As ₄ ⁺	187	BH ₃ O	H ₂ BOH	233
As ₄ O	As ₄ O	284	BH ₃ O ₃	B(OH) ₃	A326
As ₄ O	br-As ₄ O	284	BH ₃ S	II ₂ BSII	233
AuCu ₂	Cu ₂ Au	60	BH ₄ ⁻	BH ₄ ⁻	A266
AuO ₂	cyc-AuO ₂	A178	BH ₄ K	KBH ₄	A308
Au ₃	Au ₃	60,A161	BH ₄ Li	LiBH ₄	305,A307
BBrH ₂	H ₂ BBr	142	BH ₄ N ⁺	H ₂ BNH ₂ ⁺	307
BBrO	BrBO	85	BH ₄ N	H ₂ BNH ₂	309
BBrS	BrBS	85	BH ₄ Na	NaBH ₄	305,A307
BBr ₂ ⁺	BBr ₂ ⁺	A189	BH ₄ Sr	SrBH ₄	306
BBr ₂	BBr ₂	89,A192	BI ₂	BI ₂	89
BBr ₂ H ⁺	HBBBr ₂ ⁺	166	BI ₃ ⁺	BI ₃ ⁺	202
BBr ₂ H	HBBBr ₂	A238	BN ₂	NBN	67
BBr ₂ H ₂ N ⁺	NH ₂ BBr ₂ ⁺	332	BN ₂	BNN	69
BBr ₃ ⁺	BBr ₃ ⁺	202	BN ₃	NNBN	184
BCaH ₄	CaBH ₄	305	BO ₂	BO ₂	73
BClF ₃ ⁻	BF ₃ Cl ⁻	302	BO ₂ ⁻	BO ₂ ⁻	84
BClH ₂	H ₂ BCl	142	BO ₂ S	OBSO	190
BClO	ClBO	84,A189	BS ₂	BS ₂	74,A182
BClS ⁺	CIBS ⁺	74,A183	B ₂ Cl ₄ ⁺	B ₂ Cl ₄ ⁺	348
BClS	CIBS	85,A189	B ₂ F ₄ ⁺	B ₂ F ₄ ⁺	347
BCl ₂ ⁺	BCl ₂ ⁺	A189	B ₂ HN	HB ₂ NH	A232
BCl ₂	BCl ₂	89,A192	B ₂ H ₂	HBBH	A219
BCl ₂ H ⁺	HBCl ₂ ⁺	166	B ₂ H ₂ N ₂	HNBBNH	A315
BCl ₂ H	HBCl ₂	A238	B ₂ H ₃ N	H ₂ BNH	A311
BCl ₂ H ₂ N ⁺	NH ₂ BCl ₂ ⁺	332	B ₂ H ₄ ⁺	B ₂ II ₄ ⁺	306
BCl ₃ ⁺	BCl ₃ ⁺	201,A255	B ₂ H ₅ ⁺	B ₂ H ₅ ⁺	359
BCl ₃ ⁻	BCl ₃ ⁻	A261	B ₂ H ₆ ⁺	B ₂ H ₆ ⁺	A332
BCl ₄ ⁻	BCl ₄ ⁻	302	B ₂ H ₇ N	B ₂ H ₅ NH ₂	398
BFH ₂ ⁺	HBF ⁺	37	B ₂ N	cyc-BBN	62,A165
BFH ₂	H ₂ BF	142	B ₂ N	BNB	63,A165
BFO	FBO	84,A188	B ₂ N ₂	BBNN	181
BFS ⁺	FBS ⁺	74	B ₂ N ₂	BNBN	181
BFS	FBS	85	B ₂ N ₂	cyc-(BN) ₂	181
BF ₂ ⁺	BF ₂ ⁺	A189	B ₂ O	BBO	63
BF ₂	BF ₂	89,A191	B ₂ O	BOB	63
BF ₂ H ⁺	HBF ₂ ⁺	165	B ₂ O ₂ ⁺	B ₂ O ₂ ⁺	183
BF ₂ H	HBF ₂	A238	B ₂ O ₂	B ₂ O ₂	184

Formula	Structure/Name	References	Formula	Structure/Name	References
B ₃	B ₃	A162	BrF ₂ ⁻	FBrF ⁻	118
B ₃ F ₃ N ₃ ⁺	cyc-(FBNH) ₃ ⁺	399	BrF ₂ ⁻	FFBr ⁻	118
BaHO	BaOH	30	BrF ₂ P ⁺	PF ₂ Br ⁺	220
BaH ₂ N	BaNH ₂	133	BrF ₃ ⁺	BrF ₃ ⁺	226
BaH ₂ O	HBaOH	135	BrF ₃ Si ⁺	SiF ₃ Br ⁺	295
BaH ₂ O ₂	Ba(OH) ₂	251	BrF ₅ ⁺	BrF ₅ ⁺	358
BaO ₂	OBaO	A175	BrFeH	HFeBr	35
BaO ₂	cyc-BaO ₂	A175	BrGeH	HGeBr	45
Ba ₂ O ₂	cyc-(BaO) ₂	A243	BrGeH ₂	H ₂ GeBr	150
Ba ₂ O ₂	BaOBaO	A243	BrGeH ₃ ⁺	GeH ₃ Br ⁺	241
BeBr ₂	BeBr ₂	84	BrHI ⁻	BrHI ⁻	55
BeCl ₂	BeCl ₂	84	BrHO ⁺	HOBr ⁺	A155
BeF ₂	BeF ₂	83	BrHO	HOBr	52,A156
BeHO	BeOH	A138	BrHSi	HSiBr	44,A151
BeH ₂	BeH ₂	A125	BrHXe	HXeBr	A158
BeH ₂ O	HBeOH	A222	BrH ₂ N ⁺	H ₂ NBr ⁺	152
BeH ₂ O ₂	Be(OH) ₂	A281	BrH ₂ P	H ₂ PBr	A229
BeI ₂	BeI ₂	84	BrH ₃ Si ⁺	SiH ₃ Br ⁺	241
BeKrO	KrBeO	A188	BrKrXe	KrXeBr	123
BeN ₂ O ₂	NNBeO ₂	A296	BrKr ₂	Kr ₂ Br	123
BeN ₄	NNBeNN	A294	BrNO ⁺	BrNO ⁺	94
BeN ₄	NN(cyc-BeN ₂)	A294	BrNO ₂	BrNO ₂	213,A260
BeOXe	XeBeO	A188	BrNO ₂ ⁺	t-BrONO	213,A260
BeO ₂ ⁺	OBeO ⁺	A171	BrNO ₃ ⁺	BrONO ₂ ⁺	A301
BeO ₂	OBeO	A174	BrNO ₃	BrONO ₂	289,A303
BeO ₃	BeO ₃	A250	BrNS ⁺	NSBr ⁺	95
BeO ₄	O ₂ BeO ₂	A297	BrNS	NSBr	104
Be ₂ H	BeBeH	A136	BrN ₃ ⁺	BrN ₃ ⁺	197
Be ₂ H ₂	HBeBeH	A217	BrNeXe	NeXeBr	123
Be ₂ H ₂ O	HBeOBeH	A275	BrOP	BrPO	103
Be ₂ H ₃	HBeHBeH	A267	BrOPS	BrP(O)S	214
Be ₂ H ₄	HBeH ₂ BeH	A308	BrO ₂	BrOO	110,A204
Be ₂ N ₂	BeBeNN	A242	BrO ₂	OBRO	114,A206
Be ₂ N ₂	BeNNBe	A242	BrO ₂ P	PO ₂ Br	214
Be ₂ N ₂	(BeN) ₂	A242	BrPS	BrPS	104
Be ₂ N ₄	NNBeBeNN	A317	BrS ₂	SSBr	111
Be ₂ O	BeOBe	A163	BrXe ₂	Xe ₂ Br	123
Be ₂ O ₂	cyc-(BeO) ₂	A242	Br ₂ Cl ⁻	BrClBr ⁻	119
Be ₂ O ₂	BeOBeO	A242	Br ₂ Cl ⁻	ClBrBr ⁻	119
Be ₂ O ₃	OBeOBeO	A294	Br ₂ F	BrBrF	117
Bi ₃	Bi ₃	82	Br ₂ F ₂	Br ₂ F ₂	227
Bi ₄	Bi ₄	189	Br ₂ Ge ⁺	GeBr ₂ ⁺	92
Bi ₄ ⁻	Bi ₄ ⁻	197	Br ₂ GeH ₂ ⁺	GeH ₂ Br ₂ ⁺	263
BrClH ⁻	ClHBr ⁻	54	Br ₂ H ⁻	BrHBr ⁻	55,A157
BrClO	BrClO	A209	Br ₂ HN ⁺	HNBr ₂ ⁺	179
BrClO	ClBrO	A209	Br ₂ H ₂ Si ⁺	SiH ₂ Br ₂ ⁺	262
BrClO	BrOCl	A207	Br ₂ N	NBr ₂	108
BrClO ₂	BrClO ₂	A263	Br ₂ O	BrOBr	114,A207
BrClO ₂	BrOCIO	A264	Br ₂ O	BrBrO	116,A209
BrClO ₂	CIOBrO	A264	Br ₂ OP	OPBr ₂	221
BrCl ₂ ⁻	ClBrCl ⁻	119	Br ₂ P	PBr ₂	109,A203
BrCl ₂ ⁻	ClClBr ⁻	119	Br ₂ S ⁺	SBr ₂ ⁺	112
BrFH ⁻	FHBr ⁻	54	Br ₂ S	SBr ₂	115
BrFS	FSBr	A208	Br ₂ S ₂ ⁺	S ₂ Br ₂ ⁺	224
BrF ₂	BrF ₂	117	Br ₂ S ₂	SSBr ₂	226

Formula	Structure/Name	References	Formula	Structure/Name	References
Br ₂ Se ⁺	SeBr ₂ ⁺	113	CCaNO	CaNCO	182
Br ₂ Si	SiBr ₂	101	CClF	CFCI	98
Br ₃ ⁻	Br ₃ ⁻	119	CCIFOS	CIFCSO (I)	288
Br ₃ P ⁺	PBr ₃ ⁺	221	CCIFOS	CIFCSO (II)	288
Br ₃ PO ⁺	Br ₃ PO ⁺	299	CCIFS ⁺	FCICS ⁺	205
Br ₃ PS ⁺	Br ₃ PS ⁺	299	CCIF ₂ ⁺	CF ₂ Cl ⁺	210
Br ₃ Sb ⁺	SbBr ₃ ⁺	222	CCIF ₂	CF ₂ Cl	216
Br ₄ Ge ⁺	GeBr ₄ ⁺	297	CCIF ₂ NO ⁺	CF ₂ CINO ⁺	353
Br ₄ Si ⁺	SiBr ₄ ⁺	296	CCIF ₂ NO ₄	CF ₂ CIOONO ₂	425
CAI _O	AlCO	68,A172	CCIF ₃ ⁺	CF ₃ Cl ⁺	290
CAI ₂	Al ₂ C	A164	CCIF ₃ ⁻	CF ₃ Cl ⁻	303
CAsN	AsCN	71	CCIF ₄	CF ₃ CIF	356
CBF ₂ NO	F ₂ BNCO	A321	CCIN ⁺	CICN ⁺	79
CBO	BCO	68	CCIN	CINC	88
CBO ₂	OBCO	186	CCINO ⁺	CINCO ⁺	193
CBS ₂	SBCS	A249	CCINO	CICNO	199
CBaN	BaCN	62	CCINO	CINCO	198
CBeO ₂	OCBeO	A247	CCINS ⁺	CISCN ⁺	194
CBeO ₂	COBeO	A248	CCINS	CISCN	198
CBe ₂ O	BeBeCO	A241	CCINSe ⁺	ClSeCN ⁺	195
CBrCl ⁺	CClBr ⁺	91	CCIO	ClCO	90
CBrCl	CClBr	99	CCIP ⁺	CICP ⁺	81
CBrCl ₂ ⁺	CCl ₂ Br ⁺	211	CCIP	CICP	A190
CBrCl ₂	CCl ₂ Br	218	CCIS	CICS	105
CBrClF ₂ ⁻	CF ₂ ClBr ⁻	303	CCl ₂ ⁺	CCl ₂ ⁺	91
CBrF	CFBr	98	CCl ₂	CCl ₂	99
CBrF ₂ ⁺	CF ₂ Br ⁺	210	CCl ₂ ⁻	CCl ₂ ⁻	107
CBrF ₂	CF ₂ Br	216	CCl ₂ F ⁺	CFCl ₂ ⁺	210
CBrF ₂ ⁻	CF ₂ Br ⁻	225	CCl ₂ F	CFCl ₂	216
CBrF ₃ ⁺	CF ₃ Br ⁺	291	CCl ₂ FNO ⁺	CFCl ₂ NO ⁺	353
CBrF ₃ ⁻	CF ₃ Br ⁻	303	CCl ₂ FNO ₄	CFCl ₂ OONO ₂	426
CBrF ₄	CF ₃ BrF	356	CCl ₂ F ₂ ⁺	CF ₂ Cl ₂ ⁺	292,A304
CBrN ⁺	BrCN ⁺	80,A187	CCl ₂ F ₂ ⁻	CF ₂ Cl ₂ ⁻	303
CBrN	BrNC	88	CCl ₂ I	CCl ₂ I	218
CBrNO ⁺	BrCNO ⁺	A252	CCl ₂ O ⁺	Cl ₂ CO ⁺	204
CBrNO ⁺	BrNCO ⁺	193	CCl ₂ S ⁺	Cl ₂ CS ⁺	206
CBrNO	BrCNO	199,A253	CCl ₂ Se	Cl ₂ CSe	212
CBrNO	BrNCO	198	CCl ₃ ⁺	CCl ₃ ⁺	210
CBrNS ⁺	BrSCN ⁺	194	CCl ₃	CCl ₃	217
CBrNS	BrSCN	199	CCl ₃ F ⁺	CFCl ₃ ⁺	293
CBrNSE ⁺	BrSeCN ⁺	195	CCl ₃ F ⁻	CFCl ₃ ⁻	303
CBr ₂ ⁺	CBr ₂ ⁺	91	CCl ₃ NO ⁺	CCl ₃ NO ⁺	353
CBr ₂	CBr ₂	100	CCl ₃ NO ₄	CCl ₃ OONO ₂	426
CBr ₂ Cl ⁺	CClBr ₂ ⁺	211	CCl ₃ O ₂	CCl ₃ O ₂	354
CBr ₂ Cl	CClBr ₂	218	CCl ₄ ⁺	CCl ₄ ⁺	294,A305
CBr ₂ F ⁺	CFBr ₂ ⁺	210	CCl ₄	Cl ₂ CCl-Cl	302
CBr ₂ F	CFBr ₂	216	CCoO	CoCO	A167
CBr ₂ F ₂ ⁺	CF ₂ Br ₂ ⁺	292	CCrO	CrCO	A166
CBr ₂ F ₂ ⁻	CF ₂ Br ₂ ⁻	303	CCuO	CuCO	A167
CBr ₃ ⁺	CBr ₃ ⁺	211	CD ₃ O ⁺	CD ₃ O ⁺	235
CBr ₃	CBr ₃	218	CFI	CFI	99
CBr ₃ F ⁺	CFBr ₃ ⁺	293	CFI ₂ ⁺	CFI ₂ ⁺	210
CBr ₃ F ⁻	CFBr ₃ ⁻	303	CFI ₂	CFI ₂	217
CBt ₄ ⁺	CBt ₄ ⁺	294	CFN ⁺	FCN ⁺	79
CCaN	CaNC	62,A164	CFN	FNC	88

Formula	Structure/Name	References	Formula	Structure/Name	References
CFNO	FNCO	198,A253	CHBrF	HCFBr	177
CFNO ₅	FC(O)OONO ₂	A342	CHBrO	HCOBr	170,A239
CFNS ⁺	FSCN ⁺	193	CHBr ₂ ⁺	HCB ₂ ⁺	172
CFN ₂	FNCN	192	CHBr ₂	HCB ₂	178
CFO ⁺	FCO ⁺	86	CHBr ₃ ⁺	HCB ₃ ⁺	276
CFO	FCO	90,A193	CHCaO ₂	HCOOCa	268
CFO ₂	FCO ₂	203,A256	CHCl	HCCl	42,A149
CFO ₂ ⁻	FCO ₂ ⁻	212,A259	CHCl ⁻	HCCl ⁻	47
CFO ₃	FC(O)O ₂	287	CHClF ⁺	HCFCI ⁺	171
CFP ⁺	FCP ⁺	81	CHClF	HCFCI	176
CFP	FCP	A190	CHClF ₂ ⁺	HCF ₂ Cl ⁺	275
CFS	FCS	A194	CHClO ⁺	HOCl ⁺	169
CF ₂ ⁺	CF ₂ ⁺	91	CHClO	HOCl	170,A239
CF ₂	CF ₂	97,A196	CHCl ₂ ⁺	HCCl ₂ ⁺	171
CF ₂ ⁻	CF ₂ ⁻	107	CHCl ₂	HCCl ₂	177
CF ₂ I ⁺	CF ₂ I ⁺	210	CHCl ₂ F ⁺	HCFCI ₂ ⁺	275
CF ₂ I	CF ₂ I	216	CHCl ₂ NO ₄	CHCl ₂ OONO ₂	425
CF ₂ N	F ₂ CN	204	CHCl ₃ ⁺	HCCl ₃ ⁺	275
CF ₂ NOP ⁺	PF ₂ NCO ⁺	349	CHF ⁺	HCF ⁺	41
CF ₂ NP ⁺	PF ₂ CN ⁺	285	CHF	HCF	42,A148
CF ₂ NPS ⁺	PF ₂ NCS ⁺	350	CHF ⁻	HCF ⁻	47
CF ₂ N ₂ ⁺	CF ₂ N ₂ ⁺	285	CHFI ⁺	HCFI ⁺	171
CF ₂ O ⁺	F ₂ CO ⁺	204	CHFI	HCFI	177
CF ₂ OS	F ₂ CSO	288	CHFN	HFCN	168
CF ₂ O ₂	cyc-F ₂ CO ₂	A302	CHFO ⁺	HFCO ⁺	168
CF ₂ S ⁺	F ₂ CS ⁺	205	CHFS ⁺	HFCS ⁺	A237
CF ₂ Se ⁺	F ₂ CSe ⁺	206	CHF ₂ ⁺	HCF ₂ ⁺	170
CF ₃ ⁺	CF ₃ ⁺	210,A258	CHF ₂	HCF ₂	176
CF ₃	CF ₃	215,A261	CHF ₂ N ⁺	CF ₂ NH ⁺	A289
CF ₃ ⁻	CF ₃ ⁻	A263	CHF ₂ N	CF ₂ =NH	270,A289
CF ₃ I ⁺	CF ₃ I ⁺	291	CHF ₂ N	c-HFC=Nf	271
CF ₃ I ⁻	CF ₃ I ⁻	303	CHF ₂ P	CF ₂ =PH	271
CF ₃ IO	CF ₃ IO	356	CHF ₃ ⁺	HCF ₃ ⁺	274
CF ₃ IO	CF ₃ OI	356	CHI	HCI	43
CF ₃ NO ⁺	CF ₃ NO ⁺	353	CHI ⁻	HCI ⁻	48
CF ₃ NO ₃	CF ₃ ONO ₂	382,A342	CHIO	HCOI	A240
CF ₃ NO ₄	CF ₃ OONO ₂	A368	CHI ₂	HCl ₂	178
CF ₃ O	CF ₃ O	289,A303	CHN ⁺	HCN ⁺	33
CF ₃ O ⁻	CF ₃ O ⁻	302	CHN ⁺	HNC ⁺	34
CF ₃ O ₂	CF ₃ O ₂	354	CHN	HNC	38,A145
CF ₃ S	CF ₃ S	A303	CHNO ⁺	HNCO ⁺	160
CF ₄ ⁺	CF ₄ ⁺	290,A303	CHNO ⁺	HCNO ⁺	161
CF ₄ I	CF ₃ IF	356	CHNO	HNCO	163,A236
CF ₄ O ⁺	CF ₃ OF ⁺	355	CHNO	HOCN	163
CF ₄ OS	CF ₃ OSF	372	CHNO	HCNO	164
CFeO	FeCO	63,A167	CHNO	HONC	164
CFeO ⁻	FeCO ⁻	67	CHNP	HPCN	160
CGaO	GaCO	A172	CHNS ⁺	HNCS ⁺	160
CGeO	GeCO	A180	CHN ₂	HCNN	161,A235
CHBr	HCBr	43,A149	CHN ₂	HNCN	160,A235
CHBr ⁻	HCBr ⁻	47	CHO ⁺	HCO ⁺	37,A145
CHBrCl ⁺	HCClBr ⁺	171	CHO ⁺	HOC ⁺	39
CHBrCl	HCClBr	178	CHO	HCO	40,A146
CHBrClF ⁺	HCFClBr ⁺	A292	CHOP	HPCO	165
CHBrF ⁺	HCFBr ⁺	171	CHOS ⁺	HOCS ⁺	165,A236

Formula	Structure/Name	References	Formula	Structure/Name	References
CHO_2^+	HOCO^+	165	CH_2NO	H_2CNO	A283
CHO_2	<i>c</i> - HOCO	167	CH_2NOSr	HCONHSr	326
CHO_2	<i>t</i> - HOCO	167,A236	CH_2NO_2^-	CH_2NO_2^-	331,A316
CHO_2	HCO_2	A237	CH_2NO_2^-	CH_2NO_2^-	337
CHO_2^-	HCO_2^-	A239	CH_2N_2^+	CH_2N_2^+	253
CHO_2Sr	HCOOSr	268	CH_2N_2^+	<i>cyc</i> - CH_2N_2^+	254
CHO_3	HC(O)OO	270	CH_2N_2^+	NH_2CN^+	253
CHP^+	HCP^+	34	CH_2N_2	$\text{HN}=\text{C}=\text{NH}$	254,A282
CHS^+	HCS^+	38,A145	CH_2N_2	HCNNH	A282
CHS_2	<i>t</i> - HSCS	167	$\text{CH}_2\text{N}_2\text{O}$	H_2NNCO	331
CHS_2	HCS_2	168	CH_2Ni	NiCH_2	A219
CHTi	TiCH	A137	CH_2O^+	H_2CO^+	140,A223
CHV	VCH	A137	CH_2OS^+	H_2CSO^+	258
CIW	WCH	A137	CH_2OS	H_2CSO	259,A284
CH_2^+	CH_2^+	18	CH_2O_2^+	HCOOH^+	257
CH_2	CH_2	18,A130	CH_2O_2	<i>cyc</i> - H_2CO_2	A283
CH_2^-	CH_2^-	21	CH_2O_3	HC(O)OOH	336
CH_2B	HBCH	A221	CH_2S^+	H_2CS^+	141
CH_2B_2	HBCBH	245,A275	CH_2S	H_2CS	143,A224
CH_2Br^+	H_2CBr^+	147	CH_2S^-	H_2CS^-	149
CH_2Br	H_2CBr	150,A228	CH_2S_2	<i>t</i> - HCSSH	259
CH_2BrCl	$\text{H}_2\text{CCl-Br}$	264	CH_2S_2	<i>c</i> - HCSSH	259
CH_2BrF^+	CH_2FBr^+	261	CH_2Se^+	H_2CSe^+	141
CH_2BrI	$\text{H}_2\text{CBr-I}$	265	CH_2Se	H_2CSe	145,A226
CH_2Br_2^+	CH_2Br_2^+	261	CH_2Si	H_2CSI	138,A223
CH_2Br_2	$\text{H}_2\text{CBr-Br}$	264	CH_2Zn	ZnCH_2	131
CH_2CaNO	HCONHCa	326	CH_2Zn	HZnCH	131
CH_2Cl^+	H_2CCl^+	147	CH_3^+	CH_3^+	124,A213
CH_2Cl	H_2CCl	149,A228	CH_3	CH_3	125,A214
CH_2ClF^+	CH_2FCl^+	261	CH_3^-	CH_3^-	129
CH_2ClI	$\text{H}_2\text{CCl-I}$	264	CH_3Al	AlCH_3	A268
CH_2CIN	$\text{CH}_2=\text{NCI}$	258	CH_3B	$\text{H}_2\text{C}=\text{BH}$	A268
CH_2ClNO_4	$\text{CH}_2\text{ClOONO}_2$	425	CH_3BO^+	BH_3CO^+	312
CH_2ClP	$\text{CH}_2=\text{PCI}$	258	CH_3BaO	BaOCH_3	315
CH_2Cl_2^+	CH_2Cl_2^+	261,A285	CH_3Br^+	CH_3Br^+	239,A274
CH_2Cl_2	$\text{H}_2\text{CCl-Cl}$	263	CH_3Br^+	H_2CBrH^+	A274
$\text{CH}_2\text{Cl}_2\text{O}$	HICl_2OH	339	CH_3BrF	CH_3BrF	324
$\text{CH}_2\text{Cl}_2\text{O}_2$	CHCl_2OOH	368	CH_3Ca	CaCH_3	230,A267
$\text{CH}_2\text{Cl}_2\text{Si}$	$\text{CH}_2=\text{SiCl}_2$	337	CH_3CaO	CaOCH_3	314
CH_2Co	CoCH_2	A219	CH_3CaS	CaSCH_3	315
CH_2Cr	CrCH_2	A218	CH_3Cd	CdCH_3	231,A268
CH_2Cu	CuCH_2	132	CH_3Cl^+	CH_3Cl^+	239,A273
CH_2F^+	H_2CF^+	147	CH_3Cl^+	H_2CClH^+	A274
CH_2F	H_2CF	149	CH_3ClF	CH_3ClF	324
CH_2FI^+	CH_2FI^+	261	CH_3ClO^+	CH_3OCl^+	322
CH_2FO_2	CH_2FO_2	339	CH_3ClO	ClCH_2OH	323
CH_2F_2^+	CH_2F_2^+	260,A284	CH_3ClO_2	CH_2ClOOH	368
CH_2Fe	FeCH_2	131	CH_3ClSi	$\text{CH}_2=\text{SiHCl}$	321
CH_2Fe	HFeCH	132	CH_3ClSi	CH_3SiCl	321
CH_2I	H_2Cl	150	CH_3F^+	CH_3F^+	238
CH_2I_2^+	CH_2I_2^+	262	CH_3F^+	H_2CFH^+	A273
CH_2I_2	$\text{H}_2\text{Cl-I}$	265	CH_3FI	CH_3IF	324
CH_2N^+	HCNH^+	138	CH_3FO^+	CH_3OF^+	322
CH_2N	H_2CN	140	CH_3FO	CH_3OF	322
CH_2N^-	H_2CN^-	143	CH_3I^+	CH_3I^+	239

ENERGY LEVELS OF POLYATOMIC TRANSIENT MOLECULES

379

Formula	Structure/Name	References	Formula	Structure/Name	References
CH ₃ IO	CH ₃ IO	323	CH ₄ OSi	HCH ₃ SiO	363
CH ₃ IO	CH ₃ OI	323	CH ₄ O ₂	CH ₂ (OH) ₂	A325
CH ₃ IO	ICH ₂ OH	323	CH ₄ O ₃	HO ₂ CH ₂ OH	379
CH ₃ K	KCH ₃	230	CH ₄ S ⁺	CH ₃ SH ⁺	310
CH ₃ Mg	MgCH ₃	A267	CH ₄ Si ⁺	CH ₂ =SiH ₂ ⁺	308
CH ₃ N ⁺	CH ₂ NH ⁺	233	CH ₄ Si	CH ₂ =SiH ₂	309,A310
CH ₃ N	CH ₃ N	234,A269	CH ₄ Si	CH ₃ SiH	309
CH ₃ N	CH ₂ NH	234	CH ₄ Zn	CH ₃ ZnH	A309
CH ₃ NO ⁺	CH ₃ NO ⁺	318	CH ₅ BO	H ₂ B=OCH ₃	374
CH ₃ NO ⁺	CH ₂ NOH ⁺	318	CH ₅ BS	H ₂ BSCH ₃	A335
CH ₃ NO ⁺	HCONH ₂ ⁺	318	CH ₆ BN	H ₂ B=NHCH ₃	398
CH ₃ NO	CH ₃ NO	320	CH ₆ OSi	CH ₃ SiH ₂ OH	416
CH ₃ NO	CH ₂ NOH	320,A314	CIN ⁺	ICN ⁺	81
CH ₃ NO ₂	c-CH ₂ (NO)OH	367	CIN	INC	A191
CH ₃ NO ₂	t-CH ₂ (NO)OH	367	CINO ⁺	INCO ⁺	193
CH ₃ NO ₄	CH ₃ OONO ₂	425	CINO	INCO	198
CH ₃ NO ₅	HOCH ₂ OONO ₂	425	CINS ⁺	ISCN ⁺	195
CH ₃ NS ⁺	HCSNH ₂ ⁺	319	CINS	ISCN	199
CH ₃ Na	NaCH ₃	229	Cl ₃	Cl ₃	218
CH ₃ O ⁺	CH ₂ OH ⁺	235,A270	CMgN	MgCN	A164
CH ₃ O	CH ₃ O	236,A271	CMgN	MgNC	62,A164
CH ₃ O	CH ₂ OH	237,A272	CNO ⁺	NCO ⁺	71
CH ₃ O ⁻	CH ₃ O ⁻	242	CNO	CNO	82
CH ₃ OSr	SrOCH ₃	315	CNO	NCO	76,A184
CH ₃ O ₂	CH ₃ O ₂	321	CNO ⁻	NCO ⁻	85
CH ₃ O ₃	HOCH ₂ O ₂	368	CNOSr	SrNCO	182
CH ₃ P ⁺	CH ₂ PH ⁺	233	CNP	PCN	70
CH ₃ S ⁺	CH ₃ S ⁺	235,A270	CNS	NCS	77
CH ₃ S ⁺	CH ₂ SH ⁺	235	CNS ⁻	NCS ⁻	86
CH ₃ S	CH ₃ S	237,A272	CNSr	SrNC	62
CH ₃ S	CH ₂ SH	238	CN ₂	NCN	70,A180
CH ₃ S ⁻	CH ₃ S ⁻	242	CN ₂	CNN	71
CH ₃ SSr	SrSCH ₃	315	CN ₂ O ⁺	ONCN ⁺	186
CH ₃ S ₂	CH ₃ S ₂	322	CN ₂ O	NCNO	188
CH ₃ S ₂ ⁻	CH ₃ S ₂ ⁻	322	CN ₂ S ⁺	NSCN ⁺	A249
CH ₃ Si	CH ₂ SiH	232	CN ₂ S	NSCN	A252
CH ₃ Si	CH ₃ Si	232	CN ₄ ⁺	N ₃ CN ⁺	278
CH ₃ Si ⁻	CH ₂ SiH ⁻	233	CNiO	NiCO	A167
CH ₃ Si ⁻	CH ₃ Si ⁻	233	CNiO ⁻	NiCO ⁻	A171
CH ₃ Si	SiCH ₃	230,A267	COP	PCO	77
CH ₃ Te	CH ₃ Te	238	COPd	PdCO	A167
CH ₃ Zn	ZnCH ₃	230,A267	COPt	PtCO	A167
CH ₄ ⁺	CH ₄ ⁺	227	COS ⁺	OCS ⁺	78,A185
CH ₄ B	H ₂ CBH ₂	A309	COSc	ScCO	A166
CH ₄ B	CH ₃ BH	A310	COSi	SiCO	70
CH ₄ Cd	CH ₃ CdH	A309	COSn	SnCO	A180
CH ₄ Co	CH ₃ CoH	A308	COTa	TaCO	A166
CH ₄ Fe	CH ₃ FeH	306	COTi	TiCO	A166
CH ₄ Ga	CH ₃ GaH	307,A310	COU	CUO	A168
CH ₄ Hg	CH ₃ HgH	A309	COU	UCO	A168
CH ₄ N ⁺	CH ₂ NH ₂ ⁺	310	COV	VCO	A166
CH ₄ Ni	CH ₃ NiH	306	CO ₂ ⁺	CO ₂ ⁺	77,A185
CH ₄ NiO	HNI ₃ OCH ₃	362	CO ₂ ⁻	CO ₂ ⁻	90,A193
CH ₄ NiO	CH ₃ NiOH	362	CO ₂ Ti	OTiCO	A248
CH ₄ O ⁺	CH ₃ OH ⁺	310	CO ₂ U	OUCO	A248

Formula	Structure/Name	References	Formula	Structure/Name	References
CO_3	CO_3	199	C_2HF^+	HCCF^+	158
CO_3^-	CO_3^-	203	C_2HF	$\text{HFC}=\text{C}:^-$	162
CO_3Ti	O_2TiCO	A296	C_2HF^-	$\text{HFC}=\text{C}^-$	168
CO_3U	O_2UCO	A297	C_2HFO	HFCCO	269,A288
CO_4^-	CO_4^-	287	C_2HFe^-	FeCCH^-	A232
CS_2^+	CS_2^+	79,A186	C_2HI^+	HCCI^+	159
CS_2^-	CS_2^-	A194	C_2HMg	MgCCH	A231
CSi_2	Si_2C	67	C_2HN	HCCN	156,A234
CSi_3	Si_3C	181	C_2HNO	HOOCN	269,A287
C_2Al_2	AlCCAl	A242	C_2HNO_2	$\text{O}=\text{C}=\text{C}=\text{NOH}$	A317
C_2B	cyc-BCC	62,A164	C_2HNS^+	HCSCN^+	A287
$\text{C}_2\text{BFN}_2\text{O}_2$	$\text{FB}(\text{NCO})_2$	A341	C_2HNS	HCSCN	A288
C_2BeO_2	$\text{Be}(\text{CO})_2$	A293	C_2HN_2^+	HNCCN^+	268
C_2BrCl^+	CICCBBr^+	191	C_2HO^-	HCCO	156,A234
C_2BrF_3	CF_3CBr	354	C_2HS	HCCS	157,A234
C_2BrN	BrCCN	189	C_2HS	HSCC	157
C_2Br_2^+	C_2Br_2^+	191	C_2HSi	HCCSi	A233
C_2ClF_3^+	$\text{CF}_2=\text{CFCl}^+$	352	C_2HSr	SrCCH	155,A231
C_2ClF_3^-	CF_3CCl	354	C_2H_2^+	HCCH^+	133,A221
C_2ClN	CICCN	189	C_2H_2	$\text{H}_2\text{C}=\text{C}:^-$	137
C_2Cl_2^+	C_2Cl_2^+	190	C_2H_2^-	$\text{H}_2\text{C}=\text{C}^-$	139
$\text{C}_2\text{Cl}_2\text{F}_2^+$	$\text{CF}_2=\text{CCl}_2^+$	352	$\text{C}_2\text{H}_2\text{Al}$	cyc-HC=CHAl	A276
$\text{C}_2\text{Cl}_2\text{O}^+$	Cl_2CCO^+	285	$\text{C}_2\text{H}_2\text{Al}$	HCCAlH	A276
$\text{C}_2\text{Cl}_2\text{O}$	Cl_2CCO	285	$\text{C}_2\text{H}_2\text{Al}$	$(\text{C}_2\text{H}_2)\text{Al}$	A277
$\text{C}_2\text{Cl}_2\text{O}_2^+$	$t-(\text{ClCO})_2^+$	348	$\text{C}_2\text{H}_2\text{B}$	cyc-HC=CHB	A276
C_2Cl_4^+	C_2Cl_4^+	353	$\text{C}_2\text{H}_2\text{B}$	cyc-HC=CBH	A276
C_2F_2^+	C_2F_2^+	190	$\text{C}_2\text{H}_2\text{B}$	HCCBH	A276
C_2F_2	FCCF	197	$\text{C}_2\text{H}_2\text{Be}$	HBeCCH	A275
C_2F_2	$\text{F}_2\text{C}=\text{C}:^-$	197	$\text{C}_2\text{H}_2\text{Br}$	$\text{CHBr}=\text{CH}$	257
C_2F_2^-	$\text{F}_2\text{C}=\text{C}^-$	203	$\text{C}_2\text{H}_2\text{ClF}^+$	$\text{CH}_2=\text{CFCI}^+$	334
$\text{C}_2\text{F}_2\text{N}_2$	$\text{F}_2\text{C}=\text{C}=\text{N}=\text{N}$	347	$\text{C}_2\text{H}_2\text{ClF}^+$	$c\text{-CHF}=\text{CHCl}^+$	334
$\text{C}_2\text{F}_2\text{O}_2^+$	$t-(\text{FCO})_2^+$	348	$\text{C}_2\text{H}_2\text{ClF}^+$	$t\text{-CHF}=\text{CHCl}^+$	334
C_2F_3	C_2F_3	287	$\text{C}_2\text{H}_2\text{ClN}^+$	CH_2ClCN^+	331
$\text{C}_2\text{F}_3\text{O}$	CF_3CO	A322	$\text{C}_2\text{H}_2\text{Cl}_2^+$	$\text{CH}_2=\text{CCl}_2^+$	335
C_2F_4^+	C_2F_4^+	351	$\text{C}_2\text{H}_2\text{Cl}_2^+$	$c\text{-CHCl}=\text{CHCl}^+$	335
C_2F_4	CF_3CF	354	$\text{C}_2\text{H}_2\text{Cl}_2^+$	$t\text{-CHCl}=\text{CHCl}^+$	336
C_2F_5	C_2F_5	371	$\text{C}_2\text{H}_2\text{F}$	$t\text{-CHF}=\text{CH}$	257
$\text{C}_2\text{F}_5\text{P}$	$\text{CF}_2=\text{PCF}_3$	382	$\text{C}_2\text{H}_2\text{FN}^+$	CH_2FCN^+	330
C_2Fe	FeCC	A164	$\text{C}_2\text{H}_2\text{FO}$	CH_2CFO	A316
C_2Fe^-	FeCC^-	A164	$\text{C}_2\text{H}_2\text{FO}^-$	CH_2COF^-	337
C_2DFe	FeCCD	A232	$\text{C}_2\text{H}_2\text{F}_2^+$	$\text{CH}_2=\text{CF}_2^+$	332
C_2H	HCC	32,A142	$\text{C}_2\text{H}_2\text{F}_2^+$	$c\text{-CHF}=\text{CHF}^+$	333
C_2H^-	HCC $^-$	37	$\text{C}_2\text{H}_2\text{F}_2^+$	$t\text{-CHF}-\text{CHF}^+$	333
C_2HAl	HCCAl	A233	$\text{C}_2\text{H}_2\text{Fe}$	HFeCCH	245
C_2HB	HBCC	A232	$\text{C}_2\text{H}_2\text{Ga}$	$(\text{C}_2\text{H}_2)\text{Ga}$	A277
C_2HB	HCCB	A232	$\text{C}_2\text{H}_2\text{In}$	$(\text{C}_2\text{H}_2)\text{In}$	A277
C_2HB	cyc-HBC ₂	A232	$\text{C}_2\text{H}_2\text{Li}$	LiC_2H_2	244
C_2HBe	BeCCH	A231	$\text{C}_2\text{H}_2\text{N}$	H_2CCN	250,A280
C_2HBr^+	HCCBr^+	158	$\text{C}_2\text{H}_2\text{N}$	H_2CNC	250
C_2HCa	CaCCH	155,A231	$\text{C}_2\text{H}_2\text{N}^-$	H_2CCN^-	252
C_2HCl^+	HCCCl^+	158	$\text{C}_2\text{H}_2\text{N}^-$	H_2CNC^-	252
$\text{C}_2\text{HCIF}_2^+$	$\text{CHCl}=\text{CF}_2^+$	340	$\text{C}_2\text{H}_2\text{N}_2^+$	$\text{HN}=\text{CHCN}^+$	327
C_2HClO	HCICCO	269	$\text{C}_2\text{H}_2\text{N}_2^+$	$\text{H}_2\text{C}=\text{NCN}^+$	327
C_2HCl_3^+	$\text{CHCl}=\text{CCl}_2^+$	341	$\text{C}_2\text{H}_2\text{N}_2$	$c\text{-HN}=\text{CHCN}$	327
C_2HF_3^+	$\text{CHF}=\text{CF}_2^+$	340	$\text{C}_2\text{H}_2\text{N}_2$	$t\text{-HN}=\text{CHCN}$	328

Formula	Structure/Name	References	Formula	Structure/Name	References
C ₂ H ₂ N ₂	H ₂ C=NCN	328	C ₂ H ₄ Li	C ₂ H ₄ Li	360
C ₂ H ₂ N ₂ O ₂	HON=CHCNO	381	C ₂ H ₄ Li ₂	1,2-C ₂ H ₄ Li ₂	377
C ₂ H ₂ N ₂ O ₂	HON=CHNCO	381	C ₂ H ₄ N ⁺	CH ₃ CNH ⁺	362
C ₂ H ₂ Ni	(C ₂ H ₂)Ni	245	C ₂ H ₄ O ⁺	CH ₂ =CHOH ⁺	362,A324
C ₂ H ₂ Ni	NiC=CH ₂	245	C ₂ H ₄ O	CH ₂ =CHOH	363
C ₂ H ₂ O ⁺	H ₂ CCO ⁺	250,A280	C ₂ H ₄ O ₃	1,2,3-cyc-C ₂ H ₄ O ₃	422,A366
C ₂ H ₂ O	HCCOH	255	C ₂ H ₄ O ₃	1,2,4-cyc-C ₂ H ₄ O ₃	422,A367
C ₂ H ₂ OS	CHOCHS	330	C ₂ H ₄ O ₃	c-HCOOCH ₂ OH	423
C ₂ H ₂ O ₂ ⁺	(HCO) ₂ ⁺	329	C ₂ H ₄ O ₃	t-HCOOCH ₂ OH	423
C ₂ H ₂ O ₂	HOC \equiv COH	A315	C ₂ H ₄ S ⁺	CH ₃ CHS ⁺	363
C ₂ H ₂ O ₃	(HCO) ₂ O	368	C ₂ H ₄ S	CH ₃ CHS	363,A324
C ₂ H ₂ S ⁺	H ₂ CCS ⁺	253	C ₂ H ₄ Si	cyc-C ₂ SiH ₄	A324
C ₂ H ₂ S	H ₂ CCS	255,A282	C ₂ H ₅ ⁺	C ₂ H ₅ ⁺	359
C ₂ H ₂ S	HCCSH	256	C ₂ H ₅	C ₂ H ₅	359,A323
C ₂ H ₂ S	cyc-C ₂ H ₂ S	256	C ₂ H ₅ BrS	CH ₃ SBrCH ₂	418
C ₂ H ₂ S ₂	HS-CH=C=S	331	C ₂ H ₅ CIS	CH ₃ SCICH ₂	418
C ₂ H ₂ Se	H ₂ CCSe	A283	C ₂ H ₅ IO	C ₂ H ₅ IO	A364
C ₂ H ₂ Si	cyc-HC=CHSi	A278	C ₂ H ₅ IO	C ₂ H ₅ OI	A364
C ₂ H ₂ Si	HCCSiH	A279	C ₂ H ₅ IO ₂	C ₂ H ₅ IO ₂	A364
C ₂ H ₂ Si	H ₂ C=C:Si:	A279	C ₂ H ₅ IS	CH ₃ SICH ₂	418
C ₂ H ₂ Si	cyc-H ₂ SiCC	A279	C ₂ H ₅ N	c-CH ₃ CH=NH	374,A336
C ₂ H ₃ ⁺	C ₂ H ₃ ⁺	232,A268	C ₂ H ₅ N	t-CH ₃ CH=NH	374
C ₂ H ₃	C ₂ H ₃	232,A269	C ₂ H ₅ N	C ₂ H ₃ NH ₂	375,A336
C ₂ H ₃ As	HCCAsH ₂	A313	C ₂ H ₅ N	H ₂ C=NCH ₃	375
C ₂ H ₃ As	CH ₃ CAs	A313	C ₂ H ₅ NO ₂	c-CH ₃ CH(NO)OH	424
C ₂ H ₃ CaO ₂	CH ₃ COOCa	379	C ₂ H ₅ NO ₄	C ₂ H ₅ OONO ₂	425
C ₂ H ₃ Cl ⁺	CH ₂ =CHCl ⁺	319	C ₂ H ₅ O	C ₂ H ₅ O	376,A337
C ₂ H ₃ ClO	CH ₃ OCCl	366	C ₂ H ₅ O	CH ₃ CHOH	376
C ₂ H ₃ F ⁺	CH ₂ =CHF ⁺	319	C ₂ H ₅ O	HOCH ₂ CH ₂	376,A337
C ₂ H ₃ F ₂	CH ₃ CF ₂	A327	C ₂ H ₅ O ₂	C ₂ H ₅ O ₂	420
C ₂ H ₃ N ⁺	CH ₃ CN ⁺	312	C ₂ H ₅ O ₃	HOCH ₂ CH ₂ O ₂	424
C ₂ H ₃ N ⁺	CH ₃ NC ⁺	313	C ₂ H ₅ S ⁺	CH ₃ SCH ₂ ⁺	A336
C ₂ H ₃ N ⁺	CH ₂ =C=NH ⁺	313	C ₂ H ₅ S	C ₂ H ₅ S	376,A337
C ₂ H ₃ N ⁺	cyc-(HC=NCH ₂) ⁺	A311	C ₂ H ₆ Ge	(CH ₃) ₂ Ge	400
C ₂ H ₃ N	CH ₂ =C=NH	316,A312	C ₂ H ₆ Hg	C ₂ H ₅ HgH	A357
C ₂ H ₃ N	HCNCH ₂	A312	C ₂ H ₆ N ₂ O	CH ₃ NHCH ₂ NO(A)	424
C ₂ H ₃ N	cyc-(HC=NCH ₂)	A313	C ₂ H ₆ N ₂ O	CH ₃ NHCH ₂ NO(B)	424
C ₂ H ₃ NO	CH ₃ CNO	366	C ₂ H ₆ OSi	(CH ₃) ₂ SiO	416
C ₂ H ₃ NO	CH ₃ OCN	A326	C ₂ H ₆ OSi	CH ₂ =Si(OH)CH ₃	417
C ₂ H ₃ NO	HOCH ₂ CN	366	C ₂ H ₆ OSi	CH ₃ OSiCH ₃	417
C ₂ H ₃ NO ₂	CH ₃ ONCO	380	C ₂ H ₆ OSi	CH ₃ OSiH=CH ₂	417
C ₂ H ₃ O	CH ₃ CO	317	C ₂ H ₆ O ₂ Si	(CH ₃ O) ₂ Si	421
C ₂ H ₃ O	CII ₂ CHO	317,A313	C ₂ H ₆ O ₂ Si	CH ₃ SiOOCH ₃	421
C ₂ H ₃ O ⁻	CH ₂ CHO ⁻	320,A314	C ₂ H ₆ O ₂ Si	CH ₃ SiHOHCHO	A365
C ₂ H ₃ O ₂ Sr	CH ₃ COOSr	380	C ₂ H ₆ O ₃	HOCH ₂ CH ₂ OOH	424
C ₂ H ₃ O ₃	CH ₃ COO ₂	380,A339	C ₂ H ₆ O ₄	CH ₃ O ₄ CH ₃	424
C ₂ H ₃ P ⁺	CH ₃ CP ⁺	314	C ₂ H ₆ Si	(CH ₃) ₂ Si	399
C ₂ H ₃ P	CH ₃ CP	316	C ₂ H ₆ Si	CH ₃ SiH=CH ₂	399
C ₂ H ₄ ⁺	C ₂ H ₄ ⁺	307	C ₂ H ₆ Sn	(CH ₃) ₂ Sn	400
C ₂ H ₄ Al	AlC ₂ H ₄	361	C ₂ H ₆ Zn ⁺	(CH ₃) ₂ Zn ⁺	A357
C ₂ H ₄ F	FCH ₂ CH ₂	364,A325	C ₂ H ₇ ⁺	C ₂ H ₇ ⁺	383
C ₂ H ₄ Fe	HFeC ₂ H ₃	360	C ₂ H ₇ ⁺	br-C ₂ H ₇ ⁺	383
C ₂ H ₄ FeO	CH ₂ =CHFeOH	378	C ₂ H ₇ BO	H ₂ B=OC ₂ H ₅	408
C ₂ H ₄ FeO	cyc-C ₂ H ₄ OF	378	C ₂ H ₇ BS	H ₂ BSC ₂ H ₅	A361
C ₂ H ₄ In	InC ₂ H ₄	361	C ₂ H ₈ BN	H ₂ B=N(CH ₃) ₂	399

Formula	Structure/Name	References	Formula	Structure/Name	References
C ₂ H ₈ OSi	(CH ₃) ₂ SiHOH	417	C ₃ HF ₆	FeC=C=CH	A286
C ₂ I ₂ ⁺	C ₂ I ₂ ⁺	192	C ₃ HF ₆	FeCCCH	A286
C ₂ N ⁺	CNC ⁺	67	C ₃ HF ₆ ⁻	FeC=C=CH ⁻	A286
C ₂ N	CCN	68,A173	C ₃ HF ₆ ⁻	FeCCCH ⁻	A286
C ₂ N	CNC	69	C ₃ HN ⁺	HCCCN ⁺	266
C ₂ N ₂ ⁺	NCCN ⁺	183	C ₃ HN ⁺	HCCNC ⁺	267
C ₂ N ₂ ⁺	CNCN ⁺	184	C ₃ HN	HNCCC	267
C ₂ N ₂	CNCN	185,A248	C ₃ HN	HCCNC	267
C ₂ N ₂	CNNC	185	C ₃ HF ₃ S	CF ₃ H(<i>cyc</i> -CCS)	381
C ₂ N ₂ O ⁺	NCCNO ⁺	A295	C ₃ HO	HCCCO	268,A286
C ₂ N ₂ O ⁺	NCNCO ⁺	279	C ₃ HS	HCCCS	A287
C ₂ N ₂ O	NCCNO	283,A297	C ₃ H ₂ ⁺	<i>cyc</i> -C ₃ H ₂ ⁺	248
C ₂ N ₂ O	NCNCO	283	C ₃ H ₂	HCCCH	249,A278
C ₂ N ₂ O ₂ ⁺	ONCCNO ⁺	A320	C ₃ H ₂	<i>cyc</i> -C ₃ H ₂	248
C ₂ N ₂ O ₂	ONCCNO	A321	C ₃ H ₂	H ₂ C=C=C:	248,A277
C ₂ N ₂ S ⁺	S(CN) ₂ ⁺	280	C ₃ H ₂	HCCH=C:	249
C ₂ N ₂ S ⁺	NCNCS ⁺	280	C ₃ H ₂ ⁻	H ₂ C-C-C:-	A280
C ₂ N ₂ S	NCNCS	284	C ₃ H ₂ N ⁺	HCCCCNH ⁺	326,A315
C ₂ N ₂ S ₂ ⁺	(SCN) ₂ ⁺	346	C ₃ H ₂ O ⁺	H ₂ C=C=C=O ⁺	327
C ₂ N ₂ Se ⁺	Se(CN) ₂ ⁺	281	C ₃ H ₂ O	HC≡CC-OH	328
C ₂ O	CCO	69,A179	C ₃ H ₂ O	H ₂ C=C=C=O	328
C ₂ O ⁻	CCO ⁻	76,A183	C ₃ H ₂ O ₂	<i>t</i> -HCCCHO	A327
C ₂ O ₂ ⁺	<i>t</i> -OCCO ⁺	186	C ₃ H ₂ O ₂	<i>c</i> -HCCCHO	A327
C ₂ O ₂ ⁻	<i>t</i> -OCCO ⁻	190	C ₃ H ₂ O ₂	HCC(<i>cyc</i> -CHO)	A328
C ₂ O ₂ Si	Si(CO) ₂	283	C ₃ H ₂ O ₃	<i>cyc</i> -(HOC=COHC)=O	A340
C ₂ S	CCS	70	C ₃ H ₂ S	H ₂ C=C=C=S	329
C ₂ S ₂	SCCS	188	C ₃ H ₂ Se	H ₂ C=C=C=Se	329
C ₂ S ₃	S(CS) ₂	284	C ₃ H ₂ Se	HCC-CHSe	329
C ₂ Si	SiCC	66,A170	C ₃ H ₃ ⁺	CH ₂ CCH ⁺	312
C ₂ Si ₂	Si ₂ C ₂	181,A245	C ₃ H ₃	CH ₂ CCH	312,A311
C ₂ Si ₃	Si ₃ C ₂	A293	C ₃ H ₃ ⁻	CH ₂ CCH ⁻	A311
C ₂ Y	YCC	A164	C ₃ H ₃ ⁻	CH ₃ CC ⁻	A312
C ₃	C ₃	65,A168	C ₃ H ₃ Br ⁺	CH ₃ CCBr ⁺	365
C ₃ ⁻	C ₃ ⁻	68,A173	C ₃ H ₃ Cl ⁺	CH ₃ CCCl ⁺	365
C ₃ BrN ⁺	BrCCCN ⁺	282	C ₃ H ₃ N ⁺	C ₂ H ₃ CN ⁺	A325
C ₃ ClN ⁺	ClCCCN ⁺	281	C ₃ H ₃ N	H ₂ C=(<i>cyc</i> -CN=CH)	A326
C ₃ Cl ₂	<i>cyc</i> -C ₃ Cl ₂	A299	C ₃ H ₄ ⁺	H ₂ CCCH ₂ ⁺	362,A323
C ₃ Cl ₂	ClCCCCl	A299	C ₃ H ₄ Li	CH ₃ CCHLi	377
C ₃ Cl ₂ ⁺	Cl ₂ C=C=C:	A299	C ₃ H ₄ NO ₂	CH ₃ C(=NO)CHO	A367
C ₃ FN ⁺	FCCCN ⁺	281	C ₃ H ₄ O	<i>cyc</i> -(H ₂ COOC)=CH ₂	379
C ₃ F ₂	<i>cyc</i> -C ₃ F ₂	A299	C ₃ H ₄ O	CH ₃ CCOH	A338
C ₃ F ₂	FCCCCF	A299	C ₃ H ₄ S	HCH ₃ (<i>cyc</i> -CCS)	379
C ₃ F ₂	F ₂ C=C=C:	A299	C ₃ H ₅ ⁺	CH ₂ CHCH ₂ ⁺	372
C ₃ F ₂ O	F ₂ C=C=C=O	347	C ₃ H ₅ ⁺	<i>cyc</i> -C ₃ H ₅ ⁺	372
C ₃ F ₂ O	<i>cyc</i> -(CF=FCF)=O	347	C ₃ H ₅	CH ₂ CHCH ₂	372,A334
C ₃ F ₆ ⁺	C ₃ F ₆ ⁺	405	C ₃ H ₅ ⁻	CH ₂ CHCH ₂ ⁻	A335
C ₃ F ₇	<i>n</i> -C ₃ F ₇	405	C ₃ H ₅ N	H ₂ C=CH-CH=NH	408
C ₃ F ₇	<i>i</i> -C ₃ F ₇	406	C ₃ H ₅ N	H ₂ C=C=NCH ₃	408
C ₃ Fe	FeCCC	A242	C ₃ H ₅ N	H ₂ C=CHN=CH ₂	409
C ₃ Fe ⁻	FeCCC ⁻	A242	C ₃ H ₅ N	<i>cyc</i> -C ₃ H ₅ N	409
C ₃ H	HC ₃	156,A233	C ₃ H ₆ O	<i>t</i> -CH ₃ C-OCH ₃	415
C ₃ H	<i>cyc</i> -HC ₃	156,A233	C ₃ H ₆ O	<i>c</i> -CH ₃ C-OCH ₃	415
C ₃ HCl	<i>cyc</i> -C ₃ HCl	A288	C ₃ H ₆ S	(CH ₃) ₂ CS	415
C ₃ HCl	HCCCCl	A288	C ₃ H ₇ ⁺	1-C ₃ H ₇ ⁺	383
C ₃ HCl	HCIC=C=C:	A288	C ₃ H ₇ ⁺	2-C ₃ H ₇ ⁺	383

Formula	Structure/Name	References	Formula	Structure/Name	References
C ₃ H ₇	CH ₃ CH ₂ CH ₂	383	C ₄ H ₆ Ge	cyc-C ₄ H ₅ GeH(1,3)	A360
C ₃ H ₇	(CH ₃) ₂ CH	383	C ₄ H ₆ Ge	cyc-C ₄ H ₅ GeH(1,4)	A360
C ₃ H ₇ N	C ₂ H ₃ NHCH ₃	409	C ₄ H ₆ S	(CH ₃) ₂ (cyc-CCS)	419
C ₃ H ₇ O	n-C ₃ H ₇ O	416	C ₄ H ₆ Si	cyc-C ₄ H ₄ SiH ₂	A358
C ₃ H ₇ O	(CH ₃) ₂ CHO	416	C ₄ H ₆ Si	cyc-C ₄ H ₆ Si:	A358
C ₃ H ₇ O ₂	(CH ₃) ₂ CHO ₂	420	C ₄ H ₆ Si	cyc-C ₄ H ₅ SiH(1,3)	A358
C ₃ H ₈ O ₂ Si	(CH ₃) ₂ SiOHCHO	A365	C ₄ H ₆ Si	cyc-C ₄ H ₅ SiH(1,4)	A359
C ₃ H ₈ Si ⁺	(CH ₃) ₂ Si-CH ₂ ⁺	400	C ₄ H ₇ ⁺	H ₂ CC(CH ₃)CH ₂ ⁺	A343
C ₃ H ₈ Si	(CH ₃) ₂ Si=CH ₂	401	C ₄ H ₇ ⁺	cyc-C ₄ H ₇ ⁺	A343
C ₃ H ₉ Al	(CH ₃) ₃ Al	400	C ₄ H ₇	CH ₃ CHCH=CH ₂	A343
C ₃ H ₉ NSi	(CH ₃) ₃ SiN	410	C ₄ H ₇	H ₂ CC(CH ₃)CH ₂	A343
C ₃ IN ⁺	ICCCN ⁺	282	C ₄ H ₇ ⁻	H ₂ CC(CH ₃)CH ₂ ⁻	A344
C ₃ N	CCCN	183,A247	C ₄ H ₇ N	(CH ₃) ₂ C=C=NH	412
C ₃ N ₂	C(CN) ₂	278,A295	C ₄ H ₇ N	CH ₃ CC-NHCH ₃	412
C ₃ N ₂ O ⁺	CO(CN) ₂ ⁺	343	C ₄ H ₈ Si	cyc-C ₂ H ₂ Si(CH ₃) ₂	A358
C ₃ N ₂ O ₂	(CN) ₂ COO	A330	C ₄ H ₉ ⁺	t-C ₄ H ₉ ⁺	384
C ₃ O	CCCO	186	C ₄ H ₉	n-C ₄ H ₉	384,A344
C ₃ O ₂ ⁺	C ₃ O ₂ ⁺	279	C ₄ H ₉	i-C ₄ H ₉	384
C ₃ S	CCCS	186,A249	C ₄ H ₉	t-C ₄ H ₉	384
C ₃ Si ₂	SiCCCSi	278,A293	C ₄ H ₉ O ₂	t-C ₄ H ₉ O ₂	421
C ₄	C ₄	181,A245	C ₄ H ₁₀ Si	(CH ₃) ₂ Si=CHCH ₃	401
C ₄ ⁻	C ₄ ⁻	183,A246	C ₄ I ₂ ⁺	I(CC) ₂ I ⁺	345
C ₄ Br ₂ ⁺	Br(CC) ₂ Br ⁺	345	C ₄ N ₂ ⁺	NCC≡CCN ⁺	343,A319
C ₄ Cl ₂ ⁺	Cl(CC) ₂ Cl ⁺	345	C ₄ N ₂	NCCCNC	343,A320
C ₄ F ₂ ⁺	F(CC) ₂ F ⁺	344	C ₄ N ₂	CNCCNC	343
C ₄ F ₆ O	CF ₃ CCOCF ₃	418	C ₄ N ₂ O	(CN) ₂ CCO	A330
C ₄ F ₆ O	(CF ₃) ₂ (cyc-CCO)	418	C ₄ O	C ₄ O	278,A294
C ₄ Fe ⁻	FeC ₄ ⁻	A292	C ₄ OS	C ₄ OS	344
C ₄ H	C ₄ H	266,A286	C ₄ O ₂	C ₄ O ₂	344
C ₄ HBr ⁺	H(CC) ₂ Br ⁺	339	C ₄ S	C ₄ S	278,A294
C ₄ HCl ⁺	H(CC) ₂ Cl ⁺	339	C ₄ S ₂	C ₄ S ₂	344
C ₄ HF ⁺	H(CC) ₂ F ⁺	339	C ₄ Si	C ₄ Si	277,A293
C ₄ HFe ⁻	FeCCCC ⁻	A316	C ₄ Si ₂	SiC ₄ Si	A318
C ₄ HI ⁺	H(CC) ₂ I ⁺	340	C ₅ ⁺	cyc-C ₅ ⁺	277
C ₄ HO	HCCCCO	A316	C ₅	C ₅	277,A292
C ₄ HS	HCCCCS	A317	C ₅ ⁻	C ₅ ⁻	278,A293
C ₄ H ₂ ⁺	C ₄ H ₂ ⁺	325	C ₅ Br ₄	cyc-C ₅ Br ₄	A357
C ₄ H ₂	H ₂ CCCC:	326,A315	C ₅ Br ₄ O ₂	cyc-C ₅ Br ₄ O-1-O	A366
C ₄ H ₄ ⁺	cyc-C ₄ H ₄ ⁺	377,A337	C ₅ Cl ₄	cyc-C ₅ Cl ₄	A357
C ₄ H ₄ ⁺	H ₂ C=(cyc-C ₃ H ₂) ⁺	377	C ₅ Cl ₄ O ₂	cyc-C ₅ Cl ₄ O-1-O	A365
C ₄ H ₄	cyc-C ₄ H ₄	378,A338	C ₅ F ₄ ⁺	CF ₃ (CC) ₂ F ⁺	407
C ₄ H ₄	H ₂ C=(cyc-C ₃ H ₂)	377	C ₅ F ₆ O	(CF ₃) ₂ (cyc-CCO)	418
C ₄ H ₄	C ₂ H ₃ CH=C:	A338	C ₅ F ₈ O	CF ₃ (cyc-CCO)C ₂ F ₅	419
C ₄ H ₄ ⁻	C ₂ H ₃ CH=C:	A338	C ₅ H	C ₅ H	A316
C ₄ H ₄ CaN	Ca(C ₄ H ₄ N)	410	C ₅ HN ⁺	H(CC) ₂ CN ⁺	369,A328
C ₄ H ₄ CdN	Cd(C ₄ H ₄ N)	412	C ₅ H ₂ ⁺	C ₅ H ₂ ⁺	A327
C ₄ H ₄ MgN	Mg(C ₄ H ₄ N)	410	C ₅ H ₂	HC ₅ H	A327
C ₄ H ₄ NSr	Sr(C ₄ H ₄ N)	411	C ₅ H ₃ Br ⁺	CH ₃ (CC) ₂ Br ⁺	406
C ₄ H ₄ NZn	Zn(C ₄ H ₄ N)	411	C ₅ H ₃ Cl ⁺	CH ₃ (CC) ₂ Cl ⁺	406
C ₄ H ₅ N	H ₂ C=(cyc-CN=CCH ₃)	A361	C ₅ H ₃ F ₂ N ⁺	cyc-C ₅ H ₃ F ₂ N ⁺	412
C ₄ H ₆ ⁺	t-H ₂ C=CH-CH=CH ₂ ⁺	A342	C ₅ H ₃ N	cyc-C ₅ H ₃ N	412
C ₄ H ₆	C(CH ₂) ₃	A343	C ₅ H ₄ ⁺	CH ₃ (CC) ₂ H ⁺	385
C ₄ H ₆ ⁻	C(CH ₂) ₃ ⁻	A343	C ₅ H ₄	cyc-C ₅ H ₄	385
C ₄ H ₆ Ge	cyc-C ₄ H ₄ GeH ₂	A359	C ₅ H ₄ Cl	cyc-C ₅ H ₄ Cl	407
C ₄ H ₆ Ge	cyc-C ₄ H ₆ Ge:	A359	C ₅ H ₄ F	cyc-C ₅ H ₄ F	407

Formula	Structure/Name	References	Formula	Structure/Name	References
C ₅ H ₄ O	(cyc-C ₅ H ₄)O	419	C ₆ H ₄ ⁻	C ₆ H ₄ ⁻	387
C ₅ H ₄ O ₂	cyc-C ₅ H ₄ O-1-O	422,A365	C ₆ H ₄ Br	o-C ₆ H ₄ Br	428
C ₅ H ₅	cyc-C ₅ H ₅	385,A344	C ₆ H ₄ Br	m-C ₆ H ₄ Br	428
C ₅ H ₅ Ca	CaC ₅ H ₅	402,A360	C ₆ H ₄ Br	p-C ₆ H ₄ Br	428
C ₅ H ₅ Cd	CdC ₅ H ₅	403	C ₆ H ₄ Cl	o-C ₆ H ₄ Cl	427
C ₅ H ₅ Mg	MgC ₅ H ₅	401	C ₆ H ₄ Cl	m-C ₆ H ₄ Cl	427
C ₅ H ₅ Sr	SrC ₅ H ₅	402	C ₆ H ₄ Cl	p-C ₆ H ₄ Cl	427
C ₅ H ₅ Zn	ZnC ₅ H ₅	402	C ₆ H ₄ Cl ₂ ⁺	1,3-C ₆ H ₄ Cl ₂ ⁺	431
C ₅ H ₆ ⁺	cyc-C ₅ H ₆ ⁺	A345	C ₆ H ₄ Cl ₂ ⁺	1,4-C ₆ H ₄ Cl ₂ ⁺	431
C ₅ H ₆ Se	(CH ₂) ₃ C=C=Se	419	C ₆ H ₄ F	o-C ₆ H ₄ F	427
C ₅ H ₆ Si ⁺	C ₅ SiH ₆ ⁺	404	C ₆ H ₄ F	m-C ₆ H ₄ F	427
C ₅ H ₆ Si	C ₅ SiH ₆	404	C ₆ H ₄ F	p-C ₆ H ₄ F	427
C ₅ H ₆ Si	C ₅ SiH ₆ (Dewar)	405	C ₆ H ₄ F ₂ ⁺	1,2-C ₆ H ₄ F ₂ ⁺	429
C ₅ H ₁₁	n-C ₅ H ₁₁	386,A346	C ₆ H ₄ F ₂ ⁺	1,3-C ₆ H ₄ F ₂ ⁺	429
C ₅ H ₁₁	(CH ₃) ₃ CCH ₂	386	C ₆ H ₄ F ₂ ⁺	1,4-C ₆ H ₄ F ₂ ⁺	430,A369
C ₅ N	C ₅ N	A319	C ₆ H ₄ F ₂ O ⁺	2,5-F ₂ C ₆ H ₃ OH ⁺	444
C ₅ N ₂	C ₅ N ₂	A330	C ₆ H ₄ F ₂ O ⁺	3,5-F ₂ C ₆ H ₃ OH ⁺	444
C ₅ O	C ₅ O	A320	C ₆ H ₄ I	o-C ₆ H ₄ I	428
C ₅ OS	C ₅ OS	370	C ₆ H ₄ N	(cyc-C ₅ H ₄)CN	413
C ₅ O ₂	C ₅ O ₂	370	C ₆ H ₄ N ₂ O ₂	1,2-C ₆ H ₄ (NO) ₂	446
C ₅ S	C ₅ S	343	C ₆ H ₄ O	2,4-C ₆ H ₃ OH	426
C ₅ S ₂	C ₅ S ₂	370	C ₆ H ₄ S ₂	p-C ₆ H ₄ S ₂	422
C ₆	C ₆	342,A317	C ₆ H ₅ ⁺	C ₆ H ₅ ⁺	387
C ₆ ⁻	C ₆ ⁻	342,A318	C ₆ H ₅	C ₆ H ₅	388,A346
C ₆ Br ₃ F ₃ ⁺	sym-C ₆ F ₃ Br ₃ ⁺	439	C ₆ H ₅ ⁻	C ₆ H ₅ ⁻	388
C ₆ ClF ₅ ⁺	C ₆ F ₅ Cl ⁺	439	C ₆ H ₅ Cl ⁺	C ₆ H ₅ Cl ⁺	428,A368
C ₆ Cl ₃ F ₃ ⁺	sym-C ₆ F ₃ Cl ₃ ⁺	439	C ₆ H ₅ F ⁺	C ₆ H ₅ F ⁺	428
C ₆ F ₅ N	C ₆ F ₅ N:	442,A371	C ₆ H ₅ N	C ₆ H ₅ N	442,A370
C ₆ F ₅ N	bicyc-C ₆ F ₅ N	A361	C ₆ H ₅ N	3-CH(cyc-C ₅ H ₄ N)	413
C ₆ F ₆ ⁺	CF ₃ (CC ₂)CF ₃ ⁺	307	C ₆ H ₅ N	cyc-C ₆ H ₅ N	413
C ₆ F ₆ ⁺	C ₆ F ₆ ⁺	438	C ₆ H ₅ N ⁻	C ₆ H ₅ N ⁻	442,A371
C ₆ H	C ₆ H	369,A328	C ₆ H ₅ O	C ₆ H ₅ O	443,A372
C ₆ HF ₅ ⁺	C ₆ HF ₅ ⁺	437	C ₆ H ₅ O ⁻	C ₆ H ₅ O ⁻	446
C ₆ HF ₅ O ⁺	C ₆ F ₅ OH ⁺	445	C ₆ H ₅ S	C ₆ H ₅ S	443
C ₆ HN ⁺	HC ₅ CN ⁺	A340	C ₆ H ₆ ⁺	C ₆ H ₆ ⁺	388,A347
C ₆ H ₂ ⁺	C ₆ H ₂ ⁺	380,A339	C ₆ H ₆ ⁺	CH ₃ (CC ₂) ₂ CH ₂ ⁺	390
C ₆ H ₂	HC ₆ H	A340	C ₆ H ₆ F	C ₆ H ₆ F	440
C ₆ H ₂ F ₄ ⁺	1,2,3,4-C ₆ H ₂ F ₄ ⁺	435	C ₆ H ₆ Li	LiC ₆ H ₆	440
C ₆ H ₂ F ₄ ⁺	1,2,3,5-C ₆ H ₂ F ₄ ⁺	436	C ₆ H ₆ N	C ₆ H ₅ NH	443
C ₆ H ₂ F ₄ ⁺	1,2,4,5-C ₆ H ₂ F ₄ ⁺	437	C ₆ H ₆ O ⁺	C ₆ H ₅ OH ⁺	444,A373
C ₆ H ₂ F ₄ O ⁺	2,3,5,6-F ₄ C ₆ HOH ⁺	445	C ₆ H ₇	CH ₃ (cyc-C ₅ H ₄)	391
C ₆ H ₃ ClF ₂ ⁺	1,3,5-C ₆ H ₃ ClF ₂ ⁺	434	C ₆ H ₇ Ca	CaC ₅ H ₄ CH ₃	403
C ₆ H ₃ Cl ₂ F ⁺	1,3,5-C ₆ H ₃ Cl ₂ F ⁺	434	C ₆ H ₇ Cd	CdC ₅ H ₄ CH ₃	404
C ₆ H ₃ Cl ₃ ⁺	1,3,5-C ₆ H ₃ Cl ₃ ⁺	434	C ₆ H ₇ Mg	MgC ₅ H ₄ CH ₃	403
C ₆ H ₃ F ₂ N	2,6-C ₆ H ₃ F ₂ N:	A371	C ₆ H ₇ Zn	ZnC ₅ H ₄ CH ₃	404
C ₆ H ₃ F ₂ N	bicyc-C ₆ H ₃ F ₂ N	A361	C ₆ H ₇ N ⁺	C ₆ H ₅ NH ₂ ⁺	443,A372
C ₆ H ₃ F ₃ ⁺	1,2,3-C ₆ H ₃ F ₃ ⁺	432	C ₆ H ₈ ⁺	t-CH ₂ (CH) ₄ CH ₂ ⁺	391
C ₆ H ₃ F ₃ ⁺	1,2,4-C ₆ H ₃ F ₃ ⁺	432	C ₆ H ₈ Si	1-CH ₃ C ₅ SiH ₅	405
C ₆ H ₃ F ₃ ⁺	1,3,5-C ₆ H ₃ F ₃ ⁺	433	C ₆ H ₁₀	(CH ₃) ₃ CCH=C:	A348
C ₆ H ₃ F ₃ O ⁺	2,3,4-F ₃ C ₆ H ₂ OH ⁺	445	C ₆ H ₁₀	(CH ₃) ₃ CCH=C ⁻	A348
C ₆ H ₃ F ₃ O ⁺	2,4,5-F ₃ C ₆ H ₂ OH ⁺	445	C ₆ N ₂ ⁺	NC(CC) ₂ CN ⁺	414,A341
C ₆ H ₃ N ⁺	CH ₃ (CC) ₂ CN ⁺	413	C ₆ O	C ₆ O	370,A330
C ₆ H ₄ ⁺	C ₆ H ₄ ⁺	386	C ₆ O ₂ S ₄	C ₆ S ₄ O ₂	426
C ₆ H ₄	o-C ₆ H ₄	386	C ₆ S ₆	C ₆ S ₆ (A)	426
C ₆ H ₄	m-C ₆ H ₄	A346	C ₆ S ₆	C ₆ S ₆ (B)	427

Formula	Structure/Name	References	Formula	Structure/Name	References
C ₇	C ₇	369,A329	C ₉ H	C ₉ H	A353
C ₇ ⁻	C ₇ ⁻	370,A329	C ₉ HN ⁺	H(C≡C) ₄ CN ⁺	A362
C ₇ F ₈ ⁺	C ₆ F ₅ CF ₃ ⁺	442	C ₉ H ₂ ⁺	C ₉ H ₂ ⁺	A353
C ₇ H	C ₇ H	A340	C ₉ H ₂	HC ₉ H	A353
C ₇ HN ⁺	H(C≡C) ₃ CN ⁺	A362	C ₉ N ₂ ⁺	NCC ₇ CN ⁺	A363
C ₇ H ₂ ⁺	C ₇ H ₂ ⁺	A349	C ₉ O	C ₉ O	A364
C ₇ H ₂	HC ₇ H	A349	C ₁₀	C ₁₀	A353
C ₇ H ₃ F ₅ ⁺	C ₆ F ₅ CH ₃ ⁺	441	C ₁₀ ⁻	C ₁₀ ⁻	A354
C ₇ H ₄ O	(cyc-C ₅ H ₄)CCO	419	C ₁₀ H	C ₁₀ H	A354
C ₇ H ₄ O	cyc-C ₆ H ₄ C=O	419	C ₁₀ HN ⁺	HC ₉ CN ⁺	A363
C ₇ H ₅ Cl	C ₆ H ₅ CCl	441	C ₁₀ H ₂ ⁺	C ₁₀ H ₂ ⁺	A354
C ₇ H ₅ Cl	(2-ClC ₆ H ₄)CH	441	C ₁₀ N ₂ ⁺	NC(C≡C) ₄ CN ⁺	A363
C ₇ H ₅ Cl	cyc-1-C ₇ H ₅ Cl	408	C ₁₁	C ₁₁	398,A354
C ₇ H ₅ ClO	C ₆ H ₅ OCCl	446	C ₁₁ ⁻	C ₁₁ ⁻	398
C ₇ H ₅ F	C ₆ H ₅ CF	440	C ₁₁ H	C ₁₁ H	A354
C ₇ H ₅ F	cyc-1-C ₇ H ₅ F	407	C ₁₁ HN ⁺	H(C≡C) ₅ CN ⁺	A363
C ₇ H ₅ F ₃ ⁺	sym-C ₆ H ₂ F ₃ CH ₃ ⁺	441	C ₁₁ H ₂ ⁺	C ₁₁ H ₂ ⁺	A355
C ₇ H ₅ N ⁺	C ₂ H ₅ (CC) ₂ CN ⁺	414	C ₁₁ H ₂	HC ₁₁ H	A355
C ₇ H ₆	C ₆ H ₅ CH	391	C ₁₁ N ₂ ⁺	NCC ₉ CN ⁺	A363
C ₇ H ₆	cyc-C ₇ H ₆	391	C ₁₂	C ₁₂	A355
C ₇ H ₆	cyc-C ₇ H ₆ :	392	C ₁₂ ⁻	C ₁₂ ⁻	A355
C ₇ H ₆ Cl	(4-ClC ₆ H ₄)CH ₂	A370	C ₁₂ H	C ₁₂ H	A355
C ₇ H ₆ F	(2-FC ₆ H ₄)CH ₂	A369	C ₁₂ HN ⁺	HC ₁₁ CN ⁺	A363
C ₇ H ₆ F	(3-FC ₆ H ₄)CH ₂	A369	C ₁₂ H ₂ ⁺	C ₁₂ H ₂ ⁺	A355
C ₇ H ₆ F	(4-FC ₆ H ₄)CH ₂	A370	C ₁₂ N ₂ ⁺	NC(C≡C) ₅ CN ⁺	A363
C ₇ H ₇ ⁺	cyc-C ₇ H ₇ ⁺	392	C ₁₃	C ₁₃	A356
C ₇ H ₇ ⁺	C ₆ H ₅ CH ₂ ⁺	392,A349	C ₁₃ HN ⁺	H(C≡C) ₆ CN ⁺	A363
C ₇ H ₇	C ₆ H ₅ CH ₂	392,A349	C ₁₃ H ₂ ⁺	C ₁₃ H ₂ ⁺	A356
C ₇ H ₇	cyc-C ₇ H ₇	394	C ₁₃ H ₂	HC ₁₃ H	A356
C ₇ H ₇	C ₆ H ₅ CH ₂ ⁻	394	C ₁₄ ⁻	C ₁₄ ⁻	A356
C ₇ H ₈ ⁺	C ₆ H ₅ CH ₃ ⁺	394	C ₁₄ H	C ₁₄ H	A356
C ₇ N ₂ ⁺	NCC ₅ CN ⁺	A362	C ₁₄ H ₂ ⁺	C ₁₄ H ₂ ⁺	A356
C ₇ O	C ₇ O	A341	C ₁₅	C ₁₅	A357
C ₇ O ₂	C ₇ O ₂	422	C ₁₆ ⁻	C ₁₆ ⁻	A357
C ₈	C ₈	381,A340	CaHO	CaOH	27,A139
C ₈ ⁻	C ₈ ⁻	381,A340	CaHS	CaSH	28,A140
C ₈ H	C ₈ H	A351	CaH ₂	CaH ₂	14
C ₈ HN ⁺	HC ₇ CN ⁺	A362	CaH ₂ N	CaNH ₂	132,A220
C ₈ H ₂ ⁺	C ₈ H ₂ ⁺	A351	CaH ₂ O	HCaOH	134
C ₈ H ₆ N	(4-NC-C ₆ H ₄)CH ₂	A370	CaH ₂ O ₂	Ca(OH) ₂	251
C ₈ H ₈	<i>o</i> -(CH ₂) ₂ C ₆ H ₄	396	CaN ₃	CaN ₃	182
C ₈ H ₈	<i>m</i> -(CH ₂) ₂ C ₆ H ₄	A351	CaO ₂	OCaO	A174
C ₈ H ₈	<i>p</i> -(CH ₂) ₂ C ₆ H ₄	A352	CaO ₂	cyc-CaO ₂	A174
C ₈ H ₈	C ₆ H ₅ CCH ₃	396	CaO ₄	O ₂ CaO ₂	A298
C ₈ H ₈	<i>m</i> -CH ₃ C ₆ H ₄ CH:	395	Ca ₂ H ₂	Ca ₂ H ₂	130
C ₈ H ₈	<i>o</i> -CH ₃ C ₆ H ₄ CH:	395	Ca ₂ H ₂ O	HCaOCaH	246
C ₈ H ₈	<i>p</i> -CH ₃ C ₆ H ₄ CH:	395	Ca ₂ H ₂ O	HCa ₂ OH	247
C ₈ H ₈	1-CH ₃ (cyc-C ₇ H ₅)	396	Ca ₂ H ₄	HCa ₂ CaH	305
C ₈ H ₈	4-CH ₃ (cyc-C ₇ H ₅)	397	Ca ₂ O ₂	cyc-(CaO) ₂	A243
C ₈ H ₈	5-CH ₃ (cyc-C ₇ H ₅)	397	Ca ₂ O ₂	CaOCaO	A243
C ₈ H ₈ ⁻	<i>m</i> -(CH ₂) ₂ C ₆ H ₄	A352	Ca ₃ H ₂	Ca ₃ H ₂	244
C ₈ N ₂ ⁺	NC(C≡C) ₃ CN ⁺	A362	CdH ₂	CdH ₂	A127
C ₈ O	C ₈ O	A364	CdH ₂ O	HCdOH	A222
C ₉	C ₉	398,A352	CdO ₂	OCdO	A179
C ₉ ⁻	C ₉ ⁻	398,A352	Cd ₂ H	CdCdH	A136

Formula	Structure/Name	References	Formula	Structure/Name	References
Cd ₂ O ₂	CdOCdO	A244	ClO ₂	OCIO	113,A205
CeO ₂	OCeO	A179	ClO ₂	ClOO	110,A204
CIFH ⁻	FHCl ⁻	53	ClO ₂ ⁻	OCIO ⁻	115
CIFN	NFCI	108,A201	ClO ₂ P	PO ₂ Cl	213
CIFO	FCIO	116	ClO ₃	ClO ₃	A262
ClFO ₂ S ⁺	FCISO ₂ ⁺	301	ClO ₄	OCIO ₃	A306
CIFO ₃ ⁺	FCIO ₃ ⁺	300	CIPS	CIPS	104
CIFS	FSCl	115,A208	ClS ₂	SSCl	111,A205
CIFXe	XeClF	121	ClXe ₂	Xe ₂ Cl	122
CIF ₂	ClF ₂	116	Cl ₂ F	CIClF	116
CIF ₂ ⁻	FCIF ⁻	117	Cl ₂ F ⁻	CIFCl ⁻	118
CIF ₂ ⁻	FFCl ⁻	118	Cl ₂ F ⁻	FCICl ⁻	118
CIF ₂ P ⁺	PF ₂ Cl ⁺	220	Cl ₂ F ₂	Cl ₂ F ₂	227
CIF ₃ ⁺	ClF ₃ ⁺	226	Cl ₂ F ₂ S	SiCl ₂ F ₂	304
CIF ₃ P ⁻	PClF ₃ ⁻	304	Cl ₂ F ₂ Si ⁺	SiF ₂ Cl ₂ ⁺	295
CIF ₃ S	SClF ₃	304	Cl ₂ F ₃ Si ⁻	SiF ₃ Cl ₂ ⁻	357
CIF ₃ Si ⁺	SiF ₃ Cl ⁺	295	Cl ₂ Ga ⁻	GaCl ₂	90
CIF ₄ Si ⁻	SiF ₄ Cl ⁻	357	Cl ₂ GaH	HGaCl ₂	169
ClFeH	HFeCl	35	Cl ₂ Ga ₂ H ₄	H ₂ GaCl ₂ GaH ₂	A339
ClGaH ₂	GaH ₂ Cl	143	Cl ₂ Ge ⁺	GeCl ₂ ⁺	92
ClGeH	HGeCl	45	Cl ₂ Ge	GeCl ₂	102,A198
ClGeH ₂	H ₂ GeCl	150	Cl ₂ GeH ₂ ⁺	GeH ₂ Cl ₂	263
ClGeH ₃ ⁺	GeH ₃ Cl ⁺	241	Cl ₂ GeS	Cl ₂ GeS	211
ClHI ⁻	ClHI ⁻	55	Cl ₂ H ⁻	CIHCl ⁻	54,A157
ClHKr	HKrCl	A158	Cl ₂ HN ⁺	HNCl ₂ ⁺	179
ClHO ⁺	HOCl ⁺	51	Cl ₂ H ₂ Si ⁺	SiH ₂ Cl ₂ ⁺	262
ClHO	HOCl	52,A155	Cl ₂ Mn ⁺	MnCl ₂ ⁺	73
ClHO ₂	HOCIO	A241	Cl ₂ N	NCl ₂	108
ClHO ₂	HClO ₂	A241	Cl ₂ Ni ⁺	NiCl ₂ ⁺	73
ClHSi	HClSi=S	172	Cl ₂ O ⁺	Cl ₂ O ⁺	111
ClHSi	HSiCl	44,A150	Cl ₂ O	ClClO	116,A208
ClHXe	HXeCl	A158	Cl ₂ OP	OPCl ₂	220
ClH ₂ ⁺	H ₂ Cl ⁺	26	Cl ₂ OS ⁺	Cl ₂ SO ⁺	223
ClH ₂ N ⁺	H ₂ NCl ⁺	152	Cl ₂ Os ⁻	Cl ₂ SiO	211
ClH ₂ P	H ₂ PCI	A229	Cl ₂ O ₂	ClOOCl	225,A263
ClH ₃ Si ⁺	SiH ₃ Cl ⁺	240	Cl ₂ O ₂	ClClO ₂	225,A263
ClIO	OICI	116	Cl ₂ O ₂	ClOClO	A264
ClIO ₂	ClIO ₂	226	Cl ₂ O ₂ S ⁺	Cl ₂ SO ₂ ⁺	301
ClIO ₂	IClO ₂	A264	Cl ₂ O ₂ Si	cyc-Cl ₂ SiO ₂	287
ClIO ₂	IOClO	A264	Cl ₂ O ₃	Cl ₂ O ₃	302,A307
ClKrNe	NeKrCl	122	Cl ₂ O ₄	ClOClO ₃	355,A322
ClKrXe	KrXeCl	122	Cl ₂ O ₆	O ₃ ClOClO ₂	382
ClKr ₂	Kr ₂ Cl	122	Cl ₂ P	PCl ₂	108,A203
ClNO ⁺	ClNO ⁺	93	Cl ₂ S ⁺	SCl ₂ ⁺	112
ClNO ₂ ⁺	ClNO ₂ ⁺	208	Cl ₂ SSi	Cl ₂ SiS	211
ClNO ₂	ClNO ₂	212,A259	Cl ₂ S ₂ ⁺	S ₂ Cl ₂ ⁺	224
ClNO ₂	c-ClONO	213,A259	Cl ₂ S ₂	SSCl ₂	226
ClNO ₂	t-ClONO	213,A259	Cl ₂ Se ⁺	SeCl ₂ ⁺	112
ClNO ₃ ⁺	ClONO ₂ ⁺	A301	Cl ₂ Se ₂ ⁺	Se ₂ Cl ₂ ⁺	224
ClNO ₃	ClONO ₂	288,A302	Cl ₂ Si ⁺	SiCl ₂ ⁺	92
ClNO ₅	O ₂ ClONO ₂	371	Cl ₂ Si	SiCl ₂	101,A197
CINS ⁺	NSCl ⁺	95	Cl ₂ Xe	XeCl ₂	121
CIN ₃ ⁺	CIN ₃ ⁺	196	Cl ₂ Zn ⁺	ZnCl ₂ ⁺	72
CIOP	CIPO	103	Cl ₃ ⁻	Cl ₃ ⁻	119
ClO ₂ ⁺	OCIO ⁺	106	Cl ₃ FS	SiCl ₃ F	304

Formula	Structure/Name	References	Formula	Structure/Name	References
Cl ₃ FSi ⁺	SiFCl ₃ ⁺	296	FFeH	HFeF	35
Cl ₃ F ₂ Si ⁻	SiF ₂ Cl ₃ ⁻	357	FFe ₂ H	HFe ₂ F	155
Cl ₃ Ge	GeCl ₃	219	FGaO	OGaF	85
Cl ₃ HOSi	SiCl ₃ OH	342	FGeH ₃ ⁺	GeH ₃ F ⁺	241
Cl ₃ HSi ⁺	HSiCl ₃ ⁺	276	FHI ⁻	FHI ⁻	54
Cl ₃ N ⁺	NCl ₃ ⁺	219	FHN	HNF	48,A152
Cl ₃ NO ₃ Si	SiCl ₃ ONO ₂	382	FHO ⁺	HOF ⁺	51
Cl ₃ NO ₄ Si	SiCl ₃ OONO ₂	426	FHO	HOF	51
Cl ₃ OP ⁺	Cl ₃ PO ⁺	298	FHSi	HSiF	43,A150
Cl ₃ O ₂ P	OPCl ₂ OCl	355	FH ₂ ⁺	H ₂ F ⁺	26
Cl ₃ P ⁺	PCl ₃ ⁺	220	FH ₂ N ⁺	NH ₂ F ⁺	151
Cl ₃ PS ⁺	Cl ₃ PS ⁺	299	FH ₂ N	NH ₂ F	153
Cl ₃ Sb ⁺	SbCl ₃ ⁺	222	FH ₂ P	H ₂ PF	153, ^A 228
Cl ₃ Si ⁺	SiCl ₃ ⁺	211	FH ₃ N ₂	NH ₂ NHF	324
Cl ₃ Si	SiCl ₃	218	FH ₃ Si ⁺	SiH ₃ F ⁺	240
Cl ₄ FSi ⁻	SiFCl ₄ ⁻	358	FI ₂	IIF	117
Cl ₄ Ge ⁺	GeCl ₄ ⁺	297,A306	FKrXe	KrXeF	121,A211
Cl ₄ Si ⁺	SiCl ₄ ⁺	296,A306	FKr ₂	Kr ₂ F	121,A210
Cl ₄ SiO	SiCl ₃ OCl	356	FNO ⁺	FNO ⁺	93
Cl ₅ P ⁺	PCl ₅	357	FNO	FON	104
CoH ₂	CoH ₂	15,A127	FNO ₂ ⁺	FNO ₂ ⁺	208
CoH ₂ ⁻	CoH ₂ ⁻	17	FNO ₂	FONO	212
CoH ₂ O	HCoOH	136	FNO ₃ ⁺	FONO ₂ ⁺	A301
CrH ₂	CrH ₂	15	FNO ₃	FONO ₂	A302
CrH ₂ O	HCrOH	136	FNS ⁺	NSF ⁺	94
CrH ₃	CrH ₃	123	FN ₂ ⁺	FNN ⁺	87
CrH ₃ O ₂	HCr(OH) ₂	316	FN ₃ ⁺	FN ₃ ⁺	196
CrO ₂	OCrO	A175	FN ₃	FN ₃	199
CrO ₂ ⁻	OCrO ⁻	A182	FNeXe	NeXeF	121
Cr ₂ H	CrCrH	26	FNe ₂	Ne ₂ F	121
Cr ₂ H ⁻	CrCrH ⁻	26	FOP	FPO	103
CuHO	CuOH	31	FOS	FSO	111
CuH ₂ N	CuNH ₂	133	FO ₂	FOO	109
CuH ₂ O	HCuOH	137	FO ₂ S ⁻	FSO ₂ ⁻	225
CuH ₃ N	HCuNH ₂	232	FO ₃ S ⁺	FSO ₃ ⁺	289
CuO ₂ ⁺	CuOO ⁺	A172	FO ₃ S	FSO ₃	300
CuO ₂	OCuO	A178	FPS	FPS	104
CuO ₂ ⁻	CuOO	A178	FS ₂	SSF	A205
CuO ₂ ⁻	OCuO ⁻	A182	FXe ₂	Xe ₂ F	122,A211
CuO ₂ ⁻	CuOO ⁻	A182	F ₂ Ga	GaF ₂	A193
CuO ₃	CuO ₃	A251	F ₂ Ge ⁺	GeF ₂	92
CuO ₃ ⁻	CuO ₃ ⁻	A252	F ₂ Ge	GeF ₂	101,A198
CuO ₄	O ₂ CuO ₂	A299	F ₂ GeH ₂ ⁺	GeH ₂ F ₂ ⁺	263
Cu ₂ H ₂	Cu ₂ H ₂	131	F ₂ GeO	F ₂ GeO	212
Cu ₂ O	CuOCu	A163	F ₂ H ⁻	FHF ⁻	53,A156
Cu ₂ O ⁻	CuOCu ⁻	A166	F ₂ HN ⁺	HNF ₂ ⁺	178
Cu ₂ O ₂	Cu ₂ O ₂	A244	F ₂ HP ⁺	HPF ₂ ⁺	179
Cu ₂ O ₂ ⁻	Cu ₂ O ₂ ⁻	A246	F ₂ H ₂ Si ⁺	SiH ₂ F ₂ ⁺	262
Cu ₂ O ₃	Cu ₂ O ₃	A294	F ₂ H ₃ P	PH ₃ F ₂	324,A315
Cu ₂ O ₃ ⁻	Cu ₂ O ₃ ⁻	A295	F ₂ I ⁻	FIF ⁻	118
Cu ₂ O ₄	Cu ₂ O ₄	A320	F ₂ I ⁻	FFI ⁻	118
Cu ₂ O ₄ ⁻	Cu ₂ O ₄ ⁻	A320	F ₂ I ₂	I ₂ F ₂	227
Cu ₃	Cu ₃	59,A161	F ₂ Kr ⁺	KrF ₂ ⁺	117
Cu ₃ H ₂	Cu ₃ H ₂	244	F ₂ Kr	KrF ₂	120,A209
DOSc	ScOD	30	F ₂ Mg ⁺	MgF ₂ ⁺	A181

Formula	Structure/Name	References	Formula	Structure/Name	References
F ₂ Mg	MgF ₂	84	F ₆ S ⁺	SF ₆ ⁺	A331
F ₂ N ⁺	NF ₂ ⁺	103,A199	FeHI	HFeI	35
F ₂ N	NF ₂	107,A201	FeH ₂	FeH ₂	15,A126
F ₂ N ⁻	NF ₂ ⁻	A207	FeH ₂ ⁻	FeH ₂ ⁻	17
F ₂ N ₂ ⁺	t-N ₂ F ₂ ⁺	207	FeH ₂ O	HFeOH	136
F ₂ O ⁺	OF ₂ ⁺	111	FeH ₂ O ₂	Fe(OH) ₂	251
F ₂ OS ⁺	F ₂ SO ⁺	223	FeH ₃	FeH ₃	A211
F ₂ OSi	F ₂ SiO	211	FeH ₃ N	HFeNH ₂	231
F ₂ O ₂	FOOF	225	FeNO	NFeO	A171
F ₂ O ₂ S ⁺	F ₂ SO ₂ ⁺	300	FeNO	FeNO	A171
F ₂ O ₂ Si	cyc-F ₂ SiO ₂	287	FeNO ₂	NFeO ₂	A249
F ₂ O ₂ Xe	XeO ₂ F ₂	304	FeN ₂	NFeN	A166
F ₂ O ₃ Xe	XeO ₃ F ₂	358	FeN ₂	FeNN	A167
F ₂ P ⁺	PF ₂ ⁺	104	FeN ₂ O	N ₂ FeO	A248
F ₂ P	PF ₂	108,A202	FeN ₂ O ₂	N ₂ FeO ₂	A296
F ₂ S ⁺	SF ₂ ⁺	112	FeO ₂	OFeO	A176
F ₂ S	SF ₂	114,A207	FeO ₂	FeOO	A176
F ₂ S ₂ ⁺	F ₂ SS ⁺	223	FeO ₂	cyc-FeO ₂	A177
F ₂ S ₂ ⁺	FSSF ⁺	224	FeO ₂ ⁻	OFeO ⁻	A182
F ₂ Se ⁺	SeF ₂ ⁺	112	FeO ₃	FeO ₃	A250
F ₂ Se	SeF ₂	A208	FeO ₃	(cyc-O ₂ Fe)O	A251
F ₂ Se ₂	FSeSeF	A264	FeO ₃ ⁻	FeO ₃ ⁻	A252
F ₂ Se ₂	SeSeF ₂	A264	FeO ₄	(cyc-O ₂ Fe)O ₂	A298
F ₂ Si ⁺	SiF ₂ ⁺	91	FeO ₄	(cyc-O ₂ Fe)OO	A298
F ₂ Si	SiF ₂	100,A197	Fe ₂ H ₂ O	HFeOFeH	247
F ₂ Xe ⁺	XeF ₂ ⁺	117	Fe ₂ H ₂ O	HFe ₂ OH	247
F ₂ Xe	XeF ₂	120,A209	Fe ₂ H ₃ N	HFe ₂ NH ₂	311
F ₃ ⁻	F ₃ ⁻	117	Fe ₂ N	cyc-Fe ₂ N	A162
F ₃ HSi ⁺	HSiF ₃ ⁺	276	Fe ₂ N	FeFeN	A162
F ₃ N ⁺	NF ₃ ⁺	219,A262	Fe ₂ O	FeOFe	A163
F ₃ NO ⁺	F ₃ NO ⁺	298	Fe ₂ O ⁻	FeOFe ⁻	A166
F ₃ NS ⁺	F ₃ NS ⁺	298	Fe ₂ O ₂	(FeO) ₂	A244
F ₃ OP ⁺	F ₃ PO ⁺	298	Fe ₂ O ₂ ⁻	Fe ₂ O ₂	A246
F ₃ OS ⁻	SOF ₃ ⁻	304	Fe ₂ O ₃	Fe ₂ O ₃	A294
F ₃ O ₂ S ⁻	SO ₂ F ₃ ⁻	358	Fe ₂ O ₃ ⁻	Fe ₂ O ₃ ⁻	A295
F ₃ P ⁺	PF ₃ ⁺	219	Fe ₂ O ₄	Fe ₂ O ₄	A320
F ₃ PS ⁺	F ₃ PS ⁺	299	Fe ₂ O ₄ ⁻	Fe ₂ O ₄ ⁻	A320
F ₃ S	SF ₃	226	GaHO	GaOH	36
F ₃ Sb ⁺	SbF ₃ ⁺	222	GaH ₂	GaH ₂	18,A129
F ₃ Si ⁺	SiF ₃ ⁺	A258	GaH ₂ O	HGaOH	139,A223
F ₃ Si	SiF ₃	218,A262	GaH ₃	GaH ₃	A213
F ₃ Si ⁻	SiF ₃ ⁻	A263	GaInO	InGaO	64
F ₄ Ge ⁺	GeF ₄ ⁺	297	GaO ₂	cyc-GaO ₂	75
F ₄ OXe ⁺	XeOF ₄ ⁺	359	GaO ₂	OGaO	75
F ₄ P ⁻	PF ₄ ⁻	304	GaO ₃	OGaOO	190
F ₄ P ₂ ⁺	P ₂ F ₄ ⁺	355	GaP ₂	GaP ₂	A172
F ₄ P ₂ ⁻	PF ₃ =PF	355	GaSb ₂	GaSb ₂	A172
F ₄ S ⁺	SF ₄ ⁺	A307	Ga ₂ H ₂	cyc-Ga ₂ H ₂	A220
F ₄ Si ⁺	SiF ₄ ⁺	294,A305	Ga ₂ H ₂	HGaGaH	A220
F ₄ Xe ⁺	XeF ₄ ⁺	305	Ga ₂ H ₆ ⁺	Ga ₂ H ₆ ⁺	A333
F ₅ I ⁺	IF ₅ ⁺	358	Ga ₂ H ₆	Ga ₂ H ₆	372,A333
F ₅ P ⁺	PF ₅ ⁺	356	Ga ₂ O	Ga ₂ O	64
F ₅ S	SF ₅	358	Ga ₂ O ₂	GaOGaO	185
F ₅ S ⁻	SF ₅ ⁻	359	Ga ₂ O ₂	cyc-GaO ₂ Ga	185
F ₅ Si ⁻	SiF ₅	357	Ga ₂ O ₃	Ga ₂ O ₃	283

Formula	Structure/Name	References	Formula	Structure/Name	References
Ga ₂ P	Ga ₂ P	A165	HNO ₄	HOONO ₂	341,A317
Ga ₂ Sb	Ga ₂ Sb	A165	HNSc	ScNH	A138
Ga ₃ H ₂	HGa ₃ H	A275	HNSi	HNSi	38,A146
GeH ₂	GeH ₂	2,A132	HNSi	HSiN	A145
GeH ₂ I ₂ ⁺	GeH ₂ I ₂ ⁺	263	HN _Y	YNH	27
GeH ₂ O	H ₂ GeO	146	HN ₂ ⁺	HN ₂ ⁺	39
GeH ₂ O	HGeOH	146	HN ₂ O ⁺	HONN ⁺	165
GeH ₂ O ₃	H ₂ GeO ₃	338	HN ₃ ⁺	HN ₃ ⁺	161
GeH ₃ ⁺	GeH ₃ ⁺	125	HNaO ⁺	NaOH ⁺	27
GeH ₃	GeH ₃	128	HNiO	NiOH	30
GeH ₃ I ₃ ⁺	GeH ₃ I ₃ ⁺	242	HOP	HPO	46
GeH ₄ ⁺	GeH ₄ ⁺	228	HOS ⁺	HSO ⁺	A152
GeH ₄ O	GeH ₃ OH	311	HOS	HSO	50,A154
GeH ₄ S ⁺	GeH ₃ SH ⁺	311	HOSi ⁺	HOSi ⁺	39
GeI ₂ ⁺	GeI ₂ ⁺	92	HOSr	SrOH	29,A140
GeOS	OGeS	86	HOXe	XeOH	53
GeO ₂	GeO ₂	86	HOZn	ZnOH	31,A141
GeO ₂ ⁻	GeO ₂ ⁻	A194	HO ₂ ⁺	HO ₂ ⁺	47
GeS ₂	GeS ₂	86	HO ₂	HO ₂	49,A153
Ge ₂ H ₂ O	HGe ₂ OH	256	HO ₂ ⁻	HO ₂ ⁻	51
Ge ₂ O ₂	Ge ₂ O ₂	188,A251	HO ₂ P	c-HOPO	174
Ge ₂ O ₂ ⁻	Ge ₂ O ₂ ⁻	A252	HO ₃ P	HOPO ₂	273
Ge ₂ O ₃	Ge ₂ O ₃	A300	HO ₃ P	HOOPo	273
Ge ₃	Ge ₃	A171	HO ₃ P	HP(O ₂)O	273
Ge ₃ ⁻	Ge ₃ ⁻	A173	HO ₃ S	HOSO ₂	277
Ge ₃ O ₃	(GeO) ₃	346	HO ₃ Sb	HSbO ₃	274
Ge ₄	Ge ₄	A246	HO ₃ Sb	HOSbO ₂	274
HIO ⁺	HOI ⁺	A155	HPS ₂	HPSS	A241
HIO	HOI	52,A156	HPS ₃	HSPS ₂	A292
HISi	HSiI	44	HSSr	SrSH	29
HIXe	HXeI	A158	HS ₂	HS ₂	50,A154
HI ₂ ⁻	IHI ⁻	55,A158	HS ₂ ⁻	HS ₂ ⁻	53
HInO	InOH	37,A144	HXe ₂ ⁺	HXe ₂ ⁺	56
HKO ⁺	KOH ⁺	27	HXe ₂	Xe ₂ H	57
HKrXe	KrXeH	57	HZn ₂	ZnZnH	A136
HKr ₂ ⁺	HKr ₂ ⁺	56,A159	H ₂ Hf	HfH ₂	A126
HKr ₂	Kr ₂ H	57	H ₂ Hg	HgH ₂	16,A128
HLi ₂	Li ₂ H	26,A136	H ₂ Hg ₂	HHgHgH	A218
HMgO	MgOH	27,A138	H ₂ IP	H ₂ PI	A229
HNO ⁺	HNO ⁺	41	H ₂ I ₂ Si ⁺	SiH ₂ I ₂ ⁺	263
HNO	HNO	45,A151	H ₂ In	InH ₂	A129
HNO ⁻	HNO ⁻	48	H ₂ InO	HInOH	139
HNOS ⁺	HNSO ⁺	169	H ₂ Mg	MgH ₂	14,A125
HNOS	t-HONS	174	H ₂ MgO	HMgOH	134
HNOS	t-HSNO	174	H ₂ MgO ₂	Mg(OH) ₂	A282
HNOS	c-HSNO	174	H ₂ Mg ₂	HMgMgH	A217
HNOS	c-HNSO	175,A240	H ₂ Mg ₂	br-(MgH) ₂	A218
HNOS	t-HNSO	175	H ₂ Mg ₂ O	HMgOMgH	246
HNOS	c-HOSN	176	H ₂ Mg ₂ O	HMg ₂ OH	246
HNO ₂	t-HONO	172	H ₂ Mn	MnH ₂	15
HNO ₂	c-HONO	173	H ₂ Mn ⁻	MnH ₂ ⁻	17
HNO ₃ ⁺	HNO ₃ ⁺	270	H ₂ MnO	HMnOH	136
HNO ₃	HONO ₂	271,A289	H ₂ Mn ₂ O	HMn ₂ OH	247
HNO ₃	c,c-HOONO	A291	H ₂ Mn ₂ O	HMnOMnH	247
HNO ₃	t,perp-HOONO	272,A291	H ₂ Mo	MoH ₂	16

Formula	Structure/Name	References	Formula	Structure/Name	References
H ₂ N ⁺	NH ₂ ⁺	20,A132	H ₂ U ₂	cyc-U ₂ H ₂	A218
H ₂ N	NH ₂	21,A133	H ₂ V	VH ₂	14
H ₂ N ⁻	NH ₂ ⁻	26	H ₂ Xe	HXeH	A136
H ₂ NO ⁺	H ₂ NO ⁺	148	H ₂ Zn	ZnH ₂	16,A127
H ₂ NO	H ₂ NO	151	H ₂ Zn ₂	HZnZnH	130,A218
H ₂ NSr	SrNH ₂	133	H ₂ Zr	ZrH ₂	A126
H ₂ N ₂ ⁺	t-N ₂ H ₂ ⁺	142	H ₃ ⁺	H ₃ ⁺	11,A123
H ₂ N ₂	t-N ₂ H ₂	147,A227	H ₃	H ₃	12,A123
H ₂ N ₂	H ₂ NN	148	H ₃ Hf	HfH ₃	A211
H ₂ N ₂ O	NH ₂ NO	260	H ₃ Si ⁺	SiH ₃ I ⁺	241
H ₂ N ₂ O ₂	NH ₂ NO ₂	337	H ₃ In	InH ₃	A213
H ₂ Nb	NbH ₂	A128	H ₃ Mo	MoH ₃	124
H ₂ Ni	NiH ₂	16,A127	H ₃ N ⁺	NH ₃ ⁺	128,A217
H ₂ Ni ⁻	NiH ₂ ⁻	17	H ₃ NNi	HNiNH ₂	231
H ₂ NiO	HNiOH	136	H ₃ NO ⁺	NH ₂ OH ⁺	242
H ₂ NiO ₂	Ni(OH) ₂	252	H ₃ NSi	HSiNH ₂	235
H ₂ Ni ₂ O	HNi ₂ OH	248	H ₃ NSi	SiH ₂ NH	A270
H ₂ O ⁺	H ₂ O ⁺	24,A134	H ₃ N ₂ ⁺	N ₂ H ₃ ⁺	236
H ₂ OP	H ₂ PO	151,A228	H ₃ O ⁺	H ₃ O ⁺	129
H ₂ OP	HPOH	151	H ₃ OP	PH ₃ O	242
H ₂ OS	HSOH	154	H ₃ OP	c-H ₂ POH	243
H ₂ OSc	HScOH	135	H ₃ OSb	H ₃ SbO	243
H ₂ OSi	HSiOH	146	H ₃ OSb	H ₂ SbOH	244
H ₂ OSi	H ₂ SiO	145,A226	H ₃ O ₃ P	(HO) ₂ HPO	368
H ₂ OSn	HSnOH	146	H ₃ P ⁺	PH ₃ ⁺	129
H ₂ OSn ₂	HSn ₂ OH	257	H ₃ PS	H ₂ PSH	A275
H ₂ OSr	HSrOH	134	H ₃ S ⁺	H ₃ S ⁺	130,A217
H ₂ OTi	HTiOH	135	H ₃ Sb ⁺	SbH ₃ ⁺	129
H ₂ OV	HVOH	135	H ₃ Si ⁺	SiH ₃ ⁺	124,A213
H ₂ OZn	HZnOH	A222	H ₃ Si	SiH ₃	126,A215
H ₂ O ₂ ⁺	HOOH ⁺	152	H ₃ Si ⁻	SiH ₃ ⁻	129
H ₂ O ₂	HOOH	153,A229	H ₃ Th	ThH ₃	A212
H ₂ O ₂ S	HSO ₂ H	265	H ₃ Ti	TiH ₃	A211
H ₂ O ₂ Si	HSiOOH	260	H ₃ U	UH ₃	A212
H ₂ O ₃ Si	H ₂ SiO ₃	338	H ₃ Zr	ZrH ₃	A211
H ₂ P ⁺	PH ₂ ⁺	21	H ₄ Hf	HfH ₄	A265
H ₂ P	PH ₂	22,A134	H ₄ Mg ₂	HMgH ₂ MgH	A308
H ₂ PS	HPSH	A228	H ₄ N ⁺	NH ₄ ⁺	229,A266
H ₂ Pt	PtH ₂	A127	H ₄ N	NH ₄	229,A266
H ₂ S ⁺	H ₂ S ⁺	24,A135	H ₄ N ₂ ⁺	N ₂ H ₄ ⁺	311
H ₂ S ₂ ⁺	HSSH ⁺	153	H ₄ N ₂ O	NH ₂ NHOH	364
H ₂ S ₂	HSSH	155,A230	H ₄ OSi	SiH ₃ OH	311
H ₂ S ₃	c-HSSSH	A285	H ₄ P ₂ ⁺	P ₂ H ₄ ⁺	311
H ₂ S ₃	t-HSSSH	A285	H ₄ SSi ⁺	SiH ₃ SH ⁺	310
H ₂ Sb	SbH ₂	23	H ₄ Si ⁺	SiH ₄ ⁺	228
H ₂ Se ⁺	H ₂ Se ⁺	25	H ₄ Th	ThH ₄	A265
H ₂ Si ⁺	SiH ₂ ⁺	18,A130	H ₄ Ti	TiH ₄	227,A265
H ₂ Si	SiH ₂	19,A131	H ₄ U	UH ₄	A265
H ₂ Si ⁻	SiH ₂ ⁻	21	H ₄ U ₂	U ₂ H ₄	A308
H ₂ Si ₂	br-Si ₂ H ₂	138,A223	H ₄ Zr	ZrH ₄	A265
H ₂ Si ₂	Si(H)SiH	138	H ₅ ⁺	H ₅ ⁺	227
H ₂ Te ⁺	H ₂ Te ⁺	25	H ₅ NSi	SiH ₃ NH ₂	360
H ₂ Th	ThH ₂	A128	H ₅ O ₂ ⁺	H ₅ O ₂ ⁺	360,A323
H ₂ Ti	TiH ₂	14,A126	H ₅ Si ⁺	SiH ₅ ⁺	A307
H ₂ U	UH ₂	A128	H ₆ OSi ₂	SiH ₃ SiH ₂ OH	417

Formula	Structure/Name	References	Formula	Structure/Name	References
H ₇ O ₃ ⁺	H ₇ O ₃ ⁺	414	NOS ₂	SSNO	A258
H ₉ O ₄ ⁺	H ₉ O ₄ ⁺	414,A364	NOSe	SeNO	A195
HfO ₂	OHfO	A176	NOU	NUO	A172
HfO ₂ ⁻	OHfO ⁻	A182	NOU	UNO	A172
HfO ₃	OHfOO	A250	NO ₂ ⁺	NO ₂ ⁺	87,A190
Hf ₂ O ₂	cyc-(HfO) ₂	A244	NO ₂ ⁻	NO ₂ ⁻	102
IKrXe	KrXeI	123	NO ₂ U	NUO ₂	A249
INO	INO	102	NO ₃ ⁺	NO ₃ ⁺	A255
INO ₂	INO ₂	213	NO ₃	NO ₃	207,A257
INO ₃	IONO ₂	289	NO ₃ ⁻	NO ₃ ⁻	A259
INS ⁺	NSI ⁺	95	NP ₂	PNP	A187
INS	NSI	105	NS ₂ ⁺	SNS ⁺	A191
IO ₂	OIO	114,A207	NS ₂	NS ₂	94,A195
IO ₂	IOO	A204	NS ₂	NSS	94
IO ₂ ⁻	OIO ⁻	115	NS ₂ ⁻	NS ₂ ⁻	102
IXe ₂	Xe ₂ I	123	NS ₃	NSSS	208
I ₂ S	SI ₂	115	NSe ₂ ⁺	SeNSE ⁺	A191
I ₂ Si ⁺	SiI ₂ ⁺	92	NSe ₂	NSe ₂	A195
I ₃ ⁻	I ₃ ⁻	119	NSe ₂	SeSeN	A195
InO ₂	cyc-InO ₂	75	NSe ₃	SeSeNSE	A258
InO ₂	OInO	75	NSe ₃	SeSeSeN	A258
In ₂ O	In ₂ O	64	NSi ₂	SiNSi	A174
In ₂ O ₂	InOInO	185	N ₂ O ⁺	N ₂ O ⁺	83,A187
In ₂ O ₂	cyc-InO ₂ In	185	N ₂ OS	SNNO	A254
In ₂ O ₃	In ₂ O ₃	283	N ₂ O ₂ ⁺	t-(NO) ₂ ⁺	195,A252
In ₂ P	In ₂ P	A165	N ₂ O ₂	c-(NO) ₂	200,A254
In ₂ Sb	In ₂ Sb	A165	N ₂ O ₂	t-(NO) ₂	200,A254
KNO ₂	cyc-KNO ₂	A247	N ₂ O ₂	NNO ₂	A254
KNO ₂	t-KNO ₂	A247	N ₂ O ₂ ⁻	NNO ₂ ⁻	207,A256
KNO ₃	c-KOONO	A296	N ₂ O ₂ ⁻	t-(NO) ₂ ⁻	207,A257
KNO ₃	t-KOONO	A296	N ₂ O ₂ S	ON-NSO	286
LiNO ₃	c-LiOONO	A295	N ₂ O ₂ S	ONSNO	286
LiNO ₃	t-LiOONO	A295	N ₂ O ₂ Ti	O ₂ TiN ₂	A296
LiNa ₃	LiNa ₃	180	N ₂ O ₂ U ₂	(NUO) ₂	A320
Li ₂ Na ₂	Li ₂ Na ₂	180	N ₂ O ₃	O ₂ N-NO	286,A300
Li ₃	Li ₃	58,A159	N ₂ O ₃	O=N-O-N=O	286,A300
Li ₄	Li ₄	180	N ₂ O ₄ ⁺	N ₂ O ₄ ⁺	349
Li ₆	Li ₆	342	N ₂ O ₄	N ₂ O ₄	350,A321
Li ₇	Li ₇	369	N ₂ O ₄	N ₂ O ₄ (V _d)	350
Li ₈	Li ₈	381	N ₂ O ₄	ONO-NO ₂ (D)	351
MgO ₂	OMgO	A174	N ₂ O ₄	ONO-NO ₂ (D')	351
MgO ₂	cyc-MgO ₂	A174	N ₂ O ₅	O ₂ N-O-NO ₂	370,A331
MgO ₄	O ₂ MgO ₂	A298	N ₂ Pu	NPuN	A168
Mg ₂ O ₂	MgOMgO	A243	N ₂ S ⁺	NNS ⁺	83
Mn ₃	Mn ₃	61	N ₂ S	NNS	88
MoO ₂	OMoO	A175	N ₂ S ₂ ⁺	N ₂ S ₂ ⁺	196
MoO ₃	MoO ₃	A250	N ₂ S ₂	NS-SN	200
NNaO ₂	cyc-NaNO ₂	A247	N ₂ S ₄ ⁺	N ₂ S ₄ ⁺	349
NNaO ₂	t-NaONO	A247	N ₂ S ₄	cyc-N ₂ S ₄	351
NNaO ₃	c-NaOONO	A295	N ₂ Se ₂	NSENSE	A255
NNaO ₃	t-NaOONO	A296	N ₂ Si	SiNN	72
NOP	PNO	88	N ₂ Ti	TiNN	A166
NOS ⁺	SNO ⁺	A191	N ₂ U	NUN	63
NOS	SNO	93,A194	N ₃ ⁺	N ₃ ⁺	72,A181
NOS	NSO	94	N ₃	N ₃	82

<i>Formula</i>	<i>Structure/Name</i>	<i>References</i>	<i>Formula</i>	<i>Structure/Name</i>	<i>References</i>
N_3^-	N_3^-	87	O_2Si	SiO_2	86
N_3P_3	$(PN)_3$	346	O_2Si^-	SiO_2^-	A194
$N_3S_3^+$	$S_3N_3^+$	347	O_2Si_2	Si_2O_2	188
N_3Sr	SrN_3	183	O_2Sn	$OSnO$	A189
N_4^+	N_4^+	187,A250	O_2Sn_2	Sn_2O_2	A251
Na_3	Na_3	58,A160	O_2Sr	$OSrO$	A174
Na_4	Na_4	180	O_2Sr	$cyc\text{-}SrO_2$	A175
Na_5	Na_5	277	O_2Sr_2	$cyc\text{-}(SrO)_2$	A243
Na_6	Na_6	342	O_2Ta	$OTaO$	A176
Na_7	Na_7	369	O_2Tb	$OTbO$	A179
Na_8	Na_8	381	O_2Th	$OThO$	A179
Nb_3	Nb_3	A162	O_2Ti	$OTiO$	A176
NiO_2	$ONiO$	A177	O_2Ti^-	$OTiO^-$	A181
NiO_2	$cyc\text{-}NiO_2$	A177	O_2Ti_2	$cyc\text{-}(TiO)_2$	A243
NiO_2	$NiOO$	A177	O_2U	OUO	A179
NiO_2^-	$ONiO^-$	A182	O_2W	OWO	A175
NiO_2^-	$cyc\text{-}NiO_2^-$	A182	O_2Zn	$OZnO$	A178
NiO_3	$ONiOO$	A251	O_2Zn_2	$ZnOZnO$	A244
NiO_3	$cyc\text{-(}O_2Ni\text{)}O$	A251	O_2Zr	$OZrO$	A176
NiO_4	O_2NiO_2	A298	O_2Zr^-	$OZrO^-$	A182
Ni_2O	$NiNiO$	A163	O_2Zr_2	$cyc\text{-(}ZrO\text{)}_2$	A243
Ni_2O_2	$cyc\text{-(}NiO\text{)}_2$	A244	O_3^+	O_3^+	95
Ni_2O_2	$NiONiO$	A244	O_3^-	O_3^-	109,A203
Ni_3	Ni_3	61	O_3P	PO_3	209
OP_2	P_2O	88,A191	O_3Pb	$OPb(O_2)$	A253
OP_4	P_4O	284	O_3S^+	SO_3^+	209
OP_4	$br\text{-}P_4O$	284	O_3S	$OSOO$	A260
OPb_2	$PbOPb$	A181	O_3S^-	SO_3^-	223
$OSSi$	$OSiS$	86	O_3Se	SeO_3	214,A260
OS_2^+	SSO^+	97	O_3Se	$OSeOO$	A261
OS_2	SSO	105,A199	O_3Si	SiO_3	A253
OS_2^-	SSO^-	110	$O_3Si_2^-$	$Si_2O_3^-$	A300
$OSeSi$	$OSiSe$	A189	O_3Si_3	$(SiO)_3$	346
OSe_2	$SeSeO$	A201	O_3Ti	$OTiOO$	A250
OTi_2	$TiOTi$	A163	O_3U	UO_3	A251
OTl_2	Tl_2O	65	O_3W	WO_3	A250
O_2P	PO_2	93,A194	O_3Zr	$OZrOO$	A250
O_2P^-	PO_2^-	103,A199	O_4^+	$t\text{-}O_4^+$	209,A258
O_2P_2	$(PO)_2$	200	O_4^+	$cyc\text{-}O_4^+$	209,A258
O_2Pb	$OPbO$	A190	O_4^-	$t\text{-}O_4^-$	222
O_2Pb	$cyc\text{-}PbO_2$	A190	O_4Pd	O_2PdO_2	A298
O_2Pb_2	Pb_2O_2	A252	O_4Pt	O_2PtO_2	A298
O_2Ph_2	$PhPh(O_2)$	A252	O_4Rh	O_2RhO_2	A298
O_2Pd	$cyc\text{-}PdO_2$	A177	O_4S	SO_4	289
O_2Pr	$OPrO$	A179	O_4Si_2	Si_2O_4	A321
O_2Pt	$cyc\text{-}PtO_2$	A177	$O_4Si_2^-$	$Si_2O_4^-$	A321
O_2Pu	$OPuO$	A179	O_5P_2	P_2O_5	371
O_2Rh	$cyc\text{-}RhO_2$	A177	PS_2	PS_2	95
O_2S^+	SO_2^+	96,A195	PS_2^-	PS_2^-	104
O_2S	SOO	A199	P_2S	P_2S	88
O_2S^-	SO_2^-	110	P_2S_4	P_2S_4	351
O_2Se	SeO_2	A200	P_3	P_3	82
O_2Se	$SeOO$	A201	P_4^+	P_4^+	187
O_2Se^-	SeO_2^-	A204	P_4S	$cyc\text{-}P_4S$	284
O_2Se_2	$cyc\text{-(}SeOSe\text{)}=O$	A261	Pd_3	Pd_3	61

<i>Formula</i>	<i>Structure/Name</i>	<i>References</i>
Pt ₃	Pt ₃	61
S ₂ Si	SiS ₂	86
S ₃	S ₃	105,A200
S ₃ ⁻	S ₃ ⁻	110
S ₄	S ₄	214,A260
S ₄	SS ₃	214
Sb ₃	Sb ₃	82
Sb ₃ ⁻	Sb ₃ ⁻	88
Sb ₄	Sb ₄	189
Sb ₄ ⁻	Sb ₄ ⁻	197
Sc ₃	Sc ₃	61
Se ₂ Si	SiSe ₂	A189
Se ₃	Se ₃	106
Se ₄	Se ₄	214
Se ₄	SeSe ₃	215
Si ₃	Si ₃	67,A170
Si ₃ ⁻	Si ₃ ⁻	68,A173
Si ₄	Si ₄	182,A245
Si ₄ ⁻	Si ₄ ⁻	183,A247
Si ₆	Si ₆	A318
Si ₇	Si ₇	A329
Ta ₄	Ta ₄	A241
Te ₃	Te ₃	106
Te ₄	Te ₄	215
Te ₄	TeTe ₃	215