High Resolution FTIR and Diode Laser Spectroscopy of Supersonic Jets
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reprinted from

“Handbook of High-Resolution Spectroscopy”
Vol. 2, chapter 27, pages 1021–1067
M. Quack, and F. Merkt, Eds. Wiley Chichester, 2011,
Online ISBN: 9780470749593,
DOI: 10.1002/9780470749593.

with compliments from Professor Martin Quack, ETH Zürich
Abstract

Fourier transform infrared (FTIR) spectrometers and tunable diode lasers in combination with a supersonic molecular beam expansion are a perfect tool for the investigation of molecules, ions, and radicals at low temperatures. The internal degrees of freedom of the molecules are adiabatically cooled to very low temperatures and thus only low-lying energy levels are populated. The reduction of the number of populated levels at low temperatures makes the assignment of the spectra much easier as compared to the congested room-temperature spectra. Under certain conditions, the Doppler linewidths are greatly reduced, corresponding to very low effective translational temperatures. Supersonic expansion also provides a suitable method for producing and investigating van der Waals clusters and hydrogen-bonded complexes. Unstable species such as radicals and ions can be efficiently produced and studied in a molecular beam. The low rotational temperature allows for the study of nuclear spin symmetry conservation or conversion between nuclear spin isomers. A molecular beam expansion can be obtained by expanding gas through a slit or a circular nozzle. Both expansion geometries can be used in combination with a multipass optical setup and with cavity ring down spectroscopy, which enhances the effective absorption path length. Cooling of the molecules can be promoted by seeding in noble gases. This article summarizes the general aspects of the experimental technique as well as current developments. To demonstrate how powerful the combination of a molecular beam expansion with tunable diode laser and FTIR spectroscopy can be, we report results on some important current examples.

Keywords: FTIR; supersonic jet; diode laser; molecular beam; clusters; van der Waals complexes; hydrogen bonds; nuclear spin symmetry conservation; infrared spectroscopy; isotopes
1 INTRODUCTION

1.1 Brief History

The combination of spectroscopy with atomic and molecular beams can be viewed as one of the most fruitful unions in science. Its history goes back both to the Stern–Gerlach experiment (Gerlach and Stern 1924, Gerlach 1925) and the subsequent development of nuclear magnetic resonance (NMR) experiments in atomic beams by Rabi (1937) in the first half of the twentieth century (see also Bennewitz and Paul (1954) and Goldenberg et al. (1960) for further development of the experimental technology). While this research was originally curiosity driven in order to get a better understanding of nuclear structure, today the results of modern NMR spectroscopy (Ernst et al. 1987) can be admired in every chemical and analytical laboratory as well as in terms of magnetic resonance imaging (MRI) in every major hospital. The somewhat related developments in molecular beam state selection and electric resonance (Dyke et al. 1972, Bennewitz et al. 1964, Trischka 1962, Novick et al. 1973, Meerts and Dymanus 1975, Meerts et al. 1979) and beam MASER spectroscopy (Gordon et al. 1955, Dymanus 1976) as well as later Fourier Transform Microwave (FT-MW) spectroscopy of beams (Bauder 2011: Fundamentals of Rotational Spectroscopy, this handbook) had already important applications in molecular spectroscopy. The early uses of molecular beams in optical spectroscopy are connected with sensitive detection techniques such as laser-induced fluorescence (Sinha et al. 1973, Smalley et al. 1974) and this early work has been reviewed by Smalley et al. (1977) and Levy (1980).

In the classic paper, largely related to the thesis work of the late Roger Miller, Gough et al. (1977) introduced mid-infrared diode laser absorption spectroscopy of supersonic jets (reviewed in part in Miller (1992)). This technique provided high, essentially Doppler-limited resolution, but has the serious drawback of very limited scanning abilities. Scanning of a typical laser diode can range between about 10 and 50 cm$^{-1}$ at most, but scans are not continuous, often missing important parts of a spectrum. Continuous laser diode scans are frequently limited to about 2 cm$^{-1}$, which is far too small for an efficient rovibrational analysis of complex infrared bands.

Obvious infrared techniques with large scanning abilities covering the complete infrared spectral range are grating and FTIR spectrometers, which have both been used in the late 1970s and early 1980s in conjunction with supersonic jets (Kim et al. 1978, Snavely et al. 1981, 1983, 1984). These techniques reached only modest resolution of at most 0.06 cm$^{-1}$, which has the major drawback that the measured spectra are limited by the instrumental shape, resulting in only limited possibilities for rovibrational analysis and introducing artifacts concerning line strengths, which can be evaluated only if the instrumental bandwidth approaches the Doppler linewidth in the supersonic jet.

The Zürich group, therefore, in the 1980s developed high-resolution FTIR spectroscopy of supersonic jets under conditions that often satisfy the requirement...
of Doppler-limited resolution (instrumental bandwidth 0.0024 cm\(^{-1}\) or 72 MHz, unapodized) (Dübäi et al. 1984, Amrein et al. 1987a,b, 1988a,b, 1989, Quack 1990). At the same time, it was recognized that an ideal combination would be to realize the advantages of high-resolution FTIR spectroscopy and diode laser spectroscopy in the same laboratory and such a combination was realized in the Zürich laboratory, leading among other things to the first analyses of the very complex infrared spectra of the important atmospheric window bands of the chlorofluoro(hydro)carbons CHFCl\(_2\) (Snels and Quack 1991), \(\text{CF}_2\text{Cl}_2\) (D’Amico et al. 2002), and \(\text{CFCl}_3\) (Snels et al. 1995, 2001) and the complex spectra of \(\text{CF}_3\) (Hollenstein et al. 1994, He et al. 2002); see also the work on \(\text{CHClF}_2\) (Albert et al. 2010, 2004, Amrein et al. 1988a). It also led to the very first successful high-resolution rovibrational analyses of chiral molecules in the mid-1990s (Beil et al. 1994, Bauder et al. 1997), this work having significance well beyond ordinary rovibrational spectroscopy (Quack 2002, 2003, 2011, Quack et al. 2008). Until today, to our knowledge, only one other group has repeated a combination of these two spectroscopic techniques much more recently (Herman et al. 2007). Further important experimental developments concern the combination of pulsed supersonic jets with continuous scan FTIR spectroscopy (Luckhaus et al. 1995), a possibility, which was probably not anticipated by anyone, and the combination of pulsed supersonic jets with continuous laser cavity ring down spectroscopy, which provides very high sensitivity and very high resolution (1 MHz and potentially better), by Hippler and Quack (1999, 2002). The combination of pulsed IR-laser spectroscopy of pulsed supersonic jets with ionization and mass-selective detection resulted in mass-and isotope-selective spectroscopy reviewed by Hippler and Quack (2005) and Hippler et al. (2011): Mass and Isotope-selective Infrared Spectroscopy, this handbook. Such a mass-selective technique combined with supersonic-jet FTIR spectroscopy had been suggested already by Quack (1990) but has not been realized, so far, as it is technically very demanding.

1.2 Some Aspects of the Spectroscopy of Supersonic Jets and Related Developments

High-resolution spectroscopy in supersonic jets offers many advantages and opportunities as compared to experiments in static gas cells. One of the properties of a supersonic expansion is the adiabatic cooling of the internal degrees of freedom of a molecule. Whereas all degrees of freedom are in thermal equilibrium in a static gas sample, in a beam, the population of rotational and vibrational energy levels can be often approximately described by a Boltzmann distribution with different, very low rotational and vibrational temperatures. Most efficient cooling is obtained by diluting (seeding) the sample in an inert gas, typically a rare gas that promotes relaxation of rotational and vibrational levels of a polyatomic molecule. Typically rotational temperatures of 10–30 K can be routinely obtained, whereas the vibrational temperature is usually higher. In addition, because of the irregular spacing between vibrational energy levels, the population of vibration levels is not well described by a Boltzmann distribution. In some cases, bottlenecks might occur, trapping a substantial part of the population in an “isolated” vibrational level, i.e., with a high energy gap with respect to the next lower level. By varying the collision partner (He, Ne, Ar, or mixtures of noble gases), cooling can be made more efficient. At the same time, the large number of collisions in a supersonic expansion can also promote cluster formation, and thus the supersonic expansion has become an appropriate tool for generating van der Waals and hydrogen-bonded complexes. In a similar way, other unstable species, such as radicals and ions, can be generated in a supersonic expansion and successively cooled and studied. Yet another opportunity is the evaporation of solid material that can be entrained in a supersonic expansion of inert gases.

The Doppler broadening of spectral lines observed in supersonic jets can be substantial. In a supersonic expansion, the Doppler width of an absorption line is determined by the spread of the velocity component parallel to the laser beam, which is (usually) perpendicular to the direction of propagation of the jet expansion. When a gas mixture is expanded through a circular nozzle, the linewidth is determined by the average free flow velocity. In the case of a gas mixture with He as a major component, the resulting linewidth can be even larger than the Doppler width at room temperature. The linewidth can be much reduced by using planar jet expansions (slit-jet expansions).

Some disadvantages of molecular jets are the short absorption path (a few centimeters at most), the low local density \((10^{18}–10^{19} \text{ molecules cm}^{-2})\), the strong variations in local density, the nonequilibrium conditions, the large gas consumption, and the necessity for large pumps. A way to obtain an appreciable local density without the need for large pumps is to use a pulsed jet expansion. This also reduces the gas consumption, which opens up possibilities for using costly isotopic species in a supersonic-pulsed jet experiment.

Hitherto, the spectroscopic study of molecules that are in the liquid or solid state at room temperature was limited either to matrix spectroscopy or to gas-phase spectroscopy by using Knudsen cells (Wagner 1984). The advent of molecular beams opened new possibilities, such as evaporating liquids and solids and allowing the vapors to be entrained in a supersonic expansion.
Pure carbon clusters can be produced in the laboratory by laser ablation of a graphite target (Scott et al. 2001 and references therein). Followed by an adiabatic supersonic expansion, clusters grow in size and appear in a variety of structures including linear chains, rings, and three-dimensional cage forms (Kroto et al. 1985, Kroto 1997). Unstable species such as radicals and molecular ions can be readily produced by combining a discharge with a pulsed supersonic (slit) jet. The transient species are successively cooled and transported in a collisionless environment. Concentrations of $10^{12}$ radicals cm$^{-3}$ and $10^{10}$ ions cm$^{-3}$ are obtainable with these techniques (Mazzotti et al. 2008, Guennoun and Maier 2011: Electronic Spectroscopy of Transient Molecules, this handbook).

A further important development concerns the formation of liquid He droplets in molecular beams, which are then used as an almost inert matrix for molecules at very low temperatures (Toennies and Vilesov 2004). This work is reviewed in the present handbook by Callegari and Ernst (2011): Helium Droplets as Nanocryostats for Molecular Spectroscopy—from the Vacuum Ultraviolet to the Microwave Regime, this handbook. It is also possible to use gaseous He as an inert “solvent”, providing a cooling mechanism under both equilibrium and nonequilibrium conditions (Willey et al. 1989, Bauerecker et al. 2001, Albert et al. 2007). We have shown that with this more recent development FTIR spectra of gaseous molecular samples can be measured at high resolution and temperatures as low as 10 K and lower (Albert et al. 2007). This work is reviewed briefly in Albert et al. (2011): High-resolution Fourier Transform Infrared Spectroscopy, this handbook.

1.3 Organization of the Article

This article is organized as follows. In Section 2, we summarize the general aspects and experimental methods of the molecular spectroscopy of supersonic jets, including some of the basic equations for supersonic-jet expansions. We also review some of the basic experimental setups. In Section 3, we give a review of the FTIR-supersonic jet experiments. Section 4 reports the properties of a number of infrared laser sources used in supersonic-jet spectroscopy.

In Section 5, we give a tabular literature overview of FTIR supersonic-jet spectroscopic investigations at high resolution. In Section 6, we discuss some selected examples in more detail. Section 7 presents the application of high-resolution infrared spectroscopy of supersonic jets to the study of nuclear spin symmetry conservation and violation. Section 8 reports a series of studies of van der Waals clusters and hydrogen-bonded molecular complexes. Section 9 is dedicated to radicals and ions. The field has developed to an extent that an encyclopedic review is not really possible or desirable here. We have tried to concentrate on the most important aspects and applications, with presumably some weighting toward our own work, partly understandable from the historical development of the field, partly by the natural bias of specific knowledge, although we have made an effort to cover work from all groups. With apologies to those authors whose work is less than adequately covered, we mention here some relevant reviews, one that covers much of the early work (Quack 1990), and the other the general beam techniques in two volumes (Scoles 1988, 1992). The subsequent work of the group of Herman at the Université Libre de Bruxelles is well covered by the review of Herman et al. (2000). The work on ($\text{H}_2\text{O}$)$_n$ clusters can be found in reviews of Saykally and coworkers (Liu et al. 1994, 1996, 1997, Gregory et al. 1997, Paul et al. 1997, Keutsch et al. 2003). Some of the work of the Leutwyler group is covered in this handbook by Frey et al. (2011): High-resolution Rotational Raman Coherence Spectroscopy with Femtosecond Pulses, this handbook. The theoretical work related to experiment is far too exclusive to be covered here, but several articles in this handbook deal with the theoretical aspects. We draw attention to two reviews covering experimental and theoretical work on (HF)$_n$ clusters (Quack and Suhm 1997, 1998). For work on various types of clusters, we refer to Kappes and Leutwyler (1988). Work on molecular beam high-resolution laser spectroscopy with optothermal detection has been reviewed by Lehmann et al. (1994) (see also Scotoni et al. 1991), and the article by Demtröder (2011): Doppler-free Laser Spectroscopy, this handbook. The interesting applications of the results to short-time intramolecular dynamics are not covered here and we refer to Marquardt and Quack 2001, 2011: Global Analytical Potential Energy Surfaces for High-resolution Molecular Spectroscopy and Reaction Dynamics, Quack (2001, 2003) and Hippler et al. 2011: Mass and Isotope-selective Infrared Spectroscopy, this handbook and Albert et al. (2011). For sake of completeness, we draw explicit attention to all these reviews for further information.

2 GENERAL ASPECTS AND EXPERIMENTAL METHODS OF THE MOLECULAR SPECTROSCOPY OF SUPTSONIC JETS

2.1 Fundamentals of Supersonic Expansion

For the general background, we refer here to the two volumes of the handbook edited by Scoles (1988, 1992) and review here some fundamentals of supersonic expansion, as these are important for high-resolution spectroscopy (see...
An expansion of a gas or liquid through an orifice into vacuum is supersonic if the velocity of the atoms or molecules in the expansion is larger than the local velocity of sound. The ratio of the velocity of the gas \( v \) with respect to the local velocity of sound \( v_{\text{sound}} \) is also called the Mach number.

\[
M = \frac{v}{v_{\text{sound}}}
\]  

In the case of an ideal gas, the speed of sound becomes

\[
v_{\text{sound}} = \sqrt{\frac{\gamma k T}{m}}
\]

where \( \gamma \) is the ratio of heat capacities \( C_p/C_v \), \( k \) is the Boltzmann constant, \( T \) is the temperature, and \( m \) is the molecular mass. For a monoatomic gas \( \gamma = 5/3 \) and for a diatomic gas \( \gamma = 7/5 \), in a simple limit.

Now we consider an isentropic flow of a perfect (ideal) gas, from a reservoir at pressure \( P_0 \), through a circular orifice with diameter \( d \), into a vacuum chamber that is kept at a low background pressure \( P_b \). The chamber is evacuated by a turbomolecular or a diffusion pump. The expanding gas while expanding will increase its speed and will cool down, and we can define a local temperature \( T \), density \( n \), and velocity \( v \) anywhere in the expansion. Roughly, we can distinguish several regimes or zones in the expansion (Figure 1); behind the orifice or nozzle, we have a zone where many collisions occur and where cooling and cluster formation is important. Once the gas density drops and collisions become rare, we have the so-called zone of silence, where the gas is expanding undisturbed, until it reaches a distance where the local density becomes similar to the density resulting from the background pressure: the Mach disk shock. The borders of the axisymmetric expansion consist of compression waves that are limited by a barrel shock, which has the shape of a paraboloid centered on the jet axis. Most spectroscopic measurements involve the regions close to the nozzle orifice or the zone of silence where we have free molecular flow.

Therefore, it is important to know where the Mach disk is located. In an ideal expansion of a gas, the final, or terminal velocity of the molecules is given by

\[
v_t = \sqrt{\frac{2\gamma k T_0}{\gamma - 1} \frac{1}{m}}
\]

where \( T_0 \) is the temperature of the source.

Once the Mach number is known, other important parameters can be calculated, by using the equations (1)–(3) and the ideal gas law, such as the local density, pressure, and temperature \( n, P, \) and \( T \) from the equations

\[
n = n_0 F^{-1/(\gamma - 1)}
\]

\[
P = P_0 F^{-\gamma/(\gamma - 1)}
\]

\[
T = T_0 F^{-1}
\]

\[
v = M \sqrt{\frac{\gamma k T_0}{m}} F^{-1/2}
\]

where \( F = 1 + (\gamma - 1)M^2/2 \).

The Mach number in turn depends on the geometry of the expansion and can be generally expressed as a function of the distance from the nozzle \( x \) and of the orifice diameter \( d \).

\[\text{Figure 1} \quad \text{Schematic picture of a free jet expansion (After Miller 1988).} \quad P_0 \text{ and } T_0 \text{ are source pressure and temperature.} \quad M \text{ is the Mach number, as defined in the text.}\]
2.2 Axisymmetric and Planar Expansions

The most common expansion geometries are axisymmetric or planar. An axisymmetric expansion is produced when the gas expands through a circular orifice. In this case, the maximum density is found on the flow line and this decreases rapidly when the distance between source and the flow line increases. The corresponding Mach number can be described as

\[ M = \left( \frac{x}{d} \right)^{(y-1)/2} + b \left( \frac{x-x_0}{d} \right)^{-(y-1)/2} \]  

(8)

where \( A \), \( a_1 \), \( a_2 \), and \( a_3 \) are empirically determined numbers. It is easy to see that for \( M \gg 1 \) the density decreases with increasing distance from the source as \( x^{-2} \).

The Mach disk location is given by

\[ \frac{x_M}{d} = \frac{2}{3} \sqrt{\frac{P_0}{P_b}} \]  

(9)

where \( x_M \) is the distance of the Mach disk from the nozzle, \( P_b \) is the background pressure, and \( d \) is the diameter of the nozzle.

A planar jet consists of a slit nozzle opening with a length \( L \) and a width \( d \), where \( L \gg d \) (Figure 2). Planar jets are usually operated in the pulsed mode although experiments with continuous slit jets have been reported as well (Amrein et al. 1988b). In the case of an ideal slit nozzle, the gas density approximately decreases as \( n(x) \sim x^{-1} \).

The Mach number for planar jets, for \( L \gg d \) and \( x \ll L \), and an adiabatic isentropic expansion is given by

\[ M = A \left( \frac{x-x_0}{d} \right)^{(y-1)/2} + b \left( \frac{x-x_0}{d} \right)^{-(y-1)/2} \]  

(10)

where \( A \) and \( b \) are empirical parameters.

The location of the Mach disk can be expressed as (Beylich 1979)

\[ \frac{x_M}{d} = \frac{3}{2} \sqrt{\frac{P_0}{P_b}} \left( \frac{L}{d} \right)^\epsilon \]  

(11)

where \( \epsilon \) is a number between 0.47 and 0.735 and tends to smaller values for \( L \gg d \), where \( L \) is the length of the slit.

The asymptotic expressions for \( T, n \) are \( (d \ll x \ll L) \)

\[ \frac{T}{T_0} = C \left( \frac{x-x_0}{d} \right)^{(y-1)} \]  

(12)

\[ \frac{n}{n_0} = G \left( \frac{x-x_0}{d} \right)^{-1} \]  

(13)

where \( C \) and \( G \) are empirical parameters. Detailed formulas for the axisymmetric and planar free jets can be found in Hagena (1981). An experimental investigation of supersonic slit jets is reported by Beylich (1979).

One of the major limitations of the supersonic-jet expansion through a nozzle is the short effective absorption path. One way to enhance the absorption path length is multipassing the laser beam through the expansion, but this is also troublesome because of the small dimensions of the nozzle. A planar (slit) expansion produces a longer absorption path with the same gas throughput, can be used in combination with multipass optical setups, and has the additional advantages of a narrower linewidth, a slower
cooling, and a more homogeneous density distribution along the laser path. The slit nozzle has also been successfully coupled to several discharge sources in order to produce molecular ions and radicals. With respect to an axisymmetric expansion, the molecular density $n$ and the temperature decrease more slowly, providing larger two- and three-body collision rates, which enhance the cluster production. At the same time, it also guarantees a better cooling of the vibrational degrees of freedom. The Doppler width is substantially reduced because of the quenching of velocity components parallel to the slit.

### 2.3 Pulsed Jets and Continuous Jets

Both pulsed and continuous supersonic jets have been used to perform high-resolution spectroscopic measurements. Each method has its advantages and drawbacks. A continuous expansion requires large quantities of gas and elevated pumping speeds, whereas the gas consumption in pulsed experiments is limited and smaller pumping systems are sufficient to obtain a low background pressure between two gas pulses. As a consequence, pulsed experiments allow for higher reservoir pressure, which is important for cluster production and for slit nozzles with a larger surface. Generally, slit-jet expansions produce a higher molecular density behind the (slit) nozzle and eliminate most of the Doppler broadening of the absorption lines, owing to the velocity quenching parallel to the slit opening. Operating a continuous slit expansion would require very large pumps. Pulsed expansions are often combined with pulsed laser sources, for instance, in laser ablation or laser desorption experiments. In addition, the production of ions and radicals is often obtained by pulsed pyrolysis, ablation, or photolysis of precursors, which are obviously more efficiently combined with a pulsed expansion. However, when spectral scans and averaging processes require long recording times, the reduced duty cycle of a pulsed expansion with respect to a continuous jet can be a serious drawback.

### 2.4 Seeded Expansions

To obtain a more efficient cooling, a small amount of the heavier seed gas is often seeded in a lighter carrier gas. In the case of a binary mixture, one can easily obtain expressions for the average mass and velocities ($n_s$ and $n_c$ are the number densities of seed and carrier gas, respectively):

$$m_{av} = \left( \frac{n_c}{n_c + n_s} \right) m_c + \left( \frac{n_s}{n_c + n_s} \right) m_s \quad (14)$$

which for $n_c \gg n_s$ becomes

$$m_{av} \approx m_c + \left( \frac{n_s}{n_c} \right) m_s \quad (15)$$

whereas the most probable velocities become

$$v_c = \frac{1}{2} \left[ u_c + \sqrt{u_c^2 + 8kT_c/m_c} \right] \quad (16)$$

$$v_s = \frac{1}{2} \left[ u_s + \sqrt{u_s^2 + 8kT_s/m_s} \right] \quad (17)$$

At high source pressures, the seeded gas reaches the same velocity as the carrier, and for medium and low pressure, there is a velocity slip.

### 2.5 Pick-up Methods

While seeded expansions are used for expanding the seed gas diluted in a mixture with noble gas, a different solution has to be found to introduce in the expansion the molecules that cannot be mixed beforehand. This is the case of solids and liquids, which can be (laser) evaporated close to the nozzle exit and successively entrained in a supersonic expansion. The group of Giacinto Scoles developed the pick-up method to study the behavior of infrared chromophores in a solvent of noble gas clusters (Gu et al. 1990). The same authors crossed a molecular beam of CH$_3$F seeded in Ar with a second beam of HCl to study complex forming reactions in Ar clusters (Levandier et al. 1987). Refractory radicals, such as CaCl, can be entrained in a molecular expansion by crossing a noble gas expansion with a conventional effusive molecular beam (Steimle et al. 1991). In this way, the radicals can be cooled from 1200 down to 7.6 K, which simplifies their spectra enormously.

Thermally labile and nonvolatile species, such as biomolecules, can be brought in the gas phase by laser desorption and successively entrained in an expansion of He (Tembreull and Lubman 1987, Cable et al. 1987).

In the case of ions and radicals, a discharge source can be used to produce a large number of ions and radicals (Kim et al. 2005, Davis et al. 2001), which are picked up by the expansion and are cooled efficiently. To produce a variety of carbon clusters, a carbon rod is ablated by a pulsed laser and the products are introduced in the expansion. The field of the spectroscopy of molecules dissolved in cold He clusters has developed along independent lines (see the review by Toennies and Vilesov (2004) and Jäger and Xu 2011: Fourier Transform Microwave Spectroscopy of Doped Helium Clusters, this handbook).
2.6 Optical Setup

The molecular density in a molecular expansion is usually low and the absorption path very short, with respect to conventional cell measurements. One way of increasing the absorption in a jet is the so-called multipass setup, which consists of passing the laser beam several times through the supersonic expansion, possibly all at the same angle with respect to the flow velocity of the molecules and in the same (small) volume. Several schemes have been used, but we discuss only the most popular ones.

The most straightforward setup uses two flat parallel mirrors to pass the laser beam almost perpendicularly through the expanding jet (Figure 3a). This method is simple, but its main drawback is that the successive laser crossings are at increasing distances from the nozzle and hence probe different densities and populations at different temperatures.

A very interesting scheme has been proposed by Kaur et al. (1990), which consists of two concave mirrors at a distance that is slightly less than four times their focal length (Figure 3b). The laser beam is focused at the center of the cavity with a lens (or mirror) with a larger focal length than that of the two cavity mirrors and is refocused on every pass in a very small volume that intersects the molecular beam. The most advantageous configuration is the one where all successive passes are squeezed in a horizontal plane perpendicular to the flow.

Figure 3 Three different optical multipass setups used in combination with molecular jets: (a) two plane parallel mirrors; (b) two concave mirrors; and (c) White-type optics.
velocity of the jet. The alignment of this configuration is rather troublesome and requires some expertise (and patience) because of the difficulty to refocus the out-coming beam on the detector. Nevertheless, this setup has been successfully used to measure the ν6 fundamental of CF3Cl2 (Snels and Meerts 1988). The third scheme is based on compact White-type optics (White 1942). The incoming laser beam is only softly focused with a 1–2 m focal length mirror on the entrance of the White cell and is refocused several times on the field mirror by two concave mirrors at a distance of about 150 mm. Here, the familiar White cell pattern is formed and eventually a modestly diverging laser beam is focused onto a detector with an off-axis parabolic mirror. The interesting thing is that close to each of the two focusing mirrors, all odd (respectively even) passes cross in a very small volume (less than 2 × 2 × 20 mm3). The nozzle is placed close to one of the focusing mirrors and a very good orthogonal crossing of the laser with the molecular jet can be obtained (Figure 3c). The number of crossings for each setup is typically 15–20, taking into account that in the flat-mirror setup, the successive crossings are less effective due to the decreasing density further from the nozzle. The main advantages of the White configuration with respect to the confocal mirror setup are the better optical stability and the possibility to vary the number of passes without changing the direction of the output beam (see also the details of the ETH Zürich setups described in Figures 7 and 8).

### 2.7 Cavity Ring Down (CRD) Spectroscopy in Supersonic Jets

As an alternative to multiple pass setups, which allow at most 20–50 passes through the expansion, cavity ring down and cavity-enhanced spectroscopy have been used to measure absorption spectra in supersonic jets. Both techniques rely on the use of resonant cavities, consisting of two highly reflective mirrors (typically R > 99.95%), resulting in an increase of the effective absorption path length by a factor of several thousands. A key development was the combination of pulsed jets with continuous wave (cw) tunable near-infrared diode lasers for high-resolution CRD spectroscopy by Hippler and Quack (1999). The combination of high-resolution cw lasers (1 MHz bandwidth and better) with pulsed jets is nontrivial because for narrow bandwidths of the laser radiation the cavity has to be matched to the laser wavelength. When the gas pulse arrives, the resonance matching is lost because of the change of the index of refraction. This difficulty has been overcome by the Zürich group with a random wobble technique for one of the mirrors (He et al. 1998, Hippler and Quack 1999). These authors recorded very weak rovibrational lines of the ν1 + ν3 band of nitrous oxide near 7780 cm⁻¹, (HF) stretching overtones in (HF)₂ (Hippler et al. 2007), as well as an overtone level of the methane icosad (Hippler and Quack 2002). Tam et al. (2006) used a cw tunable lead-salt diode laser to measure the absorption of the CH-stretching band of methane in a supersonic jet. Other groups employed pulsed tunable lasers to investigate molecules and clusters in a supersonic-jet expansion. The group of Richard Saykally investigated a number of polycyclic aromatic hydrocarbons (PAHs) in the CH-stretching region (Schlemmer et al. 2011). They used a Nd : YAG-pumped dye laser to generate tunable pulsed visible radiation, which was then shifted by three Stokes shifts to produce infrared radiation with a bandwidth of 0.2 cm⁻¹, which could be reduced to 0.04 cm⁻¹, by using an intracavity etalon in the dye laser (Henneycutt et al. 2004). Bisson et al. (2007) generated pulsed tunable infrared laser radiation by difference frequency generation and recorded absorption spectra of the ν1 + ν3 and 2ν3 bands of several SiH4 isotopomers expanded in argon, helium, and hydrogen.

Ito and Nakanaga (2002) recorded CRD spectra of the formic acid dimer, including the OH stretching mode. The authors propose different mechanisms to explain the broad spectral features observed. See also Havenith and Birer 2011: High-resolution IR-laser Jet Spectroscopy of Formic Acid Dimer, this handbook.

CRD spectroscopy requires accurate mode matching and spatial filtering of the laser modes, in order to obtain single exponential decays. Other varieties of CRD-type spectroscopy exist with less strict requirements. One of these is the so-called cavity-enhanced absorption spectroscopy (CEAS). In the CEAS method, the optical cavity and laser wavelength are modulated in order to obtain many resonances, producing an almost continuous wavelength transmission with a long effective optical path length. Berden et al. (1999) used this CEAS technique to measure the absorption spectrum of ammonia in the 1.5 μm region in a supersonic seeded jet. The technique of Hippler and Quack (1999) has been successfully extended to the spectroscopy of ions by Birza et al. (2002).

### 2.8 Supersonic Jets Versus Alternative Techniques

The main advantages of doing absorption spectroscopy in a supersonic jet are the following. First, the efficient cooling of the internal degrees of freedom narrows the rotational distribution by populating only the lowest rotational energy levels and also removes population from most of the excited vibrational levels, transferring the population to the ground vibrational level. This leads to a significant reduction in the
The second advantage of the supersonic beam is the possibility to produce van der Waals clusters. This can also be done in a cold cell, but always within a limited temperature range, owing to the condensation limit. The collisional cooling technique is more difficult to apply for specific cluster production due to the low collision rates.

Radicals and ions can also be produced in drift tubes, using radio frequency techniques, but the temperature is usually very high (the rotational temperature is in the order of hundreds Kelvin), which renders the spectra complex and difficult to analyze (Gudeman and Saykally 1984, Sears 1987).

3 FTIR SPECTROSCOPY OF SUPersonic Jets: INSTRUMENTATION

The combination of high-resolution FTIR spectroscopy and cooling in supersonic expansions started in the 1980s. A survey of the basics of these two techniques and of the instrumentations developed in this field is given in the reviews by Quack (1990) and Herman et al. (2000), considering the main developments up to 2000. In a preliminary version of the Zürich group experiments, a jet setup was interfaced with a Bomem DA002 high-resolution spectrometer (Dubal et al. 1984). The jet assembly was placed outside of the spectrometer. The jet chamber was equipped with a circular nozzle of 100 µm diameter and was evacuated by means of a 3000 dm³ s⁻¹ oil diffusion pump backed by a 100 dm³ s⁻¹ roots pump and a 18 dm³ s⁻¹ rotary pump. By means of a transfer optical interface, the supersonic jet was crossed by the focused infrared beam in front of the nozzle. Spectra of CO₂ and CH₄, measured with apodized resolutions up to 0.006 cm⁻¹, revealed strong rotational cooling with effective rotational temperatures near 10 K in the supersonic expansions. These results encouraged further developments of this challenging technique.

In the second generation setup, the jet assembly was implemented into the sample compartment of the Bomem DA002 (Amrein et al. 1988b). A sketch of this arrangement is shown in Figure 6. The nozzle is positioned at a distance of a few millimeters in front of the focused infrared beam. An xyz translational stage allow for external adjustment of the nozzle position with respect to the infrared beam. In this version, there is no need for transfer optics to interface jet setup and spectrometer and therefore no light losses due to additional optical components and to their critical alignment. The pumping systems of the jet setup and of the spectrometer are completely separated. The latter consisted of a 14 m³ s⁻¹ oil diffusion pump and forepumps as in the
High-resolution rovibrational spectroscopy provides important information on the dynamics of coupled vibrational states and provides access to the modeling of short-time quantum dynamics in molecular systems. Supersonic-free jet expansions allow for the generation of molecules at low rotational and to a lesser extent vibrational temperatures. The combination of jet cooling with broad band FTIR spectroscopy provides a promising means to get access to a detailed analysis of complex rovibrational spectra. This was one of the main purposes in developing this challenging technique. A serious drawback of its application lies in the limited sensitivity due to the small absorption path length (in single path systems) and the low molecular densities in the detected region of the expansion. This limits the use of this technique to strong absorptions. Furthermore, because of less effective vibrational cooling, the simplification of the rovibrational structures in larger molecules is less effective due to the remaining contributions of hot-band spectra. A further limitation in the application of this technique lies in the need for large amounts of substance and in the requirements for large pumping power.

Sensitivity can be improved by using a multipass system, i.e., by increasing the absorption path length using optical tools. Several groups have reported such systems. Asselin et al. (1996) introduced a multireflection setup, which is similar to the White optics known from long-path gas cells. McNaughton et al. (1994) developed a multipass reflection setup based on a star-shaped arrangement of the light beams. A multipass system based on multiple reflections between two spherical mirrors was developed by Petry et al. (2002). A recently reported setup for investigating jet-cooled molecules, named FANTASIO (Herman et al. 2007), includes a multipass system closely analogous to the system by Petry et al. (2002).

We have developed a multipass accessory for our jet setup. Because of space limitations in the sample compartment of the spectrometer, we built a third-generation FTIR-jet setup with multipass optics and jet assembly located outside of the spectrometer. The main parts of this new version are sketched in Figure 7. We completely separated spectrometer and jet setup and therewith minimized mechanical couplings and distortions the very sensitive spectrometer may suffer from.

A casting containing the components of the optical interface is connected to the right port beam output of the Bomem DA002. This interface is based on f/4 optics similar to the optics of the spectrometer. It focuses the infrared beam into a newly designed jet assembly that contains a multipass system based on White optics. The design of this part is sketched in Figure 8. It consists of a plane mirror $M_1$ for the light input, three spherical mirrors $M_2, M_3, M_4$ of the White system, and finally a plane mirror $M_5$ for the beam output. This design is strictly similar to the original multipass optics of White (1942). The radii of the spherical mirrors corresponding to the distance between the objective mirrors $M_2, M_4$, and the field mirror $M_5$ are chosen to be 5.1 cm. This short distance allowed for a compact construction of the multipass system and a straightforward combination with parts of the previous version as the casting containing gas inlet and nozzle positioning equipment. Furthermore, the short distance between the mirrors keeps the path of the light beam outside the jet region, and therewith the fraction of signals due to warm background molecules relatively small. The aperture images on the mirrors $M_1, M_3$, and $M_5$ form a row of equally spaced spots. The nozzle is positioned such that the jet passes a few millimeter above the spot row. The number of passes may be altered in steps of four by tilting of the mirror $M_4$. The maximum number of passes is limited by the diameter of the spots and by the length of mirror $M_3$ in the direction of the spot row. This length is chosen to be 1 cm. This choice proved to be a reasonable compromise in view of several drawbacks connected with a larger length, such as the $d^{-2}$ drop of the molecular density in the expansion region, the larger fraction of warm spectra, the larger temperature span in the detected expansion region, and last but not the least, light
losses due to critical alignment of the optical components. Good conditions were obtained with the number of passes set to 4 or 8.

The combination of supersonic jets with FTIR spectroscopy usually relies on continuous expansions, in contrast to laser spectroscopy, where mostly pulsed jets are used. Pulsed jets allow for a wider range of expansion conditions, which, for instance, is useful for the investigation of molecular clusters. The coupling of FTIR spectroscopy with pulsed nozzles requires complex techniques in order to properly synchronize jet pulsation with the data acquisition of the interferograms. Jet pulsation is operated typically with a repetition rate of 50 Hz and a pulse width of about 2 ms, whereas the data-acquisition rate for a measurement in the mid-infrared region typically amounts to 15 798 Hz (derived from He–Ne laser fringes, corresponding to a mirror speed of 0.5 cm s⁻¹). A technique that uses an asynchronously pulsed supersonic jet in conjunction with an FTIR spectrometer was developed and applied in an investigation of HF-stretching spectra in HF clusters in our laboratory (Luckhaus et al. 1995). Preliminary measurements were made with NH₃, N₂O, and CH₄ in the mid-infrared region in order to test the applicability of this technique. Extended model calculations simulating the pulsed operation were performed for a synthetic interferogram corresponding to an artificial spectrum containing two lines on one hand and for a manipulated interferogram of the ν₂ band of NH₃ on the other. The results revealed that artifacts occurring in single scan experiments tend to cancel rapidly during the averaging process even for a small number of scans. In the case of the ν₂ band of NH₃, averaging over four scans leaves minor artifacts though the main features of the spectrum are clearly recognizable, and averaging over 64 scans reproduces the original (unmanipulated) spectrum almost perfectly. The possibilities of this technique merit further exploitation.
4 INFRARED LASERS FOR SUPERSONIC-JET SPECTROSCOPY

4.1 Mid-infrared Lasers

The mid-infrared spectral region (2–30 µm or about 300–5000 cm\(^{-1}\)) provides a broad field for various spectroscopic applications because it covers the fundamental vibrational modes of many polyatomic molecules except for very low frequency modes. For such spectroscopic measurements, an appropriate laser source has to be selected. The demands on such sources depend on the application. Usually broad tuning (emission), cw tuning, and sufficient optical power are required. The output power is usually in a range of microwatts to milliwatts except for CO and CO\(_2\) lasers, which emit up to kilowatts of optical power in cw operation. A narrow linewidth is important for high measurement sensitivity and selectivity. The ideal laser also has single longitudinal mode behavior and small beam divergence. Moreover, compactness, robustness, and room temperature operation are important for in situ applications. Currently, various mid-infrared sources exist. They can be distinguished according to their active laser medium into gas, semiconductor, and solid-state lasers, which directly generate mid-infrared radiation or are based on nonlinear optical paramagnetic frequency conversion of near-infrared laser sources (Tittel et al. 2003). The most frequently applied lasers for supersonic-jet techniques are semiconductor lasers.

4.1.1 Semiconductor Lasers

Lead-salt diode lasers, quantum cascade lasers (QCLs), and antimonide lasers belong to this class, which are direct laser sources sharing some common features, for instance, laser cooling and similar collimation optics (Werle et al. 2002).

Lead-salt Diode Lasers

Lead-salt diode lasers consist of various nonstoichiometric binary alloys of the Pb compounds with IV–VI elements of the periodic table. These lasers can emit infrared light from 3 to 30 µm according to their composition and are commercially available for about 500–3300 cm\(^{-1}\). Laser light is generated by electron–hole recombination in the p–n junction of n-doped and p-doped semiconductors. The optical resonator is formed by cleaved facets. Electrons and holes drift toward the depletion region on the interface between n and p semiconductors where they recombine. Electrons occupy the bottom of the conduction band and are separated from the holes present in the valence band by the band-gap energy. The photon created by the electron–hole recombination has an energy that corresponds to the band-gap size. The band gap of the lead-salt lasers is narrow (i.e., the energy separation of the conduction and valence band is small), leading to a thermal population of the conduction band at room temperature. To prevent this effect, lead-salt lasers are typically cooled in a cryostat either by liquid nitrogen or by helium. The emission wavelength of the laser changes with its temperature. Fine tuning of the wavelength can be obtained by modulating the current applied to the diode laser. Depending on the tuning rate in terms of cm\(^{-1}\) mA\(^{-1}\), an elevated stability of the injected current may be required. During the continuous wavelength tuning, mode hops may occur, which implies that the laser may emit more than one wavelength at the same time. In multimode operation of a laser, simultaneous oscillation on many modes can be observed. The transition from single mode to multimode operation can be discrete, but more continuous transitions can also be observed with the optical power being gradually redistributed. One may also obtain mode hops to higher order modes, or mode hops between modes with different polarization in lasers with polarization-independent gain. To select one wavelength during multimode operation of the laser, dispersive elements, such as a grating, can be used. Lead-salt diode lasers can be usually tuned around 100 cm\(^{-1}\) by varying the temperature and injected current. At a fixed temperature, a continuous tunability of about 0.5–2 cm\(^{-1}\) can be obtained by injection current modulation. Lead-salt diode lasers also exhibit quite strong beam divergence and astigmatism, which considerably complicate the alignment of the laser beam, in particular when multipass optics are used. This can be overcome partially by implementing special optical elements (e.g., parabolic mirrors for compensating for the beam divergence). For more information about lead-salt diode lasers, see the review from Brassington (1995) and the article by Sigrist (2011): High-resolution Infrared Laser Spectroscopy and Gas Sensing Applications, this handbook.

Antimonide Diode Lasers

Lasers based on III–V semiconductors emit in a region between 2 and 5 µm (2000–5000 cm\(^{-1}\)). The first successful mid infrared laser was based on InGaAsSb double heterostructure (DH)-active region grown by liquid epitaxy, operating at room temperature and emitting at 2.2 µm (Caneau et al. 1985). Lately, the material quality and, hence, the performance of the lasers have been improved by using molecular beam epitaxy (MBE). Because the DH-active region limits the device’s performance, such as high threshold current, low output power, and low maximum operating temperature, quantum well (QW) structures were used in the active region of the laser, which improve these properties. A QW acting as a potential well is a very
thin middle layer where the vertical variation of the electron’s wavefunction, and thus a component of its energy is quantized. Lasers containing more than one QW layer are known as multiple QW lasers. Multiple QWs improve the overlap of the gain region with the optical waveguide mode. These lasers dispose of relatively high-optical output power up to 20 mW in comparison with lead-salt diode lasers. Antimonide QW diode lasers emitting light at 2–3 µm usually can operate at room temperature and in cw regime. Sources operating at 3–5 µm usually require operation at reduced temperature. For more details, see reviews from Joullié et al. (2003) and Yin and Tang (2007).

Quantum Cascade Laser (QCL)
The QCL operates between 3 and 15 µm (about 700–3300 cm⁻¹), which includes the so-called atmospheric window absorption as well as the “fingerprint” range of the infrared and thus can be a powerful tool for applications in this spectral range. After 15 years of development, QCLs are now available for cw operation at room temperature and have high optical output power. They demonstrate a large tuning range of up to 400 cm⁻¹ (Hugi et al. 2009). On the other hand, they, like all semiconductor lasers, have quite large beam divergence and astigmatism. The main difference between QCLs and standard intersubband lasers is the cascading principle. The QCL is formed by 20–40 identical periods. An electron injected from one side passes through the first period, emits a photon, and is injected into the following period. Thus, one electron can emit more photons before leaving the structure. Moreover, no holes are involved in either electron transport or photon emission. However, the internal quantum efficiency is lowered due to the presence of nonradiative scattering mechanisms. One period is a nonperiodic alternating sequence of thin layers of two different semiconductors, several nanometers thick. Different values of the band gap of the two semiconductors provide the conduction band profile that forms multi-QWs in the direction perpendicular to the layers (growth direction). The quantum confinement gives rise to the intersubbands whose energy can be tailored by changing the QW/barrier thicknesses. The laser emission wavelength is determined by the subband spacing and, therefore, can be directly changed by changing the layer thicknesses. The electrons are injected into the upper state (subband) of the laser transition by resonant tunneling through the thick (injection) potential barrier. It undergoes radiative transitions from the upper subband to the lower subband, emitting the excessive energy in the form of photons. The electrons tunnel away through another potential (extraction) barrier and are injected into the upper state of the next period. The population inversion necessary for the lasing action is ensured by long upper and short lower lasing state lifetimes. The electron lifetimes can be tailored by the proper design of the active region. The main drawback of this approach is the fast carrier’s nonradiative relaxation from the higher states (subbands) due to longitudinal optical (LO) phonon-assisted scattering, which decreases the radiative efficiency due to the depletion of the upper lasing state, thus lowering the electron inversion. For a more detailed description of the QCLs and their operation characteristics, design, and their application, see Hofstetter and Faist (2003), Capasso et al. (2000) and Faist et al. (1994a,b). An application of QCLs for infrared spectroscopy of jet-cooled molecules and complexes is reported by Xu et al. (2009).

4.1.2 Difference Frequency Generation (DFG) and Optical Parametric Oscillator (OPO)

DFG is based on parametric frequency conversion of near-infrared sources. The nonresonant optical setup includes two laser beams (pump and signal). The emission of the pump laser at higher frequency is combined with a second laser emission (signal) with a lower frequency, in order to generate a different frequency (idler) in a nonlinear optical material such as LiNbO₃ and periodically poled lithium niobate (PPLN). Since the idler wave is build up when the beams pass through a nonlinear material, all three waves must stay in phase. Tuning of the idler wave is performed by tuning the pump, signal beam, or both together. The optical parametric oscillator (OPO) uses nonlinear optical material in a resonant cavity to generate two laser beams (idler and signal) from one pump laser beam. The cavity can be resonant for either idler or signal beam, or for both. Optical parametric generation is a second-order nonlinear process and is less efficient than DFG. An OPO can operate in the pulsed regime from picosecond to femtosecond timescales or in cw mode. As mentioned above, DFG and OPO laser sources are based on nonlinear optical parametric frequency conversion of near-infrared laser sources and can operate in a spectral range up to 2 µm. Their operation depends on spectral and spatial properties of the lasers used for generation and also on properties of the nonlinear material. Frequently used lasers for these techniques are Nd : Yag, Ti : sapphire, external cavity diode laser (ECDL), and, for DFG, OPOs can also be used as pump or signal lasers. Mid-infrared nonlinear materials and some of their optical characteristics for DFG and OPO applications are summarized in reviews by Ebrahizadeh (2003) and Fischer and Sigrist (2003) (see Sigrist 2011: High-resolution Infrared Laser Spectroscopy and Gas Sensing Applications, this handbook). Very recent developments of high-power cw OPOs allow for very high resolution Doppler
4.1.3 CO₂ and CO gas laser

The CO₂ laser operates in the 9.2–10.8 μm range and is characterized by high output power. The laser can be tuned on discrete levels of the emission spectrum of CO₂. The gas mixture in a CO₂ laser consists of CO₂, N₂, and He. The N₂ molecule is excited by an electrical discharge to a metastable long-lived state and transfers its energy by collisions to an excited vibrational state of CO₂, thus creating the population inversion necessary for the stimulated emission of photons. The lower vibrational states of CO₂ are successively depopulated through collisions with He atoms. For more details, see Patel (1965), Witteman (1967), and Repond and Sigrist (1996). By substituting the ¹²CO₂ by ¹³CO₂ or by other CO₂ isotopomers, a dense manifold of laser emissions on discrete wavelengths can be generated in the 9–12 μm range (Freed et al. 1980). In addition, N₂O (Herlemont et al. 1979) can be used to produce laser emissions in the 10.4–11 μm range. Although the CO₂ laser is essentially a line tunable laser, several possibilities exist for using it as a continuously tunable laser. In the high-pressure pulsed (TEA) CO₂ laser, the gain profile is pressure broadened, allowing for an almost continuous tuning in the range 900–1100 cm⁻¹, although with a linewidth of about 2 GHz, which can be reduced by using an intracavity prism to about 250 MHz (Duarte 1985 and Repond and Sigrist 1996). A higher resolution can be obtained by using a CO₂ waveguide laser, which can be tuned over typically 1 GHz, with a linewidth better than 1 MHz (Olafsson and Henningsen 1995). A third option is to use a GaAs waveguide modulator to create side bands on the CO₂ laser line emissions up to a few gigahertz from the line center (Cheo 1994). The CO laser emits in the spectral range 5–6.5 μm. By using the overtone emission of CO, the 2.5 μm range can also be covered (Utkin et al. 2006). This laser works similarly to the CO₂ laser. With respect to the CO₂ laser, the CO laser offers only discrete line tunability, but has higher efficiency. This efficiency increases with decreasing gas temperature. Because of this, this laser requires more sophisticated cooling.

4.2 Near-infrared Lasers

The near-infrared region (0.8–2 μm or 5000–12 000 cm⁻¹) is almost completely covered by tunable diode lasers. Compared to the lead-salt diode lasers used in the mid-infrared, the NIR diode lasers do not require cooling, and being operated at ambient temperature, they have found many applications in medicine and telecommunications. The most commonly used diode lasers (Fabry Perot lasers) are DH lasers.

In these devices, a layer of low band-gap material is sandwiched between two high band-gap layers. One commonly used pair of materials is gallium arsenide (GaAs) with aluminum gallium arsenide (AlₓGa₁₋ₓAs). Each of the junctions between different band-gap materials is called a heterostructure and, hence, the name double heterostructure laser or DH laser.

The advantage of a DH laser is that the region where free electrons and holes exist simultaneously, the active region, is confined to the thin middle layer. This means that many more of the electron–hole pairs can contribute to amplification. Relatively few are located in the poorly amplifying periphery. In addition, light is reflected from the heterojunction; hence, the light is confined to the region where the amplification takes place. Typical dimensions of these emitting regions are 2 μm × 500 μm. The DH lasers are multimode lasers and emit several modes simultaneously. The tuning can be performed by temperature and current variations and can be as much as 100 nm. Lately, vertical cavity surface emitting diode lasers (VCSELs) have been developed, which have a short cavity, thus limiting the multimode character and a very small emitting surface (5 × 5 μm typically). Single mode tunable lasers can be obtained by introducing wavelength-selecting devices inside or outside the diode laser structure. In a distributed feedback (DFB) laser, a diffraction grating is etched close to the p–n junction of the diode. This grating acts like an optical filter, causing a single wavelength to be fed back to the gain region and lase. Since the grating provides the feedback that is required for lasing, reflection from the facets is not required. In the distributed Bragg reflection (DBR) lasers, the selective regions are combined with gain regions. Both DFB and DBR lasers are single mode lasers and can be temperature tuned, typically over few (3–10) nm. They have output powers of 10–100 mW and a linewidth of 3–10 MHz typically (Buus et al. 2005).

The ECDL uses an external grating to construct a selective cavity around a Fabry–Perot laser, which has an antireflection coating applied to one surface. Several configurations (Littrow, Littman/Metcalf) are commercially available and produce a single mode output of typically 10 mW, with a linewidth better than 1 MHz and a tunability up to 150 nm (see, for example, http://www.santec.com).

The gain medium determines the wavelength of the emitted radiation: 750–1000 nm for AlGaAs, 1.2–2.0 μm for InGaAsP. The most popular wavelengths are those used for medical applications (850–980 nm) and in telecommunication (1.31 and 1.55 μm). The Zürich group had built a homemade temperature-controlled laser of this kind in free double resonance spectroscopy in the infrared (Dietiker et al. 2010).
the 1980s when these lasers were not yet available commercially, but later this laser was replaced by the more powerful laser from Radians-Innova (Ha et al. 1995, He et al. 1995). The International Telecommunication Union has defined the so-called ITU grid, which covers the O, E, S, C, and L bands (1260–1625 nm), with a spacing of 100 GHz (about 0.8 nm). Recently, a second series with a spacing of 50 GHz has been introduced. Single mode DFB lasers covering the ITU grid are commercially available and are often used for high-resolution spectroscopy in the NIR. Commercial external cavity lasers cover the range from 760 to 1050 nm and from 1260 to 1680 nm. Often the linewidth of these lasers is far below the Doppler width of a molecule at room temperature. This implies that the Doppler line shape and width of an absorption line can be measured with high accuracy, allowing one to determine the temperature with a high precision from the linewidth, as can be seen in Figure 9. Thus, high-resolution laser spectroscopy of isolated spectral lines can be used as an in situ “thermometer” both for bulk samples and for supersonic jets (He et al. 2007).

5 AN OVERVIEW OF FTIR SPECTROSCOPIC SUPERSONIC-JET INVESTIGATIONS

An early review of the work of the Zürich group has been given by Quack (1990). A literature review dedicated to FTIR absorption spectroscopy of jet-cooled species covering also the period subsequent to the early developments up to the year 2000 was included in the review article of Herman et al. (2000). In Table 1, we present an updated review, which we have restricted to high-resolution investigations. We mention the following research groups active in this field with particular topics: the Université Libre de Bruxelles (hydrocarbons (Lee et al. 2007, Hurtmans et al. 2001, Lafferty et al. 2006, Fland et al. 2001)); Université de Rennes (small clusters (Thiévin et al. 2006)), Monash University (fluorohydrocarbons (Thompson et al. 2003)); Université Pierre et Marie Curie, Paris (XY4 and XY6 spherical tops (Asselin et al. 2008, Rey et al. 2001), clusters (Asselin et al. 2006, 2007)); and our group at ETH Zürich (simple molecules and spherical tops, methane (Dübal et al. 1984, Amrein et al. 1988a,b), ammonia (Snels et al. 2006a), substituted methanes (D’Amico et al. 2002), XY6 spherical tops and Jahn–Teller molecules (Boudon et al. 2002), fluoro(hydro, chloro)carbons, hydrogenfluoride clusters, and other applications (Snels et al. 1995, Luckhaus et al. 1995, He et al. 2007, Hippler et al. 2007)) and chiral molecules (Beil et al. 1994, Bauder et al. 1997) see also Albert and Quack (2007), Albert et al. (2011), and Quack (2011).

The large spectral coverage combined with strong rotational cooling makes the FTIR-jet technique an ideal means to investigate complex rovibrational band systems of stable species. The analysis of such spectra provides access to accurate spectroscopic parameters of the states involved and, in particular, to vibrational and rovibrational coupling parameters that are of great importance for the understanding and modeling of time-dependent quantum dynamics and processes such as intramolecular vibrational and rotational vibrational redistribution (Marquardt and Quack 2001, Quack and Kutzelnigg 1995). In Table 2 we give a survey of the investigations performed in our laboratory using the FTIR-jet technique, including also the further recent work not reviewed in Quack (1990), Herman et al. (2000). The listing in Table 2 contains information about the species investigated, relevant spectrometer settings, and jet parameters, the main focus of the study, and the band systems analyzed. Instructive examples demonstrating the high value of this method are the anharmonically coupled $v_1, 2v_5$ band system of CF$_3$I, the $v_3, v_5$ Coriolis resonance band system of CHClF$_2$, and the tunneling doublets of the bending fundamentals of the ammonia isotopomers NH$_2$D and ND$_2$H. We discuss

![Figure 9](https://example.com/figure9.png)

**Figure 9** Spectral line of the first HF-stretching overtone transition of HF monomer measured in an absorption cell at room temperature (upper trace) and in a supersonic-jet expansion (middle trace); the bottom trace shows the etalon signal. [Reproduced from He et al. 2007 by permission.]
Table 1  High-resolution FTIR investigations of molecular species in supersonic jet expansions. Literature review from the year 2000 up to the present.

<table>
<thead>
<tr>
<th>Species</th>
<th>Spectral range (cm(^{-1}))</th>
<th>Resolution (cm(^{-1}))</th>
<th>(T_{\text{rot}}) (K)</th>
<th>Band(s) investigated band center(s) (cm(^{-1}))</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH(_2)D</td>
<td>1150–1750</td>
<td>0.01</td>
<td>70–100</td>
<td>(\nu_{34}(s), 1605.6404) (\nu_{44}(a), 1591.0019) (\nu_{45}(s), 1389.9063) (\nu_{45}(a), 1390.4953)</td>
<td>Snels et al. (2006a)</td>
</tr>
<tr>
<td>ND(_2)H</td>
<td></td>
<td></td>
<td></td>
<td>(\nu_{44}(s), 1233.3740) (\nu_{44}(a), 1235.8904) (\nu_{45}(s), 1461.7941) (\nu_{45}(a), 1461.9918)</td>
<td></td>
</tr>
<tr>
<td>N(_2)O</td>
<td>Near 2200</td>
<td>0.0043</td>
<td>22</td>
<td>(R(0)) line of (\nu_{1}), line profile analysis</td>
<td>Didrique et al. (2007)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>vib-vib energy transfer between (v_{2}) states of N(_2)O and OCS</td>
<td>Herman et al. (2005)</td>
</tr>
<tr>
<td>CF(_2)Cl(_2)</td>
<td>450–1450</td>
<td>0.004</td>
<td>60</td>
<td>(\nu_{3} + \nu_{7}(^{35}\text{Cl}), 888.49689) (\nu_{3} + \nu_{7}(^{37}\text{Cl}), 883.20389)</td>
<td>D’Amico et al. (2002)</td>
</tr>
<tr>
<td>C(_2)H(_2)</td>
<td>700–1400</td>
<td>0.0043</td>
<td>6–7</td>
<td>(\nu_{5}, 729.163) (\nu_{4} + \nu_{5}, 1328.081) (\nu_{3}, 3281.899) (\nu_{2} + \nu_{4} + \nu_{5}, 3294.839)</td>
<td>Lee et al. (2007)</td>
</tr>
<tr>
<td></td>
<td>3270–3310</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C(_2)H(_4)</td>
<td>700–2400</td>
<td>0.005</td>
<td>50</td>
<td>(\nu_{12}, 1442.44270) (\nu_{7} + \nu_{8}, 1888.97823) (\nu_{6} + \nu_{10}, 2047.77583)</td>
<td>Hurtmans et al. (2001)</td>
</tr>
<tr>
<td>CF(_3)CH(_2)F</td>
<td>Near 1000</td>
<td>0.0035</td>
<td>65</td>
<td>(\nu_{6}, 1104.532116)</td>
<td>Thompson et al. (2003)</td>
</tr>
<tr>
<td>(CH(_3))(_2)O</td>
<td>Near 1000</td>
<td>0.005</td>
<td>70</td>
<td>(\nu_{6}, 933.9066) (\nu_{21}, 1103.951) (\nu_{14}, 2817.385)</td>
<td>Coudert et al. (2002)</td>
</tr>
<tr>
<td></td>
<td>Near 2800</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CH(_3)CHCH(_2)</td>
<td>Near 950</td>
<td>0.005</td>
<td>80</td>
<td>(\nu_{18}, 990.77605) (\nu_{19}, 912.66776)</td>
<td>Lafferty et al. (2006)</td>
</tr>
<tr>
<td>CH(_3)CH(_2)CH(_3)</td>
<td>1300–1500</td>
<td>0.005</td>
<td>80</td>
<td>(\nu_{19}, 1338.965) (\nu_{18}, 1376.850) (\nu_{24}, 1471.874) (\nu_{4}, 1476.710)</td>
<td>Fland et al. (2001)</td>
</tr>
<tr>
<td>C(_6)H(_6), CH(_3)OH</td>
<td>Near 3000</td>
<td>0.015</td>
<td>20–25</td>
<td>CH stretch region, tests of experimental setup</td>
<td>Georges et al. (2002)</td>
</tr>
<tr>
<td>Ni(CO)(_4)</td>
<td>1950–2100</td>
<td>0.006</td>
<td>25</td>
<td>(\nu_{5}, 2061.30937)</td>
<td>Asselin et al. (2008)</td>
</tr>
<tr>
<td>V(CO)(_6)</td>
<td>1960–2020</td>
<td>0.01</td>
<td>13</td>
<td>(\nu_{6}, 1994.48, band profile analysis)</td>
<td>Rey et al. (2001)</td>
</tr>
<tr>
<td>WF(_6)</td>
<td>Near 720</td>
<td>0.0024</td>
<td>50</td>
<td>(\nu_{18}(^{182}\text{W}), 714.53819) (\nu_{18}(^{183}\text{W}), 714.21406) (\nu_{18}(^{184}\text{W}), 713.89544) (\nu_{18}(^{186}\text{W}), 713.26621)</td>
<td>Boudon et al. (2002)</td>
</tr>
<tr>
<td>ReF(_6)</td>
<td>Near 700</td>
<td>0.1, 0.5</td>
<td></td>
<td>(\nu_{3}; very complex spectrum) (combined with diode laser study)</td>
<td>Boudon et al. (2002)</td>
</tr>
<tr>
<td>H(_2)S-HF</td>
<td>Near 3700</td>
<td>0.05, 0.02</td>
<td>20</td>
<td>(\nu_{1}(\text{HF stretch}), 3724.29) (\nu_{4}(\text{DF stretch}), 2734.46)</td>
<td>Asselin et al. (2006)</td>
</tr>
<tr>
<td>D(_2)S-DF</td>
<td>Near 2700</td>
<td>0.02</td>
<td>12</td>
<td>(\nu_{1}(\text{H}^{37}\text{Cl stretch}), 2755.23) (\nu_{1}(\text{H}^{37}\text{Cl stretch}), 2753.15) (\nu_{2}(\text{D}^{37}\text{Cl stretch}), 1993.88) (\nu_{2}(\text{D}^{37}\text{Cl stretch}), 1991.05)</td>
<td>Asselin et al. (2007)</td>
</tr>
<tr>
<td>H(_2)S-HCl</td>
<td>Near 2750</td>
<td>0.02, 0.05</td>
<td></td>
<td>(\nu_{1}(\text{H}^{35}\text{Cl stretch}), 2755.23) (\nu_{1}(\text{H}^{37}\text{Cl stretch}), 2753.15) (\nu_{2}(\text{D}^{35}\text{Cl stretch}), 1993.88) (\nu_{2}(\text{D}^{37}\text{Cl stretch}), 1991.05)</td>
<td>Asselin et al. (2007)</td>
</tr>
<tr>
<td>D(_2)S-DCI</td>
<td>Near 2000</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ar-CO(_2)</td>
<td>2250–2400</td>
<td>0.01, 0.05</td>
<td>6, 9</td>
<td>(\nu_{4}(\text{CO}<em>{2}), 2348.6738) (\nu</em>{5}(\text{CO}_{2}), 2350.7716)</td>
<td>Thiévin et al. (2006)</td>
</tr>
<tr>
<td>CO(_2)-CO(_2)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CFCl(_3)</td>
<td>800–1100</td>
<td></td>
<td></td>
<td>Combined with diode laser spectra</td>
<td>Snels et al. (1995, 2001)</td>
</tr>
</tbody>
</table>
Table 2  High-resolution FTIR investigations of species in supersonic jet expansions. Investigations performed at ETH Zürich including, in part, combinations with diode laser spectroscopy.

<table>
<thead>
<tr>
<th>Species</th>
<th>Spectral range (cm(^{-1}))</th>
<th>Resolution(^{(a)}) (cm(^{-1}))</th>
<th>(p_0)(^{(b)}) (10(^5)Pa)</th>
<th>(p_c)(^{(c)}) (Pa)</th>
<th>(T_{rot}) (K)</th>
<th>Subject</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO(_2)</td>
<td>Near 2350</td>
<td>0.006</td>
<td>1.6</td>
<td>0.5</td>
<td>11</td>
<td>(\nu_2), rot. cooling</td>
<td>Dübal et al. (1984)</td>
</tr>
<tr>
<td>CH(_4)</td>
<td>Near 3030</td>
<td>0.006</td>
<td>2.0</td>
<td>0.5</td>
<td>12</td>
<td>(\nu_3), nuclear spin relaxation</td>
<td></td>
</tr>
<tr>
<td>CF(_3)Cl</td>
<td>1050–1250</td>
<td>0.004</td>
<td>4.2–4.5</td>
<td>0.8</td>
<td>35</td>
<td>(\nu_1, \nu_3 + \nu_6 - \nu_6) and (\nu_4), bands rot. analysis and vibr. cooling</td>
<td>Amrein et al. (1987a)</td>
</tr>
<tr>
<td>CO</td>
<td>2050–2230</td>
<td>0.004</td>
<td>6.75</td>
<td>0.9</td>
<td>6–13</td>
<td>Fundamental band, rot. cooling</td>
<td>Amrein et al. (1988b)</td>
</tr>
<tr>
<td>NO</td>
<td>Near 1880</td>
<td>0.004</td>
<td>6.50</td>
<td>0.6</td>
<td>17</td>
<td>Transitions in (^1\Pi_1^2) and (^2\Pi_3^2) el. states relaxation of electronic degree</td>
<td></td>
</tr>
<tr>
<td>CH(_4)</td>
<td>2960–3140</td>
<td>0.004</td>
<td>7.50; 2.75</td>
<td>0.7; 0.9</td>
<td>11; 36</td>
<td>(\nu_3) band, nuclear spin relaxation</td>
<td></td>
</tr>
<tr>
<td>C(_2)H(_2)</td>
<td>3180–3365</td>
<td>0.01</td>
<td>2.40</td>
<td>1</td>
<td>31</td>
<td>(\nu_1) fundamental and hot bands, rot. and vibr. cooling, evidence for dimers and nuclear spin conservation</td>
<td></td>
</tr>
<tr>
<td>CH(_3)CCH</td>
<td>3230–3435</td>
<td>0.015</td>
<td>2.50</td>
<td>1.5</td>
<td>50</td>
<td>(\nu_1) fundamental and hot bands, rot. and vibr. cooling</td>
<td></td>
</tr>
<tr>
<td>CF(_3)H</td>
<td>Near 1100</td>
<td>0.004</td>
<td>2.75</td>
<td>1</td>
<td>40</td>
<td>(\nu_2, \nu_4, \nu_5 + \nu_6) Coriolis–Fermi–Triad band system in (\nu_1) region Simplification of complex spectra</td>
<td></td>
</tr>
<tr>
<td>CHClF(_2)</td>
<td>1030–1380</td>
<td>0.004</td>
<td>2.3</td>
<td>0.8</td>
<td>50</td>
<td>(\nu_3) and (\nu_6) bands, rot. analysis</td>
<td>Amrein et al. (1988a)</td>
</tr>
<tr>
<td></td>
<td>2870–3175</td>
<td>0.018</td>
<td>2.37</td>
<td>0.75</td>
<td>50</td>
<td>(\nu_1) band, Q-branch features</td>
<td></td>
</tr>
<tr>
<td>N(_2)O</td>
<td>Near 1300</td>
<td>0.004</td>
<td>4.2</td>
<td>7.5</td>
<td>26</td>
<td>(\nu_1, \nu_1 + \nu_2 - \nu_2), rot. and vibr. cooling</td>
<td>Amrein et al. (1989)</td>
</tr>
<tr>
<td>CF(_3)Br</td>
<td>1020–1280</td>
<td>0.004</td>
<td>2.46</td>
<td>7.0</td>
<td>45</td>
<td>(\nu_1), rot. analysis, (\nu_1 + \nu_6 - \nu_6), rot. analysis and vibr. cooling</td>
<td></td>
</tr>
<tr>
<td>CF(_3)I</td>
<td>920–1280</td>
<td>0.004</td>
<td>2.4</td>
<td>50</td>
<td>(\nu_4), rot. analysis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Species</td>
<td>Spectral range (cm(^{-1}))</td>
<td>Resolution(^{(a)}) (cm(^{-1}))</td>
<td>(p_0) ((10^3\text{Pa}))</td>
<td>(p_0) (Pa)</td>
<td>(T_{\text{rot}}) (K)</td>
<td>Subject</td>
<td>References</td>
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<tr>
<td>CF(_3)I</td>
<td>Near 1100</td>
<td>0.005</td>
<td>2.4</td>
<td>45</td>
<td>60</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(\nu_1) band system, vib. rot. analysis including (\nu_1, 2\nu_0^3) and (\nu_3 + 3\nu_0^3) (\nu_1) band system, vib. rot. analysis including (\nu_1, 2\nu_0^3, \nu_3 + 3\nu_0^3) (\nu_1 + 3\nu_0^3) (\nu_1) band system, vib. rot. analysis including in addition (2\nu_0^2) (\nu_1, \nu_7) and (\nu_8), rot. analysis</td>
<td>Bürger et al. (1989)</td>
</tr>
<tr>
<td>CHCl(_2)F</td>
<td>700–1300</td>
<td>0.004</td>
<td>1.10</td>
<td>80</td>
<td></td>
<td>(\nu_3, \nu_7) and (\nu_8), rot. analysis</td>
<td>Snels and Quack (1991)</td>
</tr>
<tr>
<td>NO(_2), N(_2)O(_4)</td>
<td>700–1800</td>
<td>0.0024</td>
<td>4–7(^{(d)})</td>
<td>0.15–1(^{(d)})</td>
<td>20(^{(d)}), 100(^{(d)})</td>
<td>rot. analysis of (\nu_9, \nu_{11}) and Q-branches of resonance doublet (\nu_{12}, \nu_6 + \nu_{12}) of N(_2)O(_4) (\nu_1), rot. analysis</td>
<td>Luckhaus and Quack (1992)</td>
</tr>
<tr>
<td>CCl(_3)F</td>
<td>1050–1120</td>
<td>0.0024</td>
<td>1.1</td>
<td>0.8</td>
<td>80</td>
<td>(\nu_1), rot. analysis</td>
<td>Snels et al. (1995)</td>
</tr>
<tr>
<td>SPF(_3)</td>
<td>Near 1000</td>
<td>0.0024</td>
<td>2.2</td>
<td>0.6</td>
<td>80</td>
<td>(\nu_1), rot. analysis</td>
<td>Bürger et al. (1996)</td>
</tr>
<tr>
<td>CHBrClF</td>
<td>Near 1100</td>
<td>0.0024</td>
<td>0.5</td>
<td>1.0–1.3</td>
<td>100</td>
<td>(\nu_4), rot. analysis</td>
<td>Bauder et al. (1997), Beil et al. (1994)</td>
</tr>
<tr>
<td>WF(_6), ReF(_6)</td>
<td>650–780</td>
<td>0.0024</td>
<td>2</td>
<td>4</td>
<td>50</td>
<td>(\nu_3) of WF(_6), rot. analysis (\nu_3) Q-branch of ReF(_6), profile at lower resolution (\nu_3) Q-branch of ReF(_6), profile at lower resolution</td>
<td>Boudon et al. (2002)</td>
</tr>
<tr>
<td>CF(_2)Cl(_2)</td>
<td>450–1450</td>
<td>0.004</td>
<td>4.0</td>
<td>10</td>
<td>100</td>
<td>(\nu_3 + \nu_7), rot. analysis</td>
<td>D'Amico et al. (2002)</td>
</tr>
<tr>
<td>NH(_2)D, ND(_2)H</td>
<td>1150–1750</td>
<td>0.01</td>
<td>4.0</td>
<td>10</td>
<td>70–100</td>
<td>Bending fundamentals (\nu_{4a}) and (\nu_{4b}) both (s)- and (a)- tunneling components</td>
<td>Snels et al. (2006a)</td>
</tr>
</tbody>
</table>

\(^{(a)}\) Values of 0.004 and 0.0024 cm\(^{-1}\) corresponding to maximum apodized and unapodized resolution, respectively, of the Bomem DA002.

\(^{(b)}\) Stagnation pressure.

\(^{(c)}\) Residual pressure in jet chamber.

\(^{(d)}\) Seeded in Ar.

\(^{(e)}\) Neat substance.
these three examples in the following sections. In the first two cases, FTIR-jet spectra were used in the initial step of the analysis, yielding a first set of assignments and parameters, which later on formed the basis for the analysis of spectra at higher temperatures. For a summary of results specially directed at chiral molecules, we refer to Quack (2011) in this handbook.

6 DISCUSSIONS OF SPECIFIC MOLECULAR EXAMPLES

6.1 Ammonia Isotopomer Spectra

High-resolution spectroscopy of ammonia is of fundamental interest in relation to vibration–rotation tunneling dynamics. Recent interest has been concentrated on full-dimensional potential and electric dipole hypersurfaces as well as quantum wave packet dynamics (see Marquardt and Quack 2011: Global Analytical Potential Energy Surfaces for High-resolution Molecular Spectroscopy and Reaction Dynamics, this handbook). Furthermore, ammonia is an important pollutant in the Earth’s atmosphere and has been detected in planetary atmospheres and interstellar clouds. In this context, the partially deuterated species NH2D and ND2H are particularly noteworthy (see Snels et al. (2006a) and references cited therein for the above-mentioned applications).

The investigation of the bending vibrations of NH2D and ND2H (Snels et al. 2006a) was part of a systematic high-resolution analysis of the infrared spectra of deuterated ammonia isotopomers in the range of the fundamental vibrations (Snels et al. 2000, 2003, 2006b). Figure 10 shows a graphical survey of the fundamentals of various NHnDm species and the correlation between them. While for the symmetric isotopomers NH3 and ND3, the proper symmetry group for the vibration–inversion–rotation problem is a group isomorphous to D3h, the symmetry is reduced for the partially deuterated species NH2D and ND2H to a group isomorphous to C2v. The bending fundamentals, which are degenerate in the case of the symmetric isotopomers, are split into two components. Considering the large values of the rotational constants giving rise to largely extended rotational structures, furthermore the tunneling effects due to the inversion barrier, and the fact that in the case of the mixed H/D isotopomers, we have to deal with isotopic mixtures rather than pure isotopic species, the region of the bending fundamentals (1100–1800 cm⁻¹) of the mixed species appears extremely crowded in spectra taken at room temperature. Rotational cooling in supersonic expansions combined with FTIR spectroscopy was, therefore, chosen as the ideal technique in order to unravel and analyze this complex region. Because of the large amount of substance needed in continuous jet experiments, we did not scan the spectra with the maximal available spectral resolution of our FTIR spectrometer (Bomem DA002, 0.004 cm⁻¹ FWHM, apodized), but reduced it to 0.01 cm⁻¹. Furthermore, we used our multipass jet setup described in Section 3. The measurement of different isotopic mixtures obtained by mixing NH3 and ND3 in ratios 1 : 2 and 2 : 1 then allowed us to identify and assign line structures to the distinct tunneling components of the bending fundamentals of NH2D and ND2H. Peaks due to these isotopomers were sorted out and were used to establish corrected (“cleaned-up”) line spectra from the peak files where only those peaks were retained that could be definitely associated with NH2D and ND2H, respectively. Absorptions due to residual H2O and isotopic species being particularly strong in this spectral region were also eliminated by this procedure. Figure 11 shows a measured and a cleaned-up spectrum in the 1310–1470 cm⁻¹ region in comparison with a calculated spectrum obtained with best-fit parameters for a rotational temperature of 100 K.

The calculations were based on Watson-type Hamiltonians for the vibration–inversion states up to sextic terms,
and an inversion–rotation interaction Hamiltonian given by
\[
\hat{H}_{01} = F (\hat{J}_z \hat{J}_z + \hat{J}_x \hat{J}_x) + F_{\gamma} \hat{J}_z^2 (\hat{J}_x \hat{J}_x + \hat{J}_z \hat{J}_z) + F_K \left[ \hat{J}_z^2 (\hat{J}_x \hat{J}_x + \hat{J}_z \hat{J}_z) + (\hat{J}_x \hat{J}_x + \hat{J}_z \hat{J}_z) \hat{J}_z^2 \right]
\] (18)

The fitting procedure yielded reliable values for about 20 spectroscopic parameters for each fundamental. The term values obtained are listed in Table 3. Increased inversion splittings in the case of \(v_{4a}\) of ND\(_2\)H and a strongly increased splitting and an inverted order of the two inversion levels in the case of \(v_{4b}\) of NH\(_2\)D were traced to Fermi resonance interactions involving the overtone 2\(v_2\) of the inversion vibration.

The term values of the vibrational-tunneling levels allow us to draw conclusions on the mode-selective interplay of fundamental vibrational excitations with the inversion-tunneling dynamics. In Table 4, we give for all fundamentals of the mixed isotopomers a listing of the tunneling times for stereomutation defined by
\[
\tau_{L\rightarrow R} = \frac{1}{2c\Delta\tilde{v}}
\] (19)

The results obtained from the rotational analysis of the bending fundamentals of NH\(_2\)D and ND\(_2\)H completes the data set for the fundamental absorptions in all H/D isotopic species, providing a benchmark for comparisons with calculations using full-dimensional electric dipole and potential energy hypersurfaces for ammonia (Marquardt and Quack 2011: Global Analytical Potential Energy Surfaces for High-resolution Molecular Spectroscopy and Reaction Dynamics, this handbook, Lin et al. 2002, Marquardt et al. 2005). Furthermore, the completed spectroscopic data base is of importance for spectroscopic observations of...
ammonia isotopomers in planetary atmospheres and interstellar clouds.

6.2 CHClF₂

Chlorodifluoromethane (CFC₂₂) attracted our attention as an important system for the investigation of intramolecular redistribution dynamics and related topics (Albert et al. 2004, Quack 1990, Marquardt and Quack 2001, 2011: Global Analytical Potential Energy Surfaces for High-resolution Molecular Spectroscopy and Reaction Dynamics, this handbook, and references cited therein). Rovibrational spectroscopy provides important information for the understanding and modeling of such processes. This molecule represents a particularly interesting system for the investigation of the effects of vibrational and rovibrational interactions since most of its fundamentals are involved in anharmonic and/or rovibrational (Coriolis) resonances. This is illustrated in Figure 12, which shows a diagram of the vibrational levels in the region below 1450 cm⁻¹ grouped according to coupling polyads. The diagram reveals that the only isolated levels in this range are the fundamental ν₅ and to some extent its overtone 2ν₅.

The CF-stretching fundamentals ν₃ and ν₈ are very close in energy and are involved in a strong Coriolis resonance. This band system was considered to be an appropriate case for the investigation of the effects of Coriolis coupling in low-symmetry asymmetric tops (point group Cₛ). Furthermore, this band system lies in the atmospheric window for infrared radiation and since CHClF₂ is known to be a relevant atmospheric trace gas, the analysis of this band system is also of interest in this context.

The rovibrational analysis of this Coriolis resonance band has been subject of a number of investigations. The first analysis was based on FTIR-jet spectra obtained in our laboratory using the Bomem DA002 high-resolution spectrometer operated at the maximum apodized resolution of 0.004 cm⁻¹ and using expansion conditions resulting in a rotational cooling down to about 50 K (Amrein et al. 1988a). Figure 13 shows a part of the ν₃ fundamental, illustrating the effect of rotational cooling and the good agreement between FTIR-jet spectrum and model calculation.

This first high-resolution study formed a valuable basis for the analysis of spectra measured at room temperature under Doppler-limited conditions at ETH-Zürich using the Bruker 125HR Zürich prototype interferometer (maximum unapodized resolution 0.0007 cm⁻¹) (Albert et al. 2004). To unravel and fit the largely increased data set, additional interactions affecting particular regions of the spectrum turned out to be important. The additional perturbers were found to be the combination levels ν₆ + 2ν₉ and 3ν₉.

**Figure 12** Vibrational levels of CH₃⁵ClF₂ below 1450 cm⁻¹ grouped according to polyads. [After Albert et al. 2010.]

**Figure 13** Part of the ν₃ fundamental of CHClF₂. Simulated spectrum in comparison with supersonic-jet FTIR spectrum and room temperature FTIR spectrum. [Reproduced from Amrein et al. 1988a. © Elsevier, 1988.]
A closer inspection of the levels in the neighborhood of the $v_3, v_8$ dyad revealed that these perturbers belong to a Coriolis–Fermi dyad, including the six levels $3v_6, 2v_6 + v_9, v_6 + 2v_9, 3v_9, v_4 + v_6,$ and $v_4 + v_9.$ A related Coriolis–Fermi polyad including the levels $2v_6, v_4, v_6 + v_9,$ and $2v_9$ was applied in a recent rovibrational analysis of the region of the $v_4$ fundamental (Albert et al. 2006). Couplings within the polyads (intrapolad couplings) are defined to arise from the first-order terms of the usual expansion of the vibration–rotation Hamiltonian, which are (Papoušek and Aliev 1982)

$$\hat{H}_{30} = \frac{1}{6} \sum_{l,m,n} k_{lmn} q_l q_m q_n \quad (20)$$

giving rise to Fermi resonances, and

$$\hat{H}_{21} = -2 \sum_{k,l} \left( \frac{\omega_l}{\omega_k} \right)^{1/2} q_k \hat{p}_l \sum_{a} B_{al} \xi_{al} \hat{J}_a \quad (21)$$

giving rise to Coriolis resonances. Interactions between polyads (intrapolad couplings) are due to higher order terms of the expansion. The polyad structure in the low-energy region of this molecule is illustrated in Figure 12.

A revised analysis of the $v_3, v_8$ band system (Albert et al. 2010) was based on this extended polyad model and on a combination of our previous data set (Albert et al. 2004) and of the set of a closely analogous investigation performed at the Monash University, Australia (Thompson et al. 2004). The effective rotational Hamiltonians used in this treatment are diagonal operators of the Watson-type according to the A-reduction scheme and off-diagonal operators for the couplings between the levels $v'$ and $v$ given by

$$\hat{H}_{\text{interaction}}^{v',v} = \langle v' | \hat{H}_{\text{interaction}} | v \rangle \quad (22)$$

where $\hat{H}_{\text{interaction}}$ are coupling operators $\hat{H}_{30}$ and $\hat{H}_{21}.$ Figure 14 illustrates the structure ($J$-blocks) of the Hamiltonian matrix for the eight interacting levels.

In the final calculation, the data sets used comprised 10,402 transition wavenumbers for the isotopomer CH$_{35}$ClF$_2$ and 6,522 for CH$_{37}$ClF$_2.$ The fitting procedure yielded well-determined and physically reliable parameters of the polyad model, the root mean square deviations being $0.334 \times 10^{-3}$ cm$^{-1}$ for the CH$_{35}$ClF$_2$ and $0.373 \times 10^{-3}$ cm$^{-1}$ for the CH$_{37}$ClF$_2$ species (Albert et al. 2010).

### 6.3 CHCl$_2$F

The analysis of rovibrational fine structures of chlorofluorocarbons in the range 700–1400 cm$^{-1}$ is of importance for the understanding of atmospheric infrared absorption. Fluorodichloromethane (CH$_2$ClF, CFC-21) reveals three fundamentals in this range: $v_3$ near 1079 cm$^{-1}$ (CF-stretching), $v_7$ near 1239 cm$^{-1}$ (CH-bending), and $v_8$ near 807 cm$^{-1}$ (Cl-stretching). Although CHCl$_2$F is a molecule of modest complexity, spectra taken at room temperature appear extremely complex due to the presence of three isotopic species CH$_{35}$Cl$_2$F, CH$_{35}$Cl$_{37}$ClF, and CH$_{37}$Cl$_2$F, hot bands, and rotational congestion. The combination of supersonic-jet expansion with FTIR spectroscopy proved to be a particularly effective technique to investigate these bands. The simplification of the spectra due to rotational and vibrational cooling was of central importance in order to get access to a detailed rovibrational analysis. The measurements were performed using the continuous-flow supersonic-jet system as described in Amrein et al. (1988b) in combination with a Bomem DA002 high-resolution Fourier transform spectrometer, operated at the maximum available apodized resolution of 0.004 cm$^{-1}.$ For the rotational analysis of all three fundamental bands, an effective $S$-reduced Watson-type Hamiltonian including terms up to quartic proved adequate. The explicit consideration of anharmonic and rovibrational interaction was not required for an adequate fitting, which yielded well-determined spectroscopic constants for all three fundamentals of the isotopomers: CH$_{35}$Cl$_2$F and CH$_{37}$Cl$_{37}$ClF (Snels and Quack 1991).
6.4 CCl₃F

The CCl₃F molecule is important for the monitoring of the freon concentrations in the upper atmosphere, in relation to their ozone destruction and global-warming potentials. At room temperature, many vibrational levels are populated and give rise to hot-band absorptions. Heavy molecules also have small rotational constants, implying a dense manifold of rotational levels for each vibrational level. Hot bands and small rotational constants produce congested spectra with many overlapping absorption lines. In some cases, the presence of different isotopomers complicates the analysis of room temperature spectra even more. In traditional infrared spectroscopy, the spectrum of a molecule, such as CCl₃F, was considered “nonanalyzable” and its spectra resisted attempts of analysis in spite of its importance as one of the major atmospheric global pollutants. By using supersonic-jet-diode laser spectroscopy, rotational and vibrational temperatures can be reduced significantly, leading to a strong reduction of the number of the absorption lines observed. Furthermore, CCl₃F, because of the presence of three chlorine atoms, has four isotopomers of which the two major ones are evident in the spectrum (C³⁵Cl³⁷F, C³⁵Cl²³⁷F). Analysis of room temperature and cold cell spectra was prohibitive, until diode laser slit-jet spectra were recorded with a resolution of about 0.0006 cm⁻¹. In the case of the parallel band ν₁, the analysis of three isotopomers was done (C³⁵Cl³⁷F, C³⁵Cl²³⁷F and C³⁵Cl³⁷Cl²F), while the ν₄ perpendicular band was analyzed only for the symmetric top molecule C³⁵Cl³⁷F.

6.4.1 Analysis of the ν₁ Band of CCl₃F

Although cold cell spectra of CCl₃F were available in natural abundance and enriched in ³⁵Cl, analysis was prohibitive until supersonic-jet spectra were recorded (Snels et al., 1995, 2001). By using a pulsed slit expansion of 5% CCl₃F in He, a rotational temperature of about 20 K was achieved, and hot bands were efficiently cooled. The low temperature together with the sub-Doppler resolution (about 0.0006 cm⁻¹) produced well-resolved transitions of the three most abundant isotopomers (C³⁵Cl³⁷F, 43.1%; C³⁵Cl²³⁷F, 41.3%; C³⁵Cl³⁷Cl²F, 13.2%) (Figure 15). The first isotopomer is a symmetric top, and the other two are asymmetric tops. The assignment of the supersonic-jet spectra was straightforward and the analysis yielded accurate effective rotational constants. The unusual trend of the excited state rotational constants C in the three isotopomers indicates that the ν₁ band is perturbed by a nearby overtone or combination band (ν₄ + ν₂). This perturber might also be responsible for the absorption around 1085 cm⁻¹.

6.4.2 Analysis of the ν₄ band of CCl₃F

While the ν₁ band appeared as a very congested parallel band, with regular P-, Q-, and R-branches, the room temperature spectrum of the perpendicular band ν₄ has a peculiar structure, with very sharp peaks on the higher wavenumber side of the band origin and a unstructured low wavenumber side (Figure 16).

No plausible explanation was proposed to understand this unusual appearance, until the supersonic-jet spectra were measured and analyzed (Snels et al. 2001). The energy levels for a twofold degenerate vibrational mode of an...
neglecting quartic terms,

\[ E_v/hc = \tilde{v}_0 + B_v J(J + 1) + (C_v - B_v)k^2 \]
\[- 2(C\zeta)_{\nu J} J(J + 1)k^2 + \eta_k^j k^j \]
\[- D^j_J J^2(J + 1)^2 - D^j_K J(J + 1)k^2 \]
\[- D^j_K k^j \]

The expressions for \( \Delta J = +1 \) and \( \Delta K = +1 \) transitions, neglecting quartic terms, are

\[ R_{K}(J) = \tilde{v}_0 + (C_0 + B_0) - 2(C\zeta)_{\nu} - 2B_0 J \]
\[ + 2[C_0 - (C\zeta)_{\nu} - B_0]k \]
\[ + \Delta B(J + 1)k \]
\[ + (\Delta C - \Delta B)(K + 1)^2 \]

For \( \Delta J = -1 \) and \( \Delta K = -1 \) transitions, we have

\[ P_{K}(J) = \tilde{v}_0 + (C_0 + B_0) - 2(C\zeta)_{\nu} - 2B_0 J \]
\[ - 2[C_0 - (C\zeta)_{\nu} - B_0]k \]
\[ + \Delta B(J + 1)k \]
\[ + (\Delta C - \Delta B)(K + 1)^2 \]

If we assume that the Coriolis constant \( \zeta \) in the Hamiltonian is close to 1, we can show that \( R \) and \( P \) transitions form regular clusters for \( K = J - \Delta \), where \( \Delta = 0, 1, 2, 3, \ldots \)

\[ R_{K=J-\Delta}(J) = \tilde{v}_0 + (C_0 + B_0) - 2(C\zeta)_{\nu} \]
\[ - 2\Delta[C_0 - (C\zeta)_{\nu} - B_0] \]
\[ + \Delta C(\Delta - 1)^2 \]
\[ + \Delta B[2 - (\Delta - 1)^2] \]
\[ + 2[C_0 - (C\zeta)_{\nu} + \Delta C J^2 \]
\[ + \text{higher order terms} \]

\[ P_{K=J-\Delta}(J) = \tilde{v}_0 + (C_0 + B_0) - 2(C\zeta)_{\nu} \]
\[ + 2\Delta[C_0 - (C\zeta)_{\nu} - B_0] \]
\[ + \Delta C(\Delta + 1)^2 \]
\[ + \Delta B[2 - (\Delta + 1)^2] \]
\[ - 2[C_0 - (C\zeta)_{\nu} + \Delta C J^2 \]
\[ + \text{higher order terms} \]

Thus, we find clusters of \( R_{K=J-\Delta}(J) \) transitions with a regular spacing of \( 2[C_0 - (C\zeta)_{\nu} - B_0] \) between successive values of \( \Delta \) and a progression of \( (J, J - \Delta) \) lines within each cluster given by \( 2[C_0 - (C\zeta)_{\nu} + \Delta C J^2 \). The \( P_{K=J-\Delta}(J) \) clusters behave in a similar way, but here the progression goes as follows \(-2[C_0 - (C\zeta)_{\nu} + \Delta C J^2 \). For a negative value of \( \Delta C = C_4 - C_0 \), this implies that we have \( J \)-progressions to higher wavenumbers for \( R \) clusters with decreasing intervals between two adjacent transitions, eventually leading to a bandhead (Figure 17), and to lower wavenumbers for the \( P \) clusters, with increasing spacings for higher \( J \).

In the room temperature spectra, the \( R \) clusters are observed as sharp peaks and the \( P \) clusters appear as broad structureless features.

A computer simulation at a tropospheric temperature (200 K) is in very good agreement with a cold cell (203 K) FTIR spectrum as can be seen in Figure 16. In the same figure, a second series of peaks can be observed, which is assigned to the \( C^{35}\text{Cl}_{2^{37}}\text{ClF} \) isotopomer. Although the room temperature and cold cell FTIR spectrum show very similar structures for the two most abundant isotopomers, we should be aware that in one case \( C^{35}\text{Cl}_{2^{37}}\text{ClF} \) we have a symmetric top species, with one degenerate vibrational band and in the other case \( C^{35}\text{Cl}_{2^{37}}\text{ClF} \) an asymmetric top with two almost degenerate fundamental modes. The splitting between the two modes is estimated to be of the order of \( 1\text{cm}^{-1} \) from ab initio calculations. The asymmetry splitting in the rotational structure becomes negligible for higher \( K \) quantum numbers, which explains the observation of sharp band heads in the FTIR spectra. In the diode laser jet spectra, we expect that the asymmetry splitting plays an important role (as can be observed in the \( v_1 \) fundamental) and tends to smear out the spectral structures.

Figure 17  \( \text{CCl}_3\text{F}, \nu_4 \) band, detail showing the band head occurring for the \( R_{J=3}(J) \) manifold; Diode laser spectrum reported by Snels et al. (2001).
6.5 CF$_3$I

An understanding of the detailed rovibrational structure of the $v_1$ band system in the range 1060–1090 cm$^{-1}$ (symmetric CF-stretching fundamental) is of interest for a variety of applications (see, for instance, He et al. (2002) and references cited therein). We emphasize its role as a prototype system for the investigation of infrared multiphoton excitation and laser chemistry for isotope separation using excitation by CO$_2$ laser radiation (Lupo and Quack 1987; He et al. 1995). The rovibrational structure of this band system reveals highly complex features due to couplings with neighboring levels, and in spite of many attempts, a complete understanding of this band complex is still not available. In the first high-resolution study based on room temperature static cell spectra, Fermi resonance interaction with the overtone $2v_3^0$ was taken into consideration (Bürger et al. 1985). A partial analysis was based on $J$-clusters rather than on line-resolved data. A decisive progress in the understanding of this band system was achieved by means of FTIR-jet spectra obtained in our laboratory (Bürger et al. 1989). The strong rotational cooling in the supersonic expansion ($T_{\text{rot}} = 45 \pm 15$ K) allowed for a more detailed analysis and evidenced anharmonic interaction with the combination level $v_3 + 3v_6^{-3}$, in addition to the Fermi resonance with $2v_5^0$, to be important in order to achieve a better modeling of the perturbed rovibrational features. However, distinct regions of the $v_1$ band system still remained unexplained.

A further step forward was possible by the measurement of slit-jet diode laser spectra using a spectrometer setup built in our laboratory (Hollenstein et al. 1994), yielding well-resolved spectra in distinct spectral windows in the region between 1073 and 1083 cm$^{-1}$. Furthermore, we measured new FTIR-jet spectra corresponding to an effective rotational temperature of 60 K. With the help of the new data, the consideration of further levels and interactions turned out to be important for an adequate description. The additional levels are $v_3 + 3v_6^{-3}$ (anharmonically coupled to $v_1$) and $v_3 + 3v_6^{-1}$ (Coriolis interaction with $v_1$, local perturbation at level crossings). Figure 18 shows the supersonic-jet FTIR spectrum in comparison with the calculated spectrum in the central regions of the $v_1$ and $2v_5^0$ systems, and Figure 19 gives an illustrative detail of the slit-jet diode laser spectrum and calculated spectrum.

In the next step undertaken in order to get further insight into this complex system, we obtained new slit-jet diode laser spectra of CF$_3$I (neat and seeded in He) at rotational temperatures of 7, 30, and 70 K (He et al. 2002). In an extended description, anharmonic coupling between the levels $v_3 + 3v_6^{-3}$ and $v_3 + 3v_6^{-1}$ proved relevant and perturbations arising from $2v_5^{\pm 2}$ turned out to be important. The model for the description of the complex $v_1$ band system now includes a polyad with eight excited levels, which are $v_1, 2v_5^0, 2v_5^{\pm 2}, v_3 + 3v_6^{-1}$, and $v_3 + 3v_6^{-3}$. Figure 20 gives a survey of the vibrational and rovibrational interactions and couplings considered in the modeling. Furthermore, Figure 21 shows a diagram of reduced energies in the critical region defined by

$$E_{\text{red}}(J, K', v, l) = E(J, K', v, l) - h\nu \left[ B_0(J + 1) + (A_0 - B_0)K^2 \right]$$

(28)

The quantum number $K'$ is equal to $K$ for $v_1$ and the anharmonically coupled levels, but different from $K$ for the rovibrationally coupled levels, fulfilling the selection rule

$$K' - K - \sum_{i} \Delta l_i = 0 \mod 3$$

(29)

![Figure 18](image-url)  
**Figure 18** Supersonic-jet FTIR spectrum and calculated spectrum showing the central region of the $v_1$ and $2v_5^0$ systems of CF$_3$I. [Reproduced from Hollenstein et al. 1994 by permission.]
High-resolution FTIR and Diode Laser Spectroscopy of Supersonic Jets

Figure 19  Slit-jet diode laser spectrum of CF$_3$I showing the region of the $^0P(14)$ subbranch of $v_1$. [Reproduced from Hollenstein et al. 1994. © Elsevier, 1994.]

Figure 20  Coupling scheme applied in the analysis of the $v_1$ band system of CF$_3$I.

For a given value of $K$, the diagram shows the relative positions of the levels, which may interact by virtue of this selection rule. The diagram includes further levels not belonging to the polyad considered, such as $v_3 + v_4^{-1} + v_6^{-1}$, which might influence the system by anharmonic interaction and might be included in the final step of the analysis, if additional spectral information and information from high-level ab initio calculations might become available in the future.

In distinct regions of the diode laser spectra, effects arising from nuclear quadrupole interaction due to the iodine nucleus ($I = \frac{5}{2}$) are clearly visible and turned out to be important for correctly assigning the spectra. We included quadrupole hyperfine interaction in our model taking into account the first-order terms (diagonal contributions):

$$E_Q = eQq \left(\frac{3 K^2}{J(J+1)} - 1\right) Y(J, I, F)$$  \hspace{1cm} (30)

where

$$Y(J, I, F) = \frac{\frac{3}{2}C(C+1) - I(I+1)J(J+1)}{2(2J-1)(2J+3)I(2I-1)}$$  \hspace{1cm} (31)

and

$$C = F(F+1) - I(I+1) - J(J+1)$$  \hspace{1cm} (32)

with $|J-I| \leq F \leq J+I$ in integer steps, and the usual quantum numbers $J$, $K$ for rotational angular momentum, $I$ for nuclear spin, and $F$ for total angular momentum.

We considered all allowed transitions according to the selection rule $\Delta F = 0, \pm 1$ in the simulation of the spectra. Figure 22 shows the P-subbranch region for low $J$ quantum numbers of $v_1$, revealing rather strong effects arising from nuclear quadrupole interaction. In the case of $^0P(1)$, we could identify all three expected transitions. Figure 23 shows the FTIR-jet spectrum of the $v_1$ band system for an effective rotational temperature of 60 K in comparison with a calculated trace using the eight-level polyad model.

### 6.6 PFCl$_2$

All the fundamentals of the PFCl$_2$ molecules lie below 1000 cm$^{-1}$. The two most abundant isotopomers PF$^{35}$Cl$_2$
(58%) and PF$^{35}$Cl$^{37}$Cl (36%) dominate the spectra. The PF-stretching mode $\nu_1$ has the band center at about $\tilde{\nu}_0 = 836.6$ cm$^{-1}$ for PF$^{35}$Cl$_2$ and at about $\tilde{\nu}_0 = 835.5$ cm$^{-1}$ for PF$^{35}$Cl$^{37}$Cl (Horká et al. 2008). The absorption spectra of PFCI$_2$ have been measured with a diode laser spectrometer in a supersonic jet in the temperature range between 15 and 20 K and with a Bruker ZP 2001 prototype spectrometer (Albert and Quack 2007) with a resolution of 0.001 cm$^{-1}$ at room temperature.

Both isotopomers appear as asymmetric rotors. A Watson A-reduced effective Hamilton operator (Watson 1977) has been used for the analysis:

$$\hat{H}_{\text{red}} = A \hat{J}_0^2 + B \hat{J}_0^2 + C \hat{J}_z^2 - \Delta_J \hat{J}_z^4 - \Delta_{JK} \hat{J}_z^2 \hat{J}_z^2$$

$$- \Delta_K \hat{J}_z^4 - \frac{1}{2} [\delta_J \hat{J}_z^2 + \delta_K \hat{J}_z^2, \hat{J}_z^2 + \hat{J}_z^2]_+$$

$$+ \phi_J \hat{J}_z^6 + \phi_{JK} \hat{J}_z^2 \hat{J}_z^4 + \phi_{JK} \hat{J}_z^2 \hat{J}_z^2 \hat{J}_z^2 + \phi_K \hat{J}_z^6$$

$$+ \frac{1}{2} [\varphi_J \hat{J}_z^4 + \varphi_{JK} \hat{J}_z^2 \hat{J}_z^2 + \varphi_{JK} \hat{J}_z^2 \hat{J}_z^2 \hat{J}_z^2 + \varphi_{JK} \hat{J}_z^2 \hat{J}_z^2 \hat{J}_z^2]_+$$

(33)

where $\hat{J}$ is the angular momentum operator with $\hat{J}^2 = \hat{J}_0^2 + \hat{J}_z^2 + \hat{J}_z^2$. $\hat{J}_\pm = \hat{J}_z \pm i\hat{J}_z$ and $[X,Y]_+$ represents the
The hybrid $c/b$-type band structure of the PF-stretching mode of PF$_{35}$Cl$_2$ has been successfully analyzed. However, this band is strongly perturbed in both isotopomers by a Coriolis-type resonance through the $v_3 + v_5$ band. Similar Coriolis resonances were identified in PH$_2$F and PH$_2$Cl (Beckers et al. 1994). The cold diode laser spectra at 15 K simplified the spectra considerably (Figure 24) and permitted the observation and assignment of many absorption lines of the $v_1$ band of the isotopic chiral molecule PF$_{35}$Cl$_{37}$Cl. The analysis of the FTIR spectrum (room temperature spectra) resulted in a fit of 603 lines of PF$_{35}$Cl$_2$ with an RMS of $0.581 \times 10^{-3}$ cm$^{-1}$, whereas the fit of 65 lines assigned in the jet spectra (20 K) of PF$_{35}$Cl$_{37}$Cl resulted in an RMS of $0.439 \times 10^{-3}$ cm$^{-1}$ (Horká et al. 2008). Figure 25 shows very good agreement between measured and simulated spectrum.
The isotopomer PF$^{35}$Cl$^{37}$Cl is of interest as the first example of an isotopically chiral molecule, where quantitative predictions concerning a fundamentally new isotope effect due to the electroweak interactions have been made (Berger et al. 2005, Quack 2011). The high-resolution spectroscopic analysis provides the starting point for future experimental studies of this effect.

### 6.7 Larger Molecules

With the advent of increasingly sophisticated supersonic-jet techniques, spectroscopic investigation of larger molecules, with a resolution of rotational lines, permitting to obtain structural information has become possible. The group of Richard Saykally reported the first observation of a rotationally resolved spectrum of a nucleotide base, uracil (Viant et al. 1995). They used a tunable diode laser in combination with a heated pulsed slit nozzle to record the absorption spectrum of uracil around 1703 cm$^{-1}$ and analyzed the rotational structure of the fundamental $\nu_6$ mode, which is a predominant out-of-phase mixed carbonyl stretching vibration. In the same laboratory, infrared absorption spectra have been measured by using the cavity ring down technique, of jet-cooled PAHs, such as naphthalene, anthracene, phenanthracene, pyrene, and perylene with a resolution of 0.04 cm$^{-1}$ (Huneycutt et al. 2004, Schlemmer et al. 1994).

Supersonic rotationally resolved jet spectra (0.001 cm$^{-1}$) of large chlorofluorocarbons have been measured in a supersonic jet by Snels et al. (1995, 2001, 2003), D’Amico and Snels (2002, 2003). They recorded spectra of CFC142b (1,1,1-difluorochloroethane) and analyzed the $\nu_7$, $\nu_6$, and $\nu_14$ bands of both isotopomers. The same authors reported the analysis of the $\nu_{14}$ band of HFC 134a (1,1,1,2-tetrafluoroethane) (Snels and D’Amico 2003). The $\nu_6$ band of the CFC142b exhibits a remarkable splitting of all rotation–vibrational transitions due to a torsional Coriolis interaction with a highly excited torsional overtone band (Di Lauro et al. 2009). Duan and Luckhaus (2004) investigated malonaldehyde and observed the two tunnel components of the $\nu_6$ band. For heavy transition metal hexafluorides such as WF$_6$ and ReF$_6$, the room temperature spectra are heavily congested by hot bands, owing to the presence of hot bands originating from low-energy vibrational modes. By cooling WF$_6$ in a WF$_6$/He expansion, the vibrational ground-state population could be increased from 1.3% at room temperature up to about 95% at 50 K vibrational temperature. A full rotational analysis of the $\nu_3$ band has been performed for the four most abundant isotopic species. ReF$_6$ is an open-shell molecule and a most interesting Jahn–Teller system with a highly degenerate electronic ground state. It is the first time that line-resolved supersonic-jet spectra for a system of such a complexity were obtained (Boudon et al. 2002). The splitting increases the complexity of the infrared spectrum enormously and distributes the absorption strength over many transitions. For this reason, FTIR-jet spectra could not be recorded at the highest instrumental resolution. A tunable diode laser, however, combined with a supersonic slit jet and a multiple pass optical setup, permitted the recording of well-resolved ReF$_6$ spectra. The analysis of these spectra is still not complete.
7 STUDIES OF NUCLEAR SPIN SYMMETRY CONSERVATION IN JET EXPANSIONS

The principles of nuclear spin symmetry and parity conservation are among the most fundamental aspects of spectroscopy and reaction dynamics (Quack 1977, 1983, 1985) (see also Oka (2011): Orders of Magnitude and Symmetry in Molecular Spectroscopy and Quack (2011): Fundamental Symmetries and Symmetry Violations from High-resolution Spectroscopy, this handbook). Polyatomic molecules possessing identical nuclei with nonzero spin in symmetrically equivalent positions exist in several forms of nuclear spin isomers. This led to the separation of para- and ortho-H\textsubscript{2} already in 1929 through cooling in the presence of a magnetic catalyst (Bonhoeffer and Harteeck 1929). Nuclear spin isomers play an important role in fundamental research and the theory can be used in different fields such as astronomy and astrophysics to study the abundance ratio of nuclear spin isotopomers in the interstellar space, kinetics of the chemical reactions in the gas phase, NMR spectroscopy, etc. At the present time, several techniques exist for the separation of spin isomers reviewed by Chapovsky and Hermans (1999), see also Oka (2011): Orders of Magnitude and Symmetry in Molecular Spectroscopy and Quack (2011): Fundamental Symmetries and Symmetry Violations from High-resolution Spectroscopy, this handbook. One method to study nuclear spin isomers is the utilization of supersonic-jet expansion with high-resolution spectroscopy. To study this phenomenon, exact line intensities and rotational temperatures have to be derived. To obtain information about the spin symmetry conservation or spin symmetry interconversion, the line intensities have to be calculated under different hypotheses and compared with those from measurement. The intensity of the rotational lines of a spherical top molecules is given by

\[ G(J, \Gamma, i) = (\text{const}) p(J, \Gamma, i) A(J) \]  
(34)

with the rotational factor

\[ A(J) = \frac{2J' + 1}{2J + 1} \]  
(35)

where \( J \) is the angular momentum quantum number for the ground state and \( J' \) for the excited state. \( \Gamma \) denotes the nuclear spin isomer. The population of the states \( p(J, \Gamma, i) \) can be considered under two extreme situations with complete equilibration among all states (relaxed distribution) or complete nuclear spin symmetry conservation (conserved) (see Amrein et al. 1988b). The relaxed distribution is given by

\[ p_r(J, \Gamma, i) = g_1(\Gamma)(2J + 1)\exp[-E(J)/kT_{\text{rot}}]/Q_{\text{rot}}^r \]  
(36)

where \( g_1(\Gamma) \) is the nuclear spin statistical weight and \( Q_{\text{rot}}^r \) is the relaxed rotational partition function including nuclear spin at a rotational temperature \( T_{\text{rot}} \)

\[ Q_{\text{rot}}^r = \sum_{\Gamma} g_1(\Gamma)Q_{\text{rot}}(\Gamma)\exp[-E_0(\Gamma)/kT_{\text{rot}}] \]  
(37)

with

\[ Q_{\text{rot}}(\Gamma) = \sum_J N(J, \Gamma)(2J + 1) \times \exp(-[E(J) - E_0(\Gamma)]/kT_{\text{rot}}) \]  
(38)

where \( E_0(\Gamma) \) is the lowest level of nuclear spin isomer \( \Gamma \) and \( E(J) \) is the energy for a given \( J \) in the vibrational ground state. \( N(J, \Gamma) \) is the number of levels of isomer \( \Gamma \) for \( J \).

For a complete nuclear spin symmetry conservation, one gets

\[ p_c(J, \Gamma, i) = x(\Gamma)(2J + 1) \times \exp(-[E(J) - E_0(\Gamma)]/kT_{\text{rot}})/Q_{\text{rot}}(\Gamma) \]  
(39)

where \( x(\Gamma) \) is the mole fraction of the nuclear spin isomer \( \Gamma \) at the temperature \( T \) before the expansion, at which the isotopomers are assumed to be equilibrated. The relative line strength of a single rovibrational line is given by

\[ G(J) = \frac{1}{cL} \int_{\text{line}} \ln(I_0/I)\tilde{\nu}^{-1}d\tilde{\nu} \]  
(40)

where \( c \) is the concentration and \( L \) is the absorption pathlength. The interconversion of nuclear spin isomers for \( \text{CH}_3 \) has been investigated already in supersonic-jet expansions with FTIR spectroscopy by the Zürich group (Dübal et al. 1984, Amrein et al. 1988b). The measurements showed that the nuclear spin is conserved during the expansion (Figures 26 and 27). We have extended these investigations to partially deuterated methane isotopomers (Horká-Zelenková et al. 2010). To investigate the dynamical processes during the expansion, the intensities of the measured absorption lines have to be determined with high accuracy. This accuracy can only be achieved if the considered rotational lines are measured in a single scan, ensuring identical experimental conditions for all the lines. As the scanning range of the diode laser is 1–2 cm\(^{-1}\) at most and as for molecules of the type CX\(_y\)Y, the intensities of rotational lines with different \( K \) quantum numbers have to be compared, the experiments are preferentially done on the P-
or R-branch of a parallel band ($A_1 \rightarrow A_1$ transition) with $\Delta K = 0$, because for a given quantum number $J$, the lines for the different $K$ quantum numbers are close enough to be covered in a single scan. Additional complications may arise from the fact that the complete spectral range is not reachable with a single laser diode.

Different criteria have to be considered for a proper choice of the measured rotational lines. Both nuclear spin isomers must have an absorption line in the considered spectral range and at least one of the nuclear spin isomers must have more than one single line to allow for the determination of a rotational temperature. All lines with $J = 2$ or larger fulfill these criteria. On the other hand, to decide whether the nuclear spin is conserved during the expansion phase of the molecular beam or not, very low rotational temperatures have to be reached (below 15 K for CHD$_3$ (Horká-Zelenková et al. 2010)). For these temperatures, the rotational lines with $J = 3$ or larger are only weakly populated and the accuracy for the determined line intensities is poor. These two restrictions make the $P(3)$- or $R(2)$-lines of the CH-stretching vibration ($\nu_1$) an ideal choice to study spin symmetry relaxation of CHD$_3$ in the expansion zone of a molecular beam (Horká-Zelenková et al. 2010).

8 STUDIES OF CLUSTERS

By using the cooling in a supersonic-jet expansion, clusters can be produced in a more or less controlled way, by varying source pressure, nozzle temperature, mixing ratio, and carrier gas.

The spectroscopic investigation of molecular clusters is important for the study of intramolecular dynamics.
and tunneling phenomena, and it provides fundamental information on the nature and strength of the weak van der Waals forces and the much stronger hydrogen bonds in clusters, and allows to determine the structure of the complexes in vibrationally excited states.

An incomplete list of scientific publications regarding spectra of clusters resolving rotational levels is made available by Novick (2008).

It is intriguing to build clusters from isolated molecules and atoms. The structure of stable isomers sometimes can be calculated from current ab initio theory with a good precision (Marquardt and Quack 2011: Global Analytical Potential Energy Surfaces for High-resolution Molecular Spectroscopy and Reaction Dynamics, this handbook, Quack and Suhm 1997, 1998, Klopper et al. 1996, 1998 and Maerker et al. 1997), depending on the cluster size. Infrared spectra of CH4–Rg and SiH4–Rg can also be calculated theoretically and these provide a starting point for exploring infrared spectra, mostly recorded for molecular beam expansions, which contain different clusters. Sometimes several isomers are found experimentally, also depending on the production method.

8.1 Van der Waals Clusters

Some van der Waals complexes studied consist of diatomic and triatomic molecules bound to noble gas atoms, such as CO–Rg (Brookes and McKellar 1998, McKellar et al. 1999), N2O–Rg (Herrebout et al. 1998), CO2–Rg (McKellar 2006), and OCS–Rg (Hayman et al. 1989), where Rg stands for He, Ne, Ar, Kr, and Xe. All these clusters are weakly bound van der Waals complexes, with a binding energy of a few hundreds of cm⁻¹ at most. OCS–Rg and N2O–Rg were shown to be T-shape complexes, with the rare gas atom placed versus the oxygen atom. The long-range attractive forces (inductive and dispersive) scale with the polarizability of the rare gas atoms (Ne < Ar < Kr < Xe). In addition, the CO2–Rg and CO–Rg complexes behave in a similar way. Molecular complexes involving CO might occur in interstellar space due to the fact that CO is the most abundant polar molecule in outer space.

Several complexes of tetrahedral molecules with rare gases were studied, such as CH4–Rg and SiH4–Rg. The interesting point in studying these light spherical-top rare-gas complexes is the existence of internal rotation effects. CH4–He, CH4–Ar, and CH4–Kr were subject of several studies by Pak et al. (1998) and Wangler et al. (2001). Infrared spectra of SiH4–Ne and SiH4–Ar were explored in detail by Howard and coworkers (Randall et al. 1994a,b, Brookes et al. 1996, 1997). They also developed a model for the energy levels of rare gas spherical top complexes, which was successful in describing the hindered rotational structure of SiH4–Ar (Randall et al. 1994a). Analysis of SiH4–Ar is simpler since it is a much more strongly anisotropic system than CH4–Ar, and shows a behavior between free rotor and rigid molecule limit. SiH4–Ne has a smaller anisotropy than SiH4–Ar, resulting in a new angular momentum coupling scheme and from the derived rotational constant, it appears that the silane monomer is closer to the free internal rotational limit. The Coriolis model developed by Brookes et al. (1996) was also applied to CH4–Rg complexes, which are also relatively close to free internal rotation behavior.

In addition, high-resolution spectra of complexes involving heavy spherical top molecules such as SiF4 and SF6 were reported. From the analysis of the infrared spectra recorded for SF6–Rg complexes (Rg = Ne, Ar, Kr, Xe) in liquid helium droplets (Hartmann et al. 1996), it appears that the SF6–Rg complex is embedded in a layer of 10–12 He atoms, which contribute to the very small rotational constants of this embedded cluster. Urban et al. (1995) measured fully resolved rovibrational spectra of SiF4–Rg. Spectroscopic constants for the complexes SiF4–Ar and SiF4–Kr were obtained.

All complexes mentioned so far involved small molecules and rare gas atoms. The pioneering work on clusters of small molecules with noble gas atoms was gradually extended to dimers and trimers of larger molecules. Clusters involving molecules with tetrahedral and spherical symmetry, such as SF6, CH4, SiF4, have been studied by Urban and Takami (1995a, b). In these complexes, the attractive forces are due to induced dipole moments in each of the molecules. For instance in the free jet infrared spectra of SF6, SiF4, and CH4 dimers, two absorption bands have been observed and explained in terms of a resonant dipole–dipole interaction between the two monomers. In the SF6 dimer, when the ν3 threefold degenerate vibration (in the monomer) is excited, two distinct bands appear in the dimer spectrum, one is a parallel band and the other a perpendicular band, as predicted by the dipole–dipole model (Snels and Fantoni 1986).

In the SiF4 dimer spectrum, two different bands could be observed, one parallel transition and one perpendicular band, which had been observed many years before by Snels and Fantoni (1986) in low-resolution infrared predissociation measurements. Urban and Takami (1995a,b) propose three different structures for both dimers, with C3v, D2d, and C2h symmetry. They argue that the dimer with C3v symmetry should exhibit a first-order Coriolis interaction in the perpendicular band and they expect a strong–weak–weak intensity alternation in the same band, due to the threefold symmetry along the a-axis. In addition, the D2d symmetry should give rise to

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a first-order Coriolis interaction, but would show intensity alternation for even and odd \( K \), due to \( C_2 \) symmetry along the \( a \)-axis. The dimer with \( C_{2h} \) symmetry would have a degenerate perpendicular band without a first-order Coriolis interaction. The dimer spectrum of \( \text{SiF}_4 \) shows neither a Coriolis interaction nor any intensity alternation, which indicates a structure with \( C_{2h} \) symmetry. The case of the \( \text{SF}_6 \) dimer is more complicated; the perpendicular band exhibits a clear first-order Coriolis interaction. The dimer spectrum of \( \text{SiF}_4 \) shows neither a Coriolis interaction nor any intensity alternation, which excludes the \( C_{2h} \) symmetry. They further make the hypothesis of internal rotation in the \( \text{SF}_6 \) dimer.

From these relatively simple molecular clusters, the interest moved toward larger clusters, involving more noble gas atoms, up to nano droplets, which form a particular environment where superfluidity was observed. Very interesting studies of van der Waals clusters in liquid helium droplets have been performed by Scheele et al. (2005), Hartmann et al. (1996), Xu et al. (2003), McKellar (2004, 2008), Tang and McKellar (2004), Lehnig and Jäger (2006). Their results demonstrate that liquid He droplets (usually consisting of several thousands of He atoms) provide a very cold environment where molecules can rotate almost freely and that the rotational structure of the molecules trapped in the droplets produces information about the interaction between molecules and the droplet.

The \( \text{CO}_2 \) dimer was measured by the group of Brian Howard, by using a pulsed jet diode laser spectrometer (Walsh et al. 1987). The structure was found to be a parallel staggered configuration. Two distinct isomers of the \( \text{CO}_2 \) trimer have been identified by means of infrared spectroscopy. The first, which is a symmetric top with a cyclic planar structure and \( C_{3h} \) symmetry, was originally observed in the \( \text{CO}_2 \) monomer \( \nu_2 + \nu_3 \) combination band region by Fraser et al. (1987) and later in the monomer \( \nu_3 \) fundamental region by Weida et al. (1995). The second isomer, which is an asymmetric top with a \( C_2 \) symmetry and a barrel-shaped structure, was observed in the monomer \( \nu_3 \) fundamental region also by Weida and Nesbitt (1996). McKellar and coworkers (Dehghany et al. 2008) observed two combination bands \( \nu_3 + \nu_{\text{torsion}} \) for the cyclic trimer. Possible larger \( \text{CO}_2 \) cluster spectra are under study in the McKellar group.

The structure of the \( \text{N}_2\text{O} \) dimer, determined from jet absorption spectra in the \( \nu_1 \) (Ohshima et al. 1988), \( \nu_5 \) (Qian et al. 1997), and \( \nu_1 + \nu_3 \) (Huang and Miller 1988) regions, proved to be a nonpolar slipped parallel structure. Later, Miller and Pedersen also found a polar isomer in the \( \nu_1 \) band region, as well as the first trimer of \( \text{N}_2\text{O} \) (Miller and Pedersen 1998). This trimer can be thought of as a slightly distorted nonpolar dimer with a third \( \text{N}_2\text{O} \) molecule above the dimer plane, similar to the noncyclic \( \text{CO}_2 \) trimer. The same authors also measured the \( \text{N}_2\text{O} \) tetramer (Miller and Pedersen 1997) in the \( \nu_1 \) and \( \nu_1 + \nu_3 \) region. This tetramer can be considered as two dimer units stacked one on top of the other. Surprisingly, a second tetramer band has been observed in the \( \nu_1 \) region, this time a perpendicular band. Other \( \text{N}_2\text{O} \) cluster bands have also been observed but not assigned to a specific cluster up to now.

In all the clusters mentioned above, the van der Waals bonds are rather weak and can be explained in terms of dipole and quadrupole moments of the molecules involved.

### 8.2 Hydrogen Bonds

The hydrogen bond plays an important role as a significant intermolecular interaction, ranging from small molecules such as water and hydrogen fluoride to the complex biological systems as nucleic acids and proteins. The hydrogen bond results in part from a dipole–dipole force between a hydrogen atom and an electronegative atom, such as nitrogen, oxygen, or fluorine. The dynamics of hydrogen bond formation and breaking is important for the understanding of biological processes and for evaporation and condensation phenomena in hydrogen bond liquids such as water and hydrogen fluoride. The binding energies for neutral hydrogen bonds are in a range of up to a few thousand cm\(^{-1}\) and thus are one order of magnitude stronger than the bonds in van der Waals clusters. One of the simplest hydrogen bond systems can be found in the HF dimer. The vibrational spectrum shows some interesting features, such as tunneling splitting, rotational saturation, and rovibrational perturbations, of this molecule due to its nonrigidity and near linearity. The rearrangement of its hydrogen bond is governed by the coupled low-frequency modes and rotation about the F–F axis. The HF dimer has been studied by various spectroscopic methods and an accurate full-dimensional potential has been derived by ab initio theory (Quack and Suhm 1990b, 1991, Klopper et al. 1996, 1998). Our group at ETH Zürich used FTIR and supersonic-jet techniques for the measurement of the spectra, resulting in the analysis of the fundamental out-of-plane \( \text{F}–\text{H}–\text{F} \) bending vibration \( \nu_6 \), \( K = 1 \) (von Puttkamer and Quack 1987), \( K = 2 \) (von Puttkamer et al. 1989) up to \( K = 4 \) (von Puttkamer et al. 1988) and in-plane bending \( \nu_5 \) (Quack and Suhm 1990a). We also studied HF-stretching overtones in (HF)\(_2\) and HFDF around 7500–7800 cm\(^{-1}\) using both diode laser direct absorption and cavity ring down spectroscopy (He et al. 2007, Hippler et al. 2007). From the analysis of these rotationally resolved spectra, hydrogen predissociation lifetimes and tunneling switching times were obtained. Figure 28(a) and (b) shows as an example the very different appearance of the \( N_1 = 2 \) overtone band of (HF)\(_2\), which has a very small vibrational predissociation rate and line broadening and the 2\(_1\) polyad band where the very considerable line broadening is easily visible. The predissociation
rates have been analyzed quantitatively and one finds that full quantum dynamics describes the process essentially correctly, whereas classical dynamics is quite unable to provide a quantitatively satisfactory description (Manca et al. 2008). From theoretical studies, (HF)$_3$ is calculated to be cyclic and the F–F distance is shorter than in (HF)$_2$ (Suhr et al. 1993, Quack et al. 1993b, 2001). The other hydrogen halides and their isotopomers, such as (HCl)$_3$ (Färnik and Nesbitt 2004), (DCI)$_2$ (Schuder et al. 1993), (HBr)$_2$ (Castillo-Chara et al. 2004), and (HI)$_2$ (McIntosh et al. 2000), have also been studied in jet expansion with diode lasers.

Numerous studies were also done on hydrogen halides bonded via hydrogen bonds to other functional groups such as HCN–HF (Bender et al. 1987) and OC–HCl, OC–HBr, N$_2$–HBr dimer (Wang et al. 2004), etc. One can point out that the dimer (HF)$_2$ has been the prototype for high-resolution spectroscopic studies of hydrogen-bonded complexes, starting with the work of Dyke et al. (1972). We have mentioned some highlights of the spectroscopy of these complexes above and draw attention to the reviews of Quack and Suhr (1997, 1998). Detailed spectroscopic identification of large clusters of HF have been possible up to (HF)$_5$ and perhaps (HF)$_6$ (Quack et al. 1993a, Luckhaus et al. 1995). (HF)$_n$ and (DF)$_n$ as well as mixed isotopomer clusters have also been at the origin of the very first infrared spectroscopic identification of hydrogen-bonded nanoclusters in supersonic jets (Quack et al. 1997). In the mean time, this general field has had enormous development of obvious importance for atmospheric physics and astrophysics (Firanescu et al. 2006).

H$_2$O is another important molecule forming hydrogen-bonded clusters. The hydrogen bond in water involves electron pairs on the oxygen atom interacting with the hydrogen atom of other molecules. Thus, the water molecule can create hydrogen bonds with up to four other molecules. This bonding is responsible for the high boiling point, melting point, and viscosity of water in liquid phase in comparison with other similar substances. Water clusters and water-containing complexes play an important role in atmospheric processes and acid rain formation (Kolb et al. 1994) and, of course, atmospheric water vapor contributes to the greenhouse effect, in particular, due to absorption in the mid- and near-infrared range (Cess et al. 1995, Li et al. 1995, Vaida et al. 2001).

Water-containing complexes frequently display a tunneling splitting arising from proton exchange. Some of these complexes have also been studied via jet-diode laser spectroscopy as N$_2$O–H$_2$O (Ginnmiller and Havenith 2002), Ar–H$_2$O (Weida and Nesbitt 1997), CO–H$_2$O (Brookes and McKellar 1998), H$_2$O–CO$_2$, and H$_2$O–HCCH (Block et al. 1992).

Double hydrogen bonds play an important role in DNA base pairs; moreover, multiple proton transfer in hydrogen-bonded systems is one of the most fundamental processes in chemistry and biology. The formic acid dimer is one of the smallest organic complexes serving as a prototype for multiple proton transfer. The antisymmetric C–O vibrational band of this complex was studied by Ortlieb and Havenith (2007). It shows slow tunneling motion in comparison with the overall rotation and thus confirms the existence of deep local minima in the $C_{2h}$ structure. This and more recent
work are reviewed by Havenith and Birer in this handbook (Havenith and Birer 2011: High-resolution IR-laser Jet Spectroscopy of Formic Acid Dimer, this handbook).

8.3 Carbon Clusters

Laboratory spectroscopy of atomic and molecular carbon in its various forms has always been of great interest and importance for a number of reasons. The discovery of the presence of carbon-chain molecules in interstellar space intensified laboratory studies concerning the production and spectroscopic observation of pure carbon clusters.

Carbon atoms form stable covalent carbon–carbon bonds. The single, double, and triple carbon–carbon bonds permit the formation of a large variety of pure carbon molecules, ranging from linear chains to branched chains, cyclic molecules, as well as three-dimensional structures such as C_{60}. The spectroscopic investigations have been supported by high-level ab initio calculations, which predict stable linear, cyclic, and three-dimensional structures. The linear chains appear to be the most stable form for a relatively small number of carbon atoms (up to 10), larger clusters are supposed to form rings (up to 30 atoms), and still larger carbon molecules closed three-dimensional cages.

Matsumura et al. (1988) observed the rotationally resolved infrared spectrum of C_{3}. In rapid succession, other small carbon chains were produced and spectroscopically studied in the laboratory, such as C_{4} (Heath and Saykally 1991b), C_{5} (Heath et al. 1989), C_{6} (Hwang et al. 1993), C_{7} (Heath et al. 1990, Heath and Saykally 1991a), and C_{9} (Heath and Saykally 1990) using a laser ablation carbon cluster source in combination with a sensitive infrared tunable diode laser spectrometer. During the same period, C_{3} and linear C_{5} were detected in the envelope of the carbon star IRC+10216 by Hinkle et al. (1989). The largest carbon chain observed in laboratory experiments at high resolution was reported by Giesen (1989). The spectroscopic investigations have been supported by high-level ab initio calculations, which predict stable linear, cyclic, and three-dimensional structures. The linear chains appear to be the most stable form for a relatively small number of carbon atoms (up to 10), larger clusters are supposed to form rings (up to 30 atoms), and still larger carbon molecules closed three-dimensional cages.

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Kroto et al. (1985) discovered an icosahedral carbon cluster, consisting of 60 carbon atoms, which they called Buckminsterfullerene. In 1996 Curl, Kroto, and Smalley were awarded the Nobel prize in Chemistry for their discovery (Kroto 1997). C_{60} has 174 vibrational degrees of freedom, but only four F_{1u} IR-active modes (1432, 1183, 577, 528 cm^{-1}). A gas-phase IR spectrum has been observed at a temperature of 1065 K, but no rotational structure was resolved (Frum et al. 1991). Other stable carbon cage clusters (fullerenes) were discovered (C_{n}, n = 24, 28, 32, 50, 60, and 70, (Kroto 1987)). Continuing their work during 1985–1990, Curl, Kroto, and Smalley obtained further evidence for the proposed C_{60} structure. Among other things, they succeeded in producing and identifying carbon clusters that enclosed one or more metal atoms. In 1990, the group led by W. Krätschmer and D.R. Huffman (Ajie et al. 1990) produced large quantities of C_{60} using an arc discharge between two graphite rods in a helium atmosphere and extracting the carbon condensate so formed using an organic solvent. They obtained a mixture of C_{60} and C_{70}, and they also determined their structures. This confirmed the correctness of the C_{60} hypothesis. The way was thus open for studying the chemical properties of C_{60} and other carbon clusters such as C_{70}, C_{76}, C_{78}, and C_{84}. New substances were produced from these compounds, with new and unexpected properties. A new branch of “fullerene” chemistry developed, with diverse areas of interest such as astrochemistry, superconductivity, and material chemistry and physics.

9 RADICALS AND IONS

Free radicals, molecular ions, and ionic complexes are important reactive intermediates in many chemical environments, ranging from combustion processes and plasmas to atmospheric chemistry and interstellar space (Linnartz et al. 2000).

Pioneering spectroscopic work on some important molecular ions was started in the early 1980s, by using hollow cathode discharge tubes. The first infrared spectrum of the H_{2}^{+} ion was reported by Oka (1980). The analysis of the v_{2} band was accomplished by J.K.G. Watson (Oka 1980). The v_{3} band of the hydronium ion, H_{3}O^{+}, was analyzed by Saykally’s group (Begemann et al. 1983). Saykally also proposed the velocity modulation technique, which allows an efficient discrimination of spectral features due to ions with respect to neutrals (Gudeman and Saykally 1984). The v_{2} bending-inversion vibration of H_{2}O^{+} was measured and analyzed by Liu and Oka (1985), who also determined the inversion splitting in the ground state and the v_{2} = 1 state. In rapid succession, several other molecular ions were studied in discharge flow tubes, such as OH^{-} (Liu and Oka 1986), HNN^{+}, DNN^{+} (Owrutsky et al. 1986), H_{2}D^{+} (Amano and Watson 1984), and many others. We draw attention to a review paper on the early days of infrared absorption spectroscopy of molecular ions by Sears (1987).

Although discharge flow tubes have produced a wealth of spectroscopic data of molecular ions (Sears 1987), the discharge environment inevitably produces high translational, vibrational, and rotational temperatures. This can
be troublesome in the case of heavier and weakly bound instable species, because spectral congestion and population of many hot bands render the observed spectra extremely complex. In addition, the linewidths of single transitions are highly Doppler broadened due to the high translational temperatures. Both effects reduce the overall detection efficiency of the species.

The often complicated spectra of large free radicals and ions can be significantly simplified by cooling them in a supersonic expansion. A supersonic expansion provides an almost collision-free environment, enhancing the lifetime of free radicals and making their spectroscopic investigation easier. It also allows the production of weakly bound molecular ionic complexes.

Several methods for generating a sufficiently high number density of radicals and ions in a supersonic jet, allowing for infrared absorption spectroscopy have been developed. The most popular ones use an electrical discharge at the beginning of the expansion, in a way that a relatively high density of radicals and ions can be created in the high-pressure region, and successively cooled in the supersonic expansion. As an alternative to electric discharges, the methods of pulsed pyrolysis, laser ablation, and photolysis with a counterpropagating infrared laser pulse generated by an OPO. Several ionic clusters have been investigated by using this method, ranging from open-shell complexes such as He–HNH\(^+\) (Dopfer \textit{et al.} 1999a) and Ne–HNH\(^+\) (Roth \textit{et al.} 2000) to larger clusters such as Ne\(_6\)HN\(_2^+\) (\(N = 1–5\)) (Nizkorodov \textit{et al.} 1998), C\(_6\)H\(_6^+\), C\(_6\)H\(_6^+\)–N\(_2\), and C\(_6\)H\(_6^+\)–(CH\(_3\))\(_3\)– (Dopfer \textit{et al.} 1999b), C\(_3\)H\(_7^+\)–N\(_2\) (Dopfer \textit{et al.} 2002). They have also adapted the cw-CRD scheme of Hippler and Quack (1999, 2002) to the study of molecular ions (see also Birza \textit{et al.} (2002), Guennoun and Maier 2011: \textit{Electronic Spectroscopy of Transient Molecules}, this handbook for review). Spectroscopic studies of individual conformations of peptides with more than a dozen amino acids in a gas phase was investigated by the group of Rizzo \textit{et al.} (2009). They combine electrospray

The slit supersonic discharge source has been used in several laboratories to produce radicals and molecular ions (Hilpert \textit{et al.} 1994, Fukushima \textit{et al.} 1994). An alternative method to produce low-energy cooled plasmas uses electron impact ionization in a slit expansion. This technique allowed to measure the infrared spectra of weakly bound ionic complexes (Verdes \textit{et al.} 1999, Linnartz \textit{et al.} 1998, 2000).

Another method uses pulse pyrolysis to obtain free radicals and uses a Thermocoax wire to resistively heat the nozzle exit (Liu \textit{et al.} 1998, Tanaka \textit{et al.} 1999). Ions and radicals can be also prepared via laser vaporization (Heath and Saykally 1990, 1991a).

John Maier and his coworkers developed a molecular beam apparatus to study the vibrational infrared photodisassociation of mass-selected ionic complexes. The cluster ions were generated in a pulsed supersonic expansion crossed by two electron beams. After mass selection of the species of interest in a quadrupole mass spectrometer, the selected ions are injected in an octupole ion guide, where they interact with a counterpropagating infrared laser pulse generated by an OPO. Several ionic clusters have been investigated by using this method, ranging from open-shell complexes such as He–HNH\(^+\) (Dopfer \textit{et al.} 1999a) and Ne–HNH\(^+\) (Roth \textit{et al.} 2000) to larger clusters such as Ne\(_6\)HN\(_2^+\) (\(N = 1–5\)) (Nizkorodov \textit{et al.} 1998), C\(_6\)H\(_6^+\), C\(_6\)H\(_6^+\)–N\(_2\), and C\(_6\)H\(_6^+\)–(CH\(_3\))\(_3\)– (Dopfer \textit{et al.} 1999b), C\(_3\)H\(_7^+\)–N\(_2\) (Dopfer \textit{et al.} 2002). They have also adapted the cw-CRD scheme of Hippler and Quack (1999, 2002) to the study of molecular ions (see also Birza \textit{et al.} (2002), Guennoun and Maier 2011: \textit{Electronic Spectroscopy of Transient Molecules}, this handbook for review). Spectroscopic studies of individual conformations of peptides with more than a dozen amino acids in a gas phase was investigated by the group of Rizzo \textit{et al.} (2009). They combine electrospray

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**Table 5** Spectroscopic tools developed and used in the Zürieh molecular kinetics and spectroscopy group (After Quack 2003).

<table>
<thead>
<tr>
<th>Measures of Power</th>
<th>FTIR</th>
<th>Diode direct Abs.</th>
<th>NIR-Diode CW-CRD</th>
<th>ISOS/IRSIMS</th>
</tr>
</thead>
<tbody>
<tr>
<td>(R_P) (\leq 2 \times 10^6)</td>
<td>(\geq 2 \times 10^6)</td>
<td>(\geq 5 \times 10^4)</td>
<td>(\geq 6 \times 10^5)</td>
<td></td>
</tr>
<tr>
<td>(S_P) (\sim 10^7)</td>
<td>(\leq 5 \times 10^4)</td>
<td>(\geq 30 \times 10^4)</td>
<td>(\geq 30 \times 10^4)</td>
<td></td>
</tr>
<tr>
<td>(\Delta \nu) (MHz)</td>
<td>20–70</td>
<td>20–30</td>
<td>(\geq 500)</td>
<td></td>
</tr>
<tr>
<td>(\Delta \nu) (cm(^{-1}))</td>
<td>20000</td>
<td>20 (25000)</td>
<td>500</td>
<td></td>
</tr>
<tr>
<td>(L) (m)</td>
<td>100</td>
<td>100</td>
<td>500</td>
<td></td>
</tr>
</tbody>
</table>

**Powerful spectroscopic tools to apply to atmospheric analysis and quantum chemical kinetics**
ionization producing large biological molecules in a gas phase with collisional cooling in and ion trap and IR–UV double resonance techniques to simplify the spectra. Recently, the discovery of the first negative ion C$_6$H$^-$ (McCarthy et al. 2006) in interstellar clouds has brought about a revival of interest in laboratory studies of anions.

A spectroscopic data base (Vibrational and Electronic Energy Levels (VEEL) of Small Polyatomic Transient Molecules Database) including 1796 short-lived molecules is available from the Commerce Department’s National Institute of Standards and Technology.

ACKNOWLEDGMENTS

We are greatly indebted to S. Albert, K. Keppler Albert, C. Manca Tanner, E. Miloglyadov, and G. Seyfang for help and discussions as well as to Ruth Schüpbach, in particular, for secretarial help in the preparation of this article. Our research has profited from numerous collaborations cited in the references and from financial support from ETH Zürich and Schweizerischer Nationalfonds.

ABBREVIATIONS AND ACRONYMS

<table>
<thead>
<tr>
<th>Acronym</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>CEAS</td>
<td>cavity-enhanced absorption spectroscopy</td>
</tr>
<tr>
<td>cw</td>
<td>continuous wave</td>
</tr>
<tr>
<td>DBR</td>
<td>distributed Bragg reflection</td>
</tr>
<tr>
<td>DFB</td>
<td>distributed feedback</td>
</tr>
<tr>
<td>DH</td>
<td>double heterostructure</td>
</tr>
<tr>
<td>ECDL</td>
<td>external cavity diode laser</td>
</tr>
<tr>
<td>LO</td>
<td>longitudinal optical</td>
</tr>
<tr>
<td>MBE</td>
<td>molecular beam epitaxy</td>
</tr>
<tr>
<td>MRI</td>
<td>magnetic resonance imaging</td>
</tr>
<tr>
<td>NMR</td>
<td>nuclear magnetic resonance</td>
</tr>
<tr>
<td>OPO</td>
<td>optical parametric oscillator</td>
</tr>
<tr>
<td>PAH</td>
<td>polycyclic aromatic hydrocarbons</td>
</tr>
<tr>
<td>PPLN</td>
<td>periodically poled lithium niobate</td>
</tr>
<tr>
<td>QCL</td>
<td>quantum cascade lasers</td>
</tr>
<tr>
<td>QW</td>
<td>quantum well</td>
</tr>
<tr>
<td>VCSEL</td>
<td>vertical cavity surface emitting laser</td>
</tr>
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</table>

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