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UNIVERSITY OF ALBERTA

ABSOLUTE INFRARED ABSORPTION INTENSITIES OF LIQUID CHLOROBENZENE AND TOLUENE

BY

YORAM APELBLAT



A thesis submitted to the Faculty of Graduate Studies and Research in partial fulfillment of the requirements for the degree of DOCTOR OF PHILOSOPHY.

DEPARTMENT OF CHEMISTRY

Edmonton, Alberta

Spring 1996



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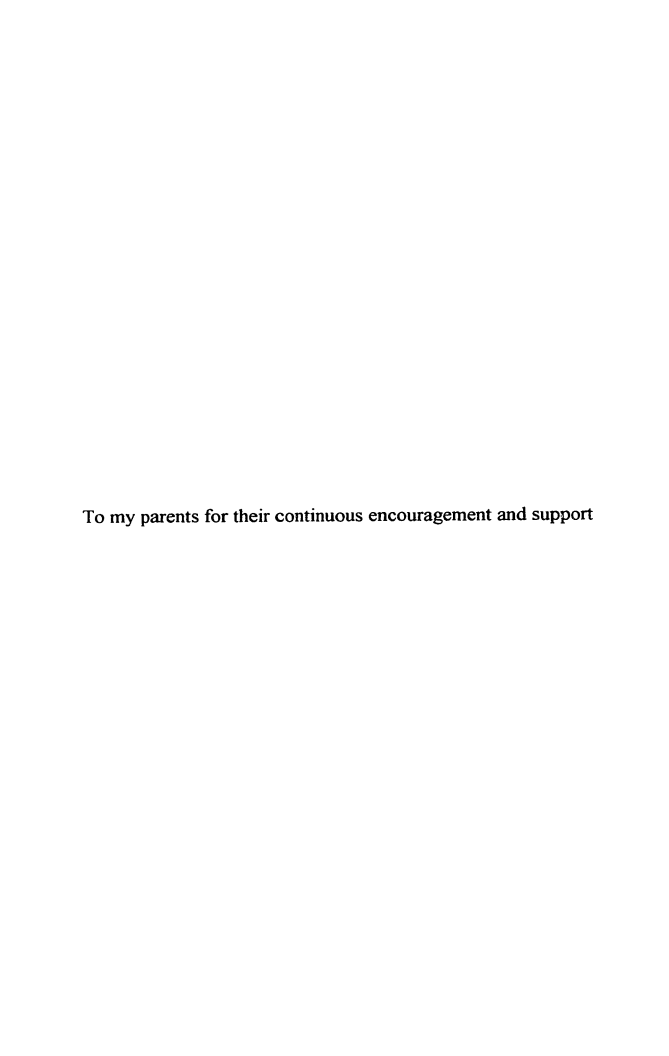
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Abstract

The optical constants of liquid toluene and chlorobenzene were measured across the entire mid-infrared region at 25°C. The estimated accuracy is $\pm 0.2\%$ for the real refractive index and $\pm 2-3\%$ for the imaginary refractive index. The optical constants were used to calculate the molar absorption coefficient and the molar polarizability spectra of these liquids.

The optical constants and the molar absorption coefficient spectra of toluene and chlorobenzene, together with those of liquid benzene and dichloromethane measured separately, were used to establish secondary infrared intensity standards for liquids.

These standards have been published by the International Union of Pure and Applied Chemistry.

Molecular properties are more directly reflected in the imaginary molar polarizability spectrum than in the imaginary refractive index spectrum or the molar absorption coefficient spectrum. In order to calculate the integrated intensities, the imaginary molar polarizability spectrum must be separated into contributions from different transitions. The separation was achieved by curve fitting the imaginary molar polarizability spectra with bands of classical damped harmonic oscillator shape, then calculating the integrated intensities from the parameters of the fitted bands. The accuracy of the integrated intensities is estimated at 3-5% for strong bands and 5-10% for weak bands. The results of this work are corrected for liquid dielectric effects and agree usually within a factor of two with literature values for the gas obtained by

experimental or by *ab initio* calculation. In these cases the agreement must mean that the vibrations in question have very similar intrinsic intensities in the liquid and gas phases.

The procedure used to determine the optical constants of the liquid from transmission measurements is exact but computationally complex. A simpler, approximate method was developed and the conditions under which it yields results of sufficient accuracy were explored.

Finally, a method for data reduction and presentation is given. The data is reduced and incorporated into a format that allows a spectrum to be tabulated in about 1/10 of the space required for a traditional table. The format allows direct retrieval of specific values, and also the retrieval, through a recovery program, of the entire spectrum without loss of intensity and line shape information.

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"Infrared Intensities of Liquids XIV: Accurate Optical Constants and Molar Absorption Coefficients Between 4800 and 450 cm⁻¹ of Chlorobenzene at 25°C from Spectra Recorded in Several Laboratories", J.E. Bertie, R. Norman Jones and Yoram Apelblat, Applied Spectroscopy, 48, 144 (1994).

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P. Norman Janes

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List of Symbols

General

- λ Vacuum wavelength.
- \widetilde{v} Vacuum wavenumber; usual unit cm⁻¹. $\widetilde{v} = 1/\lambda$.
- c Speed of light in vacuum.
- C Molar concentration; usual unit mole L⁻¹.
- $V_{\rm m}$ Molar volume: usual unit cm³ mole⁻¹.
- E Electric field strength vector.
- **D** Electric displacement vector.
- P Electric polarization vector.
- H Magnetic field strength vector.
- I Energy flux. $I = E \times H$.
- σ Conductivity
- $\varepsilon_{\rm o}$ Permittivity of vacuum
- $\mu_{\rm o}$ Permeability of vacuum
- i $\sqrt{-1}$
- o used to indicate a complex quantity.

Wavenumber dependent quantities

 $\hat{n}(\widetilde{v})$ Complex refractive index. $\hat{n}(\widetilde{v}) = n(\widetilde{v}) + i k(\widetilde{v})$.

 $n(\tilde{v})$ Real refractive index.

 $k(\vec{v})$ Imaginary refractive index, also called the absorption index.

 $\hat{\varepsilon}(\widetilde{v})$ Complex dielectric constant. $\hat{\varepsilon}(\widetilde{v}) = \varepsilon'(\widetilde{v}) + i \varepsilon''(\widetilde{v})$.

 $\varepsilon'(\widetilde{\nu})$ Real dielectric constant, usually called the dielectric constant.

 $\varepsilon''(\widetilde{\nu})$ Imaginary dielectric constant, also called the dielectric loss.

 $\hat{\alpha}_{m}(\widetilde{\nu})$ Complex molar polarizability. $\hat{\alpha}_{m}(\widetilde{\nu}) = \alpha_{m}(\widetilde{\nu}) + i \alpha_{m}''(\widetilde{\nu})$.

 $\alpha_{m}(\widetilde{v})$ Real molar polarizability.

 $\alpha_{\rm m}^{\prime\prime}(\tilde{\nu})$ Imaginary molar polarizability.

 $E_{\rm m}(\tilde{v})$ (Decadic) molar absorption coefficient.

 $K\widetilde{v}$) (Decadic) linear absorption coefficient.

Integrated intensities

- A_j Area under band j in 2.303 E_m spectrum. Usual units km mole⁻¹.
- C_i Area under band j in $\widetilde{\nu}\alpha_m''$ spectrum. Usual units km mole⁻¹.

Molecular properties

- μ_j Magnitude of the dipole moment derivative with respect to normal coordinate j.
- $\overrightarrow{R_i}$ Dipole transition moment, $\langle f|\overrightarrow{\mu}|i\rangle$, of the transition that causes band j.

Types of spectra

 $n(\tilde{v})$ vs. \tilde{v} Real refractive index spectrum.

 $k(\widetilde{v})$ vs. \widetilde{v} Absorption index spectrum.

 $E_{\rm m}(\widetilde{\nu})$ vs. $\widetilde{\nu}$ Molar absorption coefficient spectrum.

 $\alpha_m''(\widetilde{\nu})$ vs. $\widetilde{\nu}$ Imaginary molar polarizability spectrum.

Chapter 1 - Introduction

1.1 - Infrared absorption intensities

The absorbance spectrum of a liquid in the mid infrared region (8000 - 400 cm⁻¹) is due mainly to transitions between vibrational energy levels of a molecule. A similar spectrum of the gas phase may show some rotational structure in addition to the vibrational structure. In either case, the vibrational bands in the spectrum are characterised by their positions, their intensities and their band shapes¹.

The band position, measured in wavenumber units, usually cm⁻¹, is the easiest of the three to determine accurately and over the years its measurement accuracy has been improved and has reached such a high degree of accuracy that it was possible already in the early 1960s to publish tables of standard wavenumbers to calibrate spectrometers² in the 4300-600 cm⁻¹ region. The estimated accuracy of the standard wavenumbers was about 0.01-0.1 cm⁻¹. The tables were later revised³ and the range extended to 1 cm⁻¹ with an estimated accuracy of 0.001-0.005 cm⁻¹ throughout the range and < 0.0002 cm⁻¹ for the fundamental and first overtone of carbon monoxide.

The intensity and band shape are more difficult to measure accurately. Intensity measurements were done as early as the late 1920s, but their accuracy^{4,5} wasn't as good as that of the wavenumber.

From the theoretical viewpoint, it was recognized that the intensity is proportional to the dipole transition moment. For fundamental vibrations, under the double harmonic approximation, this is proportional to the square of the dipole moment

derivative with respect to the normal coordinate, $\|\partial \mu/\partial Q_j\|^2$. The intensities reflect the change in electron distribution in the molecule during the atomic displacements⁴. However, the use of the intensity was hampered by the ambiguity of the sign of $\partial \mu/\partial Q_j$ and by unreliable transformation of the observed $\partial \mu/\partial Q_j$ to related motions defined by internal coordinates, $\partial \mu/\partial R_j$.

During the 1970s the study of intensities improved considerably⁴. Advances in normal coordinate calculations combined with *ab initio* quantum mechanical calculations helped solve the sign ambiguity of the dipole moment derivatives. At the same time, the introduction of Fourier Transform Infra Red (FTIR) spectrometers based on Michelson's interferometer and the use of computer controlled instruments gave far more reproducible spectra than dispersive instruments. Later it was shown by V. Behnam and J.E. Bertie that FTIR spectrometers were also accurate in addition to being reproducible. Their studies^{5,6} of intensities of several organic liquids made with a non-calibrated FTIR spectrometer showed that their intensity values were more precise and within the error limits of R.N. Jones and co-workers' earlier work, which was made on a dispersive instrument and calibrated against primary standards⁷.

As laboratory computers, powerful software and stable FTIR spectrometers became increasingly available and inexpensive in the 1980s and 1990s, the measurement over a wide wavenumber range and interpretation of accurate absolute infrared intensities of molecules is possible today to an extent unthinkable 20 years ago. The studies reported in this thesis form part of a research program that is aimed at measuring

and using both the intensity and the wavenumber information to improve our chemical and physical knowledge of molecular vibrations in the liquid phase.

An important feature of the program is that the real and imaginary refractive indices are measured over a wide wavenumber range in the infrared. The results of any infrared experiment can be calculated from these quantities. Thus, they are fundamental physical properties, obtained as a part of the program.

1.2 - Overview of thesis

The first goal of the work for this thesis was to assist in the development of infrared secondary intensity standards. The work started in 1985 when J.E. Bertie and V. Behnam^{5,6}, showed that FTIR intensity measurements were precise to ~1%^{5,6} and probably accurate to ~3%⁵⁻⁷. Further investigation was needed to identify and eliminate the systematic errors influencing the measurements. This was done by comparing the intensities of four organic liquids⁸⁻¹¹ determined from transmission measurements made by different workers using different instruments in different laboratories. The equations relating the transmission measurement to various intensity quantities are given in Section 1.3.

The resulting good agreement led to the acceptance of the intensities of 43 selected bands as secondary standards by the International Union of Pure and Applied Chemistry (IUPAC). Later these secondary standards were published as a monograph 12.

The intensity measurements of liquid toluene and liquid chlorobenzene, which were part of this project, are given in Chapters 2 and 3, respectively.

The intensities reported as standards were the real and imaginary refractive index spectra, also known as the optical constant spectra. The molar absorption coefficient spectra of the liquids were also reported. The real and imaginary refractive indices are fundamental physical quantities, from which other measures of infrared absorption intensities such as the complex dielectric constant and the complex molar polarizability can be calculated. The relationship between these quantities is given in Section 1.3

The second goal of the work for this thesis was to determine the physical and chemical information contained in the intensities. For that purpose, the complex molar polarizability spectra of liquid toluene and chlorobenzene were calculated under the assumption of the Lorentz local field, to relate the local electric field that acts on the molecule in the liquid to the macroscopic electric field of the radiation. The imaginary molar polarizability can then be related to the transition dipole moment and under the assumption of electrically and mechanically harmonic vibration, to the square of the dipole moment derivative with respect to the normal coordinate¹³.

To relate these experimental intensities to particular vibrational modes of the molecules, fundamental frequencies must first be assigned to particular vibrations of the molecule, e.g. the CC stretch, CH bend, etc. Despite a considerable number of experimental and theoretical studies reported in the literature on toluene and chlorobenzene 18,22,23,35-47, there are still some disagreements about the assignments of the

fundamentals. In Chapter 4, the assignment of the spectra of toluene and chlorobenzene is discussed, based on our experimental spectra, previous studies, and normal coordinate calculations which relate the vibrations of benzene to those of substituted benzene derivatives of C_{2v} symmetry, chlorobenzene and toluene. For the latter molecule the CH₃ group was approximated by a point of mass 15 in the calculations in order to correlate the assignment of the vibrations of the phenyl group with those in the other molecules.

Once the fundamentals are assigned, it is possible to relate the intensities to the vibrational transitions in the different molecules. This is done in Chapter 5 where experimental and curve-fitted spectra are used to relate the intensities of toluene and chlorobenzene to the dipole moment derivatives. The curve fitting is necessary in order to separate contributions from different transitions and to make sure that the intensities in the wings of a band are included in the band area. It is noteworthy that the bands are nearly Lorentzian, so that it is necessary to integrate over more than ±16 times the full width at half height in order to include >98% of the area⁴⁸.

The procedure used to determine the optical constants of the liquid from transmission measurements was first developed by R.N. Jones and co-workers at the National Research Council of Canada⁴⁹⁻⁵³ and later improved by J.E. Bertie and C.D. Keefe⁵⁴ at the University of Alberta. During this iterative procedure, the reflection losses are calculated at all interfaces of the cell, namely air-window-liquid-window-air. The method is exact but computationally complex. In Chapter 6, an approximate

method is developed to calculate the optical constants from transmission measurements.

The conditions under which the approximation method yields results of sufficient accuracy are explored.

The presentation of quantitative spectral data for publication in the literature is not an easy task. A graph is necessary to convey the spectrum qualitatively. However, to convey the spectrum quantitatively, values of both wavenumber and intensity must be tabulated. Since the space required to publish all data points in a traditional table is very large, only selected pairs of wavenumber and intensity values have usually been given, mostly at peaks of bands^{7,55,56}. This practice was satisfactory when the intensity accuracy and precision were sufficiently low that only strong bands were reported and the shapes of bands were not considered important. Today, however, both intensity and line shapes can be precisely and accurately measured, and have gained in potential importance. The practice of reporting just band peak data is no longer valid and a method is required to allow quantitative ordinate values to be reported over the whole spectrum range in an acceptably small space. The solution to this problem is appropriate data reduction and appropriate compression of the presentation format.

In Chapter 7, a method for data reduction and presentation is given. The data is reduced and incorporated into what we call a Compact Table. The Compact Table format allows a spectrum to be tabulated in about 1/10 of the space required for a traditional table. It allows specific values to be read from the table directly and also allows, through a recovery program, the entire spectrum to be retrieved without loss of

intensity and line shape information, within the experimental error. The programs for the creation of the Compact Table and recovery of the data are given in Chapter 7 and their use is demonstrated with the spectra of chlorobenzene.

Finally, Chapter 8 contains a summary of the results of this thesis and presents some possible future extensions of the work.

1.3 - Electromagnetic theory and optical properties

When light is transmitted through a sample, part of it is absorbed by the molecules in the sample. The extent of the macroscopic response of the sample to the perturbation caused by the electric field is determined by the complex refractive index of the sample, $\hat{n} = n + ik$. The real part of the refractive index, n, is what is usually called simply the refractive index. The imaginary refractive index, k, is a measure of absorption intensity, and is sometimes called the absorption index. Together the refractive indices are frequently called the optical constants.

The fraction of light transmitted by a medium is related to the optical constants through Maxwell's equations⁵⁷. For isotropic conducting media, the curl equations relate the time and space derivatives of the magnetic, H, and electric fields, E, by

$$\nabla \times \mathbf{E} = -\mu \,\mu_0 \frac{\partial \mathbf{H}}{\partial t} \tag{1.3.1}$$

$$\nabla \times \boldsymbol{H} = \sigma \boldsymbol{E} + \varepsilon \, \varepsilon_0 \, \frac{\partial \boldsymbol{E}}{\partial t} \tag{1.3.2}$$

and the divergence conditions denote the absence of charge⁵⁷

$$\nabla \bullet E = 0 \tag{1.3.3}$$

$$\nabla \bullet \mathbf{H} = 0 \tag{1.3.4}$$

Here μ is the relative permeability, μ_0 is the permeability of vacuum, ε is the relative dielectric constant of the medium, ε_0 is the permittivity of vacuum and σ is the conductivity of the medium.

Equations 1.3.3 and 1.3.4 and the fact that $A \times (B \times C) = (A \cdot C) B - (A \cdot B) C$ give equations 1.3.5 and 1.3.6

$$\nabla \times (\nabla \times \mathbf{E}) = (\nabla \cdot \mathbf{E}) \nabla - (\nabla \cdot \nabla) \mathbf{E} = 0 - \nabla^2 \mathbf{E} = -\nabla^2 \mathbf{E}$$
 (1.3.5)

$$\nabla \times (\nabla \times \mathbf{H}) = (\nabla \cdot \mathbf{H}) \nabla - (\nabla \cdot \nabla) \mathbf{H} = 0 - \nabla^2 \mathbf{H} = -\nabla^2 \mathbf{H}$$
 (1.3.6)

Equations 1.3.5, 1.3.6, 1.3.1 and 1.3.2 give equations 1.3.7 and 1.3.8

$$-\nabla^2 \mathbf{E} = -\mu \,\mu_0 \,\sigma \frac{\partial \mathbf{E}}{\partial t} - \mu \,\mu_0 \,\varepsilon \,\varepsilon_0 \,\frac{\partial^2 \mathbf{E}}{\partial t^2} \tag{1.3.7}$$

$$-\nabla^2 \mathbf{H} = -\mu \,\mu_o \,\sigma \frac{\partial \mathbf{H}}{\partial t} - \mu \,\mu_o \,\varepsilon \,\varepsilon_o \,\frac{\partial^2 \mathbf{H}}{\partial t^2} \tag{1.3.8}$$

One solution to these equations is 58

$$E = E_0 e^{-i\omega \left(t - \frac{x}{\hat{v}}\right)}$$
 (1.3.9)

$$H = H_0 e^{-i\omega \left(t - \frac{x}{\hat{v}}\right)}$$
 (1.3.10)

which is a representation of a wave propagating in the x direction with angular frequency $\omega = 2\pi f$, f = frequency, and with complex velocity \hat{v} , that holds true provided that

$$\frac{1}{\sqrt{2}} = \mu \,\mu_0 \,\varepsilon \,\varepsilon_0 + i \,\mu \,\mu_0 \frac{\sigma}{\omega} \tag{1.3.11}$$

 \hat{v} can be written as $\frac{c}{\hat{n}}$, where \hat{n} is the complex refractive index and c is the speed of light in vacuum. Equation 1.3.11 shows that in vacuum, where $\sigma = 0$ and $\mu = \varepsilon$ = 1, the velocity of light is real and denoted by c, where $c = (\mu_0 \ \varepsilon_0)^{-1/2}$. The use of these expressions in equation 1.3.11 yields

$$\hat{n}^2 = \mu \ \varepsilon + i \frac{\sigma \mu}{\varepsilon_0 \omega} \tag{1.3.12}$$

 \hat{n} can be defined as

$$\hat{n} = n + ik \tag{1.3.13}$$

where n is the real refractive index and k is the imaginary refractive index.

Substitution of equation 1.3.13 into equation 1.3.9 and the use of $\hat{v} = \frac{c}{\hat{h}}$, gives

$$E = E_0 e^{-i\omega \left(t - \frac{x}{\hat{v}}\right)} = E_0 e^{-i\omega \left(t - \frac{x\hat{n}}{c}\right)} = E_0 e^{-i\omega \left(t - \frac{xn}{c}\right)} e^{-\left(\frac{\omega k}{c}x\right)}$$
(1.3.14)

Similarly, use of equations 1.3.13 in 1.3.10 gives

$$H = H_0 e^{-i\omega \left(t - \frac{x}{\hat{v}}\right)} = H_0 e^{-i\omega \left(t - \frac{x\hat{n}}{\hat{c}}\right)} = H_0 e^{-i\omega \left(t - \frac{x\hat{n}}{\hat{c}}\right)} e^{-\left(\frac{\omega k}{\hat{c}}x\right)}$$
(1.3.15)

Equations 1.3.14 and 1.3.15 show that the real refractive index controls the phase of the field while the imaginary part controls the attenuation of the field with propagation distance.

The energy flux, I, is given by

$$I = E \times H = I_0 e^{-2i\omega \left(t - \frac{xn}{c}\right)} e^{-2\left(\frac{\omega k}{c}x\right)}$$
(1.3.16)

where $I_o = E_o \times H_o$.

In a transmission measurement, the reduction of the energy of a light beam passing through a unit thickness of the sample is given by

$$\frac{\mathrm{d}\mathbf{I}}{\mathrm{d}\mathbf{r}} = -\mathbf{K}'\mathbf{I} \tag{1.3.17}$$

where K is the (Napierian) linear absorption coefficient. If the path length through the sample is d, equation 1.3.17 can be integrated to give

$$I = I_0 e^{-K'd}$$
 (1.3.18)

so that
$$K = -\frac{1}{d} \ln \left(\frac{I}{I_0} \right)$$
 (1.3.19)

Equation 1.3.18 is also known as Lambert's law⁵⁹. Chemists, however prefer to use the Beer-Lambert law⁵⁹

$$I = I_0 10^{-Kd} = I_0 10^{-E_m Cd}$$
 (1.3.20)

where K is the linear (decadic) absorption coefficient, $E_{\rm m}$ is the molar (decadic) absorption coefficient and C is the molar concentration. By rearrangement of equation 1.3.20, K can be written as

$$K = -\frac{1}{d}\log_{10}\left(\frac{I}{I_0}\right) \tag{1.3.21}$$

From equations 1.3.16 and 1.3.18, the energy attenuation per unit thickness or distance travelled by the wave is

$$e^{-K'd} = e^{-2\left(\frac{\omega dk}{c}\right)} \tag{1.3.22}$$

because $\omega = 2\pi f$, where f is the frequency, and $\tilde{v} = f/c$, the wavenumber in vacuum, the (Napierian) linear absorption coefficient, K, is related to the imaginary refractive index, k, by $K = 4\pi \tilde{v}k$, whereas the decadic linear absorption coefficient is given by

$$K = \frac{K'}{\ln 10} = \frac{4\pi \,\widetilde{\nu}k}{2.303} \tag{1.3.23}$$

The absorbance, A, is defined as $-\log_{10}(\frac{I}{I_o})$. From equations 1.3.21 and 1.3.23 it is given by

$$A = \frac{4\pi \,\widetilde{\nu}kd}{2.303} \tag{1.3.24}$$

It is important to note that the absorbance used here is the absorbance defined by IUPAC, namely that due only to loss of light by absorption by the sample⁶⁰. In a typical

experiment, the experimentally observed absorbance includes the absorbance by the sample and absorbance by other elements such as the windows of the cell. It is called in this laboratory the experimental absorbance, EA. The experimental absorbance must therefore be corrected for energy loss other than through absorption by the sample before it can be used in these equations. The procedure for this correction is implemented through program RNJ46A^{53,54}. The program and an alternative approximate procedure are discussed in Chapter 6. Throughout the thesis, the term absorbance is used according to the IUPAC definition, namely as due solely to absorption by the sample.

From equations 1.3.20, 1.3.23 and 1.3.24, the molar absorption coefficient is given by

$$E_m = \frac{A}{Cd} = \frac{4\pi \tilde{\nu}k}{2.303C} \tag{1.3.25}$$

Other infrared intensity quantities can be obtained from the refractive indices. Eq. 1.3.12 relates the square of the complex refractive index to the complex dielectric constant. $\hat{\epsilon}$.

$$\hat{n}^2 = n^2 + 2ink - k^2 = \mu \varepsilon + i \frac{\sigma \mu}{\varepsilon_0 \omega} = \mu (\varepsilon + i \frac{\sigma}{\varepsilon_0 \omega}) = \mu \hat{\varepsilon}$$
 (1.3.26)

 $\hat{\varepsilon}$ is defined in a similar manner to \hat{n} , as

$$\hat{E} = E' + i E'' \tag{1.3.27}$$

where the real part, ε' , is the (real) dielectric constant and the imaginary dielectric constant, ε'' , is also known as the dielectric loss.

For diamagnetics, $\mu \approx 1$, so $\hat{n}^2 \approx \hat{\varepsilon}$. From Equations 1.3.13, 1.3.26 and 1.3.27, the real and imaginary parts of the dielectric constants can be related to the refractive indices as:

$$\varepsilon' = n^2 - k^2 \tag{1.3.28a}$$

$$\varepsilon'' = 2nk \tag{1.3.28b}$$

The electric displacement, D, is given by 61

$$\mathbf{D} = \hat{\boldsymbol{\varepsilon}} \, \mathbf{E} \tag{1.3.29}$$

It is also the sum of the electric and vector and the polarization vector, P, where all of these vectors can be complex.

$$\boldsymbol{D} = \boldsymbol{E} + 4\pi \boldsymbol{P} \tag{1.3.30}$$

The polarization, i.e. the electric dipole moment per unit volume, is also given in terms of the complex polarizability^{61,62}, $\hat{\alpha}$, as

$$P = N \hat{\alpha} \left(E + \frac{4\pi}{3} P \right) \tag{1.3.31}$$

where N is the number of molecules per unit volume. $E + \frac{4\pi}{3}P$ is the Lorentz local field derived by calculating the field within a microscopic cavity in an isotropic dielectric medium^{62,63}.

Combination of equations 1.3.29, 1.3.30 and 1.3.31 yields the Lorentz-Lorenz equation⁶²,

$$\frac{\hat{n}^2 - 1}{\hat{n}^2 + 2} = \frac{\hat{\varepsilon} - 1}{\hat{\varepsilon} + 2} = \frac{4\pi}{3} N \hat{\alpha}$$
 (1.3.32)

Since $\hat{\alpha}$ is a molecular property and, thus, has a small value, the complex molar polarizability is introduced as $\hat{\alpha}_m = N_A \hat{\alpha} = \alpha_m' + i\alpha_m''$ where N_A is Avogadro's number. Multiplication of equation 1.3.32 by N_A and the use of the molar volume, $V_m = \frac{N_A}{N}$, gives the real and imaginary parts of the molar polarizability in terms of ε' and ε'' .

$$\alpha_{\rm m}' = \frac{3V_{\rm m}}{4\pi} \frac{(\varepsilon'-1)(\varepsilon'+2) + \varepsilon''^2}{(\varepsilon'+2)^2 + \varepsilon''^2} \tag{1.3.33a}$$

$$\alpha_{\rm m}^{"} = \frac{9V_{\rm m}}{4\pi} \frac{\varepsilon^{"}}{(\varepsilon'+2)^2 + \varepsilon^{"^2}} \tag{1.3.33b}$$

It is important to note that all quantities are frequency or wavenumber dependent, and thus should really be denoted as $n(\widetilde{v})$, $k(\widetilde{v})$, $E_m(\widetilde{v})$, $\varepsilon'(\widetilde{v})$, $\varepsilon''(\widetilde{v})$, $\alpha'_m(\widetilde{v})$ and $\alpha''_m(\widetilde{v})$.

1.4 - The relationship of integrated intensities to molecular properties

There are several methods to define the integrated intensity of band j. One is as the integrated intensity, $A_j^{48,64,65}$ which is the area under the $E_m(\tilde{\nu})$ spectrum

$$A_{J} = 2.303 \int E_{m}(\widetilde{\nu}) d\widetilde{\nu}$$
 (1.4.1)

Another definition of the integrated intensity is as C_j , the area under the $\widetilde{\nu}\alpha_m''(\widetilde{\nu})$ spectrum⁴⁸

$$C_{j} = \int \widetilde{v} \alpha_{m}''(\widetilde{v}) d\widetilde{v}$$
 (1.4.2)

In both equations, the integration is over the entire band j. These experimental integrated intensities must be related to molecular properties. Calculation of the complex molar polarizability of randomly oriented molecules from quantum mechanics yields 48,58,63,66

$$C_{j} = \int \widetilde{v} \alpha_{m}''(\widetilde{v}) d\widetilde{v} = \frac{N_{A}\pi}{3hc} g_{j} \widetilde{v}_{j} |R_{j}|^{2}$$
(1.4.3)

where h is Planck's constant, c is the velocity of light in vacuum, g_j is the degeneracy and $\overrightarrow{R_j}$ is the dipole transition moment. Under the assumption of mechanical and electrical harmonicity and the assumption that all of the hot bands of the fundamental contribute to the fundamental band, and the expression for the transition moment¹³ $<1|\overrightarrow{\mu}|0>$, equation 1.4.3 yields⁴⁸

$$\sum_{j} = \int \widetilde{v} \alpha_m''(\widetilde{v}) d\widetilde{v} = \frac{N_A g_j}{24\pi c^2} \mu_j^2$$
 (1.4.4)

where μ_j^2 is the square of the dipole moment derivative with respect to the j^{th} normal coordinate, e.g. $\mu_j^2 = |\partial \mu/\partial Q_j|^2$. The same result is obtained using the classical damped harmonic oscillator (CDHO) model^{48,58,63,66,67}.

Equations 1.4.3 and 1.4.4 relate the experimental integrated intensity to molecular properties through the assumption of the Lorentz local field. If an additional assumption is made, namely that the absorption bands are far apart so that the contribution across band j from all oscillators other than j is real and constant, $n(\tilde{v})$ becomes n, the constant refractive index the sample would have in the region of the band if the band were not present which is often (incorrectly) taken as the average across the band. Then Equations 1.4.1, 1.3.25 and $\hat{n}^2 = \hat{\varepsilon}$ are used to yield⁶⁸⁻⁷⁰

$$A_{j} = 2.303 \int E_{m}(\tilde{v}) d\tilde{v} = \frac{N_{A}\pi}{3c^{2}} g_{j} \frac{1}{n} (\frac{n+2}{3})^{2} \mu_{j}^{2}$$
 (1.4.5)

Values of μ_j^2 can be obtained from the rearrangement of either Equation 1.4.4 or Equation 1.4.5. However, both equations involve integration, i.e. obtaining the area under either the $\tilde{\nu}\alpha_m''$ or E_m spectra. The integration is not always easy, as the integration limits are not always clear because of overlapping bands. Equation 1.4.5 presents the further difficulties that values of either \underline{n} or $\sqrt{\underline{\varepsilon}}$ are needed. Thus, Equation 1.4.4 is preferred to 1.4.5⁴⁸. Since, for small oscillations the CDHO model yields the same results as the quantum mechanical treatment⁶⁵, it may be possible to fit the experimental α_m'' spectrum with CDHO-shaped bands and use the total area under each fitted band instead of under the experimental spectrum. The integration limits of the fitted bands are no longer uncertain and C_j values are easily obtained. This approach is used in Chapter 5 to determine the integrated intensities and dipole moment derivatives of liquid chlorobenzene and toluene.

Chapter 2 - Accurate Optical Constants and Molar Absorption Coefficients

Between 6500 and 435 cm⁻¹ of Toluene at 25°C, from Spectra Recorded in Several

Laboratories*

2.1 - Introduction

This paper continues the report of a program to measure quantitative infrared absorption intensities. It presents the agreement between absolute absorption intensities of liquid toluene at 25°C measured by different spectroscopists in this and other laboratories, using instruments made by several different manufacturers. A similar study of the absolute absorption intensities of liquid benzene was recently published and a report of the absolute absorption intensities of liquid chlorobenzene accompanies this paper. In part, this work contributes to the International Union of Pure and Applied Chemistry project to develop secondary standards for intensity measurements in infrared spectroscopy.

In the present study, experimental absorbance spectra of liquid toluene at 25°C have been measured by three different spectroscopists in this laboratory (two for the current work and one about 8 years ago) and also by three spectroscopists in other laboratories. Spectroscopists in different laboratories used instruments from different manufacturers. From these experimental absorbance spectra, the real and imaginary

^{*} A version of this chapter has been published. Bertie, Jones, Apelblat and Keefe, Appl. Spectrosc., 48, 127 (1994)

refractive index spectra have been calculated in this laboratory by the methods described previously^{1,4}.

Attention was focussed on the absorption index (imaginary refractive index) spectra, $k(\tilde{v})$ vs. \tilde{v} . For each spectroscopist the average $k(\tilde{v})$ spectrum was evaluated together with its precision. In order to determine the agreement between different spectroscopists, the peak heights and the areas beneath the bands in these average spectra were compared. The agreement was excellent. These average spectra, one from each spectroscopist, were themselves averaged, unweighted, to yield a final absorption index spectrum that is presented here as the best currently available.

This absorption index spectrum, $k(\widetilde{\nu})$ vs. $\widetilde{\nu}$, was used to calculate^{1,4} the molar absorption coefficient spectrum, $E_m(\widetilde{\nu})$ vs. $\widetilde{\nu}$, and, via Kramers-Kronig transformation, the real refractive index spectrum, $n(\widetilde{\nu})$ vs. $\widetilde{\nu}$.

The real and imaginary refractive index spectra and the molar absorption coefficient spectrum are presented as graphs and as tables. We also report the average area under individual bands or groups of bands, and the average peak heights, in the $k(\widetilde{\nu})$ spectrum obtained by each spectroscopist, as well as the weighted and unweighted averages of these averages. Further, the peak heights and the areas under the bands in the average molar absorption coefficient spectrum are reported. For certain spectral regions, the real and imaginary refractive indices and the molar absorption coefficients, as well as the areas under the molar absorption coefficient bands, will be submitted to Commission I.5 of the International Union of Pure and Applied Chemistry for

consideration as secondary absorption intensity standards.

2.2 - Method and experimental

The toluene used in this laboratory was of spectroscopic or reagent grade.

Samples were purified by fractional freezing one to three times and were checked by gas chromatography and infrared spectroscopy. No impurities were detected. Samples were kept over molecular sieve to ensure dryness.

The experimental and instrumental details of this work have been described^{1,4,5} and are summarised briefly here. All of the spectra from this laboratory were measured with a Bruker IFS 113V spectrometer. Triangular apodization was used in the early work from this laboratory⁵. The recent work in this laboratory used trapezoidal apodization¹. A Globar source, a 10 mm aperture, and a deuterated triglycine sulfate, DTGS, detector were used for all spectra measured in this laboratory. The optical retardation velocity was 0.665 cm s⁻¹. The spectra were recorded at nominal resolution of 1 cm⁻¹ and one level of zero-filling was used in the Fourier transform.

Experimental absorbance spectra of toluene were measured ^{1,4} in fixed path length cells with KBr windows and path lengths between 11 and 500 μm. To determine the linear absorption coefficients at the anchor points ^{1,4}, spectra were also measured in KBr cells with path lengths of 500 and 1500 μm, and in variable path length NaCl and CaF₂ cells with path lengths up to 5 mm. The path lengths ce the cells were determined from the fringe patterns in the experimental absorbance spectra of the empty cells by program

RNJ22A 1 . For the variable path cells, which gave weaker fringe patterns than the cells with fixed path lengths, path lengths greater than 700 μ m could not be determined in this way. They were found by calibrating the cell micrometer readings from the fringe patterns for path lengths up to 700 μ m, and assuming the calibration held for thicknesses above 700 μ m.

In addition to the experimental absorbance spectra recorded in this study, spectra reported previously 5.6 by V. Behnam were also used. Further, three spectroscopists in other laboratories kindly supplied spectra which were recorded under 1 or 2 cm⁻¹ nominal resolution with normal conditions and "good analytical laboratory technique".

Toluene used in other laboratories was generally of reagent, spectro- or HPLC grade, and spectra from other laboratories were taken to be acceptable if they showed no unexpected peaks. The spectra from other laboratories were recorded on Digilab and Nicolet spectrometers. One spectroscopist used a mercury cadmium telluride detector at 77 K, and the other two used room-temperature DTGS detectors. Happ-Genzel apodization was used in two cases, and triangular apodization in one case. In this way, we obtained from this and three other laboratories spectra recorded by six different spectroscopists on four different models of FT spectrometer by three different manufacturers.

The linear (decadic) absorption coefficient, $K(\widetilde{v})$, is the absorbance per unit length, and is related to the molar absorption coefficient by $K(\widetilde{v}) = C E_m(\widetilde{v})$, where C is the molar concentration. Values of $K(\widetilde{v})$ are needed at anchor points in the base in

order to correct the baselines^{1,4}. They were determined from experimental absorbance spectra in cells with path lengths two to ten times greater than those used for studying the nearby peaks. The experimental absorbance spectra from all sources were then converted to absorption index spectra by program RNJ46A⁴, using this anchor point information. With two exceptions, each spectrum was only used in those regions in which the bands had peak absorbances between 0.2 and 2.0. The exceptions were the two spectra of the very intense band at 729 cm⁻¹ (see later) which were used with peak absorbances between 2.0 and 2.5.

2.3 - Results

2.3.1 Imaginary refractive index spectra.

The baseline correction procedure 1,4 requires the linear absorption coefficient of the liquid at two or more anchor points in the baseline of the spectral region under study. Table 2.1 summarizes the wavenumbers of the anchor points, the measured linear absorption coefficients, $K(\tilde{v})$, and their 95% confidence limits, and the cell path lengths used to determine them. Table 2.1 includes the uncertainties in the absorption index, $k(\tilde{v})$, values at the anchor points that result from the precision of the linear absorption coefficients.

Table 2.2 shows the spectral regions that were used in the calculation⁴ of the absorption index spectra from the experimental absorbance spectra, together with the cell thicknesses used, the value of the real refractive index at high wavenumber, n_{-} , and

Table 2.1 - Linear absorption coefficients at anchor points for liquid toluene at 25°C.

| Wavenumber | Cell Pathlengths | ~ | 95 % confidence | Uncertainty in |
|---------------------|------------------|--|---------------------------|------------------------|
| (cm ⁻¹) | (mm) | $K(\widetilde{\nu})$ (cm ⁻¹) | limit (cm ⁻¹) | k(v)* |
| 6358.6 | 2 - 4.5 | \mathbf{O}_{p} | | $< 6.0 \times 10^{-7}$ |
| 5450.8 | 1.5 - 4.5 | 0.251 | 0.031 | 1.0×10^{-6} |
| 4806.7 | 1.5 - 4.5 | 0.361 | 0.019 | 7.2 x 10 ⁻⁷ |
| 4480.3 | 1.5 - 4.5 | 1.86 | 0.01 | 4.1 x 10 ⁻⁷ |
| 4146.2 | 1.5 - 4.5 | 3.39 | 0.01 | 4.4×10^{-7} |
| 3970.2 | 1.5 - 4.5 | 3.94 | 0.09 | 4.2×10^{-6} |
| 3748.0 | 1.5 - 4.5 | 1.83 | 0.01 | 4.9×10^{-7} |
| 3483.8 | 1.5 - 4.5 | 1.13 | 0.02 | 1.1 x 10 ⁻⁶ |
| 3355.5 | 1.5 - 4.5 | 1.56 | 0.01 | 5.5×10^{-7} |
| 3159.3 | 0.5 - 2.5 | 7.77 | 0.03 | 1.7×10^{-6} |
| 2759.6 | 0.5 - 3.5 | 5.46 | 0.02 | 1.3 x 10 ⁻⁶ |
| 2562.9 | 1.5 - 4.5 | 2.547 | 0.007 | 5.0×10^{-7} |
| 2219.2 | 1.5 - 4.5 | 1.612 | 0.006 | 5.0 x 10 ⁻⁷ |
| 1977.6 | ~1.5 | 5.97 | 0.01 | 9.3 x 10 ⁻⁷ |
| 1909.7 | 0.5 - 1.5 | 4.97 | 0.03 | 2.9×10^{-6} |
| 1754.9 | ~1.5 | 6.04 | 0.01 | 1.0×10^{-6} |
| 1648.4 | 0.5 - 1.5 | 7.79 | 0.03 | 3.3×10^{-6} |
| 1556.3 | 0.1 - 0.5 | 26.7 | 0.2 | 2.3×10^{-5} |
| 1398.6 | ~0.5 | 33.0 | 0.3 | 3.9×10^{-5} |
| 1231.3 | 0.5 - 1.5 | 7.49 | 0.03 | 4.5×10^{-6} |
| 919.9 | 0.5 - 1.5 | 11.11 | 0.03 | 6.0×10^{-6} |
| 776.7 | ~0.5 | 18.1 | 0.1 | 2.4 x 10 ⁻⁵ |
| 708.7 | ~0.05 | 70.2 | 0.7 | 1.8 x 10 ⁻⁴ |
| 636.9 | 0.5 - 1.5 | 5.07 | 0.03 | 8.6 x 10 ⁻⁶ |
| 508.2 | ~1.5 | 3.99 | 0.02 | 7.2 x 10 ⁻⁶ |
| 436.3 | ~1.5 | 6.21 | 0.03 | 1.3 x 10 ⁻⁵ |

a - The uncertainty in $k(\tilde{v})$, $\Delta k(\tilde{v})$, was calculated from the 95% confidence limit of $K(\tilde{v})$, $\Delta K(\tilde{v})$, by $\Delta k(\tilde{v}) = 2.303 \ \Delta K(\tilde{v})/(4\pi \tilde{v})$.

b - The value of $K(\tilde{\nu})$ was set to 0 for this anchor point because the absorbance is less than 0.01 in a 4.5 mm cell, so K is less than 0.02 cm⁻¹.

Table 2.2 - Path lengths, high-wavenumber refractive index, and number of spectra from each spectroscopist, for the regions processed.

| | Pathlengths | | | | - | | | | |
|----------------------------|-------------|----------------|-----|----|---|---|----|----|-------|
| Region (cm ⁻¹) | used (µm) | n _∞ | CDK | Α | В | С | VB | YA | Total |
| 6500 - 4500 | ~ 500 | 1.476 | 3 | 0 | 0 | 0 | 1 | 2 | 6 |
| 4500 - 3150 | ~ 500 | 1.476 | 6 | 2 | 0 | 0 | 4 | 4 | 16 |
| 3170 - 2750 | 10 - 35 | 1.471 | 8 | 6 | 0 | 2 | 11 | 3 | 30 |
| 2775 - 1970 | ~ 500 | 1.475 | 6 | 2 | 0 | 0 | 4 | 4 | 16 |
| 2000 - 1640 | 50 - 210 | 1.474 | 2 | 2 | 1 | 2 | 3 | 1 | 11 |
| 1655 - 1540 | 10 - 60 | 1.456 | 11 | 6 | 0 | 2 | 16 | 4 | 39 |
| 1565 - 1390 | 8 - 20 | 1.456 | 4 | 2 | 0 | 4 | 10 | 2 | 22 |
| 1420 - 770 | 35 - 100 | 1.470 | 8 | 10 | 0 | 4 | 6 | 2 | 30 |
| 780 - 700 | ~ 8 | 1.419 | 0 | 0 | 0 | 2 | 0 | 0 | 2 |
| 715 - 630 | 8 - 14 | 1.520 | 4 | 0 | 0 | 2 | 7 | 2 | 15 |
| 670 - 475 | ~ 500 | 1.513 | 3 | 2 | 0 | 0 | 2 | 2 | 9 |
| 510 - 435 | 8 - 20 | 1.483 | 8 | 2 | 0 | 4 | 9 | 3 | 26 |

a - $n_{_{\!\varpi}}$ is the real refractive index at the highest wavenumber in the region.

the number of spectra from each spectroscopist, for each region. The value of n_m for each region is required by the Kramers-Kronig transform in program RNJ46A, which yields the real refractive index spectrum for the region. The spectra recorded in this laboratory are labeled with the initials CDK, VB and YA, identifying C.D. Keefe, V. Behnam and Y. Apelblat, respectively. The other collaborators are identified simply by a single letter, A to C. The real refractive index at the highest wavenumber in each region, n_m , was read either from the graphs in reference 7 or, in most cases, from tables of $n(\widetilde{\nu})$ in this laboratory which were calculated from the $k(\widetilde{\nu})$ spectra in reference 5.

The peak heights and the areas under the bands were measured for each

Table 2.3 - Spectroscopist average areas under the absorption index, $k(\widetilde{v})$, bands.⁴

| Region (cm ⁻¹) | CDK | A | В | C | VB | YA |
|----------------------------|--------------|--------------|--------|-------------|--------------|--------------|
| 6307.1 - 5445.0 | 0.0393 (43) | | | | 0.0391 | 0.0362(414) |
| 4763.8 - 4518.9 | 0.0332 (10) | | | | 0.0332 (15) | 0.0333 (23) |
| 4478.4 - 4145.2 | 0.0786 (6) | 0.0781 (1) | | | 0.0786 (5) | 0.0785 (11) |
| 4145.2 - 3988.1 | 0.0685 (10) | 0.0679 (4) | | | 0.0683 (5) | 0.0684 (15) |
| 3988.1 - 3748.9 | 0.0441 (2) | 0.0442 (1) | | | 0.0442 (1) | 0.0441 (3) |
| 3748.9 - 3694.9 | 0.00632 (5) | 0.00632 (6) | | | 0.00632 (3) | 0.00633 (8) |
| 3694.9 - 3608.2 | 0.0138 (2) | 0.0138 (1) | | | 0.0138 (1) | 0.0139 (2) |
| 3608.2 - 3569.1 | 0.00378 (11) | 0.0385 (0) | | | 0.00385 (13) | 0.00382 (14) |
| 3569.1 - 3531.5 | 0.00330 (2) | 0.00336 (0) | | | 0.00334 (3) | 0.00330 (2) |
| 3531.5 - 3503.1 | 0.00204 (1) | 0.00203 (0) | | | 0.00207 (2) | 0.00205 (2) |
| 3484.3 - 3418.7 | 0.00572 (2) | 0.00569 (6) | | | 0.00570 (2) | 0.00572 (4) |
| 3418.7 - 3357.0 | 0.00592 (1) | 0.00590 (0) | | | 0.00591 (2) | 0.00592 (3) |
| 3150.1 - 2770.2 | 2.74 (2) | 2.77 (2) | | 2.83 (31) | 2.77 (1) | 2.73 (1) |
| 2759.6 - 2679.1 | 0.0412 (3) | 0.0415 (4) | | | 0.0412 (1) | 0.0411 (6) |
| 2679.1 - 2563.4 | 0.0351(1) | 0.0350 (0) | | | 0.0352 (1) | 0.0351 (1) |
| 2563.4 - 2131.0 | 0.1258 (3) | 0.1240 (3) | | | 0.1257 (2) | 0.1257 (6) |
| 2131.4 - 2095.8 | 0.00429 (1) | 0.00400 (6) | | | 0.00418 (3) | 0.00428 (3) |
| 2095.8 - 2049.5 | 0.00570 (1) | 0.00525 (13) | | | 0.00551 (3) | 0.00568 (5) |
| 1977.6 - 1910.1 | 0.1271 (6) | 0.1262 (10) | 0.1268 | 0.1273 (15) | 0.1269 (5) | 0.1270 |
| 1910.1 - 1836.9 | 0.1321 (7) | 0.1307 (9) | 0.1318 | 0.1316 (15) | 0.1322 (9) | 0.1320 |
| 1836.9 - 1754.9 | 0.1336 (0) | 0.1321 (2) | 0.1334 | 0.1337 (8) | 0.1332 (6) | 0.1336 |
| 1754.9 - 1712.5 | 0.0526 (1) | 0.0526 (5) | 0.0521 | 0.0522 (7) | 0.0525 (4) | 0.0527 |
| 1713.0 - 1686.4 | 0.0177 (1) | 0.0172 (1) | 0.0178 | 0.0178 (4) | 0.0177 (5) | 0.0177 |
| 1686.4 - 1668.1 | 0.0137 (1) | 0.0134 (1) | 0.0138 | 0.0137 (3) | 0.0138 (1) | 0.0138 |
| 1650.3 - 1555.3 | | 0.533 (7) | | 0.531 (11) | 0.535 (4) | 0.535 (4) |
| 1531.7 - 1516.3 | | 0.1004 (6) | | 0.1033 (52 | 0.1010(3) | 0.0979 (13) |
| 1513.3 - 1400.1 | | 2.03 (1) | | 2.03 (5) | 2.04 (1) | 2,02 (2) |
| 1400.5 - 1338.4 | | 0.294 (1) | | 0.292 (1) | 0.290 (2) | 0.291 (4) |
| 1265.6 - 821.0 | 2.08 (6) | 2.08 (9) | | 2.11 (12) | 2.07 (38) | 2.09 (37) |
| 790.7 - 775.7 | 0.0826 (4) | 0.0815 (4) | | 0.0820 (3) | 0.0826 (4) | 0.0828 (22) |
| 769.9 - 710.2 | | | | 7.33 (58) | | |
| 710.2 - 660.0 | 2.70 (2) | | | 2.99 (17) | 2.72 (8) | 2.72 (3) |
| 640.2 - 606.5 | 0.0580 (4) | 0.0569 (20) | | , | 0.0578 (5) | 0.0578 (18) |
| 552.0 - 498.5 | 0.0947 (16) | 0.0942 (26) | | | 0.0941 (11) | |
| 490.3 - 440.2 | 1.65 (3) | 1.84 (10) | | 1.88 (18) | 1.73 (5) | 1.65 (9) |

a - The unit of area is cm⁻¹. The numbers in parentheses are the 95% confidence limits in the last digit. In some cases only one spectrum was available, so no 95% confidence limit could be calculated, and in others no spectrum was available from that spectroscopist for that region.

Table 2.4 - Spectroscopist average absorption index peak heights, $k_{\rm max}$. ^a

| \tilde{v} (cm ⁻¹) | CDK | A | В | c | VB | YA |
|---------------------------------|---------------|----------------|---|--------------|---------------|----------------|
| 5949.9 | 0.000179 (9) | | | | 0.900181 | 0.000178 (1) |
| 4667.0 | 0.000258 (7) | | | | 0.000262 | 0.000255 (15) |
| 4637.1 | 0.000226 (6) | | | | 0.000230 (10) | 0.000224 (14) |
| 4612.1 | 0.000342 (9) | | | | 0.000347 (13) | 0.000339 (19) |
| 4573.6 | 0.000206 (4) | | | | 0.000208 (7) | 0.000204 (9) |
| 4388.7 | 0.000304 (3) | 0.000301(1) | | | 0.000304 (3) | 0.000305 (6) |
| 4311.0 | 0.000348 (5) | 0.000345 (2) | | | 0.000349 (3) | 0.000347 (8) |
| 4244.3 | 0.000341 (2) | 0.000340 (4) | | | 0.000340 (1) | 0.000340 (7) |
| 4186.3 | 0.000175 (1) | 0.000173 (1) | | | 0.000174 (1) | 0.000175 (1) |
| 4161.3 | 0.000162 (0) | 0.000161 (2) | | | 0.000163 (1) | 0.000162 (1) |
| 4132.3 | 0.000174 (0) | 0.000175 (1) | | | 0.000174 (1) | 0.000174 (1) |
| 4056.6 | 0.000829 (21) | 0.000805 (13) | | | 0.000814 (9) | 0.000830 (28) |
| 4036.3 | 0.000978 (27) | 0.000974 (11) | | | 0.000977 (20) | 0.000975 (42) |
| 3980.3 | 0.000216(1) | 0.000215 (1) | | | 0.000218 (1) | 0.000217 (1) |
| 3951.2 | 0.000212 (1) | 0.000210 (1) | | | 0.000213 (1) | 0.000212 (2) |
| 3923.8 | 0.000215 (1) | 0.000216 (0) | | | 0.000217 (1) | 0.000215 (0) |
| 3909.9 | 0.000215 (0) | 0.000214 (2) | | | 0.000215 (2) | 0.000215 (0) |
| 3869.7 | 0.000224 (3) | 0.000223 (4) | | | 0.000224 (2) | 0.000224 (4) |
| 3847.2 | 0.000251 (3) | 0.000253 (3) | | | 0.000255 (2) | 0.000251 (5) |
| 3812.2 | 0.000176 (1) | 0.000174 (5) | | | 0.000176 (1) | 0.000175 (3) |
| 3786.3 | 0.000179 (1) | 0.000177 (5) | | | 0.000179 (1) | 0.000179 (2) |
| 3763.9 | 0.000108 (0) | 0.000107 (78) | | | 0.000109 (1) | 0.000108 (1) |
| 3724.1 | 0.000140 (0) | 0.000139 (0) | | | 0.000141 (1) | 0.000141 (0) |
| 3707.0 | 0.000139 (2) | 0.000138 (2) | | | 0.000138 (1) | 0.000139 (3) |
| 3675.0 | 0.000148 (6) | 0.000144 (3) | | | 0.000144 (5) | 0.000150 (8) |
| 3649.0 | 0.000269 (2) | 0.000270 (4) | | | 0.000270 (2) | 0.000269 (3) |
| 3624.0 | 0.000161 (1) | 0.000162 (1) | | | 0.000163 (1) | 0.000161 (1) |
| 3584.6 | 0.000126 (4) | 0.000126 (1) | | | 0.000126 (20) | 0.000127 (5) |
| 3549.8 | 0.000108 (1) | 0.000108 (1) | | | 0.000109(1) | 0.000108 (1) |
| 3519.3 | 0.0000840 (1) | 0.0000845 (30) | | | 0.0000851 (10 | 0.0000842 (10) |
| 3439.7 | 0.000127 (1) | 0.000127 (1) | | | 0.000126 (0) | 0.000127 (2) |
| 3385.3 | 0.000109 (1) | 0.000109 (1) | | | 0.000109 (0) | 0.000109 (1) |
| 3167.5 | 0.000481 (2) | 0.000481 (8) | | | 0.000487 (1) | 0.000480 (2) |
| 3104.1 | 0.00364 (2) | 0.00365 (4) | | 0.00371 (4) | 0.00369 (6) | 0.00364 (4) |
| 3086.4 | 0.0115 (1) | 0.0117 (3) | | 0.0117 (1) | 0.0118 (1) | 0.0115 (1) |
| 3062.0 | 0.0133 (1) | 0.0137 (2) | | 0.0139 (2) | 0.0137 (1) | 0.0132 (0) |
| 3026.9 | 0.0305 (6) | 0.0309 (8) | | 0.0312 (3) | 0.0308 (5) | 0.0301 (0) |
| 2979.1 | 0.00821 (4) | 0.00826 (5) | | 0.00843 (13) | 0.00830 (2) | 0.00820 (3) |
| 2947.5 | 0.0101 (1) | 0.0101 (1) | | 0.0103 (2) | 0.0102 (0) | 0.0101 (0) |

Table 2.4 - Continued

| v (cm ⁻¹) | CDK | A | В | С | VB | YA |
|-----------------------|--------------|---------------|---------|----------------|-----------------|---------------|
| 2919.8 | 0.0162 (1) | 0.0165 (2) | € | 0.0166 (2) | 0.0164 (1) | 0.0161 (1) |
| 2872.2 | 0.00788 (4) | 0.00802 (4) | (| 0.00813 (10) | 0.00797 (2) | 0.00790 (6) |
| 2734.1 | 0.00173 (5) | 0.00175 (8) | | | 0.00172 (1) | 0.00171 (8) |
| 2671.7 | 0.000230 (1) | 0.000227 (2) | | | 0.000230 (0) | 0.000229 (1) |
| 2631.9 | 0.000328 (1) | 0.000325 (2) | | | 0.000329 (0) | 0.000328 (1) |
| 2604.7 | 0.000422 (2) | 0.000409(1) | | | 0.000415 (1) | 0.000421 (3) |
| 2585.9 | 0.000772 (5) | 0.000762 (9) | | | 0.000770 (1) | 0.000770 (10) |
| 2540.4 | 0.000280 (1) | 0.000271 (1) | | | 0.000276 (1) | 0.000280 (2) |
| 2509.2 | 0.000257 (0) | 0.000252 (2) | | | 0.000256 (1) | 0.000256 (1) |
| 2496.7 | 0.000254 (0) | 0.000249(1) | | | 0.000254 (1) | 0.000254 (1) |
| 2465.4 | 0.000257 (0) | 0.000254 (0) | | | 0.000259 (1) | 0.000257 (1) |
| 2412.4 | 0.000530 (2) | 0.000519 (6) | | | 0.000529 (1) | 0.000529 (4) |
| 2389.0 | 0.000442 (1) | 0.000433 (7) | | | 0.000440 (2) | 0.000441 (2) |
| 2360.6 | 0.000708 (4) | 0.000691 (12) | | | 0.000708 (1) | 0.000707 (8) |
| 2335.3 | 0.000805 (5) | 0.000779 (16) | | | 0.000789 (5) | 0.000803 (10) |
| 2312.6 | 0.000623 (2) | 0.000619 (8) | | | 0.000620 (3) | 0.000623 (4) |
| 2280.7 | 0.000387 (2) | 0.000380 (1) | | | 0.000380 (0) | 0.000386 (4) |
| 2260.5 | 0.000440 (2) | 0.000426 (2) | | | 0.000439 (1) | 0.000439 (3) |
| 2237.3 | 0.000199 (0) | 0.000194 (2) | | | 0.000198 (2) | 0.000199(1) |
| 2207.5 | 0.000278 (2) | 0.000264 (4) | | | 0.000273 (1) | 0.000277 (3) |
| 2185.3 | 0.000281 (2) | 0.000272 (1) | | | 0.000277 (1) | 0.000281 (3) |
| 2163.7 | 0.000498 (2) | 0.000483 (4) | | | 0.000493 (2) | 0.000497 (4) |
| 2116.6 | 0.000152 (0) | 0.000142 (1) | | | 0.000148 (0) | 0.000151 (1) |
| 2068.5 | 0.000170 (0) | 0.000157 (1) | | | 0.000164 (1) | 0.000169(1) |
| 2032.0 | 0.000173 (0) | 0.000162 (3) | | | 0.000169 (1) | 0.000173 (1) |
| 2008.5 | | | | | 0.000358 (1) | 0.000363 (1) |
| 1991.1 | | | | | 0.000708 (1) | 0.000702 (3) |
| 1942.1 | | 0.00430 (4) | 0.00435 | 0.00435 (6) | 0.00436 (2) | 0.00436 |
| 1872.0 | , , | 0.00266 (1) | 0.00268 | 0.00268 (3) | 0.00269 (3) | 0.00267 |
| 1857.6 | • • | 0.00423 (3) | 0.00430 | 0.00429 (5) | 0.00433 (1) | 0.00434 |
| 1802.6 | | 0.00381 (0) | 0.00387 | 0.00386 (3) | 0.00388 (3) | 0.00389 |
| 1778.5 | • • | 0.00122 (1) | 0.00124 | 0.00123 (1) | 0.00123 (1) | 0.00123 |
| 1735.6 | | 0.00221 (1) | 0.00220 | 0.00221 (4) | 0.00222 (1) | 0.00222 |
| 1696.8 | | | 0.00082 | 0.000832 (100 | o) 0.000823 (24 | 0.000833 |
| 1676.7 | - | | 0.00093 | 5 0.000925 (3) | 0.000939 (4) | 0.000946 |
| 1623.1 | - | • | | 0.00428 (8) | 0.00429 (5) | 0.00434 (7) |
| 1604. | • • | 0.0301 (41) | | 0.0302 (10) | 0.0318 (3) | 0.0321 (9) |
| 1586.7 | | | | 0.00496 (3) | 0.00491 (4) | 0.00496 (2) |
| 1572. | | | | 0.00518 (3) | 0.00519 (5) | 0.00521 (2) |

Table 2.4 - Continued

| ~ (cm ⁻¹) | CDK | A | В | С | VB | YA |
|-----------------------|--------------|--------------|-------------|--------------|--------------|---------------|
| 1550.3 | 0.00341 (4) | 0.00345 (3) | | 0.00347 (82) | 0.00359 (4) | 0.00339 (20) |
| 1523.6 | 0.00865 (5) | 0.00864 (8) | | 0.00898 (58) | 0.00879 (3) | 0.00864 (13) |
| 1495.6 | 0.118 (1) | 0.112 (4) | | 0.112 (15) | 0.114 (2) | 0.119 (1) |
| 1460.2 | 0.0276 (2) | 0.0277 (1) | | 0.0297 (27) | 0.0278 (1) | 0.0275 (0) |
| 1378.9 | 0.0134 (0) | 0.0127 (1) | | 0.0127 (0) | 0.0128 (3) | 0.0132 (1) |
| 1332.0 | 0.00206 (2) | 0.00203 (2) | | 0.00205 (0) | 0.00209 (4) | 0.00208 (17) |
| 1312.7 | 0.00219 (2) | 0.00217(1) | | 0.00219 (0) | 0.00223 (3) | 0.00221 (4) |
| 1277.6 | 0.00116 (4) | 0.00112 (1) | | 0.00114 (1) | 0.00118 (3) | 0.00121 (60) |
| 1248.7 | 0.00181 (2) | 0.00181 (0) | | 0.00182 (1) | 0.00182 (2) | 0.00185 (16) |
| 1210.1 | 0.00452 (3) | 0.00427 (3) | | 0.00434 (2) | 0.00435 (12) | 0.00451 (11) |
| 1178.6 | 0.01002 (6) | 0.00930 (8) | | 0.00949 (2) | 0.00959 (29) | 0.01006 (25) |
| 1155.9 | 0.00410 (4) | 0.00407 (3) | | 0.00417 (2) | 0.00405 (10) | 0.00413 (4) |
| 1106.5 | 0.00672 (2) | 0.00673 (4) | | 0.00681 (2) | 0.00665 (15) | 0.00674 (6) |
| 1081.4 | 0.0306 (2) | 0.0295 (3) | | 0.0298 (1) | 0.0302 (8) | 0.0308 (11) |
| 1041.4 | 0.0151 (1) | 0.0150 (1) | | 0.0153 (0) | 0.0149 (5) | 0.0151 (3) |
| 1002.3 | 0.00446 (3) | 0.00459 (7) | | 0.00460 (2) | 0.00437 (9) | 0.00448 (13) |
| 980.7 | 0.00420 (3) | 0.00439 (6) | | 0.00438 (3) | 0.00417 (6) | 0.00423 (18) |
| 966.4 | 0.00393 (2) | 0.00414 (6) | | 0.00408 (5) | 0.00391 (5) | 0.00395 (31) |
| 929.6 | 0.00292 (1) | 0.00303 (3) | | 0.00297 (2) | 0.00292 (3) | 0.00291 (8) |
| 895.4 | 0.00939 (3) | 0.00880 (8) | | 0.00916 (2) | 0.00918 (23) | 0.00943 (1) |
| 872.9 | 0.00235 (2) | 0.00196 (14) | | 0.00233 (4) | 0.00240 (4) | 0.00235 (7) |
| 842.7 | 0.00378 (3) | 0.00336 (11) | | 0.00367 (4) | 0.00380 (4) | 0.00377 (22) |
| 785.6 | 0.00867 (5) | 0.00809 (10) | | 0.00811 (2) | 0.00838 (12) | 0.00873 (17) |
| 728.9 | | | | 0.719 (60) | | |
| 694.5 | 0.354 (10) | | | 0.384 (15) | 0.360 (12) | 0.361 (13) |
| 622.0 | 0.00336 (5) | 0.00314 (13) | | | 0.00328 (4) | 0.00334 (30) |
| 537.8 | 0.00132 (2) | 0.00130 (7) | | | 0.00130 (3) | 0.00132 (13) |
| 521.0 | 0.00748 (34) | 0.00695 (34) | | | 0.00681 (3) | 0.00748 (221) |
| 464.2 | 0.288 (32) | 0.356 (370) | | 0.289 (41) | 0.284 (17) | 0.288 (112) |

a - The numbers in parentheses are the 95% confidence limits in the last digit. In some cases only one spectrum was available, so no 95% confidence limit could be calculated, and in others no spectrum was available from that group for that region.

absorption index, $k(\tilde{v})$, spectrum. For each spectroscopist the average $k(\tilde{v})$ spectrum was calculated along with the average area and peak height for each band. The average

areas and peak heights obtained by the different spectroscopists are tabulated in Tables 2.3 and 2.4, respectively. Table 2.4 includes the $k(\tilde{v})$ peak wavenumbers which were determined, with the peak heights, by fitting the top three points of the band to a parabola and determining the maximum of the parabola.

The areas and peak heights from the different spectroscopists in Tables 2.3 and 2.4 are in excellent agreement. The agreement is not always within the precision of the data of each spectroscopist, which suggests that systematic errors or inadequate statistics may still affect the results by about 2% in some regions. The quality of the agreement is shown pictorially in Figure 2.1 for the regions 3150 to 2770 cm⁻¹, 1837 to 1755 cm⁻¹, and 1650 to 1555 cm⁻¹. In Fig. 2.1, the filled symbols show the average area for each spectroscopist, and the error bars show the 95% confidence limits.

For each region, the average spectra from the different spectroscopists were themselves averaged to yield a weighted average $k(\widetilde{\nu})$ spectrum, with the weighting factor for each spectroscopist being the number of spectra which contributed to the average (Table 2.2). To check the influence of the fact that the three spectroscopists in this laboratory, CDK, VB and YA, used the same instrument and respected spectra than the other collaborators, an unweighted average $k(\widetilde{\nu})$ spectrum was also calculated. For each region, the overall average area and the overall average peak neights were measured from both the weighted and the unweighted average $k(\widetilde{\nu})$ spectra.

The overall average areas are presented in Table 2.5. Because the integration ranges have different widths for different bands, Table 2.5 includes the height of the strongest peak in the region to indicate the prominence of the absorption. For each

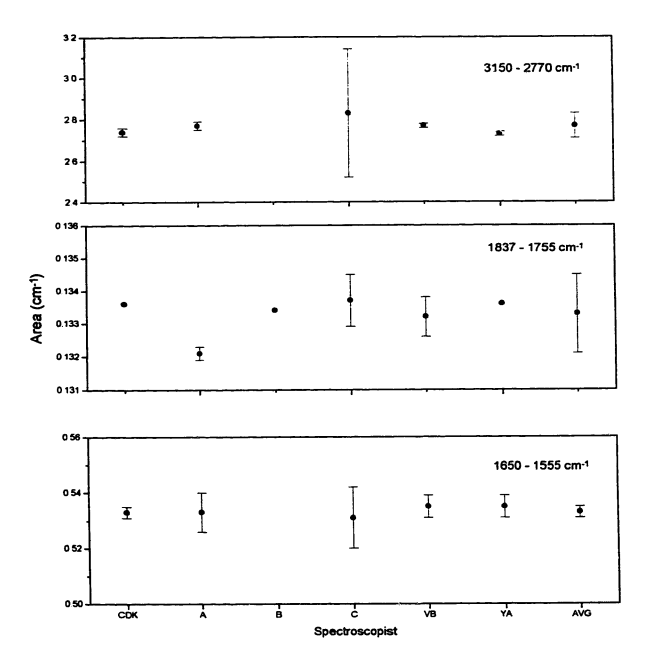


Figure 2.1 - Average areas and 95% confidence limits (vertical error bars) under the absorption index spectra in the regions 3150 to 2770 cm⁻¹ (upper), 1837 to 1735 cm⁻¹ (middle), and 1650 to 1555 cm⁻¹ (bottom) for the six spectroscopists. The values are taken from Table 2.3. Also shown for each region above the label "AVG" is the unweighted average area from Table 2.5 with the maximum deviation from it indicated by a vertical error bar.

Table 2.5 - Overall average area under the absorption index bands.

| Region (cm ⁻¹) | k _{max} ^a | Weighted Average Area ⁵ | Unweighted Average Area ^b | Maximum Deviation ^{b.c} | Anchor point Uncertainty b.d | % Estimated accuracy |
|------------------------------|-------------------------------|--|--|-------------------------------------|---------------------------------|----------------------|
| 6307.1 - 5445.0 | 0.000179 | 0.0382 | 0.0382 | ± 0.0020 | ± 0.0007 | 7.1 |
| 4763.8 - 4518.9 | 0.000344 | 0.0332 | 0.0332 | ± 0.0001 | ± 0.0001 | 0.6 |
| 4478.4 - 4145.2 | 0.000347 | 0.0785 | 0.0784 | ± 0.0003 | ± 0.0001 | 0.5 |
| 4145.2 - 3988.1 | 0.000976 | 0.0683 | 0.0683 | ± 0.0004 | ± 0.0004 | 1.2 |
| 3988.1 - 3748.9 | 0.000251 | 0.0441 | 0.0441 | ± 0.0001 | ± 0.0006 | 1.6 |
| 3748.9 - 3694.9 | 0.000140 | 0.0632 | 0.00632 | ± 0.00001 | ± 0.00004 | 0.8 |
| 3694.9 - 3608.2 | 0.000269 | 0.0138 | 0.0138 | ± 0.0001 | ± 0.00007 | 1.2 |
| 3608.2 - 3569.1 | 0.000127 | 0.00382 | 0.00383 | ± 0.00005 | ± 0.00003 | 2.1 |
| 3569.1 - 3531.5 | 0.000108 | 0.00332 | 0.00333 | ± 0.00003 | ± 0.00003 | 1.8 |
| 3531.5 - 3503.1 | 0.000084 | 0.00205 | 0.00205 | ± 0.00002 | ± 0.00002 | 2.0 |
| 3484.3 - 3418.7 | 0.000126 | 0.00571 | 0.00571 | ± 0.00002 | ± 0.00005 | 1.2 |
| 3418.7 - 3357.0 | 0.000109 | 0.00592 | 0.00591 | ± 0.00001 | ± 0.00005 | 1.0 |
| 3150.1 - 2770.2 | 0.0308 | 2.76 | 2.77 | ± 0.06 | ± 0.0006 | 2.2 |
| 2759.6 - 2679.1 | 0.00172 | 0.0413 | 0.0413 | ± 0.0002 | ± 0.0001 | 0.7 |
| 2679.1 - 2563.4 | 0.000768 | 0.035! | 0.0351 | ± 0.0001 | ± 0.0001 | 0.6 |
| 2563.4 - 2131.0 | 0.000793 | 0.1255 | 0.1253 | ± 0.0013 | ± 0.0002 | 1.2 |
| 2131.0 - 2095.8 | 0.000148 | 0.00423 | 0.00419 | ± 0.00019 | ± 0.00003 | 5.3 |
| 2095.8 - 2049.5 | 0.000165 | 0.00559 | 0.00554 | ± 0.00029 | ± 0.00003 | 5.8 |
| 1977.6 - 1910.1 | 0.00434 | 0.1269 | 0.1269 | ± 0.0007 | ± 0.0001 | 0.6 |
| 1910.1 - 1836.9 | 0.00429 | 0.1317 | 0.1317 | ± 0.0010 | ± 0.0001 | 0.8 |
| 1836.9 - 1754.9 | 0.00386 | 0.1332 | 0.1333 | ± 0.0012 | ± 0.0001 | 1.0 |
| 1754.9 - 1712.5 | 0.00221 | 0.0525 | 0.0524 | ± 0.0003 | ± 0.0001 | 0.8 |
| 1713.0 - 1686.4 | 0.000820 | 0.0176 | 0.0176 | ± 0.0004 | ± 0.00006 | 2.6 |
| 168 6 .4 - 1668.1 | 0.000931 | 0.0137 | 0.0137 | ± 0.0003 | ± 0.00004 | 2.5 |
| 1650.3 - 1555.3 | 0.0309 | 0.534 | 0.533 | ± 0.002 | ± 0.001 | 0.6 |
| 1531.2 - 1516.3 | | 0.01006 | 0.1002 | ± 0.0031 | ± 0.0005 | 3.6 |
| 1513.3 - 1400.1 | | 2.03 | 2.03 | ± 0.01 | ± 0.004 | 0.7 |
| 1400.5 - 1338.4 | | 0.292 | 0.292 | ± 0.002 | ± 0.0001 | 0.7 |
| 1265.6 - 821.0 | 0.0302 | 2.08 | 2.09 | ± 0.02 | ± 0.005 | 1.2 |

Table 2.5 - Continued

| Region (cm ⁻¹) | k _{max} * | Weighted Average Area ^b | Unweighted Average Area ^b | Maximum Deviation ^{b,c} | Anchor point Uncertainty ^{b,d} | % Estimated accuracy |
|----------------------------|--------------------|--|--|-------------------------------------|---|----------------------|
| 790.7 - 775.2 | 0.00837 | 0.0822 | 0.0823 | ± 0.0008 | ± 0.0002 | 1.2 |
| 769.9 - 710.2 | 0.719 | 7.33 | 7.33 | ± 0.58 | ± 0.006 | 8.0 |
| 710.2 - 660.0 | 0.365 | 2.75 | 2.78 | ± 0.21 | ± 0.005 | 7.7 |
| 640.2 - 606.5 | 0.00328 | 0.0577 | 0.0576 | ± 0.0007 | ± 0.0032 | 6.8 |
| 552.0 - 498.5 | 0.00716 | 0.0945 | 0.0945 | ± 0.0004 | ± 0.0005 | 1.0 |
| 490.3 - 440.2 | 0.295 | 1.73 | 1.75 | ± 0.13 | ± 0.0005 | 7.3 |

- a Height of the strongest peak in the region.
- b The unit of area is cm⁻¹.
- c The maximum deviation of the average of any one spectroscopist from the unweighted average, except for the region 769.9 710.2 cm⁻¹. In this case the maximum deviation is the 95% confidence limit of the area under the average spectrum for group C, the only spectroscopist who had spectra in this region.
- d The anchor point uncertainty is the integration range multiplied by the average of the uncertainties in $k(\tilde{\nu})$ (Table 2.1) at the two anchor points used for that range.
- e The % estimated accuracy is the sum of the maximum deviation and the anchor point uncertainty as a percentage of the unweighted average area.

band group, Table 2.5 also gives the maximum deviation of the average of any one spectroscopist from the unweighted average, which shows the agreement between the different spectroscopists. Table 2.5 includes the estimated accuracy of the overall unweighted average area of each band, which is discussed later. Again, a good sense of the quality of these results can be obtained from Fig. 2.1. Each box of Fig. 2.1 includes, above the label 'AVG', the unweighted average area and the maximum deviation.

The average agreement over the 35 band regions in Table 2.5 is $\pm 1.8\%$. For the

11 regions between 3150 and 775 cm⁻¹ with k_{max} between 0.002 and 0.112 the agreement between the areas is always better than $\pm 3.1\%$, and averages $\pm 1.1\%$. Such agreement between spectra measured in different laboratories on instruments by different manufacturers is very encouraging.

The agreement between the areas (Table 2.5) is only 5 to 8% for the very weak absorption between 6307 and 5445 cm⁻¹ and for the very strong absorption below 770 cm⁻¹. The absorption between 770 and 710 cm⁻¹ is so strong that cells with path lengths of 8 µm or less are needed to measure it accurately. There was only one spectroscopist who had such a cell and there were only two experimental absorbance spectra of suitable intensity to be measured accurately in this region. Additional measurements are needed to improve the intensities below 770 cm⁻¹.

The weighted and unweighted average areas in Table 2.5 agree to within 0.3% on average and they never disagree by more than 2.3%. This shows that the larger number of spectra from this laborate y did not unduly influence the average areas.

For each peak in the $k(\tilde{\nu})$ spectrum, the overall weighted and unweighted average peak heights are given in Table 2.6 along with the maximum deviation from the unweighted average of the averages of the different spectroscopists. Also given for each peak are the % estimated accuracy, which is discussed later, the previous value published from this laboratory⁵ and the calibrated result of Jones and co-workers⁷. The weighted and unweighted averages agree to within 0.4% on average. Few of them disagree by more than 1% and the largest disagreement is 2.2% at 872.9 cm⁻¹. Thus, the

Table 2.6 - Overall average peak heights in the absorption index spectra.

| ~ (cm ⁻¹) | Unweighted Average ^a | % Estimated Accuracy ^b | Weighted Average | Ref 5° | Ref 7 ^d |
|-----------------------|------------------------------------|-----------------------------------|---------------------|--------------|--------------------|
| 5949.9 | 0.000179 (2) | 1.6 | 0.000179 | | |
| 4667.0 | 0.000259 (4) | 1.8 | 0.000258 | | |
| 4637.1 | 0.000228 (4) | 2.0 | 0.000226 | | |
| 4612.1 | 0.000344 (5) | 1.6 | 0.000342 | | |
| 4573.6 | 0.000206 (2) | 1.2 | 0.000206 | | |
| 4388.7 | 0.000303 (2) | 0.8 | 0.000304 | | |
| 4311.0 | 0.000347 (2) | 0.7 | 0.000348 | | |
| 4244.3 | 0.000340(1) | 0.4 | 0.000340 | | |
| 4186.3 | 0.000175 (2) | 1.4 | 0.000175 | 0.000174 (5) | 0.000171 (12) |
| 4161.3 | 0.000162 (1) | 0.9 | 0.000162 | 0.000162 (5) | 0.000156 (11) |
| 4132.3 | 0.000174 (1) | 1.7 | 0.000174 | 0.000174 (5) | 0.000174 (12) |
| 4056.6 | 0.000812 (18) | 2.5 | 0.000823 | 0.000815 (8) | 0.000817 (31) |
| 4036.3 | 0.000976 (2) | 0.4 | 0.000977 | 0.000975 (5) | 0.000980 (37) |
| 3980.3 | 0.000217 (2) | 1.8 | 0.000217 | 0.000221 (5) | 0.000216 (15) |
| 3951.2 | 0.000212 (2) | 1.9 | 0.000212 | 0.000216 (4) | 0.000214 (15) |
| 3923.8 | 0.000216(1) | 1.4 | 0.000216 | 0.000219 (4) | 0.000222 (16) |
| 3909.9 | 0.000215 (2) | 1.9 | 0.000215 | | |
| 3869.7 | 0.000224 (1) | 1.3 | 0.000224 | 0.000226 (5) | 0.000227 (16) |
| 3847.2 | 0.000251 (4) | 2.4 | 0.000252 | 0.000255 (5) | 0.000257 (18) |
| 3812.2 | 0.000175 (1) | 1.7 | 0.000176 | | |
| 3786.3 | 0.000178 (1) | 1.7 | 0.000179 | 0.000177 (4) | 0.000178 (13) |
| 3763.9 | 0.000109 (2) | 3.7 | 0.000108 | | 0.000111 (8) |
| 3724.1 | 0.000140 (1) | 1.3 | 0.000140 | 0.000138 (4) | 0.000142 (10) |
| 3707.0 | 0.000139 (1) | 1.3 | 0.000139 | 0.000136 (4) | 0.000144 (10) |
| 3675.0 | 0.000146 (4) | 3.2 | 0.000147 | | 0.000160 (12) |
| 3649.0 | 0.000269 (1) | 0.7 | 0.000269 | 0.000268 (4) | 0.000269 (19) |
| 3624.0 | 0.000162 (1) | 1.1 | 0.000162 | 0.000161(4) | 0.000165 (12) |
| 3584.6 | 0.000127 (1) | 1.4 | 0.000126 | 0.000120 (9) | 0.000127 (9) |
| 3549.8 | 0.000108 (1) | 1.7 | 0.000108 | 0.000108 (4) | 0.000122 (9) |
| 3519.3 | 0.0000843 (8) | 1.9 | 0.0000844 | 0.000084 (4) | 0.000093 (8) |
| 3439.7 | 0.000126 (1) | 1.4 | 0.000127 | | 0.000128 (10) |
| 3385.3 | 0.000109 (0) | 0.7 | 0.000109 | | 0.000122 (9) |
| 3167.5 | 0.000484 (3) | 0.8 | 0.000483 | | 0.000509 (22) |
| 3104.1 | 0.00366 (5) | 1.4 | 0.00367 | | 0.00357 (9) |
| 3086.4 | 0.0116 (2) | 1.7 | 0.0117 | 0.0119 (1) | 0.0112 (4) |
| 3062.0 | 0.0136 (4) | 2.9 | 0.0136 | 0.0137 (1) | 0.0134 (4) |
| 3026.9 | 0.0308 (7) | 2.3 | 0.0307 | 0.0313 (7) | 0.0324 (6) |
| 2979.1 | 0.00830 (13) | 1.6 | 0.00827 | | 0.00799 (22) |

Table 2.6 - Continued

| v (cm ⁻¹) | Unweighted Average ^a | % Estimated Accuracy ^b | Weighted Average | Ref 5° | Ref 7 ^d |
|-----------------------|------------------------------------|-----------------------------------|---------------------|--------------|--------------------|
| 2947.5 | 0.0102 (1) | 1.0 | 0.0102 | 0.0103 (1) | 0.00962 (30) |
| 2919.8 | 0.0164 (3) | 1.9 | 0.0164 | 0.0164(1) | 0.0161 (5) |
| 2872.2 | 0.00800 (13) | 1.7 | 0.00796 | 0.00803 (11) | 0.00770 (21) |
| 2734.1 | 0.00172 (3) | 1.7 | 0.00172 | 0.00171 (1) | 0.00171 (6) |
| 2671.7 | 0.000229 (2) | 1.3 | 0.000229 | 0.000226 (3) | 0.000229 (16) |
| 2631.9 | 0.000327 (2) | 0.9 | 0.000328 | 0.000325 (3) | 0.000319 (23) |
| 2604.7 | 0.000417 (8) | 2.1 | 0.000418 | 0.000410(3) | 0.000406 (30) |
| 2585.9 | 0.000768 (6) | 0.9 | 0.000770 | 0.000766 (2) | 0.000787 (59) |
| 2540.4 | 0.000277 (6) | 2.3 | 0.000278 | 0.000272 (4) | 0.000277 (20) |
| 2509.2 | 0.000255 (3) | 1.4 | 0.000256 | 0.000252 (3) | 0.000251 (18) |
| 2496.7 | 0.000253 (4) | 1.8 | 0.000253 | | 0.000255 (18) |
| 2465.4 | 0.000257 (3) | 1.4 | 0.000257 | | 0.000260 (18) |
| 2412.4 | 0.000526 (5) | 1.0 | 0.000528 | 0.000525 (3) | 0.000512 (39) |
| 2389.0 | 0.000439 (6) | 1.5 | 0.000440 | 0.000436 (3) | 0.000438 (32) |
| 2360.6 | 0.000703 (12) | 1.8 | 0.000706 | 0.000705 (3) | 0.000769 (56) |
| 2335.3 | 0.000793 (14) | 1.8 | 0.000797 | 0.000784 (4) | 0.000818 (60) |
| 2312.6 | 0.000621 (2) | 0.4 | 0.000622 | 0.000616 (2) | 0.000614 (49) |
| 2280.7 | 0.000382 (5) | 1.4 | 0.000384 | | 0.000374 (26) |
| 2260.5 | 0.000436 (10) | 2.4 | 0.000438 | 0.000436 (3) | 0.000428 (31) |
| 2237.3 | 0.000198 (4) | 2.3 | 0.000198 | | 0.000203 (15) |
| 2207.5 | 0.000273 (9) | 3.5 | 0.000275 | 0.000270 (3) | 0.000274 (20) |
| 2185.3 | 0.000277 (5) | 2.0 | 0.000279 | 0.000274 (3) | 0.000280 (20) |
| 2163.7 | 0.000492 (9) | 2.0 | 0.000495 | 0.000491 (3) | 0.000434 (35) |
| 2116.6 | 0.000148 (6) | 4.5 | 0.000150 | | 0.000158 (13) |
| 2068.5 | • • | 4.7 | 0.000167 | 0.000163 (4) | 0.000172 (14) |
| 2032.0 | • • | 4.6 | 0.000171 | | 0.000187 (15) |
| 2008.5 | • | 2.2 | 0.000360 | | |
| 1991.1 | | 1.4 | 0.000703 | 0.000710 (3) | 0.000678 (25) |
| 1942.1 | | 1.0 | 0.00435 | 0.00441 (5) | 0.00462 (11) |
| 1872.0 | | 0.4 | 0.00268 | 0.00268 (1) | 0.00270 (10) |
| 1857.6 | | 1.4 | 0.00430 | 0.00458 (15) | 0.00460 (11) |
| 1802.6 | | 1.3 | 0.00387 | 0.00390 (4) | 0.00390 (15) |
| 1778.5 | | 0.9 | 0.00123 | | 0.00123 (5) |
| 1735.6 | | 0.6 | 0.00221 | 0.00220 (2) | 0.00222 (8) |
| 1696.8 | | | 0.000824 | 0.000832 (4) | 0.000809 (30) |
| 1676.7 | | • | 0.000934 | | |
| 1623. | · | 1.5 | 0.00430 | | 0.00426 (10) |
| 1604. | | 3.9 | 0.0315 | 0.0320(1) | 0.0304 (10) |

Table 2.6 - Continued

| $\tilde{\nu}$ (cm ⁻¹) | Unweighted Average ^a | % Estimated Accuracy ^b | Weighted Average | Ref 5° | Ref 7 ^d | |
|-----------------------------------|------------------------------------|-----------------------------------|---------------------|--------------|--------------------|--|
| 1586.7 | 0.00494 (5) | 1.3 | 0.00494 | | | |
| 1572.1 | 0.00519 (2) | 0.6 | 0.00520 | | 0.00560 (14) | |
| 1550.3 | 0.00347 (12) | 4.4 | 0.00350 | | 0.00353 (13) | |
| 1523.6 | 0.00867 (31) | 3.9 | 0.00877 | | 0.00928 (22) | |
| 1495.6 | 0.115 (4) | 3.4 | 0.115 | 0.115 (3) | 0.116 (2) | |
| 1460.2 | 0.0278 (19) | 6.9 | 0.0281 | 0.0280 (2) | 0.0265 (9) | |
| 1378.9 | 0.0129 (5) | 4.0 | 0.0129 | 0.0133 (2) | 0.0137 (3) | |
| 1332.0 | 0.00206 (3) | 2.4 | 0.00206 | | 0.00217 (8) | |
| 1312.7 | 0.00220 (3) | 2.3 | 0.00219 | | 0.00230 (9) | |
| 1277.6 | 0.00116 (5) | 6.0 | 0.00115 | | 0.00118 (4) | |
| 1248.7 | 0.00182 (3) | 2.7 | 0.00182 | | 0.00181 (7) | |
| 1210.1 | 0.00439 (13) | 3.1 | 0.00448 | | 0.00446 (16) | |
| 1178.6 | 0.00967 (39) | 4.1 | 0.00963 | 0.0100 (3) | 0.0103 (3) | |
| 1155.9 | 0.00410 (7) | 1.8 | 0.00409 | | 0.00410 (15) | |
| 1106.5 | 0.00673 (8) | 1.3 | 0.00672 | | 0.00732 (18) | |
| 1081.4 | 0.0302 (7) | 2.3 | 0.0301 | 0.0311 (2) | 0.0305 (10) | |
| 1041.4 | 0.0151 (2) | 1.3 | 0.0151 | | 0.0158 (4) | |
| 1002.3 | 0.00450 (13) | 3.0 | 0.00451 | | 0.00442 (16) | |
| 980.7 | 0.00427 (12) | 2.9 | 0.00428 | | 0.00423 (16) | |
| 966.4 | 0.00400 (14) | 3.6 | 0.00402 | | | |
| 929.6 | 0.00295 (8) | 2.9 | 0.00296 | | 0.00301 (11) | |
| 895.4 | 0.00919 (39) | 4.4 | 0.00912 | 0.00968 (33) | 0.0100 (5) | |
| 872.9 | 0.00228 (32) | 14.7 | 0.00223 | | 0.00235 (9) | |
| 842.7 | 0.00368 (32) | 9.1 | 0.00363 | | 0.00385 (14) | |
| 785.6 | 0.00837 (36) | 4.5 | 0.00845 | | 0.00839 (31) | |
| 728.9 | 0.719 (60) | 8.3 | 0.719 | | 0.658 (16) | |
| 694.5 | 0.365 (19) | 5.2 | 0.362 | 0.342 (4) | 0.380 (6) | |
| 622.0 | 0.00328 (10) | 3.3 | 0.00329 | 0.00329 (1) | 0.00320 (12) | |
| 537.8 | 0.00131 (1) | 1.4 | 0.00131 | 0.00130 (0) | 0.00126(1) | |
| 521.0 | 0.00716 (35) | 5.0 | 0.00721 | 0.00683 (5) | 0.00667 (45) | |
| 464.2 | 0.295 (39) | 13.2 | 0.292 | | 0.120 (13) | |

- a In this column the number in parentheses is the maximum deviation from the unweighted average, except for the peak at 728.9 cm⁻¹. In this case the number is the 95% confidence limit in the average spectrum for C, the only spectroscopist who had spectra in this region.
- b The % estimated accuracy is the sum of the maximum deviation and the uncertainty due to the anchor points as a percentage of the unweighted average. The uncertainty due to the anchor points is the average of the uncertainties in $k(\tilde{v})$ (Table 2.1) for the two anchor points either side of the peak.
- c In this column the number in parentheses is the 90% confidence limit in the last digit⁵.
- ${\bf d}$ In this column the number in parentheses is the evaluated uncertainty in the value 7 .

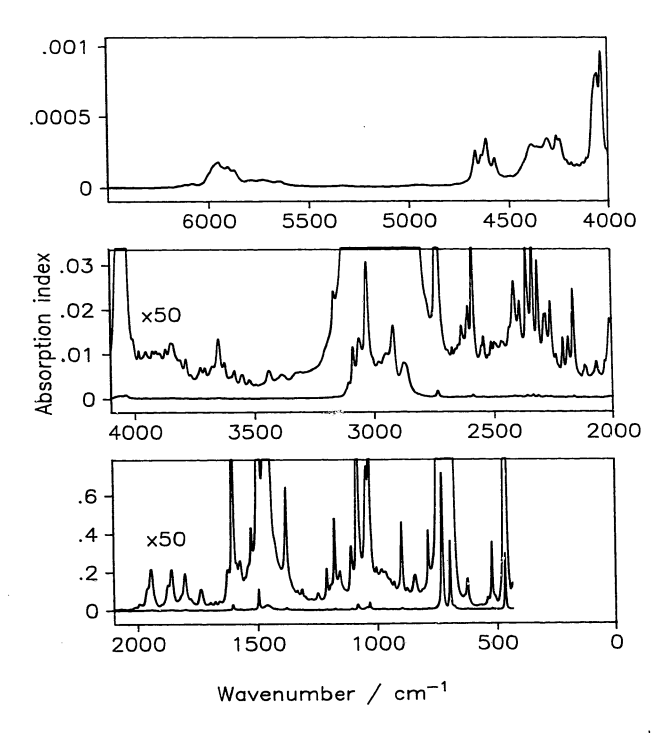


Figure 2.2 - Absorption index (imaginary refractive index), $k(\tilde{\nu})$, spectrum between 6500 and 435 cm⁻¹ of toluene at 25°C. The scale labels in the middle and bottom boxes are for the lower spectrum in the box; they must be divided by 50 for the upper spectrum in the box.

Table 2.7 - Absorption indices between 6500 and 435 cm⁻¹ of liquid toluene at 25°C.^{a,b}

| cm ⁻¹ | XF | YE | •1 | 1 | | 3 | 4 | | - 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 |
|---------------------|---------|------------------|--------------|--------------|--------------|-----------------------|---------------------|-----------------|--------------|--------------|----------------------|--------------|--------------|--------------|--------------|--------------|--------------|--------------|-----------------------|
| 6499 90 | 3 | -7 | 22 | 16 | 16 | 13 | 9 | 14 | 13 | 7 | 13 | 8 | 9 | 8 | 3 | 3 | 8 | 5 | 1 |
| 6368 76 | 3 | -7 | 5 | 1 | 1 | 1 | 2 | 4 | 3 | 1 | 2 | 1 | 1 | . 1 | 1 | 6 | 15 | 13 | 21 |
| 6237 63 6106 49 | 3 | .7 .7 | 38 194 | 28 304 | 28 238 | 30 281 | 34 305 | 40 252 | 51 205 | 1 89 | 42 208 | 63 267 | 89 334 | 141 439 | 141 517 | 150 819 | 171 979 | 209 1128 | 227 1366 |
| 5975 35 | 3 | .7 | 1513 | 1636 | 1723 | 1782 | 1721 | 1515 | 1439 | 1372 | 1357 | 1435 | 1404 | 1247 | 1181 | 1225 | 1143 | 966 | 776 |
| 5844 22 | 3 | -7 | 633 | 538 | 471 | 439 | 446 | -183 | 508 | 525 | 534 | 514 | 495 | 501 | 512 | 523 | 546 | 562 | 532 |
| 5713.08 | 3 | -7 | 502 | 485 | 443 | 416 | 393 | 379 | 394 | 424 | 439 | 436 | 418 | 350 | 280 | 246 | 227 | 193 | 165 |
| 5581 95 | 3 | -7 -7 | 172 84 | 147 81 | 117 81 | 138 81 | 121 83 | 108 85 | 126 92 | 117 90 | 108 96 | 99 107 | 113 112 | 104 111 | 87 121 | 91 128 | 83 138 | 82 144 | 81 140 |
| 5450 81 5319 68 | 3 | - <i>1</i> -7 | 118 | 120 | 121 | 98 | 100 | 96 | 84 | 81 | 77 | 74 | 79 | 39 | 85 | 75 | 74 | 73 | 69 |
| 5188 54 | 3 | -7 | 65 | 59 | 46 | 56 | 72 | 69 | 63 | 64 | 65 | 63 | 64 | 62 | 65 | 68 | 69 | 67 | 68 |
| 5057 40 | 3 | -7 | 71 | 71 | 80 | 85 | 87 | 92 | 100 | 111 | 127 | 147 | 159 | 175 | 178 | 177 | 174 | 170 | 162 |
| 4926 27 | 3 | -7 | 163 | 152 | 147 | 134 204 | 125 213 | 119 226 | 125 233 | 133 259 | 140 296 | 141 346 | 141 403 | 144 455 | 142 532 | 142 683 | 140 1027 | 137 | 138 |
| 4795 13 4686 17 | 3 | .7 .7 | 148 148 | 164 1148 | 186 1210 | 1278 | 1343 | 1409 | 1478 | 1546 | 1622 | 1696 | 1774 | 1863 | 1957 | 2056 | 2163 | 2268 | 2373 |
| 4666 89 | 2 | .7 | 2592 | 2350 | 1957 | 1714 | 1655 | 1740 | 1980 | 2224 | 2269 | 2230 | 2332 | 2572 | 2851 | 3187 | 3424 | 3275 | 2730 |
| 4601 32 | 2 | .7 | 2170 | 1818 | 1522 | 1524 | 1530 | 1677 | 1877 | .054 | 1947 | 1650 | 1437 | 1274 | 1106 | 978 | 899 | 846 | 830 |
| 4531 90 | 3 | -7 | 790 | 721 | 715 | 751 | 774 | 789 | 776 | 769 | 837 | 938 | 1055 | 1218 | 1411 | 1622 3261 | | 2067 2713 | 2453 2714 |
| 4400 76 4269 63 | 3 | -7 -7 | 2800 2983 | 2996 3668 | 3022 3376 | 2920 33 7 3 | 2849 3230 | 2847 2626 | 2847 2105 | 2817 1920 | 2809 | 2878 | 3115 | 3397 | 3449 | 3201 | 2999 | 2713 | 2714 |
| 4211 77 | 2 | -7 | 1931 | 1869 | 1765 | 1672 | 1610 | 1613 | 1708 | 1730 | 1623 | 1530 | 1488 | 1500 | 1575 | 1617 | 1599 | 1561 | 1512 |
| 4146.20 | 2 | -7 | 1497 | 1530 | 1618 | 1719 | 1741 | 1699 | 1683 | 1747 | 1879 | 2022 | 2062 | 2078 | 2180 | 2434 | 2885 | 3610 | 4674 |
| 4080 64 | 2 | -7 | 5813 | 6573 | 7023 | 7474 | 7897 | 7991 | 8093 | 7828 | 7181 | 7278 | 8272 | 9467 | 9415 | 7922 | 6015 | | 3489 |
| 4015 07 | 2 | -7 | 2992 | 2723 | 2699 | 2558 | 2293 | 2028 | 1860 | 1802 | 1916 | 2171 2044 | 1955 2143 | 1825 2098 | 1825 2026 | 1860 2033 | 1900 2049 | | 2090 1952 |
| 3949 50 3883 93 | 2 | -7 -7 | 2099 1859 | 1989 1832 | 1909 1930 | 1872 2158 | 1847 2214 | 1896 2080 | 2094 2016 | 2150 2111 | 2049 2332 | 2494 | 2509 | 2492 | 2419 | 2198 | 1994 | | 1795 |
| 3818.36 | 2 | -7 | 1728 | 1739 | 1742 | 1642 | 1487 | 1387 | 1406 | 1560 | 1766 | 1693 | 1420 | 1198 | 1077 | 1046 | 1086 | | 963 |
| 3752.80 | 2 | -7 | 911 | 894 | 903 | 944 | 1024 | 1110 | 1232 | 1381 | 1365 | 1233 | 1198 | 1306 | 1387 | 1255 | 1098 | 1064 | 1102 |
| 3687 23 | 2 | -7 | 1203 | 1305 | 1398 | 1459 | 1446 | 1-141 | 1525 | 1741 | 2118 | 2531 | 2688 | 2443 | 1982 | 1603 | 1436 | 1462 | 1597 |
| 3621 66 | 2 | -7 | 1570 | 1291 | 1044 | 947 | 941 | 959 | 979 1250 | 995 1225 | 1094 1192 | 1227 1158 | 1124 | 1093 | 1064 | 1036 | 1011 | 988 | 965 |
| 3586 47 3574 89 | -1 2 | -7 -7 | 1238 817 | 1247 745 | 1255 730 | 1263 783 | 1267 900 | 1263 1020 | 1073 | 1073 | 987 | 824 | 701 | 1093 | 675 | 738 | 829 | 819 | 728 |
| 3509 33 | 2 | -7 | 662 | 622 | 614 | 628 | 640 | 621 | 599 | 598 | 619 | 639 | 654 | 672 | 692 | 722 | 775 | 874 | 1039 |
| 3443 76 | 2 | -7 | !191 | 1261 | 1217 | 1108 | 983 | 887 | 839 | 827 | 839 | 865 | 897 | 925 | 957 | 1008 | 1058 | | |
| 3382 05 | 3 | -7 | 1079 | 1026 | 931 | 866 | 875 | 961 | 1063 | 1152 | 1167 | 1176 | 1201 | 1160 | 1171 | 1195 | 1224 | 1254 | 1307 |
| 3250 91 3179 56 | 3 | .7 -7 | 1345 3032 | 1405 3170 | 1453 3359 | 1547 3645 | 1692 4074 | 1892 4575 | 2194 4829 | 2392 4791 | 2654 4667 | 2931 4560 | 4510 | 4507 | 4586 | 4669 | 4767 | 4882 | 5025 |
| 3146 77 | i | -6 | 514 | 531 | 550 | 574 | 597 | 625 | 656 | 696 | 745 | 808 | 891 | 999 | 1131 | 1294 | | 1718 | 1911 |
| 3113 99 | i | -5 | 208 | 229 | 260 | 302 | 346 | 366 | 358 | 344 | 345 | 373 | 438 | 558 | 750 | 992 | 1152 | 1125 | 992 |
| 3079 28 | 2 | - 5 | 774 | 736 | 863 | 1134 | 1339 | 1342 | 1268 | 1172 | 1106 | 1134 | 1290 | 1700 | 2370 | 2952 | 3010 | | 1650 |
| 3013 71 | 2 2 | -5 -5 | 1234 | 1026 | 945 990 | 899 1002 | 80 7 1073 | 746 1224 | 737 1440 | 764 1621 | สว8 1 <i>5</i> 69 | 830 1270 | 822 949 | 803 724 | 788 · 593 | 827 528 | 906 514 | 958 539 | 995 593 |
| 20 18 14 2882 57 | 2 | 0 | 1017 6635 | 1007 7363 | 7906 | 7984 | 7856 | 7641 | 7130 | 6029 | 4674 | 3524 | 2710 | 2165 | 1778 | 1490 | 1283 | | 1070 |
| 281701 | 2 | -1 | 9537 | 9221 | 8845 | 7917 | 7032 | 6368 | 5771 | 5229 | 4868 | 4655 | 4476 | 4265 | 4051 | 3798 | 3646 | | 3684 |
| 2753 37 | 1 | -6 | 377 | 392 | 416 | 454 | 514 | ବେମ | 75.7 | 972 | 1273 | 1583 | 1723 | 1574 | 1267 | 977 | 761 | 612 | 510 |
| 2720 58 | 1 | -7 | 4394 | 3898 | 3536 | 3252 | 3021 | 28%2 | 75 | 2539 | 2429 | 2348 | 2273 | 2207 | 2163 1989 | 2148 2039 | 2153 2109 | | 2134 2205 |
| 2687 80 2655 02 | l l | -7 -7 | 2084 2206 | 2037 2197 | 2005 2215 | 1976 2273 | 1945 2341 | 19.1 2389 | :976 2397 | 2099 2386 | 2270 2408 | 2234 2507 | 2072 2734 | 1990 3095 | 3272 | 3161 | | 2872 | 2205 2779 |
| 2622.23 | i | -7 | 2716 | 2678 | 2677 | 2734 | 2889 | 3180 | 30°5 | 3619 | 3867 | 4164 | 3925 | 3479 | 3185 | 3110 | | 3675 | 4497 |
| 2589 45 | | -7 | 5770 | 7155 | 7658 | 6665 | 5213 | 4119 | 3408 | 2927 | 2564 | 2286 | 2088 | 1960 | 1876 | 1828 | 1821 | 1850 | 1905 |
| 2556 66 | | -7 | 1951 | 2012 | 2126 | 2273 | 2358 | 2354 | 2373 | 2522 | 2735 | 2710 | 2478 | 2255 | 2070 | 1921 | | 1788 | 1825 |
| 2523 88 | | -7 | 1901 | 1934 | 1933 | 1936 | 1978 | 2091 | 2286 | 2497 | 2533 | 2372 | 2236 | 2210 | 2281 | 2419 | | 2478 | 2442 2526 |
| 2491 10 2458 31 | l 1 | -7 -7 | 2445 2514 | 2437 2493 | 2399 2453 | 2350 2407 | 2305 2364 | 2280 2338 | 2271 2329 | 2270 2345 | 2279 2394 | 2301 2477 | 2347 2588 | 2421 2734 | 2513 2941 | 2564 3132 | | 2538 3180 | 2526 3213 |
| 2425 53 | | -, -7 | 3303 | 3460 | 3705 | 4044 | 4433 | 4804 | 5139 | 5258 | 5024 | 4634 | 4256 | 3942 | 3697 | 3533 | | 3517 | 3667 |
| 2392 74 | | -7 | 3894 | 4198 | 4387 | 4135 | 3702 | 3354 | 3084 | 2879 | 2759 | 2709 | 2705 | 2767 | 2911 | 3208 | | 4920 | 6521 |
| 2359 96 | | -7 | 6918 | 5941 | 5188 | 4755 | 4399 | 4068 | 3845 | 3814 | 4036 | 4541 | 5228 | 6106 | 7355 | 7887 | | 5063 | 4001 |
| 2327 18 | | -7 | 3393 | 3092 | 3009 | 3112 | 3436 | 4058 | 5016 | 6005 | 6061 | 5045 | 3977 | 3259 | 2827 | 2582 | | 2476 | 2577 |
| 2294 39 2261 61 | | -7 -7 | 2779 4263 | 3064 4305 | 3365 3879 | 3625 3352 | 3782 2910 | 3757 2562 | 3714 2291 | 3815 2084 | 3618 1939 | 3226 1849 | 2948 1823 | 2803 1867 | 2750 1949 | 2790 1965 | | 3289 1680 | 3 7 87 1563 |
| 2228 82 | | -7 | 1500 | 1469 | 1437 | 1392 | 1349 | ىدەدىد باد13 | 1357 | 1463 | 1660 | 1916 | 2352 | 2730 | 2372 | 1856 | | 1485 | 1507 |
| 2196 04 | | -7 | 1600 | 1761 | 1981 | 2211 | 2450 | 2711 | 2728 | 2414 | 2103 | 1915 | 1834 | 1833 | 1920 | 2155 | | 3477 | 4526 |
| 2163 26 | | -7 | 4894 | 4180 | 3232 | 2504 | 1998 | 1665 | 1459 | 1335 | 1260 | 1203 | 1154 | 1116 | 1090 | 1072 | | 1013 | 976 |
| 2130 47 | | -7 | 963 | 988 | 1070 | 1162 | 1185 | 1219 | 1348 | 1479 | 1437 | 1395 | 1400 | 1353 | 1249 | 1146 | | 1016 | 991 |
| 2097 69 | 1 | 7 | 987 | 987 | 984 | 989 | 1003 | 1027 | 1056 | 1083 | 1106 | 1134 | 1179 | 1243 | 1318 | 1402 | 1534 | 1644 | 1574 |

Table 2.7 - Continued

| | Table 2.7 - Continued. | | | | | | | | | | | | | | | | | | | |
|--|------------------------|-----|-----|------|------|------|------|-------|------|------|----------|-------|--------|------|------|------|--------|-------|--------|--------|
| 1935 1 | cm ⁻¹ | ΛZ |)E | 0 | 1 | 2 | 3 | 4 | • | | <u> </u> | | ٦ | 1.0 | 11 | 12 | 1.3 | 14 | 15 | 16 |
| 1999 1 7 7 788 243 578 645 687 682 687 682 684 646 | | | | | | | | | | | | | | | | | | | | |
| 1981 98 2 | | | | | | | | | | | | 2614 | 2888 | 3152 | 1414 | 1280 | 3547 | 3451 | 1302 | 1.1-14 |
| 1964 2 6 743 438 477 510 614 438 548 5 | | | | | | | | | | | | 26.10 | 1160 | 1371 | 1076 | 1007 | 1871 | 1 214 | 1.517 | 72.1 |
| 1898 18 2 6 1545 1896 1902 1014 1945 1946 1946 1946 1946 1946 1945 1545 1896 1946 1946 1946 1946 1947 1946 1 | | - | | | | | | | | | | | | | | | | | | |
| 1785 1886 1896 | | 2 | | | | | | | | | | | | | | | | | | |
| 1694 6 1 | | | | | | | | 930 | 831 | | 069 | | | | | | | | | |
| 166 187 | 1719 71 | 2 | -7 | 7348 | 0267 | 0084 | 0145 | 03001 | n974 | 8189 | | | | | | | | | | |
| 1.00 | | 1 | -7 | 7531 | | | | | | | 7203 | 8350 | | | 7080 | 7150 | 6714 | 0023 | いみれ | 75641 |
| 1988 1 | | | | | | | | | | | | | | | 1061 | 1181 | 1340 | 1557 | 1894 | 2325 |
| 1594 1 | | | | | | | | | | | | | | | | | | | | |
| 1555-90 1-6 3206 375 707 862 378 345 3 | | | | | | | | - | | | | | | | | | | | | |
| 1825-98 0 | | | | | | | | | | | | | | | | | | | | |
| 1490 S | | | | | | | | | | | | | | | | | | | • • | . • |
| 14150 2 | 1499 86 | 0 | -4 | 282 | 378 | 542 | 805 | 1081 | 1043 | 783 | 557 | 417 | 330 | | 234 | 209 | 185 | 165 | 152 | 144 |
| 1342 1 | 1480.58 | 2 | -5 | 1388 | 1537 | 1811 | 2180 | 2555 | 2768 | 2712 | 2430 | 2055 | 1706 | 1341 | 1011 | 838 | 70-4 | 718 | 653 | 584 |
| 1346 1 | | 2 | | | 4925 | | | | | | | | | | | | | | | |
| 131666 1 | | | | | | | | | | | | | | | | | | | | |
| 1283 1 | | | | | | | | | | | | | | | | | | | | |
| 1218 at 1 | | | | | | | | | | | | | | | | | | | | |
| 1218a | | - | | | | | | | | | | | | | | | | | | |
| 1869 1 | | | | | | | | | | | | | | | | | | | | |
| 1163 56 | | 0 | -6 | 3101 | 3295 | 3570 | 3959 | 4531 | 5402 | 6713 | 8436 | 9632 | 9066 | 7524 | 6117 | 5109 | 4441 | 4000 | 3709 | 3514 |
| 103.66 -5 559 485 | 1169 13 | 1 | -6 | 3292 | 3225 | 3269 | 3386 | 3567 | 3790 | 3992 | 4095 | 3815 | 3432 | 3113 | 2856 | 2658 | 2524 | 2423 | 2327 | 2230 |
| 1070 78 1 5 785 669 599 561 547 552 573 612 669 749 854 986 141 1344 1440 1506 1418 1038 96 0 5 1427 1471 | 1136 35 | 1 | -6 | 2159 | 2114 | 2099 | | | | | 2006 | | | | | | | | | |
| 1038 6 | | | | | | | | | | | | | | | | | | | | |
| 1002.57 0 6 7042 7342 6627 6007 5533 5170 4888 4-669 4488 4-541 4217 4121 4050 3997 3977 3707 370 3717 3733 3733 3742 3770 3788 3789 3788 3789 3781 3750 3781 3770 3733 3733 3742 3770 3788 3789 3788 3789 3781 3750 3781 3770 3770 3781 3781 37 | | | | | | | | | | | | | | | | | | | | |
| 1006 17 0 -6 4000 4056 4171 4170 4944 4407 4249 4100 3984 3891 3829 3713 3702 3774 3733 3742 3770 3784 3733 3742 3770 3784 3733 3742 3770 3888 32 3 -6 2668 2870 2938 2726 2365 2299 2224 2212 2225 2309 23181 2468 2560 2678 2869 2217 2274 2227 2333 2 -6 2012 1877 1885 2046 2410 2410 2415 2415 2410 2411 2418 2018 1985 2412 2177 2274 2227 2388 2331 2348 2426 2438 2540 2607 2700 2779 2904 3065 3259 3563 4027 2788 26 0 -6 4790 6098 7784 8339 7439 6354 5546 5036 4719 4516 4382 4298 4279 4240 4233 4283 4421 4594 4749 4896 5065 5275 5481 5648 5878 6196 6551 6024 7377 7374 6784 2376 2381 | | | | | | | | | | | | | | | | | | | | |
| 986 89 2 6 6 3946 4185 4222 3985 3915 3986 3940 3651 3388 3239 1173 3023 2781 2572 934 82 1 6 2563 2666 2870 2938 2726 2465 2299 2224 2212 2245 2309 2381 2488 264 2660 2678 2860 212 867 33 2 6 2012 1877 1885 2046 2410 3049 3613 3531 2945 2401 2041 1871 1839 1908 2042 2177 2274 2227 788 25 0 6 2199 2226 2254 2287 2331 2380 2426 2483 2540 2607 2700 2779 2904 3065 3259 1503 2482 771 87 0 6 4317 4373 4471 4594 4749 4896 5065 5275 5481 5654 5878 6196 6551 6024 7373 7946 8642 755 47 0 5 949 1048 1165 1310 1476 1651 1821 1991 2177 239 2647 2363 3319 3766 4313 15055 5991 739.08 0 4 722 887 1108 1408 1818 2371 3101 4023 5105 6214 7017 7116 6488 5421 4235 3182 2186 722.69 0 4 1747 1313 1007 790 633 519 435 371 321 282 249 222 202 189 182 183 191 706.30 0 4 207 233 273 326 3039 471 565 694 899 1298 1866 3130 2498 1821 191 689.91 0 5 8951 6555 4974 3925 3215 2724 2384 2154 2005 1919 1880 1877 1872 1830 1733 1575 1386 65712 0 6 2613 2048 1980 1863 1814 1705 1666 1574 1588 1495 1508 1433 1420 1370 1385 1311 1318 640 73 0 6 1301 1330 1335 1324 1459 1468 1492 1532 1559 1558 1556 1569 1617 1708 1850 2055 2335 62434 0 6 2681 3035 3256 3221 2943 2567 2202 1902 1557 1568 1433 1420 1370 1385 1311 1318 640 73 0 6 1301 1330 1335 1324 1459 1468 1493 4794 4796 4400 4460 4550 4619 4467 4793 3793 3720 3720 3724 3745 3786 3814 3805 4713 1385 1311 1318 640 73 0 6 1301 1330 1335 1424 1459 1468 1492 1532 1559 1558 1556 1569 1617 1708 1850 2055 2335 62343 0 6 73 881 1007 1155 1275 1303 1207 1474 1476 4793 4803 4835 4805 4715 4627 4573 1484 55877 0 7 44400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4716 4719 4714 4716 4793 4803 4835 4805 4715 4719 4714 4714 5787 4719 1471 1471 1471 1471 1471 1471 147 | | | | | | | | | | | | | | | | | | | | |
| 93.48 1 | | | | | | | | | | | | | | | | | | | | |
| 867 33 2 -6 2012 1877 1885 2046 2410 3049 3613 3531 2945 2401 2041 1871 1839 1908 2048 2139 2180 80465 0 -6 2199 2226 2254 2287 2331 2380 2426 2483 2540 2607 2700 2779 2904 3065 3259 3563 4027 788 263 0 -6 4790 6098 7784 8339 7439 6354 5546 5036 4719 4516 4382 4298 4279 4240 4233 4283 4421 771 87 0 -6 4317 4373 4471 4594 4749 4896 5065 5275 5481 5654 5878 6196 6551 6924 7373 7946 8642 755.47 0 -5 949 1048 1165 1310 1476 1651 1821 1991 2177 2393 2647 2953 3319 3766 4331 5055 5991 73998 0 -4 722 887 1108 1408 1818 2371 3101 4023 5105 6214 7017 7116 6488 5421 4234 4234 2216 2456 2456 2456 2456 2456 2456 2456 245 | 934 82 | 1 | -6 | 2563 | 2666 | 2870 | 2938 | 2726 | 2465 | 2299 | 2224 | 2212 | 2245 | 2309 | 2381 | 2468 | 2560 | 2678 | 2869 | 3213 |
| 804 65 0 -6 2199 2226 2254 2287 2331 2380 2426 2483 2540 2607 2700 2779 2904 3065 3259 3563 4027 788 26 0 -6 4790 6098 7784 8339 7439 6354 5546 5036 4719 4516 4382 4298 4279 4240 4233 4283 4121 7718 7 0 -6 4317 4373 4471 4594 4749 4896 5065 5275 5481 5664 5878 6196 6551 6924 7373 7946 6462 755.47 0 -5 949 1048 1165 1310 1476 1651 1821 1991 2177 2393 2647 2953 3319 3766 4331 5055 5991 739.08 0 -4 722 887 1108 1408 1818 2371 3101 4023 5105 6214 7017 7116 6488 5421 4235 3182 2156 722.69 0 -4 1747 1313 1007 790 633 519 435 371 321 282 249 222 202 189 182 183 191 706.30 0 -4 207 233 273 326 393 471 565 694 899 1259 1896 2861 3616 3330 2498 1765 1251 689.91 0 -5 89.91 6555 4974 3925 3215 2724 2384 2154 2005 1919 1880 1877 1872 1830 1733 1575 1346 673.52 0 -5 1202 1036 893 7777 681 607 537 484 439 3765 337 310 200 266 249 233 657.12 0 -6 2213 2048 1980 1863 1814 1705 1666 1574 1588 1495 1508 1433 1420 1370 1385 1311 1318 640.73 0 -6 1301 1330 1335 1424 1459 1468 1492 1532 1559 1558 1556 1569 1617 1708 1850 2055 2335 624.34 0 -6 2681 3035 3256 3221 2943 2267 2202 1902 1657 1464 1304 1171 1066 990 937 886 825 607.95 0 -7 750 6906 6375 5953 5613 53535 1314 4706 3962 3915 591.56 0 -7 3841 3796 3773 3760 3739 3720 3702 3744 3745 3786 3814 3867 3933 4020 4094 4211 4312 575.16 0 -7 4400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 558.77 0 -7 44457 4485 4523 4576 4570 4656 4733 1449 4760 4600 4456 4341 4361 4170 1233 1321 1440 1597 1803 552.59 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1904 1760 1653 1568 1508 1508 1508 1518 170 1908 1 -6 1462 1452 1445 1441 1394 1368 1321 1301 1334 1301 1425 1491 1606 1683 1855 1933 2038 2688 2797 2787 2965 1316 1332 1344 1140 1597 1803 552.59 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1904 1760 1653 1568 1508 1404 558.77 0 -7 4400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4803 4805 4805 4715 4627 4553 4489 4454 558.77 0 -7 4457 4485 4523 4576 4570 4656 4733 1344 1391 1425 1491 1406 406 3062 3015 408 408 408 408 408 408 | 902.04 | · I | -6 | 3823 | 4914 | 6757 | 8842 | 8711 | 6482 | 4560 | 3415 | 2763 | 2381 | 2148 | 2018 | 1985 | 2042 | | | |
| 788 26 0 | | | | | | | | | | | | | | | | | | | | |
| 771 87 0 6 4317 4373 4471 4594 4749 4896 5065 5275 5481 5654 5878 6196 6551 6924 7373 7946 8642 75547 0 5 949 1048 1165 1310 1476 1651 1821 1991 2177 2393 2647 2953 3319 3766 4331 5055 5991 739.08 0 4 722 887 1108 1408 1818 2371 3101 4023 5105 6214 7017 7116 6488 5421 4235 3182 2356 722.69 0 4 1747 1313 1007 790 633 519 435 371 321 282 249 222 202 189 182 183 191 706,30 0 4 207 233 273 326 393 471 565 694 899 1259 1896 2861 3616 3330 2498 1765 1251 689.91 0 -5 8951 6555 4974 3925 3215 2724 2384 2154 2005 1919 1880 1877 1872 1830 1733 1575 1386 673 52 0 -5 1202 1036 893 777 681 607 537 484 439 397 365 337 310 290 266 249 233 657.12 0 -6 2213 2048 1980 1863 1814 1705 1666 1574 1588 1495 1508 1433 1420 1370 1385 1311 1318 640 73 0 -6 1301 1330 1335 1424 1459 1468 1492 1532 1559 1558 1556 1569 1617 1708 1850 2055 2335 607 95 0 -7 7550 6906 6375 5953 5613 5359 5134 4934 4760 4600 4456 4341 4234 4146 4066 3962 3015 591 56 0 -7 3841 3796 3773 3760 3739 3720 3702 3744 3745 3786 3814 3867 3933 4020 4094 4211 4312 575 16 0 -7 4400 4460 4550 4619 4667 4701 4701 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 150 1508 1 -7 9312 9256 937 2868 2797 278 1806 1274 1281 1225 1144 1134 1202 1153 1690 1900 188 -1 -7 9312 9256 9572 9814 8861 8529 8349 8349 8349 936 5033 5254 536 6450 4659 4656 4733 4842 4954 5093 5258 5454 5684 5046 6275 6675 7183 5108 -1 -6 982 1022 987 1012 1124 1268 1321 1301 1334 1391 1425 1491 1606 1633 1568 1508 1477 493.69 1 -6 982 1022 987 1012 1124 1268 1321 1301 1334 1391 1425 1491 1606 1633 1568 1508 1477 4457 4485 4523 4566 4679 278 979 2787 2976 3165 3222 2078 2443 2443 2443 2443 2443 2444 1394 4176 4099 4172 9 -1 -5 524 574 638 704 777 862 963 1081 1334 1391 1425 1491 1606 1633 1568 1508 1474 4451 4479 4451 4470 4470 4470 4470 4470 4470 4470 447 | | | | | | | | | | | | | | | | | | | | |
| 755.47 0 -5 949 1048 1165 1310 1476 1651 1821 1991 2177 2393 2647 2953 3319 3766 4331 5055 5991 739.08 0 4 1722 887 1108 1408 1818 2371 3101 4023 5105 6214 7017 7116 6488 5421 4235 3182 2356 722.69 0 4 1747 1313 1007 790 633 519 435 371 321 282 249 222 202 189 183 183 191 186 607 237 266 69891 10 -5 8951 6555 4974 3925 3215 2724 2384 2154 2005 1919 1880 1877 1872 1830 1733 1575 1386 673 52 0 5 1202 1036 1814 1705 16 | | | | | | | | | | | | | | | | | | | | |
| 739.08 0 4 722 887 1108 1408 1818 2371 3101 4023 5105 6214 7017 7116 6488 5421 4235 3182 2356 722.69 0 4 1747 1313 1007 790 633 519 435 371 321 282 249 222 202 189 182 183 191 706.30 0 4 207 233 273 326 393 471 565 694 899 1259 1896 2861 3616 3330 2498 1765 1251 689.91 0 -5 8951 6555 4974 3925 3215 2724 2384 2154 2005 1919 1880 1877 1877 1872 1830 1733 1575 1386 63752 0 -5 1202 1036 893 777 681 667 537 484 439 397 365 337 310 290 266 249 233 657.12 0 -6 2213 2048 1980 1863 1814 1705 1666 1574 1588 1495 1508 1433 1420 1370 1385 1311 1318 64073 0 -6 1301 1330 1335 1424 1459 1468 1492 1532 1559 1558 1556 1569 1617 1708 1850 2055 2335 62434 0 -6 2681 3035 3256 3221 2943 2567 2002 1902 1657 1464 1304 1171 1066 990 937 886 825 60795 0 -7 7550 6906 6375 5953 5613 5399 5134 4934 4760 4600 4456 4341 4234 4144 4466 3962 3915 59156 0 -7 3841 3796 3773 3760 3739 3720 3702 3744 3745 3786 3814 3867 3933 4020 4094 4211 4312 575 16 0 -7 4400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 558.77 0 -7 4457 4485 4523 4576 4570 4656 4733 4842 4954 5093 5258 5454 5684 5946 6275 6675 7183 54238 0 -6 787 881 1007 1155 1275 1303 1247 1177 1132 1118 1133 1171 1233 1321 1440 1597 1803 52599 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1904 1760 1653 1568 1508 1474 51008 -1 -6 1462 1452 1445 1441 1394 1363 1307 1266 1274 1281 1225 1444 1134 1202 1153 1069 993 9134 93.69 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4839 4459 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4839 4459 -1 -5 524 574 638 704 774 862 963 1081 1218 1375 1568 1782 2027 2329 2693 3145 3708 469 0 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2032 1669 1322 1007 4092 4094 4451 1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2072 5408 2577 44451 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
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| 706.30 0 4 207 233 273 326 393 471 565 694 899 1259 1896 2861 3616 3330 2498 1765 1251 689.91 0 -5 8951 6555 4974 3925 3215 2724 2384 2154 2005 1919 1880 1877 1872 1830 1733 1575 1386 657.12 0 6 2213 2048 1980 1863 1814 1705 1666 1574 1588 1495 1508 1433 1420 1370 1385 1311 1318 647.12 0 -6 2681 3035 3256 3221 2943 2567 2202 1902 1657 1464 1304 1171 1066 990 937 886 825 607.95 0 -7 7550 6906 6375 5953 5613 5359 7134 < | | | | | | | | | | | | | | | | | | | | |
| 673 52 0 -5 1202 1036 893 777 681 607 537 484 439 397 365 337 310 290 266 249 233 657.12 0 -6 2213 2048 1980 1863 1814 1705 1666 1574 1588 1495 1508 1433 1420 1370 1385 1311 1318 640 73 0 -6 1301 1330 1335 1424 1459 1468 1492 1532 1559 1558 1556 1569 1617 1708 1850 2055 2335 62434 0 -6 2681 3035 3256 3221 2943 2567 2202 1902 1657 1464 1304 1171 1066 990 937 886 825 6270 0 -7 7550 6906 6375 5953 5613 5359 5134 4934 4760 4600 4456 4341 4234 4146 4166 3962 3915 591 56 0 -7 3841 3796 3773 3760 3739 3720 3702 3744 3745 3786 3814 3867 3933 4020 4094 4211 4312 575 16 0 -7 4400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 558.77 0 -7 4457 4485 4523 4576 4570 4656 4733 4842 4954 5093 5258 5454 5684 5946 6275 6675 7183 542 38 0 -6 787 881 1007 1155 1275 1303 1247 1177 1132 1118 1133 1171 1233 1321 1440 1597 1803 5259 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1994 1760 1653 1568 1508 1474 193.69 -1 -6 1462 1452 1445 1441 1394 1363 1307 1266 1274 1281 1225 1144 1134 1202 1153 1069 990 501 88 -1 -7 9312 9256 9572 9814 8861 8529 8491 8334 9016 9536 9728 9666 9111 8836 9190 9429 9337 493.69 -1 -6 982 1022 987 1012 1124 1268 1321 1301 1334 1391 1425 1491 1606 1683 1855 1983 2025 485.49 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4489 4459 1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4489 440 150 1 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2032 1649 1320 1699 1370 44551 -1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1097 902 802 45571 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 44451 -1 -6 2453 2555 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | 207 | | 273 | 326 | 393 | 471 | 565 | 694 | 899 | 1259 | 1896 | 2861 | 3616 | 3330 | 2498 | 1765 | 1251 |
| 657.12 0 -6 2213 2048 1980 1863 1814 1705 1666 1574 1588 1495 1508 1433 1420 1370 1385 1311 1318 640.73 0 -6 1301 1330 1335 1424 1459 1468 1492 1532 1559 1558 1556 1569 1617 1708 1850 2055 2335 624.34 0 -6 2681 3035 3256 3221 2943 2567 2202 1902 1657 1464 1304 1171 1066 990 937 886 825 607.95 0 -7 7550 6906 6375 5953 5613 5359 5134 4934 4760 4600 4456 4341 4234 4146 4066 3062 3915 591.56 0 -7 3841 3796 3773 3760 3739 3720 3702 3744 3745 3786 3814 3867 3933 4020 4094 4211 4312 575.16 0 -7 4400 4460 4450 4450 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 558.77 0 -7 4457 4485 4523 4576 4570 4656 4733 4842 4954 5093 5258 5454 5684 5946 6275 7183 525.99 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1904 1760 1653 1568 1508 1474 510.08 -1 -6 1462 1452 1445 1441 1394 1363 1307 1266 1274 1281 1225 1144 1134 1202 1153 1069 990 501.88 -1 -7 9312 9256 9572 9814 8861 8529 8491 8334 9016 9536 9728 9666 9111 8836 9190 9429 9337 493.69 -1 -6 982 1022 987 1012 1124 1768 1321 1301 1334 1391 1425 1491 1606 1633 1855 1983 2025 485.49 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4839 469.10 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2012 1649 1322 11901 4609 0-1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2012 1649 1322 1901 4609 0-1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1007 902 802 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 44451 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2379 2354 2383 | 689.91 | 0 | -5 | 8951 | 6555 | 4974 | 3925 | 3215 | 2724 | 2384 | 2154 | 2005 | 1919 | 1880 | 1877 | 1872 | 1830 | 1733 | 1575 | 1386 |
| 640 73 0 -6 1301 1330 1335 1424 1459 1468 1492 1532 1559 1558 1556 1569 1617 1708 1850 2055 2335 624 34 0 -6 2681 3035 3256 3221 2943 2567 2202 1902 1657 1464 1304 1171 1066 990 937 886 825 607 95 0 -7 7550 6906 6375 5953 5613 5359 5134 4934 4760 4600 4456 4341 4234 4146 4466 3962 3915 591 56 0 -7 3841 3796 3773 3760 3739 3720 3702 3744 3745 3786 3814 3867 3933 4020 4094 4211 4312 575 16 0 -7 4400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 558.77 0 -7 4457 4485 4523 4576 4570 4656 4733 4842 4954 5093 5258 5454 5684 5946 6275 6675 7183 542.38 0 -6 787 881 1007 1155 1275 1303 1247 1177 1132 1118 1133 1171 1233 1321 1440 1597 1803 5259 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1994 1760 1653 1568 1508 1474 510.08 -1 -6 1462 1452 1445 1441 1394 1363 1307 1266 1274 1281 1225 1144 1134 1202 1153 1069 990 501 88 -1 -7 9312 9256 9572 9814 8861 8529 8491 8334 9016 9536 9728 9666 9111 8836 9190 9429 9337 493.69 -1 -6 982 1022 987 1012 1124 1268 1321 1301 1334 1301 1425 1491 1606 1683 1855 1983 2025 485.49 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 44839 4454 69.0 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2012 1649 1322 1961 469.0 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2012 1649 1322 1961 460.0 -1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1007 902 802 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 4445 1 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
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| 607 95 0 -7 7550 6906 6375 5953 5613 5359 5134 4934 4760 4600 4456 4341 4234 4146 4066 3962 3915 591 56 0 -7 3841 3796 3773 3760 3739 3720 3702 3744 3745 3786 3814 3867 3933 4020 4094 4211 4312 575 16 0 -7 4400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 558.77 0 -7 4457 4485 4523 4576 4570 4656 4733 4842 4954 5093 5258 5454 5684 5946 6275 6675 7183 542 38 0 -6 787 881 1007 1155 1275 1303 1247 1177 1132 1118 1133 1171 1233 1321 1440 1597 1803 525 99 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1904 1760 1653 1568 1508 1474 510.08 -1 -6 1462 1452 1445 1441 1394 1363 1307 1266 1274 1281 1225 1144 1134 1202 1153 1069 990 50188 -1 -7 9312 9256 9572 9814 8861 8529 8491 8334 9016 9536 9728 9666 9111 8836 9190 9429 9337 493.69 -1 -6 982 1022 987 1012 1124 1268 1321 1301 1334 1391 1425 1491 1606 1683 1855 1983 2025 485.49 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4439 4459 -1 -5 524 574 638 704 774 862 965 1081 1218 1375 1568 1782 2027 2329 2693 3145 3708 469 10 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2032 1649 1322 1061 460 90 -1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1007 902 802 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 4445 1 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
| 591 56 0 -7 3841 3796 3773 3760 3739 3720 3702 3744 3745 3786 3814 3867 3933 4020 4094 4211 4312 575 16 0 -7 4400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 558.77 0 -7 4485 4523 4576 4570 4656 4733 4842 4954 5093 5258 5454 5684 5946 6275 6675 7183 542 38 0 -6 787 881 1007 1155 1275 1303 1247 1177 1132 1118 1133 1171 1233 1321 1440 1597 1803 510 08 -1 -6 1462 1452 1441 1394 1363 1307 1266 1274 1281 1225 1144 11 | | | | | | | | | | | | | | | | | | | | |
| 575 16 0 -7 4400 4460 4550 4619 4667 4701 4741 4766 4793 4803 4835 4805 4715 4627 4553 4489 4454 558.77 0 -7 4485 4523 4576 4570 4656 4733 4842 4954 5093 5258 5454 5684 5946 6275 6675 7183 542 38 0 -6 787 881 1007 1155 1275 1303 1247 1177 1132 1118 1133 1171 1233 1321 1440 1597 1803 525 99 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1904 1760 1653 1568 1508 1474 510 08 -1 -6 1462 1445 1441 1394 1363 1307 1266 1274 1281 1225 1144 1134 1202 1153 1069 990 501 88 -1 -7 9312 | | | | | | | | | | | | | | | | | | | | |
| 558.77 0 -7 4457 4485 4523 4576 4570 4656 4733 4842 4954 5093 5258 5454 5684 5946 6275 6675 7183 542 38 0 -6 787 881 1007 1155 1275 1303 1247 1177 1132 1118 1133 1171 1233 1321 1440 1597 1803 525 99 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1904 1760 1653 1568 1508 1474 510 08 -1 -6 1462 1445 1441 1394 1363 1307 1266 1274 1281 1225 1144 1134 1202 1153 1069 90 501 83 -1 -7 9312 9256 9572 9814 8861 8529 8491 8334 9016 | | | | | | | | | | | | | | | | | | | | |
| 542 38 0 -6 787 881 1007 1155 1275 1303 1247 1177 1132 1118 1133 1171 1233 1321 1440 1597 1803 525 99 0 -6 2099 2527 3186 4245 5830 7135 6291 4480 3216 2508 2132 1904 1760 1653 1568 1508 1474 510 08 -1 -6 1462 1445 1441 1394 1363 1307 1266 1274 1281 1225 1144 1134 1202 1153 1069 990 501 88 -1 -7 9312 9256 9572 9814 8861 8529 8491 8334 9016 9536 9728 9666 9111 8836 9190 9429 9337 493.69 -1 -6 982 1022 987 1012 1124 1268 1321 1301 1334 1391 1425 1441< | | | | | | | | | | | | | | | | | | | | |
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| 501 88 -1 -7 9312 9266 9572 9814 8861 8529 8491 8334 9016 9536 9728 9666 9111 8836 9190 9429 9337 493.69 -1 -6 982 1022 987 1012 1124 1268 1321 1301 1334 1391 1425 1491 1606 1683 1855 1983 2025 485.49 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4839 477.29 -1 -5 524 574 638 704 774 862 963 1081 1218 1375 1568 1782 2027 2329 2693 3145 3708 469.10 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2032 1649 1322 1061 460.90 -1 | | | | 2099 | 2527 | 3186 | 4245 | 5830 | 7135 | 6291 | | | | | | | | | | |
| 493.69 -1 -6 982 1022 987 1012 1124 1268 1321 1301 1334 1391 1425 1491 1606 1683 1855 1983 2025 485.49 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4839 477.29 -1 -5 524 574 638 704 774 862 963 1081 1218 1375 1568 1782 2027 2329 2693 3145 3708 469.10 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2032 1649 1322 1061 460.90 -1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1007 602 802 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 444.51 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
| 485 49 -1 -6 2095 2210 2303 2389 2688 2797 2787 2976 3165 3322 3443 3574 3681 3849 4106 4427 4839 477 29 -1 -5 524 574 638 704 774 862 963 1081 1218 1375 1568 1782 2027 2329 2693 3145 3708 469 10 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2032 1649 1322 1961 460 90 -1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1907 902 802 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 444.51 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
| 477 29 -1 -5 524 574 638 704 774 862 963 1081 1218 1375 1568 1782 2027 2329 2693 3145 3708 469 10 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2032 1649 1322 1061 460 90 -1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1007 602 802 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 444.51 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
| 469 10 -1 -4 439 526 640 787 979 1227 1544 1930 2362 2748 2953 2798 2443 2032 1649 1322 1961 460 90 -1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1997 992 892 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 444.51 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
| 460 90 -1 -5 8527 6935 5717 4764 4039 3473 2999 2593 2235 1928 1682 1459 1266 1120 1007 902 802 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 444.51 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
| 452.71 -1 -6 7250 6618 5960 5411 5043 4716 4349 4158 3898 3646 3523 3314 3193 3083 2978 2868 2577 444.51 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
| 444.51 -1 -6 2453 2525 2510 2400 2229 2075 2127 2296 2317 2333 2420 2345 2222 2287 2370 2354 2383 | | | | | | | | | | | | | | | | | | | | |
| 436 31 -1 -6 2463 2626 2694 2773 2989 3039 | | | | | | | | | | | | | 7 2333 | 2420 | 2345 | 2222 | 2 2287 | 237 | O 2354 | 2383 |
| | 436 3 | 1 - | 1 - | 2463 | 2626 | 2694 | 2773 | 2989 | 3039 | | | | | | | | | | | |

a - The column headed cm⁻¹ contains the wavenumber of the first $k(\tilde{\nu})$ value in the row. The columns

headed XE and YE contain the X-exponent and the Y-exponent, respectively, for the row. The columns headed $0,1,2,\cdots 16$, contain the ordinate values, and the headings give the indices of the ordinate values in the row. In a row which starts with $\widetilde{\nu}(0)$, the wavenumber corresponding to the ordinate indexed J is $\widetilde{\nu}(J) = \widetilde{\nu}(0) - \frac{15798.002}{16384} \cdot J \cdot 2^{XE}$. The $k(\widetilde{\nu})$ values in that row are the ordinate value shown times 10^{YE} . Thus the entry indexed 16 in the first row of the table shows that $k = 1 \times 10^{-7}$ at $\widetilde{\nu} = 6499.90 - \frac{15798.002}{16384} \cdot 16 \cdot 2^3 = 6376.48 \text{ cm}^{-1}$.

b - The $k(\tilde{\nu})$ values in the table can be interpolated to the original wavenumber spacing, 0.482117 cm⁻¹, and yield the original $k(\tilde{\nu})$ values accurate to 1% below 4500 cm⁻¹, 2% between 4500 and 5000 cm⁻¹ and 5% above 5000 cm⁻¹, via the 4-point spline interpolation program TRECOVER⁸.

larger number of spectra from this laboratory did not unduly influence the average peak heights. Peak heights agreed between spectroscopists to ±2.1% on average. There was no evidence that resolution affected this agreement.

The unweighted average absorption index, $k(\tilde{\nu})$, spectrum, areas and peak heights were taken as the primary absorption intensity results of this work. This $k(\tilde{\nu})$ spectrum is shown in Fig. 2.2 and is tabulated in Table 2.7 in Compact Table format⁸.

2.3.2 - Real refractive index spectrum.

The real refractive index spectrum, $n(\tilde{v})$, was determined by Kramers-Kronig transformation of the $k(\tilde{v})$ spectrum, with $n = 1.4773 \pm 0.0004$ at 8000 cm⁻¹. This value was obtained by fitting literature^{9,10} values of the refractive index at eight wavelengths in the visible region to $n^2(\tilde{v}) = a \tilde{v}^4 + b \tilde{v}^2 + c$ and extrapolating to 8000 cm⁻¹. Further, we assumed that the $k(\tilde{v})$ values are all zero between 6500 and 8000 cm⁻¹. The final real refractive index spectrum is shown in Figure 2.3 and the values are in Table 2.8.

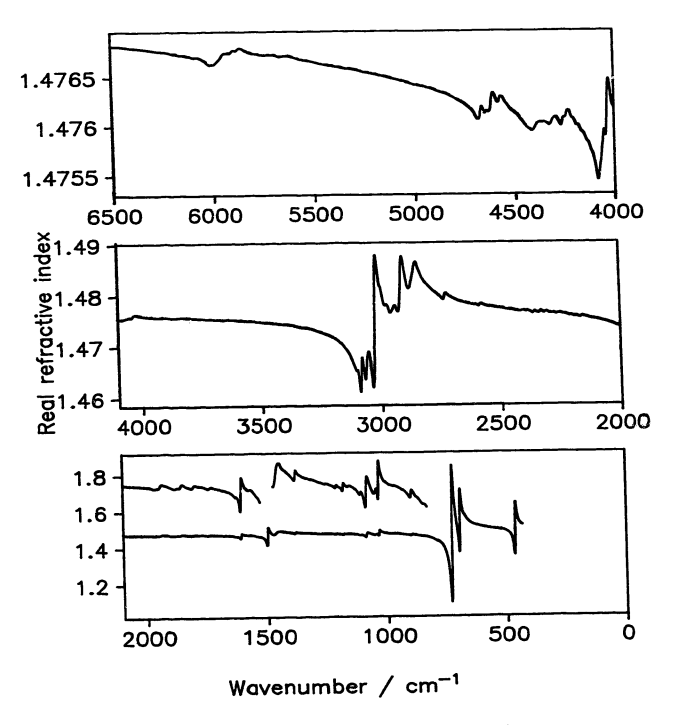


Figure 2.3 - Real refractive index, $n(\tilde{\nu})$, spectrum between 6500 and 435 cm⁻¹ of toluene at 25°C. In the bottom box, the partial spectra shown by the upper curve are $(n(\tilde{\nu}) - 1.21) \times 6.6$, to magnify and offset the curves for clarity.

Table 2.8 - Real refractive indices between 6500 and 435 cm⁻¹ of liquid toluene at 25°C. a-b

| Table 2.8 - Real ferractive mutes between 0500 and 455 cm. of riquid totaline at 25 cm. | | | | | | | | | | | | | | | | | | |
|---|----------|-------|---------|--------|---------|----------------|--------|----------------|---------|--------|---------|--------|---------|-------------|-----------------|---------|-----------------|-----------------|
| om¹ X | | 9 | 1 | 2 | 3 | 4 | | 6 | 7 | H | Ġ | 10 | 11 | 12 | 13 | 14 | 15 | 16 |
| 6499 W 3 | 3 | 14758 | 14768 | 14768 | 14768 | 4768 | 1476 | :4768 | 14768 | 4768 | 14768 | 1-4768 | 14768 | 4768 | | 14768 | | 14768 |
| 4368 76 | 3 | 14768 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 4767 | 14767 | 14767 | 14767 | 14767 |
| 623763 | 3 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14757 | 14767 | 4767 | 14767 | 14767 | 1-1/5/ | 14/6/ |
| 610649 | 3 | 14767 | 14767 | 14767 | 14767 | 14757 | 14767 | 14766 | 14766 | 14766 | 14766 | 1-1765 | 14700 | 14/00 | 1.1767 | 14767 | 1→769 1-1769 | 1.4768 |
| 5975 35 | 3 | 14766 | 14766 | 14766 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14707 | 1 1767 | 1 1767 | 1.1767 | 14767 | 14767 | 1.4767 | 14767 |
| 5844.22 3 5713.08 3 | 3 | 14757 | 14/67 | 14757 | 14757 | 14/5/ 11767 | 14767 | 14767 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14766 |
| 5713 08 3 5581 95 3 | • | 14/0/ | 14707 | 14757 | 1.1766 | 14797 | 1.1766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 |
| 545081 |) | 14790 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 |
| 531968 | , 2 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 |
| 5188 54 | í | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14765 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 |
| 5057.40 | 3 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14764 | 14763 | 14763 | 14763 | 14763 |
| 4926.27 | 3 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14763 | 14762 | 14762 | 14762 |
| 4795 13 | 3 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14761 | 14761 | 14761 | 14761 | 14761 | 14760 | 14760 | | |
| N# 17 6 | 0 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | | 14760 | |
| A | 2 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14761 | 14762 | 14762 | 14763 |
| 25.3.32 | 2 | 14763 | 14763 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 |
| 357 90 | 3 | 14762 | 14761 | 14761 | 14761 | 14761 | 14761 | 14760 | 14760 | 14760 | 14760 | 14760 | 14759 | 14759 | 14759 | 14759 | 14759 | 14759 |
| 76 رهس | | | | | | | | | | 14760 | 14760 | 14759 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 |
| 4269 63 | 3 | 14759 | 14760 | 14760 | 14760 | 14761 | 14761 | 14761 | 14760 | | | 1 4250 | 1 4750 | 14750 | 1 4750 | 1 4750 | 1.4750 | 1.1750 |
| 4211 77 | 2 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 14759 | 14759 | 14759 | 14759 | 14/59 | 14759 | 14/59 | 14/39 | 14/39 | 14/09 | 1→/38 1,1754 |
| 4146 20 | 2 | 14758 | 14758 | 14758 | 14758 | 14758 | 14757 | 14757 | 14757 | 14/57 | 14/57 | 14/50 | 14750 | 14750 | 14757 | 14751 | 14754 | 14763 |
| 4080 64 | 2 | 14754 | 14755 | 14755 | 14756 | 14757 | 14757 | 14/58 | 14759 | 14759 | 14750 | 14/38 | 14760 | 14750 | 14750 | 14750 | 14750 | 14750 |
| 4015 07 3949 50 | 2 | 14762 | 14762 | 14761 | 14761 | 14761 | 14761 | 14/01 | 14/90 | 14700 | 14700 | 1.1759 | 14759 | 14759 | 14759 | 14758 | 14758 | 1.4758 |
| 3949 50 3883 93 | 2 | 14759 | 14759 | 14/59 | 14/59 | 14/58 | 14726 | 14757 | 14757 | 14757 | 14757 | 1.1757 | 14757 | 14757 | 14757 | 14757 | 14757 | 14757 |
| 3883 93 3818 36 | 2 | 14758 | 14757 | 14/5/ | 14/5/ | 14/5/ | 14757 | 14/5/ | 14756 | 1.1756 | 1.1756 | 14757 | 14756 | 14756 | 14756 | 14756 | 14756 | 14755 |
| 3818 36 3752 80 | 2 | 14/5/ | 14757 | 14/5/ | 14757 | 14757 | 14757 | 14750 | 14754 | 14754 | 14754 | 14754 | 14754 | 14754 | 14754 | 14754 | 14754 | 14753 |
| 3687 23 | - | 11752 | 14753 | 14753 | 14753 | 14753 | 1.4753 | 14757 | 14752 | 14752 | 14752 | 14753 | 14753 | 14753 | 14753 | 14753 | 14753 | 14752 |
| 3621 66 | | | | | | | | | | | | | •• | | | | | |
| 3586 47 | 1 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 | 14751 |
| 3574 89 | , | 14751 | 14751 | 14750 | 14750 | 14750 | 14750 | 14750 | 14750 | 14750 | 14750 | 14749 | 14749 | 14749 | 14749 | 14749 | 14748 | 14748 |
| 3509 33 | 2 | 14748 | 14748 | 14748 | 14748 | 14747 | 14747 | 14747 | 14747 | 14746 | 14746 | 14746 | 14746 | 14746 | 14745 | 14745 | 14745 | 14745 |
| 3443 76 | 2 | 14745 | 14744 | 14744 | 14744 | 14744 | 14744 | 14744 | 14743 | 14743 | 14743 | 14742 | 14742 | 14742 | 14742 | 14741 | | |
| 3382 05 | 3 | 14741 | 14741 | 14740 | 14739 | 14739 | 14738 | 14737 | 14736 | 14736 | 14735 | 14734 | 14733 | 14732 | 14731 | 14730 | 14729 | 14728 |
| 3250 91 | 3 | 14727 | 14725 | 14724 | 14722 | 14721 | 14719 | 14717 | 14715 | 14712 | 14709 | | | | | | | |
| 3179 56 | ı | 14709 | 14708 | 14707 | 14706 | 14705 | 14705 | 14704 | 14704 | 14703 | 14702 | 14702 | 14700 | 14699 | 14698 | 14697 | 14696 | 1-4695 |
| 3146 77 | ì | 14694 | 14692 | 14691 | 14689 | 14688 | 14686 | 14684 | 14682 | 14680 | 14678 | 14675 | 14673 | 1-4670 | 14668 | 1-4665 | 14663 | 14661 |
| 3113 99 | 1 | 14659 | 14656 | 14653 | 14651 | 14651 | 14653 | 14653 | 14649 | 14643 | 14635 | 14625 | 14616 | 14611 | 14617 | 14639 | | 14676 |
| 3079 28 | 2 | | 14656 | 14640 | 14637 | 14656 | 14677 | 14687 | 14689 | 14679 | 14662 | 14641 | 14621 | 14632 | 14090 | 14795 | | 14875 14771 |
| | 2 | | | | 14816 | | | | | | | | 14865 | 14773 | 14/00 | 14765 | | 14814 |
| 2948 14 | 2 | 14775 | 14778 | 14778 | 14774 | 14769 | 14767 | 14776 | 14804 | 14844 | 148/1 | | | | | 14836 | | 14829 |
| 2882 57 | | | 14814 | 14821 | 14829 | 14836 | 14844 | 14854 | 14802 | 14804 | 1 4800 | 14822 | 14030 | 1.1804 | 14803 | 14801 | | 14798 |
| 2817 01 2753 37 | 2 | 14820 | 14823 | 14821 | 14820 | 14818 | 14810 | 14814 | 14012 | 14010 | 1.4702 | 14706 | 1.4800 | 14802 | 14802 | | | |
| 2753 37 2720 58 | | 14700 | 14/90 | 1.1707 | 1.4707 | 1.1704 | 14705 | 14/71 | 14704 | 14704 | 1.4702 | 14703 | 14797 | 14797 | 14791 | | | 14790 |
| 2720 58 2687 80 | • | 14799 | 14798 | 1,4790 | 14/7/ | 14720 | 14792 | 14727 | 14797 | 14797 | 14787 | 14786 | 14786 | 14786 | 14785 | 14785 | 14785 | |
| | | 14784 | 14709 | 14794 | 14783 | 14793 | 14783 | 14783 | 14782 | 14782 | 14782 | 14781 | 14781 | 14781 | 14781 | 14781 | 14781 | 14781 |
| 2622 23 | 1 | 14791 | 14780 | 14780 | 14780 | 14770 | 14779 | 14779 | 14779 | 14779 | 14779 | 14779 | 14779 | 14779 | 14778 | 14778 | 14777 | 14776 |
| 2589.45 | 1 | 14776 | 14777 | 14779 | 14781 | 14781 | 14781 | 14780 | 14780 | 14780 | 14779 | 14779 | 14778 | 14778 | 14778 | 14777 | 14777 | 14777 |
| 2556 66 | 1 | 14770 | 14776 | 14776 | 14776 | 14776 | 14775 | 14775 | 14775 | 14775 | 14775 | 14775 | 14775 | 14775 | 14775 | 14774 | 14774 | 14774 |
| 2523.88 | 1 | 14774 | 14773 | 14773 | 14773 | 14773 | 14773 | 14772 | 14772 | 14772 | 14772 | 14772 | 14772 | 14772 | 14772 | 14772 | 14772 | 1477! |
| 2491 10 | 1 | 14771 | 14771 | 14771 | 14771 | 14771 | 14771 | 14770 | 14770 | 14770 | 14770 | 14770 | 14770 | 14770 | 14769 | 14769 | 14769 | 14769 |
| 2458 31 | 1 | 14769 | 14769 | 14769 | 14769 | 14768 | 14768 | 14768 | 14768 | 14768 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 7 14766 |
| 2425 53 | 1 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14767 | 14766 | 14766 | 5 14766 |
| 2392 74 | 1 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14766 | 14765 | 14765 | 14764 | 14764 | 14763 | 14763 | 14762 | 1476 | 2 14762 |
| 2359 96 | 1 | 1476- | 14765 | 14765 | 14765 | 14765 | 14765 | 14764 | 14764 | 14763 | 14763 | 14762 | 14762 | 14763 | 14765 | 14766 | 1476 | 5 14766 |
| 2327 18 | 1 | 1476 | 14765 | 14764 | 14763 | 14763 | 14762 | 14762 | 14763 | 14764 | 14765 | 14765 | 14765 | 14764 | 14764 | 14763 | 1476 | 3 14762 |
| 2294 39 | 1 | 1476 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 14762 | 1476 | 14762 | 14761 | 14761 | 14761 | 1476 | 0 14760 |
| 2261 61 | 1 | 1476 | 1 14761 | 14762 | 14762 | 14762 | 14762 | 14761 | 14761 | 14761 | 14760 | 14760 | 14760 | 14760 | 14760 | 14760 | 1476 | 0 14759 |
| 2228 82 | 1 | 1475 | 14759 | 14759 | 14758 | 14758 | 14758 | 14758 | 14757 | 14757 | 7 14757 | 7 1475 | 7 14757 | 14757 | 7 14757 | 1 14757 | 7 1475 | 7 14756 |
| 2196.04 | 1 | 1475 | 5 14756 | 14755 | 5 14755 | 14755 | 14755 | 14756 | 14756 | 14756 | 5 1475 | 5 1475 | 5 14755 | 14754 | 4 14754 | 14753 | 3 1475 | 3 14754 |
| 2163 26 | 1 | 1475 | 5 14756 | 14756 | 5 14756 | 14756 | 14755 | 14755 | 14754 | 1475- | 1475 | 4 1475 | 4 14753 | 1475 | 3 147 5. | 1475 | 3 1475 | 2 14752 |
| 2130 47 | 1 | 1475 | 2 14752 | 14751 | 1 14751 | 14751 | 14751 | 14751 | 14751 | 1475 | 1 14750 | 0 1475 | 0 14750 | 14750 | 0 14750 | 14750 |) 1474 | 9 14749 |
| 2097 69 | 1 | 1474 | 9 14749 | 14748 | 8 14748 | 14748 | 14748 | 14747 | 7 14747 | 1474 | 7 1474 | 7 1474 | 7 14746 | 1474 | 6 1474 | 1474 | 1474 | 6 14746 |

| Table 2.8 | - Co | ntinue | d | | | | | | | | | | | | | | |
|------------------------|----------------|----------------------|------------------|------------------------|----------|--------------------|----------------|--------------------|------------------|------------------|--------------------|------------------|-------------------------------------|------------------|------------------|--------------------|-----------------------------|
| cm ⁻¹ AE | 0 | 1 | 2 | 3 | 4 | 5 | 0 | 7 | 8 | 3 | 10 | 11 | 12 | 13 | 14 | 15 | 10 |
| 2064-90 1 | 14745 | 14745 | 14745 | 14745 1 | 4744) | | 4744 3 | 4743 | 4-43 | 74743 | 14742 | : 4742 | 14742 | 14741 | 14741 | | 14*4* |
| 2032 12 1 | | | | | | | | | 4737 ! | 14737 | 14730 | 14736 | 14736 | 14736 | 14735 | 14735 | 14734 |
| 1999 34 1 | 14733 | 14732 14729 | 14732 | 14732 1 14732 1 | 4732 L | 4733 I | 4755 | 14752 13775 | | 1.1776 | 1.1735 | 1.1.15.1 | 1.1758 | 14757 | 1.1753 | 14750 | 14747 |
| 1981 98 2 | 14/51 | 1.47.11 | 1-1729 | 14/25 1 1.1725 1 | 4720 I | 4731 1 | 4723 . 4729 | 14726 | 14773 | 14723 | 14710 | 14772 | 14726 | 14725 | 14724 | 14733 | 14748 |
| 1850 84 2 | 1.1752 | 1.1751 | 14738 13736 | 14733 1 | 4736 I | 14733 I | 4731 | 14728 | 14727 | 14725 | 14722 | 14721 | 14728 | 14741 | 14748 | 14748 | 14744 |
| 1785 28 2 | 14741 | 14738 | 14737 | 14737 | 4735 | 4733 | 4731 | 14729 | 14720 | 14722 | 14719 | 14717 | 14719 | 14724 | 14729 | 14730 | 14729 |
| 1719 71 2 | | | | | | | | | | | | | | | | | |
| 1694 64 1 | 14712 | 14711 | 14710 | 14709 | 4707 | 14706 | 14704 | 14703 | 14702 | 14702 | 14702 | 14701 | 14700 | 14698 | 14090 | 140,74 | 14001 |
| 1661 85 1 | 14691 | 1-4690 | 14689 | 14688 | 14687 | 14685 | 14683 | 14681 | 14078 | 14675 | 1-1672 | 14008 | 14005 | 14001 | 14056 | 14652 | 140-48 |
| 1629 07 1 | 14645 | 1-4643 | 14645 | 14647 | 14647 | 1-16-13 | 14637 | 14626 | 14612 | 14593 | 14565 | | | | | _ | |
| 1608 82 0 | 14549 | 14533 | 14528 | 14554 | 14626 | 14713 | 14773 | 14802 | 14810 | 14807 | 1-1798 | 14788 | 14779 | 14772 | 14765 | 14759 | 14753 |
| 1591 47 1 | 14741 | 1-4732 | 14726 | 14723 | 14720 | 14715 | 14709 | 14705 | 14703 | 14702 | 14704 | 14706 | 1-4/07 | 14/06 | 1.4630 | 14/00 | 140.40 |
| 1558 68 1 1525 90 1 | 14691 | 1-4686 | 1-1681 | 14676 | 14673 | 14670 | 14605 | 1.1558 | 14024 | 1.1511 | 1-1-178 | 14043 | 14378 | 14055 | 1-40-5 | 1-40-1 | 14011 |
| 1499 86 0 | 1-100-1 | 1-012 | 14024 | 1-1018 | 1 1.152 | 1 1801 | 15112 | 151.40 | 151/35 | 15063 | 15/171 | 14983 | 14956 | 17033 | 14008 | 1-188-1 | 14861 |
| 1480 58 2 | 1.1707 | 1.4758 | 14736 | 14733 | 14755 | 14799 | 14851 | 14892 | 14915 | 14924 | 14927 | 14913 | 1-1893 | 14878 | 14869 | 14863 | 14850 |
| 1415 01 2 | | | | | | | | | | | | | | | | | |
| 1382 23 1 | 14779 | 14792 | 14835 | 14863 | 14864 | 14857 | 14850 | 14844 | 14839 | 14835 | 14831 | 14828 | 14826 | 14824 | 14821 | 14819 | 14816 |
| 1349.44 1 | 14814 | 14812 | 14810 | 14807 | 14805 | 14803 | 14801 | 14799 | 14797 | 14797 | 14797 | 14796 | 14794 | 14792 | 14790 | 14788 | 14786 |
| 1316 66 1 | 1478- | 14784 | 14786 | 14788 | 14788 | 14786 | 14784 | 14782 | 14781 | 14780 | 14779 | 14778 | 14776 | 14775 | 14773 | 14772 | 14770 |
| 1283 87 1 | 14769 | 14767 | 14766 | 14765 | 14764 | 14763 | 14762 | 14761 | 14760 | 14759 | 1-4757 | 14756 | 14754 | 14753 | 1475] | 14750 | 14748 |
| 1251 09 1 1218 31 1 | 1474 | 8 14749 | 14751 | 14751 | 14751 | 14749 | 14748 | 14746 | 1.1730 | 14743 | 14741 | 14739 | 14737 | 14733 | 14716 | 14730 | 14727 |
| 1218 31 1 | 1472 | 1 14/21 | 14/1/ | 14/13 | 14601 | 1.1600 | 14600 | 1.1607 | 14730 | 14720 | 14755 | 14757 | 14756 | 14752 | 14749 | 14745 | 14743 |
| 1169 13 1 | 1470 | 7 1-770- R 1-1733 | 14701 | 14728 | 14726 | 14726 | 14727 | 14730 | 14733 | 14733 | 14732 | 14730 | 14728 | 14725 | 14723 | 14721 | 14718 |
| 113635 1 | 1471 | 5 14713 | 14710 | 14707 | 14705 | 14701 | 14698 | 14694 | 14689 | 14684 | 1-4678 | 14672 | 14665 | 14659 | 14657 | 14665 | 14679 |
| 1103 56 1 | 1468 | 3 14680 | 14673 | 14663 | 14653 | 14640 | 14625 | 14607 | 14585 | 14561 | 14554 | 14617 | 14741 | 14801 | 14810 | 14802 | 14791 |
| 1070 78 1 | 1477 | 7 14763 | 14750 | 14738 | 14726 | 14715 | 14705 | 14695 | 14666 | 14678 | 14670 | 14669 | 14664 | 14668 | 14679 | 14691 | 14705 |
| 1038 96 | 1470 | 8 14707 | 14703 | 14695 | 14682 | 14664 | 14641 | 14620 | 14630 | 14740 | 14883 | 14936 | 14935 | 14917 | 14898 | 14880 | 14865 |
| 1022 57 | 1485 | 4 14847 | 14841 | 1-4833 | 14826 | 14819 | 14813 | 14807 | 14802 | 14798 | 14793 | 14789 | 14785 | 14782 | 1.17.16 | 14773 | 7 14772 7 14745 |
| 1006 17 | 1476 | 9 14766 | 14764 | 14762 | 14763 | 14763 | 14/63 | 14/02 | 14700 | 14728 | 14730 | 1477 | 14722 | 14721 | 14/47 | , 14/4 | 14141 |
| 986.89 | 1474 | 0 14737 9 14696 | 14/3/ | 14604 | 14/31 | 14602 | 1.1622 | 14720 | 14723 | 14/10 | 5 14673 | 1476 | 9 14665 | 14661 | 1465 | 7 14652 | 2 14645 |
| 934 82 1 902.04 1 | 1463 | 9 14090 0 1.1633 | 1.1631 | 14649 | 1.1679 | 14692 | 14692 | 14685 | 14677 | 14670 | 14663 | 1465 | 7 14650 | 146-29 | 1464 | 1 14631 | 8 14635 |
| 867.33 | 1 1463 | 8 14619 | 1.1608 | 14508 | 14587 | 14579 | 14576 | 14575 | 14570 | 14559 | 14546 | 5 1453 | 1 14515 | 1449 | 1448 | 2 1446- | 1 1445 |
| 804 65 | 1443 | 19 14434 | 14428 | 14423 | 14417 | 14410 | 14404 | 14398 | 14391 | 1438 | 4 14376 | 5 1434 | 9 14360 | 1435 | 2 1434 | 2 (433) | 2 14322 |
| 788.26 | 1431 | 0 14301 | 14303 | 14316 | 14323 | 14320 | 14311 | 14300 | 14289 | 1427 | 7 1426 | 5 1425 | 3 14240 | 1422 | 8 1421 | 4 1420 | 1 14187 |
| 771.87 | 141 | 2 14156 | 14140 | 14123 | 14105 | 14087 | 14068 | 14047 | 14026 | 1400- | 4 13979 | 9 1395 | 4 13927 | 1389 | 8 1386 | 7 1383 | 3 13798 |
| 755 47 | 0 1376 | 50 13721 | 13678 | 13633 | 13587 | 13539 | 13489 | 13433 | 13370 | 1330 | 1 1322 | 2 1313 | 5 13037 | 7 1292 | 6 1279 | 9 1265 | 6 12492 |
| 739 08 | 0 1230 | 05 12094 | 11860 | 11605 | 11345 | 11113 | 10966 | 10997 | 11346 | 1217 | 3 1358 | 2 1532 | 5 16850 |) 1790 | [1839 | 0 1840 | 2 18288 |
| 722 69 | 0 180 | 04 17692 | 17393 | 17121 | 16881 | 16669 | 16484 | 16320 | 16174 | 1604 | 2 1592 | U 1580 | H 1309: | 5 1008 5 1630 | 2 1347 4 1702 | 9 1737 9 1716 | 3 13208 A 1 <i>1</i> 074 |
| 706 30 | 0 151 | 61 15053 | 14941 | 14829 | 14717 | 14601 | 14467 | 14301 | 14093 | 9 1560 7 1500 | 8 1309 0 1597 | 3 1597 3 1597 |)/ 130 -0 0 IS 159(ii | 9 1039 R 1578 | 9 1793 8 1577 | 73 1575 | 8 15740 |
| 689 91 | U 169 | 19 16745 18 15695 | 1028 | 1 10438 1 15444 | 10513 | 12500 | 15120 | , 10043 3 5559 | 15510 | , 1392 9 1552 | 1 1550 | 4 1548 | 7 1547 | 2 1545 | 7 1544 | | |
| 0/3 32 657 12 | U 15/ U 15/ | 18 13693 05 15393 | 7 1207C 15391 | , 13040 1 15370 | 15360 | 15350 | 15340 | 15331 | 15322 | 2 1531 | 3 1530 | 5 1529 | 7 1528 | 9 1528 | 2 1527 | 4 1526 | 7 15260 |
| 640.73 | 0 152 | 53 1524 | 15730 | 15733 | 15228 | 15222 | 15217 | 7 15211 | 15200 | 6 1520 | 1 1519 | 6 1519 | 1518 | 5 1518 | 0 1517 | 75 1517 | ro 15165 |
| 674 34 | 0 151 | 62 1516 | 1516 | 2 15164 | 15165 | 15164 | 1516 | 1 15158 | 3 1515 | 4 1515 | 0 1514 | 6 1514 | 12 1513 | 7 - 1513 | 3 1513 | 30 1512 | 26 15123 |
| ≼∩7.05 | 0 151 | 19 15114 | 5 1511 | 2 15109 | 15106 | 15102 | 15099 | 15090 | 5 15093 | 3 1509 | XO 1508 | 17 150 | 34 1508 | 1 1507 | 8 1507 | 75 1507 | 72 15070 |
| 501.56 | 0 150 | 67 1506 | 4 1506 | 2 15059 | 15057 | 15054 | 15052 | 2 15049 | 1504 | 7 15% | 14 150a | 12 150 | 40 1503 | 7 1503 | 5 150 | 33 1503 | SO 15028 |
| 575 16 | 0 150 | 26 1502 | 4 1502 | 2 15020 | 15018 | 15016 | 1501 | 3 1501 | 1 1500 | 9 1500 | 7 1500 | 1501 10 110 | 03 1500 | 1 1495 | ry 1495 | 77 1495 60 1404 | 95 14993 57 14054 |
| 558 77 | 0 149 | 91 1498 | 9 1498 | 7 14984 | 14982 | 14980 | 1497 | 8 1497 | 1497 | 3 1497 | /1 1496 24 1303 |)9 149 21 140 | 97 1496 98 1 <i>4</i> 07 | M 1490 K 140 |)2 149)1 140 | 147) 18 און ענ | 57 14954 14910 |
| 542 38 | 0 149 | 152 1494 | 9 1494 0 1490 | 7 14945 5 14965 | 14944 | 14943 | 1494 | 4 1494 6 1404 |) 1403 U 1493 | 8 149: | 1491 1401 | 25 149 25 149 | 20 1491 20 1491 | 5 149 | 11 149 | 07 149 | 03 14899 |
| 525 99 610 08 | 3 145 1 149 | NJ 1490 1490 | U 1469 S 1480 | 3 14894 3 14801 | 1,489 | , 14913) 4881 | , 1473 1488 | 6 1488 | 4 148R | 1 148 | 80 148 | 78 148 | 76 1487 | 3 148 | 71 148 | 69 148 | 67 14864 |
| 501.88 | .1 145 | 262 1485 | 8 1485 | 6 14857 | 14851 | 1484 | 7 1484 | 5 1484 | 1 1483 | 8 148 | 35 148 | 32 148 | 29 1482 | 26 148 | 22 148 | 18 148 | 15 14811 |
| √a3 69 | .1 1.45 | 1480 | 3 1479 | 9 14794 | 14789 | 1478 | 1478 | 2 1477 | 8 1477 | 72 147 | 68 147 | 62 147 | 57 1479 | 51 147 | 45 147 | 39 147 | 34 14728 |
| 185 40 | .1 147 | 772 1471 | 5 1470 | 8 14700 | 14693 | 14680 | 5 1467 | 8 1466 | 8 1465 | 59 146 | 50 146 | 40 146 | 29 1461 | 17 146 | 03 145 | 90 145 | /4 14778 |
| 477 29 | .1 14 | 541 1452 | 3 1450 | 3 14484 | 1 1446 | 1 1443 | 7 1441 | 1 1438 | 3 1435 | 52 143 | 20 142 | 84 142 | 47 1420 | 3 141 | 56 141 | 02 140 | 44 13979 |
| 469 10 | .1 139 | 210 1383 | 0 1374 | 8 13660 | 1358 | 4 1352 | 2 1351 | 3 1358 | 5 1382 | 20 142 | 51 149 | 03 159 | 72 160 | 30 162 | 99 164 | 109 164 | 27 16383 |
| 460.90 | -1 16 | 317 1623 | 2 1615 | 3 1607 | 2 1600 | 1 1593 | 6 1588 | 13 1583 | 1 1578 | 87 157 | 42 157 | 03 156 | 65 156 | 31 155 | 96 155 | 68 155 | 40 15517 |
| 452 71 | -1 15 | 492 1547 | 2 1544 | 1543 | 3 1541 | 4 1539 | 9 1538 | 3 1536 | 9 1539 | 56 153 | 44 153 | 32 15 | 122 153 | 11 153 | 03 152 | 93 152 | 286 15278 |
| 444 51 | -1 15 | 269 1526 | 0 1525 | 55 1524 | 8 1524 | 3 1523 | 4 1522 | 27 1522 | 21 1521 | 18 152 | 11 152 | 157 EUS | 152 כני | ולו וע | ולו כע | ולו נעו | 88 15187 |
| -136 31 | -1 15 | 182 1518 | 52 151 | /8 1518 | 1 1517 | o 1517 | • | | | | | | | | | | |

a - The column headed cm⁻¹ contains the wavenumber of the first $n(\tilde{\nu})$ value in the row. The

column headed XE contains the X-exponent for the row. The columns headed $0,1,2,\cdots 16$, contain the $n(\tilde{\nu})$ values with the decimal point implicitly after the first digit in each value, and the headings give the indices of the $n(\tilde{\nu})$ values in the row. In a row which starts with $\tilde{\nu}(0)$, the wavenumber corresponding to the ordinate indexed J is $\tilde{\nu}(J) = \tilde{\nu}(0) - \frac{15798.002}{16384} \cdot J \cdot 2^{\lambda E}$. Thus, the entry indexed 16 in the first row of the table shows that n = 1.4768 at $\tilde{\nu} = 6499.90 - \frac{15798.002}{16384} \cdot 16 \cdot 2^3 = 6376.48 \text{ cm}^{-1}$.

b - The $n(\tilde{\nu})$ values in the table can be interpolated to the original wavenumber spacing, 0.482117 cm⁻¹. and yield the original $n(\tilde{\nu})$ values accurate to 0.1%, via the 4-point spline interpolation program TRECOVER⁸.

2.3.3 - Molar absorption coefficient spectrum.

To present the data in a form preferred by analytical chemists, we have used the final $k(\widetilde{v})$ spectrum to calculate the molar absorption coefficient spectrum¹, by $E_m(\widetilde{v}) = 4\pi \ \widetilde{v} \ k(\widetilde{v})/(2.303\ C)$, where C is the molar concentration. The $E_m(\widetilde{v})$ spectrum is shown in Figure 2.4 and the values of $E_m(\widetilde{v})$ are tabulated in Table 2.9 in Compact Table format⁸. For liquid toluene at 25°C, the molar concentration is 9.36 mol/L, as calculated from its density of 0.8623 g/ml⁹. The molar absorption coefficient is in the unit L mole⁻¹ cm⁻¹. The areas under the molar absorption coefficient bands are in Table 2.10, with their estimated accuracies. The areas are in the unit used by analytical chemists, L mole⁻¹ cm⁻². The values should be divided by 100 to put them in the km mole⁻¹ unit usually used by physical chemists. The values of the molar absorption coefficient at the peaks of the bands are listed in Table 2.11.

For selected spectral regions, the molar absorption coefficient spectrum, the

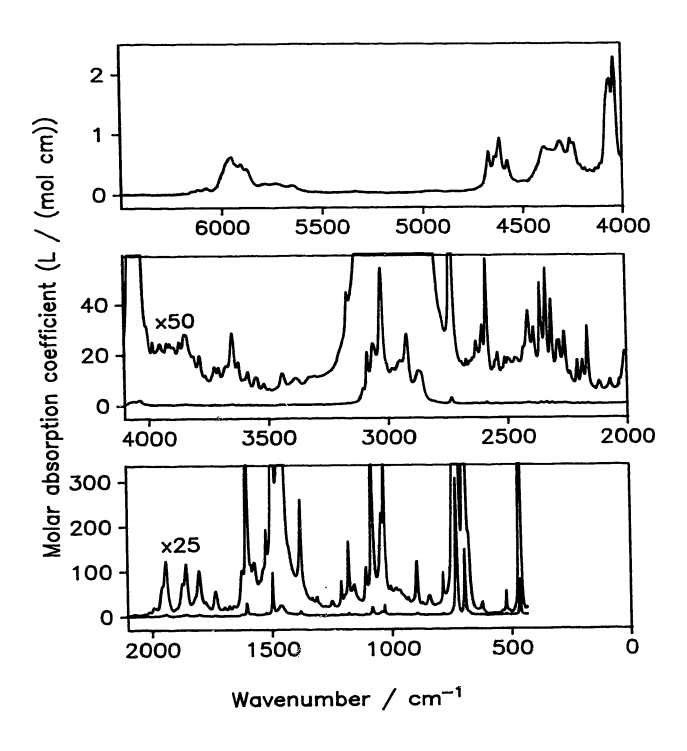


Figure 2.4 - Molar absorption index, $E_{\rm m}(\tilde{\nu})$, spectrum between 6500 and 435 cm⁻¹ of toluene at 25°C. The scale labels in the middle and bottom boxes are for the lower spectrum in the box; they must be divided by 50 or 25, as shown, for the upper spectrum in the box.

Table 2.9 - Molar absorption coefficients between 6500 and 435 cm⁻¹ of liquid toluene at 25°C. a.b.c

| Table 2 | . 7 | - N | noiar | aoson | puou | COGIII | ciciits | betwe | cu o. | 500 ai | IU 45. | CIII | OI 110 | laid i | <i></i> | | | | |
|------------------------|--------|----------------|--------------|--------------|--------------|--------------|--------------|----------------------|--------------|--------------|---------------------|--------------|--------------|---------------------------------------|--------------|--------------|----------------|------------------|--------------|
| cm ' X | Œ | YE: | 0 | 1_ | 2 | 3 | 4 | 5 | 6 | 7 | 8 | y | 10 | 11 | 12 | 13 | 14 | 15 | 16 |
| 127111 | 3 | 5 | 8321 | 5917 | 44,52 | 18.11 | 1117 | <u>5247</u> | 5020 | 2708 | 4952 | 3057 | 3309 | 3052 | 988 | 1268 | 3069 | | 372 |
| 6368.76 | 3 | 4) | 1862 | 371 | 371 | 370 | H22 | 1465 | 968 | 507 | 543 | 367 | 367 | 367 | 366 5075 | 2355 5351 | 5524 - 6096 | | 7822 8100 |
| 6237 63 | | -5 | 1389 | 1019 | 1017 | 1083 | 1233 1680 | 1461 891 | 1840 726 | 1593 567 | 152 7 732 | 2280 239 | 3195 1172 | 5043 1540 | 5035 2161 | 2866 | | | 4763 |
| 6196 49 3 5975 35 3 | | 4 4 | 691 5269 | 732 5690 | 844 5985 | 995 6180 | 5964 | 5244 | 4972 | 4735 | 4675 | 4938 | 4827 | 4282 | 4050 | 4193 | 3907 | | 2645 |
| 5844 22 | | 4 | 2158 | 1795 | 1600 | 1489 | 1511 | 1635 | 1717 | 1773 | 1800 | 1729 | 1663 | 1681 | 1715 | 1752 | 1824 | 1877 | 1772 |
| 5713 08 | | 4 | 1671 | 1614 | 1472 | 1381 | 1303 | 1255 | 1301 | 1399 | i-146 | 1436 | 1372 | 1149 | 917 | 803 | 740 | 528 | 536 |
| 5581 95 | | .5 | 5597 | 4790 | 3810 | 4477 | 3904 | 3485 | 4062 | 3756 | 3465 | 3184 | 3641 | 3331 | 2777 | 2906 | | | 2589 |
| 5450 81 | 3 | ٠5 | 2681 | 2561 | 2571 | 2569 | 2607 | 2685 | 2894 | 2846 | 3016 | 3365 | 3505 | 3463 | 3764 | 4007 | | | 4362 |
| | 3 | -5 | 3663 | 3706 | 3732 | 3030 | 3076 | 2951 | 2582 | 2498 | 2352 | 2265 | 2421 | 2712 | 2584 | 2270 2002 | 2249 2030 | | 2104 2000 |
| | 3 | -5 | 1973 | 1782 | 1400 | 1678 2492 | 2169 2546 | 2065 2699 | 1874 2914 | 1925 3227 | 1940 3704 | 1891 4270 | 1903 4609 | 1831 5069 | 1931 5164 | 5106 | 5010 | | 4657 |
| | 3 3 | -5 -5 | 2091 4692 | 2094 4349 | 2362 4196 | 3832 | 3574 | 3401 | 3554 | 3775 | 3963 | 4002 | 3993 | 4064 | 3992 | 3991 | 3943 | | 3863 |
| | 3 | -4 | 412 | 457 | 519 | 568 | 591 | 625 | 645 | 716 | 816 | 952 | 1197 | 1249 | 1458 | 1869 | 2807 | | |
| 4686 17 | | 4 | 2973 | 3135 | 3303 | 3489 | 3664 | 3844 | 4031 | 4215 | 4421 | 4623 | 4836 | 5077 | 5333 | 5601 | 5889 | | 6460 |
| 166, 99 | 2 | -1 | 7050 | 6386 | 5313 | 4649 | 1487 | 4714 | 5357 | 6015 | 6130 | 6020 | 6290 | 6932 | 7679 | 8575 | 9206 | | 7328 |
| | 2 | 4 | 5820 | 4872 | 4342 | 4077 | 1089 | 4478 | 5007 | 5476 | 5187 | 4392 | 3822 | 3384 | 2936 3650 | 2594 4190 | 2382 4697 | | 2195 6303 |
| | 3 | -1 | 2088 | 1902 | 1883 | 1972 | 2031 | 2067 | 2030 | 2006 7136 | 2181 7103 | 2440 7266 | 7739 7739 | 3157 8546) | 3659 | 8173 | 7503 | | 6765 |
| | 3 | -1 -1 | 7181 7422 | 7672 9116 | 7724 8371 | 7450 8347 | 7256 7980 | 7238 6476 | 7225 5181 | 4717 | ,103 | , 200 | ,4,500 | · · · · · · · · · · · · · · · · · · · | 4443 | | | | |
| | 2 | 4 | 4739 | 4584 | 4325 | 4093 | 3938 | 3940 | 4170 | 4219 | 3954 | 3725 | 3619 | 3645 | 3823 | 3921 | 3875 | 3780 | 3657 |
| | 2 | .3 | 362 | 369 | 390 | 414 | 419 | 409 | 404 | 419 | 451 | 484 | 494 | 497 | 521 | 581 | 688 | 860 | 1113 |
| 4080 64 | 2 | .3 | 1383 | 1562 | 1667 | 1772 | 1871 | 1892 | 1914 | 1849 | 1695 | 1716 | 1949 | 2228 | 2214 | 1861 | | 1042 | 817 |
| | 2 | -4 | 7002 | 6366 | 6304 | 5968 | 5345 | 4722 | 4328 | 4188 | 4450 | 5036 | 4531 | 4226 | 4221 4609 | 4298 4620 | 4651 | 4543 4590 | 4815 4423 |
| | 2 | -4 | 4832 | 4574 | 4386 | 4297 | 4236 4993 | 4343 4685 | 4792 4536 | 4914 4746 | 4679 5237 | 4664 5595 | 4824 5622 | 4778 5578 | 5411 | 4910 | | 4187 | 3999 |
| | 2 | 그 | 4207 3845 | 4143 3866 | 4360 3869 | 4871 3644 | 3296 | 3070 | 3110 | 3448 | 3899 | 3733 | 3127 | 2637 | 2367 | 2298 | 2382 | 2278 | 2109 |
| | 2 | 4 | 1993 | 1953 | 1972 | 2058 | 2229 | 2416 | 2677 | 2999 | 2961 | 2671 | 2592 | 2824 | 2997 | 2709 | 2367 | 2291 | 2371 |
| | 2 | -4 | 2586 | 2802 | 2998 | 3124 | 3094 | 3079 | 3257 | 3714 | 4513 | 5387 | 5715 | 5190 | 4207 | 3399 | 3040 | 3093 | 3375 |
| 3621 66 | 2 | 4 | 3314 | 2722 | 2198 | 1993 | 1977 | 2014 | 2054 | 2084 | 2290 | 2566 | | | | | | | |
| 3586 47 - | | 4 | 2589 | 2605 | 2622 | 2639 | 2646 | 2638 | 2611 | 2559 | 2489 | 2417 | 2345 | 2281 | 2220 | 2162 | 2108 | 2060 | 2012 1491 |
| | 2 | -4 | 1702 | 1550 | 1517 | 1627 | 1867 | 2113 | 2221 | 2219 | 2039 | 1700 1294 | 1444 1324 | 1356 1357 | 1388 1397 | 1515 1456 | | 1678 1758 | 2087 |
| | 2 | -1 | 1354 | 1270 2527 | 1253 2438 | 1281 2216 | 1303 1963 | 1263 1 769 | 1217 1672 | 1215 1648 | 1254 1669 | 1718 | 1781 | 1834 | 1895 | 1994 | 2091 | 1750 | 200, |
| | 2 | -÷ -4 | 2389 2126 | 2017 | 1826 | 1695 | 1709 | 1872 | 2067 | 2234 | 2259 | 2271 | 2314 | 2229 | 2245 | 2285 | | 2386 | 2483 |
| | 3 | 4 | 2548 | 2656 | 2739 | 2910 | 3176 | 3543 | 4097 | 4457 | 4932 | 5434 | | | | | | | |
| | 1 | -4 | 5619 | 5870 | 6217 | 6741 | 7531 | 8453 | 8916 | 8841 | 8607 | 8404 | 8307 | 8295 | 8436 | 8584 | | 8965 | 9221 |
| 31 46 77 | 1 | -3 | 942 | 974 | 1007 | 1050 | 1092 | 1143 | 1199 | 1270 | 1359 | 1474 | 1624 | 1820 | 2060 | 2355 | | 3122 | 3-170 |
| 3113 99 | ı | -2 | 378 | 416 | 471 | 548 | 627 | 663 | 647 | 622 | 624 | 673 | 790 | 1005 | 1351 | 1786 | | 2022 | 1782 |
| 3079 28 | 2 | -2 | 1388 | 1319 | 1545 | 2028 | 2392 | 2393 | 2258 | 2085 | 1965 | 2013 | 2286 1425 | 3008 1390 | 4189 1362 | 5212 1429 | 1562 | 4153 1650 | 2902 1713 |
| 3013.71 | 2 | -2 -2 | 2167 1747 | 1800 1728 | 1656 1697 | 1573 1714 | 1410 1834 | 1301 | 1285 2456 | 1330 2760 | 1405 2668 | 1440 2156 | 1609 | 1226 | 1003 | 892 | 866 | | 997 |
| 2948 14 2882 57 | 2 | -2 | 1115 | 1235 | 1325 | 1336 | 1313 | 1275 | 1188 | 1003 | 777 | 585 | 449 | 358 | 294 | 246 | 211 | 195 | 176 |
| 281701 | 2 | -3 | 1566 | 1512 | 1448 | 1294 | 1148 | 1038 | 940 | 850 | 791 | 755 | 725 | 690 | 654 | 613 | 587 | 584 | 592 |
| 2753 37 | ī | -3 | 604 | 628 | 667 | 727 | 822 | 970 | 1202 | 1552 | 2031 | 2524 | 2746 | 2506 | 2016 | 1553 | 1209 | | 809 |
| 2720 58 | 1 | -4 | 6967 | 6176 | 5599 | 5146 | 4776 | 4476 | 4224 | 4007 | 3830 | 3698 | 3578 | 3472 | 3401 | 3374 | 3380 | | 3345 |
| 2687 80 | l | -4 | 3264 | 3189 | 3136 | 3088 | 3038 | 3023 | 3082 | 3272 | 3536 3705 | 3477 3853 | 3223 4199 | 3093 4752 | 3089 5020 | 3165 4844 | | 3363 4396 | 3415 4250 |
| 2655 02 | ì | - 1 | 3413 | 3397 | 3422 | 3509 | 3611 4402 | 3684 4843 | 3693 5288 | 3673 5502 | 5875 | 6321 | 5954 | 5275 | 4824 | 4707 | | 5555 | 6792 |
| 2622 23 2589 45 | l l | -4 -3 | 4150 871 | 4090 1079 | 4085 1154 | 4169 1004 | 784 | 619 | 512 | 439 | 385 | 343 | 313 | 293 | 281 | 273 | 272 | | 284 |
| 2556 66 | | -4 | 2906 | 2995 | 3163 | 3380 | 3503 | 3495 | 3521 | 3738 | 4051 | 4011 | 3664 | 3332 | 3056 | 2834 | 268 | 7 2635 | 2686 |
| | i | -4 | 2796 | 2843 | 2840 | 2841 | 2900 | 3064 | 3346 | 3654 | 3703 | 3465 | 3264 | 3223 | 3324 | 3522 | | 7 3603 | 3549 |
| 2491 10 | | 4 | 3549 | 3535 | 3477 | 3404 | 3336 | 3297 | 3282 | 3278 | 3288 | 3318 | 3382 | 3485 | 3614 | 3685 | | 3642 | 3621 |
| | ı | 4 | 3602 | 3569 | 3509 | 3440 | 3377 | 3336 | 3321 | 3341 | 3409 | 3524 | | 3883 | 4174 | 4442 | | 4503 | 4546 5117 |
| 2425 53 | | 4 | | | 5229 | 5703 | 6247 | 6764 | 7229 | 7392 | 7058 | 6504 | | 5524 3824 | 5176 4021 | 4943 4427 | | 4912 | 5117 8977 |
| | | 4 | | | | 5752 652 | 5146 603 | 4658 557 | 4280 526 | 3992 522 | 3823 552 | 3750 620 | | 832 | 1002 | 1073 | | 2 688 | 543 |
| 2359 96 2327 18 | | .3 | | 816 4190 | | | 4645 | 5480 | 6770 | 8097 | 8166 | 6791 | | 4380 | 3796 | 3465 | | 5 3316 | 3449 |
| 2294 39 | | 4 | | | | 4835 | 5040 | 5002 | 4942 | 5072 | 4805 | 4281 | | 3714 | | 3690 | | 3 4343 | 4996 |
| 2261 61 | | 4 | | | | | 3823 | 3363 | 3004 | | 2538 | | | | | 2561 | | 9 2187 | 2032 |
| 2228 82 | | -4 | | | | | | 1720 | 1753 | 1889 | 2141 | 2469 | | 3513 | 3049 | 2383 | | 5 1904 | 1930 |
| 2196 04 | | -4 | | | | 2822 | 3125 | 3454 | 3473 | 3071 | 2673 | | | | | | | 5 4392 | 5712 |
| 2163 26 | | -4 | | | | | | 2090 | 1830 | | | | | | | 1336 | | 6 1260 | 1213 |
| 213047 | | -4 | | | | | | 1507 | 1665 | | | | | | | | | 8 1244 1 10#9 | 1213 1896 |
| 2097 69 | 1 | 1 | 1207 | 1205 | 1201 | 1206 | 1222 | 1250 | 1283 | 1316 | 1342 | 1375 | 1429 | 1505 | 1593 | 1694 | 193 | 1 1942 | 1020 |

Table 2.9 - Continued.

| Table 2 | .9 | <u>- C</u> | ontin | ued. | | | | | | | | | | | | | | | |
|----------------------|--------|------------|--------------|--------------|--------------------------|--------------|--------------|--------------|--------------|--------------|--------------|---------------------|--------------|--------------|--------------|--------------|-------------|-----------------|--------------|
| | _ | E | 0 | 1 | | 3 | 4 | <u> </u> | | • | 8 | ٠, | 10 | 11 | 12 | 13 | 14 | 15 | 10 |
| 2064 90 | | | 1707 | 1558 | | 1402 | | • • • | 1004 | | 1247 | | 1250 | | 1313 | 1307 | - | 1501 | 1821 |
| 2032 12 1 1999 34 | | | 2001 4294 | 1939 | 9139 1 011 | 2011 7500 | 2138 | 230 1 | -04 '411 | 2756 7167 | 3073 | ::0] | 1001 | 4054 | 4191 | 1170 | -4033 | 1901 | 4018 |
| 1981 98 | | -3 | 710 | 636 | 609 | 967 | 1484 | 2176 | 2610 | 2093 | 3(4)2 | 3037 | 4841 | 4017 | 3251 | 2107 | 1478 | 1124 | 827 |
| 191641 | | -3 | 6-11 | 549 | 530 | 577 | 081 | 713 | 755 | 3(10 | 1145 | 1564 | 2234 | 2837 | 2865 | 2800 | | 45.71 | 4153 |
| 1850 84 | 2 | -3 | 2851 | 1817 | 1291 | 1073 | 1000 | 1170 | 1274 | 1486 | 1673 | 1811 | 2170 | 2077 | 3884 | 3894 | 2935 | 2037 | 1577 |
| | 2 | -3 | 1295 | 1225 | 1266 | 1113 | 908 | 855 | 761 | 080 | c-15 | 725 | 985 | 1463 | 2012 | 2229 | 1911 | 1414) | 1 th No |
| | 2 | -1 | 7364 | 6267 | 6070 | 6117 | 6258 | 6912 | 8097 | | | | | | | | | | |
| = | 1 | -1 | 7438 | 6842 | 6374 | 6027 | 5832 893 | 5917 870 | 6252 847 | 7057 | 8180 | 9033 | 8797 933 | 7791 | 0900 | 0533 | 0437 | | 7272 |
| | l l | -3 -3 | 811 2734 | 892 3337 | 925 3861 | 911 4054 | 3973 | 3827 | 3757 | 827 3872 | 827 4294 | 871 5237 | 7340 | 1015 | 1128 | 1278 | 1483 | 1207 | 2210 |
| | ò | .2 | 933 | 1251 | 1741 | 2370 | 2835 | 2804 | 2419 | 1963 | 1564 | 1253 | 1030 | 877 | 771 | 691 | 625 | 509 | 523 |
| | l | -3 | 4713 | 4546 | 4551 | 4520 | 4211 | 3992 | 4021 | 4193 | 4421 | 4662 | 4750 | 4650 | 4372 | 4CH XX | | 3316 | Sixti8 |
| 1558 68 | ı | -3 | 2913 | 2860 | 2870 | 2983 | 3126 | 3116 | 3101 | 3193 | 3376 | 3621 | 3884 | 4076 | 4207 | 4286 | 4373 | 4580 | 5101 |
| 1525 90 | 1 | -2 | 628 | 766 | 700 | 597 | 558 | 554 | 568 | 598 | 642 | 711 | 824 | 1003 | 1325 | 1937 | | | |
| | 0 | -2 | 2465 | 3305 | 4727 | 7019 | 9428 | 9089 | 6815 | 4851 | 3627 | 2864 | 2352 | 2033 | 1815 | 1603 | 1429 | | 1247 |
| | 2 | -2 | 1197 | 1323 | 1554 | 1866 | 2182 | 2357 | 2304 | 2059 | 1736 | 1438 | 1127 | 848 | 701 | 637 | 597 | 541 | 483 |
| | 2 | -3 | 4359 | 4050 | 3831 | 3612 787 | 3527 571 | 3591 446 | 3875 375 | 4554 332 | DÚKKÓ ZUC | 277 | 260 | 247 | 234 | 219 | 204 | 193 | 183 |
| | l l | -2 -3 | 736 1763 | 948 1695 | 1018 1623 | 1558 | 1509 | 1473 | 1459 | 1469 | 300 1525 | 1599 | 1516 | 1407 | 1336 | 1290 | 1266 | 1263 | 1299 |
| | ı l | .3 | 1389 | 1553 | 1680 | 1540 | 1303 | 1161 | 1105 | 1093 | 1076 | 1038 | 981 | 923 | 877 | 847 | 831 | 823 | 823 |
| | ì | -3 | 830 | 840 | 855 | 863 | 860 | 851 | 837 | 822 | 809 | 804 | 806 | 816 | 831 | 856 | 894 | 960 | 1072 |
| | 1 | .3 | 1222 | 1321 | 1291 | 1194 | 1085 | 992 | 918 | 864 | 828 | 807 | 800 | 802 | 814 | 837 | 868 | 916 | 983 |
| 1218 31 | ı | -3 | 1078 | 1224 | 1503 | 2151 | 3050 | 2710 | 2023 | 1629 | 1487 | 1503 | 1614 | 1747 | 1830 | 1859 | 1881 | 1934 | 2050 |
| 118649 | 0 | -3 | 2145 | 2277 | 2464 | 2731 | 3123 | 3720 | 4620 | 5800 | 6617 | 6223 | 5160 | 4192 | 3498 | 3038 | 2734 | 2533 | 2198 |
| | l | -3 | 2243 | 2194 | 2220 | 2296 | 2414 | 2561 | 2693 | 2758 | 2566 | 2304 | 2086 | 1911 | 1775 | 1683 | 1613 | 1546 | 1480 |
| | 1 | -3 | 1430 | 1398 | 1385 | 1382 | 1359 | 1326 | 1306 | 1313 | 1342 | 1404 | 1511 | 1698 | 2026 | 2582 | 3413 | 4224 | 4195 |
| | l | -2 | 360 | 311 | 283 | 270 | 273 | 292 | 330 3537 | 401 3771 | 529 4112 | 772 4598 | 1224 5233 | 1796 6031 | 1799 6967 | 1356 7945 | 993 8761 | 759 9145 | 601 8987 |
| | 1 | -3 -2 | 4898 876 | 4169 851 | 3725 830 | 3483 822 | 3388 837 | 3411 896 | 1036 | 1334 | 1883 | 2393 | 2146 | 1572 | 1132 | 853 | 683 | 577 | 512 |
| 1022.57 | | -2 | 4733 | 4371 | 3942 | 3570 | 3285 | 3067 | 2897 | 2764 | 2654 | 2565 | 2490 | 2430 | 2386 | 2353 | | 2326 | 2329 |
| 1006 17 | | -3 | 2346 | 2376 | 2441 | 2555 | 2626 | 2572 | 2477 | 2388 | 2319 | 2262 | 2224 | 2194 | 2174 | 2162 | | 2163 | 2177 |
| 986 89 | 2 | -3 | 2270 | 2398 | 2410 | 2265 | 2217 | 2248 | 2213 | 2042 | 1888 | 1797 | 1754 | 1664 | 1524 | 1404 | | | |
| 934 82 | ı | -3 | 1397 | 1449 | 1557 | 1591 | 1473 | 1329 | 1237 | 1194 | 1185 | 1200 | 1232 | 1268 | 1311 | 1357 | 1417 | 1515 | 1693 |
| 902.04 | l | -3 | 2010 | 2578 | 3537 | 4619 | 4540 | 3371 | 2367 | 1768 | 1-128 | 1228 | 1105 | 1036 | 1017 | 1044 | | 1157 | 1131 |
| 867 33 | 2 | -3 | 1017 | 944 | 944 | 1020 | 1197 | 1507 | 1777 | 1729 | 1436 | 1165 | 986 | 900 | 880 | 909 | | 1009 | 1023 |
| 804.65 | | -3 | 1031 | 1043 | 1054 | 1068 | 1088 | 1109 | 1129 | 1154 | 1180 | 1210 | 1251 | 1286 | 1342 | 1415 | | 1641 | 1852 |
| | 0 | -3 | 2200 1942 | 2798 | 3567 2006 | 3817 2059 | 3401 2125 | 2901 2189 | 2529 2261 | 2294 2352 | 2147 2441 | 2052 2515 | 1988 | 1948 2749 | 1937 2903 | 1917 3064 | 1911 | 1931 3508 | 1810 1946 |
| 771 87 755 47 | | -3 -2 | 418 | 1965 461 | 512 | 574 | 647 | 722 | 796 | 869 | 949 | 1041 | 1151 | 1282 | 1439 | 1631 | 1873 | | 2584 |
| 739 08 | o | -1 | 311 | 381 | 476 | 604 | 779 | 1015 | 1325 | 1717 | 2176 | 2645 | 2983 | 3021 | 2751 | 2296 | 1791 | | 994 |
| 722.69 | o | -2 | 7358 | 5523 | 4231 | 3313 | 2652 | 2173 | 1817 | 1547 | 1339 | 1172 | 1033 | 922 | 838 | 782 | 753 | | 786 |
| | | -1 | 85 | 96 | 112 | 134 | 161 | 193 | 231 | 283 | 366 | 512 | 770 | 1160 | 1464 | 1346 | 1009 | 712 | 504 |
| 689 91 | 0 | -2 | 3599 | 2632 | 1994 | 1572 | 1286 | 1088 | 950 | 858 | 797 | 762 | 746 | 743 | 740 | 723 | 683 | 620 | 545 |
| 673 52 | | -3 | 4718 | 4060 | 3497 | 3036 | 2659 | 2365 | 2091 | 1880 | 1704 | 1539 | 1411 | 1302 | 1196 | 1116 | 1022 | | 894 |
| 657 12 | 0 | -4 | 8477 | 7830 | 7560 | 7103 | 6905 | 6481 | 6326 | 5968 | 6011 | 5649 | 5691 | 5399 | 5344 | 5147 | 5194 | | 4930 |
| 640 73 | 0 | -4 | 4857 | 4959 | 4971 | 5294 | 5414 | 5441 | 5523 | 5659 | 5752 | 5738 | 5722 | 5761 | 5927 | 6252 | 6762 | | 8510 293 |
| 624 34 | | -3 1 | 976 | 1103 | 1181 | 1167 | 1064 | 927 | 794 | 685 | 596 1665 | 525 160 7 | 467 1554 | 419 1511 | 381 1472 | 353 1439 | 334 | 1370 | 1352 |
| 607 95 591 56 | | | 2675 1324 | 2443 1307 | 2252 1296 | 2099 1290 | | 1884 1272 | 1802 1264 | 1729 1276 | 1665 1274 | 1286 | 1293 | | 1330 | | | 1416 | 1448 |
| 575 16 | | | 1475 | 1493 | | 1541 | 1554 | 1562 | 1573 | 1579 | 1585 | 1586 | 1594 | | 1549 | | | 1467 | 1453 |
| 558 77 | | | 1451 | 1458 | | 1483 | | 1503 | 1526 | 1558 | 1591 | 1633 | 1683 | | 1813 | | | 1 2118 | 2275 |
| 542 38 | | | 2486 | | | 3630 | | 4083 | 3901 | 3676 | 3526 | | 3518 | | 3813 | | | 7 4914 | |
| 525.99 | | | 643 | 773 | | 1294 | | 2167 | 1907 | 1356 | 972 | | | | 528 | | 46 | 8 450 | 439 |
| 510 08 | | | 4345 | | | 4271 | | 4032 | | 3737 | 3759 | 377 7 | 3608 | | 3334 | | | 1 3134 | |
| 501 88 | | | 2724 | 2705 | 2794 | 2862 | | 2483 | 2469 | 2421 | 2617 | | | | | | | 2 2718 | |
| 493.69 | | | 2827 | | | 2903 | | 3630 | | 3718 | | | | | | | | 3 5623 | |
| 485.49 | | | 593 | | | 674 | | | | | | | | | | | | 6 1234 | |
| 471.29 | | | | | | 195 | | | | | | | | | | | | | |
| 469 10 | | | | | | 2145 | | | | | | | | | | | | 3 3557 6 239 | |
| 460 90 452 71 | | | | | | 1276 | | | | | | | | | | | | | |
| 452.71 444.51 | | | | | | | | | | | | | | | | | | 8 59% | |
| 436 31 | | | | | | | | | | 25.73 | . ,,,,,, | , ,,,,,,, | | | | | | | , |
| 430 31 | | | | 3071 | | | | | | | | | | | | | | | |

a - The column headed cm⁻¹ contains the wavenumber of the first $E_m(\widetilde{\nu})$ value in the row. The

columns headed XE and YE contain the X-exponent and the Y-exponent, respectively, for the row. The columns headed $0,1,2,\cdots 16$, contain the ordinate values, and the headings give the indices of the ordinate values in the row. In a row which starts with $\widetilde{v}(0)$, the wavenumber corresponding to the ordinate indexed J is $\widetilde{v}(J) = \widetilde{v}(0) - \frac{15798.002}{16384} \cdot J \cdot 2^{\text{XE}}$. The $E_{\text{m}}(\widetilde{v})$ values in that row are the ordinate value shown times 10^{YE} . Thus the entry indexed 16 in the first row of the table shows that $E_{\text{m}} = 372 \times 10^{-6} = 3.72 \times 10^{-4}$ at $\widetilde{v} = 6499.90 - \frac{15798.002}{16384} \cdot 16 \cdot 2^3 = 6376.48 \text{ cm}^{-1}$.

b - The E_m(v) values in the table can be interpolated to the original wavenumber spacing, 0.482117 cm⁻¹, and yield the original E_m(v) values accurate to 1% below 4500 cm⁻¹, 2% between 4500 and 5000 cm⁻¹ and 5% above 5000 cm⁻¹, via the 4-point spline interpolation program TRECOVER⁸.
c - The unit of E_m is L mole⁻¹ cm⁻¹. Multiply the values by 1000 to convert to the unit cm² mol⁻¹.

areas under its bands, and the real and imaginary refractive index spectra will be proposed to Commission I.5 of the International Union of Pure and Applied Chemistry for consideration as secondary infrared intensity standards³.

2.4 - The accuracy of the results

As noted previously¹, the purpose of this experimental study was to determine the accuracy of the measurements, rather than to obtain the highest possible precision. One must reveal the errors in the measurements in order to reveal accuracy, and to this end as few parameters as possible were kept constant in this work, as detailed previously¹. A consequence is that the 95% confidence limits for the studies by VB, CDK and YA include the effects of factors that are usually kept constant when high precision is the goal. Thus they include errors that are usually systematic, as well as the random errors on which the use of confidence limits to indicate precision is based. They are therefore expected to reflect the accuracy of the measurements better than is usually the case. The comparison with the measurements from other laboratories should reveal

Table 2.10 - Overall average areas under molar absorption coefficient bands of liquid toluene at 25°C.

| Region (cm ⁻¹) | Area | Estimated accuracy of area ^{a,b} | Area above baseline ^{4,0} | Estimated accuracy of area above baseline ^{a,d} |
|----------------------------|-------|---|------------------------------------|--|
| 6307.1 - 5445.0 | 132 | ± 9 | 121 | ± 6 |
| 4763.8 - 4518.9 | 89.5 | ± 0.5 | 59.2 | ± 0.2 |
| 4478.4 - 4145.2 | 196.8 | ± 1.0 | 103.1 | ± 0.4 |
| 4145.2 - 3988.1 | 160.7 | ± 1.9 | 99.2 | ± 0.6 |
| 3988.1 - 3748.9 | 99.6 | ± 1.6 | 26.13 | ± 0.06 |
| 3748.9 - 3694.9 | 13.73 | ± 0.11 | 2.266 | ± 0.004 |
| 3694.9 - 3608.2 | 29.4 | ± 0.4 | 10.9 | ± 0.1 |
| 3608.2 - 3569.1 | 8.13 | ± 0.17 | 1.30 | ± 0.02 |
| 3569.1 - 3531.5 | 6.86 | ± 0.12 | 1.45 | ± 0.01 |
| 3531.5 - 3503.1 | 4.20 | ± 0.08 | 0,490 | ± 0.005 |
| 3484.3 - 3418.7 | 11.45 | ± 0.14 | 2.064 | ± 0.007 |
| 3418.7 - 3357.0 | 11.64 | ± 0.12 | 1.344 | ± 0.002 |
| 3150.1 - 2770.2 | 4830 | ± 106 | 4535 | ± 100 |
| 2759.6 - 2679.1 | 65.6 | ± 0.5 | 29.9 | ± 0.1 |
| 2679.1 - 2563.4 | 53.5 | ± 0.3 | 20.2 | ± 0.1 |
| 2563.4 ~ 2131.0 | 171.8 | ± 2.1 | 87.0 | ± 0.9 |
| 2131.4 - 2095.8 | 5.22 | ± 0.28 | 0.92 | ± 0.04 |
| 2095.8 - 2049.5 | 6.69 | ± 0.39 | 1.01 | ± 0.05 |
| 1977.6 - 1910.1 | 143.8 | ± 0.9 | 104.4 | ± 0.6 |
| 1910.1 - 1836.9 | 143.2 | ± 1.1 | 85.3 | ± 0.7 |
| 1836.9 - 1754.9 | 139.7 | ± 1.4 | 70.2 | ± 0.6 |
| 1754.9 - 1712.5 | 52.97 | ± 0.4 | 26.4 | ± 0.2 |
| 1713.0 - 1686.4 | 17.52 | ± 0.46 | 1.71 | ± 0.04 |
| 1686.4 - 1668.1 | 13.41 | ± 0.34 | 2.15 | ± 0.05 |
| 1650.3 - 1555.3 | 497 | ± 3 | 320 | ± 1 |
| 1531.7 - 1516.3 | 90.9 | ± 3.3 | 14.3 | ± 0.4 |
| 1513.3 - 1400.1 | 1730 | ± 12 | 1201 | ± 6 |
| 1400.5 - 1338.4 | 234.0 | ± 1.6 | 78.6 | ± 0.5 |
| 1265.6 - 821.0 | 1266 | ± 15 | 891 | ± 9 |
| 790.7 - 775.7 | 36.7 | ± 0.4 | 10.5 | ± 0.1 |
| 769.9 - 710.2 | 3119 | ± 250 | 2823 | ± 223 |
| 710.2 - 660.0 | 1135 | ± 87 | 911 | ± 69 |

Table 2.10 - Continued.

| Region (cm ⁻¹) | Area* | Estimated accuracy of area a.b | Area above baseline ^{a,c} | Estimated accuracy of area above baseline ^{a,d} |
|----------------------------|-------|--------------------------------|---------------------------------------|--|
| 640.2 - 606.5 | 20.8 | ± 1.4 | 8.58 | ± 0.10 |
| 552.0 - 498.5 | 27.5 | ± 0.3 | 16.48 | ± 0.07 |
| 490.3 - 440.2 | 480 | ± 36 | 456 | ± 34 |

a - Unit: L mol⁻¹ cm⁻². Divide the values by 100 to convert to the unit km mol⁻¹.

systematic errors associated with different instrument design, different laboratory, different environment, different cells, different samples, and different person.

2.4.1 - Accuracy of absorption index values.

For the above reasons, we believe the agreement between the results of the different spectroscopists gives a good indication of the accuracy of our final $k(\widetilde{\nu})$ values. We use the maximum deviation from the unweighted mean to express this agreement. However, this is not a complete description of the accuracy because we introduce a systematic error by requiring all spectra to have the same $k(\widetilde{\nu})$ value at each anchor point ^{1,4}. Accordingly, we define the estimated accuracy of our final average $k(\widetilde{\nu})$ values as the maximum deviation from the unweighted mean plus the uncertainty in the $k(\widetilde{\nu})$ values at the anchor points. The uncertainty in $k(\widetilde{\nu})$ at the anchor points (column 5 of

b - Calculated from the percent estimated accuracy of the area, which is the same for the $E_m(\tilde{\nu})$ and $k(\tilde{\nu})$ (Table 2.5) bands.

c. The baseline is a straight line drawn through the molar absorption coefficient value at each end of the integration range.

d - Calculated by scaling the maximum deviation from the corresponding area in the $k(\tilde{\nu})$ spectrum (Table 2.5), by the ratio of the $E_m(\tilde{\nu})$ and $k(\tilde{\nu})$ areas.

Table 2. 11 - Overall average peak heights in the molar absorption coefficient spectrum of liquid toluene at 25°C.

| • | | | | | | | |
|-------------------------------------|-------------------------------------|-----------------------|-----------------------------------|-----------------------|-----------------------------------|-----------------------|-----------------------------------|
| \widetilde{v} (cm ⁻¹) | $E_{\mathfrak{m}}(\widetilde{\nu})$ | v (cm ⁻¹) | $E_{\mathfrak{m}}(\widetilde{v})$ | v (cm ⁻¹) | $E_{\mathfrak{m}}(\widetilde{v})$ | v (cm ⁻¹) | $E_{\mathfrak{m}}(\widetilde{v})$ |
| 5949.9 | 0.622 (10) | 3584.6 | 0.265 (4) | 2312.6 | 0.837 (3) | 1460.3 | 23.6 (16) |
| 4667.0 | 0.705 (13) | 3549.9 | 0.224 (4) | 2280.7 | 0.507 (7) | 1378.9 | 10.4 (4) |
| 4637.2 | 0.615 (12) | 3519.3 | 0.173 (3) | 2260.5 | 0.574 (14) | 1332.0 | 1,60 (4) |
| 4612.1 | 0.924 (15) | 3439.7 | 0.253 (4) | 2237.3 | 0.258 (6) | 1312.8 | 1.68 (4) |
| 4573.6 | 0.550 (7) | 3385.4 | 0.214(1) | 2207.5 | 0.351 (12) | 1277.7 | 0,863 (51 |
| 4388.8 | 0.775 (6) | 3167.5 | 0.893 (7) | 2185.3 | 0.353 (7) | 1248.7 | 1.32 (4) |
| 4311.0 | 0.871 (6) | 3104.2 | 6.63 (9) | 2163.7 | 0.621 (12) | 1210.2 | 3.10 (10) |
| 4244.4 | 0.840 (3) | 3086.4 | 20.9 (4) | 2116.6 | 0.183 (8) | 1178.6 | 6.64 (27) |
| 4186.4 | 0.426 (6) | 3062.1 | 24.3 (7) | 2068.5 | 0.198 (9) | 1156.0 | 2.76 (5) |
| 4161.3 | 0.392 (4) | 3026.9 | 54.3 (12) | 2032.0 | 0.200 (9) | 1106.5 | 4.34 (6) |
| 4132.4 | 0.420 (7) | 2979.2 | 14.4 (2) | 2008.6 | 0.420 (9) | 1081.4 | 19.0 (4) |
| 4056.7 | 1.92 (5) | 2947.7 | 17.5 (2) | 1991.1 | 0.814 (11) | 1041.5 | 9.16 (12) |
| 4036.3 | 2.28 (1) | 2919.8 | 27.9 (5) | 1942.1 | 4.91 (5) | 1002.3 | 2.63 (8) |
| 3980.3 | 0.504 (9) | 2872.6 | 13.4 (2) | 1872.1 | 2.92 (1) | 980.8 | 2.44 (8) |
| 3951.2 | 0.489 (9) | 2734.1 | 2.75 (5) | 1857.6 | 4.65 (7) | 966.6 | 2.25 (8) |
| 3923.8 | 0.495 (7) | 2671.7 | 0.357 (5) | 1802.6 | 4.06 (5) | 929.6 | 1.60 (5) |
| 3909.9 | 0.492 (9) | 2632.0 | 0.502 (5) | 1778.5 | 1.27 (1) | 895.4 | 4.79 (21) |
| 3869.8 | 0.504 (7) | 2604.7 | 0.633 (13) | 1735.6 | 2.23 (1) | 873.0 | 1.16 (17) |
| 3847.2 | 0.564 (14) | 2585.9 | 1.16 (1) | 1696.8 | 0.811 (17) | 842.8 | 1.81 (16) |
| 3812.2 | 0.389 (7) | 2540.4 | 0.410 (9) | 1676.7 | 0.910 (16) | 785.6 | 3.84 (17) |
| 3786.3 | 0.394 (7) | 2509.2 | 0.373 (5) | 1623.1 | 4.06 (6) | 728.9 | 305 (25) |
| 3764.0 | 0.238 (9) | 2496.8 | 0.368 (7) | 1604.5 | 28.9 (11) | 694.5 | 148 (8) |
| 3724.2 | 0.305 (4) | 2465.5 | 0.369 (5) | 1586.7 | 4.57 (6) | 622.1 | 1.19 (4) |
| 3707.0 | 0.300 (4) | 2412.4 | 0.740 (7) | 1572.2 | 4.76 (3) | 537.8 | 0.410 (6 |
| 3675.1 | 0.313 (10) | 2389.0 | 0.611 (9) | 1550.4 | 3.14 (14) | 521.0 | 2.17 (11 |
| 3649.0 | • • | 2360.6 | 0.968 (17) | 1523.6 | 7.70 (3) | 464.2 | 79.9 (105 |
| 3624.0 | • • | 2335.3 | 1.08 (2) | 1495.6 | 97.4 (33) | | |

a - The unit of $E_m(\tilde{\nu})$ is L mole⁻¹ cm⁻¹. Multiply the values by 1000 to convert to the unit cm² mol⁻¹. The numbers in parentheses are the estimated accuracies in the last digit. The percent estimated accuracy is the same as that of the corresponding $k(\tilde{\nu})$ peak height.

Table 2.1) is $\sim \pm 1 \times 10^{-6}$ above 1600 cm⁻¹, and $\sim \pm 1 \times 10^{-5}$ below 1600 cm⁻¹. The estimated accuracy of $k(\widetilde{\nu})$ is expressed as a percentage of $k(\widetilde{\nu})$. The percent estimated accuracy of the $E_m(\widetilde{\nu})$ values is the same as that of the $k(\widetilde{\nu})$ values.

The percent estimated accuracies of the peak $k(\tilde{v})$ values are included in Table 2.6. They are generally good, but are poor for the six peaks at 1460, 1278, 873, 843, 728, and 464 cm⁻¹. These peaks are discussed below, and are excluded from the following general comments. The average estimated accuracy is $\pm 2.5\%$ for the heights of the 39 strong, medium and weak peaks below 4100 cm⁻¹ which have $k_{\text{max}} > 0.002$. The average estimated accuracy is $\pm 1.9\%$ for the heights of the 51 very weak peaks below 4100 cm⁻¹ which have $k_{\text{max}} < 0.002$. The average estimated accuracy is $\pm 1.3\%$ for the heights of the 11 very weak peaks above 4100 cm⁻¹, but the peaks above 4500 cm⁻¹ were measured only in this laboratory.

The heights of the peaks at 1460, 1278, 873, 843, 728, and 464 cm⁻¹ have estimated accuracies of 6.9, 6.0, 14.7, 9.1, 8.3, and 13.2%, respectively. There is no obvious reason why the peaks at 1460 and 843 cm⁻¹ should have such large uncertainties. The peaks at 1278 and 873 cm⁻¹ are very weak and form part of the baseline of surrounding peaks that are more intense. The peak at 728 cm⁻¹ was poorly determined. It is very strong, requires cells of ≤ 8 µm path, and only two spectra from one spectroscopist were available. The peak at 464 cm⁻¹ is near the transmission limit of the KBr windows but no reason for the poor accuracy is evident. It should be noted that the height of this 464 cm⁻¹ peak reported here is about 2.5 times that reported by Jones and co-workers⁷.

The peak $k(\tilde{v})$ values must be compared with the only calibrated results available, those of Jones and coworkers in Ref. 7. Of the 94 k_{max} values in Table 2.6 for which

there is a calibrated value, 29 lie outside the evaluated uncertainty limits of Ref. 7. They are at the wavenumbers 3675, 3550, 3519, 3385, 3168, 3027, 2979, 2948, 2872, 3361, 2164, 2032, 1942, 1858, 1572, 1524, 1460, 1378, 1332, 1313, 1107, 1041, 895, 843, 729, 695, 538, 521, 464 cm⁻¹. The three values at 1460, 843, and 464 cm⁻¹ had unusually poor accuracy in this work. Only the 13 peaks underlined in the above list disagree by more than the combined error limits, and only the 464 cm⁻¹ peak is in serious disagreement.

The $k(\widetilde{\nu})$ values in the baseline are not known as accurately as those in regions of significant absorption, except at the anchor points. From the agreement between different workers and the uncertainties in the anchor point values, we estimate the accuracy of the baseline $k(\widetilde{\nu})$ values to be $\pm \sim 2 \times 10^{-6}$ above 1600 cm^{-1} , and $\pm \sim 1.5 \times 10^{-5}$ below 1600 cm^{-1} . This corresponds to between ± 3 and $\pm 10\%$ of the $k(\widetilde{\nu})$ values in the baseline.

2.4.2 - Accuracy of areas.

The percent estimated accuracies of the areas under the $k(\tilde{v})$ bands are given in Table 2.5. Each was calculated by multiplying the uncertainty in $k(\tilde{v})$ at the anchor points (Table 2.1) by the integration range, adding the maximum deviation of the spectroscopist averages from the unweighted average (Table 2.5), and expressing the result as a percentage of the area. As was the situation for benzene¹, the uncertainty at the anchor points contributes about 0.5% to the percent estimated accuracies. The

percent estimated accuracy of the area under an $E_m(\tilde{\nu})$ band equals that of the corresponding $k(\tilde{\nu})$ band. The estimated accuracies, rather than the percent estimated accuracies, of the $E_m(\tilde{\nu})$ bands are given in Table 2.10.

The average accuracy of all of the band areas is $\pm 2.4\%$ and it is an excellent $\pm 1.2\%$ for the eleven regions between 3150 and 775 cm⁻¹ which contain a band with $0.002 < k_{max} < 0.112$.

The area we prefer for use as a secondary standard is the area under the molar absorption coefficient spectrum above a linear baseline drawn through the $E_{\rm m}$ values at the integration limits. This area is given in Table 2.10 with its estimated accuracy. The percent estimated accuracy of this area is the same as the maximum deviation of a spectroscopist's average area under a $k(\tilde{\nu})$ band from the unweighted average (Table 2.5) expressed as a percentage of the average.

2.4.3 - Accuracy of real refractive index values.

Three factors contribute to the accuracy of our $n(\tilde{v})$ values. First, our Kramers-Kronig transform procedure has an intrinsic accuracy of about $0.05\%^{11}$. Second, we used $n(8000 \text{ cm}^{-1}) = 1.4773 \pm 0.0004$, an accuracy of 0.03%. Third, the approximately 2.5% accuracy of the absorption indices yields an approximately 2.5% accuracy in the values of $\Delta n = n(\tilde{v})$ - $n(8000 \text{ cm}^{-1})$ from the Kramers-Kronig transform, which corresponds to an error of $\sim 0.08\%$ in $n(\tilde{v})$. The sum of these contributions gives

~0.2% as the estimated accuracy of our $n(\tilde{v})$ values.

2.5 - Summary

Transmission measurements by 6 spectroscopists in 4 laboratories have been recorded on four different instruments made by three different manufacturers. They have been converted to absorption index spectra which were compared and averaged, and the real refractive index and molar absorption coefficient spectra have been calculated from the average. The percent estimated accuracy of the absorption index and molar absorption coefficient values is on average ±2.5% at the peaks of the strong, medium and weak bands with $0.002 < k_{\text{max}} < 0.112$, and $\pm 1.9\%$ and $\pm 1.3\%$ at the peaks of the very weak bands below and above 4100 cm⁻¹, respectively. The baseline values are accurate to between 3 and 10%. The percent estimated accuracy of areas under bands or groups of bands in $k(\tilde{v})$ and $E_m(\tilde{v})$ spectra is $\pm 2.4\%$ on average, and averages $\pm 1.2\%$ for the bands between 3150 and 775 cm⁻¹ with 0.002 $k_{\text{max}} < 0.112$. The estimated accuracy of the real refractive index values is 0.2%. For long-term reference, the complete numerical data are presented in Compact Table format⁸, which is readable and allows the original spectra to be recovered by interpolation without loss of accuracy. The complete final $k(\tilde{v})$, $n(\tilde{v})$, and $E_m(\tilde{v})$ spectra obtained in this work are available on diskette from the authors. To provide continuity over the longer term, it is anticipated that they will be made available in the future on an internationally accessible data base.

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Chapter 3 - Accurate Optical Constants and Molar Absorption Coefficients

Between 4800 and 450 cm⁻¹ of Chlorobenzene at 25°C from Spectra Recorded in

Several Laboratories*

3.1 - Introduction

In previous papers^{1,2} we have reported absolute infrared absorption intensities of benzene and toluene. The quantities reported have been the spectra of the real and imaginary refractive indices and the molar absorption coefficient, and the areas under bands in the imaginary refractive index and molar absorption coefficient spectra. The imaginary refractive index is also called the absorption index³. In this paper we report these same quantities between 4800 and 450 cm⁻¹ for chlorobenzene at 25°C. The methods used were the same as have been described in detail for the spectra of benzene^{1,4} and toluene².

For selected bands, the refractive indices, molar absorption coefficients, and areas under molar absorption coefficient bands, will be submitted to Commission I.5 of the International Union of Pure and Applied Chemistry for consideration as secondary infrared intensity standards for liquids⁵.

^{*} A version of this chapter has been published. Bertie, Jones and Apelblat, Appl. Spectrosc., 48, 144 (1994).

3.2 - Methods and results

The chlorobenzene in this laboratory was of spectroscopic or reagent grade.

Samples were purified by fractional freezing one to three times and were checked by gas chromatography and infrared spectroscopy. No impurities were detected. The samples were kept over molecular sieve to ensure dryness.

Experimental absorbance spectra of chlorobenzene were obtained from fixed path length cells with KBr windows and path lengths between 11 and 1500 μm and from a variable path length cell with NaCl windows and path lengths between 100 and 500 μm. Spectra were used only in the regions where the absorbances of the strongest bands were between 0.3 and 2.0.

To determine the linear absorption coefficients at the anchor points⁴, spectra with strong baseline absorbance were also obtained in cells with path lengths from 50 μ m to 3.5 mm. Table 3.1 summarizes the wavenumbers of the anchor points, the path lengths of the cells, the measured linear absorption coefficients and their 95% confidence limits, and the uncertainty in the absorption index, $k(\tilde{\nu})$, that results from the precision of the linear absorption coefficients.

In addition to the experimental absorbance spectra recorded in our laboratory, spectra recorded in this laboratory in 1984-5 by V. Behnam, and reported previously^{6,7}, were also reprocessed by our current improved methods to yield the refractive indices. In addition, three spectroscopists in three other laboratories kindly recorded and supplied experimental absorbance spectra of chlorobenzene. One set was recorded with 2 cm⁻¹

Table 3.1 - Linear Absorption coefficients at the anchor points of liquid chlorobenzene at 25°C.

| Wavenumber | cell path lengths | | 95% confidence | Uncertainty in |
|---------------------|-------------------|--|----------------|------------------------|
| (cm ⁻¹) | (mm) | $K(\widetilde{v})$ (cm ⁻¹) | limit | k(v) |
| 4795.1 | 1.5 - 3.5 | 0.32 | 0.09 | 3.5 x 10 ⁻⁶ |
| 4475.0 | 1.5 - 3.5 | 0.617 | 0.021 | 8.6 x 10 |
| 3849.2 | 1.5 - 3.5 | 1.303 | 0.031 | 1.5×10^{-6} |
| 3333.3 | 1.5 - 3 5 | 0.682 | 0.025 | 1.4×10^{-6} |
| 2931.2 | 1.3 - 3.5 | 3,152 | 0.021 | 1.3 x 10 ⁻⁶ |
| 1908.2 | 0.7 - 3.5 | 4.723 | 0.017 | 1.6 x 10 ⁻⁶ |
| 1835.9 | 0.7 - 3.5 | 4.428 | 0.021 | 2.1 x 10 ⁻⁶ |
| 1754.9 | 0.7 - 3.5 | 5.846 | 0.018 | 1.9 x 10 ⁻⁶ |
| 1676.8 | 0.7 - 3.5 | 4.114 | 0.017 | 1.9 x 10 ⁻⁶ |
| 1630.5 | 0.5 - 1.5 | 11.64 | 0.03 | 3.6 x 10 ⁻⁶ |
| 1534.0 | 0.5 - 1.7 | 9.081 | 0.030 | 3.6 x 10 ⁻⁶ |
| 1404.8 | 0.5 - 1.7 | 7.056 | 0.032 | 4.2 x 10 ⁻⁶ |
| 1345.1 | 1.3 - 3.5 | 3.526 | 0.016 | 2.2 x 10 ⁻⁶ |
| 1251.5 | 0.5 - 3.5 | 6.235 | 0.025 | 3.7×10^{-6} |
| 1197.5 | 0.5 - 1.7 | 9.545 | 0.020 | 3.1×10^{-6} |
| 1141.6 | 0.2 - 0.7 | 17.65 | 0.10 | 1.7×10^{-5} |
| 1045.2 | 0.2 - 1.3 | 16.52 | 0.09 | 1.6 x 10 ⁻⁵ |
| 949.8 | 0.5 - 1.5 | 14.05 | 0.03 | 6.0 x 10 ⁻⁶ |
| 801.3 | 0.2 - \$.3 | 12.66 | 0.08 | 1.7 x 10 ⁻⁵ |
| 717.4 | ~ 0.05 | 65.7 | 0.6 | 1.5 x 10 ⁻⁴ |
| 630.6 | 0.5 - 1.5 | 5.46 | 0.06 | 1.6 x 10 ⁻⁵ |
| 528.4 | ~ 1.5 | 2.08 | 0.05 | 1.6 x 10 ° |
| 437.8 | 0.5 - 1.5 | 7.47 | 0.20 | 8.5 x 10 ⁻⁵ |

nominal resolution on a Nicolet 170SX spectrometer with a mercury cadmium telluride detector at 77K and Happ-Genzel apodization. For that work, the chlorobenzene was HPLC grade. The second set was recorded with 2 cm⁻¹ nominal resolution on a Nicolet 510P spectrometer with room-temperature deuterated triglycine sulfate detector and

Table 3.2 - Pathlengths, high-wavenumber refractive index, and number of spectra from each spectroscopist, for the regions processed.

| Region (cm ⁻¹) | Pathlengths (µm) | ո տ | YA | VB | Α | В | c | Total |
|-------------------------------|------------------|------------|----|----|----|----|---|-------|
| 4800 - 3990 | 200-700 | 1.501 | 18 | 5 | 0 | 1 | 2 | 26 |
| 4000 - 3325 | 500-700 | 1.503 | 18 | 5 | 0 | 0 | 0 | 23 |
| 3335 - 2925 | 30-100 | 1.495 | 24 | 6 | 15 | o | 0 | 45 |
| 2940 - 1620 | 100-200 | 1.496 | 5 | 5 | 5 | 1 | 0 | 16 |
| 1640 - 1390 | 11-13 | 1.481 | 9 | 6 | 0 | 0 | 0 | 15 |
| 1415 - 1130 | 200-700 | 1.487 | 13 | 5 | 0 | 1 | 2 | 21 |
| 1150 - 935 | 11-13 | 1.473 | 5 | 6 | 0 | O | 0 | 11 |
| 960 - 790 | 25-100 | 1.493 | 14 | 6 | 15 | 1 | 2 | 38 |
| 810 - 705 | ~8 | 1.450 | 0 | 0 | 5 | 0 | 0 | 5 |
| 725 - 610 | 8-13 | 1.596 | 5 | 4 | 5 | 0 | 0 | 14 |
| 635 - 520 | 100-500 | 1.558 | 7 | 3 | 0 | 1 | 0 | 11 |
| 535 + 420 | 11-13 | 1.520 | 5 | _3 | 0 | 0_ | 0 | 8 |

a - The spectroscopists are identified as YA, VB, A, B and C (see text).

Happ-Genzel apodization. The chlorobenzene was 99%+ grade. No experimental details are available for the third set, but they were clearly run under comparable conditions. None of the spectra showed unexpected peaks.

Table 3.2 shows the spectral regions that were used in the computations, together with the cell thicknesses used, the value of n_{∞} , and the number of spectra from each spectroscopist, for each region. The value of n_{∞} for each region is required by the Kramers-Kronig transform in program RNJ46A^{1,4}. The spectra recorded in this laboratory are labeled YA for Y. Apelblat, and VB for V. Behnam. The other collaborators are identified by A, B and C. The real refractive index at the highest

wavenumber in each region, $n_{\cdot \cdot}$, was taken from tables of $n(\vec{v})$ in this laborator which were calculated from the $k(\vec{v})$ spectra in reference 7.

3.2.1 Absorption index spectrum.

The experimental absorbance spectra of each spectroscopist were converted to absorption index, $k(\tilde{v})$, spectra by program RNJ46A^{1,4}. The peak heights, wavenumbers and areas were measured for each $k(\tilde{v})$ spectrum. For each region (Table 3.2), the $k(\tilde{v})$ spectra were averaged to give a single spectrum from each spectroscopist. Table 3.3 gives the peak wavenumber of each band and, for each spectroscopist, the average peak heights with their 95% confidence limits. The peak heights from different spectroscopists are in good agreement although, as was found for benzene and toluene, the existence of systematic errors at the few percent level is indicated by the disagreement between spectroscopists sometimes being outside the combined confidence limits.

For each region, the average spectra from the different spectroscopists were themselves averaged to yield an unweighted average $k(\widetilde{\nu})$ spectrum. The unweighted average peak heights are given in Table 3.4 with the maximum deviation of any spectroscopist's average from the unweighted average in parentheses. A weighted average $k(\widetilde{\nu})$ spectrum was also calculated, with the weighting factor from the number of spectra which contributed to the spectroscopist's average. The weighted average peak heights are listed in Table 3.4 with their 95% confidence limits in parentheses. For

Table 3.3 - Spectroscopist average absorption index peak heights.

| $\tilde{\nu}$ (cm ⁻¹) | YA | VB | Α | В | С |
|-----------------------------------|--------------|--------------|-------------|----------|----------|
| 4656.4 | 0.000495(38) | 0.000504(26) | | 0.000497 | |
| 4622.0 | 0.000163(12) | 0.000168(10) | | 0.000168 | |
| 4595,3 | 0.000108(8) | 0.000110(7) | | 0.000107 | |
| 4574.7 | 0.000243(14) | 0.000251(10) | | 0.000244 | |
| 4514.6 | 0.000075(3) | 0.000076(2) | | 0.000075 | |
| 4252.7 | 0.000111(1) | 0.000111(1) | | 0.000109 | 0.000105 |
| 4145.7 | 0.000271(3) | 0.000271(1) | | 0.000268 | 0.000262 |
| 4077.4 | 0.00120(10) | 0.00117(5) | | 0.00111 | 0.00105 |
| 4066.5 | 0.00108(5) | 0.00115(4) | | 0.00110 | 0.00105 |
| 4058.7 | 0.00129(14) | 0.00120(6) | | 0.00118 | 0.00112 |
| 4000.5 | 0.000253(4) | 0.000253(2) | | 0.000249 | 0.900246 |
| 3917.8 | 0.000103(2) | 0.000102(1) | | | |
| 3899.6 | 0.000109(1) | 0.000109(1) | | | |
| 3785.1 | 0.000354(4) | 0.000355(2) | | | |
| 3772.3 | 0.000265(2) | 0.000260(1) | | | |
| 3699.4 | 0.000521(7) | 0.000532(6) | | | |
| 3653.3 | 0.000104(2) | 0.000107(1) | | | |
| 3636.9 | 0.000137(2) | 0.000138(1) | | | |
| 3613.6 | 0.000143(2) | 0.000145(1) | | | |
| 3488.8 | 0.000080(1) | 0.000078(1) | | | |
| 3464.4 | 0.000077(1) | 0.000075(1) | | | |
| 3368.0 | 0.000065(1) | 0.000065(1) | | | |
| 3165.5 | 0.000658(7) | 0.000664(2) | 0.000671(7) | | |
| 3083.1 | 0.00659(2) | 0.00682(3) | 0.00671(3) | | |
| 3069.5 | 0.00981(2) | 0.01007(5) | 0.00989(5) | | |
| 3058.5 | 0.00811(2) | 0.00793(5) | 0.00793(5) | | |
| 3025.8 | 0.00316(1) | 0.00319(2) | 0.00318(1) | | |
| 3016.4 | 0.00279(1) | 0.00273(1) | 0.00275(1) | | |
| 2949.5 | 0.000338(4) | 0.000337(2) | 0.000343(2) | | |
| 2919.0 | 0.000439(3) | 0.000426(1) | 0.000417(1) | 0.000425 | |
| 2885.6 | 0.000270(6) | 0.000277(1) | 0.000274(1) | 0.000268 | |
| 2877.0 | 0.000284(5) | 0.000279(1) | 0.000282(1) | 0.000275 | |
| 2849.2 | 0.000260(3) | 0.000250(1) | 0.000251(1) | 0.000250 | |
| 2795.1 | 0.000102(8) | 0.000099(2) | 0.000103(1) | 0.000093 | |
| 2772.9 | 0.000302(9) | 0.000303(1) | 0.000299(1) | 0.000293 | |
| 2756.0 | 0.000128(7) | 0.000126(1) | 0.000129(1) | 0.000121 | |
| 2739.0 | 0.000111(8) | 0.000111(1) | 0.000112(1) | 0.000104 | |
| 2714.2 | 0.000087(8) | 0.000089(1) | 0.000092(1) | 0.000084 | |

Table 3.3 - Continued

| 1 able 3.3 - | Continued | | | | |
|---------------------------------|-------------|-------------|--------------|----------|---|
| \tilde{v} (cm ⁻¹) | YA | VB | A | В | С |
| 2703.0 | 0.000094(5) | 0.000096(2) | 0.000098(1) | 0.000090 | |
| 2663.5 | 0.000460(6) | 0.000464(2) | 0.000470(1) | 0.000456 | |
| 2648.1 | 0.000302(8) | 0.000299(1) | 0.000302(1) | 0.000292 | |
| 2616.3 | 0.000426(5) | 0.000420(2) | 0.000418(1) | 0.000410 | |
| 2599.8 | 0.000409(7) | 0.000415(1) | 0.000419(1) | 0.000407 | |
| 2582.2 | 0.000336(5) | 0.000328(1) | 0.000324(1) | 0.000319 | |
| 2557.4 | 0.000295(5) | 0.000295(1) | 0.000302(1) | 0.000292 | |
| 2544.3 | 0.000335(6) | 0.000336(1) | 0.000335(1) | 0.000326 | |
| 2510.9 | 0.000103(6) | 0.000107(1) | 0.000106(1) | 0.000100 | |
| 2496.0 | 0.000146(6) | 0.000148(1) | 0.000152(1) | 0.000142 | |
| 2471.4 | 0.000198(6) | 0.000199(1) | 0.000201(%) | 0.000193 | |
| 2446.6 | 0.000152(8) | 0.000154(1) | 0.000155(1) | 0.000148 | |
| 2427.3 | 0.000195(5) | 0.000197(1) | 0.000201(1) | 0.000193 | |
| 2406.4 | 0.090337(4) | 0.000338(1) | 0.000339(1) | 0.000331 | |
| 2390.3 | 0,000452(7) | 0.000452(1) | 0.000453(1) | 0.000444 | |
| 2365.7 | 0.000193(5) | 0.000196(1) | 0.000210(12) | 0.000208 | |
| 2331.0 | 0.000766(5) | 0.000770(1) | 0.000787(8) | 0.000776 | |
| 2319.8 | 0.000781(7) | 0.000768(1) | 0.000781(3) | 0.000770 | |
| 2271.8 | 0.000318(6) | 0.000321(1) | 0.000320(1) | 0.000314 | |
| 2240.5 | 0.000571(7) | 0.000569(1) | 0.000566(2) | 0.000557 | |
| 2194.0 | 0.000187(7) | 0.000189(1) | 0.000185(1) | 0.000182 | |
| 2178.7 | 0.000360(6) | 0.000363(1) | 0.000355(1) | 0.000351 | |
| 2150.5 | 0.000145(6) | 0.000148(1) | 0.000145(1) | 0.000142 | |
| 2132.5 | 0.000086(5) | 0.000090(1) | 0.000085(1) | 0.000084 | |
| 2089.1 | 0.000140(6) | 0.000145(1) | 0.000139(1) | 0.000137 | |
| 2058.2 | 0.000131(5) | 0.000138(1) | 0.600131(1) | 0.000130 | |
| 2045.6 | 0.000149(6) | 0.000149(1) | 0.000143(1) | 0.000145 | |
| 2023.7 | 0.000249(5) | 0.000248(1) | 0.000236(1) | 0.000236 | |
| 2002.5 | 0.000279(5) | 0.000289(1) | 0.000284(1) | 0.000283 | |
| 1962.2 | 0.00207(1) | 0.00207(1) | 0.00209(1) | 0.00206 | |
| 1942.9 | 0.00410(3) | 0.00406(4) | 0.00413(1) | 0:00409 | |
| 1881.4 | 0.00240(1) | 0.00240(1) | 0.00240(1) | 0.00238 | |
| 1861.9 | 0.00441(4) | 0.00444(6) | 0.00443(1) | 0.00438 | |
| 1787.8 | 0.00354(3) | 0.00357(2) | 0.00358(1) | 0.00354 | |
| 1730.5 | 0.00345(2) | 0.00349(3) | 0.00346(1) | 0.00342 | |
| 1685.9 | 0.000663(8) | 0.000649(1) | 0.000646(1) | 0.000649 | |
| 1645.5 | 0.00229(1) | 0.00227(1) | 0.00227(1) | 0.00225 | |
| 1622.2 | 0.00270(3) | 0.00263(2) | | | |

| Table 3.3 - | Continued | | | | |
|---------------------------------|------------|------------|------------|---------|---------|
| \tilde{v} (cm ⁻¹) | YA | VB | A | B | C |
| 1583.8 | 0.0826(5) | 0.0803(4) | | | |
| 1566. l | 0.0149(1) | 0.0146(1) | | | |
| 1477.4 | 0.245(3) | 0.262(22) | | | |
| 1445.4 | 0.110(1) | 0.097(1) | | | |
| 1387.1 | 0.00303(1) | 0.00304(1) | | 0.00303 | 0.00304 |
| 1370.7 | 0.00303(1) | 0.00303(1) | | 0.00302 | 0.00304 |
| 1325.2 | 0.00252(1) | 0.00254(1) | | 0.00248 | 0.00254 |
| 1298.1 | 0.00377(2) | 0.00373(1) | | 0.00369 | 0.00383 |
| 1272.5 | 0.00366(5) | 0.00385(1) | | 0.00373 | 0.00377 |
| 1235.2 | 0.00278(1) | 0.00277(1) | | 0.00276 | 0.00278 |
| 1210.9 | 0.00233(1) | 0.00235(1) | | 0.00234 | 0.00236 |
| 1171.6 | 0.00522(1) | 0.00521(1) | | 0.00512 | 0.00527 |
| 1156.4 | 0.00459(1) | 0.00459(2) | | 0.00454 | 0.00466 |
| 1122.7 | 0.0265(1) | 0.0270(1) | | | |
| 1083.2 | 0.180(1) | 0.176(1) | | | |
| 1068.1 | 0.0529(2) | 0.0513(2) | | | |
| 1022.6 | U.197(1) | 0.175(3) | | | |
| 1001.9 | 0.0461(2) | 0.0426(2) | | | |
| 963.7 | 0.00384(4) | 0.00369(6) | | | |
| 935.0 | 0.00785(1) | 0.00784(2) | 0.00780(1) | 0.00837 | 0.00787 |
| 902.9 | 0.0315(1) | 0.0317(1) | 0.0312(1) | 0.0330 | 0.0320 |
| 866.3 | 0.00304(1) | 0.00302(5) | 0.00294(1) | 0.00291 | 0.00291 |
| 830.0 | 0.00585(2) | 0.00587(6) | 0.00576(1) | 0.00594 | 0.00554 |
| 812.2 | 0.00354(2) | 0.00352(4) | 0.00349(1) | 0.00353 | 0.00346 |
| 739.5 | | | 0.747(16) | | |
| 702.2 | 0.305(2) | 0.292(4) | 0.320(2) | | |
| 684.8 | 0.372(2) | 0,353(12) | 0.395(2) | • | |
| 661.9 | 0.0170(1) | 0.0171(4) | 0.0185(1) | | |
| 614.0 | 0.00847(5) | 0.00792(2) | | 0.00776 | |
| 547.7 | 0.00100(1) | 0.00098(1) | | 0.00093 | |
| 533.7 | 0.00141(1) | 0.00138(2) | | 0.00140 | |
| 467.8 | 0.289(3) | 0.275(6) | | | |

a The numbers in parentheses are the 95% confidence limits in the last digit.

reference, we include in Table 3.4 the peak heights of chlorobenzene published previously from this laboratory⁷ and the results of the only measurements that have been made against a primary standard⁸. The latter have an evaluated uncertainty⁸ of ~6%.

Table 3.4 - Overall average absorption index peak heights.

| ~ (cm⁻¹) | Weighted average | Unweighted average | Uncertainty due to anchor points | % estimated accuracy | Ref. 8 | Ref. 7 |
|-------------|---------------------|--------------------|----------------------------------|----------------------------|---------------|-------------|
| 4656.4 | 0.000497 (28) | 0.000499 (42) | 2.2 x 10 ⁻⁶ | 1.4 | | |
| 4622.0 | 0.000165 (9) | 0.000166 (3) | 2.2 x 10 ⁻⁶ | 3.1 | | |
| 4593.3 | 0.000108 (6) | 0.000108 (2) | 2.2 x 10 ⁻⁶ | 3.9 | | |
| 4574.7 | 0.000245 (10) | 0.000246 (5) | 2.2 x 10 ⁻⁶ | 2.9 | | |
| 4514.6 | 0.000075 (3) | 0.000076 (2) | 2.2 x 10 ⁻⁶ | 5.5 | | |
| 4252.7 | 0.000110(1) | 0.000109 (4) | 1.2 x 10 ⁻⁶ | 4.8 | | |
| 4145.7 | 0.000270 (2) | 0.000268 (6) | 1.2 x 10 ⁻⁶ | 2.7 | 0.000276 (21) | |
| 4077.4 | 0.00118 (7) | 0.00113 (8) | 1.2 x 10 ⁻⁶ | 7.2 | 0.00113 (5) | |
| 4066.5 | 0.00110 (3) | 0.00110 (5) | 1.2 x 10 ⁻⁶ | 4.7 | 0.00115 (5) | |
| 4058.7 | 0.00125 (9) | 0.00120 (9) | 1.2 x 10 ⁻⁶ | 7.6 | 0.00112 (5) | |
| 4000.5 | 0.000252 (3) | 0.000250 (4) | 1.2 x 10 ⁻⁶ | 2.1 | 0.000252 (19) | |
| 3917.4 | 0.000103 (2) | (1) د 1000.0 | 1.2 x 10 ⁻⁶ | 2.1 | 0.000110 (9) | 0.000103 (1 |
| 3899.6 | 0.000109(1) | 0.000109(1) | 1.2 x 10 ⁻⁶ | 2.0 | 0.000115 (9) | 0.000109 (1 |
| 3785.1 | 0.000354 (3) | 0.000354(1) | 1.5×10^{-6} | 0.7 | 0.000352 (28) | 0,000356 (2 |
| 3772.3 | 0.000264 (2) | 0.000263 (3) | 1.5 x 10 ⁻⁶ | 1.7 | 0.000270 (20) | 0.000261 (2 |
| 3699.4 | 0.000524 (6) | 0.000527 (6) | 1.5×10^{-6} | 1.4 | 0.000541 (37) | 0,000532 (4 |
| 3653.3 | 0.000105 (1) | 0.000106 (2) | 1.5×10^{-6} | 3.3 | 0.000109 (9) | 0.000107 (2 |
| 3636.9 | 0.000137 (1) | 0.000138 (1) | 1.5×10^{-6} | 1.8 | 0.000144 (11) | 0.000138 (2 |
| 3613 | 143 (2) | 0.000144 (1) | 1.5×10^{-6} | 1.7 | 0.000146 (11) | 0.000145 (1 |
| 3488.8 | (1) 079د.۔ | 0.000079 (1) | 1.5×10^{-6} | 3.2 | 0.000083 (7) | 0.000077 (|
| 3464.4 | 0.000077 (1) | 0.000076 (1) | 1.5×10^{-6} | 3.3 | 0.000082 (7) | 0,000075 (|
| 3368.0 | 0.000065 (1) | 0.000065 (1) | 1.5 x 10 ⁻⁶ | 3.8 | 0.000071 (7) | 0,000064 (|
| 3165.5 | 0.000663 (4) | 0.000664 (7) | 1.4×10^{-6} | 1.3 | 0.000709 (32) | |
| 3083.1 | 0.00666 (3) | 0.00671 (12) | 1.4 x 10 ⁻⁶ | 1.8 | 0.00670 (30) | 0,00689 (2 |
| 3069.5 | 0.00987 (3) | 0.00992 (15) | 1.4 x 10 ⁻⁶ | 1.5 | 0.00991 (20) | 0.01015 (4 |
| 3058.5 | | 0.00799 (12) | 1.4×10^{-6} | 1.5 | 0.00796 (17) | 0,00800 (3 |
| 3025.8 | | 0.00318 (2) | 1.4 x 10 ⁻⁶ | 0.7 | 0.00321 (14) | 0.00325 (1 |
| 3016.4 | | 0.00276 (3) | 1.4 x 10 ⁻⁶ | 1.1 | 0.00278 (13) | 0.00279 (1 |
| 2949.5 | • • | 0.000339 (3) | 1.4×10^{-6} | 1.3 | 0.000364 (27) | |
| 2919.0 | • • | 0.000427 (12) | 1.5 x 10 ⁻⁶ | 3.2 | 0.000444 (35) | |

Table 3.4 - Continued

| Table 3.4 | Collandod | | | | | |
|---------------------------------------|---------------------|-----------------------|----------------------------------|----------------------|---------------|--------------|
| ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~ | Weighted average | Unweighted average | Uncertainty due to anchor points | % estimated accuracy | Ref. 8 | Ref. 7 |
| 2885.6 | 0.000274 (2) | 0.000273 (5) | 1.5 x 10 ⁻⁶ | 2.4 | 0.000300 (22) | |
| 2877.0 | 0.000274 (2) | 0.000280 (5) | 1.5 x 10 ⁻⁶ | 2.3 | 0.000300 (22) | |
| 2849.2 | 0.000281 (2) | 0.000253 (7) | 1.5 x 10 ⁻⁶ | 3,4 | 0.000268 (20) | |
| 2795.1 | 0.000101 (2) | 0.000099 (4) | 1.5 x 10 ⁻⁶ | 5.6 | 0.000117 (10) | |
| 2772.9 | 0.000301 (2) | 0.000299 (6) | 1.5 x 10 ⁻⁶ | 2.5 | 0.000301 (22) | |
| 2756.0 | 0.000301 (2) | 0.000126 (5) | i.5 x 10 ⁻⁶ | 5.2 | 0.000148 (11) | |
| 2739.0 | 0.000127 (2) | 0.000129 (5) | 1.5 x 10 ⁻⁶ | 6.0 | 0.000128 (10) | |
| 2714.2 | 0.000089 (2) | 0.000088 (4) | 1.5 x 10 ⁻⁶ | 6.3 | 0.000108 (9) | |
| 2703.0 | 0.000096 (2) | 0.000094 (4) | 1.5 x 10 ⁻⁶ | 5.9 | 0.000115 (9) | |
| 2663.5 | 0.000464 (3) | 0.000462 (8) | 1.5 × 10 ⁻⁶ | 2.1 | 0.000480 (38) | 0.000470 (2) |
| 2648.1 | 0.000300 (2) | 0.000299 (7) | 1.5 x 10 ⁻⁶ | 2.8 | 0.000310 (23) | |
| 1616.3 | 0.000421 (3) | 0.000418 (8) | 1.5 x 10 ⁻⁶ | 2.3 | 0.000437 (33) | 0.000425 (1) |
| 2599.8 | 0.000414 (3) | 0.000413 (6) | 1.5 x 10 ⁻⁶ | 1.8 | 0.000434 (33) | 0.000421 (1) |
| 2582.2 | 0.00329 (3) | 0.000327 (9) | 1.5 x 10 ⁻⁶ | 3.2 | 0.000344 (25) | 0.000333 (1) |
| 2557.4 | 0.000297 (2) | 0.000296 (6) | 1.5 x 10 ⁻⁶ | 2.5 | 0.000316 (23) | |
| 2544.3 | 0.000335 (2) | 0.000333 (7) | 1.5 x 10 ⁻⁶ | 2.6 | 0.000356 (26) | 0.000341 (1) |
| 2510.9 | 0.000105 (2) | 0.000104 (4) | 1.5 x 10 ⁻⁶ | 5.3 | 0.000124 (11) | |
| 2496.0 | 0.000148 (2) | 0.000147 (5) | 1.5 x 10 ⁻⁶ | 4.4 | 0.000161 (13) | 0.000153 (1 |
| 2471.4 | 0.090199 (2) | 0.000198 (5) | 1.5 x 10 ⁻⁶ | 3.3 | 0.000219 (30) | 0.000204 (1 |
| 2446.6 | 0.000153 (2) | 0.000152 (4) | 1.5 x 10 6 | 3.6 | 0.000171 (13) | |
| 2427.3 | 0.000197 (2) | 0.000197 (4) | 1.5 x 10 ⁻⁶ | 2.8 | 0.000216 (16) | |
| 2406.4 | 0 to 10337 (1) | 0.000336 (5) | 1.5 x 10 ⁻⁶ | 1.9 | 0.000344 (25) | 0.000342 (1 |
| 2390,3 | 0.000452 (2) | 0.000450 (6) | 1.5 x 10 ⁻⁶ | 1.7 | 0.000466 (35) | 0.000456 (2 |
| 2365.7 | 0.000200 (5) | 0.000202 (9) | 1.5×10^{-6} | 5.2 | 0.000220 (17) | |
| 2331.0 | 0.000774 (5) | 0.000775 (12) | 1.5 x 10 ⁻⁶ | 1.7 | 0.000801 (55) | 0.000775 (1 |
| 2319.8 | 0.000776 (4) | 0.000775 (7) | 1.5 x 10 ⁻⁶ | 1.1 | 0.000842 | 0.000773 (1 |
| 2271.8 | 0.000319 (2) | 0.000318 (4) | 1.5 x 10 ⁻⁶ | 1.7 | 0.000331 (24) | 0.000325 (1 |
| 2240.5 | 0.000568 (2) | 0.000566 (9) | 1.5 x 10 ⁻⁶ | 1.9 | 0.000598 (48) | 0.000573 (1 |
| 2194.0 | | 0.000186 (4) | 1.5 x 10 ⁻⁶ | 3.0 | 0.000206 (16) | 0.000192 (2 |
| 2178.7 | 0.000359 (3) | 0.000357 (6) | 1.5 x 10 ⁻⁶ | 2.1 | 0.000376 (27) | 0.000366(1 |

Table 3.4 - Continued

| <i>v</i> (cm ⁻¹) | Weighted average | Unweighted average | Uncertainty due to anchor points | % estimated accuracy | Ref. 8 | Ref. 7 |
|------------------------------|---------------------|-----------------------|---|----------------------------|---------------|--------------|
| 2150.5 | 0.000145 (2) | 0.000145 (3) | 1.5 x 10 ⁻⁶ | 3.1 | 0.000170 (14) | |
| 2132.5 | 0.000087 (2) | 0.000086 (4) | 1.5 x 10 ⁻⁶ | 6.4 | 0.000108 (10) | |
| 2089.1 | 0.000141 (2) | 0.000140 (5) | 1.5×10^{-6} | 4.6 | 0.000154 (13) | 0.000147 (1) |
| 2058.2 | 0.000133 (2) | 0.000132 (6) | 1.5 x 10 ⁻⁶ | 5.7 | 0.000149 (13) | |
| 2045.6 | 0.000147 (2) | 0.000147 (4) | 1.4 10-6 | 3.7 | 0.000164 (14) | 0.000150 (1) |
| 2023.7 | 0.000244 (3) | 0.000242 (7) | 1 f x 10 5 | 3.5 | 0.000268 (20) | 0.000249 (1) |
| 2002.5 | 0.000284 (2) | 0.000284 (5) | 19219 | 2.3 | 0.000306 (23) | |
| 1962.2 | 0.00208 (1) | υ.00207 (2) | 1.5 × 10 ⁻⁶ | 1.0 | 0.00213 (8) | |
| 1942.9 | 0.00410 (2) | 0.00409 (4) | 1.5 x 10 ⁻⁶ | 1.0 | 0.00441 (11) | |
| 1881.4 | 0.00240 (1) | 0.00239 (1) | 19 x 10 ⁻⁶ | 0.5 | 0.00243 (9) | 0.00239 (1) |
| 1861.9 | 0.00442 (2) | 0.00442 (4) | 1.9 x 10 ⁻⁶ | 0.9 | 0.00465 (12) | 0.00445 (3) |
| 1787.8 | 0.00356 (1) | 0.00355 (3) | 2.0 x 10 ⁻⁶ | 0.9 | 0.00357 (13) | 0.00355 (2) |
| 1730.5 | 0.00346 (1) | 0.00345 (3) | 1.9 x 10 ⁻⁶ | 0.9 | 0.00347 (13) | 0.00348 (1) |
| 1685.9 | 0.000652 (4) | 0.000652 (11) | 1.9×10^{-6} | 2.0 | 0.000673 (25) | 0.00652 (1) |
| 1645.5 | 0.00227 (1) | 0.00227 (2) | 2.8 x 10 ⁻⁶ | 1.0 | 0.00233 (8) | 0.00230 (1) |
| 1622.2 | 0.00267 (3) | 0.00267 (4) | 3.6 x 10 ⁻⁶ | 1.6 | 0.00267 (10) | 0.00264 (1) |
| 1583.8 | 0.0816 (7) | 0.0814 (12) | 3.6 x 10 ⁻⁶ | 1.5 | 0.0811 (20) | 0.0805 (5) |
| 1566.1 | 0.0148 (1) | 0.0147 (2) | 3.6 x 10 ⁻⁶ | 1.4 | 0.0141 (3) | |
| 1477.4 | 0.252 (9) | 0.254 (9) | 3.9 x 10 ⁻⁶ | 3.5 | 0.323 (8) | |
| 1445.4 | | 0.103 (7) | 3.9 x 10 ⁻⁶ | 6.8 | 0.109 (3) | 0.098 (1) |
| 1387.1 | 0.00303 (1) | 0.00303 (1) | 3.2 x 10 ⁻⁶ | 0.4 | 0.00315 (111) | |
| 1370.7 | | 0.00303 (1) | 3.2 x 10 ⁻⁶ | 0.4 | 0.00313 (12) | |
| 1325.2 | 0.00253 (1) | 0.00252 (4) | 3.0×10^{-6} | 1.7 | 0.00268 (10) | |
| 1298.1 | | 0.00375 (8) | 3.0 x 10 ⁻⁶ | 2.2 | 0.00390 (15) | |
| 1272.5 | | 0.00375 (10) | 3.0 x 10 ⁻⁶ | 2.7 | 0.00402 (15) | |
| 1235.2 | | 0.00277 (1) | 3.4 x 10 ⁻⁶ | 0.5 | 0.00285 (11) | |
| 1210.9 | | 0.00235 (2) | 3.4 x 10 ⁻⁶ | 1.0 | 0.00246 (9) | |
| 1171.6 | | 0.00520 (8) | 1.0 x 10 ⁻⁵ | 1.7 | 0.00532 (20) | |
| 1156.4 | | 0.00459 (7) | 1.0 x 10 ⁻⁵ | 1.7 | 0.00480 (18) | |
| 1122.1 | | 0.0268 (3) | 1.7 x 10 ⁻⁵ | 1.2 | 0.0258 (5) | |

Table 3.4 - Continued

| v (cm ⁻¹) | Weighted average | Unweighted average | Uncertainty due to anchor points | estimated accuracy | Ref 8 | Ref 7 |
|-----------------------|---------------------|-----------------------|---|-----------------------|--------------|------------|
| 1083.2 | 0.178 (2) | 0.178 (2) | 1.7×10^{18} | 1.1 | 0.179 (5) | 0.176 (1) |
| 1068.1 | 0.0520 (6) | 0.0521 (8) | 1.7×10^{15} | 1.6 | 0.0515 (10) | 0.0517 (3) |
| 1022.6 | 0.185 (8) | 0.186 (11) | 1.1×10^{-5} | 5.9 | 0.194 (5) | 0.175 (2) |
| 1001.9 | 0.0442 (12) | 0.0444 (17) | 1.1×10^{3} | 3.9 | 0.0434 (8) | 0.0427 (4) |
| 963.7 | 0.00376 (6) | 0.00376 (8) | 1.1×10^{-8} | 2 4 | 0 00395 (15) | |
| 935.0 | 0.00784 (3) | 0.00795 (42) | 1.2×10^{-5} | 5.4 | 0.00812 (24) | |
| 902.9 | 0.0315(1) | 0.0319 (11) | 1.2 x 10 ⁻⁵ | 3.5 | 0.0304 (6) | |
| 866.3 | 0.00299 (2) | 0.00296 (8) | 1.2 x 10 ⁻⁵ | 3.1 | 0.00314 (12) | |
| 830.0 | 0.00582 (2) | 0.00579 (25) | 1.2×10^{-5} | 4.5 | 0,00609 (23) | |
| 812.2 | 0.00352 (1) | 0.00351 (5) | 1.2 x 10 ⁻⁵ | 1.8 | 0.00366 (14) | |
| 739.5 | 0.747 (16) | 0.747 (16) | 8.4 x 10 ⁻⁵ | 2.2 | 0.785 (20) | |
| 702.2 | 0.307 (7) | 0.306 (14) | 8.4 x 10 ^{.5} | 4.6 | 0.311 (8) | 0.293 (3) |
| 684.8 | 0.375 (10) | 0.373 (22) | 8.3 x 10 ⁻⁵ | 5.9 | 0.389 (11) | 0.355 (6) |
| 661.9 | 0.0176 (4) | 0.0175 (10) | 8.3 x 10 ⁻⁵ | 6.2 | 0.0175 (4) | |
| 614.0 | 0.00826 (20) | 0.00805 (42) | 1.6 x 10 ⁻⁵ | 5.4 | 0,00902 (30) | |
| 547.7 | 0.00099 (1) | 0.00097 (4) | 1.6 x 10 ⁻⁵ | 5.8 | 0.00101 (4) | |
| 533.7 | 0.00140(1) | 0.00140 (2) | 1.6 x 10 ⁻⁵ | 2.6 | 0.00144 (5) | |
| 467.8 | 0.284 (6) | 0.282 (7) | 5.1 x 10 ⁻⁵ | 2.5 | 0.263 (39) | |

a - In this column the number in parentheses is the 95% confidence \text{\text{imit} in the last digit.}

b - In this column the number in parentheses is the maximum deviation from the unweighted average, except for the 739.5 cm⁻¹ band where it is the 95% confidence limit in the last digit of the data of A, the only spectroscopist to measure that band.

c - The uncertainty is calculated as the average uncertainty in the anchor points (Table 3.1) at each side of the band.

d - The estimated accuracy is the sum of the maximum deviation from the unweighted average and the uncertainty due to the anchor points, as a percentage of the unweighted average.

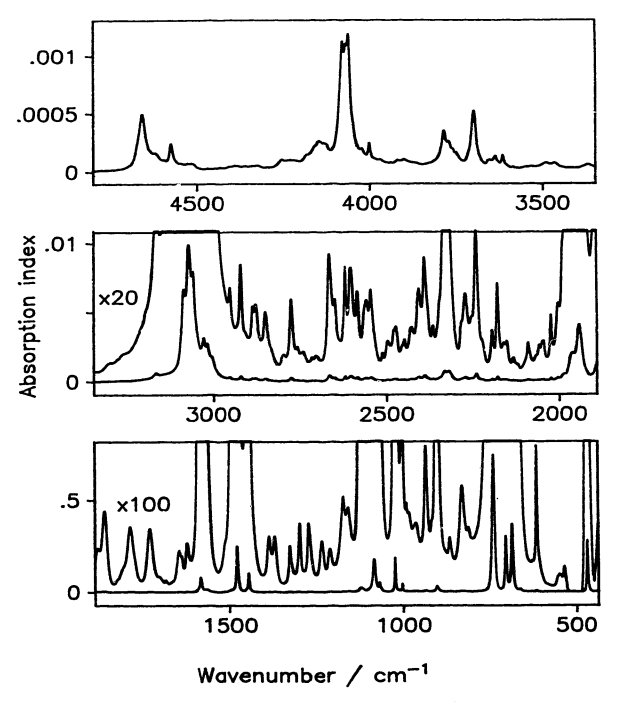


Figure 3.1 - Absorption index, $k(\tilde{\nu})$, spectrum between 4800 and 435 cm⁻¹ of chlorobenzene at 25°C. The ordinate scale labels in the middle and bottom boxes are for the lower spectrum in the box; they must be divided by 20 and 100, as shown, for the upper spectrum in the box.

Table 3.5 - Absorption indices between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C ab

| | | | .0301 | phon | marce | 3 000 | · · · · · | | 111G T. | · · · · · · | | iquid | CHIQIC | JOCHZ | ene a | 125 (| | | |
|--------------------|--------|------------------|--------------|--------------|--------------|--------------|--------------|--------------|-----------------------|-----------------------|--------------|---------------------|--------------|-------------------|--------------|--------------|-------------|--------------|---|
| em ⁻¹ | | YE | .) | 1 | | :_ | 4 | | ٠ | 7 | 8 | 3 | 1-3 | 11 | 12 | 13 | 14 | 15 | 10 |
| 1790 95 | 0 | -7 | 124 | 127 | 126 | 127 | 123 | 118 | 122 | 125 | 121 | 1.54 | 1.20 | 137 | 141 | 132 | 111 | 117 | 1.41 |
| 4783 56 4766 20 | 0 1 | -7 -7 | 139 152 | 143 | 143 | 147 | 142 | 150 | 154 | 155 | 157 | 154 | 153 | 155 | 1,00 | 154 | 1 > 4 | 120 | 124 |
| 4731 49 | 2 | -7 | 230 | 156 248 | 151 257 | 161 273 | 162 298 | 173 | 180 | 191 | 40g | 198 | 203 | 204 | 210 | 207 | 208 | 208 | 214 |
| 4665 92 | - | -7 | 3361 | 4:45 | 4873 | 4821 | A H NO | 3178 | 2563 | 2132 | 1882 | 550 1770 | 614 [643 | 1649 | 1030 | 1129 | 1 20.2 | 2105 | 2*23 |
| 4600 35 | = | .7 | 1040 | 1070 | 1035 | 205 | 1.130 | 1420 | 2206 | 3340 | 1700 | 1311 | 1003 | 1144 | 822 | 1203 | 1420 | 1250 | (093 (07 |
| 4534 79 | 2 | -7 | 675 | 673 | 725 | 721 | 718 | 740 | 728 | 202 | 547 | 437 | 370 | 140 | 314 | 282 | 267 | 255 | 250 |
| 4469 22 | 2 | -7 | 281 | 280 | 262 | 257 | 272 | 297 | 298 | 30% | 3.48 | 387 | 4-20 | 3cm) | 140 | 145 | 149 | 180 | 415 |
| 4403 65 | 2 | -7 | 463 | 457 | 467 | 200 | 546 | 542 | 5.10 | 516 | 473 | 461 | 475 | 401 | 5(3) | 488 | 473 | 488 | 512 |
| 4338 08 | 2 | -7 | 511 | 518 | 541 | 554 | 502 | 5 3() | 460 | 422 | 101 | 388 | 16/2 | 402 | 405 | 44 | 415 | 414 | -Mari |
| 4272 51 | 2 | -7 | 526 | 609 | 708 | 839 | 987 | 1088 | 1043 | 997 | 987 | 17:313 | 1008 | 1018 | 1037 | 1028 | 1023 | 1008 | 287 |
| 4206.95 | 2 | -7 | 956 | 935 | 940 | 985 | 10% | 1253 | 1419 | 1529 | 1534 | 1588 | 1711 | 1911 | 2147 | 2257 | 2384 | 2591 | 26.74 |
| 4141 38 | i | -6 -0 | 259 1131 | 249 1108 | 247 1056 | 248 1935 | 242 1054 | 234 1094 | 200 | 189 | 185 | 193 | 216 | 256 | 328 | 430 | 613 | 828 | Librici |
| 4044.95 | i | -0 -7 | 4230 | 3708 | 3220 | 2848 | 2592 | 2400 | 1109 | 2091 2091 | 1302 | 1158 | 1195 1972 | 1114 | 921 | 739 | CK14 | 119 | - Ano |
| 4012.17 | ì | -7 | 1527 | 1487 | 1495 | 1589 | 1826 | 2217 | 2497 | 2293 | 1896 | 1588 | 1305 | 2025 1284 | 2021 1234 | 1923 | 1.788 | 1678 | 1597 |
| 3977.46 | 2 | -7 | 1048 | 1037 | 1063 | 1030 | 940 | 870 | 827 | 816 | 785 | 769 | 821 | 780 | 7eres | NIH. | H 7H | 1000 | 1188) |
| 3911 89 | 2 | -7 | 966 | 959 | 1007 | 1086 | 1056 | 980 | 912 | 881 | 839 | 824 | 788 | 762 | 758 | 730 | 672 | 647 | 621 |
| 3846 32 | 2 | -7 | 621 | 639 | 670 | 709 | 7-40 | 771 | 810 | 861 | 931 | 1013 | 1005 | 1196 | 1412 | 1797 | 2309 | ion Z | |
| 3786 54 | 1 | -7 | 3438 | 3534 | 3325 | 2990 | 2715 | 2579 | 2575 | 2017 | 2610 | 2525 | 2375 | 2196 | 2058 | 2002 | 1979 | 1894 | 1765 |
| 3753 76 | 1 | -7 | 1663 | 1641 | 1665 | 1640 | 1536 | 1424 | 1312 | 1211 | 1128 | 1068 | 1017 | 974 | 1744 | 929 | 926 | 950 | 1004 |
| 3720 97 | l | -7 | 1103 | 1267 | 1475 | 1670 | 18-14 | 2066 | 2410 | 2933 | 3621 | 4110 | 5021 | 5253 | \$138 | 4621 | 1330 | rini) | 2486 |
| 3688.19 | 1 | -7 | 2080 | 1794 | 1584 | 1411 | 1259 | 1130 | 1024 | 941 | 874 | 816 | 762 | 723 | 711 | 747 | 830 | 952 | 1012 |
| 3655 40 3622.62 | 1 1 | .7 -7 | 1041 763 | 1055 785 | 10-45 887 | 1013 | 990 | 997 | 1043 | 1127 | 1241 | 1351 | 1364 | 1246 | 1089 | *285 | 915 | 880 | Sen |
| 3589 84 | l | -7 | 374 | 388 | 405 | 1098 426 | 1358 395 | 1424 367 | 1223 353 | 982 341 | 797 333 | 692 326 | 634 | 594 | 149 | 49K | 450 | 412 | 185 |
| 3557 05 | ì | -7 | 365 | 378 | 391 | 404 | 420 | 431 | 431 | 430 | 428 | 432 | 122 439 | 323 453 | 400 323 | 325 476 | 331 475 | 3.18 | 151 -159 |
| 3522.34 | 2 | -7 | 445 | 453 | 478 | 508 | 548 | 607 | 667 | 724 | 770 | 782 | 718 | 685 | 663 | 677 | 730 | 762 | 721 |
| 3456.77 | 2 | -7 | 626 | 533 | 471 | 439 | 409 | 376 | 348 | 326 | 316 | 321 | 320 | 312 | 313 | 313 | 316 | 124 | 3-30, |
| 3391 20 | 2 | -7 | 384 | 435 | 480 | 519 | 567 | 623 | 647 | 624 | 570 | 517 | 465 | 420 | 393 | 179 | 374 | 175 | • |
| 3332.39 | 0 | -7 | 376 | 370 | 372 | 375 | 383 | 394 | 396 | 405 | 408 | 415 | 423 | 433 | 418 | -1-18 | 458 | 470 | 479 |
| 3315.99 | 0 | -7 | 489 | 516 | 524 | 537 | 549 | 556 | 578 | 593 | 601 | 616 | 639 | (-1 6 | 656 | 668 | 671 | 682 | |
| 3299.60 | 1 | -7 | 699 | 706 | 6 97 | 701 | 696 | 700 | 708 | 720 | 737 | 752 | 775 | アソソ | 822 | 824 | 843 | 867 | 'N K' |
| 3261.03 | 3 | -7 | 1007 | 1041 | 1078 | 1162 | 1303 | 1476 | 1693 | 1939 | 2248 | 2609 | 2989 | 4020 | | | | | |
| 3174.25 3141.47 | 1 | -7 -7 | 4429 5986 | 4871 6095 | 5361 | 5969 | 6535 | 6604 | 6288 | 5914 | 5633 | 5444 | 5341 | 5320 | 5349 | 5424 | | 5670 | 2844 |
| 3108 68 | ì | - <i>1</i> -6 | 1034 | 1101 | 6180 1183 | 6261 1292 | 6359 1439 | 6479 1641 | 5616 1912 | 67 7 9 2280 | 6971 2771 | 7188 3325 | 7441 4036 | 7705 5061 | 8019 | X377 | | 9245 | 9767 |
| 3075 90 | i | -6 | 6569 | 7592 | 9004 | 9870 | 9732 | 8988 | 8160 | 7685 | 7740 | 7989 | 7561 | 1448 | 6140 5351 | 4028 | | 6381 3819 | 0230 1425 |
| 3043.12 | 1 | -6 | 3112 | 2860 | 2658 | 2504 | 2421 | 2435 | 2543 | 2738 | 3008 | 3174 | 2989 | 2687 | 2554 | 2659 | 2754 | | 2289 |
| 3010 33 | 1 | -6 | 2033 | 1883 | 1842 | 1840 | 1761 | 1593 | 1393 | 1205 | 1050 | 927 | 829 | 747 | 676 | 616 | 563 | 517 | 479 |
| 2977 55 | 1 | -7 | 4474 | 4204 | 39 77 | 3790 | 3626 | 3487 | 3348 | 3223 | 3099 | 2991 | 2906 | 2869 | 2993 | 3079 | 3356 | 3305 | 2999 |
| 2944 76 | 1 | -7 | 2550 | 2301 | 2149 | 2059 | 1985 | 1949 | 1945 | 1963 | 2007 | 2083 | 2218 | 2510 | 3178 | 4147 | | | |
| 2918 73 | 0 | -7 - | 4248 | 3947 | 3473 | 3021 | 2657 | 2380 | 2181 | 2043 | 1947 | 1887 | 1843 | 1813 | 1791 | 1776 | 1748 | 1721 | 1691 |
| 2902.34 | | -7 | 1664 | 1636 | 1611 | 1596 | 1585 | 1581 | 1580 | 1588 | 1597 | 1615 | 1643 | 1690 | 1759 | 1874 | | 2288 | 2558 |
| 2868.59 | 0 | -7 -7 | 2713 1799 | 2695 1641 | 2605 1553 | 2534 1507 | 2514 | 2551 1522 | 2621 | 2698 | 2766 | 2798 | 2787 | 2727 | 2624 | 2494 | 2348 | | 2044 |
| 2835.81 | 1 | - <i>1</i> -7 | 1313 | 1259 | 1193 | 1116 | 1496 1035 | 951 | 1599 872 | 1752 799 | 2003 745 | 233 3 700 | 2524 666 | 2417 (40 | 2186 627 | 1949 626 | 1722 | | 1396 |
| 2803.02 | ì | -7 | 782 | 876 | 949 | 979 | 992 | 982 | 958 | 927 | 909 | 923 | 974 | 1086 | 1282 | 1636 | 431 2224 | 656 2855 | 700 2936 |
| 2770 24 | | -7 | 2508 | 2023 | 1626 | 1366 | 1222 | 1175 | 1200 | 1251 | 1239 | 1150 | 1064 | 1024 | 1033 | 1043 | | 1073 | 1089 |
| 2737.45 | | . 7 | 1078 | 1028 | 950 | 874 | 810 | 768 | 741 | 730 | 727 | 740 | 758 | 823 | 880 | 844 | 817 | | 874 |
| 2704 67 | | -7 | 928 | 943 | 929 | 902 | 870 | 835 | 805 | 786 | 780 | 780 | 796 | 822 | 869 | 949 | | 1245 | 1505 |
| 2671 89 | 1 | -7 | 1887 | 2443 | 3226 | 4096 | 4585 | 4530 | 4139 | 3660 | 3231 | 2914 | 2760 | 2810 | 2966 | 2891 | | 2078 | 180% |
| 2639.10 | | -7 | 1635 | 1551 | 1516 | 1503 | 1490 | 1477 | 1526 | 1680 | 1944 | 2318 | 2894 | 3745 | | | | | |
| 2616 92 | | -7 | 4094 | 4164 | 3899 | 3469 | 3006 | 2745 | 2548 | 2440 | 2407 | 2439 | 2535 | 2701 | 2957 | 3293 | 3652 | 3929 | 4111 |
| 2599.57 | | -7 | 4123 | 4024 | 3663 | 3148 | 2705 | 2447 | 2364 | 2444 | 2819 | 3265 | 2829 | 2205 | 1826 | 1634 | 1591 | | 1963 |
| 2566 78 | | -7 | 2306 | 2580 | 2716 | 2801 | 2907 | 2955 | 2850 | 2669 | 2557 | 2625 | 2900 | 3231 | 1305 | 3000 | | 2273 | 21/1/ |
| 2534.00 | | -7 - | 1769 | 1554 | 1361 | 1181 | 1032 | 915 | 835 | 788 | 761 | 754 | 788 | 916 | 1040 | 941 | 866 | | 986 |
| 2501.22 | | -7 | 1138 | 1316 | 1439 | 1463 | 1422 | 1363 | 1315 | 1288 | 1289 | 1328 | 1431 | 1636 | 1832 | 1823 | 1865 | 1958 | 1950 |
| 2468.43 | | -7 | 1811 | 1565 | 1298 | 1134 | 1073 | 1087 | 1158 | 1242 | 1298 | 1328 | 1390 | , | | , | | | , |
| 2448.18 2430.83 | | -7 -7 | 1449 1842 | 1507 1941 | 1515 1963 | 1447 1924 | 1355 1854 | 1282 1774 | 1237 | 1214 | 1212 | 1219 | 1237 | 1268 | 1312 | 1375 | | 1551 | 1655 |
| 2398 04 | | -7 | 2671 | 2816 | 3296 | 4077 | 4499 | 4217 | 1 <i>7</i> 27 3740 | 1743 3310 | 1855 2941 | 2086 2606 | 2457 2273 | 2917 1962 | 3279 1737 | 3344 1627 | | 2920 1774 | |
| 2365.26 | | -7 | 2014 | 1864 | 1675 | 1596 | 1625 | 1728 | 1872 | 2072 | 2374 | 2739 | 2952 | 3092 | 3451 | 4078 | | 5818 | |
| 2332.48 | | -7 | 7550 | 7728 | 7373 | 6912 | 6765 | 7082 | 7621 | 7674 | 7025 | 6110 | | 4416 | 3695 | 3088 | | 2159 | |
| 2299 69 | | -7 | 1583 | 1384 | 1260 | 1197 | 1172 | 1220 | 1395 | 1718 | 2038 | 2157 | | 2350 | 2622 | | | 3149 | |

Table 3.5 - Continued

| um i | A.E. | }'F: | ٠. ٠, | i | 2 | 3 | | <u> </u> | - 5 | 7 | × | 9 | !0 | 1! | 12 | 13 | 14 | 15 | 16 |
|---------|------|----------|---------|-------|------|-------|---------------|-------------------------|-------|-------|------|---------------------|-------|--------------|--------------|------|--------------|--------------|--------------|
| 2266.91 | 1 | 7 | 2636 | 2365 | 2179 | 2010 | 2115 | 2213 | 2311 | 2345 | 2362 | 2483 | 2834 | 3481 | 4412 | 5370 | | 1847 | 3820 |
| 29412 | i | , | فاهن | 2316 | 1677 | 1528 | 1525 | نواو باشد. نواو باشد | (45th | ربعوا | 1219 | ار واست المواول: | 919 | 598 | 736 | 599 | 546 | 727 | 798 |
| 291.34 | i | 7 | 921 | 1112 | 1383 | 1714 | 1845 | 15144 | 1316 | 1199 | 1276 | 1590 | 2264 | 3224 | 3537 | 2895 | | 1660 | 1416 |
| Del 50 | i | 7 | 1283 | 1214 | 1222 | 1.254 | 1378 | 1356 | 1343 | 1326 | 1370 | 1438 | 1418 | 1256 | 1012 | 822 | 718 | 673 | 572 |
| 135.77 | 1 | 7 | 72% | 823 | 856 | 772 | 4. 34.0 | 4,643 | 4,47 | 636 | 524 | 595 | 5.49 | 511 | 487 | 482 | 491 | 521 | 560 |
| 102 7 | ì | 7 | والإيوم | 666 | 723 | 770 | 328 | 925 | 1147 | | | | | | | | | - - . | . 5.5 |
| 45 (16) | 11 | 7 | 1294 | 1392 | 1381 | 1292 | 1290 | 1130 | 1973 | 1918 | 963 | 906 | 852 | 810 | 779 | 759 | 747 | 7-11 | 740 |
| 074 00 | ń | . 7 | 7-14 | 756 | 777 | 809 | 443 | 876 | 898 | 914 | 929 | 942 | 959 | 979 | 1997 | 1053 | | 1228 | 1313 |
| 057.57 | į, | .7 | 1312 | 1252 | 1208 | 1199 | 1223 | 1273 | 1336 | 1395 | 1433 | 1447 | 1447 | 1448 | 1458 | 1461 | 1413 | | 1187 |
| 2041-28 | g. | 7 | 1981 | 997 | '443 | 91)T | 445 | 873 | 869 | 870 | 877 | 893 | 923 | 972 | 1044 | 1136 | 1237 | | 1698 |
| 2024 89 | 9 | .7 | 2102 | 2415 | 2310 | 1984 | ۱ <i>۲</i> /۰ | 1519 | 1407 | 1340 | 1306 | 1295 | 1308 | 1341 | 1392 | 1455 | 1542 | | 1783 |
| 2008 49 | ŋ | .7 | 1952 | 2144 | 2344 | 2530 | 2683 | 791 | 2838 | 2822 | 2773 | 2727 | 2697 | 2690 | 2719 | 2789 | 2915 | | 3-400 |
| 1992 10 | 9 | .7 | 3783 | 4231 | 4645 | 4927 | 5079 | 51/42 | 5329 | 5481 | 5594 | 5642 | 5639 | | | 5771 | | | |
| 974 74 | 1 | • | 833 | 1/42 | 1297 | 1547 | 7949 15044 | 1081 | 2002 | 2061 | 2028 | 2002 | 2028 | 5629 2141 | 5661 2365 | 2731 | 5994 3220 | 3711 | 6842 4945 |
| | | | | 3797 | | | | | | | | | | | | | | | |
| 941.96 | i | | 412X | | 3351 | 2827 | 2316 | 1872 | 1519 | 1262 | 1084 | 950 | 834 | 732 | 647 | 578 | 527 | 496 | 472 |
| 1907.25 | 2 | ٠, | 456 | 512 | 420 | 821 | 1198 | 1800 | 2291 | 2377 | 2251 | 2433 | 3199 | 4112 | 4374 | 3455 | | 1206 | 746 |
| 1841 68 | 2 | 6 | 532 | 448 | 456 | 550 | 100 | 738 | 883 | 1068 | 1248 | 1389 | 15-48 | 1869 | 2478 | 3226 | 3553 | | 2710 |
| 1775 11 | 2 | -6 | 2066 | 1434 | 1154 | 860 | 697 | 618 | 623 | 716 | 953 | 1425 | 2205 | 3124 | 3435 | 2911 | 2105 | 1445 | 1051 |
| 1710 54 | 2 | -7 | 8674 | 8392 | 8091 | 7158 | 6164 | 5304 | | | | | | | | | | | |
| 1689 33 | 1 | .7 | 5373 | 6089 | 6452 | 5664 | 5313 | 4872 | 1191 | 4520 | 4698 | 5026 | 53-48 | 5584 | 5784 | 6082 | 6509 | 70-19 | 7916 |
| 1656 55 | 1 | -6 | 929 | 1138 | 1427 | 1732 | 2028 | 2223 | 2263 | 2174 | 2023 | 1918 | 1953 | | | | | | |
| 1636-30 | 0 | -6 | 1938 | 1816 | 1631 | 1471 | 1362 | 1306 | 1308 | 1345 | 1431 | 1576 | 1778 | 2028 | 2293 | 2518 | 2638 | | 2586 |
| 161991 | 0 | -6 | 2456 | 2301 | 2139 | 2002 | 1893 | 1800 | 1744 | 1724 | 1717 | 1739 | 1785 | 1842 | 1906 | 1996 | | 2247 | 2425 |
| 1603 51 | ŋ | -5 | 264 | 291 | 323 | 362 | 411 | 477 | 567 | 692 | 861 | 1077 | 1333 | 1604 | 1866 | 2118 | 2384 | | 3166 |
| 1587 12 | 0 | -5 | 3882 | 5116 | 6860 | 8035 | 8017 | 7346 | 6287 | 5116 | 4054 | 3195 | 2555 | 2103 | 1802 | 1617 | 1518 | | 1435 |
| 1569 77 | t | ٠5 | 1374 | 1415 | 1473 | 1341 | 1978 | 822 | 623 | 468 | 357 | 278 | 224 | 186 | 161 | 144 | 132 | 122 | 116 |
| 153598 | ŧ | 6 | 1087 | 1969 | 1103 | 1163 | 1225 | 1296 | 1398 | 1516 | 1705 | 1920 | 2188 | 2460 | 2678 | 2819 | 2949 | 3196 | 3633 |
| 1504 20 | t | -5 | 441 | 569 | 749 | 931 | 1046 | 1119 | 1230 | 1417 | 1678 | 2067 | 2746 | | | | | | |
| 1483 95 | 0 | -4 | 331 | 414 | 543 | 756 | 1121 | 1713 | 2365 | 2467 | 2135 | 1490 | 1013 | 735 | 577 | 463 | 363 | 274 | 209 |
| 1467 56 | 0 | - 5 | 1662 | 1371 | 1173 | 1031 | 928 | 855 | 897 | 783 | 778 | 79 6 | 837 | 904 | 999 | 1127 | 1291 | 1489 | 1713 |
| 1451 17 | 0 | -4 | 196 | 225 | /265 | 337 | 493 | 800 | 1033 | 783 | 504 | 340 | 245 | 192 | 163 | 137 | 109 | 86 | 70 |
| 1433.81 | 1 | -6 | 5156 | 4193 | 3559 | 3047 | 2608 | 2224 | 1893 | 1624 | 1414 | 1271 | 1162 | 1056 | 983 | 941 | 916 | 919 | 940 |
| 1401 03 | 1 | -6 | 976 | 1047 | 1180 | 1410 | 1766 | 2252 | 2755 | 3022 | 2941 | 2655 | 2335 | 2112 | 2058 | 2214 | 2559 | 2927 | 3016 |
| 1368 24 | ı | -6 | 2718 | 2224 | 1739 | 1339 | 1042 | 837 | 703 | 615 | 557 | 519 | 496 | 483 | 480 | 485 | 497 | 517 | 546 |
| 1335.46 | 1 | 6 | 594 | 676 | 840 | 1139 | 1753 | 2464 | 2300 | 1808 | 1537 | 1327 | 1144 | 1032 | 992 | 1008 | 1074 | 1201 | 1430 |
| 1303 64 | 0 | -6 | 1612 | 1869 | 2231 | 2701 | 3227 | 3645 | 3737 | 3455 | 2962 | 2436 | 2005 | 1688 | 1469 | 1329 | 1244 | 1194 | 1166 |
| 1287 25 | 0 | -6 | 1149 | 1139 | 1135 | 1137 | 1150 | 1174 | 1213 | 1271 | 1352 | 1472 | 1647 | 1916 | 2330 | 2910 | 3470 | 3738 | 3699 |
| 1269 89 | 1 | -6 | 3311 | 2893 | 2443 | 1982 | 1614 | 1345 | 1150 | 1019 | 942 | 911 | 922 | 980 | 1092 | 1275 | | 1897 | 2289 |
| 1237 11 | 1 | -6 | 2629 | 2773 | 2634 | 2317 | 1969 | 1647 | 1407 | 1277 | 1249 | 1325 | 1512 | 1795 | 2104 | 2309 | 2331 | | 1968 |
| 1204 32 | i | -6 | 1743 | 1575 | 1483 | 1454 | 1471 | 1524 | 1610 | 1729 | 1888 | 2093 | 2349 | 2674 | 3077 | 3531 | | 4439 | 4945 |
| 1171 54 | 1 | -ó | 5201 | 4965 | 4573 | 4297 | 4193 | 4235 | 4373 | 4530 | 4591 | 4454 | 4142 | 3786 | 3458 | 3162 | 2955 | | 2825 |
| 1138 75 | ı | -5 | 293 | 322 | 376 | 479 | 655 | 960 | 1455 | 2129 | 2636 | 2558 | 2241 | 2051 | 1875 | 1587 | | 1145 | 1073 |
| 1105 97 | i | 4 | 108 | 118 | 137 | 167 | 208 | 252 | 303 | 382 | 504 | 717 | 1095 | 1595 | 1764 | 1417 | 950 | 606 | 401 |
| 1074 15 | 0 | -5 | 3391 | 2992 | 2804 | 2872 | 3277 | 4161 | 5137 | 4936 | 3874 | 3070 | 2577 | 2182 | 1807 | 1469 | 1199 | 990 | 829 |
| 1057 76 | 0 | -6 | 7024 | 6034 | 5284 | 4744 | 4297 | 3932 | 3606 | 3377 | 3214 | 3097 | 3012 | 2974 | 2926 | 2896 | 2912 | 2966 | |
| 1041 37 | | -5 | 310 | 320 | 330 | | | | 409 | | | | | | | | | | 3008 |
| 1024 97 | 0 | -3 | 552 | 1031 | | 340 | 358 | 379 667 | | 450 | 500 | 570 | 664 | 795 | 980 | 1235 | 1604 | 2219 | 3310 |
| | | | | | 1772 | 1726 | 1104 | 662 | 412 | 273 | 193 | 150 | 126 | 115 | 121 | 126 | 107 | 86 | 76 |
| 1008 58 | 0 | -5 | 704 | 703 | 733 | 826 | 1088 | 1616 | 3063 | 4433 | 2907 | 1500 | 1005 | 734 | 613 | 537 | 484 | 466 | 451 |
| 992 19 | | -6 | 4491 | 4756 | 4851 | 4544 | 4141 | 4385 | 4301 | 4271 | 4302 | 4324 | 4316 | 4242 | 4148 | 4013 | | 3769 | 3656 |
| 974 83 | | -6 | 3437 | 3417 | 3510 | 3566 | 3631 | 3713 | 3744 | 3697 | 3518 | 3273 | 3014 | 2846 | 2693 | 2709 | | 2937 | 3256 |
| 042 05 | ı | -6 | 3809 | 4727 | 6125 | 7578 | 7845 | 6726 | 5316 | 4244 | 3593 | 3293 | 3318 | 3608 | 3 875 | 3949 | | 4920 | 6298 |
| 909 27 | | -5 | 907 | 1451 | 2362 | 3134 | 2949 | 2215 | 1541 | 1050 | 719 | 515 | 399 | 327 | 283 | 256 | | 230 | 223 |
| 876 48 | 1 | -6 | 2161 | 2112 | 2091 | 2210 | 2565 | 2942 | 2862 | 2604 | 2342 | 2107 | 1933 | 1825 | 1772 | 1776 | 1839 | 1954 | 2110 |
| 843 70 | | -6 | 2343 | 2663 | 3057 | 3571 | 4199 | 4901 | 5501 | 5787 | 5628 | 5144 | 4575 | 4090 | 3703 | 3446 | 3328 | 3372 | 3494 |
| 810 91 | 1 | -6 | 3456 | 3256 | 3075 | 2962 | 2909 | 2895 | 2889 | 2913 | 2955 | 3013 | 3094 | 3191 | 3332 | 3481 | 3609 | 3812 | 4105 |
| 778 13 | 1 | -4 | 45 | 49 | 54 | 60 | 67 | 76 | 86 | 100 | 118 | 142 | 176 | 227 | 305 | 433 | 657 | 1077 | 1858 |
| 745 35 | | 4 | 3172 | 4938 | 6619 | 7450 | 6104 | 3723 | 2025 | 1120 | 676 | 450 | 326 | 253 | 209 | 183 | | | |
| 719 31 | | -4 | 175 | 170 | 168 | 169 | 173 | 180 | 192 | 209 | 233 | 269 | 321 | 398 | 518 | 709 | 1012 | 1492 | 2179 |
| 702 92 | 0 | -4 | 2870 | 3019 | 2581 | 1976 | 1440 | 1056 | 802 | 638 | 545 | 510 | 524 | 576 | 652 | 753 | 908 | 1186 | 1691 |
| 686 53 | 0 | 4 | 2511 | 3425 | 3690 | 3083 | 2232 | 1564 | 1099 | 784 | 577 | 441 | 349 | 286 | 242 | 210 | 186 | | 155 |
| 670 14 | | -5 | 1457 | 1399 | 1376 | 1362 | 1371 | 1420 | 1513 | 1642 | 1745 | 1717 | 1546 | 1321 | 1123 | 961 | 826 | | 620 |
| 053 74 | | -6 | 5387 | 4749 | 4276 | 3849 | 3500 | 3203 | 2971 | 2742 | 2533 | 2391 | 2275 | 2146 | 2026 | 1946 | | 1798 | 1757 |
| 637 35 | | -6 | 1745 | 1695 | 1654 | 1648 | 1645 | 1608 | 1623 | 1584 | 1599 | 1625 | 1658 | 1680 | .,J2J | | . 0.55 | | . , , , |
| 626 26 | | -ó | 1679 | 1679 | 1675 | 1670 | 1674 | 1678 | 1693 | 1708 | 1737 | 1766 | 1814 | 1862 | 1937 | 2013 | 2126 | 2239 | 2416 |
| 618 07 | | -0 | 2594 | 2886 | 3178 | 3697 | 4217 | 5100 | 5982 | 6969 | 7957 | | | | 4864 | | | | |
| | | -0 -6 | 2476 | 2318 | 2219 | | | | | | | 7581 | 7205 | 6035 | | 4105 | | 2989 | 2633 |
| 609 87 | | | -4/0 | . 118 | 19 | 2120 | 2022 | 1924 | 1826 | 1728 | 1652 | 1577 | 1516 | 1456 | 1406 | 1356 | 1308 | 1260 | 1213 |

Table 3.5 - Continued

| em¹¹ | ΛŒ | IE | :_o | 1 | 2 | 3 | 4 | · | 6 | - | S | 3 | | 11 | 12 | 13 | 14 | 15 | 10 |
|--------|----|----|------|------|------|-------|-------|-------|------|------|------|-------|-------|---------|-------|-------|------|---------|-------|
| 601 19 | 0 | -ó | 1123 | 1040 | 905 | 897 | 835 | 776 | 721 | 671 | 627 | 588 | | | 120 | 4 7 7 | 457 | .1.1. | |
| 284 80 | 1) | -7 | 4262 | 4222 | 4151 | 4039 | 3007 | 3-130 | 3710 | 3045 | 3588 | 3:40 | 1470 | 3412 | 3323 | 3241 | 3174 | 3130 | 3137 |
| 568 41 | G | -7 | 3142 | 3140 | 3160 | 32(14 | 3270 | 4417 | 3581 | 3700 | 4070 | 4305 | 4718 | 5202 | 5833 | | | 8089 | 83.00 |
| 552 02 | 0 | -7 | 8660 | 8854 | 9125 | 2133 | 36-11 | 3633 | 2507 | 9358 | 9129 | 8804 | 8400 | 8204 | SUSS | | • | | |
| 539 97 | -1 | -6 | 808 | 812 | 816 | 831 | 845 | 884 | 922 | 985 | 1:47 | 1130 | 1212 | 1287 | 1362 | 1374 | 1387 | 1315 | 1243 |
| 531 77 | -1 | ٠ć | 1144 | 1045 | 368 | 802 | 830 | 78.7 | 750 | 722 | 533 | 5.453 | 454 | 3.4 | 328 | 2 18 | 100 | | *6 |
| 516 34 | 3 | 0 | С | 0 | •) | a | (1) | | | | | | | | | - " | • | • • • • | • • |
| 484 52 | 0 | _1 | 0 | 2 | 0 | 10 | 17 | 25 | 35 | 51 | 71 | 1343 | 141 | 202 | 200 | 462 | 758 | 1302 | 2119 |
| 468.13 | 0 | 4 | 2767 | 2566 | 1798 | 1121 | 7 | 454 | 310 | 238 | 191 | 150 | 113 | So | 00 | 52 | 40 | 30 | 21 |
| 451 74 | Ú | -6 | 1585 | 1048 | 978 | 1003 | 800 | 500 | 1029 | 1100 | 1257 | 1455 | 1.727 | 1 non-4 | 2024 | 2425 | | 1190 | 3487 |
| 435 35 | o | -ó | 3616 | 4147 | | | | | | | | | - | | - • • | | | | . • |

Note: Footnotes follow Table 3.7

The average agreement between the weighted and unweighted average peak heights is 0.6% which shows that, as was found previously^{1,2}, the fact that more spectra were run in this laboratory than elsewhere did not have an important influence on the averages. The agreement between spectroscopists is shown by the maximum deviations from the unweighted average peak heights, which average 2.4% for the 108 peak heights in Table 3.4.

The unweighted average $k(\tilde{v})$ spectrum, is taken as the primary intensity result of this work. It is shown in Fig. 3.1 and tabulated in Compact Table format in Table 3.5.

3.2.2 - Real refractive index spectrum.

The real refractive index spectrum $n(\tilde{v})$ was obtained by Kramers-Kronig transformation of the unweighted average $k(\tilde{v})$ spectrum^{1,10} with the value of n at 8000 cm⁻¹ equal to 1.5043±0.0005. This value was found by fitting the literature values¹¹⁻¹³ of

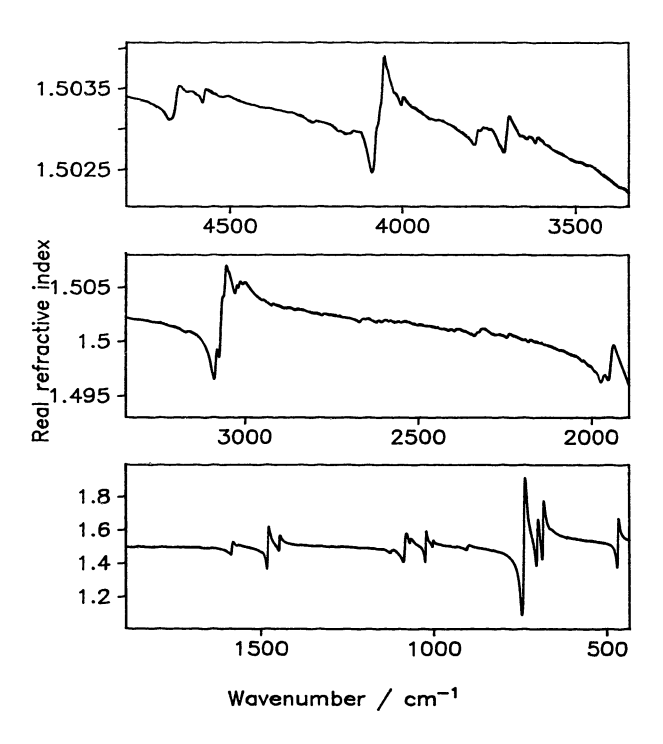


Figure 3.2 - Real refractive index, $n(\tilde{\nu})$, spectrum between 4800 and 435 cm⁻¹ of chlorobenzene at 25°C.

Table 3.6 - Real refractive indices between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C. Ab

| em¹. | λΈ | | 1 | | | 4 | | | | × | 3 | | | | | | | |
|--------------------|------------|-------|----------------|-------|---------|-------|-------|--------|---------|--------|---------|--------|---------|----------------|-------------|-----------|----------------|--------------------|
| | | 15033 | 15033 | | | 15033 | | 15033 | 15.137 | | | 15022 | 15033 | 12 | 13 | 15022 | 15033 | 10 |
| 4783 56 | | | 15033 | | | | | | 15033 | | | | | | | | | 15033 |
| | | | 15033 | | | | | | 15033 | | | - | | | | | 15033 | |
| | | - | 15033 | | | 15032 | | 15032 | | | | | 15031 | | | | | 18031 |
| 4665 92 | 2 | 15031 | 15031 | 15032 | 15034 | 15035 | | | 15035 | | | | | | | | 15034 | |
| 4600 35 | 2 | 15034 | 15034 | 15034 | 15033 | 15033 | 15033 | 55033 | 15034 | 15034 | 15034 | 15034 | 15034 | 15034 | 15034 | 15034 | 15034 | 15034 |
| 4534 79 | 2 | 15034 | 15034 | 15033 | 15033 | 15033 | 15033 | 15034 | 15034 | 15034 | 15033 | 15034 | 15033 | 15033 | 15033 | 15033 | 15033 | 18033 |
| 4469 22 | 2 | 15033 | 15033 | 15033 | 15033 | 15033 | 15033 | 15033 | 15033 | 15032 | 15032 | 15032 | 15032 | [5032 | 15032 | 18032 | 15032 | 18032 |
| 4403.65 | 2 | 15032 | 15032 | 15032 | 15032 | 15032 | 15032 | 15032 | 15032 | | | | | | | | 15032 | 18032 |
| 4338 08 | 2 | | 15032 | | | 15032 | | 15031 | | | | | 15031 | | • | | 12031 | 15031 |
| 4272 51 | | | 15030 | | | | | | 15030 | | | | | | | | | |
| 4206 95 | 2 | | | 15030 | | | | | | | | | 15029 | | 15029 | | 174:20 | 150,50 |
| 4141.38 4077.74 | 1 | | 15029 15029 | | | | | | | | | | 15026 | | | | 15024 | 15026 |
| 4044 95 | i | | 15037 | | | | | | 15035 | | | - | - | | | | 15038 | 15037 |
| 4012.17 | - | | 15033 | | | | | | 15033 | 15034 | | | 15033 | | | | | 15013 |
| 3977.46 | | | 15032 | | | | | 15032 | 15032 | | | | 15031 | | | | | 15031 |
| 3911 89 | | | 15031 | | | | | 15031 | 15031 | | | | 15030 | | | | 15030 | |
| 3846 32 | 2 | 15030 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15028 | 15028 | 15028 | 15028 | 15028 | 15028 | |
| 3786.54 | l | 15028 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15030 | 15030 | 15030 | 15030 | 15030 | 15030 | 15030 | 15030 | 15030 |
| 3753.76 | 1 | | 15029 | | | | 15029 | | 15029 | 15029 | 15029 | | 15029 | | | 15028 | 15028 | 18028 |
| 3720 97 | - | | 15028 | | | | | | | | 15027 | | 15029 | | | 15031 | 15031 | 15031 |
| 3688.19 | | | 15031 | | | | | | 15030 | 15030 | 15029 | | 15029 | | | 15029 | 15029 | 15029 |
| 3655.40 | | | | | | | 15028 | | | 15028 | | | 15029 | | | 15029 | 15028 | 15028 |
| 3622.62 3589 84 | | | 15028 15028 | | | | | | 15027 | 15029 | 15028 | | 15028 | | | | 15028 | 15028 15027 |
| 3557.05 | 1 | | 15027 | | | | 15028 | | | | 15026 | 15026 | | | | | 15027 | 15026 |
| 3522.34 | _ | | | | | | 15026 | | | | 15025 | | 15025 | | | 15025 | | |
| 3456.77 | | | 15025 | | | | | | 15024 | 15024 | | | 15024 | | | | | |
| 3391 20 | | | 15023 | | | | | | | | 15022 | | 15022 | | | 15021 | 15021 | , |
| 3332.39 | 0 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15021 | 15020 | 15020 | 15020 |
| 3315 99 | 0 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | 15020 | |
| 3299.60 | 1 | 15020 | 15020 | 15020 | 15019 | 15019 | 15019 | 15019 | 15019 | 15019 | 15019 | 15019 | 15019 | 15912 | 15018 | 15018 | 15018 | 15018 |
| 3261.03 | 3 | 15018 | 15017 | 15017 | 15016 | 15015 | 15015 | 15014 | 1:013 | 15012 | 15011 | 15010 | 15009 | | | | | |
| 3174.25 | | 15009 | | | 15008 | | | | | 15009 | | | | | 15007 | | | |
| 3141.47 | - | | 15005 | | | | | | | | | | 14998 | | 14996 | | | |
| 3108 68 | | 14991 | | | 14985 | | | | | | | | | | 14981 | | 14992 | |
| 3075 90 | - | | | | 15007 | _ | | | | | 15050 | | 15069 | | 15067 | | | |
| 3043.12 3010.33 | | | 15058 15054 | | | | | | | | 15048 | | 15050 | 15050 15040 | | | 15054 15047 | |
| 2977 55 | | | 15044 | | | | | | | | 15040 | | 15039 | | | | | |
| 2944.76 | | 15038 | | | 15037 | | | | | | | | 15033 | | | | 1 /// 16 | 1, 113 |
| 2918.73 | | | 15035 | | | | | | | | | | 15033 | | | | 15033 | 15033 |
| 2902.34 | 0 | 15033 | | | 15032 | | | | | | 15031 | | 15031 | 15031 | 15031 | 15030 | 15030 | 15030 |
| 2385.95 | 0 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 | 15031 |
| 2868.59 | 1 | 15031 | 15030 | 15030 | 15030 | 15030 | 15029 | 15029 | i 5029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 | 15029 |
| 2835.81 | | 15029 | | 15029 | | | | | 15028 | | | | | | | | | |
| 2803.02 | | | 15026 | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | | 15024 |
| | | | | | | | | | | | | | | | | | | 2 15022 2 15019 |
| | | | | | | | | | | | | | | | | | | 1 15621 |
| | | | 15020 | | | | | | | | | | | | | 1.7721 | 1 1112 | 1.0361 |
| | | | | | | | | | | | | | | | 8 15018 | 15018 | 15011 | 8 15018 |
| | | | | | | | | | | | | | | | | | | 8 15017 |
| | | | | | | | | | | | | | | | | | | 8 15018 |
| 2534.00 |) 1 | 15018 | 3 15018 | 15018 | 15018 | 15018 | 15017 | 15017 | 7 15017 | 1501 | 7 15016 | 5 1501 | 5 15016 | 15016 | 6 15016 | 5 15016 | 5 15019 | 5 15015 |
| 2501.22 | 2 1 | 1501 | 5 15015 | 15015 | 15015 | 15015 | 15015 | 15015 | 5 15015 | 1501 | 4 15014 | 4 150i | 4 15014 | 15014 | 4 15014 | 15014 | 1 15014 | 4 15014 |
| 2468 43 | 3 1 | 15014 | 15014 | 15014 | 15014 | 15014 | 15013 | 15013 | 3 15013 | 1501 | 3 15013 | 3 1501 | 3 | | | | | |
| | | | | | | | | | | | | | | | | | | 1 15011 |
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| | | | | | | | | | | | | | | | | | | 5 15005 |
| | | | | | | | | | | | | | | | | | | 1 15010 |
| 2299.69 | <u>/ 1</u> | 1501 | U 15009 | 1500 | × 12008 | 12008 | 15007 | (1500) | / 15007 | 1 1500 | 0 1500 | טעכו ס | 0 1200 | 1500 | טווכז פ | לונוכן ני | (אולבי ה | 6 15007 |

Table 3.6 - Continued

| 11 11 12 13 14 14 14 14 14 14 14 | | | | | | | | | | | | | | | | | | | |
|--|-------------|----|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|--------|-------|-------|
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| 1 | | | | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | | | |
| 1969 | 2135 77 | ı | 14997 | 14997 | 14997 | 14997 | 14996 | 14996 | 14996 | 14996 | 14996 | 14995 | 14995 | 14995 | 14995 | 14994 | 14994 | 14994 | 14993 |
| . 1976 1 1976 1976 1976 1976 1978 1979 1979 1979 1978 | 2102 99 - 1 | i | 14993 | 14993 | 14993 | 14992 | 14992 | 14992 | 14992 | | | | | | | | | | |
| 1995 1996 | 2090 45 (| 'n | 18992 | 14992 | 14992 | 14992 | 14992 | 14992 | 14992 | 14991 | 14991 | 14991 | 14991 | 14991 | 14991 | 14991 | 14990 | 14990 | 14990 |
| 1948 1949 | 2074-06 (| Ú | 14990 | 14990 | 14990 | 14990 | 14989 | 14989 | 14989 | 14989 | 14389 | 14989 | 14989 | 14988 | 14988 | 14988 | 14988 | 14988 | 14988 |
| | | | | | | | | | | | | | | | | | | | |
| 1997 1997 1998 1999 1999 1999 1999 1999 1999 1999 1999 1999 1999 1999 1999 1999 1999 | | | | | | | | | | | | | | | | _ | | | |
| | | | | | | | | | | | | | | | | | | | |
| 1997 1998 1998 1996 1996 1996 1996 1996 1998 1998 1998 1998 1998 1998 1998 1998 1999 1999 1994 1999 1994 1995 1996 1998 | _ | • | | • | | | | | | | | | | | | | | | |
| 949 19, 1 1498 14998 14996 14996 14996 14996 14996 14996 14986 14986 14997 149 | | - | | | | | | | | | | | | | | | | | |
| 1997 2 1997 1997 1996 1996 1996 1996 1996 1996 1996 1996 1996 1997 1997 1997 1997 1997 1997 1997 1997 1997 1997 1997 1997 1998 1994 1999 1994 199 | | | | | | | | | | | | | | | - | | | | |
| | | | | | | | | | | | | | | | | | | | |
| 1776 1 2 1477 1477 1477 1477 1477 1478 | | - | | _ | | | | | | | | | | | | | | | |
| 1999 1990 | 1776 11 | 2 | 14275 | 14974 | | | | | | | | | | | | | | | |
| 1666 1667 1867 | 1710 54 | 2 | 14952 | 14948 | 14946 | 14943 | 14940 | 14936 | | | | | | | | | | | |
| 1666 1687 1887 | 1689 33 | 1 | 14934 | 14933 | 14932 | 14931 | 14929 | 14927 | 14925 | 14923 | 14921 | 14919 | 14916 | 14914 | 14912 | 14910 | 14907 | 14905 | 14902 |
| 1619 9 | 1656 55 | 1 | 14898 | 14895 | 14893 | 14891 | 14891 | 14892 | 14893 | 14893 | 14892 | 14889 | 14887 | | | | | | |
| 1693 5 0 14768 14758 1476 14756 1476 14723 14708 14673 14673 14673 14673 14673 14673 14673 14673 14673 14673 14673 14673 14673 14673 14673 14768 1 | 1636 30 (| 0 | 14887 | 14887 | 14885 | 14883 | 14880 | 14877 | 14874 | 14871 | 14867 | 14864 | 14861 | 14859 | 14858 | 14857 | 14857 | 14858 | 14858 |
| 1847 1 | 1619 91 (| () | 14857 | 14855 | 14853 | 14849 | 14846 | 14842 | 14837 | 14833 | 14828 | 14823 | 14817 | 14812 | 14806 | 14799 | 14792 | 14785 | 14776 |
| 1.569 1. 1.5971 1. 1.5971 1. 1.5971 1. 1. 1. 1. 1. 1. 1. | 1603 51 (| 0 | 14768 | 14758 | 14747 | 14736 | 14723 | 14708 | 14691 | 14673 | 14655 | 14638 | 14625 | 14615 | 14608 | 14600 | 14588 | 14570 | 14544 |
| 1 | 1587 12 (|) | 14512 | 14495 | 14562 | | | | 15208 | 15264 | 15278 | 15265 | 15238 | 15206 | 15174 | 15145 | 15121 | 15103 | 15091 |
| 1 | 1569 77 | | | | | | | | | | | | | 14995 | 14983 | | | | |
| 1483 9 | | | | | | | | | | - | | | | 14825 | 14814 | 14802 | 14787 | 14770 | 14749 |
| 1487 1486 1487 1488 1484 1488 1484 1488 1484 1488 | | | | | | | | | | | | | | | | | | | |
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| 1433 81 | | | | | | | | | | | | | | | | | | | |
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| 1.586 1.507 1507 1507 1506 1506 1506 1506 1505 | | | | | | | | | | | | | | | | | | | |
| 1335 46 | | | | | | | | | | | | | | | | | | | |
| 1303 64 0 14986 | | | | | | | | | | | | | | | | | | | |
| 1.287 2.5 | | | | | | | | | | | | | | | | | | | |
| 1498 1 1498 1498 1498 1498 1498 1498 1498 1497 1497 1497 14968 14964 14960 14957 14953 14949 14946 14944 14943 14941 14940 14936 14951 1497 14946 1497 14946 | | | | | | | | | | | | | | | | | | | |
| 1237 1 14945 14948 14950 14951 14950 14951 14950 14951 14950 14984 14940 14936 14931 14972 14924 14923 14923 14924 14924 14923 1204 14924 14 | | | | | | | | | | | | | | | | | | | |
| 1171 54 | 1237 11 | 1 | 14945 | 14948 | | | | | | | | | | | | | | | |
| 1138 75 | 1204 32 | 1 | 14920 | 14916 | 14912 | 14908 | 14903 | 14899 | 14895 | 14890 | 14886 | 14881 | 14876 | 14872 | 14868 | 14865 | 14864 | 14862 | 14863 |
| 105 97 1 14671 14627 14580 14532 14486 14441 14382 14306 14213 14113 14088 14388 15078 15634 15803 15745 15610 1074 15 0 15535 15460 15386 15312 15247 15221 15296 15429 15489 15478 15478 15458 15441 15422 15398 15372 15344 15318 10577 0 15293 15299 15247 15226 15207 15189 15172 15155 15139 15124 15109 15094 15080 15066 15056 15052 15038 15024 10141 37 0 15009 14994 14978 14962 14944 14924 14903 14880 14853 14823 14789 14749 14702 14644 14571 14474 14344 1024 97 0 14175 14068 14544 15495 15927 15904 15786 15666 15554 15481 15415 15360 15320 15306 15297 15271 15241 1008 58 0 15211 15182 15151 15114 15072 15018 15009 15199 15377 15361 15310 15276 15248 15227 15209 15194 15181 992 19 0 15169 15160 15156 15150 15143 15137 15132 15126 15121 15117 15114 15110 15107 15104 15100 15096 15092 1974 83 1 15084 15075 15068 15066 15056 15051 15047 15043 15039 15035 15028 15022 15014 15006 14998 14980 942 05 1 14970 14962 14958 14966 14982 14992 14991 14983 14972 14960 14947 14936 14926 14915 14899 14879 14854 909 27 1 14825 14799 14809 14901 15021 15081 15092 15081 15061 15039 15019 15002 14987 14963 14924 14988 14992 14991 14983 14972 14960 14947 14936 14926 14915 14899 14879 14854 843 70 1 14804 14794 14785 14776 14769 14763 14761 1476 | 1171 54 | 1 | 14867 | 14870 | 14869 | 14865 | 14860 | 14856 | 14852 | 14849 | 14847 | 14845 | 14842 | 14837 | 14829 | 14821 | 14810 | 14798 | 14785 |
| 1074 15 0 15335 15460 15386 15312 15247 15221 15296 15429 15489 15478 15 | 1138 75 | ı | 14769 | 14751 | 14731 | 14707 | 14680 | 14652 | 14632 | 14640 | 14697 | 14762 | 14786 | 14792 | 14800 | 14799 | 14779 | 14748 | 14712 |
| 1057 76 0 15293 15269 15247 15226 15207 15189 15172 15155 15139 15124 15109 15094 15080 15066 15052 15038 15024 104137 0 15009 14994 14978 14962 14944 14924 14903 14880 14853 14823 14789 14749 14702 14644 14571 14474 14344 10249 7 0 14175 14068 14544 15495 15927 15904 15786 15666 15564 15481 15415 15360 15320 15306 15297 15271 15241 1008 58 0 15211 15182 15151 15114 15072 15018 15009 15199 15377 15361 15310 15276 15248 15227 15209 15194 15181 1992 19 0 15169 15160 15156 15150 15143 15137 15132 15126 15121 15117 15114 15110 15107 15104 15100 15096 15092 1974 83 1 15084 15075 15068 15062 15056 15051 15047 15043 15039 15035 15028 15022 15014 15006 14998 14989 14980 14920 1 14970 14962 14995 14966 14982 14992 14991 14983 14972 14960 14947 14936 14926 14915 14899 14879 14854 14905 1 14935 14927 14918 14908 14900 14897 14895 14890 14884 14877 14868 14860 14851 14842 14832 14823 14814 843 70 1 14804 14794 14785 14769 14769 14763 14761 14761 14761 14761 14769 14660 14215 14165 14110 14049 13979 13899 13807 13699 13571 13415 13223 12983 12673 12270 11763 11217 745 35 1 10897 11291 12770 15432 18216 19110 18712 18021 17420 16958 16603 16320 16084 15880 14910 15785 15692 15722 15722 15722 15723 15723 15724 15750 14055 16065 16055 16564 16485 16416 16334 670 14 16069 16299 16249 16205 16166 16130 16097 16097 16092 17092 17092 17092 16686 16762 16555 16564 16485 16416 16334 670 14 16097 15904 15901 1 | 1105 97 | i | 14671 | 14627 | 14580 | 14532 | 14486 | 14441 | 14382 | 14306 | 14213 | 14113 | 14088 | 14388 | 15078 | 15634 | 15803 | 15745 | 15610 |
| 1041 37 0 15009 14994 14978 14962 14944 14924 14903 14880 14853 14823 14789 14749 14702 14664 14571 14474 14344 1024 97 0 14175 14068 14544 15495 15927 15904 15786 15666 15564 15481 15415 15360 15320 15306 15297 15271 15241 1008 58 0 15211 15182 15151 15114 15702 15018 15009 15199 15377 15361 15310 15276 15248 15227 15209 15194 15181 992 19 0 15169 15160 15156 15150 15143 15137 15132 15166 15151 15114 15110 15107 15104 15100 15096 15092 1974 83 1 15084 15075 15068 15062 15056 15051 15047 15043 15039 15035 15022 15014 15006 14998 14998 14989 14989 14920 14930 | 1074 15 | 0 | | | 15386 | 15312 | 15247 | 15221 | 15296 | 15429 | 15489 | 15478 | 15458 | 15441 | 15422 | 15398 | 15372 | 15344 | 15318 |
| 1024 97 0 14175 14068 14544 15495 15927 15904 15786 15666 15564 15481 15415 15360 15320 15306 15297 15271 15241 1508 58 0 15211 15182 15151 15114 15072 15018 15009 15199 15377 15361 15310 15276 15248 15227 15209 15194 15181 1992 19 0 15169 15160 15165 15150 15143 15137 15132 15126 15121 15117 15114 15110 15107 15104 15100 15096 15092 1748 1 15084 15075 15068 15062 15056 15051 15047 15043 15039 15035 15028 15022 15014 15006 14998 14989 14989 14980 1942 05 1 14970 14962 14958 14966 14982 14992 14991 14983 14972 14960 14947 14936 14926 14915 14899 14879 14854 14909 27 1 14825 14799 14809 14901 15021 15081 15092 15081 15061 15039 15019 15002 14987 14963 14953 14954 14837 14864 14794 14785 14776 14769 14763 14761 14761 14761 14761 14768 14761 14764 14765 14769 14696 14915 14980 14994 14978 14978 14978 14978 14978 14978 14979 14989 14899 14 | | | | | | | | | | | | | | | | 15066 | | 15038 | 15024 |
| 1008 58 0 15211 15182 15151 15114 15072 15018 15009 15199 15377 15361 15310 15276 15248 15227 15209 15194 15181 1992 19 0 15169 15160 15156 15150 15143 15137 15132 15126 15121 15117 15114 15110 15107 15104 15100 15096 15092 1974 83 1 15084 15075 15068 15062 15056 15051 15047 15043 15039 15035 15028 15022 15014 15006 14998 14980 14980 142 05 1 14970 14962 14958 14966 14982 14992 14991 14983 14972 14960 14947 14936 14926 14915 14899 14879 14854 14902 14918 14988 14900 14897 14895 14890 14884 14877 14868 14860 14851 14842 14832 14814 84370 1 14804 14794 14785 14776 14769 14763 14761 14761 14761 14758 14751 14742 14731 14718 14705 14692 14680 14810 1 14609 14656 14641 14625 14608 14591 14572 14552 14531 14509 14486 14460 14433 14404 14373 14338 14300 778 13 1 14260 14215 14165 14110 14049 13979 13899 13807 13699 13571 13415 13223 12983 12673 12270 11763 11217 14535 1 10897 11291 12770 15432 18216 19110 18712 18021 17420 16958 16603 16320 16084 15880 14913 14908 14909 16540 14656 14641 14625 14608 14591 14502 15018 14895 | | | | | | | | | | | | | | | | | | | |
| 992 19 0 15169 15160 15165 15150 15150 15143 15137 15132 15126 15121 15117 15114 15110 15107 15104 15100 15096 15092 974 83 1 15084 15075 15068 15062 15056 15051 15047 15043 15039 15035 15028 15022 15014 15006 14998 14989 14980 942 05 1 14970 14962 14955 14966 14982 14992 14991 14983 14972 14960 14947 14936 14926 14915 14899 14879 14854 909 27 1 14825 14799 14809 14901 15021 15081 15092 15081 15061 15039 15019 15002 14987 14974 14963 14953 14954 876 48 1 14935 14927 14918 14908 14900 14897 14895 14890 14884 14877 14868 14860 14851 14842 14832 14832 14814 84370 1 14804 14794 14785 14776 14769 14769 14769 14761 14761 14761 14761 14758 14751 14742 14731 14718 14705 14692 14686 14091 1 14669 14666 14661 14625 14608 14591 14572 14552 14531 14509 14486 14460 14433 14404 14373 14338 14300 14818 14906 14761 | | | | | | | | | | | | | | | | | | | |
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| 942 05 | | | | | | | | | | | | | | | | | | | |
| 909 27 | | | | | | | | | | | | | | | | | | | |
| 876 48 | | | | | | | | | | | | | | | | | | | |
| 843 70 | | | | | | | | | | | | | | | | | | | |
| 810 91 | 843 70 | · | 14804 | 14704 | 14785 | 14776 | 14760 | 14763 | 14093 | 14761 | 14761 | 140// | 14008 | 14600 | 14031 | 14719 | 14034 | 14600 | 14614 |
| 778 13 | | | | | | | | | | | | | | | | | | | |
| 745 35 | | | | | | | | | | | | | | | | | | | |
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| 686 53 0 14308 14992 16271 17318 17725 17728 17583 17392 17202 17032 16886 16762 16655 16564 16485 16416 16354 670 14 0 16299 16249 16205 16166 16130 16097 16070 16070 16052 16048 16053 16055 16046 16030 16011 15991 15971 15951 653 74 0 15932 15913 15894 15877 15861 15845 15830 15816 15803 15790 15777 15765 15754 15743 15732 15721 15711 637 35 0 15702 15693 15604 15607 | 702 92 | 0 | 14554 | 15526 | 16281 | 16616 | 16640 | 16504 | 16314 | 16107 | 15897 | 15692 | 15490 | 15322 | 15155 | 14076 | 14760 | 14507 | 14280 |
| 670 14 0 16299 16249 16205 16166 16130 16097 16070 16052 16048 16053 16055 16046 16030 16011 15991 15971 15951 653 74 0 15932 15913 15894 15877 15861 15845 15830 15816 15803 15790 15777 15765 15754 15743 15732 15721 15711 637 35 0 15702 15693 15684 15675 15667 15659 15651 15643 15635 15628 15621 15614 15607 15601 15607 15604 15509 15535 15532 15532 15532 15533 15547 15561 15570 15570 15570 15576 155 | 686 53 | 0 | 14308 | 14992 | 16271 | 17318 | 17725 | 17728 | 17583 | 17392 | 17202 | 17032 | 16886 | 16762 | 16655 | 16564 | 16485 | 16416 | 16354 |
| 653 74 0 15932 15913 15894 15877 15861 15845 15830 15816 15803 15790 15777 15765 15754 15743 15732 15721 15711 637 35 0 15702 15693 15684 15675 15667 15659 15651 15643 15635 15628 15621 15614 626 26 -1 15611 15607 15604 15601 15597 15594 15590 15586 15583 15579 15576 15572 15568 15564 15560 15556 15552 618 07 -1 15548 15544 15539 15535 15532 15532 15532 15533 15547 15561 15570 15579 15578 15576 15576 15576 15576 15576 15576 | 670 14 | 0 | 16299 | 16249 | 16205 | 16166 | 16130 | 16097 | 16070 | 16052 | 16048 | 16053 | 16055 | 16046 | 16030 | 16011 | 15991 | 15971 | 15951 |
| 637 35 0 15702 15693 15684 15675 15667 15659 15651 15643 15635 15628 15621 15614 626 26 -1 15611 15607 15604 15601 15597 15594 15590 15586 15583 15579 15576 15572 15568 15564 15560 15556 15552 618 07 -1 15548 15544 15539 15535 15532 15532 15539 15531 15533 15547 15561 15570 15579 15578 15576 15576 15576 15576 15576 15576 | | | | | | | | | | | | | | | | | | | |
| 626 26 -1 15611 15607 15604 15601 15597 15594 15590 15586 15583 15579 15576 15572 15588 15564 15560 15556 15552 618 07 -1 15548 15544 15539 15535 15532 15529 15531 15533 15547 15561 15570 15579 15578 15576 15572 15567 15563 | | | | | | | | | | | | | | | | , | | | |
| 618 07 -1 15548 15544 15539 15535 15532 15529 15531 15533 15547 15561 15570 15579 15578 15576 15572 15567 15563 | 626 26 - | 1 | 15611 | 15607 | 15604 | 15601 | 15597 | 15594 | 15590 | 15586 | 15583 | 15579 | 15576 | 15572 | 15568 | 15564 | 15560 | 15556 | 15552 |
| 609 87 -1 15558 15555 15551 15548 15546 15543 15540 15538 15535 15533 15530 15528 15525 15523 15521 15518 15516 | 618.07 - | 1 | 15548 | 15544 | 15539 | 15535 | 15532 | 15529 | 15531 | 15533 | 15547 | 15561 | 15570 | 15579 | 15578 | 15576 | 15572 | 15567 | 15563 |
| | 609 87 - | 1 | 15558 | 15555 | 15551 | 15548 | 15546 | 15543 | 15540 | 15538 | 15535 | 15533 | 15530 | 15528 | 15525 | 15523 | 15521 | 15518 | 15516 |

Table 3.6 - Continued

| cm ⁻¹ | Æ | <u> </u> | 1_ | _:_ | ; | _ + | <u> </u> | | | 8 | ٠, | 10 | 11 | 12 | 13 | 14 | 15 | 10 |
|------------------|----|----------|-------|-------|-------|-------|----------|-------|-------|--------|-------|-------|-------|-------|-------|-------|-------|----------|
| 601 19 | o | 15512 | 15508 | 15503 | 15400 | 15495 | 15401 | 15487 | 15483 | 15.179 | 15476 | 15472 | 15408 | 15405 | 15461 | 15458 | 15454 | 12471 |
| 584 80 | 0 | 15447 | 15444 | 15441 | 15438 | 15435 | 15432 | 15429 | 15426 | 15423 | 15420 | 15417 | 15414 | 15412 | 1447 | 15416 | 15403 | 15-8(4) |
| 568 41 | o | 15397 | 15395 | 15392 | 15389 | 15386 | 15384 | 15381 | 15378 | 15375 | 15373 | 15370 | 15307 | 15305 | 15362 | 15300 | 15358 | 15357 |
| 552 02 | 0 | 15355 | 15353 | 15351 | 15349 | 15347 | 15346 | 15344 | 15342 | 15340 | 15338 | 15336 | 14333 | 15441 | | | | |
| 539 97 | -1 | 15330 | 15328 | 15327 | 15326 | 15324 | 15323 | 15322 | 15320 | 15319 | 15318 | 15318 | 15317 | 15317 | 15317 | 15317 | 15318 | 15317 |
| 531 77 | -1 | 15317 | 15316 | 15315 | 15314 | 15313 | 15312 | 15310 | 15310 | 15300 | 15308 | 15300 | 15305 | 15303 | 15302 | 15300 | 15298 | 15290 |
| 516 34 | 3 | 15268 | 15239 | 15201 | 15146 | 15045 | | | | | | | | | | | | |
| 484 52 | 0 | 15025 | 15001 | 14976 | 14947 | 14914 | 14877 | 14833 | 14784 | 14723 | 14652 | 14563 | 14453 | 14312 | 14137 | 13011 | 13787 | 1 517424 |
| | | | | | | | 16390 | | | | | | | | | | | |
| | | | | | | | 15555 | | | | | | | | | | | |
| | | 15471 | | | | | | | | | | | | | . ••• | | | |

Note: Footnotes follow Table 3.7

the refractive index at five wavelengths in the visible region to $n^2(\widetilde{v}) = a \widetilde{v}^4 + b \widetilde{v}^2 + c \widetilde{v}$ and extrapolating to 8000 cm⁻¹. $k(\widetilde{v})$ is less than 5 x10⁻⁵ above 4800 cm⁻¹, apart from a doublet of height 2.3 x40⁻⁶ sees 5000 cm⁻¹. We did not make reliable measurements in this region, so we assumed that $k(\widetilde{v}) = 0.0$ between 4300 cm⁻¹. The real refractive index spectrum is shown in Figure 3.2 and tabulated in Compact Table format in Table 3.6.

3.2.3 - Molar absorption coefficient spectrum.

The molar absorption coefficient spectrum was calculated from the unweighted average $k(\tilde{\nu})$ spectrum. The molar concentration of liquid chlorobenzene at 25°C is 9.783 mole L⁻¹, as calculated from the density¹² 1.10118 g cm⁻³. The $E_m(\tilde{\nu})$ spectrum is shown in Figure 3.3 and the values are tabulated in Table 3.7. The molar absorption coefficient peak heights are listed in Table 3.8.

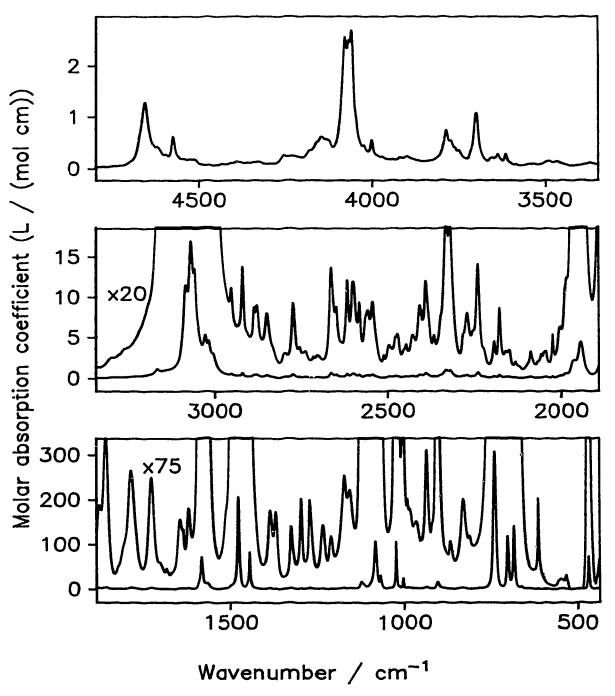


Figure 3.3 - Molar absorption coefficient, $E_m(\tilde{\nu})$ spectrum between 4800 and 435 cm⁻¹ of chlorobenzene at 25°C. The ordinate scale labels in the middle and bottom boxes are for the lower spectrum in the box; they must be divided by 20 and 75, as shown, for the upper spectrum in the box.

Table 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients between 4800 and 435 cm⁻¹ of liquid chlorobenzene at 25°C abservable 3.7 - Molar absorption coefficients at 3.7 - Molar absorption c

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|--|------------------|-------------|----|--------|-------|------|------|---------|------|------|---------|-------|---------|-------|--------|--------|-------|-------|------|---------|
| . 1973 56 0 | cm ⁻¹ | λE | ΥE | 0 | 1 | | 3 | 4 | 5 | - 6 | 7 | 8 | ١) | 10 | 11_ | 12 | 13 | 1-4 | 15 | 10 |
| | | | | | | | | | | | | | 3340 | | 3520 | 3501 | 3535 | 3551 | 1640 | 3252 |
| 1971 1972 1972 1973 1974 1975 1974 1975 1974 1975 1974 1975 1974 1975 | | | | | | | | | | | | | | | | | | | | |
| 1466 1667 1668 | | | | | | | | | | | | | | | | | | | | |
| 1400 1500 1700 | | | | | | | | | | | | | | | | | | | | |
| 4534 92 2 4 1706 1709 1830 1820 1810 1875 1832 1832 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1832 1833 1833 | | | | | | | | | | | | | | | | | | | | |
| 1499 1499 1499 1599 1591 1599 1591 1599 | | | | | | | | | | | | | | | | | | | | |
| 4-049 1-24 138 11-20 11-49 13-56 13-79 13-75 13-79 | | | | | | | | | | | | | | | | | | | | |
| 1472 1472 1473 1474 | 4403.65 | 2 | -4 | 1138 | 1120 | 1146 | | | 1325 | 1311 | | | | | | | | | | |
| | 4338 08 | 2 | -4 | 1235 | 1251 | 1308 | 1336 | 1355 | 1277 | 1128 | 1015 | 945 | 932 | 937 | 130, 3 | .>,,, | 967 | 100 | 1030 | 1114 |
| 441418 2 3 5 5 5 5 5 5 5 5 5 | | | | | | | 1995 | 2343 | 2580 | 2471 | 2362 | | 2361 | 2380 | 2403 | 2445 | 2421 | 2408 | 2369 | 2318 |
| 40479 5 3 2918 2918 240 235 2392 2483 2514 2477 2497 2497 2497 2494 2497 | | | | | | | | 2561 | | | | | | 3978 | 4436 | 4982 | 5232 | | | 6183 |
| Model Mode | | | | | | | | | | | | | | | | | | | | |
| 90117 | | | | | | | | | | | | | | | | | | | | |
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| 3946.32 2 4 1307 2996 2193 236.2 2998 2192 9197 999 818 178 1798 1044 1578 1444 1548 1518 319 2418 319 319 319 319 319 319 319 319 319 319 | | | | | | | | | | | | | | | | | | | | |
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| 1785 1 | | | | | | | | | | | | | | | | | | | | , , , , |
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| 3655-60 1 4 1213 2149 2128 2062 2739 2520 2742 2766 2525 772 3789 2770 8188 1790 2730 772 38984 1 5 7498 7760 8108 8522 7892 7374 7647 6798 6637 6086 6410 6433 6417 6020 6737 6700 9100 9110 9111 1600 6377 6700 6787 6700 9112 1000 9488 917 1000 9488 917 1000 9488 917 1000 9488 918 1000 9488 905 8088 917 8788 987 9788 1088 11717 1121 1121 1170 1008 9088 9871 9010 9878 1088 11717 1121 1170 1008 908 9888 918 902 1014 1007 1008 9089 9889 9899 1014 | 3720.97 | 1 | -3 | 229 | 263 | 306 | 346 | 382 | 428 | 499 | 606 | 748 | 912 | 1037 | 1084 | 1060 | 953 | 790 | 632 | |
| \$\frac{9}{352} \frac{1}{2} \ | | I | 4 | 4278 | 3688 | 3254 | 2898 | 2585 | 2319 | 2100 | 1929 | 1790 | 1671 | 1560 | 1479 | 1454 | 1526 | 1714 | 1944 | 2004 |
| 1889 1 | | | | | | | | | - | | | | | | | | 1007 | | 1780 | 1630 |
| S550 1 | | | | | | | | | | | | | | | | | | | | |
| 352.42 2 .4 873 890 937 994 072 1185 1302 1411 1500 1522 1435 1329 1285 1312 1413 1472 191 3557 2 .4 1207 1026 906 843 786 722 666 623 605 614 611 976 767 776 776 777 777 778 778 778 778 7 | | | | | | | | | | | | | | | | | | | | |
| 3391 20 2 4 1207 1026 906 938 786 722 966 978 1068 1171 1215 1170 1068 908 809 809 785 714 706 607 6 | | | | | | | | | | | | | | | | | | | | |
| 3331 29 | | | | | | | | | | | | | | | | | | | | |
| 3315 99 0 | | | | | | | | | | | | | | | | | | | | 1, 1-4 |
| 33199 0 | 3332 39 | 0 | -5 | | _ | | | | | | | | | | | | | | | 8871 |
| 3261 03 3 | 3315 99 | 0 | -4 | 905 | 953 | 968 | 992 | 1014 | 1027 | 1066 | 1094 | 1108 | 1137 | 1178 | 1191 | | 1230 | | | |
| 3174.25 1 -3 784 862 948 1055 1154 1166 1109 1043 992 939 940 936 940 953 972 995 1025 314147 1 -3 1049 1067 1081 1095 1111 1132 1155 1183 1215 1252 1252 1266 1341 1355 1456 1524 1645 1694 3108.68 1 -2 179 191 205 224 249 284 330 394 478 573 2666 872 1057 1150 1137 11096 1071 307590 1 -2 1127 1302 1543 1690 1665 1537 1395 1313 1321 1363 1289 1099 911 788 717 649 882 3043.12 1 -3 5282 4851 4506 4242 4099 4119 4299 4626 5035 5357 5040 4529 4301 4476 4612 4341 8845 3010.33 1 -3 3414 3160 3088 3084 2949 2666 2329 2015 1754 1584 1382 1246 1126 1025 936 800 800 92918.73 0 4 748 4188 3776 3525 3375 3251 3191 3182 3209 3270 3401 3618 4093 5170 6752 2918.73 0 4 6916 6424 5650 4914 4319 3866 3543 3317 3162 3063 2991 2040 2055 2870 2833 2788 2789 2902.34 0 4 2693 2647 2607 2581 2562 2555 2552 2564 2579 2607 2651 2725 2836 3020 3296 3686 4110 2885.95 0 4 3289 2642 482 2407 2386 2440 4100 4211 4333 4440 4490 4472 4374 4207 3996 3762 3514 3272 2836.002 1 4 2879 2624 2482 2407 2386 2490 1374 1490 1491 3819 3409 3491 2730 2830.02 1 4 2879 2624 2482 2407 2386 2490 1374 1490 1491 3819 3409 3491 2730 2431 2210 2835.81 1 4 2076 1991 1885 1762 1633 1499 1373 1298 1484 3490 4472 4374 4207 3996 3762 3514 3272 2830.02 1 4 2879 2604 1993 1766 1632 1569 1583 1596 1611 1640 1644 2773 45 1 4 1646 1568 1489 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1596 1611 1640 1644 2773 45 1 4 1646 1568 1489 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1590 1583 1596 1611 1640 1644 2773 45 1 4 1646 1568 1489 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1596 1511 1510 1175 1235 1238 1427 1235 1238 1427 1235 1238 1340 2400 1 4 2407 2832 2832 283 283 2136 1168 1127 1109 1104 1122 1149 1246 1333 1597 1305 1238 1427 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1238 1327 1235 1239 1336 1330 1331 1331 1331 1336 1369 1305 1309 1301 1301 1301 1301 1301 1301 1301 | 3299 60 | | | 1287 | 1299 | 1281 | 1288 | 1278 | 1285 | 1298 | 1319 | 1351 | 1376 | 1418 | 1461 | 1503 | 1505 | 1540 | 1581 | 1651 |
| 314147 1 - 3 1049 1067 1081 1095 1111 1132 1155 1183 1215 1252 1296 1341 1195 1456 1524 1605 1694 1310888 1 - 2 179 191 205 224 249 284 330 394 478 573 606 872 1057 1150 1137 1096 1071 307590 1 - 2 1127 1302 1543 1690 1665 1537 1395 1313 1321 1363 1289 1099 111 788 717 649 3013 1492 30133 1 - 3 3414 3160 3088 3084 2949 2666 2329 2015 1754 1548 1382 1246 1126 1025 946 800 796 2977 55 1 4 7830 6977 6596 6283 6066 5773 5339 5328 5121 499 4795 4731 4779 5070 5523 5415 4781 299476 1 - 4 4188 3776 3525 3375 3251 3191 3182 3209 3279 3401 3618 4993 5179 6773 291873 0 - 4 6916 6424 5650 4914 4319 3866 3543 3317 3162 3063 2991 2940 2005 2870 2870 2873 29885 95 0 - 4 4367 4336 4190 4074 4042 4100 4211 4333 4440 4490 4472 4374 4207 3996 3706 3616 4119 2835 81 1 - 4 2076 1991 1885 1762 1633 1499 1373 1492 1493 1513 1685 1994 1999 1027 10706 2835 81 1 - 4 2076 1991 1885 1762 1633 1499 1373 1492 1434 1433 1513 1685 1989 1495 3700 3444 4418 4418 4440 2737 3996 1027 2704 67 1 - 4 1646 1568 1449 1332 1234 1168 1127 1109 1104 1102 1109 1333 1277 1235 1258 1327 2704 67 1 - 4 1646 1568 1449 1332 1255 1269 1180 1169 1170 1192 1230 1300 1418 1507 1885 1272 1270 467 1 - 4 1646 1568 1449 1332 1258 1266 2377 2460 2875 3533 3533 3412 3525 3548 3686 1927 4297 4784 4300 4472 4374 4207 3996 1077 1876 2235 1456 1531 1492 1443 1414 1433 1513 1685 1988 1277 1235 1258 1347 2704 67 1 - 4 1646 1568 1449 1332 1255 1269 1180 1169 1170 1192 1230 1300 1418 1507 1858 1244 2519 2704 67 1 - 4 1646 1568 1449 1332 1255 1269 1180 1169 1170 1192 1230 1300 1418 1507 1858 1244 2519 2704 67 1 - 4 1646 1568 1449 1332 1254 1662 3800 3710 3553 3502 3548 3686 1927 4297 4784 4303 5704 5944 2599 27 1 - 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 1260 1448 1317 1271 129 1290 1300 1418 1507 1858 1244 1260 1254 1448 1449 1449 1449 1449 1449 1449 144 | | | | | | | | 2348 | | | | 4011 | | | | | | | | |
| 3108.68 1 - 2 179 191 205 224 249 284 330 394 478 573 696 872 1057 1150 1137 1096 1071 307590 1 - 2 1127 1302 1543 1690 1665 1537 1395 1313 1321 1363 1289 1099 911 788 717 649 582 304312 1 - 3 5282 4851 4506 4242 4099 4119 4299 4626 5085 5375 5040 4529 4301 4476 4612 4414 1845 3010 33 1 - 3 3414 3160 3088 3084 2949 2666 2329 2015 1754 1548 1382 1246 1126 1025 936 800 796. 2971 555 1 - 4 7430 6977 6596 6283 6006 5773 5539 5328 5121 4939 4795 4731 4779 5070 5523 5415 4781 2994.76 1 - 4 4188 3776 3525 3375 3351 3191 3182 3209 3279 3401 3618 4993 31179 6753 2918.73 0 - 4 6916 6424 5650 4914 4319 386° 3543 3317 3162 3063 2991 2940 2905 2870 2833 2788 2719 2902.34 0 - 4 2693 2647 2607 2581 2562 2555 2552 2564 2579 2607 2651 2725 2836 3020 3279 38086 4119 2885.95 0 - 4 3367 4336 4409 4472 44100 4211 44100 4211 4433 34140 4490 4472 4374 4272 2408 2835.95 0 - 4 3467 4336 4490 4472 44100 4211 4430 4411 4433 3413 1685 1988 233 3444 4418 4540 2835 24 4182 223 1368 1482 1528 1546 1531 1492 1443 1414 1433 1513 1685 1988 2534 3444 4418 4540 2770 24 1 - 4 2879 2624 2482 2407 2386 1546 1531 1492 1443 1414 1433 1513 1685 1988 2535 3444 4418 4540 2770 467 1 - 4 1646 1568 1449 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1429 1374 51 1 - 4 1646 1568 1449 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1429 1370 1363 1444 4118 4540 2773 45 1 - 4 1640 1421 1339 1357 1308 1255 1299 1180 1169 1170 1192 1230 1300 1448 1597 1858 2244 2671.89 1 - 4 2879 2695 6075 5887 5088 4455 4000 3710 3553 3502 3548 3686 3027 4297 4784 5303 5704 5944 2259 2791 270467 1 - 4 1640 1421 1339 1357 1308 1255 1299 1180 1169 1170 1192 1230 1300 1448 1597 1858 2244 2569 2791 270467 1 - 4 1640 1421 1399 1357 1308 1255 1299 1180 1169 1170 1192 1240 1340 14418 1597 1858 2244 2569 2791 270467 1 - 4 1640 1421 1399 1357 1308 1255 1299 1180 1169 1170 1192 1240 1346 1333 1427 1235 1258 1327 1255 1258 1327 1256 1258 1327 1256 1258 1327 1256 1258 1327 1256 1258 1327 1256 1258 1327 1256 1258 1327 1256 1258 1327 1256 1258 1357 1257 1257 1257 1257 1257 1259 1258 125 | | | | | | | | | | | | _ | | | | | | - | | |
| 3075 90 1 2 1 127 1302 1543 1690 1665 1537 1395 1313 1321 1363 1289 1099 911 788 717 649 582 3043 12 1 3 5282 4851 4506 4242 4099 4119 4299 4626 6080 5357 5040 4529 4301 4476 4612 4411 1845 3010 33 1 -3 3414 3160 3088 3084 2949 2666 2329 2015 1754 1548 1382 1246 1126 1025 946 8669 2977 55 1 4 7430 6977 6596 6283 6006 5773 5539 5328 5121 4939 4795 4731 4779 5079 5523 5415 4781 2944 76 1 4 4188 3776 3525 3375 3251 3191 3182 3209 3279 3401 3618 4993 5179 6753 2918 73 0 4 6916 6424 5650 4914 4319 386° 3543 3317 3162 3063 2991 2940 2905 2870 2813 2788 2719 2902 34 0 4 2693 2647 2607 2581 2562 2555 2552 2564 2559 2607 2651 2725 2836 3020 3296 3686 4119 2885 95 0 4 3367 4336 4190 4074 4042 4100 4211 4333 4440 4490 4472 4470 4470 3996 3760 3514 3272 2868 59 1 4 2879 2624 2482 2407 2386 2427 2549 2791 3188 3710 4011 3839 3469 3091 2730 2411 2210 2883 81 1 4 2076 1991 1885 1762 1633 1499 1373 1258 1172 1100 1046 1005 944 991 9991 1078 999 1079 1096 2873 74 1 4 1222 1368 1482 1528 1546 1531 1492 1443 1414 1433 1513 1685 1988 2535 3444 4418 4540 2770 24 1 4 3874 3124 2509 2106 1883 1809 1846 1924 1903 1766 1632 1596 1583 1596 1611 1640 1642 2773 45 1 4 1646 1568 1449 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1277 1235 1258 1327 2704 67 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4984 4154 4982 4268 3671 3063 2641 2639 10 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4984 4154 4382 4268 3671 3063 2641 2639 10 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4984 4154 4382 4268 3671 3063 2641 2536 40 1 4 2907 2281 2228 2288 2186 2166 2237 2460 2845 3389 4229 5468 2636 50 1 4 5977 5830 5303 4555 3911 3535 3412 3525 4063 3639 3732 4429 4404 4487 4478 4471 5470 2481 2536 40 1 4 2977 2526 2667 1574 1845 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1291 1376 2501 22 1 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 1795 1284 1456 1317 1211 1291 1376 2504 44 1 4 14 14 14 14 14 14 14 14 14 14 1 | | | | | | | | | | | | | | | | | | | | |
| 3043 12 1 -3 5282 4851 4506 4242 4099 4119 4299 4626 5080 5357 5040 4529 4101 4476 4632 4141 1845 3010 33 1 -3 3414 3160 3088 3084 2949 2666 2329 2015 1754 1548 1382 1246 1126 1025 936 800 706, 2977 555 1 -4 7430 6977 6596 6283 6006 5773 5539 5328 5121 4939 4795 4731 4779 5070 5523 5435 4781 2944 76 1 -4 4188 3776 3525 3375 3251 3191 3182 3209 3279 3401 3618 4093 5179 6773 2918 73 0 -4 6916 6424 5650 4914 4319 386° 3543 3317 3162 3063 2991 2940 2905 2870 2833 2788 2719 2903.34 0 -4 2693 2647 2607 2581 2562 2555 2552 2564 2579 2607 2651 2725 2836 3020 1296 30.66 4119 2885 95 0 -4 4367 4336 4190 4074 4042 4100 4211 4333 4440 4490 4472 4374 4207 3996 3762 3514 3272 2883 95 1 -4 2879 2624 2482 2407 2386 2427 2549 2791 3188 3710 4011 3839 3469 3091 2730 2431 2210 2835 81 1 -4 2076 1991 1885 1762 1633 1499 1373 1258 1172 1100 1046 1005 984 981 989 1027 1096 2803 02 1 -4 1222 1368 1482 1528 1528 1546 1531 1492 1443 1414 1433 1513 1685 1988 2535 3444 4418 4440 27770 24 1 -4 3874 3124 2509 2106 1883 1809 1846 1924 1903 1766 1632 1569 1583 1596 1611 1640 1644 26718 9 1 -4 1400 1421 1399 1357 1308 1255 1209 1104 1102 1149 1246 1333 1277 1235 1258 1127 2704 67 1 -4 1400 1241 1399 1357 1308 1255 1209 1104 1102 1149 1246 1333 1277 1235 1258 1127 2704 67 1 -4 4007 2281 2282 2208 2186 2166 2237 2460 2845 3389 4229 5468 2535 3611 1690 1490 1490 1490 1490 1490 1490 1490 14 | | | | | | | | _ | | | | | | | | | | | | |
| 3010 33 | | | | | _ | | | | | | | | | | | | | | | |
| 2977 55 1 4 7430 6977 6596 6283 6006 5773 5539 5328 512! 4939 4795 4731 4779 5070 5523 5435 4781 2944 76 1 4 4188 3776 3525 3375 3251 3191 3182 3209 3279 3401 3618 4493 1679 6753 5229 1876 6550 4914 4319 386° 3543 3317 3162 3063 2991 2940 2905 2870 2833 2788 2739 290.34 0 4 2693 2647 2607 2581 2562 2555 2552 2564 2579 2607 2651 2725 2836 3020 3296 3666 4119 2885.95 0 4 4367 4336 4190 4074 4042 4100 4211 4333 4440 4490 4472 4374 4207 3996 3762 3514 3272 2868 59 1 4 2879 2624 2482 2407 2386 2427 2549 2791 3188 3710 4011 3839 3469 3091 2730 2431 2210 2835.81 1 4 2076 1991 1885 1762 1633 1499 1373 1258 1172 1100 1046 1075 984 981 989 1027 1076 2833 2778 2830 20 1 4 1222 1368 1482 1528 1546 1531 1492 1433 1414 1433 1513 1685 1988 2535 3444 4418 4440 2770.24 1 4 3874 3124 2509 2106 1883 1809 1846 1924 1903 1766 1632 1569 1583 1596 1611 1640 1664 2737 45 1 4 1646 1568 1449 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1277 1235 1258 1327 2704.67 1 4 1400 1421 1399 1357 1308 1255 1209 1180 1169 1170 1192 1230 1300 1418 1597 1858 2244 2671.89 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4784 4154 4354 4362 2569 20 4 5976 6075 5687 5058 4455 4000 3710 3553 3502 3548 3686 3927 4297 4784 5303 5704 5942 2566 8 1 4 5977 5830 5303 4555 3911 5353 3412 3525 1209 180 1169 1170 1192 1230 1300 1418 1597 1363 2601 2566 8 1 4 5977 5830 5303 4555 3911 5353 3412 3525 4062 3803 3639 3732 4120 4588 4689 4253 3710 3245 2842 253400 1 4 2407 2811 299 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2504 2488 3 1 4 2493 2153 1785 1557 1472 1490 1587 1701 1707 1816 1899 224 4404 4449 4441 4441 1439 1246 1337 1264 2447 2283 2440 2448 18 0 4 1979 2056 2667 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2448 18 0 4 2497 2659 2657 2603 2505 2396 2331 2350 2499 2808 3305 3202 4404 4404 4404 4447 4241 3912 3459 23864 1 4 2657 2456 2207 2101 2136 2270 2458 2718 2719 368 8762 4042 4508 3307 331 277 2346 2332 480 233248 1 4 3 2657 2456 2207 2101 2136 2270 2458 2718 2719 368 8769 672 568 476 397 | | | | | | | | | | | | | | | | | | | | |
| 2944 76 | | | | | | | | | | | | | | | | | | | | |
| 2918.73 0 | 2944.76 | 1 | 4 | 4188 | | | | | | | | | | | | | | | | |
| 2885.95 0 4 4367 4336 4190 4074 4042 4100 4211 4333 4440 4490 4472 4374 4207 3996, 3762 3514 3272 2868.59 1 4 2879 2624 2482 2407 2386 2427 2549 2791 3188 3710 4011 3839 3469 3091 2730 2431 2210 2835.81 1 4 2076 1991 1885 1762 1633 1499 1373 1258 1172 1100 1046 1095 984 981 989 1027 1096 2803.02 1 4 1222 1368 1482 1528 1546 1531 1492 1443 1414 1433 1513 1685 1988 2535 3444 4418 4540 2770.24 1 4 3874 3124 2509 2106 1883 1809 1846 1924 1903 1766 1632 1569 1583 1596 1611 1640 1664 2737.45 1 4 1646 1568 1449 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1277 1235 1258 1327 2704.67 1 4 1400 1421 1399 1357 1308 1255 1209 1180 1169 1170 1192 1230 1300 1418 1597 1858 2244 2671.89 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4084 4154 4382 4268 3671 3063 2661 2639.10 1 4 2407 2281 2228 2208 2186 2166 2237 2460 2845 3389 4229 5468 261692 0 4 5976 6075 5687 5058 4455 4000 3710 3553 3502 3548 3686 3927 4297 4784 5303 5704 5944 2599.57 1 4 5977 5830 5303 4555 3911 3535 3412 3525 4063 4703 4071 3171 2622 2347 2283 2430 2812 2566.78 1 4 3302 3691 3883 4001 4150 4215 4062 3800 3639 3732 4120 4588 4689 4253 3710 3245 2842 2534.00 1 4 2501 2194 1920 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2501.22 1 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2709 2695 2488.43 1 4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1899 2448 4404 44044 44044 44044 44044 44044 4404 | 2918.73 | 0 | 4 | 6916 | 6424 | 5650 | 4914 | | 386° | 3543 | 3317 | 3162 | 3063 | 2991 | 2940 | 2905 | 2870 | 2833 | 2788 | 2739 |
| 2868 59 1 4 2879 2624 2482 2407 2386 2427 2549 2791 3188 3710 4011 3839 3469 3091 2730 2431 2210 2835 81 1 4 2076 1991 1885 1762 1633 1499 1373 1258 1172 1100 1046 1095 984 981 989 1027 1096 2803 02 1 4 1222 1368 1482 1528 1546 1531 1492 1443 1414 1433 1513 1685 1988 2535 3444 4418 4540 2770 24 1 4 3874 3124 2509 2106 1883 1809 1846 1924 1903 1766 1632 1569 1583 1596 1611 1640 1644 2737 45 1 4 1646 1568 1449 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1277 1235 1258 1127 2704.67 1 4 1400 1421 1399 1357 1308 1255 1209 1180 1169 1170 1192 1230 1300 1418 1597 1858 2244 2671.89 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4084 4154 4382 4268 3671 3063 2661 2639.10 1 4 2407 2281 2228 2208 2186 2166 2237 2460 2845 3389 4229 5468 261692 0 4 5976 6075 5687 5058 4455 4000 3710 3553 3502 3548 3686 3927 4297 4784 5303 5704 5044 2556 78 1 4 3302 3691 3883 4001 4150 4215 4062 3800 3639 3732 4120 4588 4689 4253 3710 3245 2842 2534.00 1 4 2501 2194 1920 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2501.22 1 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2709 2605 2468.43 1 4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1859 1224 1783 1868 1976 2105 2245 24308 1 4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 3059 2389 441 448 1799 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 3059 2389 441 4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2501 2501 2504 2334 2332.48 1 4 3 6577 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4588 5323 6371 7581 8849 2332.48 1 4 3 6577 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4508 5323 6371 7581 8849 2332.48 1 4 3 6577 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4508 5323 6371 7581 8849 2332.48 1 4 3 6577 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4568 5323 6371 7581 8849 2332.48 1 4 3 6577 2456 2207 2101 213 | 2902.34 | 0 | -4 | 2693 | 2647 | 2607 | | 2562 | 2555 | 2552 | 2564 | 2579 | 2607 | 2651 | 2725 | 2836 | 3020 | 3296 | 3686 | 4119 |
| 2835.81 | 2885.95 | 0 | 4 | 4367 | 4336 | 4190 | 4074 | 4042 | 4100 | 4211 | 4333 | 4440 | 4490 | 4472 | 4374 | 4207 | 3996 | 3762 | 3514 | 3272 |
| 2803.02 | | | | | | | | | | | | | | | | | | | | |
| 2770 24 1 4 3874 3124 2509 2106 1883 1809 1846 1924 1903 1766 1632 1569 1583 1596 1611 1640 1664 2737.45 1 4 1646 1568 1449 1332 1234 1168 1127 1109 1104 1122 1149 1246 1333 1277 1235 1258 1327 2704.67 1 4 1400 1421 1399 1357 1308 1255 1209 1180 1169 1170 1192 1230 1300 1418 1597 1858 2244 2671.89 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4/84 4154 4382 4268 3671 3063 2661 2639.10 1 4 2407 2281 2228 2208 2186 2166 2237 <td< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></td<> | | | | | | | | | | | | | | | | | | | | |
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| 2704.67 1 4 1400 1421 1399 1357 1308 1255 1209 1180 1169 1170 1192 1230 1300 1418 1597 1858 2244 2671.89 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4084 4154 4382 4268 3671 3063 2601 2639.10 1 4 2407 2281 2228 2208 2186 2166 2237 2460 2845 3389 4229 5468 2616.92 0 4 5976 6075 5687 5058 4455 4000 3710 3553 3502 3548 3686 3927 4297 4784 5303 5704 5904 2599 57 1 4 5977 5830 5303 4555 3911 3535 3412 3525 4063 4703 4071 3171 2624 2347 2283 2430 2812 2566.78 1 4 3302 3691 3883 4001 4150 4215 4062 3800 3639 3732 4120 4588 4689 4253 3710 3245 2842 2534.00 1 4 2501 2194 1920 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2501.22 1 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2700 2695 2448.18 0 4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 3059 2398.04 1 4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2301 2153 2175 2344 2574 2332.48 1 -3 982 1005 958 897 877 918 987 993 908 789 672 568 476 397 331 277 236 | | | | | | | | | | | | | | | | | | | | |
| 2671.89 1 4 2812 3639 4800 6091 6812 6727 6142 5427 4788 4314 4/84 4154 4382 4268 3671 3063 2661 2639.10 1 4 2407 2281 2228 2208 2186 2166 2237 2460 2845 3389 4229 5468 2616.92 0 4 5976 6075 5687 5058 4455 4060 3710 3553 3502 3548 3686 3927 4297 4784 5303 5704 5704 2599.57 1 4 5977 5830 5303 4555 3911 3535 3412 3525 4063 4703 4071 3171 2624 2347 2283 2430 2812 2566.78 1 4 3302 3691 3883 4001 4150 4215 4062 3800 3639 3732 4120 4588 4689 4253 3710 3245 2842 2534.00 1 4 2501 2194 1920 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2501.22 1 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2700 2695 2468.43 1 4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1859 2448.81 0 4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 3659 2398.04 1 4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2301 2153 2175 2344 2574 2332.48 1 -3 982 1005 958 897 877 918 987 993 908 789 672 568 476 397 331 277 236 | | | | | | | _ | | | | | | | | | | | | | |
| 2639.10 1 4 2407 2281 228 2208 2186 2166 2237 2460 2845 3389 4229 5468 2616.92 0 4 5976 6075 5687 5058 4455 4060 3710 3553 3502 3548 3686 3927 4297 4784 5303 5704 5904 2599.57 1 4 5977 5830 5303 4555 3911 3535 3412 3525 4063 4703 4071 3171 2624 2347 2283 2430 2812 2566.78 1 4 3302 3691 3883 4001 4150 4215 4062 3800 3639 3732 4120 4588 4689 4253 3710 3245 2842 2534.00 1 4 2501 2194 1920 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2501.22 1 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2700 2695 2448.43 1 4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1859 2448.18 0 4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 3659 2332.48 1 -3 982 1005 958 897 877 918 987 993 998 789 672 568 476 397 331 277 236 | | | | | | | | | | | | | | | | | | | | |
| 2616.92 0 .4 5976 6075 5687 5058 4455 4060 3710 3553 3502 3548 3686 3927 4297 4784 5303 5704 5994 2599.57 1 .4 5977 5830 5303 4555 3911 3535 3412 3525 4063 4703 4071 3171 2624 2347 2283 2430 2812 2566.78 1 .4 3302 3691 3883 4001 4150 4215 4062 3800 3639 3732 4120 4588 4689 4253 3710 3245 2842 2534.00 1 .4 2501 2194 1920 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2501.22 1 .4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2700 2695 2468.43 1 .4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1859 2448.18 0 .4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 .4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 1659 2398.04 1 .4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2301 2153 2175 2344 2574 2332.48 1 .3 982 1005 958 897 877 918 987 993 998 789 672 568 476 397 331 277 236 | | | | | | | | | | | | | | | | 7,112 | -4130 | | | 2.571 |
| 2599.57 4 5977 5830 5303 4555 3911 3535 3412 3525 4063 4703 4071 3171 2624 2347 2283 2430 2812 2536.08 4 3302 3691 3883 4001 4150 4215 4062 3800 3639 3732 4120 4588 4689 4253 3710 3245 2842 2534.00 4 2501 2194 1920 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2501.22 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2700 2695 2468.43 4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1859 2448.18 0 4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 1659 2398.04 4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2301 2153 2175 2344 2574 2365.26 4 3 2657 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4508 5323 6371 7581 8849 2332.48 1 3 982 1005 958 897 877 918 987 993 998 789 672 568 476 397 331 277 236 | | | | | | | | | | | | | | | | 4297 | 4784 | 5303 | 5704 | 5904 |
| 2566.78 1 -4 3302 3691 3883 4001 4150 4215 4062 3800 3639 3732 4120 4588 4689 4253 3710 3245 2842 2534.00 1 -4 2501 2194 1920 1665 1454 1288 1175 1108 1069 1058 1105 1284 1456 1317 1211 1239 1376 2501.22 1 -4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2700 2695 2468.43 1 -4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1859 2448.18 0 -4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 -4 | | | | | | | | | _ | | | | | | | | | | | |
| 2501.22 1 4 1588 1835 2004 2037 1978 1895 1826 1787 1787 1840 1981 2263 2533 2517 2574 2700 2695 2468.43 1 4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1859 2448.18 0 4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 3659 2398.04 1 4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2301 2153 2175 2344 2574 2365.26 1 4 2657 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4508 5323 6371 7581 8849 2332.48 1 3 982 1005 958 897 877 918 987 993 908 789 672 568 476 397 331 277 236 | | | | | | | | | | | | | | | | | | | | |
| 2468.43 1 4 2493 2153 1785 1557 1472 1490 1587 1701 1776 1816 1899 2448.18 0 -4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 -4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4494 4487 4241 3912 3659 2398.04 1 -4 3573 3764 4401 5440 5998 5618 4978 4402 3999 3461 3016 2601 2301 2153 2175 2344 2574 2365.26 1 -4 2657 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4942 4508 5323 6371 7581 8849 2332.48 1 -3 | 2534.00 | 1 | 4 | 2501 | 2194 | 1920 | | | | | | | | | 1284 | 1456 | 1317 | 121 | 1239 | 1376 |
| 2448.18 0 -4 1979 2056 2067 1974 1847 1747 1685 1653 1650 1659 1683 1724 1783 1868 1976 2105 2245 2430.83 1 -4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 3659 2398.04 1 -4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2301 2153 2175 2344 2574 2365.26 1 -4 2657 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4508 5323 6371 7581 8849 2332.48 1 -3 982 1005 958 897 877 918 987 993 998 789 672 568 476 397 331 277 236 | | | 4 | 1588 | 1835 | 2004 | 2037 | 1978 | 1895 | 1826 | 1787 | 1787 | 1840 | 1981 | 2263 | 2533 | 2517 | 2574 | 2700 | 2695 |
| 2430.83 1 -4 2497 2629 2657 2603 2505 2396 2331 2350 2499 2808 3305 3920 4404 4487 4241 3912 3659 2398.04 1 -4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2301 2153 2175 2344 2574 2365.26 1 -4 2657 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4508 5323 6371 7581 8849 2332.48 1 -3 982 1005 958 897 877 918 987 993 908 789 672 568 476 397 331 277 236 | | | | | | | | | | | | 1776 | 1816 | | | | | | | |
| 2398.04 1 -4 3573 3764 4401 5440 5998 5618 4978 4402 3909 3461 3016 2601 2301 2153 2175 2344 2574 2365.26 1 -4 2657 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4508 5323 6371 7581 8849 2332.48 1 -3 982 1005 958 897 877 918 987 993 908 789 672 568 476 397 331 277 236 | | | | | | | | | | | | | | | | | | | | |
| 2365.26 1 -4 2657 2456 2207 2101 2136 2270 2458 2718 3112 3586 3862 4042 4508 5323 6371 7581 8849 2332.48 1 -3 982 1005 958 897 877 918 987 993 908 789 672 568 476 397 331 277 236 | | | | | | | | | | | | | | | | | | | | |
| 2332.48 1 -3 982 1005 958 897 877 918 987 993 908 789 672 568 476 397 331 277 236 | | | | | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | | | | |
| | | | | | 1774 | | | 1499 | 1559 | 1781 | 2191 | | | | | | | | | |

Table 3.7 -Continued

| Table | | -c. | | | | | | | | | | | | | | | | | |
|--------------------|-----|----------|--------------|--------------|--------------|--------------|--------------|--------------|--------------|-----------------------|--------------|--------------|--------------|--------------|--------------|--------------|------------|--------------|-------------|
| tm ^t | XI: | YI | | 1_ | 2 | 3 | 4 | 5 | -5 | 7 | - 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | <u>!6</u> |
| 2266.91 | 1 | 4 | 3325 | 2988 | 2750 | 2543 | 25/5 | 2786 | 2907 | 2947 | 2967 | 3115 | 3553 | 4360 | 5522 | 5714 | 7001 | 5050 | 4764 |
| 2234 12 | 1 | -4 | 3692 | 2883 | 2335 | 2024 | 1894 | 1850 | 1805 | 1685 | 1508 | 1316 | 1135 | 997 | 908 | 861 | 857 | 894 | 981 |
| | 1 | 4 | 1131 | 130 | 1695 | 2099 | 2257 | 1948 | 1607 | 1463 | 1556 | 1937 | 2756 | 3921 | 4297 | 3513 | 2585 | 2011 | 1715 |
| 2158 56 2135 77 | 1 | न न | 1552 465 | 1468 979 | 1475 1918 | 1561 917 | 1661 830 | 1676 783 | 1616 766 | 1594 753 | 1645 737 | 1725 704 | 1700 649 | 1504 | 1211 | 983 | 858 | 803 | 401 5 59 |
| 2102 99 | i | 4 | 715 | 780 | 847 | 901 | 998 | 1080 | 1338 | 793 | 757 | //)-4 | 6449 | 603 | 574 | 568 | 578 | 613 | 6.58 |
| 2090 45 | Ġ | 4 | 1500 | 1622 | 1609 | 1505 | 1396 | 1314 | 1248 | 1183 | 1119 | 1051 | 989 | 940 | 903 | 880 | 865 | 858 | 356 |
| 2074-06 | á | -4 | 861 | 874 | 898 | 934 | 974 | 1012 | 1036 | 1054 | 1070 | 1085 | 1104 | 1127 | 1159 | 1211 | 1296 | | 1507 |
| 2057-07 | 6 | -4 | 1505 | 1436 | 1385 | 1374 | 1401 | 1457 | 1528 | 1595 | 1638 | 1654 | 1653 | 1653 | 1664 | 1667 | | 1490 | 1352 |
| 2041-28 | 6 | 4 | 1231 | 1135 | 1073 | 1031 | 1006 | 992 | 986 | 987 | 995 | 1012 | 1046 | 1101 | 1182 | 1285 | | 1595 | 1918 |
| 2024 89 | O | -1 | 2374 | 2715 | 2606 | 2238 | 1921 | 1711 | 1585 | 1508 | 1469 | 1457 | 1470 | 1507 | 1563 | 1633 | 1729 | 1850 | 1999 |
| 2008 49 | O | -4 | 2187 | 2400 | 2623 | 2831 | 2999 | 3119 | 3170 | 3151 | 3095 | 3042 | 3007 | 2998 | 3028 | 3105 | 3244 | 3462 | 3779 |
| 1992 10 | O | 4 | 4204 | 4699 | 5156 | 5466 | 5633 | 5755 | 5904 | 6069 | 6191 | 6241 | 6235 | 6221 | 6253 | 6372 | 6615 | 6998 | 7543 |
| 1974 74 | 1 | - 3 | 916 | 1146 | 1426 | 1721 | 1985 | 2171 | 2258 | 2258 | 2217 | 2186 | 2212 | 2332 | 2574 | 2969 | | 4028 | 1386 |
| 1941.96 | 1 | . 3 | 4396 | 4108 | 3622 | 3053 | 2499 | 2018 | 1636 | 1357 | 1165 | 1020 | 895 | 784 | 692 | 618 | 563 | 529 | 503 |
| 1907 25 | 2 | -3 | 485 | 543 | 656 | 868 | 1264 | 1895 | 2407 | 2493 | 2356 | 2541 | 3334 | 4277 | 4540 | 3579 | | 1244 | 768 |
| 1841 68 | 2 | -3 .3 | 546 | 459 | 4/6 | 562 | 673 | 750 | 896 | 1081 | 1261 | 1400 | 1557 | 1876 | 2481 | 3223 | | 3264 | 2690 |
| 1776 11 1710 54 | 2 | ! | 2047 8276 | 1615 7988 | 1138 7684 | 846 6783 | 585 5828 | 605 5004 | 609 | 699 | 928 | 1384 | 2137 | 3021 | 3314 | 2802 | 2022 | 1385 | 1005 |
| 1689 33 | 1 | -4 | 5063 | 5731 | 6065 | 5318 | 4983 | 4564 | 4205 | 4225 | 4386 | 4687 | 4982 | 5195 | 5376 | 5649 | 6035 | 6528 | 7322 |
| 1656 55 | i | -3 | 858 | 1050 | 1315 | 1595 | 1865 | 2042 | 2077 | 1992 | 1851 | 1753 | 1783 | -133 | J.J. 10 | | 5000 | 0040 | 1000 |
| 1636 30 | ò | .3 | 1769 | 1656 | 1487 | 1340 | 1241 | 1188 | 1189 | 1223 | 1300 | 1431 | 1613 | 1839 | 2078 | 2280 | 2388 | 2405 | 2338 |
| | 0 | - 3 | 2219 | 2078 | 1930 | 1805 | 1706 | 1621 | 1570 | 1551 | 1544 | 1563 | 1603 | 1653 | 1710 | 1790 | | 2012 | 2170 |
| 1603.51 | O | -2 | 236 | 260 | 288 | 323 | 367 | 425 | 505 | 617 | 767 | 958 | 1185 | 1425 | 1657 | 1879 | | 2402 | 2804 |
| 1587 12 | 0 | -2 | 3437 | 4526 | 6065 | 7099 | 7079 | 6483 | 5545 | 4510 | 3571 | 2813 | 2248 | 1849 | 1584 | 1420 | 1332 | 1290 | 1258 |
| 1569 77 | 1 | -2 | 1203 | 1237 | 1286 | 1170 | 939 | 715 | 541 | 406 | 309 | 240 | 194 | 161 | 139 | 124 | 114 | 105 | 99 |
| 1536 98 | ı | -3 | 932 | 915 | 943 | 994 | 1046 | 1104 | 1189 | 1288 | 1447 | 1627 | 1853 | 2080 | 2261 | 2377 | 2484 | 2688 | 3052 |
| 1504 20 | 1 | -2 | 370 | 477 | 627 | 778 | 873 | 933 | 1024 | 1178 | 1394 | 1714 | 2274 | | | | | | |
| | 0 | -1 | 274 | 343 | 449 | 624 | 926 | 1413 | 1950 | 2032 | 1758 | 1226 | 833 | 604 | 473 | 380 | 298 | 224 | 171 |
| | 0 | -2 -2 | 1360 1588 | 1122 | 959 | 842 | 757 3979 | 697 | 658 | 638 | 634 | 648 | 680 | 735 | 811 | 914 | | 1206 | 1387 |
| 1433 81 | 1 | -3 | 4123 | 1818 3348 | 2145 2838 | 2723 2427 | 2075 | 6457 1766 | 8329 1501 | 6310 1286 | 4057 1118 | 2734 1004 | 1968 917 | 1539 832 | 1309 773 | 1103 739 | 874 719 | 687 720 | 560 736 |
| 1401 03 | i | -3 | 763 | 817 | 920 | 1098 | 1372 | 1747 | 2135 | 2339 | 2273 | 2049 | 1799 | 1626 | 1582 | 1699 | 1961 | 2240 | 736 2305 |
| 1368 24 | i | -3 | 2074 | 1694 | 1324 | 1017 | 791 | 634 | 532 | 464 | 420 | 391 | 373 | 363 | 360 | 363 | 371 | 386 | 408 |
| 1335 46 | i | -3 | 443 | 503 | 616 | 845 | 1298 | 1822 | 1698 | 1333 | 1131 | 975 | 839 | 756 | 726 | 737 | 784 | 875 | 1041 |
| 1303 64 | 0 | -3 | 1172 | 1358 | 1620 | 1959 | 2339 | 2641 | 2705 | 2499 | 2141 | 1760 | 1447 | 1217 | 1059 | 957 | 895 | 858 | 837 |
| 1287 25 | 0 | -3 | 825 | 817 | 813 | 815 | 823 | 840 | 867 | 308 | 965 | 1049 | 1174 | 1365 | 1658 | 2069 | 2465 | | 2624 |
| 1269 89 | 1 | -3 | 2345 | 2046 | 1725 | 1398 | 1136 | 945 | 807 | 714 | 659 | 636 | 643 | 682 | 760 | 885 | 1071 | 1313 | 1582 |
| 1237 11 | 1 | -3 | 1814 | 1911 | 1812 | 1591 | 1350 | 1127 | 962 | 871 | 851 | 902 | 1027 | 1218 | 1424 | 1561 | 1573 | 1477 | 1324 |
| 1204 32 | 1 | -3 | 1171 | 1056 | 993 | 972 | 982 | 1016 | 1071 | 1148 | 1252 | 1386 | 1552 | 1765 | 2027 | 2322 | 2609 | 2910 | 3237 |
| 1171 54 | 1 | -3 | 3399 | 3239 | 2978 | 2794 | 2721 | 2745 | 2829 | 2926 | 2960 | 2868 | 2662 | 2429 | 2215 | 2022 | 1886 | 1817 | 1797 |
| 1138 75 | 1 | -2 | 186 | 204 | 238 | 302 | 414 | 605 | 915 | 1336 | 1652 | 1600 | 1399 | 1278 | 1·167 | 986 | 815 | 709 | 663 |
| 1105 97 | 1 | -1 | 67 | 73 | 84 | 103 | 127 | 154 | 185 | 232 | 307 | 436 | 664 | 965 | 1065 | 854 | 571 | 364 | 241 |
| 1074 15 | 0 | -2 | 2032 | 1791 | 1677 | 1716 | 1956 | 2482 | 3061 | 2939 | 2304 | 1825 | 1530 | 1294 | 1071 | 870 | 709 | 585 | 489 |
| 1057 76 | 0 | -3 | 4144 | 3556 | 3112 | 2791 | 2526 | 2309 | 2116 | 1980 | 1882 | 1812 | 1761 | 1737 | 1707 | 1688 | 1696 | | 1748 |
| 1041 37 1024 97 | 0 | -2 -1 | 180 316 | 186 589 | 191 1011 | 197 984 | 207 629 | 219 377 | 236 234 | 260 155 | 288 110 | 328 85 | 382 71 | 457 65 | 563 68 | 709 71 | 919 60 | | 1894 |
| 1008 58 | | -2 | 396 | 395 | 412 | 463 | 610 | 905 | 1713 | 2477 | 1623 | 836 | 560 | 408 | 341 | 298 | 269 | 48 258 | 43 250 |
| 992 19 | | -3 | 2485 | 2629 | 2679 | 2507 | 2448 | 2415 | 2366 | 2348 | 2362 | 2372 | 2365 | 2322 | 2269 | 2193 | | 2055 | 1992 |
| 974 83 | | -3 | 1869 | 1854 | 1901 | 1927 | 1959 | 1999 | 2012 | 1982 | 1882 | 1748 | 1607 | 1514 | 1429 | 1435 | | 1549 | 1714 |
| 942 05 | | -3 | 2001 | 2479 | 3205 | 3957 | 4088 | 3498 | 2759 | 2198 | 1837 | 1698 | 1708 | 1853 | 1986 | 2020 | | 2506 | 3201 |
| 909 27 | 1 | -2 | 460 | 734 | 1193 | 1579 | 1483 | 1111 | 772 | 524 | 359 | 256 | 198 | 162 | 140 | 126 | 118 | | 109 |
| 876 48 | 1 | -3 | 1057 | 1030 | 1018 | 1073 | 1243 | 1422 | 1381 | 1254 | 1125 | 1010 | 924 | 870 | 843 | 843 | 871 | 924 | 995 |
| | l | -3 | 1103 | 1250 | 1432 | 1669 | 1958 | 2280 | 2553 | 2680 | 2600 | 2371 | 2103 | 1876 | 1695 | 1573 | 1516 | 1532 | 1584 |
| 810 91 | | -3 | 1563 | 1469 | 1384 | 1330 | 1303 | 1294 | 1288 | 1295 | 1311 | 1333 | 1366 | 1406 | 1464 | 1526 | | 1663 | 1786 |
| 778 13 | | -2 | 194 | 211 | 232 | 258 | 289 | 324 | 369 | 426 | 502 | 601 | 745 | 957 | 1284 | 1817 | 2753 | 4499 | 7745 |
| 745 35 | | -1 | 1319 | 2047 | 2737 | 3073 | 2511 | 1528 | 829 | 457 | 275 | 183 | 132 | 102 | 84 | 74 | | | |
| 719 31 | | -2 | 704 | 682 | 672 | 674 | 689 | 718 | 763 | 830 | 926 | 1066 | 1269 | 1574 | 2046 | 2794 | | 5866 | 8554 |
| 702.92 | | -1 | 1125 | 1182 | 1009 | 772 | 561 | 411 | 312 | 248 | 211 | 197 | 202 | 222 | 251 | 290 | 349 | | 648 |
| 686.53 | | -l | 961 | 1309 | 1409 | 1175 | 850 | 595 | 417 | 297 | 218 | 167 | 132 | 108 | 91 | 79 | 70 | | 58 |
| 670 14 653 74 | | -3 -3 | 5446 | 5223 | 5128 | 5070 | 5094 | 5269 | 5606 | 6075 | 6446 | 6333 | 5695 | 4861 | 4124 | 3524 | | 2609 | 2262 |
| 637.35 | | -3 -4 | 1964 6205 | 1729 6016 | 1554 | 1397 | 1269 | 1159 | 1074 | 990 | 913 | 860 | 817 | 770 | 726 | 696 | 655 | 641 | 625 |
| 626 26 | | → | 5866 | 6016 5860 | 5863 5840 | 5830 5821 | 5811 5830 | 5672 5840 | 5719 5886 | 55 7 0 5933 | 5615 6029 | 5698 | 5806 6287 | 5872 | 6706 | 6060 | 7215 | 7722 | 9227 |
| 618 07 | | -3 | 894 | 994 | 1094 | 1272 | 1449 | 1751 | 2053 | 2389 | 2726 | 6125 2595 | 2464 | 6449 2063 | 6705 1661 | 6960 1401 | | 7732 1018 | 8337 896 |
| 609 87 | | 4 | 8421 | 7879 | 7536 | 7194 | 6857 | 6520 | 6183 | 5846 | 5585 | 5325 | 5118 | 4911 | 4737 | 4564 | | 4234 | 4075 |
| | | | | | | | | | | | - 2.00 | - 343 | 2113 | 7714 | | | | 72.57 | 7012 |

Table 3.7 -Continued

| em 1 | λΈ | E | ij | 1 | 2 | ; | 4 | 5 | | 7 | S | ٠, | 1.3 | 11 | 12 | 13 | 14 | : 5 | 10 |
|--------|----|----------|------|-------|------|-------|-------|---------|------|-------|------|------|------|-------|------|-------|------|---------|--------|
| 501 19 | Ò | -4 | 3765 | 3481 | 3227 | 2004 | 2781 | 2581 | 1304 | 2220 | 2076 | 1944 | 1828 | 1222 | 1031 | 1202 | | 1-4-4-1 | 1.4(*) |
| 584 80 | 0 | 4 | 1390 | 1375 | 1349 | 1311 | 1200 | 1228 | 1201 | 1175 | 1155 | 1139 | 1116 | 1089 | 1062 | 1035 | 1011 | 13438 | 12020 |
| 568 41 | 0 | 4 | 300 | 994 | 998 | 1011 | 1.432 | 1071 | 1124 | 11-20 | 1273 | 1363 | 1470 | 1618 | 1812 | 2048 | 2301 | 2400 | 2(43%) |
| 552 02 | 0 | 4 | 2666 | 2721 | 2800 | 2889 | 2048 | 3-3-74) | 2800 | 2846 | 2771 | 2008 | 2561 | 2477 | 2438 | | | | |
| 539.96 | -1 | 4 | 2433 | 2442 | 2452 | 2495 | 2537 | 2050 | 2762 | 2047 | 3131 | 3374 | 3617 | 3838 | 4088 | 4441 | 4124 | 343(17 | 1001 |
| 531 77 | -1 | → | 3394 | 3.107 | 2867 | 2638 | 2481 | 2324 | 2229 | 2129 | 1864 | 1614 | 1334 | 1(55) | 30.2 | 814 | 583 | 1.15 | 223 |
| 510 34 | 3 | 0 | 1,1 | 1.2 | n | ò | 9 | | | | | | | | - | | | • | •• |
| 484 52 | 0 | -2 | o | 7 | 1.5 | 27 | 45 | 67 | -94 | 135 | 189 | 263 | 372 | 513 | '88 | 1217 | 1992 | 1414 | >>-14 |
| 468 13 | 0 | -2 | 7224 | 6686 | 4676 | 29/19 | 1806 | 1173 | 815 | 613 | 490 | 384 | 288 | 219 | 100 | 133 | 102 | 25 | 37 |
| 451 74 | 0 | 4 | 3994 | 2635 | 2454 | 2512 | 2163 | 1305 | 2501 | 2745 | 3113 | inve | 4258 | 4115 | 4970 | 50.40 | | 7.705 | 8486 |
| 435 35 | 0 | -3 | 878 | 1005 | | | | | | | | | | | | • | , , | | |

Footnotes to Tables 3.5, 3.6 and 3.7

- a The column headed cm⁻¹ contains the wavenumber of the first ordinate value in the row. The columns headed XE and YE contain the X-exponent and the Y-exponent, respectively, for the row. The columns headed $0,1,2,\cdots 16$, contain the ordinate values, and the headings give the indices of the ordinate values in the row. In a row which starts with $\widetilde{V}(0)$, the wavenumber corresponding to the ordinate indexed J is $\widetilde{V}(J) = \widetilde{V}(0) \frac{15798.002}{16384} \cdot J \cdot 2^{XE}$. For Tables 3.5 & 3.7, the $k(\widetilde{V})$ and $E_m(\widetilde{V})$ values in that row are the ordinate value shown times 10^{YE} . In Table 3.6, the $n(\widetilde{V})$ values are given directly with the decimal point implicitly after the first digit. Thus the entry indexed 16 in the first row of Tables 3.5, 3.6 and 3.7 shows that at $\widetilde{V} = 4799.95 \frac{15798.002}{16384} \cdot 16 \cdot 2^0 = 4784.52$ cm⁻¹ the ordinate values are: $k = 141 \times 10^{-7} = 1.41 \times 10^{-5}$, n = 1.5033 and $E_m = 3752 \times 10^{-5} = 3.752 \times 10^{-5}$
- b The 4-point spline interpolation program TRECOVER⁹, interpolated the $k(\tilde{\nu})$ and $E_m(\tilde{\nu})$ values in the table to the original wavenumber spacing, 0.482117 cm⁻¹, and yielded the original values accurate to 1% below 4000 cm⁻¹, and to 1-2% between 4000 and 4800 cm⁻¹. The original $n(\tilde{\nu})$ values were similarly recovered accurate to 0.1%.
- c The unit of the E_m values is L mole⁻¹ cm⁻¹. Multiply the values by 1000 to change the unit to cm² mole⁻¹.

3.2.4 - Areas under $k(\widetilde{\nu})$ and $E_m(\widetilde{\nu})$ bands.

Table 3.9 contains the average area under the $k(\tilde{v})$ spectrum for each spectroscopist in the integration ranges given in Table 3.1. The areas were measured

Table 3.8 - Peak heights in the molar absorption coefficient spectrum of liquid chlorobenzene at 25°C. a.b

| $\widetilde{\mathbf{v}}$ (cm ⁻¹) | $E_m(\widetilde{\nu})$ | v (cm ⁻ⁱ) | $E_{\rm m}(\widetilde{v})$ | v (cm ⁻ⁱ) | $E_{\mathfrak{m}}(\widetilde{\nu})$ | \widetilde{v} (cm ⁻¹) | $E_{\rm m}(\widetilde{ u})$ |
|--|------------------------|-----------------------|----------------------------|-----------------------|-------------------------------------|-------------------------------------|-----------------------------|
| 4656.4 | 1.29(2) | 3016 3 | 4.64(5) | 2331.0 | 1.01(2) | 1370.7 | 2.32(1) |
| 4622.4 | 0.429(13) | 2949.8 | 0.559(8) | 2319.8 | 1.00(1) | 1325.2 | 1.86(3) |
| 4595.3 | 0.276(11) | 2919.0 | 0.694(22) | 2271.8 | 0.403(7) | 1298.1 | 2.72(6) |
| 4574.7 | 0.627(18) | 2885.6 | 0.438(11) | 2240.5 | 0.706(13) | 1272.5 | 2.66(7) |
| 4514.6 | 0.189(10) | 2877.0 | 0.449(10) | 2194.0 | 0.227(7) | 1235.2 | 1.91(1) |
| 4252.9 | 0.258(12) | 2849.2 | 0.401(14) | 2178.7 | 0.434(9) | 1210.9 | 1.58(2) |
| 4145.8 | 0.619(17) | 2795.2 | 0.155(9) | 2150.5 | 0.173(5) | 1171.6 | 3.40(6) |
| 4077.4 | 2 58(19) | 2773.0 | 0.463(12) | 2132.5 | 0.103(7) | 1156.5 | 2.96(5) |
| 4066.5 | 2.52(12) | 2755.9 | 0.194(10) | 2089.1 | 0.163(7) | 1122.7 | 16.8(2) |
| 4058.7 | 2.71(21) | 2738.9 | 0.167(10) | 2058.2 | 0.152(9) | 1083.3 | 107.(1) |
| 4000.5 | 0.558(12) | 2714.3 | 0.133(8) | 2045.6 | 0.167(6) | 1068.1 | 31.0(5) |
| 3917.7 | 0.225(7) | 2703.1 | 0.142(8) | 2023.7 | 0.273(10) | 1022.6 | 106.(6) |
| 3899.5 | 0.237(7) | 2663.5 | 0.687(14) | 2002.5 | 0.317(7) | 1001.9 | 24.8(10) |
| 3785.1 | 0.748(10) | 2648.1 | 0.441(12) | 1962.3 | 2.27(2) | 964.0 | 2.01(5) |
| 3772.3 | 0.552(9) | 2616.3 | 0.610(14) | 1942.9 | 4.43(4) | 935.0 | 4.14(22) |
| 3699.4 | 1.09(2) | 2599.8 | 0.598(11) | 1881.5 | 2.51(1) | 902.9 | 16.1(6) |
| 3653.4 | 0.215(11) | 2582.2 | 0.470(15) | 1862.0 | 4.58(4) | 866.3 | 1.43(4) |
| 3636.9 | 0.279(7) | 2557.4 | 0.428(11) | 1787.9 | 3.54(3) | 830.1 | 2.68(12) |
| 3613.6 | 0.290(9) | 2544.3 | 0.472(12) | 1730.6 | 3.33(3) | 812.3 | 1.59(3) |
| 3488.8 | 0.153(7) | 2510.9 | 0.146(8) | 1685.9 | 0.611(12) | 739.7 | 308.(7) |
| 3464.1 | 0.147(5) | 2496.0 | 0.204(9) | 1645.5 | 2.08(2) | 702.2 | 119.(5) |
| 3368.5 | 0.122(7) | 2471.4 | 0.272(9) | 1622.1 | 2.41(4) | 684.8 | 142.(8) |
| 3165.4 | 1.17(2) | 2446.6 | 0.208(7) | 1583.8 | 71.9(11) | 662.1 | 6.47(40) |
| 3083.1 | 11.5(2) | 2427.3 | 0.266(7) | 1566.1 | 12.9(2) | 614.1 | 2.74(15) |
| 3069.5 | 17.0(3) | 2406.4 | 0.451(9) | 1477.6 | 207.(7) | 547.7 | 0.296(17) |
| 3058.5 | 13.6(2) | 2390.3 | 0.600(10) | 1445.4 | 83.3(57) | 533.4 | 0.414(11) |
| 3025.8 | 5.36(4) | 2365.7 | 0.266(14) | 1387.1 | 2.35((1) | 467.8 | 73.5(18) |

a - The unit of $E_{\rm m}(\widetilde{\nu})$ is L mole⁻¹ cm⁻¹. Multiply the values by 1000 to change the unit to cm² mole⁻¹.

above zero ordinate. The 95% confidence limit is given in parentheses except when only one spectrum was available. These spectroscopist average areas were themselves averaged to give an overall unweighted average area and an overall weighted average

b - The number in parentheses is the estimated accuracy in the last digit. The % estimated accuracy is the same as for the corresponding $k(\tilde{\nu})$ value.

Table 3.9 - Spectroscopist average areas under the absorption index bands^a.

| Region (cm ⁻¹) | YA | VB | A | В | С |
|----------------------------|------------|------------|------------|---------|--------|
| 4785.0-4475.0 | 0.0328(26) | 0.0343(22) | | 0.0337 | |
| 4301.9-3850.2 | 0.0981(33) | 0.0978(15) | | | |
| 3850.2-3665.5 | 0.0320(3) | 0.0326(2) | | | |
| 3239.8-2930.3 | 0.5355(19) | 0.5423(21) | 0.5430(25) | | |
| 2930.3-2895.1 | 0.00809(8) | 0.00796(2) | 0.00804(1) | 0.00798 | |
| 2895.1-2811.2 | 0.0144(5) | 0.0142(1) | 0.0145(1) | 0.0140 | |
| 7 11.2-2688.3 | 0.0133(9) | 0.0134(2) | 0.0137(1) | 0.0127 | |
| 2688.3-2517.1 | 0.0407(11) | 0.0412(2) | 0.0419(1) | 0.0402 | |
| 2517.1-2111.7 | 0.0880(21) | 0.0893(2) | 0.0902(5) | 0.0879 | |
| 2000.3-1905.3 | 0.1377(6) | 0.1363(29) | 0.1390(2) | 0.1372 | |
| 1905.3-1835.4 | 0.1335(8) | 0.1324(5) | 0.1342(2) | 0.1327 | |
| 1835.4-1754.4 | 0.1262(6) | 0.1258(3) | 0.1269(2) | 0.1256 | |
| 1754.4-1690.3 | 0.0933(5) | 0.0931(3) | 0.0936(1) | 0.0928 | |
| 1605.4-1545.2 | 1.080(5) | 1.073(5) | | | |
| 1530.2-1410.2 | 2.772(14) | 2.756(40) | | | |
| 1405.9-1345.1 | 0.1013(2) | 0.1014(1) | | 0.1017 | 0.1013 |
| 1345.1-1252.1 | 0.1491(4) | 0.1500(1) | | 0.1487 | 0.1496 |
| 1252.1-1198.5 | 0.0948(1) | 0.0951(1) | | 0.0946 | 0.0950 |
| 1198.5-1141.7 | 0.1959(3) | 0.1967(3) | | 0.1952 | 0.1975 |
| 1140.2-1050.1 | 2.988(15) | 3.002(13) | | | |
| 1045.2- 970.0 | 1.308(5) | 1.266(6) | | | |
| 950.3- 851.9 | 0.6002(4) | 0.6028(32) | 0.5958(5) | 0.6195 | 0.5942 |
| 799.6- 717.1 | | | 8.354(56) | | |
| 719.3- 649.9 | 5.065(29) | 4.997(46) | 5.442(31) | | |
| 630.6- 569.9 | 0.0941(2) | 0.0931(3) | • | 0.0898 | |

a - The unit of area is cm⁻¹. The numbers in parentheses are the 95% confidence limits in the last digit.

area for each band. The weighting was the number of spectra that contributed to the spectroscopist's average. These average areas are listed in Table 3.10. The weighted and unweighted average areas agree well, and the unweighted average areas are taken as the integrated intensities that result from this study.

The agreement between different spectroscopists is shown in Table 3.10 by the

Table 3.10 - Overall average areas under the absorption index bands.

| Region (cm ⁻¹) | k _{max} ⁴ | Weighted Average Area ^b | Unweighted Average Area ^b | Maximum Deviation ^{b,c} | Anchor point Uncertainty ^{b,d} | % Estimated accuracy ° |
|----------------------------|--------------------|--|---|-------------------------------------|--|------------------------|
| 4785.0-4475.0 | 0.000474 | 0.0332 | 0.0336 | ±0.0008 | ±0.0014 | 6.55 |
| 4301.9-3850.2 | 0.00128 | 0.0981 | 0.0980 | ±0.0002 | ±0.0011 | 1.33 |
| 3850.2-3665.5 | 0.000521 | 0.0321 | 0.0323 | ±0.0003 | ±0.00027 | 1.76 |
| 3239.8-2930.3 | 0.00993 | 0.5389 | 0.5403 | ±0.0048 | ±0.00042 | 0.97 |
| 2930.3-2895.1 | 0.000424 | 0.00802 | 0.00802 | ±0.00007 | ±0.00005 | 1.50 |
| 2895.1-2811.2 | 0.000279 | 0.0143 | 0.0143 | ±0.0003 | ±0.00012 | 2.94 |
| 2811.2-2688.3 | 0.000298 | 0.0134 | 0.0133 | ±0.0006 | ±0.00018 | 5.86 |
| 2688.3-2517.1 | 0.000461 | 0.0412 | 0.0410 | ±0.0009 | ±0.00025 | 2.80 |
| 2317.1-2111.7 | 0.000774 | 0.0891 | 0.0889 | ±0.0013 | ±0.0006 | 2.14 |
| 2000.3-1905.3 | 0.00409 | 0.1376 | 0.1375 | ±0.0015 | ±0.00015 | 1.20 |
| 1905.3-1835.4 | 0.00442 | 0.1333 | 0.1332 | ±0.0010 | ±0.00013 | 0.85 |
| 1835.4-1754.4 | 0.00355 | 0.1263 | 0.1261 | ±0.0008 | ±0.00016 | 0.76 |
| 1754.4-1690.3 | 0.00345 | 0.0933 | 0.0932 | ±0.0004 | ±0.00012 | 0.56 |
| 1605.4-1545.2 | 0.0813 | 1.077 | 1.077 | ±0.004 | ±0.00022 | 0.39 |
| 1530.2-1410.2 | 0.254 | 2.765 | 2.764 | ±0.008 | ±0.00047 | 0.31 |
| 1405.9-1345.1 | 0.00303 | 0.1014 | 0.1014 | ±0.0003 | ±0.0002 | 0.49 |
| 1345.1-1252.1 | 0.00375 | 0.1493 | 0.1493 | ±0.0007 | ±0.00027 | 0.65 |
| 1252.1-1198.5 | 0.00277 | 0.0949 | 0.0949 | ±0.00025 | ±0.00018 | 0.45 |
| 1198.5-1141.7 | 0.00520 | 0.1962 | 0.1963 | ±0.0012 | ±0.0011 | 1.37 |
| 1140.2-1050.1 | 0.178 | 2.996 | 2.995 | ±0.007 | ±0.0015 | 0.28 |
| 1045.2- 970.0 | 0.186 | 1.285 | 1.287 | ±0.021 | ±0.00083 | 1.70 |
| 950.3- 851.9 | 0.0319 | 0.5990 | 0.6025 | ±0.0170 | ±0.00011 | 2.84 |
| 799.6- 717.1 | 0.747 | 8.354 | 8.354 | ±0.056 ^f | ±0.014 | 0.84 |
| 719.3- 649.9 | 0.373 | 5.181 | 5.168 | ±0.274 | ±0.0058 | 5.41 |
| 630.6- 569.9 | 0.00796 | 0.0934 | 0.0923 | ±0.0026 | ±0.0010 | 3.90 |

a - Height of the strongest peak in the region.

b - The unit of area is cm⁻¹.

c - Maximum deviation from the unweighted average.

d - The anchor point uncertainty is the integration range multiplied by the average of the uncertainties in $k(\tilde{\nu})$ (see Table 3.1) at the two anchor points used for that range.

e - The estimated accuracy is taken as the sum of the maximum deviation and the anchor point uncertainty.

f - This value is the 95% confidence limit of the measurements of A, the only spectroscopist who measured this band.

maximum deviation of any spectroscopist's average from the unweighted average. The average agreement over all bands is 1.9%. The average agreement for strong, medium and weak bands with $k_{\text{max}} > 0.002$ is 1.3%. The average agreement is 3.1% for very weak bands with $k_{\text{max}} < 0.002$.

The areas under the molar absorption coefficient, $E_{\rm m}(\tilde{\nu})$, spectrum are more widely used in analytical chemistry. They are listed in Table 3.11. In addition to the area above zero ordinate, Table 3.11 contains the area above a linear baseline drawn through the $E_{\rm m}$ values at the integration limits. This latter area is the more accurate, and is the area we recommend as an intensity calibration standard (vide infra).

3.3 - Accuracy of results

3.3.1 - Accuracy of absorption indices and molar absorption coefficients.

We have described previously^{1,2} our reasons for regarding the agreement between different spectroscopists as a good measure of the accuracy of our results. We must also consider the systematic error in our absorption index values due to the uncertainties in the values at the anchor points^{1,2,4}. Thus, the estimated accuracy of our $k(\tilde{v})$ values is taken as the sum of the maximum deviation from the unweighted average (in parentheses in column 3 of Table 3.4) plus the uncertainty due to the anchor points. For the peak heights, the latter is given in column 4 of Table 3.4 and was calculated as the average of the uncertainties in $k(\tilde{v})$ at the anchor points on either side of the band (Table 3.1).

Table 3.11 - Overall average areas under molar absorption coefficient bands of liquid chlorobenzene at 25°C

| Region (cm ⁻¹) | Area ^a | Estimated accuracy of area ^{a,b} | Area above baseline ^a | Estimated accuracy of area above baseline ^{a,c} |
|----------------------------|-------------------|---|----------------------------------|--|
| 4785.0-4475.0 | 86.66 | 5.68 | 71.14 | 1.69 |
| 4301.9-3850.2 | 218.9 | 2.9 | 167.8 | 0.3 |
| 3850.2-3665.5 | 67.41 | 1.19 | 41.62 | 0.39 |
| 3239.9-2930.3 | 923.8 | 9.0 | 841.8 | 7.5 |
| 2930.3-2895.1 | 13.03 | 0.20 | 2.80 | 0.02 |
| 2895.1-2811.2 | 22.74 | 0.67 | 7.84 | 0.16 |
| 2811.2-2688.3 | 20.41 | 1.20 | 7.20 | 0.32 |
| 2688.3-2517.1 | 59.53 | 1.67 | 40.44 | 0.89 |
| 2517.1-2111.7 | 115.0 | 2.5 | 81.98 | 1.20 |
| 2000.3-1905.3 | 149.5 | 1.8 | 110.8 | 1.2 |
| 1905.3-1835.4 | 139.1 | 1.2 | 105.2 | 0.8 |
| 1835.4-1754.4 | 126.0 | 1.0 | 83.36 | 0.53 |
| 1754.4-1690.3 | 89.72 | 0.50 | 54.49 | 0.23 |
| 1605.4-1545.2 | 948.8 | 3.7 | 849.5 | 3.2 |
| 1530.2-1410.2 | 2266 | 7 | 2159 | 6 |
| 1405.9-1345.1 | 77.98 | 0.38 | 45.19 | 0.13 |
| 1345.1-1252.1 | 107.7 | 0.7 | 61.30 | 0.29 |
| 1252.1-1198.5 | 64.79 | 0.29 | 21.68 | 0.06 |
| 1198.5-1141.7 | 127.8 | 1.5 | 48.72 | 0.30 |
| 1140.2-1050.1 | 1817 | 5 | 1650 | 4 |
| 1045.2- 970.0 | 729.7 | 12.4 | 593.8 | 9.7 |
| 950.3- 851.9 | 304.3 | 8.6 | 192.4 | 5.4 |
| 799.6- 717.1 ^d | 3451 | 29 | 3120 | 21 |
| 719.3- 649.9 | 1994 | 108 | 1705 | 90 |
| 630.6- 569.9 | 31.45 | 1.23 | 11.34 | 0.32 |

a - The unit of area is L mole⁻¹ cm⁻². Divide the values by 100 to obtain areas in the unit km mole⁻¹.

b - Calculated from the percent estimated accuracy of the area, which is the same for $E_{\rm m}(\widetilde{\nu})$ and $k(\widetilde{\nu})$ (Table 3.10) bands.

c - The percent estimated accuracy of the area under $E_{\rm m}(\widetilde{\nu})$ above baseline equals the maximum deviation from the unweighted average $k(\widetilde{\nu})$ area (Table 3.10) as a percentage of the unweighted average $k(\widetilde{\nu})$ area (Table 3.10). This percentage, multiplied by the area under $E_{\rm m}(\widetilde{\nu})$ above baseline gives the numbers in this column.

d - These results are from 5 spectra from spectroscopist A.

The percent estimated accuracies of the $k(\tilde{v})$ values at the peak heights are included in Table 3.4. For the 108 peaks, the average accuracy is $\pm 2.9\%$. This is somewhat larger than the 2.4% agreement between spectroscopists, because of the large number of very weak bands for which the uncertainty due to anchor points is comparable to the maximum deviation from the unweighted average. Thus, for the 65 peaks which have $k_{\text{max}} < 0.002$, all except 3 of which are above 2000 cm⁻¹, the average accuracy is $\pm 3.3\%$ while the average agreement is $\pm 2.4\%$. The heights of the remaining 43 peaks with $k_{\text{max}} > 0.002$ are accurate to $\pm 2.4\%$ on average and the agreement between spectroscopists is 2.3%.

There are only 19 peaks for which the estimated accuracy is no better than 5%, and the poorest accuracy is 7.6%. 13 of these 19 peaks are very weak peaks with k_{max} <0.002, and 9 of them have k_{max} <0.0002. Three of the 19 peaks are among the 10 strong peaks which have k_{max} >0.05, and which have an average accuracy of 3.6%. If these 3 poorly known peaks are excluded, the accuracy of the remaining 7 strong peaks is 2.4%.

The $k(\widetilde{\nu})$ values at the peaks can be compared with those of Jones and coworkers⁸ which are the only calibrated measurements available. The calibrated results were obtained from samples held at either 20°C or 28°C⁸, not at the 25°C of this work. The density change for a 5°C temperature change affects the $k(\widetilde{\nu})$ values by about 0.5%, and it is unlikely that a 5°C temperature difference would affect the $k(\widetilde{\nu})$ values by more

than 2% overall. Köser's results¹⁴ indicate that the heights of the peaks at 1477 and 1023 cm⁻¹ decrease by about 2% when the temperature increases from 20 to 25°C, while the peak at 740 cm⁻¹ is almost independent of temperature.

There are 102 peak heights reported in both this paper and ref. 8. Forty two of our values lie outside of the evaluated uncertainty limits of Ref. 8, but only twenty values lie outside of the combined error limits of the two studies. Ten of these are very weak peaks between 2100 and 2800 cm⁻¹. In fact only two peaks disagree substantially, those at 1477 and 614 cm⁻¹. The estimated accuracy of the 1477 cm⁻¹ peak is 3.5%, but the band was really too intense to yield good measurements in the cells available. This may contribute to the disagreement. As noted above, Köser's results¹⁴ argue that sample temperature is not the cause of our 1477 cm⁻¹ peak being 27% lower than the calibrated value⁸.

The $k(\tilde{v})$ values in the baseline are not known as accurately as those in regions of significant absorption, except at the anchor points. From the agreement between different workers and the uncertainties in the anchor point values, we estimate the accuracy of the baseline $k(\tilde{v})$ values to be approximately $\pm 2 \times 10^{-6}$ above 3000 cm⁻¹, $\pm 6 \times 10^{-6}$ between 3000 and 1650 cm⁻¹, and $\pm 6 \times 10^{-5}$ below 1650 cm⁻¹. These values corresponds to about $\pm 5\%$ and $\pm 2.5\%$ of the weakest $k(\tilde{v})$ values above and below 3000 cm⁻¹, respectively.

The percent estimated accuracy of the molar absorption coefficient, $\mathcal{E}_{m}(\tilde{\nu})$,

values is that of the corresponding $k(\tilde{v})$ value. The accuracies are given in Table 3 8 in parentheses as absolute values, not as percentages.

3.3.2. - Accuracy of areas.

To calculate the estimated accuracy of the areas under the $k(\tilde{v})$ bands given in Table 3.10, the average uncertainty in $k(\tilde{v})$ at the anchor points (Table 3.1) on each side of the integration range was multiplied by the integration range and added to the maximum deviation from the unweighted average area (Table 3.10). The estimated accuracy is given in Table 3.10 for each band, as a percentage of the unweighted area. The average accuracy over all 25 spectral regions is an excellent 1.9%. Of the 17 bands, for which $k_{\text{max}} > 0.002$ the average accuracy is 1.3%. The only regions for which the accuracy of the area is poor are the regions of very weak absorption from 4785 to 4475 cm⁻¹, 2811 to 2688 cm⁻¹, and 719 to 650 cm⁻¹. The latter may have been influenced by the very strong neighbouring band at 739.5 cm⁻¹, for which spectra were only available from one spectroscopist.

A comparison was made with the areas under three $k(\tilde{v})$ bands reported by Köser from dispersive transmission measurements¹⁵. For the integration ranges 1550.0 - 1350.0, 1100.0 - 825.0, and 825.0 - 615.0 cm⁻¹, the areas are 2.89, 4.53 and 13.65 cm⁻¹ from our spectrum and 2.97, 1.86 and 13.71 cm⁻¹ from Köser's spectrum. Two regions show excellent agreement. The agreement in the 1100-825 cm⁻¹ region is extremely

poor However, the integration range 1100 - 825 cm⁻¹ includes a large region of weak absorption, and it is probable that baseline accuracy plays a large part in the disagreement.

The percent estimated accuracy of the area under a molar absorption coefficient band equals that of the corresponding absorption index band (Table 3.10). The accuracy is given in Table 3.11 as an absolute value, not as a percentage.

Table 3.11 includes the areas under the molar absorption coefficient spectrum above a linear baseline drawn through the molar absorption coefficients at the integration limits. These areas are free from error due to the anchor points, so their estimated accuracy simply reflects the agreement between spectroscopists that is shown by the maximum deviation from the unweighted average area under the corresponding $k(\tilde{v})$ band (Table 3.10). The percent accuracy of the area above the baseline thus equals the maximum deviation as a percentage of the unweighted mean area (Table 3.10). In Table 3.11 we report the actual estimated accuracy of the area above the baseline, not the percent accuracy.

For selected regions, these $E_{\rm m}(\tilde{\nu})$ areas above baseline and the real and imaginary refractive index and molar absorption coefficient values, will be submitted to Commission I.5 of the International Union of Pure and Applied Chemistry for consideration as secondary intensity standards for infrared spectroscopy of liquids.

3.3.3 - Accuracy of real refractive indices.

The real refractive indices, $n(\widetilde{v})$, were calculated by adding the value $n=1.5043\pm0.0005$ at 8000 cm⁻¹ to the value of $\Delta n(\widetilde{v})$ that is calculated by the Kramers-Kronig transform¹⁰ of the $k(\widetilde{v})$ spectrum. The accuracy of $n(\widetilde{v})$ is, then, the sum of the 0.05% accuracy intrinsic to our Kramers-Kronig transform¹⁰, the 0.03% accuracy of the value 1.5043 at 8000 cm⁻¹ plus the ~0.08% accuracy of $\Delta n(\widetilde{v})$ which results from the accuracy of the $k(\widetilde{v})$ values. The accuracy of the $n(\widetilde{v})$ values is, thus, ~0.2%.

3.4 - Summary

Transmission spectra of liquid chlorobenzene at 25°C that were measured by 5 spectroscopists in 4 different laboratories have been transformed to absorption index spectra. These $k(\widetilde{\nu})$ spectra were compared and averaged to yield spectra between 4800 and 450 cm⁻¹ of the absorption index, the real refractive index and the molar absorption coefficient that are believed to be the most accurate currently available. Absorption index, $k(\widetilde{\nu})$, and molar absorption coefficient, $E_m(\widetilde{\nu})$, values are believed accurate to $\pm 2.4\%$ on average at the peaks of stronger bands, and $\pm 3.3\%$ at weaker bands (k<0.002). The baseline $k(\widetilde{\nu})$ values are known to ~5% above 3000 cm⁻¹ and $\pm 2.5\%$ below 3000 cm⁻¹. The areas under 17 bands or band groups in $k(\widetilde{\nu})$ and $E_m(\widetilde{\nu})$ spectra for which $k_{max} > 0.002$ are known to $\pm 1.3\%$ on average. The real refractive

index, $n(\tilde{v})$, values are believed accurate to 0.2%.

For future reference, the complete numerical data is presented in Compact Table format⁹ to allow the original spectra to be recovered by interpolation without loss of accuracy. The complete final $k(\tilde{v})$, $n(\tilde{v})$, and $E_m(\tilde{v})$ spectra obtained in this work are available on diskette from the authors. To make them available over the longer term, it is planned to place them on an internationally accessible data base.

3.5 - References

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Chapter 4 - Assignments of Vibrations for Chlorobenzene and Toluene

Despite numerous experimental and theoretical studies reported in the literature on toluene¹⁴⁻³⁶ and chlorobenzene^{18,22,23,35-47} there are still some disagreements about the assignments of the fundamental vibrations, especially for toluene. While some of the disagreements are simply due to the use of different notation systems, others are due to different choice of wavenumbers or the assignment of similar wavenumbers to different symmetry modes.

4.1 - Symmetry and vibrational notations for benzene and monosubstituted benzenes

The differences in notation arise from two main numbering schemes for the vibrations of the benzene molecule: Wilson's 13,71 and Herzberg's 72. A third notation system of using letters for the assignments was used by Whiffen 37 for halogen substituted benzenes but was not used afterwards.

Wilson's notation was introduced first and has only been used for benzene and substituted derivatives of benzenes. Herzberg's notation was introduced later and is in general use. In both numbering schemes, vibrations are numbered according to their symmetry species. Within each symmetry species, Herzberg numbers the vibrations in order of decreasing wavenumber. Wilson, on the other hand, numbers them in order of increasing wavenumber with one or two exceptions. An additional difference between the two systems is the order in which the symmetry species for D_{6h} molecules such as

Table 4.1 - Assignments of benzene using Herzberg and Wilson notation schemes.

| Symmetry | Assignment ^{a,b} | Herzberg's | Wilson's |
|----------------------------|--|-------------------|-------------------|
| Species | | Notation | Notation |
| \mathbf{A}_{lg} | CH stretch | \mathbf{v}_1 | V ₂ |
| A_{1g} | CC stretch | \mathbf{v}_2 | $\mathbf{v_i}$ |
| A_{2g} | HCC bend | $\mathbf{v_3}$ | V3 |
| $\mathbf{A}_{2\mathrm{u}}$ | oop HCC bend (75). CCC deformation (25) | V_4 | \mathbf{v}_{11} |
| \mathbf{B}_{lu} | CH stretch | V_5 | \mathbf{v}_{13} |
| $\mathbf{B_{lu}}$ | CCC deformation | $\mathbf{v_6}$ | V ₁₂ |
| B_{2g} | oop HCC bend (65), CCC deformation (35) | V ₇ | V ₅ |
| B_{2g} | oop CCC deformation (85), HCC bend (15) | $\mathbf{v_8}$ | ν., |
| $\mathbf{B}_{2\mathbf{u}}$ | CC stretch (75), HCC bend (25) | V9 | V _{1.4} |
| $\mathbf{B}_{2\mathrm{u}}$ | HCC bend (65), CC stretch (35) | \mathbf{v}_{10} | V ₁₅ |
| E_{lg} | oop HCC bend | v_{11} | \mathbf{v}_{10} |
| E_{1u} | CH stretch | V ₁₂ | V ₂₀ |
| E_{lu} | HCC bend (70), CC stretch (30) | ν ₁₃ | V ₁₉ |
| E_{lu} | CC stretch (65), HCC bend (35) | ν ₁₄ | V ₁₈ |
| E_{2g} | CH stretch | ν ₁₅ | V ₇ |
| E_{2g} | CC stretch (65), HCC bend (25), CCC deformation (20) | ν ₁₆ | V8 |
| E_{2g} | HCC bend (80), CC stretch (20) | v_{17} | Vo |
| E_{2g} | CCC deformation | $\mathbf{v_{18}}$ | v ₆ |
| E_{2u} | oop HCC bend (80), CCC deformation (20) | | |
| | | V ₁₉ | V ₁₇ |
| E_{2u} | oop CCC deformation (85), HCC bend (15) | V ₂₀ | V ₁₆ |

a - Potential energy distribution in percent is given in brackets. Contributions over 90% are omitted.

benzene are listed. Wilson's character table 13 lists the species in the following order: A_{1g} , A_{2g} , B_{1g} , B_{2g} , E_{1g} , E_{2g} , A_{1u} , A_{2u} , B_{1u} , B_{2u} , E_{1u} and E_{2u} . Herzberg 69 lists them in the following order: A_{1g} , A_{1u} , A_{2g} , A_{2u} , B_{1g} , B_{1u} , B_{2g} , B_{2u} , E_{1g} , E_{1u} , E_{2g} and E_{2u} . Table 4.1 lists the vibrational assignments of benzene under both, Wilson's and Herzberg's

b - oop indicate out-of-plane displacements. The remaining vibrations are in-plane.

Table 4.2 - Correlation table between D_{6h} and C_{2v} symmetry species for monosubstituted benzenes

| D _{6h} | C _{2v} a | D_{6h} | C _{2v} ^a |
|-------------------|-------------------|-----------------|------------------------------|
| A_{lg} | \mathbf{A}_1 | В _{2ь} | B ₂ |
| A_{2g} | \mathbf{B}_2 | E_{1g} | $A_2 + B_1$ |
| A_{2u} | $\mathbf{B_1}$ | E_{1u} | $A_1 + B_2$ |
| $\mathbf{B_{lu}}$ | $\mathbf{A_{1}}$ | E_{2g} | $A_1 + B_2$ |
| \mathbf{B}_{2g} | $\mathbf{B_{i}}$ | E_{2u} | $A_2 + B_1$ |

a - The A₁ and B₂ displacements are in the plane of the ring.

notations. The potential energy distributions were obtained from earlier normal coordinate calculations of benzene made by C.D. Keefe⁷³.

Monosubstituted benzenes have C_{2v} symmetry which indicates a lower symmetry than benzene. The lower symmetry has the effect of splitting some of the degenerate vibrations in benzene into non degenerate vibrations in the substituted benzenes. The symmetry species for C_{2v} symmetry are: A₁, A₂, B₁ and B₂. The A₁ and A₂ vibrations are the in-plane and out-of-plane modes, respectively, that are symmetric with respect to the 2-fold rotation axis. The B₁ and B₂ modes are antisymmetric with respect to the 2-fold rotation and they depend on the coordinate system used. In a coordinate system in which the plane of molecule is the YZ plane, the B₁ modes are the out-of-plane modes and the B₂ are the in-plane modes. This notation is used throughout the thesis, as recommended by the Joint Commission for Spectroscopy of IUPAP and IUPAC⁷⁴. These C_{2v} species can be related to the D_{6h} species through the correlation table¹³ given

in Table 4.2.

The difference in symmetry species for D_{6h} and C_{2v} should result in different numbering of related vibrations for the substituted benzenes. Chlorobenzene, a representative of the monosubstituted benzenes, has 12 atoms which leads to 12x3-6=30 vibrational modes. These 30 vibrational modes are distributed among the C_{2v} symmetry species as $11A_1+3A_2+6B_1+10B_2$. If either Wilson's or Herzberg's notation is used, the 11 A_1 vibrations would be numbered 1 through 11, the 3 A_2 vibrations would be numbered 12 through 14, the 6 B_1 vibrations would be numbered 15 through 20 and the 10 B_2 vibrations would be numbered 21 through 30. The vibration numbers would be different from those of the parent vibration of benzene. Thus, under Herzberg's notation, v_{15} (E_{2g}) of benzene, is split under C_{2v} symmetry into an A_1 and a B_2 vibration. The vibration numbers would be between 1 and 11 for the A_1 mode and between 21 and 30 for the B_2 mode.

Clearly, such a rigorous application of Herzberg's or Wilson's notation makes comparison of similar vibrations in different molecules very difficult. Differences of this nature always occur between benzene and monosubstituted benzenes, and even occur among the monosubstituted benzenes themselves as the number given to a vibration within a symmetry species depends on its wavenumber. The wavenumber of a particular vibration, such as the A₁, CX stretching vibration, may change significantly with the change of mass of the substituent X, and this may result in the CX stretching mode being in a different position in the sequence of the A₁ vibrations ordered according to

wavenumber. For example, the A_1 CD stretching vibration of monodeuterobenzene is v_4 while the CCl stretching vibration of chlorobenzene is regarded as v_{11} .

It is clear from the above example that there is an incentive to use a similar numbering scheme for substituted benzenes to that used for benzene. By doing so, the comparison of similar vibrations in different molecules is simplified. The practice of most authors is to use either Wilson's or Herzberg's notation for benzene. Whenever a degenerate vibration in benzene is split, the benzene vibration number is kept and the letter "a" or "b" is added. All "a" vibrations in monosubstituted benzenes are symmetric with respect to 2-fold rotation, and so are either A₁ or A₂ species under C_{2v}, while all "b" vibrations in monosubstituted benzenes are antisymmetric with respect to 2-fold rotation and so are either B₁ or B₂ species. Thus, under Herzberg's notation and the above practice, v_{15} (E_{2g}), the CH stretch in benzene, is split under C_{2v} symmetry into v_{15a} (A₁) and v_{15b} (B₂). Under Wilson's notation, the same vibration in benzene is labeled v_7 (E_{2g}) and is split into $v_{7a}(A_1)$ and $v_{7b}(B_2)$ under C_{2v} symmetry. These notations do not follow either Wilson's or Herzberg's notations for C2v molecules, but are commonly used. Throughout the thesis they are termed either pseudo-Wilson or pseudo-Herzberg notation.

It must be realized that the pseudo-Wilson and pseudo-Herzberg notations only approximately indicate the relation between a vibration of the $C_{2\nu}$ molecule and the parent vibration of benzene. To understand this, consider the A_1 vibrations under $C_{2\nu}$. The reduction of symmetry from D_{6h} to $C_{2\nu}$ causes a mixing of all the benzene vibrations

that correlate with A_1 vibrations under $C_{2\nu}$. In most cases the vibration under $C_{2\nu}$ can not be described by just one vibration under D_{6h} , which is what the pseudo-Wilson or pseudo-Herzberg notations attempt. Nevertheless, these notations have been widely used and the pseudo-Herzberg notation will be used, together with the true Herzberg notation in the thesis.

The disagreements that result from the use of different notations can be easily resolved by a) adopting a single notation scheme⁷⁵, or b) quoting previous authors and pointing out the correlation between the previous authors assignments and the current assignments. Throughout the thesis, the recommendation by Miller⁷⁵, to use the Herzberg notation, is followed. In addition, the pseudo-Herzberg notation is preferred to the pseudo-Wilson and consequently assignments using the pseudo-Wilson notation made by various authors are converted to the pseudo-Herzberg notation in this thesis using Table 4.3.

In order to relate the true Herzberg notation to the pseudo-Herzberg notation, the relation between the form of the $C_{2\nu}$ vibration and that of the D_{6h} vibration must be known. The origin of the relation used must be stated with each table in which the two notations appear.

The disagreements due to notation are not important to the understanding of the vibrations and therefore will be considered to have been resolved. Other differences in assignments are discussed in Section 4.2 for chlorobenzene and in Section 4.4 for toluene.

Table 4.3 - Conversion table for the pseudo notations for the assignments of vibrations of monosubstituted benzenes.

| pseudo-Herzberg | pseudo-Wilson | pseudo-Herzberg | pseudo-Wilson |
|------------------|-------------------|-------------------------|----------------------------|
| v_1 | \mathbf{v}_2 | $ u_{13b}$ | V _{19b} |
| v_2 | \mathbf{v}_{1} | V_{14a} | V_{18a} |
| v_3 | \mathbf{v}_3 | V_{14b} | v_{18b} |
| V ₄ | \mathbf{v}_{11} | v_{15a} | v_{7a} |
| V ₅ | v_{13} | v_{15b} | $\mathbf{v}_{7\mathrm{b}}$ |
| V_6 | v_{12} | V _{16a} | V 8a |
| V ₇ | v_5 | V_{16b} | v_{8b} |
| v_8 | V_4 | V_{17a} | V9a |
| V9 | V_{14} | v_{17b} | V _{9b} |
| v_{10} | v_{15} | V_{18a} | V_{6a} |
| v_{11a} | V_{10a} | $ u_{18b}$ | v_{6b} |
| ν_{11b} | V_{10b} | V_{19a} | ν_{17a} |
| V_{12a} | V _{20a} | $ u_{19b}$ | V _{17b} |
| v_{12b} | V _{20b} | V _{20a} | v_{16a} |
| V _{13a} | V _{19a} | V _{20b} | V _{16b} |

4.2 - Previous assignments of the chlorobenzene vibrations

All of the 30 fundamental vibrations of chlorobenzene are Raman active, and all except the 3 A₂ vibrations are infrared active. In principle, theoretical calculations^{72,76-79} of band contours of asymmetric top molecules allow the determination of the symmetry species of the vibrations which cause bands in the observed gas phase infrared spectrum. These calculations yield three different types of band for each C_{2v} molecule, A, B and C, due to to total rotational transitions associated with the vibrational transitions. The band

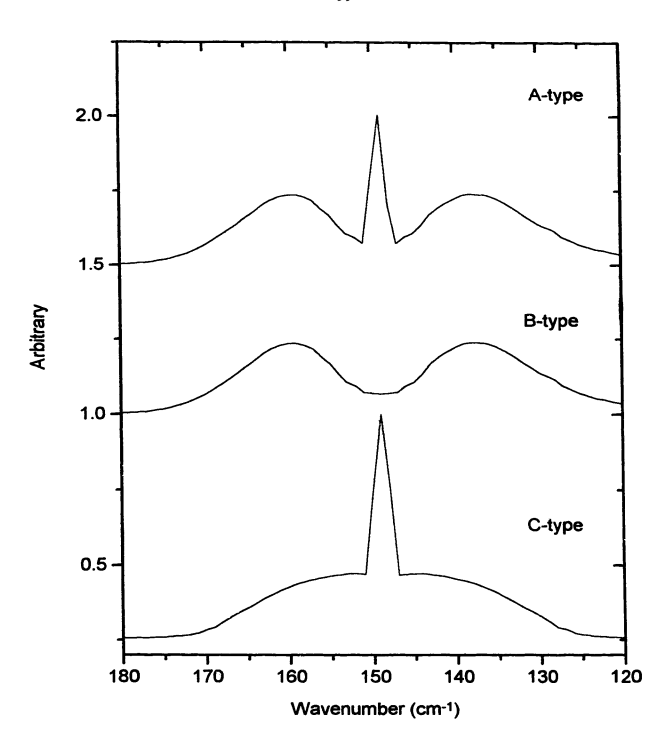


Figure 4.1 - Qualitative band contour shapes for chlorobenzene.

shapes depend on the moments of inertia. For chlorobenzene $^{80-82}$ these are $I_a = 89.1 \pm 0.1$ a.m.u Å^2 , $I_b = 320.5 \pm 0.1$ a.m.u Å^2 and $I_c = 409.7 \pm 0.1$ a.m.u Å^2 and for toluene 83 are $I_a = 89.2$ a.m.u Å^2 , $I_b = 200.8$ a.m.u Å^2 and $I_c = 289.1$ a.m.u Å^2 . According to Ueda and Shimanouchi 79 the moments of inertia yield the three band shapes shown in Fig. 4.1, which are qualitatively the same for chlorobenzene and toluene. In type A, a strong sharp central peak due to Q branch transitions is observed as well as medium side-bands due to P and R branches. In type B, the Q branch is absent and only the unresolved P and R branches are observed. In the C type, a strong Q branch is observed with weak P and R branches. For chlorobenzene and toluene, an A-type band indicates an A_1 vibration, a B-type band indicates a B_2 vibration and a C-type band indicates a B_1 vibration.

In practice, a complete assignment based just on the gas phase infrared spectrum is not possible. First, some wavenumbers of fundamental and/or combination transitions are very similar which results in overlapping and distortion of band shapes. Second, the A_2 vibrations are infrared inactive and are unobserved in the gas phase spectrum. However, in the liquid phase, where adjacent molecules can induce a dipole moment change, the A_2 vibrations can be observed with weak intensity. Further information about the symmetry of vibrations can be obtained from the Raman spectrum. According to Herzberg⁷², the A_1 modes are polarized in the Raman spectrum, with depolarization ratio for linearly polarized incident light $\rho_{\ell} \leq 0.75$, while the other modes are depolarized, with depolarization ratio $\rho_{\ell} = 0.75$. Cross correlation of observed

wavenumbers from infrared and Raman measurements permitted several authors to present a complete assignment for the fundamental vibrations of chlorobenzene. The assignment of the fundamental vibrations made by several authors are summarized in Tables 4.4 to 4.7. The differences between them are not due to notation but rather to the use of different wavenumbers or different symmetry species. It is important though, to clarify the notation used by each author and his data source.

The first major study of chlorobenzene since 1950 was by Whiffen³⁷. For the 30 vibrations he used the letters a through y for the assignment of 25 vibrations and z₁ through z₅ for the remaining 5 vibrations, the CH stretches. Whiffen's letters define a C_{2v} symmetry coordinate through Fig. 1 in his paper. His notation was quoted in later studies but was not adopted. He used wavenumbers from experimental infrared and Raman spectra collected from various studies cited in his paper. He took the plane of the molecule as XZ and, thus, his B₁ species are interchanged with our B₂ species.

In 1960, Schmid, Brandmuller and Nonnenmacher¹⁸ performed normal coordinate calculations with a constant force field and obtained the wavenumbers of benzene and monosubstituted benzenes including chlorobenzene and toluene. In addition to the calculated wavenumbers, experimental wavenumbers obtained from gas and liquid phase infrared and liquid phase Raman measurements were also given, some measured by the authors and some taken from earlier studies. The authors use a true Herzberg notation but their B₁ and B₂ species are interchanged with ours. The authors also gave the pseudo-Wilson notation.

In the early 1970s, Bist and co-workers^{40,42} reported vibrational data for gaseous chlorobenzene from measurements of the infrared spectrum and the UV electronic spectrum. The authors used the pseudo-Wilson notation.

In the early 1970's, two books were published which, *inter alia*, summarized the vibrations of monosubstituted benzenes. Sverdlov, Kovner and Krainov²² selected without explanation fundamental wavenumbers from a variety of authors, several of whom published in unobtainable USSR journals. The true Herzberg notation was used with the B₁ and B₂ vibrations interchanged with ours. The second book, written by Varsanyi²³, is specific to benzene and its derivatives. Varsanyi again selected without explanation fundamental frequencies from a variety of authors including Sverdlov, Kovner and Krainov²². The pseudo-Wilson notation was used.

In 1992-3, Pulay and co-worker. ⁴⁷ reported the fundamental vibrational wavenumbers obtained from scaled *ab initio* calculations. The experimental wavenumbers which guided the scaling were taken from Bist *et al.* ^{40,42} and Whiffen³⁷. The notation used is a true Herzberg notation.

Several other authors have reported vibrational studies on chlorobenzene.

However, the studies either gave only partial wavenumbers and assignments^{38,41,44-46} or, as is the case for some theoretical studies^{39,43}, used wavenumbers and assignments from previous studies without change. These studies will be cited when deemed necessary.

In the following sub-sections, the assignment of the fundamentals by these authors is reviewed. The differences will be pointed out, however, a critical evaluation

of the authors' choices is left to Section 4.3 where an assignment is proposed, based on all available data, including data measured for this thesis.

4.2.1 - The 11 A₁ vibrations of chlorobenzene

Previously reported wavenumbers and assignments for the 11 A₁ vibrations of chlorobenzene are given in Table 4.4. The first column gives the true Herzberg notation, and, for convenience, the pseudo-Herzberg notation is given in the second column. In Table 4.4 and in the later Tables 4.5 to 4.7, the pseudo notation was taken from Schmid *et al.*¹⁸, who determined the relation between the C_{2v} vibrations and those of benzene in the early 1960's. Schmid's pseudo-Wilson notation was converted to the pseudo-Herzberg notation in column 2.

The remaining columns in Table 4.4 contain each author's wavenumbers and assignments. Bist's^{40,42} and Varsanyi's²³ pseudo-Wilson notations have been converted to pseudo-Herzberg notation in Table 4.4. No assignment is given for Pulay⁴⁷, Schmid¹⁸ and Sverdlov²². Pulay and Sverdlov used only the true Herzberg notation which is shown in column 1 of Table 4.4. The experimental wavenumbers listed in the table are those from infrared measurement in the liquid phase except for those of Bist. When the Raman wavenumber is used, it is denoted by the letter R next to the wavenumber.

The agreement on the fundamental wavenumbers is generally very good among the experimental studies. In several cases, Bist's wavenumber is high by up to 8 cm⁻¹, but this is likely due to differences in the wavenumber calibration. There are however

Table 4.4 - Previous assignments of the A₁ vibrations of chlorobenzene

| | | | Theoretical | | | | | | | |
|------------------------|-------------------|------------------|-----------------------|------------------|---------------------------|--------------------------------|-----------------------------------|--------------------|-----------------------------------|----------------------------------|
| Herzberg | Herzberg notation | | Whiffen a [Ref 37] | | st ^{b,c} [40] | Sverdlov ^d [Ref 22] | Varsanyi ^b [Ref 23] | | Schmid ^{d,e} [Ref 18] | Pulay ^{d.e} [Ref 47] |
| true | pseudo | cm ⁻¹ | | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ |
| v_l | v_{12a} | 3071 | Z_1 | 3082 | v_{12a} | 3086 | 3086 | \mathbf{v}_{i} . | 3075 | 3087 |
| v_2 | ν_{l} | 3050 | Z_2 | 3054 | v_{l} | 3050R | 3071 | v_l | 3069 | 3071 |
| V ₃ | V ₅ | 3029 | Z_3 | 3031 | v_5 | 3029 | 3029 | V ₅ | 3053 | 3047 |
| V ₄ | V _{16a} | 1580 | k | 1586 | v_{16a} | 1580 | 1580 | V _{16a} | 1632 | 1590 |
| V ₅ | v_{13a} | 1477 | m | 1482 | v_{13a} | 1475 | 1477 | v_{13a} | 1513 | 1475 |
| v_6 | v_{15a} | 1174 | а | 1153 | v_{17a} | 1175 | 1174 | V _{17a} | 1225 | 1179 |
| V ₇ | v_{17a} | 1085 | q | 1093 | V _{15a} | 1085 | 1085 | v_2 | 1140 | 1097 |
| Vg | v_{14a} | 1026 | b | 1026 | v_{14a} | 1027 | 1026 | Vija | 1023 | 1017 |
| V9 | V ₆ | 1003 | p | 1004 | v_2 | 1002 | 1003 | v_6 | 1000 | 989 |
| v_{10} | v_{lg_a} | 701 | r | 706 | v ₆ | 702 | 701 | v_{18a} | 708 | 698 |
| v ₁₁ | V ₂ | 415 | t | 417 | V _{18a} | 418 | 420 | V _{15a} | 389 | 413 |

a - Author used his own notation scheme.

three significant wavenumber differences. Bist $et\ al.^{40}$ reported v_6 at 1153 cm⁻¹ while Whiffen³⁷, Sverdlov $et\ al.^{22}$ and Varsanyi²³, all put it at 1174±1 cm⁻¹. Other wavenumber differences occur for two CH stretching vibrations, v_1 and v_2 . v_2 is given as

b - Assignments in pseudo-Wilson notation were converted to pseudo-Herzberg notation using Table 4.3.

c - Wavenumbers of the infrared spectrum of the gas.

d - Authors used the true Herzberg notation in column 1.

e - Calculated wavenumbers.

3052±2 cm⁻¹ by Whiffen³⁷, Bist⁴⁰ and Sverdlov²², while Varsanyi²³ places it about 20 cm⁻¹ higher at 3071 cm⁻¹. v₁ is given as 3071 cm⁻¹ by Whiffen³⁷, while Bist⁴⁰, Sverdlov²² and Varsanyi²³ placed it at 3084±2 cm⁻¹.

There are also deviations from the pseudo-Herzberg notation. Varsanyi²³ and Bist⁴⁰ disagree with each other and with Schmid¹⁸ about the pseudo-Herzberg description of v_6 , v_7 , v_9 , v_{10} and v_{11} . v_6 is assigned as v_{17a} , HCC deformation and CC stretch (Table 4.1), by both Varsanyi and Bist, while Schmid¹⁸ assigns it as v_{15a} , the E_{2g} CH stretch which, near 1160 cm⁻¹, means it is related to the CX stretching mode. Miller⁷⁵ has noted that v_9 derives from v_6 of benzene not v_2 . However, in the other cases, the authors give little evidence to help evaluate the other assignments which, in any case, probably arise from the limitations of the pseudo-Herzberg description.

The computed frequencies by Schmid et al. 18 and Pulay et al. 47 agree reasonably well with the experimental wavenumbers. However both are to some extent fitted to their chosen experimental data, so can not really settle disagreements.

4.2.2 - The 3 A₂ vibrations of chlorobenzene

The wavenumbers and assignments of the A₂ vibrations are given in Table 4.5.

The table is arranged in a similar manner to Table 4.4. All wavenumbers and assignments from the different experimental studies are in excellent agreement, and the computed wavenumbers are in satisfactory agreement with experiment.

Table 4.5 - Previous assignments of the A₂ vibrations of chlorobenzene

| | | | | Theoretical | | | | | | |
|-----------------|------------------|--------------------|---|------------------|------------------|--------------------------------|-----------------------------------|------------------|--------------------------------|----------------------------------|
| Herzberg | g notation | Whiffen a [Ref 37] | | Bist [Ref | | Sverdlov ^d [Ref 22] | Varsanyi ^b [Ref 23] | | Schmid ^{d,e} [Ref 18] | Pulay ^{d,e} [Ref 47] |
| true | pseudo | cm ⁻¹ | | cm ⁻¹ | | cın ⁻¹ | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ |
| v_{12} | v_{14a} | 965 | h | 961 | v_{14a} | °65 | 965 | v_{14a} | 979 | 956 |
| v_{13} | v_{11a} | 830 | g | 832 | Vila | 830 | 830 | ν_{lla} | 848 | 821 |
| V ₁₄ | V _{20a} | 400 | w | 400R | ν _{20a} | 400R | 400 | v _{20a} | 405 | 404 |

a - Author used his own notation scheme.

Table 4.6 - Previous assignments of the B₁ vibrations of chlorobenzene*

| | | | Theoretical | | | | | | | |
|-------------------|-----------------------|------------------|-------------|------------------|------------------|--------------------------------|-----------------------------------|------------------|-----------------------------------|----------------------------------|
| Herzberg notation | | Whiff [Ref : | - | | | Sverdlov ^d [Ref 22] | Varsanyi ^b [Ref 23] | | Schmid ^{d,e} [Ref 18] | Pulay ^{d,e} [Ref 47] |
| true | pseudo | cm ⁻¹ | - | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | · | cm ⁻¹ | cm ⁻¹ |
| v_{15} | V 7 | 985 | j | 981 | V 7 | 984 | 985 | V ₇ | 984 | 981 |
| V ₁₆ | V _{19b} | 902 | i | 902 | V _{19b} | 903 | 902 | V _{19b} | 919 | 899 |
| v_{17} | v_{11b} | 740 | f | 740 | v_{11b} | 740 | 740 | V ₄ | 752 | 740 |
| v_{18} | v ₈ | 682 | v | 685 | ν8 | 682 | 682 | ν | 700 | 679 |
| V ₁₉ | V ₄ | 467 | y | 467 | V _{20b} | 469 | 467 | V _{20b} | 490 | 466 |
| V ₂₀ | V _{20b} | 196R | x | 19 8 R | V ₄ | 196R | 196 | v_{l1b} | 224 | 187 |

^{* -} See footnotes to Table 4.5

b - Assignments in pseudo-Wilson notation were converted to pseudo-Herzberg notation using Table 4.3.

c - Wavenumbers of the infrared spectrum of the gas.

d - Authors used the true Herzberg notation in column 1.

e - Calculated wavenumbers.

4.2.3 - The 6 B₁ vibrations of chlorobenzene

The wavenumbers and assignments of the B₁ vibrations are given in Table 4.6 which is arranged in a similar manner to Table 4.4. All wavenumbers are in excellent agreement from the different experimental studies and the computed wavenumbers are in satisfactory agreement with the experimental values.

Differences in assignment or description of the vibration by the pseudo-Herzberg notation exist for v_{17} , v_{19} and v_{20} with different authors assigning different vibrations to v_4 , v_{11b} and v_{20b} . However, these pseudo-Herzberg description all indicate a mixture of out-of-plane HCC and CCC deformations, so the differences are of detail, not principle.

4.2.4 - The 10 B₂ vibrations of chlorobenzene

The wavenumber and assignments of the B₂ vibrations are given in Table 4.7 which is arranged in a similar manner to Table 4.4.

The only significant wavenumber disagreement is for v_{21} , v_{22} and v_{23} . For the CH stretching modes, there is no agreement within 25 cm⁻¹ about the wavenumber of v_{21} , and Bist⁴⁰ preferred 3067 cm⁻¹ to 3052 cm⁻¹ for v_{22} . For v_{23} , Bist's wavenumber is 18 cm⁻¹ higher than that of Whiffen³⁷, Sverdlov²² and Varsanyi²³. This difference is at the high end of what can be expected from calibration errors.

The only disagreement in notation is Varsanyi's assignment of v_{28} and v_{30} to v_{14b} and v_{10} instead of the reverse. v_{14b} and v_{10} are, however, both mixtures of HCC in-plane

Table 4.7 - Previous assignments of the B2 vibrations of chlorobenzene

| | | <u>Experimental</u> | | | | | | | Theoretical | |
|------------------------|-----------------------|---------------------|----|---------------------|------------------|-----------------------|-----------------------|-------------------|-----------------------|----------------------|
| | | Whiffen * | | Bist ^{b,c} | | Sverdlov ^d | Varsanyi ^b | | Schmid ^{d.e} | Pulay ^{d,e} |
| Herzberg notation | | [Ref 37] | | [Ref 40] | | [Ref 22] | [Ref 23] | | [Ref 18] | [Ref 47] |
| true | pseudo | cm ⁻¹ | | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ |
| \mathbf{v}_{21} | v_{12b} | 3071 | Z4 | 3096 | V _{12b} | 3086 | 3050 | v_{12b} | 3079 | 3082 |
| V ₂₂ | v_{15b} | 3052R | Zs | 3067 | v_{15b} | 3052R | 3052 | V _{15b} | 3044 | 3058 |
| V ₂₃ | V _{16b} | 1580 | l | 1598 | v _{16b} | 1580 | 1570 | $ u_{16b}$ | 1592 | 1596 |
| V ₂₄ | V_{13b} | 1445 | u | 1448 | V _{13b} | 1445 | 1445 | V13b | 1445 | 1438 |
| v ₂₅ | V9 | 1326 | o | 1326 | V9 | 1326 | 1326 | V 9 | 1324 | 1320 |
| V ₂₆ | v ₃ | 1271 | е | 1271 | v ₃ | 1271 | 1271 | $\mathbf{v_3}$ | 1298 | 1272 |
| v ₂₇ | V _{17b} | 1157 | c | | v_{17b} | 1157R | 1157 | v_{17b} | 1141 | 1162 |
| V ₂₈ | \mathbf{v}_{10} | 1068 | d | 1068 | v_{10} | 1068 | 1068 | V _{14b} | 1064 | 1069 |
| V ₂₉ | V _{18b} | 616 | s | 615 | v_{18b} | 617 | 616 | V _{18b} | 629 | 618 |
| V ₃₀ | V _{14b} | 297 | u | 295 | V _{14b} | 296 | 297 | \mathbf{v}_{10} | 321 | 291 |

a - Author used his own notation scheme.

bend and CC stretching, so the difference is again one of detail not principle.

b - Assignments in pseudo-Wilson notation were converted to pseudo-Herzberg notation using Table 4.3.

c - Wavenumbers of the infrared spectrum of the gas.

d - Authors use the true Herzberg notation in column 1.

e - Calculated wavenumbers.

4.3 - Assignments of the chlorobenzene vibrations

There is good but not complete agreement between the assignments of the fundamentals of chlorobenzene in the literature. Unfortunately very little experimental evidence has been cited in support of any of these assignments, so it is impossible to evaluate the disagreements. For this reason, it was decided to collect some evidence in this work to help to evaluate the assignments. Three types of evidence were collected in addition to the quantitative infrared spectrum of the liquid reported in Chapter 3.

First, the infrared spectrum of the gas was recorded in order to observe the contours of the different bands, and, hence, deduce the symmetry of the vibration that causes the band. As discussed previously, for chlorobenzene an A-type indicates an A₁ vibration, B-type indicates a B₂ vibration and a C-type indicates a B₁ vibration. A₂ vibrations are unobserved in the gas phase spectrum.

Second, the Raman spectrum of the liquid was recorded under parallel (I_1) and perpendicular (I_1) polarizations, with linearly polarized incident light. Frequently bands that are weak in the infrared are strong in the Raman so additional vibrations can sometimes be observed in the Raman. Further, values of the depolarization ratio, $\rho_{\ell} = \frac{I_1}{I_1}$ that are less than 0.75 indicate that the band arises from an A_1 vibration under $C_{2\nu}$. For A_2 , B_1 and B_2 vibrations under $C_{2\nu}$, $\rho_{\ell} = 0.75$.

Third, the selection of fundamental vibrations was guided by a very simple normal coordinate calculation that showed the changes in fundamental wavenumbers that

must occur simply from the change in mass of the substituent atom. Thus the latest force field and geometry for liquid benzene⁷³ were used without change, and only the mass of the substituent atom was changed. This was done for C_6H_3D , $C_6H_3CH_3$ and C_6H_3Cl , where a single atom of mass 15 was used to substitute the CH_3 group in toluene. These calculations indicated the approximate wavenumber of the fundamentals in the monosubstituted benzenes, with C_6H_3Cl being of interest in this section. Further, when the symmetry coordinates under D_{6h} that were written for benzene, were used in the calculations for the molecules of C_{2v} symmetry, the relation between the C_{2v} vibration and the D_{6h} symmetry coordinates was directly given by the eigenvectors. Thus, the pseudo-Herzberg notation for the vibration was made by choosing the D_{6h} symmetry coordinate that contributed the most to the C_{2v} vibration.

The experimental data and the results of $12 \, \omega$ calculations are presented below so as to form a basis for the evaluation of the assignments. The liquid phase real and imaginary refractive index spectra, obtained for the secondary intensity standards project and reported in Chapter 3, were converted into the real and imaginary molar polarizability spectra by use of Equations 1.3.33 and 1.3.28. The wavenumbers assigned in this section are the wavenumbers of peaks in the imaginary molar polarizability, α_m'' , spectrum. The experimental spectra of chlorobenzene are shown in Figures 4.2 to 4.8. First, the entire infrared α_m'' spectrum of the liquid is given in Fig. 4.2. Figures 4.3 to 4.8 show the same infrared spectrum, the infrared bands of the gas, and the parallel and perpendicular Raman spectra for smaller wavenumber ranges, and thus provide a better view of the various features of the spectra. In addition, Table 4.8

gives information from all the three spectra for every feature observed in the regions where the fundamentals are expected, i.e. $3150 - 2950 \text{ cm}^{-1}$ and $1700 - 175 \text{ cm}^{-1}$. The first three columns give the wavenumber, a brief description and the peak height of the band in the infrared α_m'' spectrum of the liquid. When the feature is a shoulder its position and height are estimated and are indicated by a \sim . In columns 4 and 5 the infrared gas phase band's wavenumber and shape are given. The labels? and ?? are used to indicate uncertainty and great uncertainty, respectively, in the determination of the band shape. The wavenumber of a B-type vibration was taken at the minimum of the R and P branches while the peak of the Q branch is given for the A-type and C-type bands. In columns 6 and 7 the Raman wavenumber and the depolarization ratio are given for the liquid. When the feature is too weak or noisy to obtain a reliable ratio, the labels p and dp are used to indicate a polarized or depolarized band.

The computed wavenumbers, eigenvectors and potential energy distribution

(PED) for chlorobenzene are given in Table 4.9. Only the main contributions are given in the eigenvector and PED columns. It is clear from this table that the fundamental vibrations are a mixture of several symmetry coordinates. Hence, attempts in the past to describe them as being derived from a single vibration in benzene, which is what the pseudo notations do, were bound to fail and create disagreements in notations.

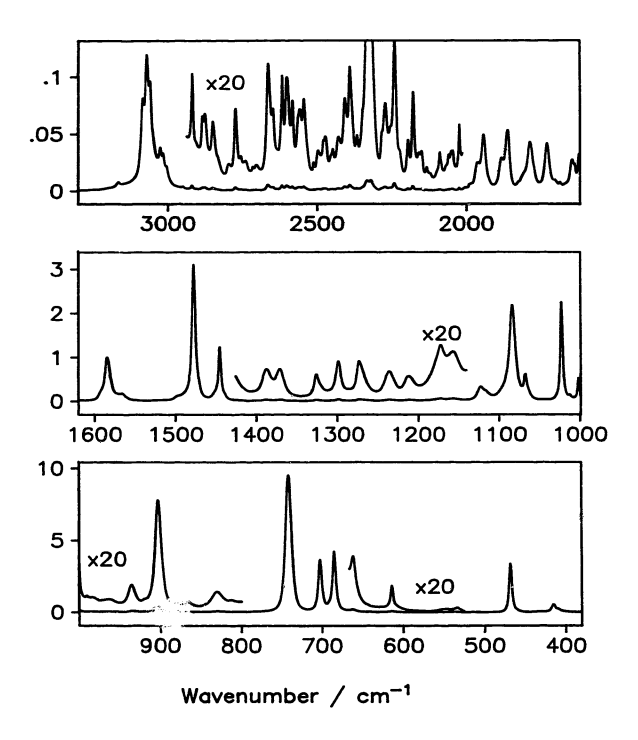


Figure 4.2 - Imaginary molar polarizability, α_m^n , spectrum of liquid chlorobenzene. Units are cm³ mol⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by 20 for the upper curve.

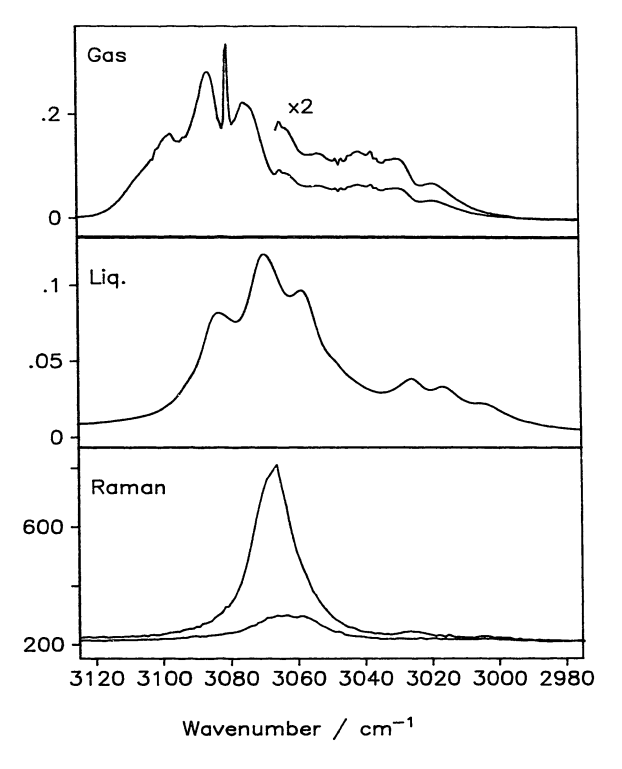


Figure 4.3 - Infrared spectra of gas and liquid chlorobenzene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 3125 and 2975 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

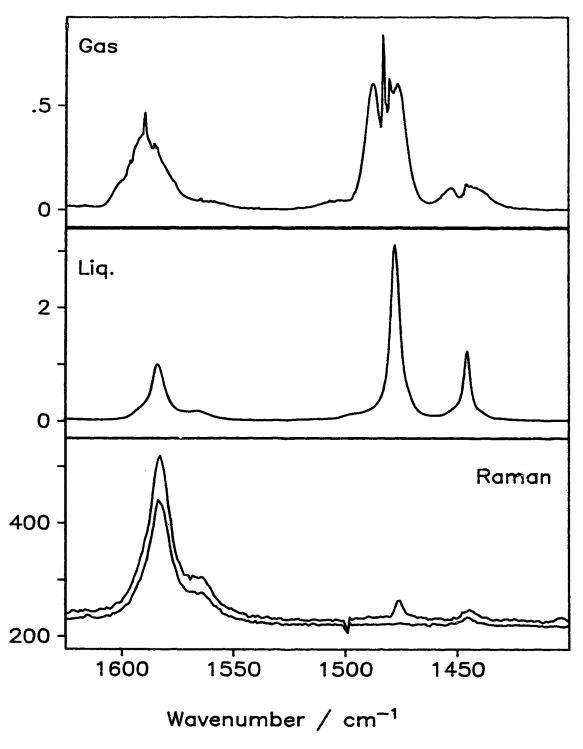


Figure 4.4 - Infrared spectra of gas and liquid chlorobenzene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 1625 and 1400 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

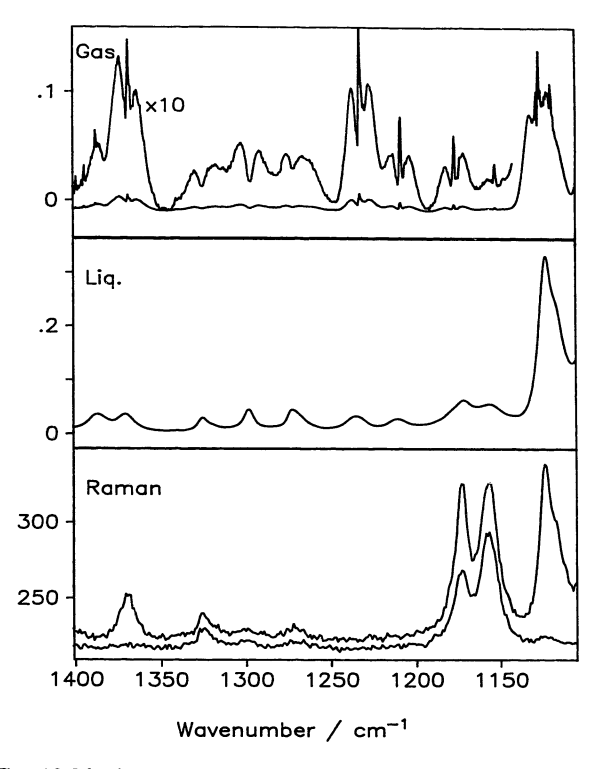


Figure 4.5 - Infrared spectra of gas and liquid chlorobenzene and Raman parallel (upper)and perpendicular (lower) spectra of the liquid between 1400 and 1105 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

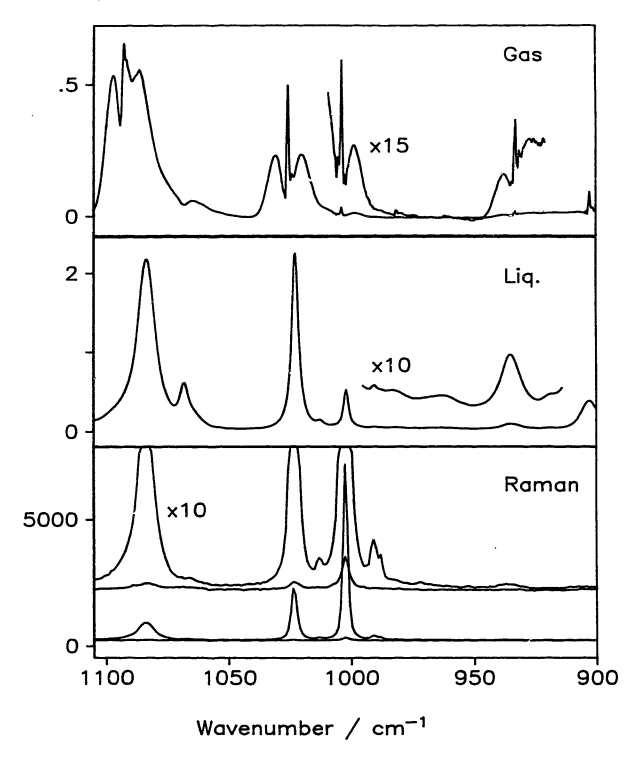


Figure 4.6 - Infrared spectra of gas and liquid chlorobenzene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 1105 and 900 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

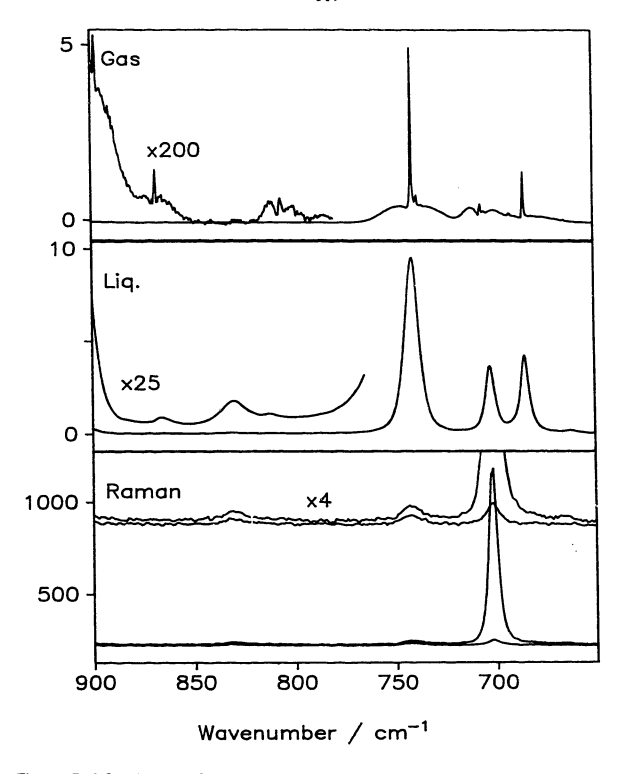


Figure 4.7 - Infrared spectra of gas and liquid chlorobenzene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 900 and 650 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

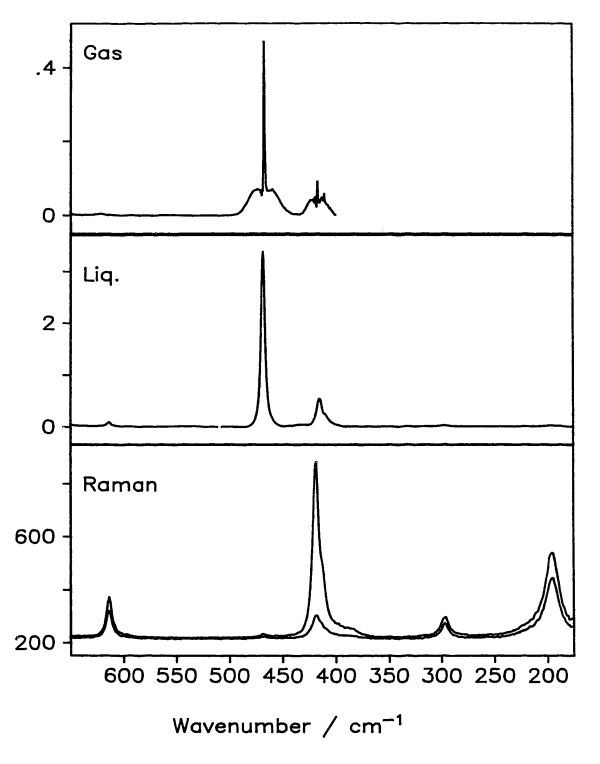


Figure 4.8 - Infrared spectra of gas and liquid chlorobenzene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 650 and 175 cm⁻¹.

Table 4.8 - Features in the experimental infrared spectra of gas and liquid chlorobenzene and Raman spectra of liquid chlorobenzene

| In | frared liquio | i° | Infrar | ed gas d | Raman liq | uid ^b | |
|------------------|---------------|--------------------------------|------------------|------------|------------------|----------------------------|-------------|
| cm ⁻¹ | Desc.b | $lpha_{ m m}^{\prime\prime}$ c | cm ⁻¹ | Contour | cm ⁻¹ | $\rho_{\ell}^{\ \epsilon}$ | comments |
| 3083.1 | m | 0.0817 | ~3095 | B? | | | |
| 3069.5 | S | 0.121 | 3080 | Α | 3066 s | 0.20 | |
| 3058.5 | m br | 0.0968 | 3065 | A?? | | | |
| ~3050 | m sh | 0.0545 | ~3055 | B?? | | | |
| 3025.8 | w | 0.0385 | ~3037 | A?? | 3027 vw | 0.07 | |
| 3016.3 | w | 0.0334 | ~3022 | B ? | | | |
| ~3004 | w sh | ~0.02 | | | 3005 vw | 0.41 | |
| ~1703 | vw sh | ~0.01 | | | | | |
| 1685.9 | vw | 0.008 | | | | | |
| 1645.5 | w | 0.0278 | | | | | |
| 1636.9 | w p/sh | 0.0240 | | | | | |
| 1622.1 | w br | 0.0327 | | | | | |
| ~1591 | s sh | ~0.25 | 1589 | A? | | | |
| 1583.9 | vs | 1.00 | ~1586 | B?? | 1583 m | 0.76 | |
| 1566.1 | s br sh | 0.178 | | | ~1565 w sh | dp | |
| ~1496 | s sh | 0.13 | | | | | |
| 1477.8 | vs | 3.11 | 1483 | Α | 1476 vw | 0.05 | |
| 1445.5 | vs | 1.24 | ~1449 | В | 1445 vw | 0.78 | |
| 1387.1 | w | 0.0367 | | | | | |
| 1370.7 | w | 0.0367 | 1369 | Α | 1370 vw | 0.05 | |
| 1325.2 | w | 0.0306 | 1325 | B ? | 1326 vw | 0.79 | |
| 1298.1 | m | 0.0456 | 1297 | В | 1300 | dp | |
| 1272.5 | m | 00457 | ~1271 | B ? | 1272 vw | dр | |
| 1235.2 | w | 0.0339 | 1232 | Α | | | |
| 1210.9 | w | 0.0287 | 1208 | Α | | | |
| 1171.6 | m | 0.0639 | 1176 | A | 1173 w | 0.42 | |
| 1156.4 | m | 0.0565 | | | 1157 w | 0.79 | |
| 1122.7 | S | 0.332 | 1126 | A ? | 1124 w | 0.03 | |
| | | | 1128 | A ? | | | |
| ~1116 | s sh | 0.244 | ~1119 | C?? | | | |
| 1083.6 | vs | 2.18 | 1092 | Α | 1084 s | 0.04 | |
| 1068.1 | S | 0.617 | ~1068 | B? | ~1069 vw | dp | |
| 1022.8 | vs | 2.27 | 1025 | Α | ~1023 vs | 0.03 | |
| 1012.6 | S | 0.151 | | | 1013 w | 0.00/3 | |
| 1001.9 | S | 0.532 | 1004 | A ? | 1002 vs | 0.04 | |
| 990.5 | m | 0.0585 | | | 991 | р | |

Table 4.8 - Continued

| Į, | nfrared liquid | a | Infra | red gas d | Raman li | quid ^b | |
|------------------|----------------|----------|------------------|---------------|------------------|----------------------------|------------|
| cm ⁻¹ | Desc.b | α,,, c | cm ⁻¹ | Contour | cm ⁻¹ | ρ_{ℓ}^{e} | comments |
| 983.2 | m br sh/p | 0.0521 | 982 | ?? | 972? vw | p | |
| 963.7 | m br | 0.0454 | | | | | |
| 935.0 | m | 0.0967 | 933 | Α | 937 vw | 0.17 | |
| ~919 | m sh/p | ~0.0469 | | | | | |
| 903.0 | S | 0.389 | 903 | C? A? | ~904? vw | dр | |
| 866.3 | w | 0.0363 | 868 | A? C ? | | | |
| 830.0 | m | 0.0717 | | | 831 vw | 0.71 | |
| 812.2 | m | 0.0437 | 806 | A ? | | | |
| | | | 811 | A?? | | | |
| 741.2 | vs | 9.52 | 741 | ·C | 743 vw | 0.78 | |
| 702.5 | vs | 3.67 | 706 | A? C? | 702 vs | 0.04 | |
| 685.2 | vs | 4.25 | 685 | C | | | |
| | | | 692 | | | | |
| 662.1 | s | 0.197 | 665 | C?? | | | |
| | | | 666 | B?? | | | |
| 614.1 | m | 0.0933 | 615 | В | 614 w | 0.73 | |
| ~552 | w sh | ~0.0102 | | | | | |
| 547.7 | w | 0.0115 | | | | | |
| 533.4 | w | 0.0165 | | | | | |
| 468.1 | vs | 3.39 | 467 | С | 468 vw | dp | |
| ~434 | m br sh/p | 0.049 | | | | | |
| 415.0 | S | 0.548 | 417 | A | 418 s | 0.15 | |
| ~400 | m sh | ~0.0549 | | | | | literature |
| 297.5 | 7.5 w 0.0292 | | | | 296 vw 0.73 | | |
| 195.9 | w | 0.0326 | | | 195 m | 0.72 | |

a - Features are from the α_m'' spectrum.

b - Abbreviations used: v- very, w- weak, m- medium, s- strong, sh- shoulder, br- broad, sh/p- shoulder or a peak.

c - Peak heights in the α_m'' spectrum in cm³ mol⁻¹.

d - Peak positions in the absorbance spectrum of the gas. Wavenumbers of B-type bands are measured at the minimum between the R and P branches. ? and ?? indicate degrees of uncertainty in the determination of the band shape.

e - Depolarization ratio are $\rho_{\ell} = \frac{I_{\perp}}{I_{\parallel}}$. When the feature is too weak or noisy to obtain a reliable ratio, p or dp are used to indicate polarized and depolarized, respectively.

Table 4.9 - Computed wavenumbers, eigenvectors and potential energy distributions for chlorobenzene.

| v_1 | cm ⁻¹ | Eigen | vectors4 | | PED ^b |
|---------------------------|------------------|---|------------------------|----------------------|-----------------------------|
| $\mathbf{v}_{\mathbf{l}}$ | 3177 | S ₅ -0.84 S ₁ -0.59 | | | СН |
| \mathbf{v}_2 | 3171 | S ₁ -0.73 S _{15a} -0.55 | S ₅ 0.42 | | СН |
| v_3 | 3158 | S _{12a} -0.80 S _{15a} 0.62 | | | СН |
| v_4 | 1595 | S_{18a} 0.85 S_{16a} 0.43 | $S_{1^{-}a} = 0.39$ | | CC (75), HCC (25) |
| \mathbf{v}_{5} | 1541 | $S_{14a} - 1.17$ $S_{13a} = 0.27$ | | | HCC (60), CC (40) |
| v_6 | 1270 | $S_{18a} 0.69 S_{14a} -0.59$ | | | HCC (55), CC (25), CCl (20) |
| V 7 | 1148 | $S_{18a} = 0.79 \qquad S_{14a} = 0.30$ | | | HCC (55), CCI (30), CC (15) |
| v_8 | 1029 | S_{14a} 0.49 S_6 -0.31 | $S_{13a} = 0.20$ | | CC (55), HCC (22), CCC (22) |
| V9 | 1001 | S ₆ -0.47 S ₂ -0.20 | | | CCC (50), CC (50) |
| v_{10} | 773 | S_{17a} -0.31 S_6 0.18 | | | CCC (50), CC (30), CCl (20) |
| \mathbf{v}_{11} | 398 | S_{17a} 0.20 S_{18a} -0.10 | | | CCC (60), CCl (40) |
| v_{12} | 971 | S_{19a} 1.25 S_{20a} 0.44 | | | оор Н |
| v_{13} | 852 | S _{11a} 1.03 | | | оор Н |
| v_{14} | 403 | $S_{20a} = 0.70 \qquad S_{19a} = 0.26$ | | | oop C |
| v_{15} | 987 | S ₇ 1.13 S _{19b} 0.64 | S ₈ -0.59 | | oop H (70), oop C (30) |
| v_{16} | 915 | S _{19b} -0.84 S _{11b} -0.67 | S ₇ 0.49 | | оор Н |
| v_{17} | 748 | $S_8 0.72 S_4 0.61$ | S _{11b} -0.53 | S ₇ -0.45 | oop H (67), oop C (33) |
| v_{18} | 698 | S ₈ 1.04 S ₄ -0.52 | S ₇ -0.41 | | oop C (67), oop H (33) |
| V ₁₉ | 481 | S _{20b} 0.66 S _{19b} 0.39 | S ₄ -0.32 | | oop C (50), oop C1 (50) |
| v_{20} | 230 | S _{20b} 0.24 S ₄ 0.13 | S _{11b} 0.11 | | oop C (60), oop Cl (40) |
| v_{21} | 3157 | S_{15b} 0.84 S_{12b} -0.59 | | | СН |
| v_{22} | 3151 | S_{12b} -0.85 S_{15b} -0.59 | | | СН |
| ν_{23} | 1584 | S _{18b} -0.80 S _{16b} -0.43 | S_{17b} -0.37 | | CC (75), HCC (25) |
| V ₂₄ | 1452 | S _{14b} 1.09 S ₃ -0.54 | | | HCC (65), CC (35) |
| v_{25} | 1330 | S ₃ -1.08 S ₁₀ -0.66 | • | | HCC (70), CC (30) |
| V ₂₆ | 1293 | S ₃ 0.59 S ₁₀ -0.49 | S ₉ 0.40 | S_{18b} -0.35 | CC (75), HCC (25) |
| V ₂₇ | 1158 | S ₁₀ -0.80 S _{18b} -0.61 | | | HCC (65), CC (35) |
| V ₂₈ | 1073 | S _{14b} 0.44 S ₁₀ 0.39 | S _{18b} -0.38 | S_{13b} 0.22 | CC (55), HCC (45) |
| V ₂₉ | 615 | S _{17b} 0,40 | | | CCC |
| V ₃₀ | 329 | S _{18b} 0.20 S _{14b} 0.18 | S ₁₀ -0.15 | S ₃ 0.11 | НСС |

a - The numbers are the $\partial S_i/\partial Q_j$ elements of the indicated symmetry coordinate in the particular vibration. Only the dominant terms are shown.

b - CH, CC or CCl are stretches, HCC is CH bend, CCC is ring deformation. Out-of-plane vibrations are denoted by oop. **Cent contributions are given in brackets..

4.3.1 - Assignments of the 11 A₁ vibrations of chlorobenzene

There are five CH stretches in chlorobenzene: three A_1 vibrations and two B_2 vibrations. According to the calculations all are expected within ~30 cm⁻¹ and allowing for anharmonicity should be at 3100 - 3050 cm⁻¹, about 100 cm⁻¹ lower than the calculated values.

The strong A-type band at 3080 cm⁻¹ in the infrared spectrum of the gas, has a counterpart in the strong infrared band in the liquid at 3070 cm⁻¹ (Fig 4.3). The wavenumber shift between the gas and liquid is in agreement with observations of benzene⁶⁹. There is a strong polarized Raman band at 3066 cm⁻¹. The Raman wavenumber calibration is such that this band may coincide with the infrared band at 3070 cm⁻¹ or there may be two A₁ vibrations at 3070 cm⁻¹ and 3066 cm⁻¹.

No other obvious A_1 vibrations are observed. The peak at 3058 cm⁻¹ in the infrared spectrum of the liquid is often assigned to an A_1 fundamental and we do the same, clearly for lack of a better alternative.

Calculations predict that v_4 is near 1595 cm⁻¹ with v_{23} (B₂) nearby at 1584 cm⁻¹. In the Raman spectra a depolarized band is observed at 1583 cm⁻¹ and a depolarized, weak shoulder at 1565 cm⁻¹ (Fig. 4.4). The gas's infrared band contour is not easy to interpret, but has a Q branch at 1589 cm⁻¹. In the infrared spectrum of the liquid there is a strong band at 1584 cm⁻¹, a peak at 1566 cm⁻¹ and a shoulder at ~1590 cm⁻¹. The evidence is not clear, but the gas phase Q branch and complex band shape lead us to assign v_4 (A₁) to the Q branch at 1589 cm⁻¹ in the gas and at 1584 cm⁻¹ in the liquid, in

both cases very close to v_{23} (B₂).

The calculated wavenumber for v_5 (A₁) is 1541 cm⁻¹ and that of v_{24} (B₂) is 1452 cm⁻¹. Both are due to the splitting of the strong ring mode of benzene. v_5 (A₁) is clearly at 1483 cm⁻¹ in the infrared spectrum of the gas and 1478 cm⁻¹ in the liquid where the liquid infrared and polarized Raman band essentially coincide (Fig. 4.4).

The next four A_1 vibrations are expected near 1270 cm⁻¹ for v_6 , 1150 cm⁻¹ for v_7 , 1030 cm⁻¹ for v_8 and 1000 cm⁻¹ for v_9 . The A-type contours in the infrared spectrum of the gas are located at 1232 cm⁻¹, 1208 cm⁻¹, 1176 cm⁻¹, 1126 cm⁻¹, 1092 cm⁻¹, 1025 cm⁻¹ and probably at 1004 cm⁻¹ (Fig. 4.5 and 4.6). All these bands have a corresponding band within 5 cm⁻¹ to low wavenumber in the infrared spectrum of the liquid. The polarized Raman bands in this region are located at 1173 cm⁻¹, 1124 cm⁻¹, 1084 cm⁻¹, 1023 cm⁻¹ and 1002 cm⁻¹. The bands at 1023 cm⁻¹ and 1002 cm⁻¹, for which the calculated wavenumbers agree with those from all three experimental spectra, are assigned to v_8 and v_9 respectively. The very strong band at 1084 cm⁻¹ in the liquid and 1092 cm⁻¹ in the gas is assigned to v_7 . We follow Whiffen³⁷ by assigning 1172 cm⁻¹ in the liquid and 1176 cm⁻¹ in the gas to v_6 and assigning the stronger band at 1123 cm⁻¹ in the infrared spectrum of the liquid to the combination $v_{10}+v_{11}$.

The last two A_1 vibrations are expected near 775 cm⁻¹ and 400 cm⁻¹. There are two strong polarized Raman bands at 702 cm⁻¹ and 418 cm⁻¹ (Fig. 4.7 and 4.8). A weak A-type band is observed in the spectrum of the gas at 706 cm⁻¹ and a very strong band in the liquid is at 703 cm⁻¹. It is assigned to v_{10} . v_{11} is assigned to the infrared and Raman

bands of the liquid near 415 cm⁻¹ correlated with the A-type band at 417 cm⁻¹ in the spectrum of the gas.

4.3.2 - Assignment of the 3 A2 vibrations of chlorobenzene

The three A₂ vibrations are expected near 970, 850 and 400 cm⁻¹. These vibrations are inactive in the infrared but may appear as very weak bands in the liquid. They are active in the Raman spectra and should be depolarized but are possibly weak.

The only weak, depolarized Raman band without a corresponding band in the infrared spectrum of the gas is found at 831 cm⁻¹ (Fig. 4.7). The band is essentially coincident with the 830 cm⁻¹ band in the infrared spectrum of the liquid and is assigned to v_{13} .

The weak Raman bands at 991 cm⁻¹ and 972 cm⁻¹ are clearly polarized and can not be assigned to v_{12} , while the strong, polarized Raman band at 418 cm⁻¹ masks the 400 cm⁻¹ region. Without evidence, we follow Whiffen³⁷ in assigning v_{12} at 964 cm⁻¹ and v_{14} at 400 cm⁻¹ at 400 cm⁻¹.

4.3.3 - Assignment of the 6 B₁ vibrations of chlorobenzene

The computed wavenumbers of the six B_1 fundamentals of chlorobenzene are: 987 cm⁻¹ for v_{15} , 915 cm⁻¹ for v_{16} , 748 cm⁻¹ for v_{17} , 698 cm⁻¹ for v_{18} , 481 cm⁻¹ for v_{19} and 230 cm⁻¹ for v_{20} . These bands should be depolarized in the Raman spectrum and should

have a C-type contour in the infrared gas spectrum.

Three strong bands at 903, 741 and 468 cm⁻¹ are observed in all experimental spectra (Fig. 4.6 to 4.8) with the appropriate contour and depolarization. They are assigned to v_{16} , v_{17} and v_{19} , respectively. The band at 685 cm⁻¹ is very strong in the liquid and has a C-type contour in the spectrum of the gas. Although no Raman band is observed nearby, it is assigned to v_{18} .

As the gas spectrum was not recorded below 400 cm⁻¹, the choice for v_{20} is based on the Raman bands and the computed wavenumbers. Of the two depolarized Raman bands at 296 and 195 cm⁻¹, v_{20} is assigned at 195 cm⁻¹ because its computed wavenumber is 100 cm⁻¹ lower than that of v_{30} (B₂).

 v_{15} was previously^{22,23,37,42} assigned at ~985 cm⁻¹. A band at 983 cm⁻¹ is observed in the infrared spectrum of the liquid. A band of undetermined shape is observed in the gas at 982 cm⁻¹ but the Raman shoulder observed in that area is clearly polarized. For lack of better evidence, we follow the previous assignments and assign v_{15} at 983 cm⁻¹.

4.3.4 - Assignment of the 10 B₂ vibrations of chlorobenzene

The computed wavenumbers for the two B₂ CH stretches are 3157 and 3151 cm⁻¹. Allowing for anharmonicity the observed wavenumbers should be about 100 cm⁻¹ lower. A possible B-type contour is observed at ~3095 cm⁻¹ in the infrared spectrum of

the gas corresponding to a band at 3083 cm⁻¹ in the infrared spectrum of the liquid (Fig 4.3). These features are assigned to v_{21} . More questionable B-type contours are observed at 3067, 3055 and 3022 cm⁻¹. The observed infrared bands of the liquid are a shoulder at ~3050 cm⁻¹ and peaks at 3059 (v_3 A₁), 3026 and 3016 cm⁻¹. There is a depolarized Raman band at ~3060 cm⁻¹ that is not coincident with the strong polarized band at 3066 cm⁻¹. As the computation suggest that all five CH stretches should be within 30 cm⁻¹, we assign v_{22} at ~3060 cm⁻¹. Thus we assign v_3 (A₁) and v_{22} (B₂) nearly coincident at 3060 cm⁻¹.

 v_{23} is expected near 1585 cm⁻¹. As was discussed previously, the depolarized Raman band at 1583 cm⁻¹ and the infrared band at 1584 cm⁻¹ can be assigned to v_{23} . v_{23} (B₂) and v_4 (A₁) combine to give the confused gas phase band in Fig. 4.4.

 v_{24} is expected at 1450 cm⁻¹. It is assigned to the very strong band at 1446 cm⁻¹ in the infrared spectrum of the liquid, the corresponding depolarized Raman band at 1445 cm⁻¹, and the B-type contour at ~1449 cm⁻¹ in the infrared spectrum of the gas.

v₂₅ is expected near 1330 cm⁻¹. It is assigned to the weak band at 1325 cm⁻¹ in the infrared spectrum of the liquid that is essentially coincident with a depolarized Raman band and a B-type band in the infrared spectrum of the gas.

 v_{26} is expected at 1290 cm⁻¹. Two bands are observed in the infrared spectrum of the liquid, at 1298 cm⁻¹ and 1272 cm⁻¹. Both bands correspond to depolarized Raman bands and B-type contours in the infrared spectrum of the gas. We follow Whiffen³⁷ in assigning v_{26} at 1272 cm⁻¹ and the combination $v_{14}+v_{16}$ at 1299 cm⁻¹.

 v_{27} is expected near 1160 cm⁻¹. It is assigned to the band at 1156 cm⁻¹ in the infrared spectrum of the liquid that is essentially coincident with a depolarized Raman band.

v₂₈ is expected near 1075 cm⁻¹. It is assigned to the band at 1068 cm⁻¹ in the infrared spectrum of the liquid coincident with a B-type band in the infrared spectrum of the gas. A probable depolarized Raman band at 1069 cm⁻¹ is observed on the side of the strong polarized Raman band at 1084 cm⁻¹.

v₂₉ is expected near 615 cm⁻¹. It is assigned to the medium-weak band at 614 cm⁻¹ in the infrared spectrum of the liquid and the corresponding depolarized Raman band and B-type gas phase band.

Finally, v_{30} is expected near 330 cm⁻¹. As discussed in Section 4.3.3, the depolarized Raman band at 296 cm⁻¹ is assigned to v_{30} while that at 196 cm⁻¹ is assigned to v_{20} (B₁).

4.3.5 - Summary of the assignments of chlorobenzene

The proposed assignments are given in Table 4.10. The true Herzberg description is given in the first column followed by the proposed wavenumber. When a wavenumber was chosen with imperfect or no supporting evidence from the experimental data gathered for this thesis, it is indicated by ? or ??, respectively.

In the pseudo-Herzberg or pseudo-Wilson notations for the vibrations of the

Table 4.10 - Assignments of the fundamental vibrations of chlorobenzene

| true Herzberg | cm ^{-1 a} | pseudo-Herzberg ^b | Description ^c |
|-----------------------|--------------------|--|--------------------------|
| v_1 | 3069.5 | v_5, v_1 | СН |
| v_2 | 3069 ?? | v_1, v_{15a}, v_5 | СН |
| v ₃ | 3058.5 ?? | v_{12a}, v_{15a} | СН |
| v_4 | 1584 ?? | v_{18a}, v_{16a} | HCC, CC |
| V ₅ | 1477.8 | V_{14a} | НСС |
| ν ₆ | 1171.6? | v_{18a}, v_{14a} | HCC |
| v ₇ | 1083.6 | v_{18a} | HCC |
| v ₈ | 1022.8 | v_{14a}, v_6 | HCC, CCC |
| V 9 | 1001.9 | $\mathbf{v_6}$ | ccc |
| v_{10} | 702.5 | v_{17a} , v_6 | ccc |
| v_{11} | 415.0 | v_{17a} , v_{18a} | CCC, HCC |
| v_{12} | 963.7 ?? | v_{19a} | oop H |
| V ₁₃ | 830.0 | v_{l1a} | оор Н |
| v_{14} | 400 ?? | V _{20a} | oop C |
| v ₁₅ | 983.2 ?? | v_7 , v_{19b} , v_8 | oop H. oop C |
| V ₁₆ | 903.0 | v _{19b} , v _{11b} , v ₇ | oop H |
| V ₁₇ | 741.2 | v_8, v_4, v_{11b}, v_7 | oop C, oop H |
| $\mathbf{v_{ls}}$ | 685.2 | v_8, v_4 | oop C, oop H |
| v_{19} | 468.1 | v_{19b}, v_4 | оор С, оор Н |
| v_{20} | 195.9 | V _{20b} , V ₄ | оор С, оор Н |
| v_{21} | 3083.1 ? | v_{15b}, v_{12b} | СН |
| v ₂₂ | 3060 ?? | v_{12b}, v_{15b} | СН |
| V ₂₃ | 1583.9 | v_{18b}, v_{16}, v_{17b} | HCC, CC, CCC |
| V ₂₄ | 1445.5 | v_{14b} | HCC |
| V ₂₅ | 1325.2 | v_3 , v_{10} | НСС |
| V ₂₆ | 1272.5 ? | $v_3, v_{10}, v_9, v_{18b}$ | HCC, CC |
| V ₂₇ | 1156.4 | v_{10}, v_{18b} | нсс |
| V ₂₈ | 1068.1 | $v_{14b}, v_{10}, v_{18b}, v_{13b}$ | HCC |
| V ₂₉ | 614.1 | v_{17b} | ccc |
| v ₃₀ | 297.5 | V _{18b} , V _{14b} , V ₁₀ , V ₃ | нсс |

a - Peak wavenumbers in the imaginary molar polarizability spectrum of liquid chlorobenzene.

b - Contributions greater than 50% of the largest contribution.

c - CH, CC are stretches, HCC is CH bend, CCC is ring deformation and oop is out-of-plane.

monosubstituted benzenes the symbol v_j has been used to relate the vibration to one symmetry coordinate of benzene, not to one vibration of benzene⁶⁸. We follow this practice, except that it is clear from Table 4.9 that most of the chlorobenzene vibrations are mixtures of more than one symmetry coordinate of benzene and therefore more than one v_j is required in the pseudo-Herzberg notation. In column 3, v_j is given for each symmetry coordinate that contributed ≥ 50 % of the largest contribution. Note that v_2 and v_{13a} , both CC stretching displacements, do not appear in this table. Although these displacements contribute to many vibrations they are always minor contributors. The last column gives a chemical description of the vibration.

A question arises from the table. Where are the C-Cl stretching vibration and the Cl-C-C in-plane and out-of-plane bending vibrations? From the unsymmetrized eigenvectors, $\frac{\partial R_{C-Cl}}{\partial Q_k}$, the C-Cl stretch contributes with HCC, CC and CCC to the following A₁ vibrations: v_4 (1595), v_5 (1541), v_6 (1773), v_7 (1148), v_{10} (773) and v_{11} (398). It does not make the major contribution to any of these vibrations. From the potential energy distribution (PED), the C-Cl stretch is among the three main contributors to v_6 (1270), v_7 (1148), v_{10} (773) and v_{11} (398) but is the major contributor to none of these. Therefore we conclude that no single vibration can be identified as the C-Cl stretch.

Both the unsymmetrized eigenvectors and the PED show that v_{30} (B₂) is the Cl-C-C in-plane deformation vibration. They also show that v_{19} and v_{20} are both heavy

mixtures of the Cl-C-C and C-C-C out-of-plane deformations, so the Cl-C-C oop deformation can not be assigned to any one vibration.

4.4 - Previous assignments of the toluene vibrations

Toluene, has 15 atoms and 15x3-6 = 39 fundamental vibrations. Technically the symmetry of the molecule is at most C_* but often the CH₃ group is considered to be freely rotating which means that the CH₃ group can be approximated by a point of mass 15, and thus the molecule has C_{2v} symmetry. Assuming C_{2v} symmetry, the 39 vibrational modes are distributed among the symmetry species as $13A_1+4A_2+9B_1+13B_2$. As is the case for chlorobenzene, all of the vibrations in toluene are Raman active, and all except the $4A_2$ vibrations are infrared active. Of the 39 vibrations, 30 vibrations are similar to the 30 vibrations of chlorobenzene and 9 vibrations are related to the CH₃ vibrations. The CH₃ related modes form the representation $2A_1+1A_2+3B_1+3B_2$ under C_{2v} .

Although the CH₃ group can be approximated in simple calculations as an atom of mass 15, the 9 vibrations of the CH₃ group cause problems in notation. If true Herzberg notation is used, the A₂, B₁ and B₂ vibrations would be assigned different number ranges compared with other monosubstituted benzenes. Therefore, the common practice is to treat the phenyl group vibrations separately and to number them using either the true Herzberg notation, v_1 to v_{30} , or the pseudo-Wilson or pseudo-Herzberg notations. The CH₃ vibrations are considered then extra and are usually

labeled as v_s , v_a , δ_s , δ_a , and r, where v, δ and r indicate CH bond stretching, HCH angle deformation and CH₃ rocking vibrations, and the "s" and "a" designate symmetric and antisymmetric vibrations, respectively. v_a and δ_a are sometimes marked with 'or" to differentiate between the B₁ and B₂ species.

As was the case for chlorobenzene, the use of different spectra is necessary in order to determine the symmetry species of toluene. However, the difference between shapes⁷⁴ of A-type and C-type bands in the gas phase infrared spectrum of toluene, is less obvious than for chlorobenzene, so the assignment of symmetry species is more ambiguous. The infrared and Raman spectra of the liquid provide similar information to that obtained for chlorobenzene. Comparison of observed wavenumbers in infrared and Raman spectra permitted several authors to present a complete assignment of the fundamental vibrations of toluene.

The first major study of toluene since 1950, was by Wilmshurst and Bernstein¹⁵. The authors reported spectra of toluene and toluene-α-d₃. They also used literature data for toluene-p-d. They included figures of their infrared spectra of the gas and liquid and their Raman spectra of the liquid, and tabulated all the wavenumbers they observed. They reported gas-phase band types and whether the Raman bands were polarized or depolarized. They assigned wavenumbers to fundamental vibrations and to overtone and combination transitions. On the basis of the CH₃ to CD₃ isotope shifts, they related some of the fundamentals to the vibrations of benzene using the pseudo-Wilson notation, which was converted in this thesis to the pseudo-Herzberg notation.

They took the plane of the molecule as XZ. Thus, their B_1 species are interchanged with our B_2 species.

In 1960, Fuson *et al.*¹⁷ reported the spectra of C₆H₅CH₃, C₆H₅CD₃ and C₆D₅CD₃. The authors tabulated all the wavenumbers they observed in the infrared spectra of the gas and liquid phases and included Raman data from Wilmshurst and Bernstein¹⁵ which was confirmed by measurements in their laboratory. For the phenyl group vibrations the authors used the pseudo-Wilson notation, which was converted to the pseudo-Herzberg notation throughout this thesis. Their B₁ species are interchanged with our B₂ species. For the CH₃ group vibrations, the authors used ' and " to differentiate between the B₁ and B₂ species. The authors also compared their assignments and those of Wilmshurst and Bernstein¹⁵ and those of Kovner¹⁶.

In 1960, Schmid, Brandmuller and Nonnenmacher¹⁸ performed normal coordinate calculations on toluene with the same force field as for benzene and obtained the wavenumbers of the 30 phenyl group vibrations in toluene. The CH₃ group was approximated in the calculations by an atom of mass 15. In addition to the calculated wavenumbers, experimental wavenumbers obtained from gas and liquid phase infrared and liquid phase Raman measurements were also given. Their B₁ species are interchanged with our B₂ species. and for the 30 ring vibrations the authors used the true Herzberg notation. Alongside, a pseudo-Wilson notation is also given.

In 1971, La Lau and Snyder¹⁹ derived a valence force field for methyl substituted benzenes from 303 observed wavenumbers of benzene, toluene, p-xylene, m-xylene,

mesitylene and some of their deuterated derivatives. For toluene, they gave figures of the infrared spectrum of the crystalline phase and tabulated the observed fundamental wavenumbers and their computed wavenumbers together with their potential energy distribution. In a separate table, they gave the wavenumbers of fundamentals below 1606 cm⁻¹ of the Raman spectrum of the liquid with the polarization of the band.

As was the case for chlorobenzene, the books by Sverdlov et al.²² and by Varsanyi²³ give the wavenumbers and assignments for toluene without explanation. In the book by Sverdlov et al., the B₁ and B₂ species are interchanged with our B₂ and B₁ species. Sverdlov et al. used the true Herzberg notation system for all 39 vibrations. In the book by Varsanyi, the pseudo-Wilson notation is used.

In 1985, Draeger²⁵ reported experimental and theoretical wavenumbers for benzene, toluene and some of their deuterated derivatives. The experimental wavenumbers were obtained from Raman and infrared measurements of the gas. The wavenumbers were classified according to symmetry species and a potential energy distribution was given.

In 1986, Xie and Boggs²⁶ reported fundamental wavenumbers obtained from scaled *ab initio* calculations. The experimental wavenumbers used in the scaling were those of Wilmshurst and Bernstein¹⁵, Kovner *et al.*¹⁶, Fuson *et al.*¹⁷ and Draeger²⁵. Xie and Boggs numbered the wavenumbers of all 39 vibrations in order of increasing wavenumber regardless of symmetry species.

In 1995, Schrotter and co-workers^{32,33} reported experimental Raman

wavenumbers of the gas and liquid phases and wavenumbers obtained from normal coordinate calculation based on the experimental wavenumbers. The B₁ and B₂ species were interchanged with our notation and the authors used the true Herzberg notation for all vibrations. However, one A₂ vibration was omitted and, consequently, the numbering scheme is off by one for the subsequent vibrations.

In the following sub-sections, the wavenumbers and assignments of the 30 phenyl group vibrations by these authors are given for each symmetry species, followed by the assignments and wavenumbers of the 9 CH₃ vibrations. Differences in assignment will be pointed out. However a critical evaluation of the authors' assignment is left to Section 4.5 where the assignment is evaluated on the basis of all available data, including data measured for this thesis.

Other studies either gave only partial wavenumbers and assignments^{27,29} or used wavenumbers and assignments from previous studies without change^{28,30,31,34}. These studies will be quoted and used when deemed necessary.

4.4.1 - Previous assignments of the 11 A_1 phenyl group vibrations of toluene

The wavenumbers and assignments of the 11 A₁ phenyl group vibrations of liquid toluene are given in Table 4.11. The first column gives the true Herzberg notation for the 30 phenyl vibrations under the qualification that the CH₃ group vibrations are numbered 31 to 39 regardless of symmetry species. The pseudo-Herzberg notation is given in the second column. In Table 4.11 and later in Tables 4.12 to 4.14, the pseudo

notation is taken from Schmid et al. 18 who determined the relation between the vibrations of substituted benzenes and those of benzene. Schmid's original pseudo-Wilson notation was converted to the pseudo-Herzberg notation through use of Table 4.3. The remaining columns contain the wavenumbers and assignments of the different authors. The experimental wavenumbers listed in the table are those from the infrared measurement of the liquid phase. When only the Raman band was observed, it is denoted by the letter R next to the wavenumber.

In general, the agreement between the different experimental studies, on the A_1 phenyl wavenumbers is very good, with differences that can be attributed to calibration errors, often within ± 5 cm⁻¹ of each other. There are, however, two major differences in wavenumbers and several differences in the pseudo-Herzberg notation. The latter involve interchange of v_1 , v_5 , v_{123} and v_{15a} , all CH stretches, and interchange of v_2 and v_{18a} , a CC stretch and a HCC deformation, respectively.

Two assigned wavenumbers for the three aromatic A₁ CH stretching vibrations are in good agreement among the different experimental studies, 3063±4 cm⁻¹ and 3055±1 cm⁻¹. The third A₁ CH stretching vibration was assigned by Wilmshurst and Bernstein¹⁵, Varsanyi²³ and Schrotter *et al.*³² at 3003 cm⁻¹. Sverdlov *et al.*²² placed it at 3060 cm⁻¹. Fuson *et al.*¹⁷ placed it at 3087 cm⁻¹, above their other two CH stretches, and Draeger²⁵ placed it at 3039 cm⁻¹, below the other CH stretches. Consequently the true Herzberg notation, for, e.g., 3063 cm⁻¹ is different for different workers (Table 4.11).

The second major difference is the wavenumber of v4. Wilmshurst and

Table 4.11 - Previous assignments of the 11 A₁ phenyl vibrations of toluene

| | | | | | | Experime | ental | | | | Theoretical | | | |
|----------------|------------------|-------------------------|------------------|--------------------------------|-----------------------------|-----------------------------------|------------------|------------------|----------------------------------|---------------------------------|---------------------------------|---------------------------------|--------------------------------|--|
| | rzberg lation | Wilmshurst* [Ref 15] | | Fuson ^a [Ref 17] | | Sverdlov ^b [Ref 22] | Vars [Ref | • | Draeger ^c [Ref 25] | Schrotter ^d [Ref 32] | Schmid ^e [Ref 18] | Snyder ^c [Ref 19] | Boggs ^f [Ref 26] | |
| true | pseudo | cm ⁻¹ | | cm ^{-t} | | cm ⁻¹ | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ | |
| v_{i} | v_{12a} | 3067 | ν_{12a} | 3087 | V _{15a} | 3060 | 3063 | v_{12a} | 3067 | 3065R | 3075 | 3057 | 3088 | |
| V ₂ | v_i | 3056 | $\mathbf{v_i}$ | 3063 | v_{i2a} | 3060 | 3055 | v_l | 3056 | 3055R | 3069 | 3056 | 3063 | |
| v_3 | V5 | 3003 | V5 | 3055R | $\mathbf{v_i}$ | 3060 | 3003 | ν_{15a} | 3039 | 3003R | 3053 | 3055 | 3049 | |
| V4 | Vion | 1585R | Y'(oe | 1605 | $\mathbf{v}_{\mathrm{ios}}$ | 1608 | 1605 | Vide | 1611 | 1606R | 1656 | 1614 | 1620 | |
| V5 | V13a | 1491 | V _{13a} | 1494 | V _{13a} | 1500 | 1494 | V13a | 1500 | 1495R | 1524 | 1503 | 1504 | |
| V ₆ | V _{15a} | 1209 | Visa | 1208 | V5 | 1218 | 1208 | ν5 | 1212 | 1210R | 1299 | 1205 | 1214 | |
| V 7 | V _{17a} | 1178 | V _{17a} | 1175 | V ₁₇₈ | 1184 | 1175 | V _{17a} | 1178 | 1180R | 1157 | 1182 | 1189 | |
| v ₈ | V _{14a} | 1029 | V14a | 1030 | V ₁₄₈ | 1030 | 1030 | V _{14a} | 1030 | 1030R | 1023 | 1026 | 1034 | |
| Vq | V ₆ | 1002 | V ₆ | 1003 | V ₆ | 1004R | 1003 | V 6 | 1004 | 1005R | 1000 | 1005 | 1001 | |
| Vio | V18a | 785 | V ₂ | 784 | V ₂ | 774 | 784 | \mathbf{v}_2 | 786 | 787R | 778 | 793 | 784 | |
| VII | $\mathbf{v_2}$ | 521 | V _{18a} | 521 | Viga | 521 | 521 | V18a | 521 | 521R | 506 | 526 | 519 | |

a - Assignments in pseudo-Wilson notation were converted to pseudo-Herzberg notation using Table 4.3.

Bernstein placed it at 1585 cm⁻¹ with a B₂ vibration at 1604 cm⁻¹. Other experimental studies reverse these assignments.

The wavenumbers calculated by Schmid et al. 18 and Boggs et al. 26 are in

b - Authors used the true Herzberg notation for 39 vibrations.

c - No notation was used.

d - Authors used the true Herzberg notation for 38 vibrations. See Section 4.4.

e - Authors used the true Herzberg notation for 30 ring vibrations.

f - Authors numbered all 39 vibrations in order of increasing wavenumber.

Table 4.12 - Previous assignments of the 3 A₂ phenyl vibrations of toluene

| | | | Theoretical | | | | | | | | | | |
|------------------|------------------|------------------|-------------|------------------|------------------|-----------------------------------|-----|-----------------------------|----------------------------------|------------------------------------|---------------------------------|---------------------------------|--------------------------------|
| | zberg ation | Wilms [Ref | | Fuso | | Sverdlov ^b [Ref 22] | | sanyi [*] f 23] | Draeger ^c [Ref 25] | Schrotter ⁴ [Ref 32] | Schmid ^e [Ref 18] | Snyder ^s [Ref 19] | Boggs ^t [Ref 26] |
| true | pseudo | cm ⁻¹ | | cm ⁻¹ | | cm ⁻¹ | ₊m¹ | | em ^{-t} | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ |
| ν_{12} | v_{i9a} | 994R | V19a | 964 | v_{lo_a} | 970R | 964 | Viva | 964R | 990R | 979 | 975 | 965 |
| V ₁ 3 | v_{11a} | 842 | Vila | 844 | V118 | 836 | 843 | V314 | 843R | 844R | 848 | 847 | 836 |
| V14 | V _{20a} | 407 | V20a | 408R | V _{20a} | 414R | 408 | V20a | 405R | 408R | 405 | 403 | 401 |

a - Assignments in pseudo-Wilson notation were converted to pseudo-Herzberg notation using Table 4.3.

satisfactory agreement with the experimental values, while the wavenumbers computed by La Lau and Snyder¹⁹, are in excellent agreement with the different experimental values with the exception of the wavenumbers of the CH stretches.

4.4.2 - Previous assignments of the 3 A2 phenyl group vibrations of toluene

The wavenumbers and assignments of the A_2 vibrations are given in Table 4.12 which is arranged similarly to Table 4.11. One major disagreement is the assignment of v_{12} . Wilmshurst and Bernstein¹⁵ and Schrotter *et al.*³² placed it at 992±2 cm⁻¹, while the other experimental studies^{17,22,23,25} placed it at about 25 cm⁻¹ lower at 967±3 cm⁻¹. The

b - Authors used the true Herzberg notation for 39 vibrations.

c - No notation was used.

d - Authors used the true Herzberg notation for 38 vibrations. See Section 4.4.

e - Authors used the true Herzberg notation for 30 ring vibrations.

f - Authors numbered all 39 vibrations in order of increasing wavenumber.

Table 4.13 - Previous assignments of the 6 B₁ phenyl vibrations of toluene

| | | | | | | | Theoretical | | | | | | |
|----------------------|------------|----------------------|----------------|--------------------|------------------|-----------------------------------|-----------------------------------|------------------|----------------------------------|---------------------------------|---------------------------------|---------------------------------|--------------------------------|
| Herzberg notation | | Wilmshurst* [Ref 15] | | Fuson* {Ref 17} | | Sverdlov ^b [Ref 22] | Varsanyi ^a [Ref 23] | | Draeger ^c [Ref 25] | Schrotter ^d [Ref 32] | Schmid ^e [Ref 18] | Snyder ^c [Ref 19] | Boggs ^f [Ref 26] |
| true | pseudo | cm ⁻¹ | | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ |
| V15 | V 7 | 966 | V7 | 978 | V 7 | 993R | 978 | V7 | 980 | 974R | 984 | 989 | 976 |
| V ₁₆ | V19b | 895 | VIIb | 893 | V19b | 892 | 893 | Visb | 894 | 897R | 920 | 909 | 892 |
| V ₁₇ | VIIb | 729 | V4 | 728 | V4 | 716 | 728 | V4 | 730 | 731R | ⁻ 54 | 734 | 721 |
| VĮS | V | 695 | Vq | 495 | Vş | 702 | 695 | Vg | 695 | | 700 | 698 | 688 |
| V19 | V4 | 464 | : 2° ⋅. | ó4 | V _{20b} | 465 | 464 | V _{20b} | 462 | 466R | 497 | 463 | 457 |
| V ₂₀ | V20b | 216 | V196 | 217R | VIIb | 217R | 217 | VIII | 205 | 217R | 254 | 210 | 201 |

a - Assignments in pseudo-Wilson notation were converted to pseudo-Herzberg notation using Table 4.3.

computed wavenumbers ^{18,19,26} of this vibration range from 965 to 979 cm⁻¹. The wavenumbers of the remaining two A₂ phenyl vibrations are in excellent agreement among the different experimental studies with the exception of Sverdlov's which are off by 6-8 cm⁻¹. The computed wavenumbers for these two vibrations are in very good agreement with the experimental values.

b - Authors used the true Herzberg notation for 39 vibrations.

c - No notation was used.

d - Authors used the true Herzberg notation for 38 vibrations. See Section 4.4.

e - Authors used the true Herzberg notation for 30 ring vibrations.

f - Authors numbered all 39 vibrations in order of increasing wavenumber.

4.4.3 - Previous assignments of the 6 B1 phenyl group vibrations of toluene

The wavenumbers and assignments of the B₁ vibrations are given in Table 4.13 which is arranged in a similar manner to Table 4.11. There are several disagreements about wavenumber assignments of the B₁ vibrations.

V₁₅ is assigned by Wilmshurst and Bernstein¹⁵ at 966 cm⁻¹, while Fuson *et al.*¹⁷, Varsanyi²³, Draeger²⁵ and Schrotter *et al.*³² placed it around 977±3 cm⁻¹. Sverdlov *et al.*²² placed it at 993 cm⁻¹ based on Raman measurement. The computed wavenumbers^{18,19,26} for this vibration range from 976 to 991 cm⁻¹.

In two other cases, one study disagrees with the remaining experimental studies.

All experimental studies placed v_{17} at 729±2 cm⁻¹ with the exception of Sverdlov *et al.*who placed it at 716 cm⁻¹. All experimental studies assigned v_{20} at 217±1 cm⁻¹ with the exception of Draeger²⁵ who placed it at 205 cm⁻¹

The pseudo notations v_4 , v_{11b} , v_{19b} and v_{20b} are often interchanged in different studies ^{15,17,18,23}. However these differences are minor as all of these displacements are essentially mixtures of out-of-plane H and out-of-plane C displacements.

4.4.4 - Previous assignments of the 10 B₂ phenyl group vibrations of toluene

The wavenumbers and assignments of the 10 B₂ vibrations are given in Table
4.14 which is arranged in a similar manner to Table 4.11. There are several
disagreements on the wavenumbers of the B₂ vibrations. These disagreements cause a

Table 4.14 - Previous assignments of the 10 B₂ phenyl vibrations of toluene

| | | | | | | Theoretical | | | | | | | |
|----------------------|------------------|----------------------------------|------------------|------------------|------------------|-----------------------------------|-----------------------|------------------|----------------------------------|---------------------------------|---------------------------------|---------------------------------|--------------------------------|
| Herzberg notation | | Wilmshurst ^a [Ref 15] | | Fuson* [Ref 17] | | Sverdlov ^b [Ref 22] | Varsanyi* [Ref 23] | | Draeger ^c [Ref 25] | Schrotter ^d [Ref 32] | Schmid ^e [Ref 18] | Snyder ^c [Ref 19] | Boggs ^f [Ref 26] |
| true | pseudo | cm ⁻¹ | | cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | | em ⁻¹ | cm ^{-t} | cm ⁻¹ | cm ⁻¹ | cm ⁻¹ |
| V21 | Vi2b | 3090 | V12b | 3039R | VISB | 3074 | 3039 | V _{15b} | 3056 | 3065R | 3079 | 3055 | 3074 |
| V ₂₂ | v_{15b} | 3032 | V15b | 3029 | ν_{12b} | 3034 | 3029 | v_{12b} | 3039 | 3034R | 3044 | 3054 | 3052 |
| V ₂₃ | V _{16b} | 1600 | Vi6b | 1586 | v_{lob} | 1581 | 1586 | Viób | 1585 | 1586R | 1593 | 1589 | 1602 |
| V24 | V13b | 1455 | V _{13b} | 1494 | V9 | 1439R | 1468 | V:Di | 1463 | 1441R | 1444 | 1445 | 1468 |
| V25 | \mathbf{v}_3 | 1155 | Vo | 1460 | v_{13b} | 1330 | 1331 | v_3 | 1330 | 1332R | 1308 | 1326 | 1338 |
| V20 | V9 | 1155 | V ₁₀ | 1312 | ν3 | 1278R | 1318 | Vo | 1278 | 1308R | 1280 | 1290 | 1192 |
| V27 | V17b | 1105 | V3 | 1154 | V _{17b} | 1162 | 1154 | V17b | 1155 | 1156R | 1147 | 1168 | 1123 |
| V28 | Vio | 1081 | V _{14b} | 1080 | v_{14b} | 1086 | 1080 | v_{14b} | 1083 | 1082R | 1092 | 1081 | 1082 |
| V29 | v_{18b} | 623 | VISb | 623R | V _{18b} | 623 | 623 | v_{18b} | 623 | 623R | 632 | 616 | 628 |
| V30 | V _{14b} | 345 | V17b | 344R | VIO | 347 | 344 | V ₁₀ | 341 | 347R | 388 | 341 | 341 |

a - Assignments in pseudo-Wilson notation were converted to pseudo-Herzberg notation using Table 4.3.

different wavenumber ordering which leads to different true Herzberg notations for the same wavenumber in different studies. Thus, ~ 1155 cm⁻¹ is v_{26} (and v_{25}) for Wilmshurst and Bernstein¹⁵ but v_{27} for most other works^{17,22,23,25,32}.

One of the two aromatic CH stretching B2 vibrations, is generally agreed to be at

b - Authors used the true Herzberg notation for 39 vibrations.

c - No notation was used.

d - Authors used the true Herzberg notation for 38 vibrations. See Section 4.4.

e - Authors used the true Herzberg notation for 30 ring vibrations.

f - Authors numbered all 39 vibrations in order of increasing wavenumber.

 $3032 \pm 3 \text{ cm}^{-1}$ with the exception of Draeger²⁵ who placed it at 3039 cm⁻¹. There is no agreement on the assignment of v_{21} which has been placed from 3090 to 3039 cm⁻¹.

 v_{23} was assigned by Wilmshurst and Bernstein¹⁵ at 1600 cm⁻¹, while the other experimental studies^{17,22,23,25,32} placed it at 1585±5 cm⁻¹. The two groups of authors interchanged the assignment of v_{23} and v_4 (A₁) vibration.

Different choices were made for the B_2 phenyl group vibrations between 1500 and 1100 cm⁻¹. Consequently, the wavenumbers and true Herzberg notation of v_{24} to v_{27} disagree in the different studies. The only agreement is that one fundamental is near 1155 cm⁻¹ and a second is near 1455 cm⁻¹.

All authors agree on the assignment of v_{28} to v_{30} .

The differences in the pseudo notation are not serious. Under the pseudo-Herzberg notation, v_{12b} and v_{15b} are both CH stretching displacements, while v_3 , v_9 , v_{10} , v_{13b} , v_{14} , v_{17b} and v_{18b} are HCC deformation and CC stretching displacements which mix in the normal vibration.

4.4.5 - Previous assignment of the 9 CH₃ group vibrations of toluene

The wavenumbers and assignments of the 9 CH₃ group vibrations are given in Table 4.15 which is arranged in a similar manner to Table 4.11. The vibrations are arranged in order of symmetry species, i.e. 2 A₁, 1 A₂, 3B₁ and 3 B₂, and are numbered 31 to 39 using the true Herzberg notation.

Table 4.15 - Previous assignments of the 9 CH₃ group vibrations of toluene

| | | | Experimental | | | | | | | | | Theoretical | | | |
|------------------|------------------|------------------|----------------|---------------------------------|---------------------------|---|---|---------------------|--|--|---|---|--|--|--|
| True Herzberg | Sym. | Wilmsh [Ref I | | Fus [Ref cm ⁻¹ | | Sverdlov ^b [Ref 22] cm ⁻¹ | Varsanyi ^a [Ref 23] cm ^{.1} | | Draeger ^c [Ref 25] cm ⁻¹ | Schrotter ^d [Ref 32] cm ⁻¹ | Schmid ^e [Ref 18] cm ⁻¹ | Snyder ^c [Ref 19] cm ⁻¹ | Boggs ^f [Ref 26] cm ⁻¹ | | |
| V31 | Aı | 2923 | V _s | 2921 | Vs | 2910 | 2921 | V _s | 2921 | 2920R | | 2900 | 2919 | | |
| V ₃₂ | A_i | 1377 | δs | 1379 | δs | 1380 | 1379 | δ_{s} | 1384 | 1379R | | 1379 | 1384 | | |
| V ₃₃ | A ₂ | | | | t | | | | 15 | | | 44 | | | |
| V34 | $B_{\mathbf{i}}$ | 2930 | v | 2979 | v _a "' | 2976 | 2979 | Vas | 2933 | | | 2949 | 2976 | | |
| V ₃₅ | Bı | 1436 | δ′ | 1460 | $\delta_{\textbf{a}}{''}$ | 1462 | 1460 | δ_{as} + | 1453 | 1441R | | 1435 | 1454 | | |
| V ₃₆ | Bı | 1041 | r | 1040 | ra" | 1040 | | | 1043 | | | 104(5 | 1043 | | |
| V ₃₇ | B ₂ | 2952 | v | 2952 | v _a ' | 2976 | 2952 | v_{as} | 2933 | 2982R | | 29 % i | 2990 | | |
| V38 | B ₂ | 1460 | δ′ | 1460 | $\delta_{a}{}'$ | 1462 | | | 1463 | | | 1463 | 1468 | | |
| V39 | B ₂ | 1081 | г | 1040 | ra' | 970R | 1040 | δ ₃₅ - | 980 | | | 976 | 983 | | |

a - Authors used their own notation.

There is general agreement on the wavenumbers of the A_1 vibrations. Only Draeger²⁵ has assigned the A_2 CH₃ torsion. There are several disagreements in the wavenumbers of the B_1 and B_2 vibrations.

There is disagreement over the wavenumbers, particularly of the two antisymmetric CH₃ stretching vibrations, and over the wavenumber of v_{35} (B₁), one of

b - Authors used the true Herzberg notation for 39 vibrations.

c - No notation was used.

d - Authors used the true Herzberg notation for 38 vibrations. See Section 4.4.

e - Authors used the true Herzberg notation for 30 ring vibrations.

f - Authors numbered all 39 vibrations in order of increasing wavenumber.

the two antisymmetric CH_3 deformations, and its proximity to v_{38} , the B_2 deformation. The wavenumber of the CH_3 rocking modes is also not settled. The most reliable assignment appears to be that of Fuson et al¹⁷ who found that some of Wilmshurst and Bernstein's doublet bands were actually due to noise, and who had the spectra of toluene- α -d₃ to guide their determination of the bands due to the CH_3 vibration.

4.5 - Assignment of the vibrations of toluene

The methods used to evaluate the assignment of the toluene vibrations are similar to those used for the chlorobenzene assignment. Thus, the infrared spectrum of the gas and the Raman spectrum of the liquid recorded under parallel and perpendicular polarizations with linearly polarized incident light were obtained and a simple normal coordinate calculation was performed for toluene.

There are however two minor differences. First, the CH₃ group was approximated in the calculation by an atom of mass 15. Therefore no wavenumbers, eigenvectors or PED calculations are available for the 9 CH₃ vibrations. Second, the difference between A-type and C-type contours is not as clear as for chlorobenzene because of smaller differences between the moments of inertia of toluene.

Although Wilmshurst and Bernstein¹⁵ gave figures of the infrared spectra of the gas and liquid phases and tabulated, as did Fuson *et al.*¹⁷, all of the observed features in their infrared and Raman spectra, the experimental data and the results of the calculations obtained for this thesis will be presented on expanded scales, so as to form a

basis for the evaluation.

The liquid phase refractive index spectra of toluene reported in Chapter 2 were converted into the liquid phase molar polarizability spectra by use of Equations 1.3.33 and 1.3.28. The entire imaginary molar polarizability, α_m'' , spectrum of toluene is shown in Figure 4.9. Figures 4.10 through 4.14 show the infrared absorbance spectrum of the gas, the α_m'' spectrum, and the parallel and perpendicular Raman spectra of the liquid for smaller wavenumber ranges, and thus provide a better view of the features in these spectra.

Table 4.16 contains the information for every feature observed in the three different spectra of toluene for the regions where the fundamentals are expected, i.e. 3150 - 2850 cm⁻¹ and 1700 - 200 cm⁻¹. The table is arranged in a similar manner to Table 4.8 of chlorobenzene.

The computed wavenumbers, eigenvectors and potential energy distribution (PED) for the 30 phenyl group vibrations are given in Table 4.17. The table is arranged in a similar manner to Table 4.9 of chlorobenzene.

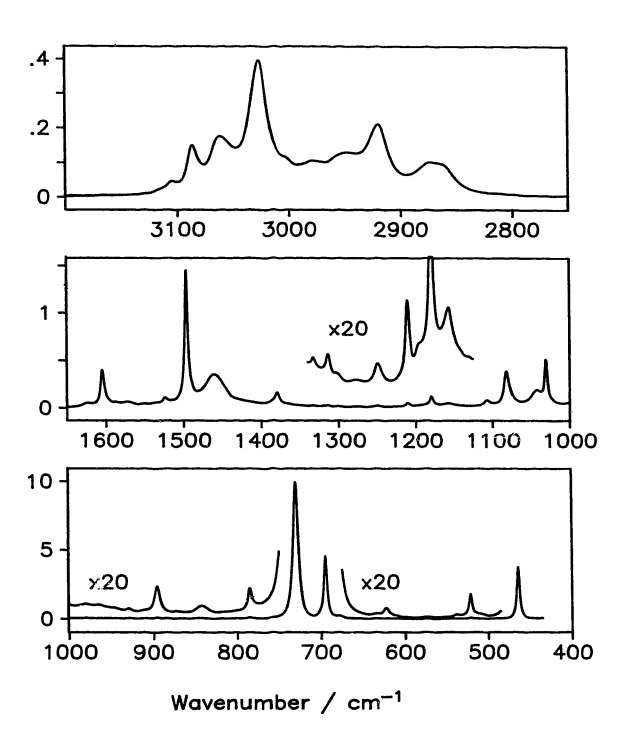


Figure 4.9 - Imaginary molar polarizability, α''_{m} , spectrum of liquid toluene. Units are cm³ mol⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by 20 for the upper curve.

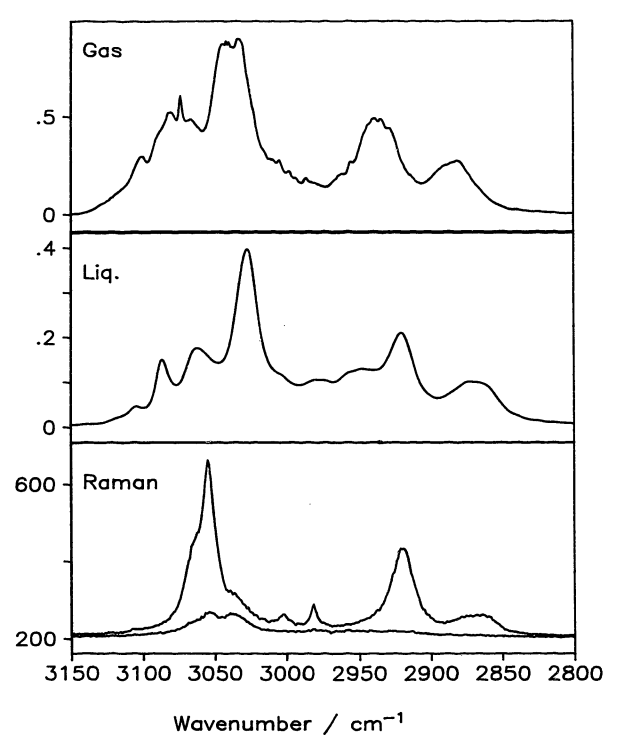


Figure 4.10 - Infrared spectra of gas and liquid toluene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 3150 and 2800 cm⁻¹.

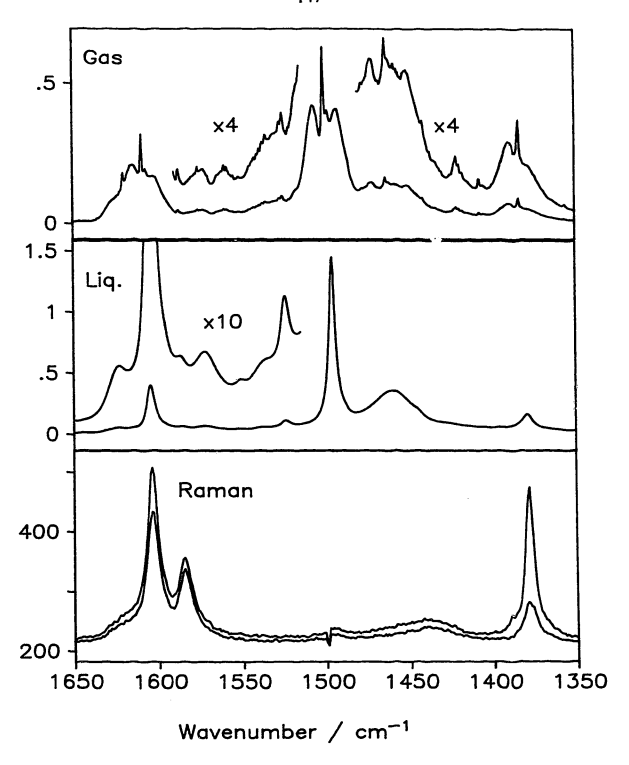


Figure 4.11 - Infrared spectra of gas and liquid toluene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 1650 and 1350 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

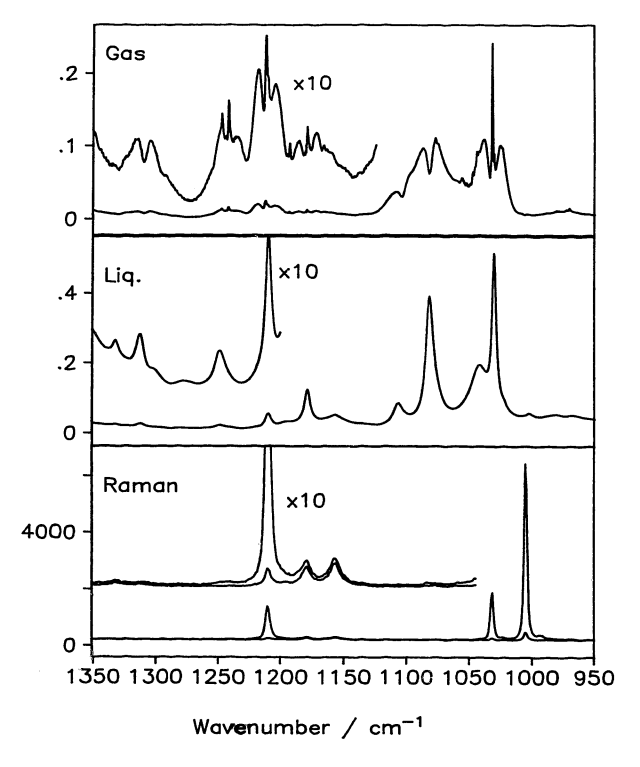


Figure 4.12 - Infrared spectra of gas and liquid toluene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 1350 and 950 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

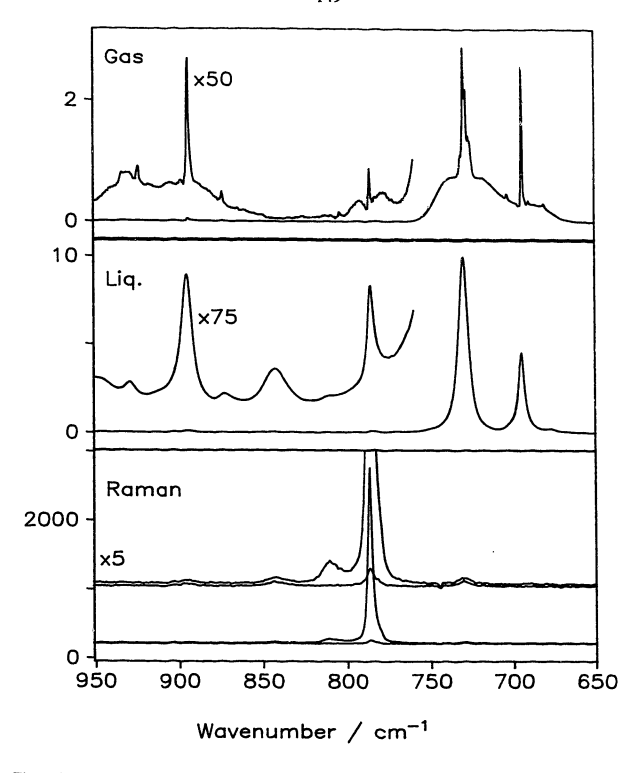


Figure 4.13 - Infrared spectra of gas and liquid toluene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 950 and 650 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

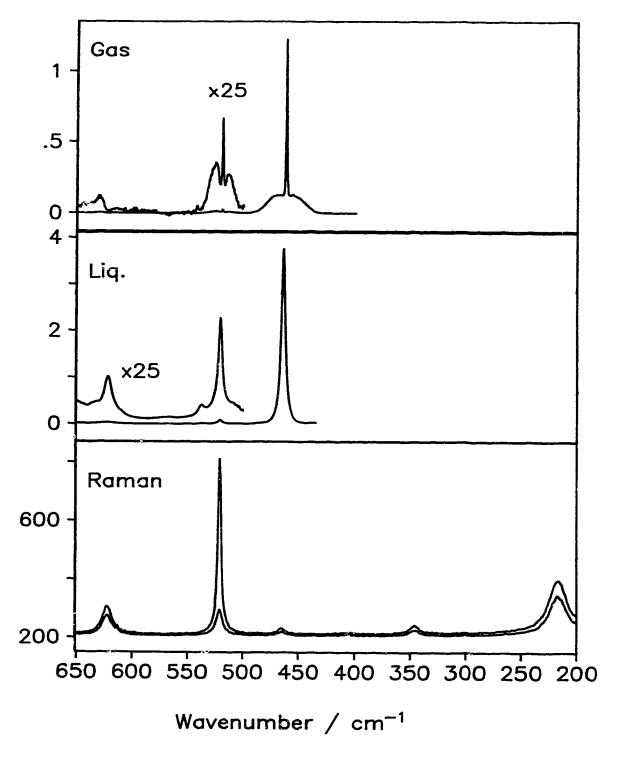


Figure 4.14 - Infrared spectra of gas and liquid toluene and Raman parallel (upper) and perpendicular (lower) spectra of the liquid between 650 and 200 cm⁻¹. In each box, the ordinate scale is for the lower curve. It needs to be divided by the multiplication factor for the upper curve.

Table 4.16 - Spectral features in infrared spectrum of gaseous toluene and infrared and spectra of the liquid

| Ir | nfrared liquid | | Infra | ed gas ^d | Raman liq | uiđ ^b | |
|------------------|----------------|----------------------|------------------|---------------------|------------------|------------------|----------|
| cm ⁻¹ | Desc.b | $a_{\rm m}^{\rm nc}$ | cm ⁻¹ | Contour | cm ⁻¹ | p_i^e | comments |
| ~3203 | vw br sh | ~0.0029 | | | 3205 vw | 0.16 | |
| 3167.5 | vw | 0.00629 | 3175 | ·?•? | 3168 vw | p | |
| ~3115 | w br sh | ~0.026 | ~3120 | 22 | | | |
| 3104.1 | m | 0.0478 | | | | | |
| 3086.4 | s | 0.152 | 3096 | В | | | |
| 3062.1 | s br | 0.177 | 3073 | Α | ~3065 sh | | |
| ~3054 | s br sh? | 0.157 | | | 3055 ms | 0.08 | |
| | | | | | ~3038 vw | dр | |
| 3027.0 | s | 0.398 | ~3040 | B ? | | | |
| ~3003 | s sh | 0.118 | | | 3003 vw | p | |
| | | | 2998 | ?? | | | |
| | | | 2993 | ?? | | | |
| 2979.1 | s br | 0.107 | 2986 | C ?? | 2981 vw | 0.10 | |
| ~2950 | s br | 0.131 | ~2959 | B ??? | | | |
| 2919.9 | s | 0.211 | ~2934 | A ??? | 2920 m | р | |
| 2872.3 | s br | 0.103 | 2881 br | | 2870 br vw | р | |
| 1696.8 | w | 0.0106 | | | | | |
| 1676.7 | w | 0.0121 | | | | | |
| 1657.8 | w br | 0.0124 | | | | | |
| 1623.1 | m br sh/p | 0.056 | 1620 | A? C? | | | |
| 1604.6 | s | 0.402 | 1609 | Α | 1604 m | 0.82 | |
| 1586.7 | m p/sh | 0.0641 | | | 1585 w | dр | |
| 1572.1 | m | 0.0674 | 1576 | C?? | | | |
| 1550.3 | m br sh/p | 0.0452 | ~1560 | ?? | | | |
| ~1537 | m br sh | ~0.06 | | | | | |
| 1523.6 | s br | 0.113 | 1526 | B?? | | | |
| 1495.7 | vs | 1.45 | 1500 | Α | | | |
| 1460.3 | s br | 0.358 | 1467 | B ?? | | | |
| | | | 1463 | C ?? | | | |
| | | | 1454 | 9 ?? | ~1442 br vw | dp | |

Table 4.16 - Continued

| I. | nfrared liquid | * | Infra | red gas ^d | Raman lic | luid ^b | |
|------------------|----------------|----------------------------------|----------------------|----------------------|------------------|---------------------|----------|
| cm ⁻¹ | Desc.b | $a_{\mathrm{m}}^{\prime\primec}$ | cm ⁻¹ | Contour | cm ⁻¹ | ρ_i^{ϵ} | comments |
| ~1422 | m sh | ~0.08 | 1421 | ?? | | | |
| 1378.9 | S | 0.166 | 1384 | Α | 1378 m | 0.33 | |
| 1332.0 | w | 0.0266 | 1330 | B? | 1332 vw | 0.73 | |
| 1312.7 | w | 0.0284 | 1310 | В | 1313 vw | dр | |
| ~1302 | w sh | 0.0184 | | | | | |
| 1277.6 | w br | 0.0150 | | | | | |
| 1248.7 | w | 0.0236 | 1242 | A? | ~1245 vw | p | |
| | | | 1247 | C?? | | | |
| 1210.2 | m | 0.0569 | 1213 | Α | 1210 s | 0.05 | |
| ~1193 | w br sh | 0.0345 | 1193 | A? C? | | | |
| 1178.6 | s | 0.125 | 1179 | A | 1179 vw | 0.70 | |
| 1155.9 | m | 0.0531 | | | 1156 vw | 0.73 | |
| ~1130 | w br sh/p | 0.0272 | | | | | |
| 1106.5 | m | 880.0 | 1104 | E. | | | |
| 1081.4 | s | 0.392 | 1082 | R | 1083 vw | p | |
| 1041.4 | s | 0.196 | ~1043 | ?? | | | |
| 1030.1 | s | 0.516 | 1032 | Α | 1031 s | 0.02 | |
| ~1023 | s sh | 0.102 | | | 1023 vw | p | |
| 1002.3 | m | 0.0581 | | | 1003 vs | 0.02 | |
| | | | | | ~993 w | p | |
| 980.7 | m | 0.0553 | | | | | |
| 966.4 | m br | 0.0519 | 970-5 | ?? | ~971 vw | p | |
| ~947 | w br sh | 0.0211 | | | | | |
| 929.6 | w | 0.0383 | ~924 - 933 | A??C?? | | | |
| ~910 | w br sh? | 0.0107 | 913 | B?? | | | |
| 895.4 | S | 0.120 | 894 | С | 896 vw | 0.77 | |
| 872.9 | w | 0.0297 | 874 | С | | | |
| 842.7 | m | 0.0482 | | | 842 vw | dp | |
| 810.4 | w br sh | 0.0282 | | | 810 vw | p | |
| 785.6 | s | 0.111 | 786 | Α | 786 vs | 0.28 | |

Table 4.16 - Continued

| I | Infrared liquida | | | red gas ^d | Raman lic | quid ^b | |
|------------------|------------------|--------------------------------|--------|----------------------|------------------|-------------------|-------------------------|
| cm ⁻¹ | Desc.b | $a_{\mathrm{m}}^{\mathrm{re}}$ | CPRI-1 | Contour | cm ⁻¹ | ρ_i^e | comments |
| 729.9 | vs | 9.97 | 730 | С | 731 vw | 0.78 | |
| 694.8 | vs | 4.56 | 694 | C | | | |
| ~678 | s br p/sh | 0.219 | ~681 | ?? | | | |
| 633.0 | w sh | 0.0195 | | | | | |
| 622.0 | m | 0.0411 | ~630 | ?? | 622 m-w | 0.76 | |
| 565.2 | vw br | 0.0061 | | | | | |
| 537.8 | w | 0.0166 | | | | | |
| 521.0 | m | 0.091 | 520 | A | 521 m-s | 0.22 | |
| ~508 | vw sh | 0.00342 | | | | | |
| 464.4 | vs | 3.77 | 462 | С | 465 vw | 0.75 | |
| | | | • | | 405 vvw | dp | |
| | | | | | 346 w | 0.73 | |
| | | | | | 217 mw | 0.75 | |
| <100 | | | | | | | CH ₃ torsion |

a - Features are from the α_m'' spectrum.

b - Abbreviations used: v- very, w- weak, m- medium, s- strong, sh- shoulder, br- broad, sh/p- shoulder or a peak.

c - Peak heights in the α_m'' spectrum in cm³ mol⁻¹.

d - Peak positions in the absorbance spectrum of the gas. B-type position is measured at the minimum between the R and P branches. ? and ?? indicate degrees of uncertainty in the determination of the band shape.

e - Depolarization ratio, $\rho_{\ell} = \frac{I_{\perp}}{I_{\parallel}}$. When the feature is too weak or noisy to obtain a reliable ratio, p or dp indicate polarized and depolarized, respectively.

Table 4.17 - Computed wavenumbers, eigenvectors and potential energy distributions for toluene.

| | cm ⁻¹ | | Eigenv | ectors | | PED ⁶ |
|-----------------|------------------|-------------------------|------------------------|------------------------|------------------------|-----------------------------|
| $\mathbf{v_l}$ | 3177 | S ₅ -0.84 | S ₁ -0.59 | | | СН |
| v_2 | 3171 | S ₁ -0.73 | S_{15a} -0.55 | S ₅ 0.42 | | СН |
| V ₃ | 3158 | S_{12a} 0.80 | S_{15a} -0.62 | | | СН |
| V4 | 1597 | S_{18a} 0.80 | S_{16a} 0.42 | $S_{17a} = 0.39$ | S _{14a} -0.27 | CC (75), HCC (25) |
| V5 | 1556 | S _{i-4a} -1.08 | $S_{88a} = 0.41$ | S _{13a} -0.25 | | HCC (50), CC (35), CX (15) |
| V6 | 1314 | S _{14a} -0.73 | $S_{18_{h}}$ 0.55 | S ₆ -0.26 | | HCC (55), CX (45) |
| V 7 | 1157 | S_{18a} 0.89 | $S_{44a} = 0.23$ | | | HCC (60), CC (25), CX (15) |
| V8 | 1030 | S_{14a} 0.50 | S ₆ -0.29 | S_{13a} 0.21 | | CC (60), HCC (25), CCC (15) |
| V 9 | 1001 | S ₆ -0.48 | S ₂ -0.19 | | | CCC (55), CC (45) |
| v_{10} | 824 | S ₆ 0.24 | S_{17a} -0.24 | S ₂ -0.12 | | CCC (40), CC (40), CX (20) |
| v_{11} | 490 | $S_{17a} = 0.29$ | S_{18a} -0.13 | | | CCC (75), CX (25) |
| V12 | 971 | S_{19a} 1.25 | $S_{20a} = 0.44$ | | | oop H |
| v_{13} | 852 | $S_{11a} = 1.03$ | | | | oop H |
| v_{14} | 403 | $S_{20a} = 0.70$ | S ₁₉ 0.26 | | | oop C (85), oop H (15) |
| v_{15} | 987 | S ₇ 1.14 | S _{19b} 0.64 | S ₈ -0.59 | | oop H (85), oop C (15) |
| v_{16} | 916 | S _{19b} -0.85 | S _{11b} -0.66 | $S_7 = 0.49$ | | oop H |
| V17 | 749 | $S_8 	 0.71$ | S ₄ 0.60 | S _{11b} -0.54 | S ₇ -0.45 | oop H (67), oop C (33) |
| v_{18} | 698 | S ₈ 1.04 | S_4 -0.51 | S ₇ -0.41 | | oop C (67), oop H (33) |
| v_{i9} | 489 | S _{20b} 0.63 | S_{19b} 0.39 | S ₄ -0.34 | | oop X (50), oop C (50) |
| V ₂₀ | 260 | S_{20b} 0.30 | $S_4 = 0.14$ | S _{11b} 0.12 | | oop C (65), oop X (35) |
| v_{21} | 3157 | S _{15b} 0.86 | S_{12b} -0.57 | | | СН |
| V22 | 3151 | S _{12b} -0.87 | S _{15b} -0.57 | | | СН |
| V ₂₃ | 1584 | S _{18b} -0.80 | S _{16b} -0.43 | S _{17b} -0.37 | | CC (75), HCC (25) |
| V ₂₄ | 1452 | S _{14b} 1.10 | S ₃ -0.53 | | | HCC (65), CC (35) |
| V ₂₅ | 1330 | $S_3 - 1.09$ | S ₁₀ -0.65 | | | HCC (70), CC (30) |
| V ₂₆ | 1294 | $S_3 0.58$ | S ₁₀ -0.50 | S ₉ 0.40 | S _{18b} -0.34 | CC (75), HCC (25) |
| V27 | 1158 | S_{10} -0.80 | S _{18b} -0.61 | | | HCC (65), CC (35) |
| V_{28} | 1073 | S _{14b} 0.44 | S_{10} 0.39 | S ₁₈₅ -0.38 | S _{13b} 0.22 | CC (55), HCC (45) |
| V29 | 617 | S _{17b} 0.40 | | | | CCC |
| _ <u>V30</u> | 399 | S _{18b} 0.25 | S _{14b} 0.22 | S ₁₀ -0.18 | S ₃ 0.13 | XCC |

a - The numbers are the $\partial S_i/\partial Q_j$ elements of the indicated symmetry coordinate in the particular vibration. Only the dominant terms are shown.

b - CH, CC or CX are stretches, HCC and XCC are bends, CCC is ring deformation. Out-of-plane vibrations are denoted by oop. Percent contributions are given in brackets..

4.5.1 - Assignments of the 11 A1 phenyl group vibrations of toluene

Five aromatic CH stretching fundamentals are expected in toluene: three A_1 and two B_2 vibrations. As was the case for chlorobenzene, they are expected between 3100 and 3050 cm⁻¹.

One obvious A₁ vibration yields the A-type band at 3073 cm⁻¹ in the infrared spectrum of the gas and the corresponding band at 3062 cm⁻¹ in the infrared spectrum of the liquid and the polarized Raman shoulder at about 3065 cm⁻¹. A second A₁ vibration is observed as the polarized Raman band at 3055 cm⁻¹. There is a hint of a shoulder in the infrared spectrum of the liquid and a band centered at 3054 cm⁻¹ was needed to obtain a good curvefit with CDHO bands.

The third A₁ CH vibration is not accounted for. Wilmshurst and Bernstein¹⁵, Varsanyi²³ and Schrotter³² assigned 3003 cm⁻¹. While a polarized Raman band is observed near 3003 cm⁻¹, the calculations suggest that this wavenumber is rather low as all of the aromatic CH stretching vibrations are expected within a 30 cm⁻¹ range.

Overall the band at 3003 cm⁻¹ is probably due to combination or overtone transitions. Fuson et al¹⁷ assigned this fundamental at 3087 cm⁻¹. However the corresponding gasphase band is a B-type, so a B₂ vibration is assigned to it. Draeger²⁵ assigned it to a polarized Raman band at 3039 cm⁻¹ but we find the band to be depolarized (Fig. 4.10).

v₄ is expected near 1600 cm⁻¹ and is assigned to the strong band in the infrared spectrum of the liquid at 1605 cm⁻¹ which corresponds to the A-type band at 1609 cm⁻¹ in the gas. A weak, depolarized Raman band is observed at 1604 cm⁻¹ and this could be

a case where the depolarization ratio of an A₁ band is equal to 0.75.

v₅ is expected near 1550 cm⁻¹ and is assigned as the very strong band in the infrared spectrum of the liquid at 1496 cm⁻¹ which corresponds to the A-type band at 1500 cm⁻¹ in the gas.

The remaining A₁ vibrations are predicted near 1310 cm⁻¹ (v_6), 1160 cm⁻¹ (v_7), 1030 cm⁻¹ (v_8), 1000 cm⁻¹ (v_9), 825 cm⁻¹ (v_{10}) and 490 cm⁻¹ (v_{11}). The assignment of v_8 , v_9 , v_{10} and v_{11} is clearly shown by the polarized Raman bands at 1031, 1003, 786 and 521 cm⁻¹. The assignments are confirmed by A-type bands in the spectrum of the gas at 1030 (s), 786 (vw) and 520 (w) cm⁻¹ and strong or medium bands in the α''_m spectrum of the liquid at 1030, 1002, 786 and 523 cm⁻¹.

The assignment $\frac{1}{4}$ $\frac{1}{4}$ and $\frac{1}{4}$ $\frac{1}{4}$ less clear. They are expected near 1310 and 1160 cm⁻¹ from the approximate potential coordinate calculation. A-type bands are observed in the gas at 1380, possibly 1242, 1213 and 1179 cm⁻¹ (Figs. 4.11 and Polarized Raman bands are at 1378 (m), ~1245 (vvw), 1210 (s) and (possibly polarized) 1179 (vw) cm⁻¹. The corresponding infrared bands of the liquid are at 1379 (s), 1249 (w), 1210 (m) and 1179 (s) cm⁻¹. Based on the infrared spectrum of the gas and the Raman spectrum, the most natural assignment is v_6 at 1379 cm⁻¹ and v_7 at 1210 cm⁻¹. However, the spectrum of $C_6H_5CD_3$ shows¹⁷ that the 1379 cm⁻¹ band is due to the symmetric CH₃ deformation, so v_6 and v_7 are apparently both between 1250 and 1175 cm⁻¹. Accordingly, the assignment of previous authors is accepted, with v_6 at 1210 cm⁻¹ and v_7 at 1179 cm⁻¹.

4.5.2 - Assignments of the 3 A2 phenyl group vibrations of toluene

The three A_2 vibrations are expected n_{eaf} 970, 850 and 400 cm⁻¹. These vibrations are inactive in the infrared of the gas but may appear in the spectrum of the liquid. Usually the intensity in the liquid is very weak but as toluene is not a pure C_N molecule, the A_2 modes may interact with the B_1 modes and their intensity might be stronger. The vibrations give depolarized Raman bands but these are usually very weak for A_2 vibration of C_{2v} molecules.

The assignments of v_{13} and v_{14} were in agreement among previous experimental studies and the present evidence is consistent with the assignments. Thus, the very weak, depolarized Raman band at 842 cm⁻¹, coincident with a medium band in the infrared spectrum, is assigned to v_{13} and the very weak Raman band, probably depolarized, which is observed at about 405 cm⁻¹, is assigned to v_{14} .

Fuson¹⁷, Svendlov²², Varsanyi²³ and Draeger²³ assigned v₁₂ at 967±3 cm⁻¹. While a weak Wilmshurst and Bernstein¹⁵ and Schrotter³² assigned it as 992±2 cm⁻¹. While a weak Raman band is observed at 993 cm⁻¹ it is clearly polarized and can not be assigned to v₁₂, Therefore we assign v₁₂ to the medium, oroad hand at 966 cm⁻¹ in the infrared spectrum of the liquid, which corresponds to a band of a complex contour at 975-970 cm⁻¹ for the gas.

4.5.3 - Assignments of the 6 B₁ phenyl group vibrations of toluene

The computed wavenumbers for the six B_1 vibrations of toluene are: 987 cm⁻¹ for v_{15} , 916 cm⁻¹ for v_{16} , 749 cm⁻¹ for v_{17} , 698 cm⁻¹ for v_{18} , 489 cm⁻¹ for v_{19} and 260 cm⁻¹ for v_{20} . The fundamental bands should be depolarized in the Raman spectrum and should have a C-type contour in the infrared gas spectrum.

Three strong bands at 895, 730 and 464 cm⁻¹ are observed in all experimental spectra with the appropriate contour and polarization and are assigned to v_{16} , v_{17} and v_{19} , respectively. The band at 695 cm⁻¹ in the liquid is very strong and exhibits a C-type contour in the gas spectrum. Although no Raman band is observed nearby, it is assigned to v_{18} .

As the gas spectrum was not recorded below 400 cm⁻¹, the assignment of v_{20} is based on the Raman bands and on the computed wavenumbers. Of the two depolarized Raman bands at 346 and 217 cm⁻¹, v_{20} is assigned to the band at lower wavenumber, in agreement with its computed wavenumber being 140 cm⁻¹ lower than that of v_{30} (B₂).

 v_{15} was previously 15,17,22,23,25,32 assigned at 980 ± 15 cm⁻¹. A band at 981 cm⁻¹ is observed in the infrared spectrum of the liquid. A very weak band of undetermined shape is observed in the gas at about 980 cm⁻¹. The Raman band at 993 cm⁻¹ is clearly polarized. For lack of better evidence, we follow the previous authors and assign v_{15} to 981 cm⁻¹.

4.5.4 - Assignments of the 10 B2 phenyl group vibrations of toluene

The computed wavenumbers of the two B_2 CH stretches are 3157 and 3151 cm⁻¹. Allowance for anharmonicity places the observed wavenumbers about 100 cm⁻¹ lower. In this region, the strongest band in the spectrum of the gas has a B contour at about 3040 cm⁻¹ and corresponds to the strongest band in the spectrum of the liquid, at 3027 cm⁻¹. It is assigned to v_{22} .

A B-type contour is observed at 3096 cm⁻¹ in the infrared spectrum of the gas, corresponding to 3086 cm⁻¹ in the infrared spectrum of the liquid. A band at similar wavenumber for chlorobenzene was assigned to v_{21} although the wavenumber is higher than expected. It is assigned as v_{21} .

v₂₃ is expected near 1585 cm⁻¹. It is assigned to the depolarized Raman band observed at 1587 cm⁻¹, nearly coincident with a medium shoulder in the infrared spectrum of the liquid.

v₂₄ is expected near 1450 cm⁻¹. The B₁ and B₂ antisymmetric CH₃ deformations are also expected in this vicinity. A strong and very broad band at 1460 cm⁻¹ is observed in the infrared spectrum of the liquid (Fig. 4.11). There is a corresponding possible B-type contour at 1467 cm⁻¹ and a possible C-type band at 1463 cm⁻¹ in the spectrum of the gas. Another possible B-type contour is at 1454 cm⁻¹ which corresponds to the very weak and broad depolarized Raman band at about 1442 cm⁻¹. Balfour and Fried³⁰ gave in their figure 1, the transmittance spectra of liquid toluene, toluene-o-d, toluene-m-d, toluene-p-d and toluene-α-d. It is clear from their figure, that

a broad band remains when the isotopic substitution is in the ring, suggesting it is due to the antisymmetric CH₃ deformations. When the isotope substitution is in the methyl group the broad band disappears and a sharp band appears at a slightly lower wavenumber, suggesting it is due to the phenyl B₂ fundamental. Therefore we conclude that the antisymmetric CH₃ deformations are at 1460 cm⁻¹ and v₂₄ is within 20 cm⁻¹ of them, perhaps at 1442 cm⁻¹.

The remaining B₂ vibrations v₂₅, v₂₆, v₂₇, v₂₈, v₂₉ and v₃₀ are expected near 1330, 1300, 1160, 1070, 620 and 400 cm⁻¹. Their assignment is fairly clear from depolarized Raman bands that are coincident with infrared bands of the liquid and, in several cases, B-type bands in the gas. v₃₀ is assigned at 346 cm⁻¹ based solely on the Raman spectrum, while v₂₅, v₂₆, v₂₇, v₂₈ and v₂₉ are assigned at 1332, 1313, 1156, 1081 and 622 cm⁻¹ based on the infrared and Raman spectra of the liquid. B-type bands in the spectrum of the gas at 1330 (vw), 1310 (vw) and 1082 (s) cm⁻¹ support the assignments of v₂₅, v₂₆, v₂₇ and v₂₈. The clear B-type band at 1104 cm⁻¹ (Fig. 4.12) is assigned to a combination band, following Fuson et al¹⁷.

4.5.5 - Assignments of the 9 CH₃ vibrations of toluene

There are three methyl CH stretching vibrations in toluene. They are expected at about 100 cm⁻¹ lower than the aromatic CH stretching vibrations. The symmetric stretch is expected about 50 cm⁻¹ below the two antisymmetric stretching vibrations.

A strong band is of served at 2920 cm⁻¹ in the infrared spectrum of the liquid

about 2934 cm⁻¹ in the spectrum of the gas and is perhaps related. As the Raman and infrared band of the liquid are the strongest bands in the region they are assigned to the A₁ fundamental, the symmetric CH₃ stretching vibration, v₃₁.

The antisymmetric CH stretching vibrations, v_{34} (B₁) and v_{37} (B₂), are expected approximately 50 cm⁻¹ higher than the symmetric stretch. Bands are observed in the spectrum of the liquid at 2950 cm⁻¹ and 2979 cm⁻¹, both broad, and there is a shoulder at ~2965 cm⁻¹. In the spectrum of the gas, a Q branch may occur at 2986 cm⁻¹ (Fig. 4.10) and an even more uncertain B-type band may be observed near 2959 cm⁻¹. The only Raman band in the vicinity is very weak and polarized at 2981 cm⁻¹, so it can not arise from a B₁ or a B₂ vibration. For lack of better evidence we tentatively assign both v_{34} (B₁) and v_{37} (B₁) at 2950 cm⁻¹.

v₃₂ (A₁), the symmetric CH₃ deformation, is assigned to the A-type band at 1384 cm⁻¹ in the spectrum of the B. Dlarized Raman band at 1378 cm⁻¹ and the band at 1379 cm⁻¹ in the infrared spectrum of the liquid. It is clear from the spectra of Balfour and Fried³⁰ that the band remains when the phenyl ring is isotopically substituted, and almost disappears when the isotopic substitution occurs at the methyl group.

As discussed in the previous section, the antisymmetric CH₃ deformations are assigned to the very broad band at 1460 cm⁻¹ in the liquid. In addition to the spectra provided by Balfour³⁰ which indicate that the band is due to a methyl vibration, a complex contour is observed in the spectrum of the gas with a possible B-type band at

1467 cm⁻¹ and a possible C-type band at 1463 cm⁻¹. We are unable to separate the two vibrations and use the same wavenumber for both.

The methyl rocking vibrations, v_{36} (B₁) and v_{39} (B₂) are both assigned to the same wavenumber in the infrared spectrum of the liquid. at 1041 cm⁻¹. It is clear from the spectra of Balfour and Fried³⁰ that the band remains when the ring is isotopically substituted, and disappears when the isotopic substitution occurs at the methyl group. Furthermore, two CDHO bands at 1040 cm⁻¹ and 1043 cm⁻¹ are needed to obtain a good fit of the α_m'' band of the liquid.

v₃₃, the A₂ methyl torsion vibration was not observed in this study. Draeger²⁵ reported an observed and a calculated wavenumber for this vibration, both at 15 cm⁻¹. La Lau and Snyder¹⁹ calculated the wavenumber at 44 cm⁻¹. In their calculations, the authors used a small value (1x10⁻⁴ mdyne-Å/rad) so that their calculated wavenumber is low enough to prevent it from coupling with other vibrations. Hameka and Jensen³⁴, based on La Lau and Snyder data, calculated it at 28.9 cm⁻¹. It is clear that the wavenumber for this vibration is well below 100 cm⁻¹.

4.5.6 - Summary of the assignments of toluene

The fundamental wavenumbers are collected in Table 4.18 which corresponds to Table 4.10. The table includes the pseudo-Herzberg assignments obtained from Table 4.17 by including all symmetry coordinates with eigenvectors at least half of the largest. It is clear that the pseudo-Herzberg assignment is complicated for most of the vibrations.

Table 4.18 - Assignments of the fundamental vibrations of liquid toluene

| true Herzberg | cm ^{-1 a} | pseudo-Herzberg ^b | Description ^e |
|------------------------------------|--------------------|--|---------------------------------------|
| \mathbf{v}_1 | 3062.1 | v ₅ , v ₁ | СН |
| \mathbf{v}_2 | 3055.0 | V ₁ , V _{15a} , V ₅ | СН |
| \mathbf{v}_3 | 3038 ??? | V _{12a} , V _{15a} | СН |
| V4 | 1604.6 | $V_{18a}, V_{16a}, V_{17a}$ | HCC, CC |
| v_5 | 1495.7 | V_{14a} | нсс |
| v_6 | 1210.2 | V_{14a} , V_{18a} | HCC |
| \mathbf{v}_7 | 1178.6 | v_{18a} | HCC |
| v_8 | 1030.1 | v_{14a}, v_6 | HCC, LLU |
| \mathbf{v}_{9} | 1002.3 | v_6 | CC4. |
| \mathbf{v}_{10} | 785.6 | v_6 , v_{17a} , v_2 | CCC, CC |
| \mathbf{v}_{11} | 521.0 | v_{17a} | CCC, CC |
| v_{12} | 966.4 ?? | v_{19a} | ούρ Η |
| $\mathbf{v_{13}}$ | 842.7 | v_{i1a} | оор Н |
| V ₁₄ | ~405? | V _{20a} | оор С |
| \mathbf{v}_{15} | 980.7 ?? | v_7, v_{19b}, v_8 | оор Н, оор С |
| \mathbf{v}_{16} | 895.4 | v_{19b} , v_{11b} , v_7 | оор Н, оор С |
| \mathbf{v}_{17} | 729.9 | v ₈ , v ₄ , v _{11b} , v ₇ | oop C, oop H |
| $ u_{18}$ | 694.8 | V ₈ , V ₄ | oop C, oop H |
| \mathbf{v}_{19} | 464.4 | V _{20b} , V _{19b} , V ₄ | oop C, oop H |
| \mathbf{v}_{20} | 217 | V _{20b} , V ₄ | oop C, oop H |
| v_{21} | 3086.4 | V _{15b} , V _{i2b} | СН |
| v_{22} | 3027.0 | V _{12b} , V _{15b} | СН |
| v_{23} | 1586.7 | ν _{18b} , ν _{16b} | HCC, CC, CCC |
| V ₂₄ | ~ 1442 ?? | v_{14b} | HCC, CC |
| V ₂₅ | 1332.0 | v_3 , v_{10} | нсс |
| V ₂₆ | 1312.7 | $v_3, v_{10}, v_9, v_{18b}$ | HCC, CC |
| \mathbf{v}_{27} | 1155.9 | v_{10}, v_{18b} | нсс |
| v_{28} | 1081.4 | V _{14b} , V ₁₀ , V _{18b} , V _{13b} | HCC, CC |
| \mathbf{v}_{29} | 622.0 | V _{17b} | CCC |
| v_{30} | 346 | V_{18b} , V_{14b} , V_{10} , V_3 | HCC, CC |
| v ₃₁ (A ₁). | 2919.9 | | symmetric CH ₃ stretch |
| y32 (A1) | 1378.9 | | symmetric CH ₃ deformation |

Table 4.18 - Continued

| true Herzberg | cm ^{-1 a} | pseudo-Herzberg ^b | Description ^c |
|-----------------------------------|--------------------|------------------------------|---|
| $y_{33}(A_2)$ | < 100 | | CH ₃ torsion |
| $v_{34}(B_1)$ | ~2950 ?? | | antisymmetric CH ₄ stretch |
| $v_{35}(B_1)$ | 1460.3 ? | | antisymmetric CH ₃ deformation |
| $v_{36}(B_1)$ | 1041.4 ? | | CH ₃ rock |
| $v_{37}(B_2)$ | ~2950 ?? | | antisymmetric CH ₃ stretch |
| $v_{38}(B_2)$ | 1460.3 ? | | antisymmetric CH ₃ deformation |
| v ₃₉ (B ₂) | 1041.4 ? | | CH ₃ rock |

a - Peak wavenumbers of the imaginary molar polarizability spectrum of liquid toluene.

Note that v_{13a} , a C-C stretching displacement, does not appear in Table 4.18. It is a minor contributor to several vibrations.

A question arises from the table, where are the C-CH₃ stretch and the C-CH₃ inplane and out-of-plane bends?

From the unsymmetrized eigenvectors, the C-CH₃ stretch contributes with HCC, CC and CCC to the following A₁ vibrations: v_4 (1597), v_5 (1556), v_6 (1314), v_7 (1157), v_{10} (824) and v_{11} (490). It does not make the major contribution to any of them. From the potential energy distribution, the C-CH₃ stretch is among the three main contributors to v_5 (1556), v_6 (1314), v_7 (1157), v_{10} (824) and v_{11} (490) but is the major contributor to none of these. Therefore we conclude that no single vibration can be called to the C-CH₃ stretch.

The unsymmetrized eigenvectors and the PED agree that the CC- CH₃ in-plane

b - Contributions greater than 50% of the largest contribution.

c - CH, CC are stretches, HCC is CH bend, CCC is ring deformation and oop is out-of-plane.

bending displacement is the dominant contributor to v₃₀, calculated at 399 cm⁻¹ and observed at 346 cm⁻¹ and is, therefore, the CC- CH₃ in-plane bend.

The unsymmetrized eigenvectors and the PED show that the out-of-plane CC-CH₃ bending displacement mixes with the CCC out-of-plane displacement to form v_{19} and v_{20} . v_{19} is about an equal mixture of the two while the CCC deformation dominates by a factor of 2 in v_{20} .

Chapter 5 - Infrared Integrated Intensities for Liquid Chlorobenzene and Toluene

In Chapter 4, the fundamental vibrations of chlorobenzene and toluene were assigned. While it is possible to determine infrared intensities without a complete assignment, it is far more meaningful to determine transition moments and dipole moment derivatives with some knowledge of the nature of the transition.

Bertie *et al.*⁴⁸ have shown that if the Lorentz local field is valid, molecular properties and behavior are more directly reflected in the imaginary molar polarizability spectrum, α_m'' , than in the imaginary refractive index spectrum, k, the molar absorption coefficient spectrum, E_m , or the imaginary dielectric constant spectrum, ε'' .

Accordingly, the refractive indices of toluene and chlorobenzene, given in Chapters 2 and 3 respectively, were used to calculate the imaginary molar polarizability spectra of toluene and chlorobenzene using Eqs. 1.3.28a, 1.3.28b and 1.3.33b. The imaginary molar polarizability spectra are shown in Figs. 5.1 and 5.9 and the integrated intensities of chlorobenzene and toluene are presented in Sections 5.1 and 5.2, respectively.

In order to calculate the integrated intensities, the α_m''' spectrum must be separated into contributions from the different transitions. The integration limits can be chosen arbitrarily so that the integrated intensity is taken as the area between these limits and is assumed to be solely due to a single specific transition. This approach can give good results in optimal circumstances, namely when the band is single, well defined, and fairly isolated from other bands. But even for such a band, the choice of integration limits can introduce large uncertainty to the intensity. To obtain 98% of the area under a

Lorentzian band one must integrate over 16 times the full-width at half-height on each side of the band. Further, when, as is usual, bands are close or actually overlap, any arbitrary separation could introduce considerable error into the integrated intensity value.

The alternative method is to fit the α_m'' spectrum with bands of appropriate and well-defined shape, and calculate the integrated intensities from the areas under the fitted bands. As discussed in Section 1.4, the general theory of infrared spectra of liquids⁴⁸, indicates that the bands in the α_m'' spectrum have the shapes of classical damped harmonic oscillator (CDHO) bands. Thus, the α_m'' spectra of chlorobenzene and toluene were fitted with CDHO bands.

The integrated intensities, C_j , (Eqs. 1.4.2 to 1.4.4) are the areas under the $\tilde{\nu}\alpha_m''$ spectra of the bands that fitted the original α_m'' spectra. The j^{th} CDHO band in the imaginary molar polarizability is described by

$$\alpha_{\rm m}''(\widetilde{\nu}) = \frac{S_j \, \Gamma_j \, \widetilde{\nu}_j}{(\widetilde{\nu}_i^2 - \widetilde{\nu}^2)^2 + \Gamma_i^2 \, \widetilde{\nu}_i^2} \tag{5.1.1}$$

Program CURVEFIT yields the parameters, S_j , Γ_j and $\widetilde{\nu}_j$. The area C_j under the corresponding $\widetilde{\nu}\alpha_m''$ band is given by $C_j = S_j \pi/2$. Provided a good fit can be obtained, the uncertainty introduced by the choice of integration limits is, then, eliminated as the limits are effectively zero and infinity.

There are, however, two major disadvantages to curve fitting. First, the fit can be artificially improved by the addition of extra bands. Some of these extra bands may not be actually present in the original spectrum and they will affect the integrated

addition of extra bands is limited and can be justified. In the curve fitting performed in this work, the minimum number of bands was used to obtain a good fit. Extra bands were added only when the observed band was clearly asymmetric. With the exception of two or three cases, all bands used for the fit could be justified based on band asymmetry or a distinct shoulder observed in the original spectrum.

Second, the fit within a region is heavily influenced by bands that are outside the region. The best solution for this problem would be to fit the entire spectrum. This solution, however, is often in conflict with available computer programs for curve fitting. These programs usually require large memory space and execution times increase manyfold with increase of the number of bands to be fitted. Therefore, the spectrum is usually divided into several regions and a fit is performed for each region. If the curve fitting procedure is done properly, the contributions from adjacent regions are included and taken into account during the fit.

Galactic Industries' program CURVEFIT was modified in our laboratory by C.D. Keefe and S.L. Zhang to use CDHO bands. It was also written in FORTRAN and compiled with Microway's NDP FORTRAN under OS/2 2.1 in order to provide access to 16 Megabyte of memory and to obtain a tenfold decrease in execution time⁷³. The program was limited to 80 peaks in the fitted spectrum and the input spectrum was limited to 8192 points.

For benzene⁷³, chlorobenzene and toluene, each with well over 150 bands of various intensity in the spectrum, 80 peaks are not enough and the spectrum must be divided into two or three regions. Furthermore, at the start of this work, some of the 80 peaks, had to be reserved for bands outside the region that contributed more than 10⁻⁶ cm³ mole⁻¹ to the fit inside the region. For benzene⁷³, the number of regions had to be increased to 10 to 15 as the number of required outside peaks increased as more regions were fitted. The fit in each region was refined iteratively until the bands in the entire spectrum were constant.

Thus, a serious drawback to this FORTRAN program at the start of this work was the need to include among the 80 peaks that would be refined an increasingly large number of peaks outside the region being fitted. An improvement was made for the work of this thesis.

In the improved program, once a region has been fitted, its bands are considered constant for the adjacent region. Therefore the contribution can be calculated first and then subtracted from the region of the spectrum to be fitted. This resulting region is then fitted with bands within the region. This allows all of the 80 peaks that can be refined by the program to be within the region. Each region is fitted in this way, then, the whole procedure is repeated several times, each time calculating the initial contribution from the last fit of the adjacent regions. Refinement of the fit is continued until all bands in all regions remain constant. The new program allows 80 bands within the region to be fitted and 200 constant bands outside of the region to be included. This

allows larger regions to be fitted, and, to accommodate this, the number of points in the input spectrum was increased to 16384.

With this revised program only three regions were needed to fit an entire spectrum and, with the inclusion of all contributions from all of the outside bands, only three to four cycles of refinement were needed to fit the α_m'' spectra of chlorobenzene and toluene.

5.1 - Integrated intensities and dipole moment derivatives of liquid chlorobenzene

The imaginary molar polarizability, α_m'' , spectrum of liquid chlorobenzene was fitted with 193 CDHO bands between 4475 and 350 cm⁻¹. An entire spectrum, called the fitted spectrum in this thesis, was obtained by adding the 193 bands, each of which extended from 4475 to 350 cm⁻¹. This fitted spectrum and the experimental α_m'' spectrum are shown in Figure 5.1.

The fit is very good and the two spectra nearly coincide. As can be seen from the magnified curves in Fig 5.1, the fit is not as good in the wings of sharp bands as elsewhere. The CDHO band does not fit the observed bandshape very well, and, the addition of extra CDHO bands did not improve the fit in these wings, and, consequently these extra bands were not used in the final fit.

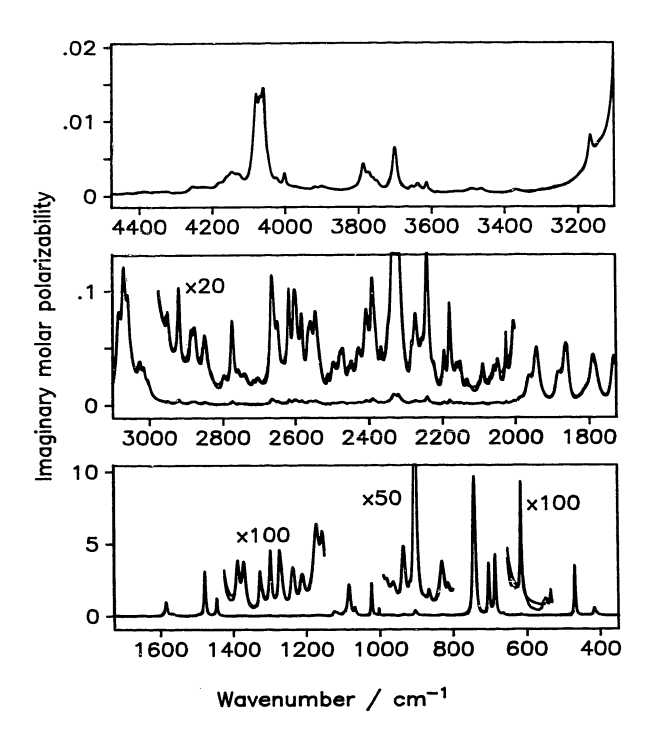


Figure 5.1 - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid chlorobenzene. Units are cm³ mole⁻¹. In each box, the ordinate scale is for the lower curves. The labels must be divided by the multiplication factor for the upper curves.

The overall average percent difference between the fitted and experimental $\alpha_{\rm m}^{\prime\prime\prime}$ peak heights is 2.5%. The difference is on average greater for weaker peaks and decreases for stronger peaks. For peak heights less than < 0.01 cm³ mole⁻¹, between 0.01 and 0.1 cm³ mole⁻¹, between 0.1 and 1 cm³ mole⁻¹, and > 1 cm³ mole⁻¹, the average percent differences are ~3%, 2.8%, 1.7% and 0.6%, respectively. Thus, the peak heights are fitted to 0.6% on average for the peaks that are usually observed in an infrared spectrum that is recorded to show the peak maxima.

An important check on the quality of the fit is to compare the total area under the experimental spectrum to that under the fitted spectrum over a wide wavenumber range. Between 4475 and 800 cm⁻¹, the area under the fitted spectrum is greater by 0.45% and between 800 and 500 cm⁻¹, it is greater by 2.6%. Overall, the area under the fitted spectrum is 1.7% greater than that under the experimental α_m'' spectrum.

The parameters of the fitted CDHO bands are given in Table 5.1. In this table, the peak positions in the α''_m spectrum of the liquid and the band symmetry of the spectrum of the gas are given in the first two columns. The fitted peak positions are given in column 3, and the CDHO parameter Γ_j , which is essentially the full width at half height (FWHH) of the band, is given in column 4. It is followed by the integrated intensity, C_j , in km mole⁻¹ in column 5 and by the dipole transition moment, $|\overrightarrow{R_j}|$, in Debye in column 6. The C_j values were calculated using Eq. 1.4.2 while the $|\overrightarrow{R_j}|$ values were calculated using Eq. 1.4.3.

Table 5.1 - Integrated intensities and dipole moment transitions of liquid chlorobenzene.

| Exp $a_m^{"}$ ab | Gasc | | Fitted | Γ_{j} | С, | $ \overrightarrow{R_j} $ | Assignment ^d |
|------------------|------|---------|------------------|------------------|----------|--------------------------|---|
| cm ⁻¹ | ··· | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| 4468.1 vw | | | 4469.5 | 26.5 | 0.000225 | 0.00126 | |
| 4448.9 vw | | | 4449.1 | 15.1 | 0.000091 | 0.00080 | |
| | | | 4436.8 | 11.2 | 0.000054 | 0.00062 | |
| 4432.5 vw | | { | 4432.1 | 8.7 | 0.000045 | 0.00057 | |
| | | | 4426.8 | 34.3 | 0.000367 | 0.00162 | |
| 4404.1 vw | | | 4405.7 | 14.1 | 0.000141 | 0.00100 | v ₂₁ +v ₂₅ (4408) |
| 4387.4 vw br | | | 4384.9 | 36.4 | 0.00106 | 0.00276 | v_1+v_{25} (4395) or v_3+v_{25} (4385) |
| 4359.2 vw | | | 4358.4 | 20.4 | 0.000272 | 0.00140 | v ₂₁ +v ₂₆ (4356) |
| 4340.8 vw | | | 4341.0 | 23.7 | 0.000318 | 0.00152 | $v_1 + v_{26}$ (4341) |
| 4323.9 vw br | | | 4323.2 | 26.0 | 0.000610 | 0.00211 | $v_3 + v_{26}$ (4331) or $v_{22} + v_{26}$ (4332) |
| ~4295 vw br | | | 4295.3 | 16.9 | 0.000078 | 0.00076 | |
| 4252.8 vw br | | <u></u> | 4254.5 | 18.2 | 0.000534 | 0.00199 | v ₂₁ +v ₆ (4255) |
| 42.J2.G VW UI | | l | 4250.8 | 44.7 | 0.000698 | 0.00227 | |
| 4226.8 vw br | | | 4223.6 | 59.4 | 0.00306 | 0.00478 | |
| 4178.6 vw | | | 4180.8 | 21.6 | 0.00109 | 0.00287 | |
| ~4161 vw sh | | | 4161.3 | 21.5 | 0.00126 | 0.00309 | |
| 4145.8 vw br | | | 4145.8 | 26.7 | 0.00321 | 0.00494 | $v_3 + v_7$ (4142) or $v_{22} + v_7$ (4144) |
| | | | | | | | or $v_{21}+v_{28}$ (4151) |
| 4131.4 vw br | | _ | 4126.2 | 26.4 | 0.00257 | 0.00443 | |
| 4077 4 | В | { | 4082.6 | 20.4 | 0.00651 | 0.00709 | |
| 4077.4 w | Ð | l | 4077.5 | 11.5 | 0.00433 | 0.00578 | $v_{22}+v_8$ (4083) or $v_{21}+v_9$ (4085) |
| 4066.5 w | Α | | 4067.1 | 15.0 | 0.00701 | 0.00737 | v_1+v_9 (4071) |
| 4058.7 w | В | | 4057.8 | 11.2 | 0.00647 | 0.00709 | v ₂₂ +v ₉ (4062) |
| ~4045 vw sh | | | 4045.5 | 18.4 | 0.00205 | 0.00400 | |
| 4022.8 vw br | | | 4021.9 | 19.6 | 0.00121 | 0.00308 | $v_{22}+v_{12}$ (4024) or v_3+v_{12} (4023) |
| 4000.5 vw | | | 4000.3 | 8.4 | 0.000921 | 0.00269 | 4v ₉ (4008) |
| 3968.9 vw br | | | 3977.5 | 70.8 | 0.00320 | 0.00503 | $v_1 + v_{16}$ (3973) |
| ~3950 vw sh | | | | | | | |
| 3938.4 vw | | | | | | | |
| 3917.7 vw | | | 3917.8 | 11.3 | 0.000189 | 0.00123 | v ₂₁ +v ₁₃ (3913) |

Table 5.1 - Continued

| Exp $\alpha_{\rm m}^{\prime\prime}$ a,b | Gas ^c | Fitted | Γ_j | C_{j} | $ \overrightarrow{R_i} $ | Assignment ^d |
|---|------------------|------------------|------------------|----------|--------------------------|--|
| cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| 3899.5 vw br | | 3899.5 | 10.8 | 0.000148 | 0,00109 | v ₁ +v ₁₃ (3899) |
| | | 3894.3 | 75.0 | 0.00303 | 0.00495 | $v_{21}+v_{13}$ (3890) or v_3+v_{13} (3889) |
| ~3878 vw sh | | | | | | |
| ~3867 vw sh | | | | | | |
| ~3804 vw sh | | 3796.6 | 73.1 | 0.00279 | 0.00481 | $v_{1,2}+v_{17}$ (3810) |
| 3785.1 vw | | 3785.2 | 15.3 | 0.00276 | 0.00479 | $v_{21}+v_{10}$ (3786) |
| 3772.2 vw br | | 3772.9 | 12.5 | 0.000331 | 0.00166 | $v_1 + v_{10}$ (3772) |
| | | 3768.3 | 8.9 | 0.000797 | 0.00258 | $v_{21}+v_{18}$ (3768) |
| ~3759 vw sh | | 3759.1 | 10.4 | 0.000421 | 0.00188 | |
| 3749.3 vw br | | 3747.8 | 17.0 | 0.000901 | 0.00275 | |
| 3699.4 vw | | 3699.5 | 16.6 | 0.00572 | 0.00698 | $v_{21}+v_{29}$ (3697) |
| ~3685 vw sh | | 3685.5 | 191.7 | 0.00178 | 0.00390 | $v_1 + v_{29}$ (3684) |
| 3653.4 vw br | | 3654.3 | 12.0 | 0.000356 | 0.00175 | |
| 3636.9 vw | | 3636.8 | 16.8 | 0.00102 | 0.00297 | |
| ~3628 vw sh | | | | | | |
| 3613.6 vw | | 3613.3 | 9.0 | 0.000613 | 0.00231 | 4v ₁₆ (3612) |
| ~3602 vw sh | | 3601.8 | 10.9 | 0.000091 | 0.00089 | |
| 3583.9 vw | | 3583.7 | 5.3 | 0.000031 | 0.00052 | |
| 3546.9 vw br | | 3549.0 | 16.0 | 0.000089 | 0.00089 | $v_{21}+v_{19}$ (3551) |
| 3531.1 vw | | 3531.5 | 27.2 | 0.000254 | 0.00151 | v_1+v_{19} (3538) or v_3+v_{19} (3527) or ? $v_{22}+v_{19}$ (3528) |
| 3488.8 vw | | 3490.7 | 33.9 | 0.00109 | 0.00314 | |
| 3464.1 vw | | 3463.3 | 18.6 | 0.000463 | 0.00205 | |
| ~3445 vw sh | | 3446.1 | 9.9 | 0.000025 | 0.00048 | |
| 3420.4 vw | | | | | | |
| 3368.4 vw br | | 3369.7 | 17.2 | 0.000325 | 0.00174 | $v_1 + v_{30}$ (3367) |
| 3297.5 vw br | | | | | | |
| ~3261 vw sh | | | | | | $v_1 + v_{20}$ (3265) or $v_3 + v_{20}$ (3254) |
| 3165.4 vw br | | 3165.9 | 11.7 | 0.00183 | 0.00427 | 2v ₂₃ (3168) or 2v ₄ (3168) |
| | | | | | | or v ₂₃ +v ₄ (3168) |
| ~3140 vw sh | | 3144.3 | 89.5 | 0.0153 | 0.0124 | |

Table 5.1 - Continued

| Exp $\alpha_m^{\prime\prime}$ a,b | Gasc | | Fitted | Γ_{j} | C, | $ \overrightarrow{R_j} $ | Assignment ^d |
|-----------------------------------|------|---------|------------------|------------------|----------|--------------------------|--|
| cm ⁻¹ | | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| 3083.1 m | В | | 3084.0 | 12.5 | 0.0319 | 0.0181 | v_{2l} |
| 3069.5 s | Α | | 3069.5 | 12.4 | 0.0482 | 0.0222 | $v_1, v_2 ??$ |
| 3058.5 m br | A?? | | 3057.9 | 9.0 | 0.0163 | 0.0130 | v_3 (3059), v_{22} (3060??) |
| 5056.5 III 01 | | l | 3056.3 | 40.9 | 0.0703 | 0.0269 | $\int_{0}^{1} \sqrt{3} \left(3039\right), \sqrt{22} \left(3000\right) \left(1\right)$ |
| 3025.8 w | | | 3025.8 | 10.6 | 0.00962 | 0.0100 | $v_4 + v_{24}$ (3029) or $v_{23} + v_{24}$ (3029) |
| 3016.3 w | | | 3015.7 | 9.1 | 0.00633 | 0.00813 | |
| ~3004 w sh | | | 3003.7 | 19.5 | 0.0112 | 0.0108 | 3v ₉ (3006) |
| 2949.8 vw | | | 2949.9 | 12.9 | 0.000802 | 0.00293 | 2v ₅ (2956) |
| 2919.0 vw | В | | 2918.8 | 7.9 | 0.00121 | 0.00361 | v ₅ +v ₂₄ (2923) |
| ~2906 vw sh | | | | | | | $v_4 + v_{25}$ (2909) or $v_{23} + v_{25}$ (2909) |
| 2885.6 vw db? | Α | | 2882.8 | 20.1 | 0.00162 | 0.00421 | 2v ₂₄ (2891) |
| 2877.0 vw db? | | | 2874.9 | 9.0 | 0.000433 | 0.00218 | |
| 2849.2 vw | | | 2847.7 | 18.6 | 0.00166 | 0.00429 | $v_4 + v_{26}$ (2856) or $v_{23} + v_{26}$ (2856) |
| ~2831 vw sh | | | | | | | |
| 2795.2 vw br | | | 2795.6 | 12.7 | 0.000275 | 0.00176 | v ₅ +v ₂₅ (2803) ?? |
| 2772.9 vw | | | 2772.7 | 9.9 | 0.00127 | 0.00380 | $v_{24}+v_{25}$ (2771) |
| 2755.9 vw | | | 2755.9 | 14.0 | 0.000348 | 0.00199 | $v_4 + v_6$ (2756) or $v_{23} + v_6$ (2756) |
| 2738.9 vw br | ** | | 2738.8 | 21.1 | 0.000607 | 0.00264 | $v_4 + v_{27}$ (2740) or $v_{23} + v_{27}$ (2740) |
| 2714.2 vw | | | 2714.7 | 9.0 | 0.000099 | 0.00107 | $v_{24}+v_{26}$ (2718) |
| 2703.1 vw br | | | 2701.9 | 16.7 | 0.000340 | 0.00630 | 3v ₁₆ (2709) |
| 2663.5 vw | | \[| 2664.7 | 10.1 | 0.00146 | 0.00415 | $v_4 + v_7$ (2668) or $v_{23} + v_7$ (2668) |
| 2003.3 VV | | l | 2659.4 | 12.6 | 0.00108 | 0.00305 | |
| 2648.1 vw | | | 2647.7 | 10.5 | 0.000838 | 0.00316 | $v_5 + v_6$ (2649) or $2v_{25}$ (2650) |
| ~2632 vw sh | | | 2625.8 | 57.8 | 0.00185 | 0.00471 | v ₅ +v ₂₇ (2634) |
| 2616.2 vw | | | 2616.4 | 7.9 | 0.00111 | 0.00366 | v ₂₄ +v ₆ (2517) |
| 2599.8 vw br | | | 2602.3 | 7.7 | 0.000576 | 0.00264 | v ₂₄ +v ₂₇ (2602) |
| 2377.6 VW UI | | Ì | 2597.5 | 11.1 | 0.00119 | 0.00380 | $v_{25}+v_{26}$ (2598) |
| | | | 2590.1 | 21.0 | 0.000542 | 0.00257 | |
| 2582.2 vw | | | 2581.9 | 7.1 | 0.000648 | 0.00281 | $v_{23}+v_9$ (2586) or v_4+v_9 (2586) |
| ~2563 vw sh | | | 2564.9 | 9.7 | 0.000461 | 0.00238 | |
| 2557.4 vw | | | 2557.3 | 14.1 | 0.00122 | 0.00388 | |

Table 5.1 - Continued

| Exp $\alpha_m^{"a,b}$ | Gas ^c | | Fitted | I_j^r | C, | $\overrightarrow{R_i}$ | Assignment ^d |
|-----------------------|------------------|---|------------------|------------------|----------|------------------------|--|
| cm ⁻¹ | | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| 2544.3 vw asym | | | 2544.1 | 11.4 | 0.00116 | 0,00379 | |
| | | | 2535.7 | 16.7 | 0.000604 | 0.00274 | |
| 2510.9 vw | | | 2511.0 | 4.1 | 0.000064 | 0.00090 | v ₂₄ +v ₂₈ (2514) |
| 2496.0 vw | | | 2495.5 | 16.2 | 0.000709 | 0.00299 | v ₂₅ +v ₆ (2497) or v ₅ +v ₈ (2501) |
| 2477.5 vw br | | | 2477.6 | 14.5 | 0.000656 | 0.00289 | v ₅ +v ₉ (2480) or v ₂₅ +v ₂₇ (2482) |
| 2471.3 vw | | | 2469.7 | 10.7 | 0.000494 | 0.00251 | v ₂₄ +v ₈ (2468) |
| ~2453 vw sh | | | 2453.9 | 10.5 | 0.000215 | 0.00166 | |
| 2446.6 vw | | | 2446.6 | 8.5 | 0.000246 | 0.00178 | $v_{24}+v_9$ (2447) or $v_{26}+v_6$ (2444) |
| 2427.3 vw br | | | 2428.0 | 18.2 | 0.00110 | 0.00378 | v ₂₆ +v ₂₁ (2429) |
| 2406.4 vw | | | 2406.3 | 14.3 | 0.00162 | 0.00461 | |
| 2390.3 vw asym | | | 2390.3 | 9.9 | 0.00133 | 0.00419 | $v_{25}+v_{28}$ (2393) |
| | | | 2382.9 | 14.2 | 0.000877 | 0.00340 | $v_5 + v_{16}$ (2381) |
| 2365.6 vw | | | 2365.7 | 9.7 | 0.000385 | 0.00226 | |
| ~2346 vw sh | | | 2347.7 | 12.7 | 0.000475 | 0.00252 | $v_{25}+v_8$ (2348) or $v_{24}+v_{16}$ (2349) |
| 2331.0 vw | | | 2331.8 | 15.6 | 0.00428 | 0.00760 | |
| 2319.8 vw | | | 2319.1 | 12.1 | 0.00273 | 0.00609 | |
| | | | 2311.3 | 12.1 | 0.000724 | 0.00314 | |
| ~2282 vw sh | | | 2283.1 | 6.0 | 0.000182 | 0.00158 | |
| 2271.8 vw | | | 2271.8 | 15.1 | 0.00159 | 0.00470 | |
| ~2255 vw sh | | | 2255.6 | 13.3 | 0.000488 | 0.00261 | v ₆ +v ₇ (2255) or v ₂₆ +v ₁₅ (2256) |
| 2240.5 vw | B ? | | 2240.3 | 11.5 | 0.00244 | 0.00586 | v ₆ +v ₂₈ (2240) or v ₂₇ +v ₇ (2240) |
| ~2223 vw sh | | | 2221.9 | 10.6 | 0.000251 | 0.00189 | $3v_{17}$ (2224) of $v_{27}+v_{28}$ (2225) |
| 2194.0 vw | | | 2194.3 | 8.1 | 0.000423 | 0.00246 | v ₆ +w ₈ (2194) |
| 2178.7 vw | | ſ | 2179.0 | 6.8 | 0.000712 | 0.00321 | |
| 2170.7 *** | | J | 2176.1 | 7.8 | 0.000240 | 0.00186 | |
| ~2167 vw sh | | | 2167.5 | 12.3 | 0.000151 | 0.00148 | 2v; (2167) |
| 2159.7 vw | | | 2159.3 | 9.3 | 0.000236 | 0.00186 | v ₂₇ +v ₉ (2158) |
| 2150.5 vw | | | 2150.1 | 9.6 | 0.000334 | 0.00221 | |
| 2132.5 vw | | | 2131.3 | 9.9 | 0.000128 | 0.00138 | 2v ₂₈ (2136) ? |
| ~2123 vw sh | | | | | | | $v_{27} + v_{12}$ (2120) |
| ~2100 vw sh | | | 2098.7 | 3.9 | 0.000013 | 0 00044 | $v_{16}+v_{13}$ (2103) |

Table 5.1 - Continued

| Exp α_m^* ab | Gas | | Fitted | Γ, | С, | $ \overrightarrow{R_i} $ | Assignment ^d |
|---------------------|-----|----------|------------------|------------------|----------|--------------------------|--|
| cm ⁻¹ | | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| 2089.1 vw | | | 2088.6 | 9.4 | 0.000306 | 0.00215 | v_5+v_{29} (2092) or $v_{28}+v_8$ (2091) or v_7+v_9 (2086) |
| ~2067 vw sh | | | 2067.4 | 7.1 | 0.000044 | 0.00082 | $v_{28}+v_9$ (2070) or v_7+v_{15} (2067) or $v_{25}+v_{17}$ (2066) |
| 205@.2 vw | | | 2058.0 | 7.9 | 0.000152 | 0.00153 | $v_{27}+v_{16}$ (2059) or $v_{24}+v_{29}$ (2060) |
| 2045.5 vw db? | | | 2047.8 | 7.8 | 0.000206 | 0.00178 | |
| 2023.7 vw | | | 2023.7 | 3.1 | 0.000159 | 0.00157 | $v_8 + v_9$ (2025) or $v_{25} + v_{10}$ (2028) |
| 2002.5 vw br | | | 2003.3 | 6.1 | 0.000218 | 0.00185 | $2v_9$ (2004) or v_8+v_{15} (2006) or ? v_6+v_{13} (2002) |
| ~1989 vw sh | | | 1989.2 | 5.3 | 0.000314 | 0.00223 | |
| 1983.0 vw | | | 1984.8 | 3.6 | 0.000151 | 0.00155 | v_9+v_{15} (1985) or $v_{23}+v_{14}$ (1984) or ? v_4+v_{14} (1984) |
| 1962.2 w br | Α | | 1963.9 | 16.7 | 0.00921 | 0.0122 | 2v ₁₅ (1966) |
| 1942.9 m | В | _ | 1942.5 | 17.4 | 0.0247 | 0.0200 | $v_{15}+v_{12}$ (1947) |
| 1881.4 w br | Α | \ | 1885.4 | 11.5 | 0.00496 | 0.00910 | $v_{23}+v_{30}$ (1881) or $v_{26}+v_{29}$ (1887) |
| | | ĺ | 1878.6 | 13.4 | 0.00508 | 0.00923 | or v ₁₅ +v ₁₆ (i886) |
| 1861.9 m | В | Į | 1865 3 | 12.4 | 0.0113 | 0.0138 | $v_{24}+v_{11}$ (1861) or $v_{27}+v_{10}$ (1859) |
| | | Į | 1859.0 | 10.6 | 0.00953 | 0.0127 | |
| ~1825 vw sh | | | 1816.0 | 12.8 | 0.00141 | 0.00495 | $v_7 + v_{17}$ (1825) or $v_{15} + v_{13}$ (1813) |
| ~1806 w sh | | , | 1806.4 | 15.5 | 0.00265 | 0.00680 | $2v_{16}$ (1806) or ? $v_{28}+v_{17}$ (1809) |
| 1787.8 m br | Α | Į | 1789.4 | 17.1 | 0.0152 | 0.0164 | v_7+v_{10} (1786) |
| | | Į | 1780.3 | 18.4 | 0.00841 | 0.0122 | |
| | | | 1736.7 | 9.6 | 0.00288 | 0.00723 | |
| 1730.5 m br | В | { | 1731.1 | 9.8 | 0.00632 | 0.0107 | $v_{16}+v_{13}$ (1733) |
| | | | 1725.1 | 13.7 | 0.00699 | 0.0113 | J |
| ~1703 vw sh | | | 1703.3 | 14.7 | 0.00168 | 0.00557 | $v_9 + v_{10}$ (1704) or $v_{12} + v_{17}$ (1705) |
| 1685.9 vw | | | 1686.0 | 4.2 | 0.000270 | 0.00225 | |
| | | | 1682.1 | 2.0 | 0.000029 | 0.00074 | |
| 1645.5 w | | | 1644.6 | 15.6 | 0.00891 | 0.0131 | $v_{16}+v_{17}$ (1644) or $v_{12}+v_{18}$ (1649) |
| 1636.9 w br | | | | | | | v ₈ +v ₂₉ (1637) |
| 1622.1 w br | | | 1622.3 | 6.1 | 0.00301 | 0.00764 | $v_{25}+v_{30}$ (1623) or $v_{27}+v_{19}$ (1625) |

Table 5.1 - Continued

| Exp α _m " ab | Gasc | | Fitted | <i>I</i> ; | С, | $ \overrightarrow{R_i} $ | Assignment ^d |
|-------------------------|------------|-------------|------------------|------------------|----------|--------------------------|---|
| cm ⁻¹ | | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | - |
| ~1591 s sh | A?? | | 1592.0 | 4.0 | 0.00750 | 0.0122 | 4 . 4 % (1.588) or v4 ?? |
| 1583.9 vs | B?? | | 1583.5 | 7.4 | 0.179 | 0.0597 | v_{23} and v_{23} |
| 1566.1 s br sh | | | 1566.1 | 11.7 | 0.0386 | 0.0279 | $v_{26}+v_{30}$ (1570) |
| ~1496 s sh | | | 1494.4 | 13.8 | 0.0249 | 0.0229 | v-+v ₁₁ (1499) or v ₈ +v ₁₉ (1491) |
| 1477.8 vs | Α | | 1477.6 | 4.9 | 0.354 | 0.0869 | V ₅ |
| | | ſ | 1446.0 | 11.6 | 0.0594 | 0.0360 | |
| 1445.5 vs | В | 1 | 1445.4 | 3.0 | 0.0677 | 0.0384 | \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ |
| 1387.1 w | | | 1386.7 | 9.6 | 0.00588 | 0.0116 | $v_{10}+v_{18}$ (1387) |
| 1370.7 w | A | | 1370.6 | 10.9 | 0.00696 | 0.0126 | $v_{16}+v_{19}$ (1371) or $2v_{18}$ (1370) |
| 1325.2 w | B ? | | 1325.3 | 5.7 | 0.00270 | 0.00801 | v ₂₅ |
| | | | 1319.1 | 9.9 | 0.00159 | 0.00616 | $v_8 + v_{30}$ (1320) or $v_{16} + v_{11}$ (1318) |
| 1298.1 m | В | | 1298.2 | 8.0 | 0.00634 | 0.0124 | $v_9 + v_{30}$ (1299) or $v_{16} + v_{14}$ (1303) |
| | | | | | | | or v ₁₃ +v ₁₉ (1298) |
| 1272.5 m | B? | \ | 1273.0 | 5.0 | 0.00194 | 0.00693 | $\left. \begin{array}{l} \left. \left. \right\rangle \right. \left. \left. \left\langle \right. \right\rangle \right. \left. \left. \left\langle \right. \right\rangle \right. \left. \left. \left\langle \right. \right\rangle \right. \left. \left\langle \left. \left. \left\langle \right. \right\rangle \right. \left. \left\langle \left. \left. \left\langle \right. \right\rangle \right. \left. \left\langle \left. \left\langle \left. \right\rangle \right. \left. \left\langle \left. \left\langle \right. \right\rangle \right. \left. \left\langle \left. \left\langle$ |
| 1373.3 111 | | l | 1269.3 | 12.9 | 0.00698 | 0.0132 |] *26 |
| 1235.2 w | • | } | 1237.7 | 10.0 | 0.00329 | 0.00915 | $v_{13}+v_{14}$ (1230)? |
| 1233.2 W | Α | l | 1232.7 | 9.3 | 0.00272 | 0.00834 | \(\frac{1}{3}\tau_{14}(1230)\)? |
| 1210.9 w | Α | | 1210.6 | 13.5 | 0.00491 | 0.0113 | $v_{17}+v_{19}$ (1209) |
| 1171 6 m | • | | 1172.4 | 18.7 | 0.0167 | 0.0212 | v ₆ |
| 1171.6 m | Α | | 1171.1 | 1.5 | 0.000040 | 0.00104 | $v_{10}+v_{19}$ (1171) |
| 1156.4 m | | | 1155.9 | 11.3 | 0.00632 | 0.0131 | V ₂₇ |
| 1122.7 s | A ? | | 1123.2 | 7.5 | 0.0336 | 0.0307 | $v_{10}+v_{11}$ (1118) |
| ~1116 s sh | | | 1116.6 | 9.9 | 0.0244 | 0.0262 | v ₄ -v ₁₉ (1116) |
| ~1093 s sh | | | 1094.1 | 13.5 | 0.0282 | 0.0285 | $v_{16}+v_{20}$ (1099) |
| 1092 6 | | [| 1083.9 | 7.9 | 0.256 | 0.0863 | \ |
| 1083.6 vs | A | ĺ | 1080.8 | 6.6 | 0.0417 | 0.0349 | \\ \ ⁷ |
| 1068.1 s | В | ſ | 1068.0 | 3.7 | 0.0279 | 0.0287 | |
| | | 1 | 1064.2 | 4.5 | 0.00796 | 0.0153 | } ∨ ₂₈ |
| | | ĺ | 1023.1 | 2.4 | 0.0382 | 0.0343 |) |
| 1022.8 vs | A | Į | 1022.3 | 3.3 | 0.0749 | 0.0480 | \v_8 |
| 1022.8 VS | A | | 1020.0 | 3.2 | 0.00360 | 0.0105 | |
| | | | | | | | , |

| Table 5.1 - Continue |
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| Table 5.1 - Cont Exp $\alpha_m^{"}$ *b | Gasc | | Fitted | Γ, | С, | $ \overrightarrow{R_i} $ | Assignment ^d | |
|---|------------|----------|------------------|------------------|----------|--------------------------|--|--|
| cm ⁻¹ | | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | | |
| 1012.6 s | | | 1012.3 | 4.4 | 0.00486 | 0.0123 | v ₂₉ +v ₁₄ (1014) | |
| 1001.9 s | A ? | | 1001.9 | 2.6 | 0.0206 | 0.0254 | V9 | |
| 990.5 m | | | 990.6 | 11.4 | 0.00428 | 0.0117 | v ₁₀ +v ₃₀ (1000) | |
| 983.2 m br | | | 980.7 | 13.4 | 0.00498 | 0.0126 | v_{15} | |
| 963.7 m br | | | 963.0 | 18.7 | 0.00852 | 0.0167 | \mathbf{v}_{12} | |
| 935.0 m | Α | | 935.0 | 10.6 | 0.0125 | 0.0205 | $v_{17}+v_{20}$ (937) or $2v_{19}$ (936) | |
| ~919 m sh | | | 919.4 | 5.6 | 0.000896 | 0.00554 | v ₂₉ +v ₃₀ (912) | |
| 903.0 s asym | C | _ | 903.5 | 6.9 | 0.0293 | 0.0320 | | |
| 203.0 s asym | C | l | 900.1 | 8.5 | 0.0153 | 0.0231 | \right\{\nu_{16}} | |
| ~880 w sh ? | | | 878.5 | 7.5 | 0.000495 | 0.00421 | v ₁₈ +v ₂₀ (881) | |
| 866.3 w | | | 865.4 | 9.9 | 0.00270 | 0.00991 | v ₁₉ +v ₁₄ (868) | |
| 830.0 m | | | 830.2 | 12.8 | 0.00968 | 0.0192 | v_{13} | |
| 812.2 m | ?? | | 813.0 | 6.7 | 0.00132 | 0.00715 | $v_{11}+v_{14}$ (815) or $v_{29}+v_{20}$ (810) | |
| | | | 743.9 | 5.6 | 0.259 | 0.105 | | |
| 741.2 vs | С | 1 | 741.2 | 4.9 | 0.329 | 0.118 | v_{17} | |
| | | | 738.6 | 5.6 | 0.221 | 0.0971 | } | |
| | | Ì | 702.9 | 4.0 | 0.122 | 0.0739 |) | |
| 702.5 vs | Α | 1 | 701.0 | 4.7 | 0.0647 | 0.0539 | \v_{10} | |
| 685.2 vs | С | | 685.1 | 4.8 | 0.217 | 0.100 | v_{18} | |
| 662.1 s | ?? | | 661.9 | 6.3 | 0.00847 | 0.0201 | v ₁₉ +v ₂₀ (664) | |
| 614.1 m | В | | 614.0 | 3.7 | 0.00280 | 0.0120 | V ₂₉ | |
| | D | ſ | 549.2 | 2.8 | 0.000103 | 0.00243 |) | |
| 547.7 w | | 1 | 544.9 | 2.8 | 0.000105 | 0.00243 | $v_{17}-v_{20}$ (545) | |
| 533.4 w | | (| 533.8 | 2.5 | 0.000222 | 0.00234 | | |
| 468.1 vs | С | | 468.0 | 3.9 | 0.0983 | 0.0813 | V ₁₉ | |
| ~434 m br sh | | | 433.9 | 3.3 | 0.000549 | 0.00631 | v ₁₆ -v ₁₉ (435) | |
| 415.0 s | Α | | 415.1 | 5.5 | 0.0185 | 0.00031 | V ₁₁ | |
| ~400 m sh | Λ | | 408.8 | 7.3 | 0.00621 | 0.0373 | V ₁₄ | |

a - Wavenumbers of peaks in the imaginary molar polarizability spectrum of liquid chlorobenzene.

b - Abbreviations used: v-very, w-weak, m-medium, s-strong, br-broad, sh-shoulder and db-doublet.

c - Band contours in the infrared spectrum of the gas.

d - Calculated wavenumbers are given in brackets.

In the last column of Table 5.1, an assignment is given. The assignments of the fundamentals were discussed in Chapter 4. The remaining peaks were assigned as overtone or combination transitions based on the band shape of the gas spectrum when observed, and on the agreement of the observed wavenumber with sums or differences of fundamental wavenumbers.

The accuracy of the C_j values is estimated from the sum of the accuracy of the fit and the accuracy of the real and imaginary refractive indices, n and k, that are used in the calculation of α_m^m . The accuracy for the refractive indices was reported in Chapter 3 and is 0.2% for n and on the order of 2 to 3% for k. The accuracy of the fit is estimated from the accuracies of peak heights and areas under the α_m^m spectrum, and is about 0.6 to 3% depending on the intensity of the band. Therefore, we estimate that the C_j values are accurate to better than 3-5% for strong peaks and to better than 5-10% for weak peaks.

5.1.1 - Integrated intensities and dipole moment derivatives of the fundamental vibrations of liquid chlorobenzene.

The integrated intensities, C_j , and the absolute values of the dipole moment derivatives with respect to normal coordinates, $|\mu_j|$, of the fundamental vibrations of liquid chlorobenzene are given in Table 5.2. The $|\mu_j|$ were calculated from the C_j using Eq. 1.4.4. As discussed earlier, the accuracy of C_j for strong fundamentals is estimated to be 3-5% and 5-10% for weak bands. Thus, in general, the percent accuracy of $|\mu_j|$ for

Table 5.2 - Dipole moment derivatives of liquid chlorobenzene

| | · · · · · · · · · · · · · · · · · · · | | | A _{l,gas} b | | |
|-------------------|---------------------------------------|---------------------|---------------------------|------------------------|------------------------------|--|
| ν, | Fitted cm ^{-1 a} | $C_{j,lig}$ b | <i>µ</i> _j c | This work ^d | Pulay [Ref. 47] ^e | |
| v_1 | 3069.5 f | 0.0482 ^f | 0.300 ^f | 3.81 ^f | 2.51 | |
| V ₂ | 3069.5 ^f | t | f | f f | 15.89 | |
| V3 | 3057.9 | 0.0163 | 0.175 | 1.29 | 0.00 | |
| V ₄ | 1592.0 ⁸ | 0.0075 ^g | 0.118 ^g | 0.59 ^g | 2.12 | |
| V5 | 1477.6 | 0.354 | 0.813 | 27.95 | 44.83 | |
| V ₆ | 1172.4 | 0.0167 | 0.177 | 1.32 | 0.05 | |
| V7 h | 1083.9 | 0.298 | 0.971 | 23.50 | 40.23 | |
| /8 h | 1022.3 | 0.117 | 0.723 | 9.21 | 10.98 | |
| V 9 | 1001.9 | 0.0206 | 0.196 | 1.63 | 2.06 | |
| ′10 h | 702.9 | 0.187 | 0.825 | 14.74 | 22.76 | |
| vii | 415.1 | 0.0185 | 0.186 | 1.46 | 3.46 | |
| V12 | 963.0 | 0.00852 | 0.126 | 0.67 ⁱ | i | |
| V ₁₃ | 830.2 | 0.00968 | 0.134 | 0.76 | i | |
| V ₁₄ | 408.8 | 0.00621 | 0.108 | 0.49 1 | <u> </u> | |
| 15 | 980.7 | 0.00498 | 0.096 | 0.39 | 0.08 | |
| 16 h | 903.5 | 0.0446 | 0.403 | 3.52 | 3.94 | |
| 17 h | 741.2 | 0.809 | 2.123 | 63.88 | 46.75 | |
| V18 | 685.1 | 0.217 | 0.637 | 17.13 | 42.45 | |
| 119 | 468.0 | 0.0983 | 0.429 | 7.76 | 8.99 | |
| / 20 | (196.0) | | | | 0.09 | |
| / 21 | 3084.0 | 0.0319 | 0.244 | 2.52 | 8.80 | |
| V ₂₂ | (3060.0 ??) ^յ | | | | 8.77 | |
| V ₂₃ | 1583.5 ⁸ | 0.179 ⁸ | 0.578 ^g | 14.13 ^g | 19.07 | |
| v ₂₄ h | 1445.4 | 0.127 | 0.689 | 10.04 | 10.79 | |
| V ₂₅ | 1325.3 | 0.00270 | 0.071 | 0.21 | 0.58 | |
| v ₂₆ h | 1269.3 | 0.00892 | 0.174 | 0.70 | 0.34 | |
| V ₂₇ | 1155.9 | 0.00632 | 0.109 | 0.50 | 0.00 | |
| V ₂₈ h | 1068.0 | 0.0359 | 0.350 | 2.83 | 2.03 | |
| V ₂₉ | 614.0 | 0.0028 | 0.072 | 0.22 | 0.17 | |
| V ₃₀ | (297.5) | | | | 0.18 | |

a - $\,$ Integrated intensities were not obtained for ν_{20} and ν_{30} which are below the experimental

wavenumber range of this work.

- b Units are km mole⁻¹.
- c Units are Debye A-1 amu-12.
- d Integrated intensities for the gas, $A_{j,gas}$ were calculated as $A_{j,gas} = 8\pi^2 C_{j,hq}$ with the assumption that $\mu_{j,hq}^2 = \mu_{j,gas}^2$.
- e $A_{l,ab}$ in values obtained from ab initio calculations reported in Ref. 47.
- f The same wavenumber was assigned to v_1 and v_2 . All quantities are assigned to v_1 in the table.
- g 1584 cm⁻¹ was assigned to both v₄ and v₂₃ although the evidence for v₄ is less conclusive. The intensity of the fitted band at 1583.5 cm⁻¹ was assigned in the table to v₂₃. The reported values for v₄ are those of the much weaker band at 1592 cm⁻¹ which may possibly be v₄.
- h Several CDHO bands were used to fit the observed band. The wavenumber reported is that of the prominent contributing band. All related quantities for this vibration are the sums from all contributing bands as shown in Table 5.1.
- i A_2 vibrations in chlorobenzene are infrared inactive in the spectrum of the gas, i.e $A_{j,gas}$ =0 and were not calculated in Ref. 47. Our values are derived from $C_{j,liq}$ as described in footnote d.
- j The wavenumber was assigned based on Raman spectra.

strong fundamentals is about 1.5-2.5% and about 2.5-5% for weak fundamentals.

The only integrated intensities for chlorobenzene in the literature are those reported by Pulay *et al.*⁴⁷. In their study the authors used scaled ab initio calculations to obtain wavenumbers and infrared integrated intensities of the fundamental vibrations of chlorobenzene. In order to compare our results with Pulay's, our integrated intensities of the liquid are converted to the integrated intensities the corresponding band would have in the gas phase if the dipole moment derivative were the same in the liquid and gas phases.

Equations 1.4.4 and 1.4.5 relate the square of the dipole moment derivative to the integrated intensities C_i and A_i . For a gas \underline{n} is 1 and Equation 1.4.5 simplifies to

$$A_{j,\text{gas}} = \frac{N_{\text{A}}\pi}{3c^2} g_j \,\mu_j^2 \tag{5.1.2}$$

This equation and Equation 1.4.4 give

$$\frac{A_{IRas}}{C_{I^{liq}}} = 8\pi^2 \left(\frac{\mu_{IRas}}{\mu_{I^{liq}}}\right)^2 \tag{5.1.3}$$

Thus, if the dipole moment derivative is the same in the gas and liquid phases, $A_{j,gas}$ can be calculated from our measured C_j values by

$$A_{j,gas} = 8\pi^2 C_j = 78.96 C_j \tag{5.1.4}$$

The resulting $A_{j,gas}$ are included in Table 5.2 under the heading " $A_{j,gas}$ this work". The integrated intensities, $A_{j,ab\ m}$, obtained from ab initio calculation by Pulay et al.⁴⁷ are given in column 6 of Table 5.2 under the heading " $A_{j,gas}$ Pulay".

In the following subsections, figures of the curve fitted and experimental α_m^m spectra and the CDHO bands required in the fit, are given for regions of small wavenumber range. The intensities of fundamentals in these regions will be discussed and compared with Pulay's *ab initio* intensities.

5.1.1a - Intensities of the CH stretching vibrations

The experimental and curve fitted α_m'' spectra of liquid chlorobenzene for the region of the CH stretching vibrations are shown in Figure 5.2. There are seven bands required to fit this region, centred at 3084, 3069.5, 3057.9, 3056.3, 3025.8, 3015.7 and 3003.7 cm⁻¹. The fit in this region is very good. The area under the $\tilde{\nu}\alpha_m'''$ spectrum of

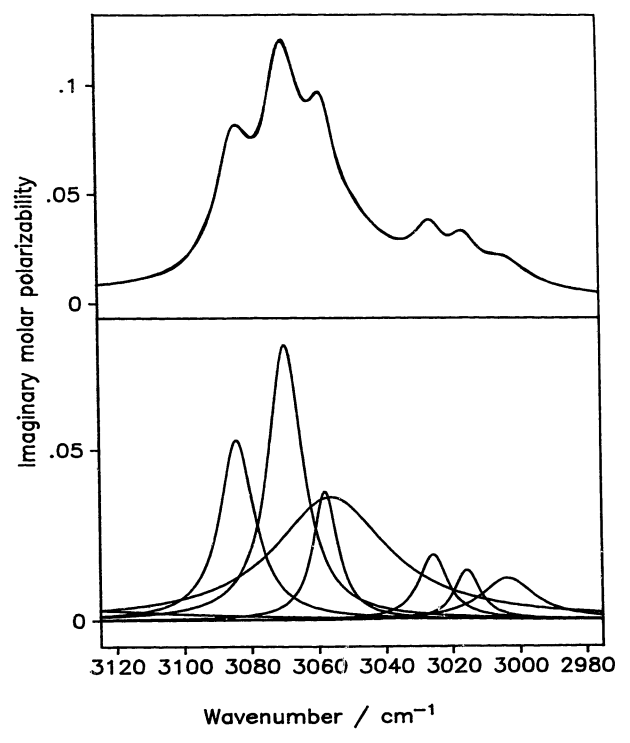


Figure 5.2 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid chlorobenzene between 3125 and 2975 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Contribution from the 3140 cm⁻¹ band can be seen as a tail inside the shown region from 3125 to ~3060 cm⁻¹. Units for both boxes are cm³ mole⁻¹.

the fitted bands is 0.1% greater than the area under the $\tilde{\nu}\alpha_m^{\prime\prime}$ experimental spectrum. The peak heights are fitted on average to 0.3% and to within 0.7%.

In Chapter 4, three of the five CH stretching fundamentals were assigned to bands in the infrared spectrum, v_1 , v_3 and v_{21} , and v_2 and v_{22} were assigned to features in the Raman spectrum. It is not clear whether v_2 and v_{22} contribute to the infrared spectrum, although they well may do so. It is also not clear whether all of the intensity in the CH stretching region should be assigned to the active fundamentals, or just the intensity in the band required to fit the prominent features that were assigned to the fundamentals. It is probable that all of the intensity in the region should be so assigned, on the grounds that some of the fundamental intensity is lent to the many overtone and combination transitions that contribute to the extensive absorption in this region, through Fermi resonance between the fundamental state and the various overtone and combination states. The intensity of the weak overtone and combination bands near the CH stretching fundamentals is noticeably greater than the intensity of the weak features further removed, such as those between 2 900 and 2000 cm^{-1} .

Accordingly, the intensity is assigned in two ways. First, the intensities of the fitted bands at 3084, 3069.5 and 3058 cm⁻¹ are assigned to v_{21} , v_1 , and v_3 , respectively, and no infrared intensity is assigned to v_2 and v_{22} . The total intensity assigned to the fundamentals in this way is, from Table 5.1, $\Sigma C_{CH} = 0.0964$ km mole⁻¹. The intensity left to be assigned to overtone and combination bands in this region is 0.0974 km mol⁻¹, greater than that assigned to the fundamentals.

As a consequence, an alternative assignment of the intensity is considered, namely that the total intensity of the bands between 3125 and 2975 cm⁻¹, $\Sigma C_j = 0.194$ km mole⁻¹ is assigned to the CH stretching fundamentals as a group. With this assignment it is recognized that the harmonic and anharmonic interactions are sufficiently complex that assignment of the intensity to individual normal vibrations is not possible, and possibly not even meaningful.

The equivalent intensity $A_{I\,gas}$ of the fundamentals in these two assignments is 7.61 km mol⁻¹ and 15.32 km mol⁻¹, respectively. The sum of the intensity of the fundamentals calculated by Pulay⁴⁷ is 35.97 km mol⁻¹. Unfortunately, Pulay did not calculate the intrinsic intensity of overtone and combination bands. Clearly, Pulay's calculated intensities are 4.7 or 2.4 times those found for the liquid, depending on the assignment used. For benzene⁷³ the ratio between the actual intensities of the gas and those of the liquid, both expressed as $A_{I\,gas}$, is 1.72. Thus, part of the difference between Pulay's calculations and our results may be a significant difference between the intensities of the CH stretching bands of chlorobenzene in the gas and liquid phases.

5.1.1b - Intensities of the v_4 , v_5 , v_{23} and v_{24} vibrations

The experimental α_m'' and curve fitted α_m'' spectra of liquid chlorobenzene between 1650 and 1400 cm⁻¹ are shown in Figure 5.3. The fit in this region is good. The area

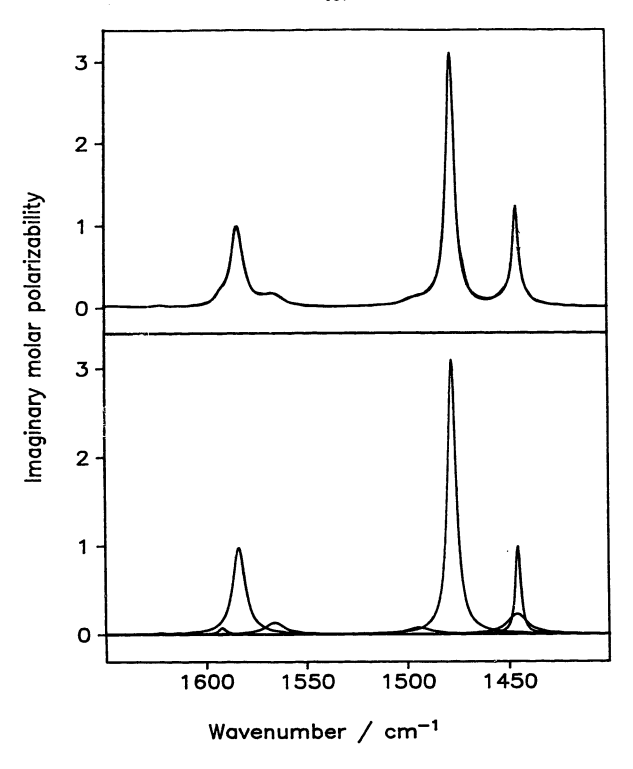


Figure 5.3 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid chlorobenzene between 1650 and 1400 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Units for both boxes are cm³ mole⁻¹.

under the curve fitted α_m'' spectrum is greater by 0.75% than the area under the experimental α_m'' spectrum. The peak heights of the fundamentals are reproduced to better than 0.32%. Those of the weak band at 1622 cm⁻¹ and the shoulders at 1566 and 1496 cm⁻¹ are reproduced to 5-6%. All of these three bands are due to combination transitions.

In Chapter 4, v_{23} and v_4 were assigned at the same wavenumber, 1584 cm⁻¹. The assignment of v_{23} was supported by the various experimental data. On the other hand, the assignment of v_4 was less conclusive and relied on a possible Q branch at 1589 cm⁻¹ in the spectrum of the gas phase. Therefore, we assign the intensity of the strong CDHO band at 1583.5 cm⁻¹ to v_{23} . A weak CDHO band centered at 1592 cm⁻¹ was needed to fit the 1591 cm⁻¹ shoulder. We tentatively assign its intensity to v_4 .

The intensity of the CDHO band at 1477.6 cm⁻¹ is assigned to v_5 while those of the two CDHO bands at 1446.0 and 1445.4 cm⁻¹ which were required to fit the 1445.5 cm⁻¹ peak are assigned to v_{24} .

The *ab initio* calculated $A_{j,ab\,in}$ of Pulay et al⁴⁷ is greater by 7% for v_{24} , 60% for v_{5} and 35% for v_{23} . They are 3.6 times larger for v_{4} . However, if the intensities of v_{4} and v_{23} are combined, since the assignment of v_{4} is not as clear as that of the others, Pulay's value is only 45% greater than $A_{j,gas}$.

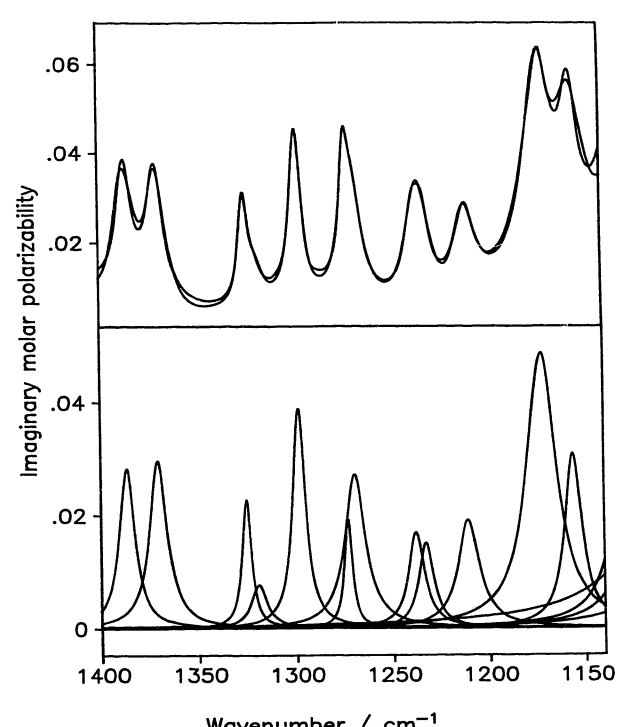
5.1.1c - Intensities of the v_6 , v_{25} , v_{26} and v_{27} vibrations

The experimental α_m''' and curve fitted α_m''' spectra of liquid chlorobenzene between 1400 and 1140 cm⁻¹ are shown in Figure 5.4. The absorption is weak and, as a consequence, the fit in this region is not good as in the previous regions. Large contributions from the strong bands at 1123 and 1084 cm⁻¹ influenced the fitting in this region and the addition of more CDHO bands didn't improve the fit significantly. Thus, the fit presented involves the minimum number of bands needed to obtain a decent, but far from perfect, fit. The area under the curve fitted α_m'' is smaller by 0.44% than the area under the experimental α_m'' spectrum. The peak heights of the 9 peaks in the region are reproduced on average to 2.5%. The reproduction of only one fundamental, v_{27} at 1156 cm⁻¹, exceeds this average with 4.25%.

The intensity of the CDHO band at 1325.3 cm⁻¹ was assigned to v_{25} while that of the close band at 1319.1 cm⁻¹ that is needed to fit the distinct shoulder in the α''_m spectrum is assigned as a combination transition. Similarly, the intensity of the CDHO band at 1172.4 cm⁻¹ is assigned to v_6 while that of the hardly visible (Fig. 5.2), much weaker band at 1171.1 cm⁻¹ is assigned to a combination transition.

The intensities of the two CDHO bands at 1273.0 and 1269.3 cm⁻¹ are assigned to v_{26} while that of the band at 1155.9 cm⁻¹ is assigned to v_{27} .

For v_{25} , Pulay's⁴⁷ reported $A_{j,ab\ in}$ equals 0.58 km mole⁻¹, greater by a factor of 2.8 than our value of 0.21 km mole⁻¹. Pulay's reported values for the other three fundamentals in this region are much smaller than the values obtained in this work. For



Wavenumber / cm⁻¹
Figure 5.4 - Top - Experimental imaginary molar polarizability, α_m^m , and curve fitted α_m^m spectra of liquid chlorobenzene between 1400 and 1140 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m^m spectrum. Strong contributions from 4 bands at lower wavenumbers are clearly seen as tails from 1140 to about 1300 cm⁻¹. Units for both boxes are cm³ mole⁻¹.

 v_{26} , Pulay's value is about half of ours, 0.34 versus 0.70 km mole⁻¹. For v_6 , Pulay's value is about 4% of our value, and for v_{27} Pulay's reported value is zero while our value is 0.50 km mole⁻¹.

The fact that for v_6 , v_{26} and v_{27} , our A_j values for the gas are much greater than those reported by Pulay is perhaps an indication that for these three weak vibrations, the small dipole moment derivatives are influenced by the liquid forces and are greater in the liquid than in the gas.

5.1.1d - Intensities of the v_7 , v_8 , v_9 and v_{28} vibrations

The experimental α_m'' and curve fitted α_m'' spectra of liquid chlorobenzene between 1140 and 990 cm⁻¹ are shown in Figure 5.5. The fit in this region is very good. 12 bands were needed to fit the nine features visible in the spectrum. Two of the three unsuspected bands were needed to fit the 1022.8 cm⁻¹ peak and the third was needed to fit the 1084 cm⁻¹. The area under the curve fitted α_m'' is greater by 0.49% than the area under the experimental α_m'' spectrum. Peak heights in this region are reproduced on average to better than 1%.

Two CDHO bands at 1083.9 and at 1080.8 cm⁻¹ were needed to obtain a good fit of the very strong band observed at 1083.6 cm⁻¹ in the α_m'' spectrum of the liquid. Their combined intensities are assigned to ν_7 . Similarly the intensities of the two CDHO bands at 1068.0 and at 1064.1 cm⁻¹ are assigned to ν_{28} and those of the bands at 1023.1,

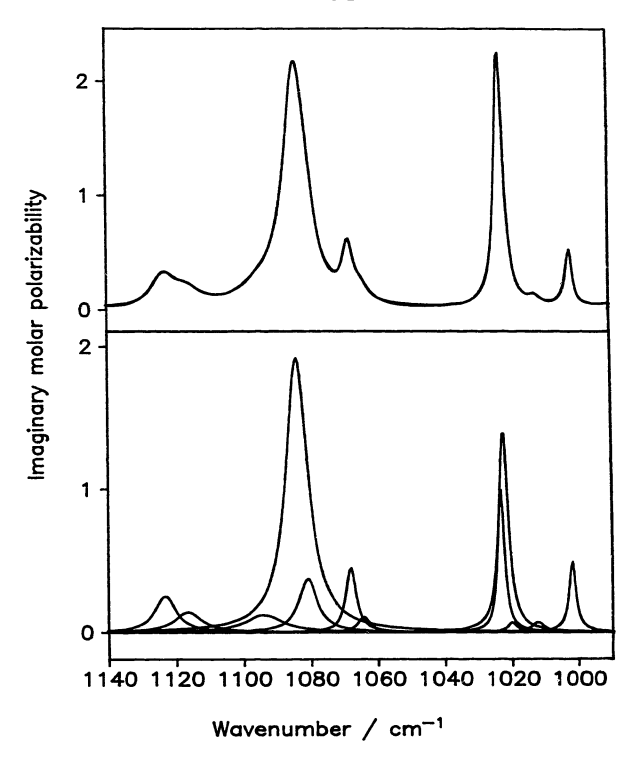


Figure 5.5 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid chlorobenzene between 1140 and 990 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Units for both boxes are cm³ mole⁻¹.

1022.3 and 1020.0 cm⁻¹ are assigned to v_8 . The v_9 band at 1001.9 cm⁻¹ was fitted nearly perfectly by a single fitted band

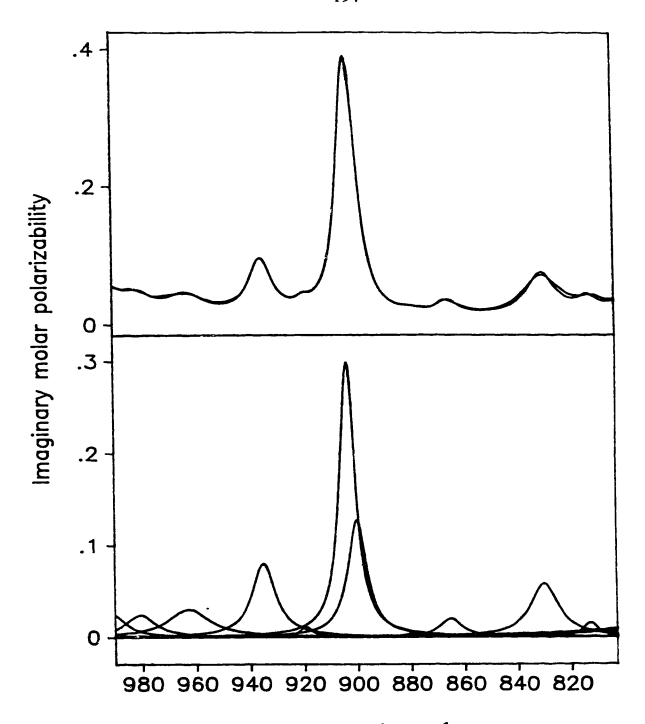
Pulay's⁴⁷ reported $A_{j,ab\ in}$ values for v_7 , v_8 and v_9 are greater than the assigned $A_{j,gas}$ values by 71%, 19% and 26%, respectively (Table 5.2). Pulay's reported $A_{j,ab\ in}$ for v_{28} , is smaller than the assigned value by about 18%.

5.1.1e - Intensities of the v_{12} , v_{13} , v_{15} and v_{16} vibrations

The experimental α_m''' and curve fitted α_m''' spectra of liquid chlorobenzene between 990 and 800 cm⁻¹ are shown in Figure 5.6. The fit between 950 and 850 cm⁻¹ is very good but in the remaining, weak, regions it is not good. At the low wavenumber end this is partly because the spectrum is influenced by strong bands outside the region, the tails of which can be clearly seen in Fig 5.6. The area under the curve fitted α_m'' is smaller by 1.32% than the area under the experimental α_m'' spectrum. Peak heights in this region are reproduced on average to 2.5%.

Two CDHO bands at 903.5 and at 900.1 cm⁻¹ were needed to obtain a good fit of the strong asymmetric band observed at 903.0 cm⁻¹ in the α_m'' spectrum of the liquid. Their combined intensities were assigned to v_{16} .

 v_{12} and v_{13} are A_2 vibrations and as such are infrared inactive in the spectrum of the gas and their intensities were not calculated by Pulay⁴⁷. The values reported by us in Table 5.2 for the $A_{j,gas}$ of the A_2 vibrations were calculated from the $C_{j,lig}$ following



Wavenumber / cm⁻¹
Figure 5.6 - Top - Superimposed experimental imaginary molar polarizability, α_m^n , and curve fitted α_m^n spectra of liquid chlorobenzene between 990 and 800 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m^n spectrum. Strong contributions from bands outside the shown region can be seen as tails on either sides of the region. Units for both boxes are cm³ mole⁻¹.

the same procedure as for the other modes.

For the two B₁ vibrations in this region, v_{15} and v_{16} , Pulay's reported $A_{j,ab\ in}$ is smaller than our assigned value for v_{15} by 79% (0.05 vs. 0.39 km mole⁻¹) and for v_{16} is greater by 12% (3.94 vs. 3.52 km mole⁻¹). Pulay's calculated relative intensities support the assignment of these two modes, particularly that of v_{15} to an extremely weak infrared feature.

5.1.1f - Intensities of the v_{10} , v_{17} and v_{18} vibrations

The experimental α_m'' and curve fitted α_m'' spectra of liquid chlorobenzene between 800 and 650 cm⁻¹ are shown in Figure 5.7. The area under the curve fitted α_m'' is greater by 1.66% than the area under the experimental α_m'' spectrum. The peak height of v_{17} at 741 cm⁻¹ is reproduced to 1.55% while those of v_{10} and v_{18} at 703 and 685 cm⁻¹, respectively, are reproduced to better than 0.15%.

Three CDHO bands at 743.9, 741.2 and 738.6 cm⁻¹ were needed to obtain a good fit of the very strong band observed at 741.2 cm⁻¹ in the α''_m spectrum of the liquid. Their combined intensities were assigned to v_{17} . Pulay's⁴⁷ reported $A_{j,ab~in}$ of 46.75 km mole⁻¹ is lower by 27% the assigned intensity of 63.88 km mole⁻¹.

Two CDHO bands at 702.9 and 701.0 cm⁻¹ were needed to obtain fit the v_{10} band at 702.5 cm⁻¹ and their combined intensities were assigned to v_{10} . Pulay's⁴⁷ reported $A_{f,ab\ in}$ of 22.76 km mole⁻¹ is 55% greater than the assigned value of 14.74 km mole⁻¹.

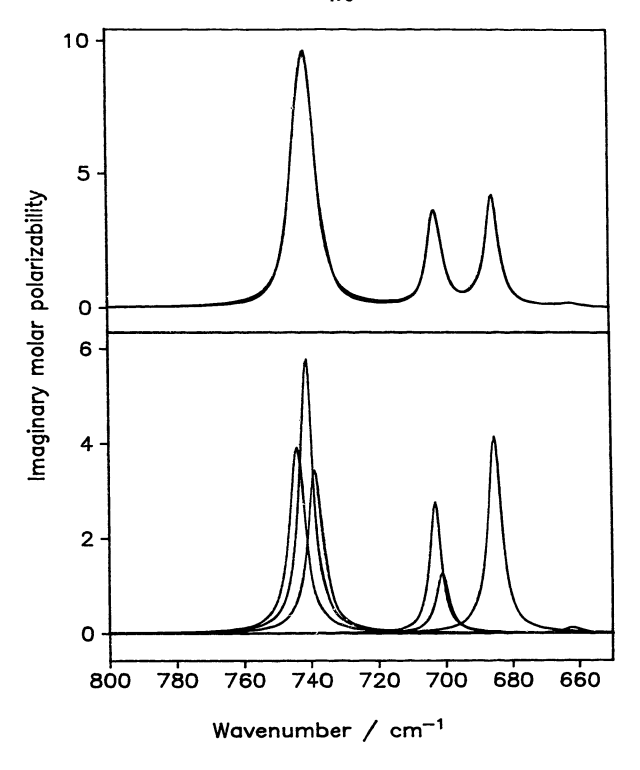


Figure 5.7 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid chlorobenzene between 800 and 650 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Units for both boxes are cm³ mole⁻¹.

 v_{18} at 685 cm⁻¹ was fitted by a single of C_j =0.217 km mole⁻¹ and corresponding $A_{j,gas}$ = 17.13 km mole⁻¹. Pulay's value 42.45 km mole⁻¹, greater by a factor of 2.5

5.1.1g - Intensities of the v_{11} , v_{14} , v_{19} , v_{20} , v_{29} and v_{30} vibrations

The experimental α_m'' and curve fitted α_m'' spectra of liquid chlorobenzene between 650 and 350 cm⁻¹ are shown in Figure 5.8. The fit in this region is not as good as in other regions. The area under the curve fitted α_m'' is greater by 7.8% than the area under the experimental α_m'' spectrum. Peak heights are reproduced to 2-4%. The poor fit is due partly to the curve fitted wings of the band at 468 cm⁻¹ being higher than the wings of that band in the experimental spectrum, and partly to poorer experimental data between 521 and 484.5 cm⁻¹ for which k was set to zero (Table 3.5).

The bands at 614 cm⁻¹, due to v_{29} , and 468 cm⁻¹, due to v_{19} , were each fitted by a single CDHO band. The two fitted bands at 415.1 and 408.8 cm⁻¹ were assigned to v_{11} and v_{14} , respectively, the latter very tentatively.

No comparison is made for three vibrations. The wavenumbers of two fundamentals, v_{20} and v_{30} , are below the experimental range so their intensities were not measured. The intensity of v_{14} , an A_2 vibration, is zero for the gas, so was not calculated by Pulay *et al.*⁴⁷.

Pulay's⁴⁷ reported intensities, $A_{j,ab\ in}$, for v_{11} , v_{19} and v_{29} are greater by a factor of 2.4, greater by 16% and smaller by 23%, respectively, than our assigned intensities, $A_{j,gas}$,

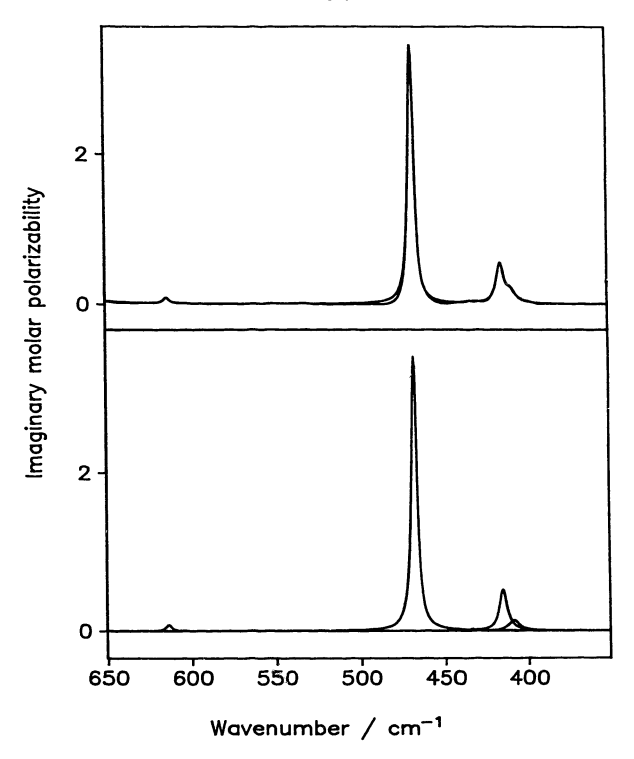


Figure 5.8 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid chlorobenzene between 650 and 350 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Units for both boxes are cm³ mole⁻¹.

for these vibrations. However, the magnitude of Pulay's intensities clearly support the assignments.

5.1.2 - Summary of intensity assignment of chlorobenzene

The assignment of the fundamental vibrations to the experimentally observed infrared bands of liquid chlorobenzene is given in Table 4.10, and the assignment of the intensity of the fitted bands is summarized in Table 5.2.

The comparison of the $A_{j,gas}$ values, which were calculated from the C_j values of the liquid under the assumption that the dipole moment derivatives are the same in the gas and liquid phases, with the $A_{j,ab,in}$ values, which were calculated by Pulay⁴⁷ by ab initio quantum mechanics is of interest. The agreement is extremely good in the sense that Pulay's calculations reproduce the pattern of intensities extremely well. He calculates weak bands to be week and intense bands to be intense. The numerical agreement is within a factor of two for most cases, which is remarkably good. In principle there is no reason for agreement between intensities calculated for the gas and those observed for the liquid. The agreement observed must mean that the vibrations in question have very similar intensities in the two phases, and that the ab initio calculations do a remarkably good job of calculating them.

There are notable exceptions. As discussed in Section 5.1.1a, the CH stretching intensities in Table 5.2 were obtained by assigning the intensity of the fitted band to a specific fundamental, and the total is nearly 5 times smaller than Pulay calculated. If the

total intensity in the region is assigned to the CH stretching fundamentals collectively, the measured intensities are smaller than Pulay's by a factor of 2.4. This can be compared with a factor of 1.7 between the intensities measured for gaseous and liquid benzene, to suggest that the CH stretching vibrations of chlorobenzene may also be much weaker in the liquid than in the gas phase. However, firm conclusions must await the measurement of the gas phase intensities.

Another exception to the factor of two generality is for v_{18} at 685 cm⁻¹, which is \sim 2.5 times weaker than calculated. But the experimental intensity of v_{17} and v_{18} combined is \sim 81 km mol⁻¹, just 9% smaller than Pulay's calculated value, 89.2 km mole⁻¹. Thus, with reference to Table 4.10, the difference probably arises because the calculations for the gas gave different contributions to these two vibrations from the two sets of out-of-plane displacements, the CCCC torsion and the oop hydrogen motion, than actually occur in the liquid. A similar explanation may well be correct for the weaker example of v_{25} and v_{26} for which the total intensity calculated is within 2% of that observed, but the distribution between the two vibrations is different.

Thus, the experimental study and assignment of the liquid and the *ab initio* calculations of Pulay are thought to support each other strongly.

The assignment of nearly all of the other fitted bands that are classified as "weak" or stronger was made from the sums of the fundamental wavenumbers with little difficulty. They will provide experimental evidence against which to compare calculations of the intensities of the overtone and combination bands, which are only

active through the anharmonic parts of the potential energy and the non-linearity of the dipole moment with respect to internuclear distance.

Many of the very weak bands above 2000 cm⁻¹ have not been assigned in this way, mainly because there are frequently many possible assignments for them.

5.2 - Integrated intensities and dipole moment derivatives of liquid toluene

The imaginary molar polarizability, α_m'' , spectrum of liquid toluene was fitted with 166 CDHO bands between 4800 and 442 cm⁻¹. An entire spectrum, called the fitted spectrum, was computed through the summation of all the 166 band spectra, each of which expanded over the full range, 4800-442 cm⁻¹. This fitted spectrum and the α_m'' spectrum are shown in Figure 5.9.

The two spectra almost coincide and, in general, the fit is very good, especially for peak heights and areas under the spectra. As can be seen from the magnified curves in Fig 5.9, the fit is not as good at the wings of sharp bands.

For peak of heights $< 0.01 \text{ cm}^3 \text{ mole}^{-1}$, between 0.01 and 0.1 cm³ mole⁻¹, between 0.1 and 1 cm³ mole⁻¹, and greater than 1 cm³ mole⁻¹, the average percent difference between the fitted and measured spectra are $\pm 3\%$, $\pm 2.3\%$, $\pm 1.1\%$ and $\pm 0.5\%$, respectively.

For the region between 4800 and 800 cm⁻¹, the area under the fitted spectrum is smaller than the area under the α_m'' spectrum by 0.45% and for the region between 800

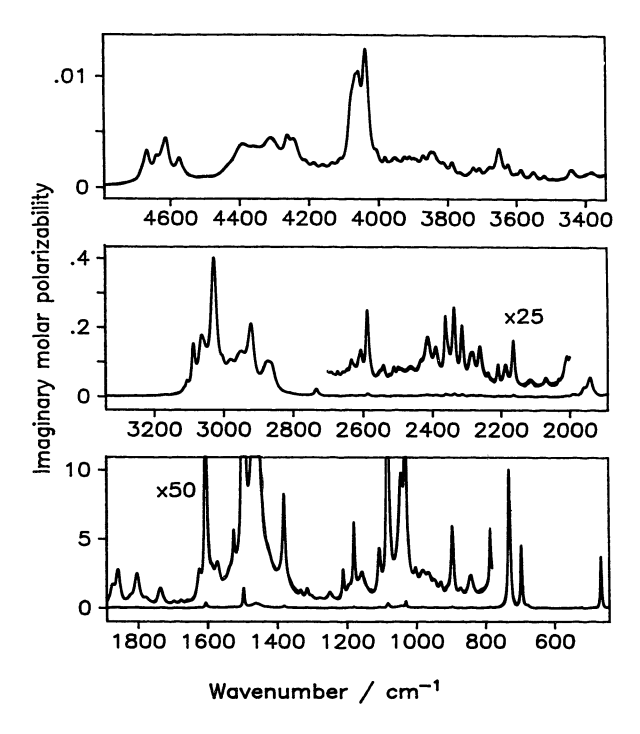


Figure 5.9 - The experimental imaginary molar polarizability, α_m'' , spectrum and curve fitted α_m'' spectrum of liquid toluene. Units are cm³ mole⁻¹. In each box, the ordinate scale is for the lower curves. It needs to be divided by the multiplication factor for the upper curves. The two spectra coincide over most of the range.

Table 5.3 - Integrated intensities of toluene.

| Exp α_m'' a,b | Gas ^c | Fitted | Γ_{j} | C_{j} | $ \overrightarrow{R_j} $ | Assignment ^d |
|----------------------|------------------|------------------|------------------|----------|--------------------------|--|
| cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| ~4716 vw br sh | | 4713.9 | 95.4 | 0.00867 | 0.00761 | |
| ~4682vw br sh | | 4681.4 | 18.4 | 0.000568 | 0.00195 | v ₂₁ +v ₄ (4691) |
| 4667.0 vw | | 4667.2 | 20.8 | 0.00378 | 0.00505 | $v_1 + v_4$ (4667) or $v_{21} + v_{23}$ (4673) |
| 4637.1 vw | | 4639.0 | 21.7 | 0.00234 | 0.00399 | v_3+v_4 (4643) or v_2+v_{23} (4642) |
| ~4623 vw sh | | 4622.9 | 20.9 | 0.00138 | 0.00307 | v ₃ +v ₂₃ (4625) |
| 4612.1 vw | | 4611.3 | 21.6 | 0.00481 | 0.00573 | v ₂₂ +v ₂₃ (4614) |
| 4573.6 vw br | \[| 4573.5 | 18.2 | 0.00153 | 0.00325 | $v_{21}+v_5$ (4582) ? |
| | l | 4570.2 | 61.2 | 0.00348 | 0.00490 | |
| ~4536 vw br sh | | | | | | |
| ~4495 vw br sh | | 4501.9 | 59.0 | 0.00120 | 0.00290 | |
| ~4435vw br sh | | 4434.7 | 53.1 | 0.00198 | 0.00375 | $v_2 + v_{32}$ (4434) or $v_1 + v_{32}$ (4441) |
| 4388.7 vw | | 4393.3 | 57.1 | 0.0101 | 0.00851 | |
| ~4357 vw | | 4356.0 | 51.8 | 0.00474 | 0.00585 | |
| 4311.0 vw br | | 4308.1 | 63.4 | 0.0141 | 0.0102 | |
| 4261.3 vw | | 4261.9 | 21.6 | 0.00293 | 0.00465 | |
| 4244.3 vw br | | 4241.0 | 33.1 | 0.00583 | 0.00658 | |
| ~4211 vw sh | | 4208.6 | 27.3 | 0.00155 | 0.00341 | |
| 4186.3 vw | | 4185.4 | 20.5 | 0.00108 | 0.00285 | v ₂₂ +v ₂₇ (4183) |
| 4161.3 vw br | | 4160.2 | 32.2 | 0.00202 | 0.00391 | |
| 4132.3 vw | | 4132.9 | 25.4 | 0.00144 | 0.00331 | |
| 4106.7 vw br sh | | 4110.7 | 22.5 | 0.00129 | 0.00314 | |
| | | 4078.3 | 22.0 | 0.00635 | 0.0070 | |
| ~4065 w vvbr | | 4065.4 | 21.3 | 0.00681 | 0.00726 | |
| | | 4055.2 | 15.9 | 0.00397 | 0.00555 | |
| 4036.3 w | | 4035.7 | 21.8 | 0.0144 | 0.0106 | |
| ~4007 vw sh | | 4003.7 | 19.1 | 0.00149 | 0.00342 | |
| 3980.3 vw | | 3980.0 | 15.8 | 0.000979 | 0.00278 | |
| 3951.2 vw br | | 3951.7 | 41.7 | 0.00460 | 0.00606 | |
| 3923.8 vw | | 3923.6 | 19.6 | 0.00142 | 0.00338 | $v_{31}+v_9$ (3922) or $v_{22}+v_{16}$ (3922) |
| 3909.9 vw | | 3909.5 | 15.6 | 0.000938 | 0.00275 | |

Table 5.3 - Continued

| Exp $\alpha_m^{"}$ a,b | Gas ^c | | Fitted | I_j^* | C, | $ \overrightarrow{R_j} $ | Assignment ^d |
|------------------------|------------------|----------|------------------|------------------|----------|--------------------------|---|
| cm ⁻¹ | | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| 3895.9 vw br db ? | | \ | 3897.7 | 17.9 | 0.000898 | 0.00269 | $v_{31}+v_{15}$ (3901) or v_2+v_{13} (3898) |
| | | Į | 3888.1 | 19,9 | 0.00102 | 0.00287 | v ₄₁ +v ₁₂ (3886) |
| 3869.7 vw | | | 3870.0 | 19.3 | 0.00167 | 0.00369 | v_{22} r v_{13} (3870) or v_{21} + v_{10} (3872) |
| 3847.2 vw br db | | \ | 3850.5 | 20.6 | 0.00149 | 0.00349 | |
| | | l | 3838.0 | 29.1 | 0.00302 | 0.00498 | |
| 3812.2 vw br | | | 3811.6 | 29.1 | 0.00203 | 0.00410 | |
| 3786.3 vw | | | 3785.7 | 17.1 | 0.00133 | 0.00333 | $v_2 + v_1$, (3785) |
| 3763.9 vw | | | 3762.3 | 23.1 | 0.000717 | 0.00245 | $v_{31}+v_{13}$ (3763) |
| ~3738 vw br sh | | | 3738.0 | 22.9 | 0.000461 | 0.00197 | |
| 3724.1 vw | | | 3724.4 | 16.7 | 0.00867 | 0.00856 | $v_{22}+v_{18}$ (3722) |
| 3707.0 vw | | | 3706.7 | 16.2 | 0.000797 | 0.00260 | $v_{21}+v_{29}$ (3708) or $v_{31}+v_{10}$ (3705) |
| 3675.0 vw | | | 3678.2 | 28.8 | 0.00162 | 0.00372 | v ₂ +v ₂₉ (3677) |
| 3649.0 vw | | | 3649.1 | 21.9 | 0.00344 | 0.00545 | $v_{31}+v_{17}$ (3650) or $v_{22}+v_{29}$ (3649) |
| 3624.0 vw | | | 3623.1 | 15.0 | 0.000935 | 0.00285 | |
| 3598.9 vw br | | | 3600.6 | 17.9 | 0.00346 | 0.00550 | $v_{21}+v_{11}$ (3607) |
| 3584.6 vw | | | 3585.1 | 15.1 | 0.000744 | 0.00256 | v_1+v_{11} (3583) |
| 3549.8 vw br | | | 3550.5 | 21.1 | 0.00963 | 0.00924 | |
| 3519.3 vw | | | 3518.5 | 13.6 | 0.00305 | 0.00523 | v ₂ +v ₁₉ (3519) |
| 3494.7 vw br | | | 3494.9 | 14.4 | 0.000099 | 0.00094 | |
| ~3472 vw br sh | | | 3471.4 | 14.2 | 0.000057 | 0.00072 | v _{34,37} +v ₁₁ (3471) |
| 3439.7 vw | | | 3439.4 | 21.6 | 0.00105 | 0.00310 | $v_{31} + v_{11}$ (3441) |
| ~3404 vw sh | | | 3397.5 | 20.1 | 0.000280 | 0.00161 | $v_2 + v_{30}$ (3401) or $v_1 + v_{30}$ (3408) |
| 3385.3 vw br | | | 3381.3 | 17.9 | 0.000376 | 0.00187 | $v_3 + v_{30}$ (3384) or $v_{31} + v_{19}$ (3384) |
| ~3325 vw br sh | | | 3330.0 | 10.2 | 0.000127 | 0.00110 | $v_{31}+v_{14}$ (3325) |
| 3306,5 vw | | | 3317.5 | 9.0 | 0.000079 | 0.00087 | $v_{22}+v_{20}$ (3303) |
| ~3203 vw br sh | | | | | | | 2v ₄ (3209) |
| 3167.5 vw | | | 3168.0 | 3.1 | 0.000184 | 0.00135 | 2v ₂₃ (3173) or v ₃₄ +v ₂₀ (~3167) |
| ~3115 w or sh | | | 3114.4 | 9.3 | 0.00335 | 0.00582 | |
| 3104.1 m | | | 3105.3 | 8.2 | 0.00878 | 0.00944 | v ₄ +v ₅ (3100) |
| 3086.4 s | В | | 3086.3 | 10.9 | 0.0617 | 0.0251 | v_{21} |
| 3062.1 s br | Α | | 3063.8 | 18.4 | 0.0982 | 0.0318 | v_1 |

Table 5.3 - Continued

| Exp a'' ab | Gas ^c | Fitted | <i>I</i> , | C, | $ \overrightarrow{R_j} $ | Assignment ^d |
|------------------|------------------|------------------|------------------|----------|--------------------------|---|
| cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| ~3055 s br sh | | 3054.0 | 19.9 | 0.0510 | 0.0229 | v_2 and ?? v_3 |
| 3027.0 s | В | 3027.3 | 19.0 | 0.335 | 0.0590 | V ₂₂ |
| ~3003 s sh | | 3002.4 | 15.0 | 0.0234 | 0.0157 | 3v ₉ (3007) |
| 2979.1 s br | | 2980.6 | 29.1 | 0.0709 | 0.0274 | v ₄ +v ₃₂ (2984) |
| ~2950 s sh | | 2950.1 | 39.4 | 0.167 | 0.0422 | ? v_{34} and ?? v_{37} |
| 2919.9 s | A?? { | 2920.7 | 21.8 | 0.149 | 0.0401 | \rightarrow \v_{31} |
| | 1 | 2916.1 | 11.5 | 0.0139 | 0.0123 | J * 31 |
| 2872.3 s vbr | | 2875.4 | 27.7 | 0.0825 | 0.0301 | v ₅ +v ₃₂ (2875) |
| | | 2860.2 | 21.4 | 0.0503 | 0.0235 | |
| ~2824 w br sh | | | | | | |
| ~2811w br sh | | 2808.9 | 8.5 | 0.000599 | 0.00259 | |
| ~2780 vw br sh | | | | | | v ₄ +v ₇ (2783) |
| 2734.1w | A ? | 2734.0 | 10.8 | 0.00897 | 0.0102 | v ₃₄ -v ₂₀ (2733) ?? |
| ~2692 vw br sh | | 2689.1 | 6.8 | 0.000051 | 0.00077 | v ₃₂ +v ₂₆ (2692) |
| 2671.7 vw | | 2671.6 | 6.5 | 0.000155 | 0.00135 | |
| ~2657 vw br sh | | 2656.0 | 10.6 | 0.000232 | 0.00166 | 2v ₂₅ (2664) |
| ~2644 vw br sh | | 2644.4 | 15.7 | 0.000578 | 0.00262 | v ₄ +v _{36,39} (2646) |
| 2631.9 vw | | 2632.0 | 10.2 | 0.000687 | 0.00287 | |
| ~2624 vw br sh | | 2624.1 | 14.1 | 0.000463 | 0.00236 | $v_{23} + v_{36,39}$ (2628) or $2v_{26}$ (2625) |
| ~2609 vw br sh | | 2609.3 | 15.9 | 0.00125 | 0.00388 | |
| 2604.7 vw | | 2603.9 | 8.0 | 0.000585 | 0.00266 | v ₄ +v ₉ (2607) |
| 2585.9 vw | A? | 2586.0 | 10.2 | 0.00331 | 0.00636 | |
| ~2576 vw | | 2576.4 | 17.5 | 0.000400 | 0.00221 | v ₅ +v ₂₈ (2577) |
| 2548.2 vw | | 2550.6 | 19.2 | 0.000819 | 0.00318 | $v_{23}+v_{12}$ (2553) |
| 2540.4 vw | | 2539.6 | 12.5 | 0.000795 | 0.00314 | $v_{38}+v_{28}$ (2542) or $v_{25}+v_6$ (2542) |
| ~2521 vw br sh | | 2521.8 | 15.4 | 0.000415 | 0.00228 | |
| 2509.2 vw | | 2509.4 | 10.5 | 0.000558 | 0.00837 | v ₂₅ +v ₇ (2511) |
| 2496.7 vw | | 2497.2 | 12.0 | 0.000434 | 0.00233 | |
| ~2490 vw br sh | | 2487.5 | 22.1 | 0.000918 | 0.00341 | |
| 2465.4 vw br db | | 2462.2 | 36.9 | 0.00277 | 0.00595 | |
| 2431.2 vw br | | 2432.6 | 19.4 | 0.00113 | 0.00383 | v ₂₃ +v ₁₃ (2429) |

Table 5.3 - Continued

| Exp α_{m}^{n} ab | Gas ^c | Fitted | Γ_{j} | C_j | $ \overrightarrow{R_i} $ | Assignment ^d |
|-------------------------|------------------|------------------|------------------|----------|--------------------------|---|
| cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| 2412.4 vw | | 2412.5 | 20.4 | 0.00382 | 0.00706 | |
| 2389.0 vw | | 2388.8 | 17.1 | 0.00216 | 0.00534 | |
| 2360.6 vw | | 2360.7 | 6.3 | 0.00106 | 0.00376 | $v_{25}+v_8$ (2362) or v_6+v_{27} (2366) |
| ~2354 vw br sh | | 2355.6 | 24.2 | 0.00319 | 0.00653 | |
| 2335.3 w | | 2335.5 | 10.3 | 0.00298 | 0.00634 | |
| 2312.6 vw | A? | 2312.6 | 10.4 | 0.00233 | 0.00563 | |
| 2286.1 vw db | | 2287.6 | 18.4 | 0.00175 | 0.00491 | |
| 2280.7 vw db | | 2279.6 | 12.5 | 0.000716 | 0.00315 | |
| 2260.5 vw | В | 2260.3 | 13.3 | 0.00176 | 0.00495 | $v_7 + v_{28}$ (2260) |
| ~2240 vw | | 2240.4 | 63.6 | 0.00214 | 0.00549 | $v_6 + v_8$ (2240) |
| 2237.3 vw | | 2236.6 | 6.8 | 0.000121 | 0.00131 | v ₂₇ +v ₂₈ (2237) |
| ~2225 vw br sh | | | | | | |
| 2207.5 vw | | 2207.7 | 7.1 | 0.000507 | 0.00269 | |
| 2185.3 vw br | В | 2186.0 | 13.4 | 0.00105 | 0.00389 | $v_{27} + v_8$ (2186) |
| 2163.7 vw | Α | 2163.4 | 10.1 | 0.00178 | 0.00509 | v ₃₂ +v ₁₀ (2165) |
| ~2138 vw sh | | | | | | |
| ~2123 vw sh | | | | | | |
| 2116.6 vw | | 2114.9 | 24.3 | 0.000698 | 0.00322 | |
| ~2111 vw sh | | | | | | |
| 2068.5 vw | | 2069.5 | 12.3 | 0.000356 | 0.00233 | |
| 2032.0 vw | | | | | | v ₈ +v ₉ (2032) |
| ~2014 vw sh | | 2014.6 | 7.7 | 0.000222 | 0.00186 | v ₅ +v ₁₁ (2017) |
| 2008.5 vw br | C? | 2008.2 | 8.9 | 0.000469 | 0.00271 | v ₈ +v ₁₅ (2011) |
| 1991.1 vw | Α | 1991.6 | 9.4 | 0.00157 | 0.00498 | v ₆ +v ₁₀ (1996) |
| 1983.6 vw br | C? | 1983.8 | 5.2 | 0.000334 | 0.00230 | v ₉ +v ₁₅ (1983) |
| ~1958 w sh | Α | 1959.4 | 14.6 | 0.00812 | 0.0114 | $v_7 + v_{10}$ (1964) or $2v_{15}$ (1960) |
| 1942.1m | В | 1942.1 | 16.7 | 0.0267 | 0.0208 | $v_{15}+v_{12}$ (1947) or $v_{27}+v_{16}$ (1942) |
| 1897.9 vw br | | 1899.9 | 4.0 | 0.000147 | 0.00156 | $v_9 + v_{16}$ (1898) or $v_{32} + v_{11}$ (1900) |
| 1872.0 w br | A | 1873.7 | 15.9 | 0.0108 | 0.0135 | v ₁₅ +v ₁₆ (1876) |
| 1857.6 m | В | 1857.3 | 14.3 | 0.0201 | 0.0185 | $v_{12}+v_{16}$ (1862) or $v_{25}+v_{11}$ (1853) |
| ~1820 w sh | В? | 1822.1 | 17.0 | 0.00367 | 0.00797 | v ₁₅ +v ₁₃ (1823) |

Table 5.3 - Continued

| Exp $\alpha_{\rm m}^{\prime\prime}$ a,b | Gas ^c | Fitted | Γ_{j} | С, | $ \overrightarrow{R_i} $ | Assignment ^d |
|---|------------------|------------------|------------------|----------|--------------------------|--|
| cm ⁻¹ | | cm ⁻¹ | cm ⁻¹ | km / moi | Debye | |
| 1802.6 m | Α | 1802.5 | 17.0 | 0.0215 | 0.0194 | |
| 1778.5 w br | | 1776.1 | 18.0 | 0.00358 | 0.00797 | $v_{27}+v_{29}$ (1778) or $3A_2$ options |
| 1735.6 w | | 1735.3 | 16.2 | 0.0110 | 0.0141 | $v_{16}+v_{13}$ (1738) |
| 1696.8 w | | 1697.4 | 9.3 | 0.000968 | 0.00424 | |
| 16 7 6.7 w | | 1675.8 | 18.3 | 0.00197 | 0.00609 | |
| 1657.8 w br | | | | | | |
| 1623.1 m br | Q? | 1623.6 | 12.7 | 0.0117 | 0.0151 | $v_{16}+v_{17}$ (1625)? |
| 1604.6 s | A { | 1604.9 | 4.7 | 0.0319 | 0.0250 | } _{v4} |
| 1001.00 | [| 1602.8 | 7.4 | 0.0249 | 0.0221 | |
| 1586.7 m br | | 1590.0 | 26.9 | 0.0207 | 0.0203 | V ₂₃ |
| 1572.1 m | Q? | 1571.4 | 16.2 | 0.0141 | 0.0168 | $2v_{10}$ (1571) or $v_{13}+v_{17}$ (1573) |
| 1550.3 m br | Q? | | | | | v_8+v_{11} (1551) or v_6+v_{30} (1556) |
| ~1537 m br sh | | 1530.0 | 45.1 | 0,0358 | 0.0271 | $v_{13}+v_{18}$ (1538) |
| 1523.6 s br | B?? | 1523.5 | 5.2 | 0.00571 | 0.0109 | $v_7 + v_{30}$ (1525) |
| 1495.7 vs | A . | 1495.7 | 4.0 | 0.113 | 0.0488 | } _{v5} |
| | | 1494.3 | 10.5 | 0.0467 | 0.0314 | J., |
| 1460.3 s br | | 1459.8 | 31.9 | 0.250 | 0.0734 | v_{35} ??, v_{38} ?? and v_{24} ?? |
| ~1422 m sh | | 1419.7 | 18.8 | 0.00653 | 0.0120 | $v_{17}+v_{18}$ (1425) or v_6+v_{20} (1427) |
| | | | | | | or $v_{28}+v_{20}$ (1427) |
| 1378.9 s | A ∫ | 1382.3 | 44.4 | 0.0356 | 0.0285 | } _{v32} |
| 15 / 5 5 | [| 1379.0 | 7.1 | 0.0170 | 0.0197 | J * 32 |
| 1332.0 w | B ? | 1332.0 | 10.0 | 0.00174 | 0.00641 | V ₂₅ |
| 1312.7 w | В | 1312.7 | 10.0 | 0.00331 | 0.00891 | V ₂₆ |
| ~1302 w sh | | 1300.7 | 12.2 | 0.00127 | 0.00555 | v ₁₆ +v ₁₄ (1300) |
| 1277.6 w br | | 1277.6 | 22.6 | 0.00236 | 0.00763 | v ₅ -v ₂₀ (1279) |
| 1248.7 w | | 1248.2 | 22.0 | 0.00593 | 0.0122 | |
| 1210.2 m | A | 1210.1 | 6.3 | 0.00480 | 0.0112 | V ₆ |
| ~1193 w br sh | Q? | 1193.4 | 18.7 | 0.00602 | 0.0126 | $v_{17}+v_{19}$ (1194) or $v_{15}+v_{20}$ (1198) |
| 1178.6 s | | 1178.6 | 5.7 | 0.0103 | 0.0166 | V ₇ |
| 1155.9 m | | 1156.8 | 31.5 | 0.0219 | 0.0244 | V ₂₇ |
| ~1130 w br sh | | | | | | $v_{10}+v_{30}$ (1132) or $v_{17}+v_{14}$ (1135) |

Table 5.3 - Continued

| Exp $\alpha_m^{"}$ ab | Gas ^c | Fitted | $\Gamma_{\!\scriptscriptstyle J}$ | C, | $ \overrightarrow{R_i} $ | Assignment ^d |
|-----------------------|------------------|------------------|-----------------------------------|----------|--------------------------|--|
| c m -1 | | cm ⁻¹ | cm ⁻¹ | km / mol | Debye | |
| 1106.5 m | В | 1106.5 | 9.4 | 0.0102 | 0.0170 | v ₁₈ +v ₁₄ (1100) |
| 1081.4 s | В | 1081.1 | 8.6 | 0.0521 | 0.0390 | V ₂₈ |
| 1041.4 s | ?? { | 1042.6 | 11.4 | 0.0145 | 0.0209 | v ₃₆ ?? and v ₃₀ ?? |
| 2012.10 | [| 1040.2 | 37.2 | 0.0604 | 0.0428 | 30 and v ₃ y |
| 1030.1 s | A | 1030.0 | 3.9 | 0.0261 | 0.0283 | v_8 |
| 1002.3 m | | 1001.5 | 10.8 | 0.00394 | 0.0111 | V9 |
| 980.7 m | | 980.9 | 23.2 | 0.0115 | 0.0192 | v ₁₅ ?? |
| 966.4 m br | | 964.2 | 15.7 | 0.00591 | 0.0139 | v ₁₂ ?? |
| ~947 w br sh | | 947.4 | 17.7 | 0.00559 | 0.0136 | $v_{17} + v_{20}$ (947) |
| 929.6w | | 929.0 | 12.1 | 0.00366 | 0.0111 | 2 ₁₉ (929) or v ₁₁ +v ₁₄ (926) |
| ~910 w br sh | | 910.4 | 13.9 | 0.00211 | 0.00854 | $v_{18}+v_{20}$ (912) |
| 895.4 s | С | 895.4 | 8.6 | 0.0126 | 0.0211 | v_{16} |
| 872.9 w | | 872.3 | 14.2 | 0.00264 | 0.00976 | v ₁₉ +v ₁₄ (869) or 4v ₂₀ (868) |
| 842.7 m | | 842.8 | 17.4 | 0.00783 | 0.0171 | v_{l3} |
| 810.4 w br sh | | | | | | $v_{19}+v_{30}$ (810) or $2v_{14}$ (811) |
| 785.6 s | Α | 785.5 | 3.7 | 0.00337 | 0.0116 | v_{10} |
| 729.9 vs | С | 729.9 | 6.7 | 0.777 | 0.183 | v_{17} |
| | ſ | 694.9 | 4.1 | 0.189 | 0.0926 | |
| 694.8 vs | c { | 693.1 | 3.2 | 0.0151 | 0.0262 | v_{18} |
| | Į | 690.9 | 4.2 | 0.0115 | 0.0229 | |
| ~678 s br sh | | 677.5 | 6.8 | 0.00857 | 0.0200 | v ₁₉ +v ₂₀ (681) |
| 633.0 w sh | | | | | | |
| 622.0 m | | 621.9 | 6.6 | 0.00161 | 0.00903 | V ₂₉ |
| 565.2 vw br | | | | | | v ₃₀ +v ₂₀ (563) |
| 537.8 w | | 537.5 | 4.5 | 0.000342 | 0.00448 | |
| 521.0 m | Α | 521.0 | 4.4 | 0.00297 | 0.0134 | v_{11} |
| ~508 vw sh | | 509.6 | 12.7 | 0.000341 | 0.00459 | |
| 464.4 vs | С | 464.4 | 4.1 | 0.110 | 0.0864 | V ₁₉ |

a - Wavenumbers of peaks in the imaginary molar polarizability spectrum of liquid toluene.

b - Abbreviations used: v-very, w-weak, m-medium, s-strong, br-broad, sh-shoulder and db-doublet.

c - Band contours in the infrared spectrum of the gas.

d - Calculated wavenumbers are given in brackets.

and 440 cm⁻¹, it is greater by 2.8% than that of the α_m'' spectrum. Overall, the area under the fitted spectrum is greater by 1.5% than that under the α_m'' spectrum.

The parameters of the 166 CDHO bands required to fit the α_m'' spectrum of toluene are given in Table 5.3. The table is arranged similarly to Table 5.1 of chlorobenzene. The accuracy of the integrated intensities, C_j , is estimated from the sum of the accuracy of the fit and the accuracy of the real and imaginary refractive indices, n and k, that are used in the calculation of the α_m''' spectrum of toluene. The accuracy of the refractive indices was reported in Chapter 2 and is 0.2% for n and on the order of 2 to 3% for k. The accuracy of the fit is about 0.5 to 3% depending on the intensity of the band. Therefore, the accuracy of C_j is estimated to be better than 3-5% for strong peaks and 5-10% for weak peaks.

5.2.1 - Integrated intensities and dipole moment derivatives of the fundamental vibrations of liquid toluene

The intensities of the bands fitted to the experimental α_m'' spectrum are assigned in the following sub-sections to the fundamental vibrations of toluene. Table 5.4 summarized integrated intensities, C_j , of the fundamental vibrations and the dipole moment derivatives with respect to the normal coordinates, $|\mu_j|$, calculated from them through Eq. 1.4.4. The estimated errors in the intensities are discussed above, and the errors in the dipole moment derivatives are estimated to be about 2% for strong fundamentals and about 4% or more for weak ones. In some cases it will be clear that

the uncertainty in the assignment is considerable, making the estimated error in the values potentially very large.

The only integrated intensities of toluene reported in the literature are those published by Boggs²⁶ and by Galabov²⁹. Boggs et al. used scaled ab initio calculations to obtain wavenumbers and infrared integrated intensities, $A_{l,ab,m}$, of the fundamental vibrations of toluene. The latter are given in column 6 of Table 5.4. Galabov et al. used ab initio calculation to obtain the sign of the dipole moment derivatives with respect to symmetry coordinates but refined their calculated intensities through a least squares refinement procedure using experimental integrated intensity data for 4 bands of benzene, 12 bands of toluene and 16 bands of toluene-d₈. Galabov used an empirical force field developed for methyl benzenes by La Lau and Snyder¹⁹, so, in contrast to Boggs, their wavenumbers and eigenvectors were not determined from ab initio calculations. The problems of separating the contributions of overlapping bands to the total intensity are more severe for the generally broader gas-phase bands than for the liquid-phase bands. Galabov et al. acknowledged the problem, but gave no information that allows the error of their experimental intensities to be estimated. Galabov's computed, $A_{j,ab in}$, and experimental, $A_{j,exp}$, intensities are given in columns 7 and 8 of Table 5.4. For comparison, our integrated intensities of the gas, $A_{l,gas}$, were calculated from the C_i of the liquid using Eq. 1.5.4 under the assumption that the $|\mu_i|$ values are the same in the gas and in the liquid phases, and are given in column 5 of Table 5.4.

Table 5.4 - Dipole moment derivatives and integrated intensities of toluene

| | | This v | vork | | Boggs ²⁶ | Galabov ²⁹ | | |
|-----------------|---------------------------|--------------------|---------------------------|------------------------|--------------------------|-----------------------|----------------------|--|
| ν, | Fitted cm ^{-1 a} | $C_{j,lig}$ b | <i>µ</i> _j c | A _{J.gas} b,d | A _{J,ab in} b.e | Aj,ab in b.f | A _{j,exp} b | |
| $\mathbf{v_1}$ | 3063.8 | 0.0982 | 0.428 | 7.75 | 14.52 | 0.49 | g | |
| V ₂ | 3054.0 | 0.0510 | 0.309 | 4.03 | 14.62 | 4.03 | 26.45 ⁸ | |
| ν_3 | (3038 ???) h | 0.0 | 0.0 | 0.0 | 5.57 | 3.40 | g | |
| V ₄ | 1604.9 1 | 0.0568 | 0.460 | 4.49 | 6.31 | 0.61 | 7.26 ^J | |
| ν ₅ | 1495.7 1 | 0.160 | 0.755 | 12.61 | 15.95 | 11.50 | 14.87 | |
| ν ₆ | 1210.1 | 0.00480 | 0.095 | 0.38 | 0.03 | | | |
| V ₇ | 1178.6 | 0.0103 | 0.139 | 0.81 | 0.22 | | | |
| v_8 | 1030.0 | 0.0261 | 0.221 | 2.06 | 1.41 | 2.30 | | |
| V9 | 1001.5 | 0.00394 | 0.086 | 0.31 | 0.13 | | | |
| v_{10} | 785.5 | 0.00337 | 0.079 | 0.27 | 0.35 | 0.72 | k | |
| v_{11} | 521.0 | 0.00297 | 0.074 | 0.23 | 0.77 | 80.0 | e . | |
| v ₁₂ | 964.2 | 0.00591 | 0.105 | 0.47 ^m | 0.00 | | | |
| v_{13} | 842.7 | 0.00783 | 0.121 | 0.62 ^m | 0.00 | | | |
| v_{14} | (405 ?) | | | | 0.00 | | | |
| V ₁₅ | 980.9 | 0.0115 | 0.147 | 0.91 | 0.22 | | | |
| V ₁₆ | 895.4 | 0.0126 | 0.153 | 0.99 | 0.39 | | | |
| V ₁₇ | 729.9 | 0.777 | 1.205 | 61.35 | 47.18 | 34.48 | 40.96 ^k | |
| v_{18} | 694.9 1 | 0.216 | 0.909 | 17.02 | 27.99 | 18.18 | k | |
| V _{l9} | 464.4 | 0.110 | 0.453 | 8.69 | 8.47 | 6.00 | 6.24 ' | |
| V ₂₀ | (217) | | | | 2.48 | | | |
| v ₂₁ | 3086.3 | 0.0617 | 0.340 | 4.87 | 43.01 | 19.16 | 38.18 ⁸ | |
| V ₂₂ | 3027.3 | 0.335 | 0.791 | 26.45 | 3.61 | 31.85 | 8 | |
| V ₂₃ | 1590.0 | 0.0207 | 0.197 | 1.63 | 0.83 | 0.95 | J | |
| V ₂₄ | 1459.8 ⁿ | 0.250 ⁿ | 0.683 ⁿ | 19.74 ⁿ | 14.56 | 1.16 | 6.54 ° | |
| V ₂₅ | 1332.0 | 0.00174 | 0.057 | 0.14 | 0.00 | 0.02 | P | |
| V ₂₆ | 1312.7 | 0.00331 | 0.079 | 0.26 | 0.03 | | | |
| V ₂₇ | 1156.8 | 0.0219 | 0.202 | 1.73 | 0.20 | | | |
| V28 | 1081.1 | 0.0521 | 0.312 | 4.11 | 3.70 | 3.52 | 4.30 | |
| V ₂₉ | 621.9 | 0.00161 | 0.055 | 0.13 | 0.09 | 0.01 | k | |
| V ₃₀ | (346) | | | | 0.42 | | | |

Table 5.4 - Continued

| | | This | Boggs ²⁶ | Galabo | ov ²⁹ | | |
|-----------------|---------------------------|--------------------|---------------------|------------------------|------------------|-------------|--------------------|
| <u></u> | Fitted cm ⁻¹ a | $C_{j,lig}$ b | 141° | A _{s.gas} b.d | Anab in be | Ajab in b.f | A,exp |
| v_{31} | 2920.7 | 0.149 | 0.528 | 11.76 | 27.05 | 10.42 | 11.05 |
| V ₃₂ | 1382.3 1 | 0.0526 | 0.436 | 4.15 | 0.39 | 3.26 | 3.08 ^p |
| v_{33} | (<100) | | | | | | |
| V34 | 2950.1 ^q | q | q | q | 28.25 | 5.50 | 18.52 ^r |
| V ₃₅ | 1459.8 ⁿ | n | n | n n | 5.15 | 2.45 | o |
| V ₃₆ | 1042.6 * | 0.0145 3 | 0.165 3 | 1.14 * | 9.24 | | |
| V ₃₇ | 2950.1 ^q | 0.167 ^q | 0.559 ^q | 13.19 ^q | 18.46 | 12.94 | t |
| V ₃₈ | 1459.8 ⁿ | n | n n | ⁿ | 0.15 | 3.78 | o |
| V ₃₉ | 1040.2 3 | 0.064 * | 0.336 * | 4.77 * | 0.02 | | |

- a Integrated intensities were not obtained for v_{14} , v_{20} , v_{30} and v_{33} which are below the experimental wavenumber range for this work.
- b Units are km mole⁻¹.
- c Units are Debye Å⁻¹ amu^{-1/2}.
- d Integrated intensities for the gas, $A_{j,gas}$ were calculated as $A_{j,gas} = 8\pi^2 C_{j,liq}$ with the assumption that $\mu_{j,liq}^2 = \mu_{j,gas}^2$.
- e $A_{j,gas}$ values obtained from ab initio calculations reported in Ref. 26.
- f $A_{j,gas}$ values reported in Ref. 29. Intensities were obtained from *ab initio* calculations refined with experimental intensities of 4 bands of benzene, 12 bands in toluene and 16 bands in toluene- d_8 .
- g Total aromatic CH stretching intensity is 64.63 km mole⁻¹. Galabov gave the two values 26.45 and 38.18 given in the table but did not say how the separation was achieved.
- h This vibration has not been assigned with confidence.
- i More than one CDHO band was used in the fit. The wavenumber listed is that of the prominent contributing band. All related quantities for this vibration are the sums from all contributing bands.
- j Experimental $A_{j,gas}$ was given for v_4 and v_{23} collectively.
- k Experimental $A_{j,gas}$ was given for v_{10} , v_{17} , v_{18} and v_{29} collectively.
- ℓ Experimental $A_{J,gas}$ was given for v_{11} and v_{19} collectively.
- m A_2 vibrations in chlorobenzene are infrared inactive in the spectrum of the gas, i.e $A_{j,gas} = 0$. Our values are derived from $C_{j,liq}$ as described in footnote d.
- n The same wavenumber was assigned to v_{24} , v_{35} and v_{38} . All quantities are assigned in this table to v_{24} .
- o Experimental $A_{j,gas}$ was given for v_{24} , v_{35} and v_{38} collectively.

- **p** Experimental $A_{j,gas}$ was given for v_{25} and v_{32} collectively.
- q The same wavenumber was assigned to v_{37} and v_{34} . All quantities are assigned to v_{37} in this table.
- r The value given for v_{34} includes the intensity of and v_{37} .
- s 1041 cm⁻¹ was assigned to both v_{36} and v_{39} . The intensities of the fitted bands at 1042.6 and 1040.2 cm⁻¹ were assigned arbitrarily to v_{36} and v_{39} respectively.

-5.2.1a - Integrated intensities of the CH stretching vibrations

The experimental α_m''' and curve fitted α_m''' spectra of liquid toluene in the region of the CH stretching vibrations are shown in Figure 5.10. The fit in this region is very good. The area under the curve fitted α_m'' is smaller by 0.1% than the area under the experimental α_m''' spectrum. Peak heights are reproduced on average to 0.6% and no fundamental exceeds 1.1%.

The existence of many relatively intense overtone and combination bands in the CH stretch region again makes it impossible to make a unique assignment of the intensity to the different normal vibrations. The possible assignments are presented and discussed below, starting with the assignment of the intensity of specific fitted bands to specific fundamentals.

The intensity assignments of four of the five aromatic CH stretching vibrations, v_{21} , v_1 , v_2 and v_{22} are straightforward. The intensities of the single CDHO bands at 3086, 3064, 3055 and 3027 cm⁻¹, are assigned to these vibrations, respectively. In section 4.5.1, v_3 was assigned to a Raman band at 3038 cm⁻¹ with no infrared counterpart identified. No additional CDHO band was needed to improve the fit in this region, and no integrated intensity is assigned to this vibration.

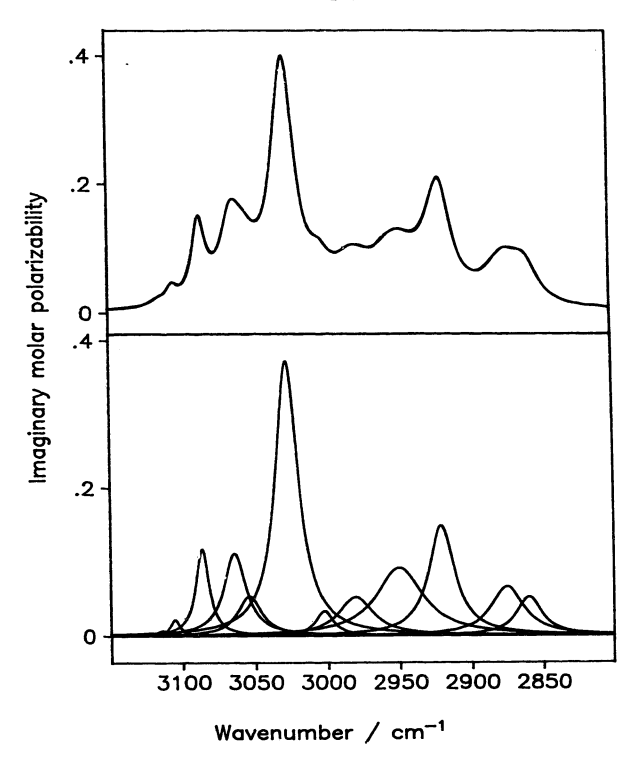


Figure 5.10 - Top - Superimposed experimental imaginary molar polarizability, α_m^m , and curve fitted α_m^m spectra of liquid toluene between 3150 and 2800 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m^m spectrum. Units for both boxes are cm³ mole⁻¹.

The combined intensities, $A_{j,gas}$, of the aromatic CH stretching vibrations, calculated from the assigned $C_{j,liq}$ is 43.1 km mole⁻¹. Galabov's²⁹ combined experimental value, $A_{j,exp}$, is 64.6 km mole⁻¹, about 50% greater than our value. Boggs'²⁶ value, $A_{j,ab\ in}$, of 81.3 km mole⁻¹ is about a factor of 2 larger than our value.

The intensity of the fitted band at 2920.7 cm⁻¹ is assigned to the symmetric CH₃ stretching vibration. This yields an $A_{J,gas}$ value of 11.76 km mole⁻¹ which is within 6% of Galabov's experimental value for the gas, 11.05 km mole⁻¹. Boggs' calculated value²⁶, $A_{J,ab\ in}$ is 27.05 km mole⁻¹, 2.3 times greater than either of these experimental values.

The assignment of the antisymmetric CH₃ stretching vibrations v_{34} and v_{37} is very uncertain. They were both tentatively assigned in section 4.5.5 to the broad peak at 2950 cm⁻¹. This peak was fitted by a single band of intensity $C_j = 0.167$ km mole⁻¹, which is equivalent to $A_{j,gas} = 13.2$ km mole⁻¹. Galabov's combined experimental value for these two vibrations is 18.5 km mole⁻¹ and Boggs' combined *ab initio* value is 46.7 km mole⁻¹. These values are greater than our value by 40% and by a factor of 3.5, respectively. Under this assignment the intensities $A_{j,gas}$ of the aromatic and aliphatic CH stretching vibrations are 43.1 and 25.0 km mole⁻¹, respectively, while those of Galabov are 64.6 and 29.5 km mole⁻¹ and those of Boggs 81.3 and 73.7 km mole⁻¹. The agreement is not impressive.

A second assignment is to note that the CDHO band at 2980.6 cm⁻¹ is assigned in Table 5.3 to the combination transition v_4+v_{32} because the polarized band in the Raman spectrum excludes it from being assigned as a B₁ or a B₂ vibration (section 4.5.5). If,

however, its intensity is added to that of the 2950 cm⁻¹ band, the combined $A_{p,gas}$ is then 18.78 km mole⁻¹, within 1% of Galabov's experimental value. Boggs' value is then greater by a factor of 2.5 than our value, which is similar to the discrepancy observed in the aromatic CH stretching vibrations.

A third alternative assignment, and possibly the most probable, is to assign to the CH stretching fundamentals all of the intensity in the region where the overtone and combination bands are unusually strong, namely between 3200 and 2775 cm⁻¹. This intensity is $C_J = 1.08$ km mole⁻¹ which transforms to $A_{J,gas} = 84.1$ km mole⁻¹. The sum of all of the intensity in this region reported by Galabov or Boggs is 94.2 or 155 km mole⁻¹, respectively, but there is no indication in the case of Galabov whether he has removed any of the experimental intensity as due to overtone or combination bands.

In a somewhat arbitrary attempt to refine this assignment by separating the aromatic and aliphatic CH stretching intensities, the total area under the spectrum was divided at 3000 cm⁻¹ to obtain $\Sigma C_j = 0.581$ km mole⁻¹ between 3200 and 3000 cm⁻¹ and 0.498 km mole⁻¹ between 3000 and 2775 cm⁻¹. These values convert to $\Sigma A_j = 45.9$ km mole⁻¹ for the aromatic CH and 39.3 km mole⁻¹ for the aliphatic CH, which compares with 43.1 for the aromatic and 25.0 for the aliphatic under the first assignment. Clearly, the first and third assignments are not very different for the aromatic CH stretching vibrations but the third assignment increases the intensity of the aliphatic CH stretches by 56%.

These assignments of the CH stretching vibrations are discussed again later in a

comparison of the intensities of different molecules.

5.2.1b - Integrated intensities of the v_4 , v_5 , v_{23} , v_{24} , v_{32} , v_{35} and v_{38} vibrations

The experimental α_m'' and curve fitted α_m'' spectra of liquid toluene between 1650 and 1350 cm⁻¹ are shown in Figure 5.11. The fit in this region is very good. The area under the curve fitted α_m'' is smaller by 0.36% than the area under the experimental α_m'' spectrum. Peak heights are reproduced on average to 0.8% and no fundamental exceeds 1.0%.

The intensities of the two CDHO bands at 1604.9 and 1602.8 cm⁻¹ are assigned to v_4 , while that of the CDHO band at 1590.0 cm⁻¹ is assigned to v_{23} . The combined intensity, $A_{j,gas}$, of v_4 and v_{23} is 6.12 km mole⁻¹. Galabov's²⁹ combined experimental value, $A_{j,exp}$, is 7.26 km mole⁻¹, about 19% greater than our value, while Boggs'²⁶ value, $A_{j,ab\ m}$, of 7.14 km mole⁻¹ is about 17% greater than our value.

The intensities of the two CDHO bands at 1495.7 and 1494.3 cm⁻¹ are assigned to v_5 and combine to give $A_{j,gas} = 12.61$ km mole⁻¹. It is smaller by 18% than Galabov's reported experimental value of 14.87 km mole⁻¹, and 26% smaller than Boggs'²⁶ computed $A_{j,ab~in}$ of 15.95 km mole⁻¹.

In sections 4.5.4 and 4.5.5, v_{35} and v_{38} were both tentatively assigned at 1460 cm⁻¹, with v_{24} in their proximity, perhaps at 1442 cm⁻¹. Only one CDHO band was required to obtain a good fit of the strong broad band at 1460 cm⁻¹ in the α_m'' spectrum, and its intensity is assigned to these three vibrations. Our calculated, $A_{j,gas}$, is 19.74 km

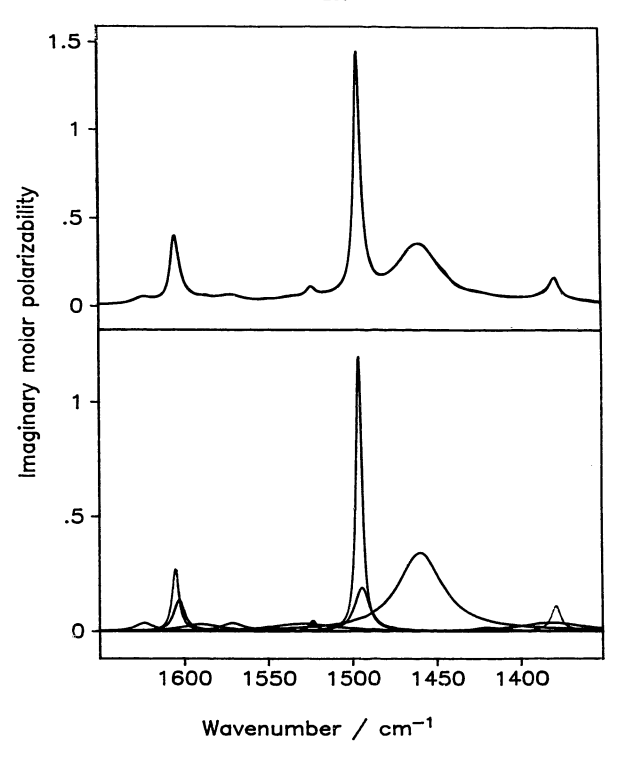


Figure 5.11 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid toluene between 1650 and 1350 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Units for both boxes are cm³ mole⁻¹.

km mole⁻¹, in good agreement with Boggs⁻²⁶ computed value of 20.08 km mole⁻¹, but far larger than Galabov's²⁹ experimental value, 6.54 km mole⁻¹.

The intensities of the two CDHO bands at 1382.3 and 1379.0 cm⁻¹ are assigned to the symmetric CH₃ deformation v_{32} . Galabov²⁹ reported the combined experimental intensity, $A_{J,exp}$, of v_{32} and v_{25} (1332 cm⁻¹) as 3.08 km mole⁻¹. Boggs⁻²⁶ computed intensities for these vibrations are 0.39 and 0.00 km mole⁻¹. The intensity assigned to v_{25} is (next section) $C_J = 0.00174$ km mole⁻¹, i.e. $A_{J,gas} = 0.14$ km mole⁻¹. Thus, for v_{32} and v_{25} combined we find $A_{J,gas} = 4.29$ km mole⁻¹, which is 40% greater than Galabov's experimental value and 11 times greater than Boggs' reported values. It is noteworthy that based on our work and the theoretical values of Boggs and Galabov, the intensity of v_{32} is much greater than that of v_{25} , and therefore Galabov's $A_{J,exp}$ of 3.08 km mole⁻¹ is largely due to v_{32} rather than to v_{25} .

5.2.1c - Integrated intensities of the v_6 , v_7 , v_8 , v_9 , v_{25} , v_{26} , v_{27} , v_{28} , v_{36} and v_{39} vibrations

The experimental α_m'' and curve fitted α_m'' spectra of liquid toluene between 1350 and 990 cm⁻¹ are shown in Figure 5.12. The fit in this region of weak absorption is not as good as in other regions. The area under the curve fitted α_m'' is smaller by 1.6% than the area under the experimental α_m'' spectrum. Peak heights are reproduced on average to 3.1%.

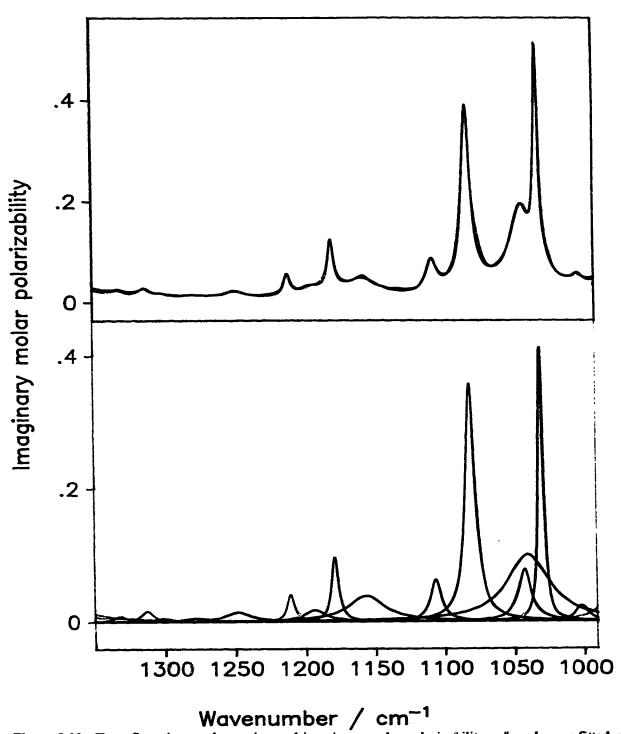


Figure 5.12 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid toluene between 1350 and 990 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Strong contributions from bands outside the region can be seen as tails on both ends of the region. Units for both boxes are cm³ mole⁻¹.

The intensity assignments of v_{25} , v_{26} , v_6 , v_7 , v_{27} , v_{28} , v_8 and v_9 are straightforward. They are assigned the intensities of the single CDHO bands at 1332, 1313, 1210, 1179, 1156, 1081, 1030 and 1002 cm⁻¹, respectively. In section 4.5.5, v_{36} and v_{39} were both assigned to the same wavenumber at 1040 cm⁻¹. Two CDHO bands were used in the fit, one at 1042.6 and a sharper band at 1040.2 cm⁻¹. Their intensities are arbitrarily assigned to v_{36} and v_{39} , respectively. Their combined intensity will be compared with literature values.

The experimental intensity of v_{25} was discussed with that of v_{32} in section 5.2.1b. Galabov gave No experimental $A_{j,exp}$ are given for v_6 , v_7 , v_9 , v_{26} , and v_{27} . Boggs²⁶ reported *ab initio* values are smaller than our $A_{j,gas}$ by factors of 12.7, 3.7, 2.4, 8.7 and 8.7, respectively for these weak vibrations.

 $A_{j,gas}$ of v_{28} , is 4.11 km mole⁻¹ and is in good agreement with Galabov's²⁹ $A_{j,exp}$ of 4.30 km mole⁻¹. Boggs'²⁶ calculated value is 3.70 km mole⁻¹, about 11% smaller than our value.

Galabov²⁹ gives the combined intensity of v_{36} , v_{39} and v_8 as 2.67 km mole⁻¹. Our $A_{J,gas}$ values are 5.91 km mole⁻¹ for v_{36} and v_{39} combined and 3.06 for v_8 , for a total of 7.97 km mole⁻¹. Boggs²⁶ ab initio</sup> values for these vibrations are 9.26 and 1.41 km mole⁻¹ for a total of 10.67 km mole⁻¹, about 35% greater than our total value.

5.2.1d - Integrated intensities of the ν_{10} , ν_{12} , ν_{13} , ν_{15} and ν_{16} vibrations

The experimental α_m''' and curve fitted α_m''' spectra of liquid toluene between 990 and 750 cm⁻¹ are shown in Figure 5.13. The fit in this region is not as good as in other regions probably because the absorption is extremely weak. The area under the curve fitted α_m''' is greater by 3.8% than the area under the experimental α_m''' spectrum. The large difference which can be seen in Fig 5.13 at the low wavenumber edge of the region, is mainly due to the strong bands at 730 and 695 cm⁻¹. Peak heights are reproduced on average to 1.5% with only one peak, that of v_{10} at 786 cm⁻¹, exceeding this value with 4.3%.

The intensities of the single CDHO bands at 981, 964, 895, 843 and 786 cm⁻¹, are assigned to v_{15} , v_{12} , v_{16} , v_{13} and v_{10} , respectively. v_{12} and v_{13} are A_2 vibrations and as such are inactive in the infrared spectrum of the gas, so their intensities were not given by Galabov²⁹ and were given as 0.00 by Boggs²⁶.

The $A_{J,gas}$ values for the two B₁ vibrations, v_{15} and v_{16} , are about equal at 0.91 and 0.99 km mole⁻¹. Boggs reported *ab initio* values that are much smaller and less equal, namely 0.02 and 0.39 km mole⁻¹ for v_{15} and v_{16} .

 $A_{j,gas}$ for v_{10} is 0.27 km mole⁻¹ and, considering the weakness and the far from perfect fit of this band (FIg. 5.13) is in good agreement with Boggs' $A_{j,ab\ m}$ value, 0.35 km mole⁻¹.

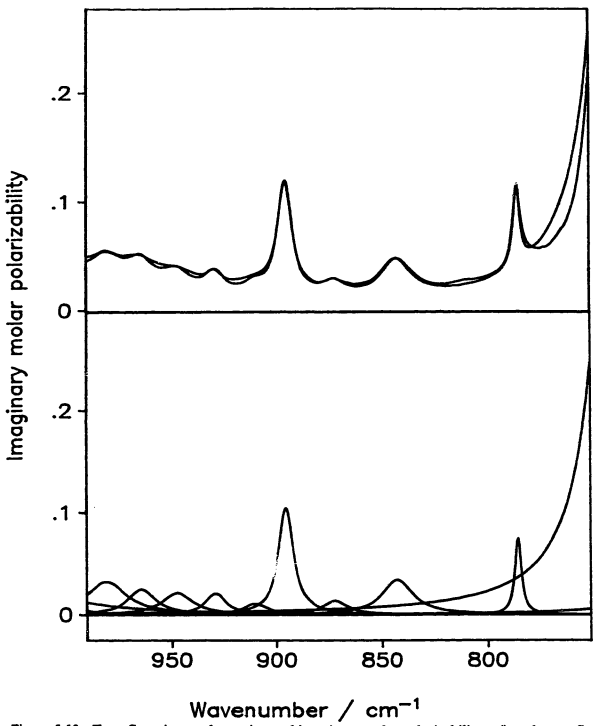


Figure 5.13 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid toluene between 990 and 750 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Strong contributions from the strong bands at 730 and 695 cm⁻¹ is clearly visible. Units for both boxes are cm³ mole⁻¹.

5.2.1e - Integrated intensities of the ν_{17} and ν_{18} vibrations

The experimental α_m'' and curve fitted α_m'' spectra of liquid toluene between 750 and 650 cm⁻¹ are shown in Figure 5.14. The fit in this region is good except for the wings of the strong band at 730 cm⁻¹. Here, and elsewhere, the experimental bands fall off slightly faster than the CDHO bands, undoubtedly due to a small Gaussian contributions. This accounts for the area under the fitted curve being greater by 2.1% than the area under the experimental α_m'' spectrum. The peak height of v_{17} is reproduced to 1.2% and that of v_{18} to 0.1%.

Three CDHO bands, all within 5 cm⁻¹, were needed to fit the strong band at 695 cm⁻¹ and their combined intensity is assigned to v₁₈. Only one CDHO band was needed for the stronger band at 730 cm⁻¹. The fit in the wings was not improved by the addition of extra CDHO bands.

 $A_{j,gas}$ for v_{17} is 61.35 km mole⁻¹ and for v_{18} is 17.02 km mole⁻¹ (Table 5.4). Boggs⁻²⁶ ab initio</sup> intensities for these two vibrations are 47.18 and 27.99 km mole⁻¹, respectively. Although the values for each vibration are different the combined intensities differ by less than 5% (78.37 to 75.17 km mole⁻¹). Galabov's²⁹ experimental value is 40.96 km mole⁻¹ for v_{17} and v_{18} combined and also includes the intensities of v_{10} (at 785 cm⁻¹) and v_{29} (at 622 cm⁻¹) as well, although according to his theoretical calculations the calculated results of his fitting procedure indicated that their contributions are small.

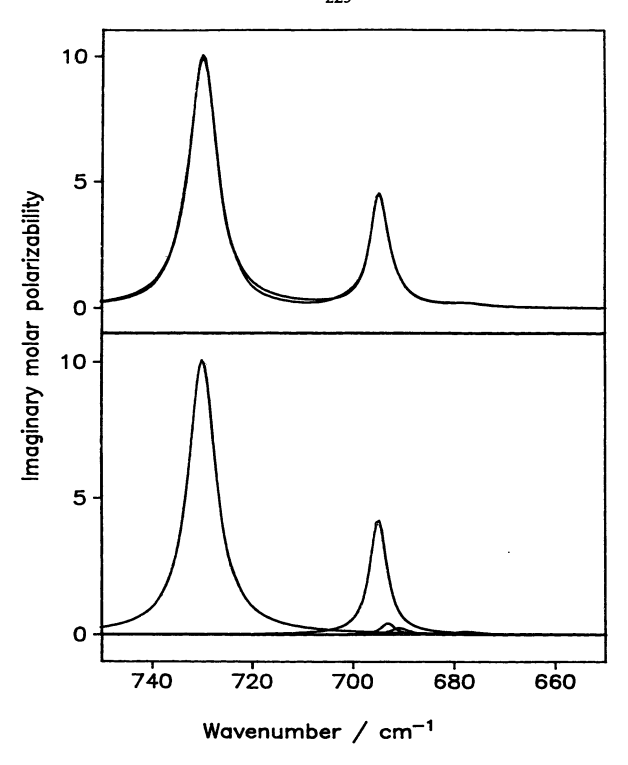


Figure 5.14 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid toluene between 750 and 650 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Units for both boxes are cm³ mole⁻¹.

5.2.1f - Integrated intensities of the v_{11} , v_{14} , v_{19} , v_{20} , v_{29} , v_{30} and v_{33} vibrations

The experimental α_m'' and curve fitted α_m'' spectra of liquid toluene between 650 and 440 cm⁻¹ are shown in Figure 5.15. The area under the curve fitted α_m'' is greater by 3.8% than the area under the experimental α_m'' spectrum. Peak heights of fundamentals are reproduced on average to 1.05%.

The wavenumbers of v_{14} , v_{20} , v_{30} and v_{33} are below the experimental range studied in this work, and, thus, no intensities were obtained for them.

The intensity of the CDHO band at 621.9 cm⁻¹ is assigned to v_{29} . Its $A_{J,gas}$ is 0.13 km mole⁻¹, which is small but in good agreement with Boggs⁻²⁶ ab initio value of 0.09 km mole⁻¹.

The intensity of the CDHO band at 521.0 cm⁻¹ is assigned to v_{11} . Its $A_{j,gas}$ value of 0.23 km mole⁻¹ is about 3 times smaller than Bcggs²⁶ value 0.77 km mole⁻¹. The intensity of the CDHO band at 464.4 cm⁻¹ is assigned to v_{19} . Its $A_{j,gas}$ of 8.69 km mol⁻¹ is in good agreement with Boggs' reported $A_{j,ab\ m}$ of 8.47 km mole⁻¹. Galabov²⁹ gives the combined experimental intensity of v_{11} and v_{19} as 6.24 km mole⁻¹.

5.2.2 - Summary of intensity assignments of toluene

The assignments in the previous sections are summarized in Table 5.4, with the intensities from the literature. As noted previously, in order to compare our results for the liquid with the computed, fitted or measured values for the gas, the C_l values (Eq.

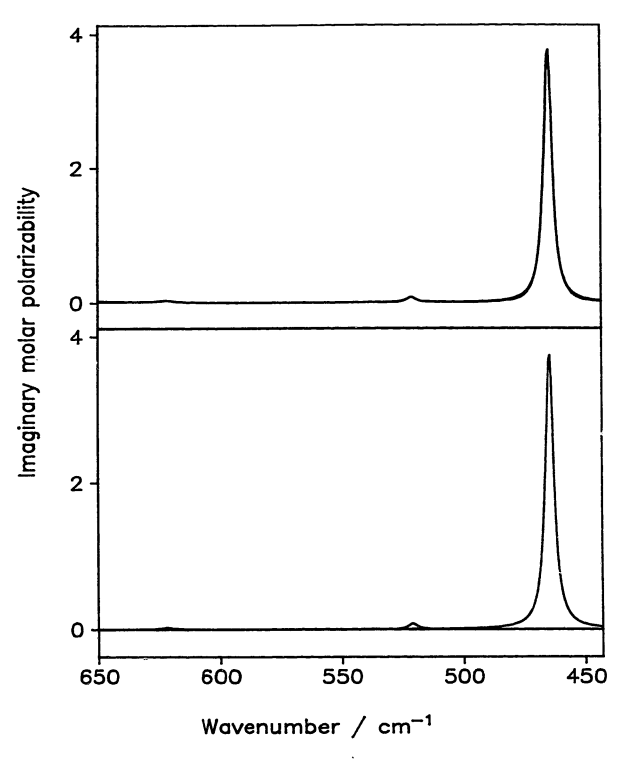


Figure 5.15 - Top - Superimposed experimental imaginary molar polarizability, α_m'' , and curve fitted α_m'' spectra of liquid toluene between 650 and 442 cm⁻¹. Bottom - The prominent CDHO bands in this region that were used to fit the experimental α_m'' spectrum. Units for both boxes are cm³ mole⁻¹.

1.4.2) assigned for the liquid were converted to A_j values (Eq. 1.4.1) for the gas by multiplying them by $8\pi^2$, i.e, by 78.96 (Eq. 5.1.4). This conversion gives the A_j intensity the gas molecules would have if the intrinsic intensity of the vibration were the same in the gas and liquid phases.

In this section, the A_f values are used to compare the present and literature measurements. Boggs' values²⁶ are from *ab initio* calculations of isolated molecules, and it is of interest to observe how well such calculations agree with the values measured for the liquid. Galabov²⁹ gives two sets of values, those measured for gaseous molecules and those calculated through the eigenvectors obtained from normal coordinate calculations from the dipole moment parameters calculated by fitting the measured intensities. His experimental values are the most fundamentally valuable. Unfortunately, the many studies of the intensities of gaseous benzene have shown that experimental gas phase intensities are difficult to measure and are frequently not very reproducible between laboratories. In the case of benzene, the agreement between $A_{f,gas}$ values of the four infrared active fundamentals from different laboratories of is within $\pm 15\%$ for each band.³⁴

The vibrations of the phenyl group are considered first. Four of these, v_3 , v_{14} , v_{20} and v_{30} were not measured in this work, and v_{12} , v_{13} and v_{14} have A_2 symmetry and are not infrared active in the gas. Of the remaining 24 vibrations, 12 have intensity $A_{J,gas}$ greater than 2.0 km mole⁻¹ and 12 have intensity less than 2.0 km mole⁻¹.

Of the 12 more intense bands, the *ab initio* calculated intensities agree with the observed intensities for the liquid to better than a factor of 2 for all except v_2 , v_{21} and v_{22} . In the case of the B_1 CH stretching modes, v_{21} and v_{22} , the sum of their intensities agrees with the sum calculated by Boggs to well within a factor of 2, and in fact the individual intensities agree within a factor of 2 if the assignments are reversed. It seems most probable that this difference is due to differences in the calculated wavenumbers or eigenvectors, rather than to a difference in the simplest molecular intensity quantities, the dipole moment derivatives. v_2 is also a CH stretching mode, of symmetry A_1 . However, neither the addition of the intensities of v_1 , v_2 and v_3 , nor any interchange of assignments, improves the agreement with the *ab initio* values to within a factor of 2 for these A_1 CH stretching modes.

There are 12 vibrations with $A_{j,gas}$ intensities less than 2 km mole⁻¹, and 10 of them differ by a factor of 2 or more from the *ab initio* results. This is not a surprising result, given the weakness of the bands and the imperfection of the experiments, assignments, and calculations.

Overall, the agreement between the experimental and *ab initio* results is very good, sufficiently good to indicate fairly clearly that the liquid forces do not cause the gas phase intensities to change in a major way for these liquids which do not interact strongly in solution.

The question of the extent to which the individual vibrational intensities actually do change between the gas and liquid phase is a more detailed question that requires a

Table 5.5 - Experimental integrated intensities of toluene

| | A_j | ab | |
|-----------------------------------|-----------|-----------------------------|--|
| ν_j | This work | Galabov et al ²⁹ | |
| v_1, v_2, v_3 | 11.8 | 26.5 | |
| v_{21}, v_{22} | 31.3 | 38.2 | |
| v_{31} | 11.8 | 11.1 | |
| V ₃₄ , V ₃₇ | 13.2 | 18.5 | |
| v_4, v_{23} | 6.1 | 7.3 | |
| V ₅ | 12.6 | 14.9 | |
| v_{24}, v_{35}, v_{38} | 19.7 | 6.5 | |
| v ₂₅ , v ₃₂ | 4.3 | 3.1 | |
| V ₂₈ | 4.1 | 4.3 | |
| $v_{10}, v_{17}, v_{18}, v_{29}$ | 79 | 41 | |
| v_{11}, v_{19} | 8.9 | 6.2 | |

a- $A_{j,gas}$ values for this work are taken from column 5 of Table 5.4, and Galabov's experimental values are taken from column 8 of Table 5.4.

b- Units are km mole⁻¹.

comparison of actual measured intensities. For this we compare our $A_{j,gas}$ values in column 5 of Table 5.4 with Galabov's 11 measured values in column 8. Because many of Galabov's values are for the sum of several vibrations, Table 5.5 conveniently compares them with appropriate sums of our $A_{j,gas}$ values. It is clear that there are cases of excellent agreement, that in the majority of cases the values lie within a factor of 2, and that there are many cases of significant disagreement.

As noted above, there are uncertainties in our intensity assignments, but the intensities of the major bands are believed to be accurate to a few percent. Although there is no intention to question the care with which Galabov *et al.* did their

measurements, it must be noted that they provided no spectra or other evidence to allow readers to judge the probable accuracy of their experimental values. The fact that there are many factors that can introduce error into the measurement of intensities of gases suggests that it is wisest to await other measurements of the gas phase intensities before any attempt is made to interpret these differences.

5.3 - Comparison of integrated intensities of liquid chlorobenzene and liquid toluene

As discussed in Chapter 4, it is not possible to describe the fundamental vibrations of chlorobenzene and toluene as arising from a single symmetry coordinate of benzene. The majority of the vibrations are rather due to mixture of coordinates.

Nevertheless, as the wavenumbers of many fundamentals of the two compounds do not differ by more than 15 cm⁻¹, it is interesting to compare the integrated intensities of liquid chlorobenzene and toluene because the two compounds have the same molecular skeleton with the exception of the chlorine atom and the methyl group.

Table 5.6 lists the integrated intensities, $C_{j,liq}$, of the fundamentals of chlorobenzene, the 30 phenyl group fundamentals of toluene and some of the fundamentals of benzene. The values for the first two liquids are from Tables 5.2 and 5.4, and those of benzene are taken from Ref. 73. The assignment for chlorobenzene and toluene is given first, followed by the wavenumber and intensities $C_{j,liq}$ for these liquids. This is followed by the ratio of the $C_{j,liq}$ of the two liquids, and the

Table 5.6 - Integrated intensities of liquid chlorobenzene, toluene and benzene

| | Chlorobenzene | | Toluene | | | Benzene [Ref. 73] | | |
|-----------------------------------|----------------------------|--------------------|----------------------------------|---------------------|--------------------------------------|--------------------|------------------------------|------------------|
| ν ₎ ^a | cm^{-1} b $C_{j,liq}$ c | | cm ^{-1 b} $C_{j,lig}$ c | | $\frac{C_{i,C6H5C1}}{C_{j,C6H5CH3}}$ | cm ^{-1-b} | v _j d | $C_{j,liq}^{-c}$ |
| $v_1, v_2, v_3, v_{21}, v_{22}$ | ~3070 ° | 0.119 ° | ~3060 ° | 0.546 ° | 0.22 ° | ~3040 ° | V12 ° | 0.539 ° |
| ν ₄ ν ₂₃ | ~1592 ^f 1584 | 0.187 ⁸ | 1605 1587 | 0.0775 ^g | 2.4 | 1586 | V ₁₆ | 0.00782 |
| v_5 | 1478 | 0.354 | 1496 | 0.160 | 2.2 | ~1475 | V ₁₃ | ~0.25 ' |
| V ₂₄ | 1445 | 0.127 | 1460 ^h | 0.250 ^h | 0.51 | ~14/3 | | ~0.23 |
| v_5 and v_{24} | | 0.481 | | 0.410 | 1.17 | | | |
| V ₂₅ | 1325 | 0.00270 | 1332 | 0.00174 | 1.55 | ~1345 ¹ | $\mathbf{v_3}$ | |
| ν ₆ k | 1172 | 0.0167 | 1179 | 0.0103 | 1.62 | 1177 | V ₁₇ | 0.0233 |
| v ₂₇ | 1156 | 0.00632 | 1156 | 0.0219 | 0.29 | 11,, | - 1, | 0.0233 |
| v_6 and v_{27} | | 0.0230 | | 0.0322 | 0.71 | | | |
| v_{28} | 1068 | 0.0359 | 1081 | 0.0521 | 0.69 | 1025 | | 0.130 |
| v_8 | 1023 | 0.117 | 1030 | 0.0261 | 4.5 | 1037 | v_{14} | |
| v_{28} and v_8 | | 0.153 | | 0.0782 | 2.0 | | | |
| V 9 | 1002 | 0.0206 | 1002 | 0.00394 | 5.2 | 1010 | V ₆ | 0.0050 |
| V ₁₅ | 983 | 0.00498 | 981 | 0.0115 | 0.43 | 971 | Ven | 0.0163 |
| v_{12} | 964 | 0.00852 | 966 | 0.00591 | 1.44 | 9/1 | V ₁₉ | 0.0163 |
| v_{15} and v_{12} | | 0.0135 | | 0.0174 | 0.78 | | | |
| v_{16} | 903 | 0.0446 | 895 | 0.0126 | 3.5 | 850 | v_{11} | 0.015 |
| v_{13} | 830 | 0.00968 | 843 | 0.00783 | 1.24 | | | |
| v_{16} and v_{13} | | 0.0543 | | 0.0204 | 2.7 | | | |
| v_{17} | 741 | 0.809 | 730 | 0.777 | 1.04 | 703 ^J | Vg | |
| v_{18} | 685 | 0.217 | 695 | 0.216 | 1.00 | 675 | V ₄ | 1.275 |
| V_{29} | 614 | 0.00280 | 622 | 0.00161 | 1.74 | 606 ^J | v ₁₈ ^ℓ | |
| v_{19} | 468 | 0.0983 | 464 | 0.110 | 0.89 | 403 | V ₂₀ | 0.000 |
| v_{14} | ~400 | 0.00621 | ~405 | m | | 105 | 2.7 | 5.750 |

Table 5.6 - Continued

| | Chlorobenzene | | | luene | | Benzene [Ref. 73] | | |
|-----------------|--------------------|-----------------|--------------------|---------------|--------------------------------------|--------------------|-------------------------------|---------------|
| ν, * | cm ^{-1 b} | $C_{j,liq}^{c}$ | cm ^{-1 b} | $C_{j,liq}$ c | $\frac{C_{i,C6H5C1}}{C_{j,C6H5CH3}}$ | cm ^{-1 b} | v_j^{-d} | $C_{j,liq}$ c |
| x-sensitive | | | | | | | | |
| V ₂₆ | 1270 | 0.00892 | 1312 | 0.00331 | 2.7 | 1310 | v ₉ ? " | 0.00342 |
| V7 k | 1084 | 0.298 | 1210 | 0.00480 | 62.1 | | ? n | |
| v_{1o} | 702 | 0.187 | 786 | 0.00337 | 55.5 | 993 | v ₂ ? ⁿ | 0.00642 |
| v_{11} | 415 | 0.0185 | 521 | 0.00297 | 6.2 | 60 6 ¹ | v_{is} | |
| V ₃₀ | 297 | m | 345 | m | | 1148 | v_{10} ? ⁿ | |
| v_{20} | 196 | m | 217 | m | | 994 | v ₇ ? ⁿ | |

- a Assignments for chlorobenzene and toluene.
- b Observed wavenumber in the imaginary molar polarizability spectrum of the liquid.
- c Units are km mole-1.
- d Assignments were taken from Ref. 73. Correlation between vibrations of benzene and those of chlorobenzene and toluene is based on wavenumber proximity and symmetry species.
- e The integrated intensities of the aromatic CH stretching vibrations are taken collectively. In benzene, $C_{j,liq}$ is assigned to the active vibration v_{12} . 3 inactive vibrations, v_1 , v_5 and v_{15} are also assigned wavenumbers in the proximity of v_{12} .
- f The wavenumber of the CDHO band assigned to v_4 .
- g Combined intensity for v_4 and v_{23} .
- h The same wavenumber was assigned to v_{24} , v_{35} and v_{38} . $C_{j,liq}$ of the CDHO band at 1460 cm⁻¹ is given.
- i Average value of the two values given in Ref. 73.
- j Wavenumber not observed in the spectrum and taken from Painter and Koening85.
- $k v_6$ in chlorobenzene is v_7 in toluene and vice versa.
- ℓ The degenerate v_{18} vibration in benzene was correlated with v_{29} and v_{11} in chlorobenzene and toluene.
- m Wavenumber below the experimental range of this study. thus, no integrated intensity is available.
- n Uncertain correlation between the benzene vibration and those of chlorobenzene and toluene.

should be noted that v_{11} to v_{20} are degenerate vibrations in gaseous benzene that are each split in C_{2v} molecules into two vibrations. A tentative correlation between the vibrations of benzene and those of chlorobenzene and toluene has been made in Table 5.6 based on wavenumbers and symmetry species of the fundamentals of the three compounds.

Of the 30 vibrations, the wavenumbers of 24 vibrations are not sensitive to the different group substitution, in that their wavenumbers in chlorobenzene and toluene differ by less than 15 cm⁻¹. The integrated intensities of these vibrations are listed in order of decreasing wavenumber. Six vibrations can be considered X-sensitive³⁷, in that the different substitutions cause the wavenumbers to be more than 15 cm⁻¹ apart. The integrated intensities for these six vibrations are given at the end of the table.

The ratio, $\frac{C_i \ cohscl}{C_j \ cohsch3}$, gives an indication whether the intensity of a fundamental vibration in liquid chlorobenzene is significantly different from that in liquid toluene. For qualitative evaluation, a difference is not considered significant if the ratio of the integrated intensities is $0.5 < \frac{C_i \ cohsch}{C_j \ cohsch3} < 2.0$, i.e. the intensities differ by less than a factor of 2.

For the vibrations under 990 cm⁻¹ that are not sensitive to substitution, it is found that for 6 out of 8 vibrations $0.5 < \frac{C_{I} \, CoH3CI}{C_{J} \, CoH3CH3} < 2.0$. Thus, for these 6 vibrations, the change in intensity between the two molecules is not significant.

The two exceptions are v_{15} and v_{16} , where the ratios are 0.43 and 3.5. The difference in v_{15} can be explained when the splitting of the degenerate v_{19} in benzene is considered. v_{19} in benzene is assigned to 971 cm⁻¹ and is split in C_{2v} molecules into v_{15} (~982 cm⁻¹) and v_{12} (~965 cm⁻¹). The ratio for the combined intensities for these vibrations in chlorobenzene and toluene is 0.78. Thus, it appears that a different distribution of intensities occurs in chlorobenzene and toluene but the total intensity does not change significantly.

The ratio for the combined intensities of v_{16} and v_{13} which result from the split of v_{11} in benzene is 2.7. The total intensity increases from 0.0155 km mole⁻¹ in benzene to 0.0204 km mole⁻¹ in toluene to 0.0543 km mole⁻¹ in chlorobenzene. The individual ratios for v_{16} and v_{13} are 3.5 and 1.24. Thus, the intensity of v_{16} changes significantly upon substitution.

The non sensitive vibrations above 990 cm⁻¹, largely derive from the splitting of degenerate modes in benzene, except v_{25} and v_9 . The intensity of v_{25} does not change significantly between chlorobenzene and toluene, while that of v_9 in chlorobenzene is 5.2 times more intense than in toluene and 4.1 times more intense than the corresponding intensity in benzene (v_6).

There are three pairs of vibrations, in which the intensity distribution among the two vibrations that result from the splitting of a degenerate vibration in benzene is significantly different in chlorobenzene and toluene. Yet, the ratio of the combined intensities does not indicate a significant difference. These vibration pairs are: v_5 and

 v_{24} , which arise from v_{13} in benzene, v_6 and v_{27} , which arise from v_{17} in benzene and v_{28} and v_8 , which arise from v_{14} in benzene.

A fourth pair, v_4 and v_{23} , shows significant change, as C_{hlq} of chlorobenzene is 2.4 greater than that of toluene which is itself about 10 times greater than the intensity of v_{36} the corresponding degenerate vibration in benzene.

The combined intensities $C_{j,liq}$ of the aromatic CH stretching vibrations in toluene, v_1 , v_2 , v_3 , v_{21} and v_{22} , is 0.546 km mole⁻¹ which is very similar to the intensity assigned to the infrared active vibration in benzene⁷³, v_{12} , at 0.539 km mole⁻¹. It is greater by a factor of 4.6 than the corresponding intensity of chlorobenzene. It appears that the chlorine substitution shifts intensity from the CH stretching vibrations to other vibrations.

Of the six X-sensitive vibrations which have wavenumbers that differ by more than 15 cm⁻¹ in chlorobenzene and toluene, v_{30} and v_{20} , are below the experimental range of the study, so no integrated intensities are available. The ratios $\frac{C_1 \cos H3CI}{C_1 \cos H3CH3}$ were calculated for the other four vibrations, v_{26} , v_7 , v_{10} and v_{11} and all showed integrated intensities in liquid chlorobenzene that are significantly greater than those in toluene, by factors of 2.7, 62.1, 55.5 and 6.2, respectively. It is quite possible that the gain in intensity in these vibrations, and in particular v_7 and v_{10} , is associated with the presence of the electronegative chlorine and with the loss of intensity of the CH stretching vibrations.

A satisfactory further analysis of these intensity differences between benzene, chlorobenzene and toluene requires a combination of *ab initio* calculations and a simple normal coordinate analysis for these molecules. These calculations will allow proper account to be taken of the different combinations of atomic displacements that occur in the normal vibrations of the different molecules, i.e., of the different eigenvectors in the different molecules, which, in turn, will allow the intensities to be analysed in terms of more basic properties such as the change in molecular dipole moment during a particular symmetry coordinate or a particular bond or angle displacement. Such calculations are not ideal for the interpretation of such data but, when guided by *ab initio* calculations, are the best that is available.

In order to understand the change caused by the reduction of symmetry from the D_{6h} of renzene to the C_{2v} of chlorobenzene and toluene without the added complication of the different chemical nature of the substituent, the intensities of C_6H_5D have been measured in this laboratory. Thus, a further analysis of the data presented in this thesis will be made in the future with the wavenumber and intensity values of liquid C_6H_6 , C_6D_6 , C_6H_5D , C_6H_5Cl , and $C_6H_5CH_3$.

Chapter 6 - A simple and effective approximate method for the calculation of infrared optical constant spectra of liquids from transmission measurements*

6.1 - Introduction

Earlier papers from this laboratory reported the absolute infrared absorption intensities of four liquids, benzene¹, toluene², chlorobenzene³ and dichloromethane⁴, obtained from transmission measurements. Intensities from this work were accepted by the International Union of Pure and Applied Chemistry (IUPAC) as secondary absorption intensity standards⁵. The purpose of this paper is to describe a much simpler but approximate computational procedure for obtaining such quantitative intensities from transmission spectra of liquids. This approximate procedure yields results of very nearly the same accuracy as the more complex but exact procedure used to date, although we recommend caution if the refractive index of the sample in non-absorbing infrared regions differs from that of the windows by more than ~0.15.

The primary result of transmission spectroscopy is the transmittance of the cell plus sample, $\frac{I_t}{I_o}$, where I_o is the intensity incident on the cell and I_t is the intensity which leaves the cell, both measured outside of the cell. For quantitative studies this is always converted to the "absorbance", by taking $-\log_{10} \left\{ \frac{I_t}{I_o} \right\}$. The term "absorbance" is given

^{*} A version of this chapter has been accepted for publication. Bertie and Apelblat, Appl. Spectrosc., 1996.

in quotation marks here, because it is not the quantity that is equated with E_mCd , the product of the molar absorption coefficient, E_m , the concentration, C, and the path length through the liquid, d, in the Beer-Lambert law. The absorbance, A_{10} , as defined by $IUPAC^6$ and as equated with E_mCd , is a measure of energy lost from the radiation beam due solely to absorption by the sample, a liquid in our case. In contrast, the "absorbance" that is calculated as described above, and reported by the spectrometer in most transmission experiments, is a measure of the energy lost due to absorption by the sample, plus the energy lost due to reflection from the interfaces of the cell, plus the energy lost due to unexplained baseline errors.

The IUPAC definition⁶ of absorbance, A_{10} , is used in this laboratory. The term experimental absorbance⁷ is used for the "absorbance" calculated as described above, and the term apparent absorbance⁷ is used for the contributions to the experimental absorbance from processes other than absorption by the sample. Thus the experimental absorbance is the sum of the absorbance, A_{10} , plus the apparent absorbance due to reflection plus the apparent absorbance due to baseline errors. It should be noted that the latter may be positive or negative.

In Refs. 1 to 4, the imaginary refractive index spectrum, $k(\tilde{v})$, of the liquid was calculated by an iterative procedure that fully corrects the experimental absorbance, EA, spectrum for the non-absorption processes. The real refractive index spectrum, $n(\tilde{v})$, and the molar absorption coefficient spectrum $E_m(\tilde{v})$ were calculated from the $k(\tilde{v})$ spectrum.

The method used previously¹⁻⁴ to correct the experimental absorbance spectrum for the apparent absorbance due to reflection losses is exact for an ideal experiment⁷⁻⁹. Thus Fresnel's equations are applied to each interface, specifically, the interface between the air and the first window, the interface between the first window and the thin liquid film, the interface between the liquid film and the second window, and the interface between the second window and the air. The interference fringes generated by multiple reflections within the liquid layer are calculated. In contrast, the calculation averages over the multiple reflections within the windows and, thus, averages over the interference fringes generated in the windows, as is appropriate for ~5mm alkali halide windows and 1 cm⁻¹ resolution. Further, the calculation corrects for the convergence and polarization of the incident beam. It is a calculation that is exact in the ideal case, but is sufficiently complex to discourage new users.

In the present paper, a very simple calculation of the apparent absorbance due to reflection is presented. This greatly simplifies the correction of the experimental absorbance spectrum for the apparent absorbance due to reflection and, thus, greatly simplifies the calculation of optical constant spectra from transmission spectra. Although simpler, the method gives imaginary refractive index, k, values which agree within 1% with those from the exact calculation for all but the strongest infrared absorptions, provided that the real refractive indices of the sample and windows are not too different.

6.2 - Method

The EA spectrum of a cell full of liquid is obtained by dividing the intensity spectrum through the full cell by that through the empty sample compartment and taking the negative logarithm of the result.

$$EA = -\log_{10} T = -\log_{10} \frac{I_{t}(\widetilde{\nu})}{I_{o}(\widetilde{\nu})}$$
(6.1)

In the method used previously¹⁻⁴, the refractive index spectra are calculated from the *EA* spectrum by program RNJ46A. The program was developed by Jones and his coworkers in the National Research Council of Canada and was later improved in this laboratory. It is described elsewhere⁷⁻¹⁰, the algorithm is outlined in the appendix, and the salient features for the present purpose are given briefly next.

Experimental absorbance spectra are used in two parts of the procedure. In the first part, EA spectra of samples with several different very long path lengths are used to determine the linear absorption coefficients, $K(\widetilde{v})$, at so-called *anchor points* in the regions of baseline absorption. $K(\widetilde{v}) = \ell^{-1} A_{10}$, where ℓ is the path length through the liquid and A_{10} is the absorbance of the sample. The K values are calculated from the EA spectra by program ANCHORPT FOR which is outlined in the appendix. The baseline error is assumed to be zero in this calculation, which is a good approximation if the experimental absorbance is about 0.3 or greater and which is evaluated later by the agreement seen in the K values calculated from cells with different path lengths. With this assumption, $EA = A_{10} + AA_{R}$, where AA_{R} is the apparent absorbance due to

reflection. At each anchor point AA_R is calculated exactly, by applying Fresnel's equations to each interface and using the known real refractive index of the windows, for an approximately correct value of the real refractive index of the sample.

The K values thus produced are then used in the second part of the procedure to remove the apparent absorbance due to baseline errors in spectra of samples with normal path lengths. This is done by calculating where the baseline should be at each anchor point, through $EA = K\ell + AA_R$, and shifting the spectrum to move the baseline to that correct place. Linear interpolation of the required shift is used between the anchor points.

In the second part of the procedure, the *EA* spectrum is used in program RNJ46A to calculate the refractive index spectra by an iterative procedure, after the baseline is corrected. During this part, the apparent absorbance due to reflection is recalculated in each iteration cycle.

In the first calculation cycle, a first approximation to the imaginary refractive index, k, spectrum is calculated. For this purpose, the reflection loss is calculated, for all wavenumbers in the EA spectrum, from the known refractive index spectrum of the windows and a constant value of the real refractive index of the sample. From the approximate k spectrum thus generated, a real refractive index spectrum, n, is calculated by a Kramers-Kronig (KK) transform. In subsequent iteration cycles, the wavenumber-dependent real refractive index spectrum of the sample is used in the calculation of the reflection losses and of the baseline-correction of the EA spectrum.

In the second part of the procedure, the n and k spectra of the sample are used with the n spectrum of the windows, to calculate the EA spectrum. The calculated and the baseline-corrected experimental EA spectra are compared, the k spectrum is adjusted to improve the fit and the n spectrum is recalculated. This cycle is repeated as the k and n spectra are refined by iteration⁷⁻⁹ to convergence. The current n spectrum is used in the calculation of AA_R and the baseline correction. The convergence of the radiation beam can be included in the calculation, as can be the polarization of the beam. The latter two factors are not significant for the beam convergence and polarization in the Bruker IFS 113V spectrometer in this laboratory.

In the approximate method presented here, the reflection losses are calculated by considering that the cell is simply a single window. The transmittance of the window is given by $T(\tilde{v}) = \frac{2n(\tilde{v})}{n^2(\tilde{v})+1}$, where n is the real refractive index of the window. Thus, the

apparent absorbance due to reflection losses at wavenumber \tilde{v} , $AA_R(\tilde{v})$ is simply

$$AA_{R}(\widetilde{\nu}) = -\log_{10} \frac{2n(\widetilde{\nu})}{n^{2}(\widetilde{\nu}) + 1}$$
(6.2)

The approximate method uses this simple formula in both parts of the procedure outlined above, which eliminates the need for an iterative calculation in the second part. The method is implemented in the program WIND8.FOR. The program incorporates the functions of both ANCHORPT and RNJ46A. Program WIND8 is outlined in the appendix, where its relation to ANCHORPT and RNJ46A is emphasized.

It is convenient to consider here the approximation made in this simple procedure. The discontinuity in the real refractive index at the window-liquid boundary is neglected. Fig, 6.1 shows an absorption band in the k spectrum and the anomalous dispersion¹¹ in the real refractive index, n, that accompanies it. The real refractive index of the liquid is not constant near an absorption band, so the neglected reflection losses are not constant. Whether these losses are greater to high or to low wavenumber of the absorption band depends on whether the (nearly constant) refractive index of the window is larger or smaller than the average n value of the sample through the absorption band. The size of the deviations of n from the average value is directly proportional to the height of the absorption band in the k spectrum. Thus, the simple procedure is expected to be most accurate for weak absorptions, and the largest deviations from the correct procedure are expected for strong absorptions.

The approximation involved in the simpler procedure can be seen directly in Figure 6.2, in which the bottom box shows the standard real refractive index spectrum of chlorobenzene^{3,5} between 1700 and 1100 cm⁻¹, and the top and middle boxes show the apparent absorbance due to reflection that is calculated by the approximate and exact methods for two different cases.

The top box shows the case of 100 μ m thick chlorobenzene between KBr windows. The refractive indices of the window and sample are very close in this case, with $n \sim 1.51$ for chlorobenzene between strong absorptions and $n \sim 1.52$ for KBr. The smooth bottom curve is that calculated from Eq. 6.2. The top curve is the result of the exact calculation, which used the standard n spectrum of chlorobenzene^{3,5}. The

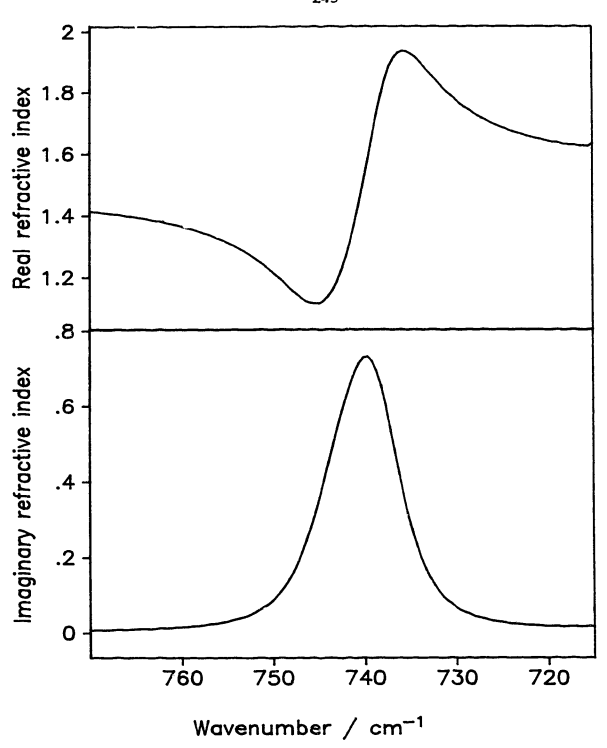


Figure 6.1 - The real (upper box) and imaginary (lower box) refractive index spectra of the strongest absorption band in liquid chlorobenzene³.

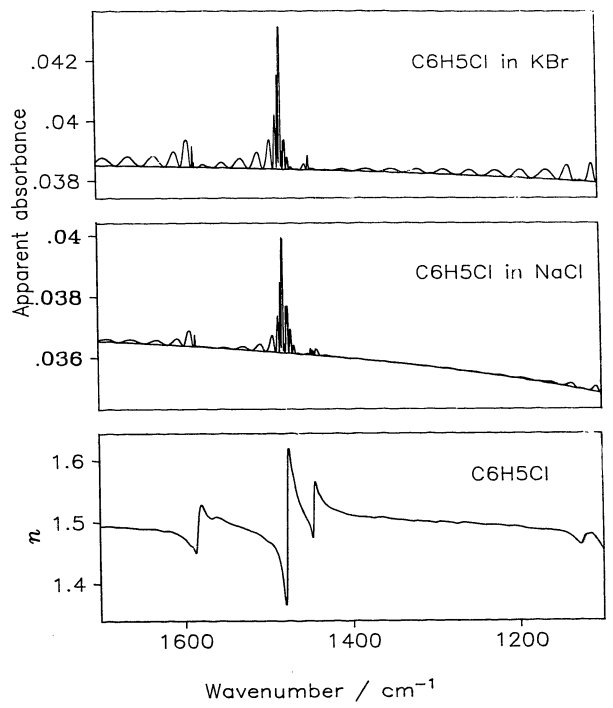


Figure 6.2 - Top and middle boxes: The apparent absorbance due to reflection calculated by the exact method (upper, featured curve) and the approximate method (lower unfeatured curve) for $100~\mu m$ of chlorobenzene(ℓ) between KBr windows (top box) and NaCl windows (middle box). Bottom box: The real refractive index spectrum of liquid chlorobenzene^{3,5} at 25°C.

interference fringes arise from the multiple reflections that are ignored in the approximate method. The middle box shows the case of 100 μ m thick chlorobenzene between NaCl window, for which $n \sim 1.50$, again with the smooth bottom curve being that from the approximate calculation. The effect of the refractive index of the windows is evident in the lower apparent absorbance and in the different shape of the apparent absorbance spectrum from the exact calculation. The effect of the wavenumber dependence of the refractive index of the sample is also evident, in that no interference fringes are visible in regions where the refractive indices of sample and window are very close, but fringes are visible in other regions where the refractive index of the sample is different because of the anomalous dispersion.

6.3 - Results

The EA spectra were processed with programs ANCHORPT and RNJ46A to calculate k spectra by the exact method. The same EA spectra were processed with program WIND8 to calculate k spectra by the approximate method. The K values at anchor points in the baseline were calculated and used to adjust the baseline in both cases; it is essential to do this for the best quantitative results. In the exact method the molar absorption coefficient, E_m spectrum is calculated from the final, converged, k spectrum through

$$E_{\rm m} = \frac{4\pi \widetilde{\nu} k}{2.303C} \tag{6.3}$$

where C = the molar concentration. In the approximate method the E_m spectrum is calculated directly from the absorbance spectrum (after correction for the apparent absorbance due to baseline errors and reflection losses) through

$$E_{\rm m} = \frac{A_{10}}{C\ell} \tag{6.4}$$

The approximate method was tested with spectra of chlorobenzene, several mixtures of chlorobenzene and toluene, and, to a limited extent, with spectra of acetonitrile. To demonstrate its value, results are first presented for four regions of the spectrum of liquid chlorobenzene. These are the region between 805 and 715 cm⁻¹ which contains a single band of very strong absorption, and three regions which contain multiple bands of strong (1635 - 1400 cm⁻¹), medium (3200 - 2930 cm⁻¹) and weak (1405 - 1140 cm⁻¹) absorption. In Figs. 6.3 and 6.4 these regions are labeled A, B, C and D, respectively. Comparison is made of the peak heights, the areas under the $E_m(\tilde{\nu})$ spectrum, and the K values at the anchor points, that are calculated by the two methods. The peak heights and areas are compared for E_m spectra instead of for k spectra, because the molar absorption coefficient is more familiar to chemists, and because the two quantities are related by Eq. 6.3 so the percent differences are essentially the same in both cases. Note that regions B and D are in the range of Fig. 6.2, but these figures can not be compared directly because the two path lengths, and hence the interference fringe patterns, are different.

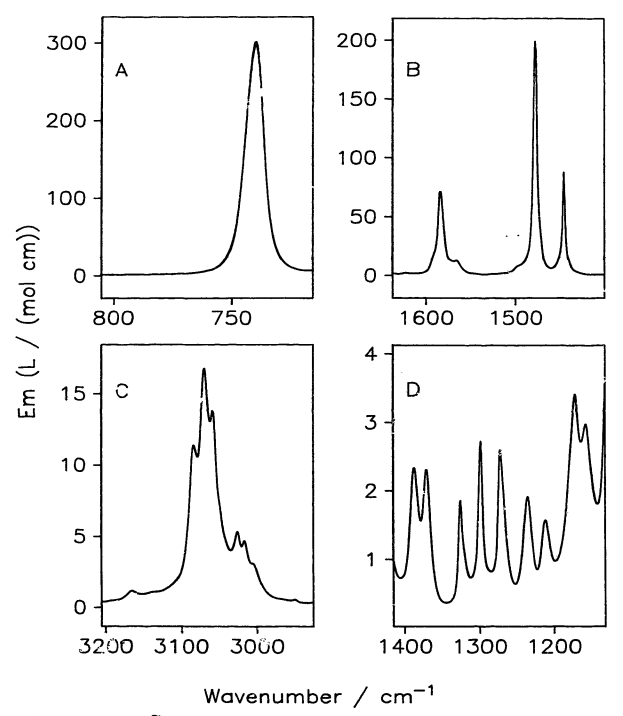


Figure 6.3 - The $E_{\rm m}(\widetilde{\nu})$ spectrum of chlorobenzene(ℓ) in four regions. The spectra calculated by the correct and approximate methods are shown in each box, and essentially coincide. The transmission spectra, from which these $E_{\rm m}$ spectra were calculated, were recorded with KBr windows and chlorobenzene thicknesses of: Box A 8.70 μ m; Box B 11.02 μ m; Box C 34.39 μ m; Box D 533.0 μ m.

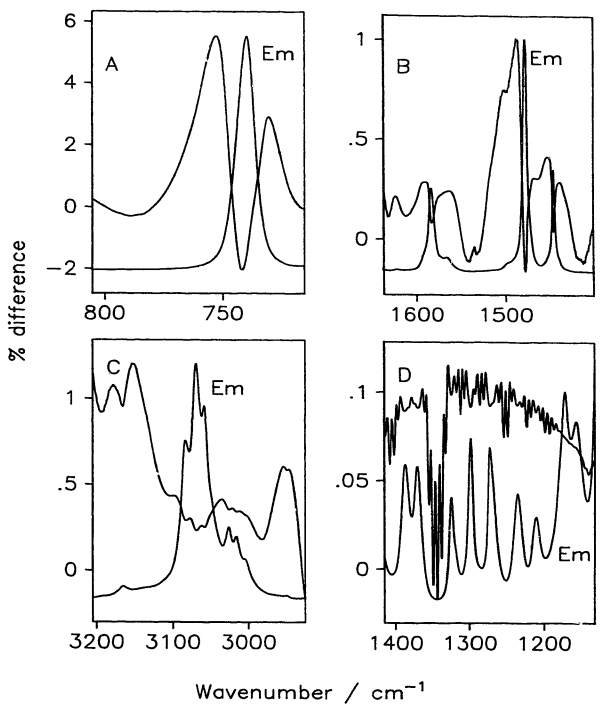


Figure 6.4 - The percent difference between the $E_{\rm m}$ spectra, $\frac{{\rm approximate}\;E_{\rm m} - {\rm exact}\;E_{\rm m}}{{\rm exact}\;E_{\rm m}} \times 100\%$, of chlorobenzene(ℓ) in the four regions of Fig. 6.3. For reference, the exact $E_{\rm m}$ spectrum is included, on an arbitrary scale, in each box.

Figure 6.3 shows the approximate and correct molar absorption coefficient spectra in the four regions. The spectra are essentially coincident. This shows that the band shapes are not much distorted by the approximate calculation. Bandshape distortion is seen more clearly in the spectra of the % difference, defined as $\frac{\text{approximate } E_{\text{m}} - \text{exact } E_{\text{m}}}{\text{exact } E_{\text{m}}} *100.$ These spectra are shown in Figure 6.4 for the four regions. For reference, the exact E_{m} spectrum is included in each box of Fig. 6.4, on an arbitrary scale. The % difference spectra show that the line shape is slightly distorted, but the distortion is ~1% or less for all but the strong band at 740 cm⁻¹, for which the distortion reaches several percent in the wings of the band.

Values of the linear absorption coefficients at the anchor points, K, computed by the two methods are given in Table 6.1. They are averages from several spectra and their 95% confidence limits are given in parentheses. The values from the approximate method agree with those from the correct method to within 0.05%, well within the 95% confidence limits of the values. The two sets of 95% confidence limits also agree. The one exception is for the highest anchor point K value, at 717.4 cm⁻¹, which is less precise from the approximate method although its precision is still 1.2%.

The peak heights calculated by the two methods, and their percent differences, are given in Table 6.2. The peak heights agree to better than 1%, except for 1.5% for the very strong 740 cm⁻¹ band.

As noted above, small changes in band shape result from the approximate

Table 6.1 - Linear absorption coefficients. $K(\tilde{\nu})$, of liquid chlorobenzene at anchor points in the baseline.

| Region | Wavenumber (cm ⁻¹) | Approximate $K(\widetilde{\nu})^4$ | Exact K(V) | % differenceb |
|--------|---------------------------------|------------------------------------|------------|---------------|
| Α | 801.3 | 12.66 (8) | 12.66(8) | +0.05 |
| | 717.4 | 65.9(8) | 65.9(6) | -0.002 |
| В | 1630.5 | 11.65(3) | 11.64 (3) | +0.03 |
| | 1534.0 | 9.08 (3) | 9.08 (3) | +0.04 |
| | 1404.8 | 7.06(3) | 7.06(3) | +0.05 |
| С | 3200.3 | 3.98(2) | 3.98(2) | +0.01 |
| | 2931.2 | 3.15 (2) | 3.15 (2) | +0.02 |
| D | 1404.8 | 7.06(3) | 7.06(3) | +0.05 |
| | 1345.1 | 3.53(2) | 3.53(2) | +0.01 |
| | 1251.5 | 6.24(3) | 6.24(3) | +0.02 |
| | 1197.5 | 9.55(2) | 9.55(2) | +0.02 |
| | 1141.6 | 17.7(1) | 17.6 (1) | +0.04 |

a - The number in parentheses is the 95% confidence limit in the last digit.

method, most noticeably for the strong peak in region A, for which the anomalous dispersion in the real refractive index is large (Fig. 6.1). For this band, deviations of ~5% and ~2.5% occur on the high- and low- wavenumber sides of the band (Fig. 6.4). For lineshape studies of strong bands, the exact method is clearly preferred. The bands in region B are weaker but are still quite intense for organic compounds. For these bands the distortion is of the order of 1%, which is probably too small to be significant at the present time. The bands in groups C and D are maken weaker, and the distortion within the band is well below 1%, although the error is ~1% in the baseline near 3200

b - The % difference is $\frac{\text{approximate } K - \text{exact } K}{\text{exact } K}$ *100, calculated before the values were truncated for presentation.

Table 6.2 - Peak heights in the molar absorption coefficient spectrum, $E_m(\tilde{\nu})$, of liquid chlorobenzene.

| Region | Wavenumber (cm ⁻¹) | Approximate $E_{m}(\tilde{\nu})^{a}$ | Exact $E_{\rm m}(\widetilde{\nu})^{\rm a}$ | % difference ^b |
|--------|--------------------------------|--------------------------------------|--|---------------------------|
| A | 739.7 | 297.7 | 302.3 | -1.52 |
| В | 1622.0 | 2.428 | 2.423 | +0.21 |
| | 1583.8 | 72.89 | 72.82 | +0.10 |
| | 1566.1 | 12.93 | 12.90 | +0.23 |
| | 1477.1 | 199.5 | 199.8 | -0.15 |
| | 1445.4 | 87.89 | 87.85 | +0.05 |
| С | 3165.2 | 1.193 | 1.182 | +0.93 |
| | 3083.2 | 11.46 | 11.43 | +0.26 |
| | 3069.5 | 16.82 | 16.78 | +0.24 |
| | 3058.4 | 13.84 | 13.81 | +0.22 |
| | 3025.7 | 5.358 | 5.339 | +0.36 |
| | 3016.3 | 4.699 | 4.683 | +0.34 |
| | 2949.7 | 0.561 | 0.558 | +0.54 |
| D | 1386 9 | 2.341 | 2.339 | +0.09 |
| | 1370 6 | 2.318 | 2.315 | +0.13 |
| | 232% 2 | 1.870 | 1.868 | +0.11 |
| | 1255.0 | 2.733 | 2.731 | +0.07 |
| | 1272.5 | 2.605 | 2.603 | +0.08 |
| | 1235.1 | 1.918 | 1.916 | +0.10 |
| | 1210.8 | 1.575 | 1.574 | +0.06 |
| | 1171.5 | 3.413 | 3.411 | +0.06 |
| | 1156.4 | 2.980 | 2.978 | +0.07 |

a - The unit of $E_{\rm m}$ is L mol⁻¹ cm⁻¹.

cm⁻¹ for the bands in region C.

The areas under bands in the exact and approximate $E_m(\tilde{\nu})$ spectra are given in Table 6.3. Two areas are given for each band, the area above zero ordinate and the area

 $b - \frac{\text{approximate } E_{\text{m}} - \text{exact } E_{\text{m}}}{\text{exact } E_{\text{m}}} * 100.$

Table 6.3 - Areas under the molar absorption coefficient spectrum, $E_{\rm m}(\tilde{\nu})$, of chlorobenzene.

| | | AREA | | | | | |
|--------|---------------------------|----------------------------------|-------|---------------------------|-----------------|-------|---------------------------|
| | Integration | Above zero ordinate ^a | | | Above baseline* | | |
| Kegion | range (cm ⁻¹) | Approximate | Exact | % difference ^b | Approximate | Exact | % difference ^b |
| Α | 801.3-717.9 | 3430 | 3433 | -0.09 | 3093 | 3096 | -0.10 |
| В | 1605.4-1410.7 | 3226 | 3220 | +0.20 | 2939 | 2933 | +0.22 |
| | 1605.4-1545.7 | 949 | 947 | +0.18 | 844 | 842 | +0.20 |
| | 1530.2-1410.7 | 2261 | 2257 | +0.22 | 2158 | 2153 | +0.23 |
| С | 3200.3-2931.2 | 909 | 906 | +0.37 | 810 | 808 | +0.34 |
| D | 1405.9-1345.1 | 77.9 | 77.8 | +0.09 | 44.9 | 44.8 | +0.11 |
| | 1345.1-1252.5 | 107.3 | 107.2 | +0.10 | 60.7 | 60.6 | +0.12 |
| | 1252.5-1198.5 | 65.1 | 65.0 | +0.09 | 21.1 | 21.1 | +0.11 |
| | 1198.5-1141.7 | 127.7 | 127.6 | +0.07 | 47.9 | 47.9 | +0.08 |

a - The unit of area is L mol⁻¹ cm⁻². Multiply the area by 100 to change the unit to ken mol⁻¹.

above a linear baseline drawn between the ordinates at the integration limits. The latter is useful when the baseline intensities are not of interest. Table 6.3 shows that the exact and approximate methods yield the same areas to better than 0.4% in all ranges, and to better than 0.25% for all except the CH stretching region, 3200.3 to 2931.2 cm⁻¹. It is noteworthy that region shows the largest differences for the peak heights and lineshapes but shows nearly the smallest difference for the areas.

Most transmission measurements in this laboratory have been made with KBr and NaCl windows. Thus the refractive index mismatch between sample and windows is usually small, because most liquids have refractive indices near 1.5. Exceptions exist,

b - approximate area - exact area *100, calculated before the values were truncated for presentation.

however, and the probable effect of refractive index mismatch on the usefulness of the approximate method has been explored by applying both methods to spectra of 23.26 μ m of acetonitrile, $n \sim 1.32$, between KBr windows. The top box of Figure 6.5 shows the apparent absorbance due to reflection calculated by the approximate and exact methods. The approximate method fails to calculate interference fringes that are ~ 0.006 absorbance units peak-to-peak, about 10 times larger than for chlorobenzene between KBr. The impact of this on the molar absorption coefficients calculated by the two methods is shown in the middle and lower boxes of Fig. 6.5. The approximate method gives E_m values that are smaller than the exact values in this case, and fails to correct for the interference fringes that are visible in the EA spectrum but are correctly eliminated by the exact calculation. The percent difference between the E_m values from the two methods is $\leq \sim 2\%$ near the peaks and rises to 30% in the baseline due to the neglect of the fringes. The percent difference between the areas under the peaks is 2% between 1700 and 1200 cm⁻¹ and 4% between 1200 and 900 cm⁻¹.

We have not explored the effect of refractive index mismatch to a much greater extent than this, because in practice it is not common. It is clear that the effect is significant, but that acceptable accuracy in peak heights and reas for peaks of medium intensity can be obtained when, as for CH₃CN between KBr, the refractive index mismatch is ~0.2. It is recommended that results calculated with the approximate method be regarded with caution if the refractive indices of the sample and windows differ by more than 0.15.

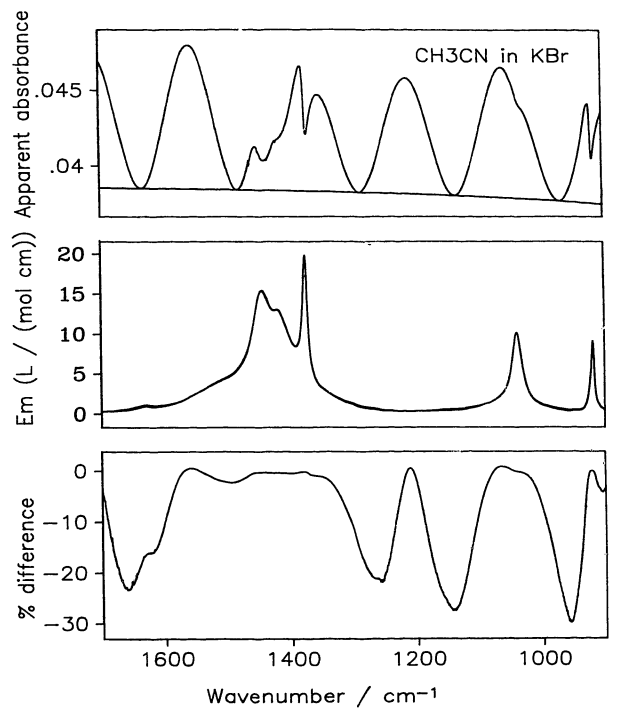


Figure 6.5 - Top box: The apparent absorbance due to reflection calculated by the exact method (upper, featured curve) and the approximate method (lower unfeatured curve) for 23.26 μm of CH₃CN(ℓ) between KBr windows. Middle box: The E_m spectra of CH₃CN(ℓ) calculated by the exact and approximate methods. Lower box: The % difference between the E_m spectra, defined as for Fig. 6.4, of CH₃CN between KBr windows. The refractive indices of CH₃CN(ℓ) and KBr in this spectral region are ~1.32 and ~1.52.

6.4 - Conclusions

An approximate method for the calculation of the optical constant spectra of liquids from transmission spectra is presented in this paper. The method calculates the reflection losses by treating the cell as a single window, instead of applying Fresnel's equations to each interface. In spite of this drastic approximation the method yields, for all but the strongest bands, imaginary refractive index and molar absorption coefficient values that are within ~1% of those calculated by the exact method. The line shape distortion is minimal for all but the strongest bands, and the integrated intensities agree with those from the exact method to better than 0.4% throughout. The agreement is excellent in regions of very weak baseline absorption, with the K values at the anchor points within 0.05% of those from the exact method.

This approximate method is much easier to understand and program than the more complex method, and a well written program runs much faster. It is clearly the method of choice except when one needs to obtain the exact bandshapes and peak heights without risk of distortion, and except when the spectrum contains bands whose intensities are comparable to or greater than that of the 740 cm⁻¹ band of chlorobenzene. The effect of the size of the refractive index mismatch has been explored to some extent with CH₃CN between KBr windows, for which the mismatch is $\Delta n \sim 0.2$. It is recommended that results from the approximate method be regarded with caution if the refractive indices of the sample and windows differ by more than 0.15.

6.5 - Appendix I

6.5.1. - Program ANCHORPT

ANCHORPT calculates the linear absorption coefficient, $K(\tilde{v})$, at anchor points in the baseline from the experimental absorbance, EA, spectrum of the liquid in a cell with long path length. Usually several EA spectra recorded with different path lengths are provided, so that an average value of K can be calculated, with its precision, for each anchor point. The program contains equations from which the real refractive indices of the common window materials can be calculated at infrared wavenumbers. Other required input is: the approximate real refractive index spectrum of the liquid; the number of anchor points and their wavenumbers; for each EA spectrum, the path length and the window material.

Procedure

- 1. For the first anchor point wavenumber, the real refractive indices of the liquid and the windows are used to calculate the transmission of the cell full of liquid, T_{abc} , and hence the apparent absolute due to reflection, $AA_{R} = -\log_{10} (T_{abc})$. T_{abc} , is calculated as in Ref. 9, corrected as described by Ohta and Ishida¹². The equations are given in Appendix II. For this calculation it is assumed that the liquid does not absorb at the anchor point wave number.
- 2. K at this wave number is calculated by subtracting AA_R from the EA value at this wave number in the first EA spectrum, and dividing the result by the path length.

- 3. Steps 1 and 2 are repeated for each anchor point.
- 4. Steps 1 to 3 are repeated for each EA spectrum.
- 5. The average value of K at each anchor point is calculated, with its standard deviation and 95% confidence limit.

6.5.2 - Program RNJ46A

RNJ46A is a modified version of the National Research Council of Canada program⁸ XLVI. The modifications have been described⁷, and the general procedure has been outlined^{1,10}. RNJ46A calculates the real, n, and imaginary, k, refractive index spectra of a liquid from an experimental absorbance spectrum. The program contains equations from which the real refractive indices of the common window materials can be calculated at infrared wavenumbers. Other required input is: the anchor point wavenumbers and linear absorption coefficients, K, the window material, the path length of the cell, ℓ , and the real refractive index of the liquid at the high-wavenumber limit of the $\frac{1}{2}$ spectrum, $m(\tilde{v}_{max})$. The half-cone angle of the non-parallel beam and the polarization discrimination, i.e., the relative intensities in the two polarizations of the incident beam, can be supplied if desired.

Procedure

1. The transmittance spectrum, T_{abc} , of the cell full of liquid is calculated as in Ref. 9, corrected as described by Ohta and Ishida¹². The equations are given in Appendix II. If

the half-cone angle of the beam and the polarization discrimination are supplied, the calculation uses these factors. Then $-\log_{10} T_{\rm abc}$ is calculated.

- 2. The first requirement is to calculate the approximate transmission spectrum that results solely from reflection losses. It is, therefore, first assumed that the imaginary refractive index is zero at all wavenumbers and the real refractive index of the liquid is constant. The value $n(\tilde{\nu}_{max})$ is used. With these assumptions, step 1 yields an approximate apparent absorbance due to reflection, AA_R . AA_R is subscatted from the EA spectrum to yield the approximate absorbance spectrum, AS.
- 3. The absorbance at the anchor points is calculated as $A_{AP} = K \times \ell$.
- 4. The baseline correction to be applied to the AS spectrum at the anchor points is calculated as A_{AP} AS, and the corrections required between the anchor points are found by linear interpolation. The AS spectrum is corrected to give the AS_I spectrum.
- 5. The baseline of the EA spectrum is corrected by calculating the correct EA at the anchor points, as $EA_c = AA_R + A_{AP}$, calculating the correction to EA at the anchor points as $EA_c EA$, calculating the correction to EA between the anchor points by linear interpolation, and adding the corrections to the EA spectrum. The resulting baseline-corrected EA spectrum has been called the ideal experimental absorbance spectrum. It is here denoted EA_I . The EA_I spectrum is used in step 9.
- 6. An approximate k spectrum is calculated from the corrected AS spectrum from step 4

as
$$k = \frac{2.303}{4\pi \ell \tilde{v}} AS_1$$
.

- 7. An approximate n spectrum is calculated from the k spectrum and $n(\tilde{v}_{max})$ by the Kramers-Kronig transformation.
- Thus, in this step one need not take the usual care to ensure that all the bands in the k spectrum are on scale and that all absorption beyond the ends of the EA spectrum is included in the transform. The n spectrum that is reported for the liquid is calculated later from a k spectrum that is assembled from many different EA spectra.
- 8. Step 1 is repeated, but this time using the k and n spectra calculated in steps 6 and 7. The result of this calculation is the calculated ideal experimental absorbance spectrum,

 EAC.
- 9. The calculated, $EA_{\rm C}$ and experimental, $EA_{\rm I}$, ideal experimental absorbance spectra are compared, and the k spectrum is adjusted by the addition of $\Delta k = \frac{2.303}{4\pi \ell \tilde{\nu}} \{EA_{\rm I} EA_{\rm C}\}$.

Here k, $EA_{\rm I}$ and $EA_{\rm C}$ are at wavenumber \tilde{v} .

10. The original baseline correction was not accurate, because the n spectrum of the liquid was assumed constant in step 1. Accordingly, the K values at the anchor points are used to correct the new k spectrum, where $K=4\pi \tilde{\nu} k/2.303$, again using linear interpolation of the correction between the anchor points. Further, the experimental EA spectrum is again baseline corrected, as in step 5 except that now the AA_R values at the anchor points are calculated using the n spectrum calculated in step 7.

- 11. A new *n* spectrum is calculated from the new, corrected, *k* spectrum and $n(\tilde{v}_{max})$, as in step 7.
- 12. Steps 8 to 11 are repeated until the magnitude of the correction in step 9 averaged over all spectral points is less than 2×10^{-5} .

6.5.3 - Program WIND8

WIND8 combines the functions of ANCHORPT and RNJ46A except that the *n* spectrum of the liquid is not calculated. The program makes a simplified calculation of the reflection losses in which Eq. 6.2, is used instead of the equations in Appendix II. The program contains equations from which the real refractive indices of the common window materials can be calculated at infrared wavenumbers.

Thus, WIND8 calculates the values of K at the anchor points in the baseline from EA spectra of cells with long path lengths, and calculates the imaginary refractive index, k, and molar absorption coefficient, $E_{\rm m}$, spectra of a liquid from an experimental absorbance spectrum in a cell of normal path length. Usually a large number of EA spectra in long path length cells are supplied.

The following input is required: EA spectra in long path length cells, an EA spectrum in a cell of normal path length, the window material and path lengths of these cells, the molar volume of the liquid, the number of anchor points and their wavenumbers.

Procedure

- 1. The K values at the anchor points are calculated exactly as in program ANCHORPT except that Eq. 6.2 is used to calculate AA_R instead of the equations in appendix II.
- 2. The calculation of the k spectrum from the EA spectrum in a cell of normal length is started by following steps 2 to 4 of program RNJ46A, except that AA_R is calculated by Eq. 6.2. The result is the baseline-corrected absorbance spectrum, AS_I , which is the approximation to A_{10} given by this method.
- 3. The k spectrum is calculated, as in step 6 of program RNJ45A, as $k = \frac{2.303}{4\pi \ell \tilde{v}} AS_1$.
- 4. The $E_{\rm m}$ spectrum is calculated as $E_{\rm m} = \frac{AS_{\rm I}}{C\ell}$, where ℓ is the molar concentration and ℓ is the path length of the cell.

Comments about the use of the programs

Both RNJ46A and WIND8 yield useful k values in those regions of the EA spectrum in which $0.2 \le EA \le 2.0$. Other regions of the EA spectrum are processed by the programs but do not yield useful values of k. The final k spectrum is assembled by averaging and merging results from EA spectra in cells of sufficiently different lengths to give useful k values throughout. The final n spectrum is calculated from the final k spectrum by Kramers-Kronig transformation and the value of n at the high-wavenumber limit of the k spectrum or, ideally, when it is known¹³, the value of $n_{\rm cl}$ at each wave number, where $n_{\rm cl}$

is due solely to electronic absorption¹³.

6.6 - Appendix II

$$T_{abc} = \frac{T_a T_{ba}}{1 - R_a R_{ba}}$$

Where: $T_{ba} = \frac{T_a T_b}{1 - R_a R_b}$ $R_{ba} = R_b + \frac{R_a T_b^2}{1 - R_a R_b}$

$$T_a = \frac{4n_w}{(1+n_w)^2}$$
 $R_a = \left\{\frac{1-n_w}{1+n_w}\right\}^2$

$$R_{b} = \left| \frac{r_{12} (1 - e^{-i\delta})}{1 - r_{12}^{2} e^{i\delta}} \right|^{2} \delta = 4\pi \tilde{v} \ell \hat{n}_{s} \qquad r_{12} = \frac{\hat{n}_{s} - n_{w}}{\hat{n}_{s} + n_{w}}$$

$$T_{b} = \left| \frac{t_{12} t_{21}}{1 - r_{12}^{2} e^{i\delta}} \right|^{2} \qquad t_{12} = \frac{2 \hat{n}_{s}}{\hat{n}_{s} + n_{w}} \qquad t_{21} = \frac{2 n_{w}}{n_{w} + \hat{n}_{s}}$$

 \hat{n}_{\bullet} and n_{w} are the complex refractive index of the sample liquid and the real refractive index of the window, respectively. ℓ is the path length of the cell.

6.7 - References

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Chapter 7 - A Compact Table for the Publication of Infrared Spectra That Are

Ouantitative on Both Intensity and Wavenumber Axes*

7.1 - Introduction

Infrared spectroscopists have traditionally reported their results as a spectrum in graphical form and as a table of the wavenumbers of the peaks and other spectral features with their assignment and points of interest. The ordinate values are usually measures of the percentage of the incident light that is transmitted or reflected, or the logarithmic forms of these quantities, such as the absorbance. These ordinate values have typically been put on record simply by noting, next to the wavenumber, that a feature is strong (s) or weak (w), etc. This approach has usually been employed because accurate knowledge of the wavenumber axis has been possible since the publication of the IUPAC "green" and "blue" books but the ordinate axis has been trusted only to give a qualitative indication of the relative intensities. The ordinate axis could not be trusted to give an accurate measure of the amount of radiation absorbed or reflected.

Several workers have made quantitative intensity measurements. Even in these cases it has been usual for the results to be presented as figures of the spectra and tables of the wavenumber and intensity values at the peaks and other spectral features³, and

^{*} A version of this chapter has been published. Bertie, Jones and Apelblat, Appl. Spectrosc., 47, 1989 (1993).

sometimes tables of the areas under the bands in the spectra. Although some authors have presented tables of pairs of wavenumber and intensity values over specific bands in the spectrum⁵, and several authors⁶ have presented such data over the entire spectrum at relatively large intervals, it has not been the practice to do so. The graphs are needed to allow one to see the form of the spectrum quickly, but quantitative values can only be recovered with much labour and very limited accuracy unless they are tabulated.

As long as the accuracy of the intensity information was low, these practices were acceptable, if annoying, to those who wanted to recover and use quantitative information that was available only as graphs. However, today it is possible to measure intensities with an estimated accuracy of 1 or 2%. To place such information on the record for future use requires that both wavenumber and intensity information be given numerically throughout the spectrum. This is particularly the case if the real and imaginary refractive index spectra are measured. These are fundamental physical properties of the sample, and all other optical properties, such as the dielectric constants of the sample, or the result of any spectroscopic experiment on the sample, can be calculated if these two refractive index spectra are available in numeric form.

Clearly the most compact way of presenting spectra numerically is in digital form, either on diskettes or on a generally accessible data base. However, the organization of and confidence in long-term storage by such digital means have not reached the state that the scientific record can dispense with the paper copy.

The difficulty with paper copy is that a spectrum that is digitized at 0.5 cm⁻¹

intervals from 5000 to 500 cm⁻¹ contains 9000 spectral points, i.e. 9000 wavenumber, \widetilde{v} , and intensity, $Y(\widetilde{v})$, pairs. Scientific journals can print sixty lines on each page, and six double columns \widetilde{v} and $Y(\widetilde{v})$ can be put on each line, to give 360 spectral points per page. The table for each spectrum thus requires 25 journal pages. If real, $n(\widetilde{v})$, and imaginary, $k(\widetilde{v})$, refractive index spectra are both to be presented, four triple columns \widetilde{v} , $k(\widetilde{v})$, and $n(\widetilde{v})$ can be put on each line, to give 240 data triplets per page. The table then requires 38 journal pages. These tables are clearly too large to be published in a spectroscopic journal. For publication in a journal for the scientific record, the spectral information must be compacted in such a way that the table is short and readable but allows the original spectrum to be recovered with at least its experimental accuracy.

We describe in this paper a procedure that enables quantitative spectroscopic data to be published over the entire spectrum, by compressing spectral data into a Compact Table. This table contains sufficient information to allow the spectrum to be recovered without loss of accuracy, yet is sufficiently concise that journals can realistically be expected to print it. Further, the table is readable, so that individual numerical values can be recovered from it with minimal effort. We also present a Fortran program for creating the table from a digital spectrum, and a Fortran program that reads the tabulated data and recreates the original spectrum. The creation of the Compact Table, and the recovery of the spectrum from it, is demonstrated for the real and imaginary refractive index spectra, $n(\tilde{v})$ vs. \tilde{v} and $k(\tilde{v})$ vs. \tilde{v} and the molar

absorption coefficient spectrum, $E_m(\tilde{v})$ vs. \tilde{v} , of liquid chlorobenzene⁷. Compact Tables have been used recently to present spectra of the refractive indices and molar absorption coefficient of liquid benzene⁸ and the refractive indices of liquid methanol⁹. The effectiveness of the Compact Table is particularly well illustrated by Table 2 of reference 9. The one-page table contains all of the information needed to recover the absorption intensities of liquid methanol at 1 cm⁻¹ intervals between 8000 and 2 cm⁻¹, accurate to about 2%.

7.2 - Construction of a Compact Table

There are two aspects to the procedure. First, the number of spectral points that are reperied in the contains the reduced spectrum.

The major consideration in reducing the number of spectral points is to ensure that the tabulated information allows recovery of the original spectrum without reducing the accuracy below that of the measurements. Our experimental values of the imaginary refractive index or molar absorption coefficient are believed accurate within $\sim \pm 2.5\%$, so that the original spectrum must be recovered from the tabulated data within $\pm 2.5\%$. In fact a stricter criterion is applied which ensures recovery of nearly all of the spectrum to $\leq \pm 1\%$.

In the construction of the table, the number of spectral points is first reduced by a factor of two by ignoring every second point¹⁰. The ordinate values are truncated to the

appropriate number of significant figures and are tabulated. The tabulated values are then read into the recovery program, interpolated back to the original wavenumber spacing and compared with the original spectrum. The same procedure is repeated for reduction factors of 4, 8, 16, 32, etc.. Then the spectrum is divided into regions of constant wavenumber spacing in such a way that the number of spectral points in each region is reduced by the largest factor that allows recovery to ≤1%.

The original real and imaginary refractive index spectra and the original molar absorption coefficient spectrum of chlorobenzene⁷ each contains 9024 points between 4800 and 450 cm⁻¹. The method described above allows each spectrum to be reduced to 2437 points, in 35 regions with different spacing in adjacent regions. Even after such a reduction of the number of spectral points by a factor of almost four, seven pages are needed for a normal ($\tilde{\nu}$ Y) table with 60 lines per page and 6 $\tilde{\nu}$ -Y pairs per line. Such tables are still too long to be published in most journals. In Table 7.1, we present a small part of such a table, from 2008.49 to 1352.82 cm⁻¹ which contains 360 spectral points.

When the spectral points are known to be uniformly spaced, it is wasteful in Table 7.1 to give the wavenumber every time an ordinate value is given. Instead of giving all of the wavenumber values in the spectrum, we divide the table into regions of constant wavenumber spacing, and give the first wavenumber of each line and the wavenumber spacing for that line. A similar procedure with just one region per spectrum has been used for digital data files for nearly two decades, e.g. in files in the

Table 7.1 - Reduced data of imaginary refractive index of chlorobenzene in XY format.

| cm ⁻¹ | k | cm ⁻¹ | k | cm ⁻¹ | k | cm ⁻¹ | k | cm ⁻¹ | k | cm ⁻¹ | ٨ |
|--------------------|---------------------|--------------------|--------------------|--------------------|------------|--------------------|------------|--------------------|----------------------------------|--------------------|------------|
| 2008 49 | 0001953 | 1924 61 | 0009501 | 1706 69 | 0008302 | 1010.27 | (4)17846 | 143313 | 0011030 | | ili-seleci |
| 2007.53 | 0002144 | 1922 68 | 0008343 | 1702 83 | 0008091 | loop to | 0018416 | 1531-20 | OULTOSS | 1450.21 | 0224716 |
| 2006 57 | 0002344 | 1920 75 | 0007323 | 1608 98 | 0007158 | 10018 24 | (4)19(6)5 | 1529.27 | (901,2259) | | 0265342 |
| 2005 60 | 0002531 | 1918 82 | 0006468 | 1695 12 | (A)()()(A) | 12012.38 | 1,01,000 | 1527.34 | (41 <u>1, 2</u> 064) | | 01110 |
| 2004 64 | 0002683 | 1916 89 | 0005778 | 1691 26 | 0005305 | 1000.41 | 0021089 | 1525.42 | 0013976 | | 0403107 |
| 2003 68 | 0002791 | 191497 | 0005273 | 1689 34 | 0005373 | 10015-15 | 0022471 | 1323.40 | 0015155 | | 118141344 |
| 2002 71 | 0002838 | 1913 04 | 0004955 | 1687 41 | (KAC)Q(KA) | 104448 | 0024248 | 1521.56 | out ast | 1.445 38 | 1011180 |
| 2001.75 | 0002822 | 1911 11 | 0004722 | 1685 48 | 0006452 | 1003 52 | (1026-442 | 1519 63 | .60[9]99 | 1-1-11-45 | 0783155 |
| 2000 78 | 0002773 | 1907 25 | 0004558 | 1583.55 | (10)1500-4 | 1602.55 | (10)29(6)9 | 1517.70 | 0021886 | 1-1-1-1-0 | 0503902 |
| 1999 82 | 0002727 | 1903 40 | 0005119 | 1681 62 | 0005313 | 1601 59 | 0032295 | 1515 77 | 0024005 | 1442 49 | 113 PARS 7 |
| 1998.85 | 0002697 | 1899 54 | 0006196 | 1679 69 | 0004872 | 1000 63 | 0036169 | 1513.85 | (4)26776 | 1441 53 | 0244700 |
| 1997 89 | .0002690 | 1895 68 | 0008208 | 1677 76 | 0004494 | 1500 66 | 0041135 | 1511/92 | 0028190 | 1440 56 | 0191549 |
| 1996.93 | .0002719 | 1891 82 | 0011981 | 1675 84 | 0004521 | 1598 70 | (4)47675 | 150000 | 0020493 | [439.60 | 0163045 |
| 1995 96 | 0002790 | 1887 97 | 0017996 | 1673 91 | 0004698 | 1597 73 | 0056709 | 1508.06 | 0031955 | 1438 63 1437 67 | 0137411 |
| 1995.00 | 0002915 | 1884 11 | 0022907 | 1671 98 | 0005026 | 1596 77 | 0069249 | 1506 13 | (N)3の33か (N)4 (N) | 1430 71 | 01085718 |
| 1994 03 | 0003113 | 1880 25 | 0023773 | 1670 05 | 0005349 | 1595 81 1594 84 | 0086147 | 1504-20 1502-27 | (M. 644) | 1435 74 | (KRIPA) |
| 1993.07 | 0003400 | 1876 40 | 0022514 | 1668.12 | 0005784 | 1593.88 | 0133269 | 1500 35 | (1074934 | 1413-81 | (XX)2X/2 |
| 1992.10 | 0003784 | 1872.54 | 0024329 | 16/6 19 | 0006082 | 1593.66 | 0160376 | 1498 42 | CKNIF CKNI | 1431 89 | (RFI1926 |
| 1991.14 | 0004231 | 1868 68 | 0031992 0041122 | 1664.27 1662.34 | 0006509 | 1501.95 | 0186616 | 1496 49 | 0104603 | 1429.96 | (M)35584 |
| 1990.18 1989.21 | .0004645 0004927 | 1864 83 1860 97 | 0043739 | 1660 41 | 0007049 | 1500.08 | 0211753 | 1494 56 | 0111881 | 1428 03 | Costal CKI |
| 1989 21 | 0004927 | 1857 11 | 0034551 | 1658 48 | 0007049 | 1590 02 | 0238395 | 1492 63 | 0122996 | 1426 10 | 0026083 |
| 1987 28 | 0005080 | 1853.26 | 0021042 | 1656 55 | 0009286 | 1589 06 | 0270981 | 1490 70 | 0141691 | 1424 17 | 0022237 |
| 1986.32 | .0005192 | 1849.40 | 0012062 | 1654 62 | 0011378 | 1588 09 | 0316606 | 1-188 78 | 0167832 | 1422-24 | 0018924 |
| 1985.36 | .0005481 | 1845 54 | 0007463 | 1652 69 | 0014265 | 1587 12 | 0388300 | 1486 85 | 0206693 | 1420 31 | 0016236 |
| 1984 39 | .0005594 | 1841.68 | 0005319 | 1650 77 | 0017320 | 1586 16 | 0511551 | 1484 92 | 0274626 | 1418 39 | 0014137 |
| 1983.43 | .0005594 | 1837 83 | 0004480 | 1648 84 | 0020280 | 1585 20 | 0686019 | 1483 95 | 0330786 | 1416.46 | 0012710 |
| 1982.46 | 0005639 | 1833 97 | 0004555 | 1646 91 | 0022225 | 1584 23 | 0803482 | 1482 99 | 0414312 | 1414 53 | 0011619 |
| 1981.50 | 0005629 | 1830 11 | 0005505 | 1644 98 | 0022633 | 1583 27 | 0801655 | 1482 03 | 0543342 | 1412 60 | 0010560 |
| 1980.53 | 0005661 | 1826.26 | .0006608 | 1643 05 | 0021739 | 1582 31 | 0734641 | 1481 06 | 0755927 | 1410.67 | 0009829 |
| 1979.57 | 0005771 | 1822.40 | 0007380 | 1641.12 | 0020225 | 1581 34 | 0628704 | 1480 10 | 1121719 | 1408 74 | ондачко |
| 1978.61 | 0005994 | 1818.54 | .0008833 | 1639 20 | 0019179 | 1580 38 | 0511596 | 1479 13 | 1712698 | 1406.82 | 0009162 |
| 1977.64 | 0006344 | 1814.69 | .0010684 | 1637 27 | 0019527 | 1579 41 | 0405318 | 1478 17 | 2365395 | 1404 89 | 0000155 |
| 1976.68 | 0006842 | 1810.83 | 0012481 | 1636 30 | 0019380 | 1578 45 | 0319523 | 1477 20 | 2466504 | 1402.96 | (иинти) |
| 1974.75 | 0008319 | 1806.97 | 0013892 | 1635 34 | 0018158 | 1577 48 | 0255474 | 1476 24 | 2134474 | 1401 03 | 0009762 |
| 1972.82 | .0010416 | 1803.12 | .0015481 | 1634 37 | 0016314 | 1576 52 | 0210303 | 1475.28 | 1490597 | 1399 10 | 0010468 |
| 1970.89 | 0012972 | 1799 26 | 0018692 | 1633 41 | 0014709 | 1575 56 | 0180213 | 1474 31 | 1012828 | 1397 17 | 0011805 |
| 1968.96 | | 1795 40 | 0024781 | 1632 45 | 0013624 | 1574 59 | 0161671 | 1473 35 | 0734671 | 1395.24 | 0014104 |
| 1967.04 | | 1791 54 | 0032260 | 1631.48 | 0013055 | 1573 63 | 0151757 | 1472 38 | 0576492 | 1393 32 | 0017663 |
| 1965.11 | 0019811 | 1787.69 | .0035527 | 1630 52 | 0013077 | 1572 66 | 0147124 | 1471 42 | 0463420 | 1391.39 | 0022515 |
| 1963.18 | | 1783.83 | 0032802 | 1629 55 | 0013453 | 1571 70 | 0143540 | 1470 45 | 0363176 | 1389 46 | 0027548 |
| 1961.25 | | 1779.97 | .0027101 | 1628 59 | 0014313 | 1569 77 | 0137425 | 1469 49 | 0273849 | 1387 53 | 0030219 |
| 1959.32 | | 1776.11 | .0020662 | 1627.62 | 0015765 | 1567 84 | 0141497 | 1468 53 | 0208967 | 1385 60 | 0029408 |
| 1957 39 | | 1772.26 | 0016335 | 1626 66 | 0017782 | 1565 91 | 0147272 | 1467 56 | 0166134 | 1383 67 | 0026547 |
| 1955.46 | | 1768.40 | 0011537 | 1625 70 | 0020285 | 1563 99 | 0134075 | 1466 60 | 0137121 | 1381-75 | 0023349 |
| 1953.54 | .0021407 | 1764.55 | .0008596 | 1624.73 | 0022934 | 1562 06 | 0107829 | 1465 63 | 0117276 | 1379 82 | 0021122 |
| 1951.61 | .0023646 | 1760.69 | .0006972 | 1623 77 | 0025175 | 1560 13 | 0082193 | 1464 67 | 0103099 | 1377 89 | 0020582 |
| 1949.68 | | 1756.03 | .0006177 | 1622.80 | 0026380 | 1558 20 | 0062247 | 1463 71 | 0092746 | 1375 96 | |
| 1947.75 | .0032198 | 1752.98 | 0006227 | 1621.84 | 0026583 | 1556 27 | | 1462 74 | | 1374 03 | 0025594 |
| 1945.82 | .0037111 | 1749.12 | 0007161 | 1620.88 | 0025862 | 1554 34 | | 1461 78 | | 1372 10 | 0029269 |
| 1943.89 | .0040454 | 1745.26 | .0009533 | 1619 91 | 0024562 | 1552 41 | 0027762 | 1460 81 | 0078293 | 1370 17 | 0030159 |
| 1941.96 | .0040582 | 1741.40 | 0014245 | 1618 95 | 0023009 | 1550 49 | | 1459 85 | | 1368 24 | 0027181 |
| 1940.04 | .0037969 | 1737.55 | 0022052 | 1617.98 | | 1548 56 | | 1458 88 | | 1366 32 | |
| 1938.11 | .0033506 | 1733.69 | 0031242 | 1617 02 | 0020018 | 1546 63 | | 1457 92 | | 1364 39 | |
| 1936.18 | .0028268 | 1729.83 | .0034346 | 1616.05 | 0018928 | 1544 70 | 0014374 | 1456 96 | | 1362 46 | |
| 1934.25 | .0023159 | 1725.98 | .0029111 | 1615.09 | 0017998 | 1542 77 | | 1455 99 | | 1360 53 | |
| 1932.32 | .0018723 | 1722.12 | .0021053 | 1614.13 | 0017440 | 1540 84 | 0012203 | 1455 03 | | 1358 60 | |
| 1930.39 | .0015193 | 1718.26 | .0014449 | 1613.16 | .0017242 | 1538 92 | 0011566 | 1454 06 | | 1356 68 | |
| 1928.47 | .0012618 | 1714.41 | 0010508 | 1612.20 | 0017166 | 1536.99 | 0010871 | 1453 10 | 9148860 | 1354 75 | |
| 1926.54 | 4 .0010844 | 1710.55 | .0008674 | 1611.23 | .0017394 | 1535 06 | 0010687 | 1452 13 | 0171278 | 1352 87 | 0005569 |

| Table 7.2 - The Compact | Table of the imaginary | refractive index of chlorobenzene.4 |
|-------------------------|----------------------------|-------------------------------------|
| Table ' 2 - The Combact | I acic of the initiation i | icitactive index of childrene. |

| um i | XE | YE. | . 4 | _1 | 2 | 3 | _4 | 5 | 6 | 7 | 8 | 9 | 19 | 11 | 12 | 13 | 14 | 15 | 16 |
|----------|---------|-----|------|--------|-------|------|-------|---------|------|--------|------|------|------|-------|------|------|------|------|-------|
| 21/18/49 | 9 | -7 | 1953 | 2144 | 2344 | 2531 | 2683 | 2791 | 2838 | 2822 | 2773 | 2727 | 2697 | 2690 | 2719 | 2790 | 2915 | 3113 | 3-490 |
| 1702.10 | 9 | 7 | 3784 | 4231 | 21.45 | 1727 | sogn | 51/92 | 5309 | 5.48] | 5594 | 5642 | 5639 | 5629 | 500 | 5771 | 5994 | 6344 | <845 |
| 1974.75 | 1 | .6 | 832 | 1942 | 1297 | 1567 | 1809 | 1981 | 2062 | 2064 | 2028 | 2002 | 2028 | 2141 | 2365 | 2731 | 3220 | 3711 | 4045 |
| 1941 96 | 1 | ., | V358 | 3797 | 3351 | 2827 | 2316 | 1872 | 1519 | 1262 | 1684 | 950 | 834 | 732 | 547 | 578 | 527 | 496 | 472 |
| 1907.25 | <u></u> | 1, | 440 | 5.2 | 0.20 | 821 | 11.48 | زاران ز | 2291 | 2377 | 2251 | 2433 | 3199 | 4112 | 4374 | 3455 | 21/4 | 1206 | 746 |
| 1841 68 | 2 | .14 | 532 | 448 | 456 | 550 | 161 | 738 | 883 | 1068 | 1248 | 1389 | 1548 | 1869 | 2478 | 3226 | 3553 | 3280 | 2710 |
| 1776 11 | 2 | 1, | 2066 | 1634 | 1154 | 860 | 697 | 618 | 623 | 716 | 953 | 1425 | 2205 | 3124 | 3435 | 2911 | 2105 | 1445 | 1051 |
| 1710.55 | 2 | -7 | 8674 | 8392 | 8090 | 7157 | 0164 | 53()4 | | | | | | | | | | | |
| 1689 33 | 1 | .7 | 5373 | (1090) | 6452 | 5664 | 5313 | 4872 | 4494 | 4521 | 4698 | 5026 | 5348 | 5584 | 5784 | 6082 | 6509 | 7049 | 7916 |
| 1656 55 | 1 | -6 | 929 | 1138 | 1426 | 1732 | 2028 | 2223 | 2263 | 2174 | 2023 | 1918 | 1953 | | | | | | |
| 1636 30 | O | .1) | 1938 | 1816 | 1631 | 1471 | 1362 | 1306 | 1308 | 1345 | 1431 | 1576 | 1778 | 2028 | 2293 | 2517 | 2638 | 2658 | 2586 |
| 1619 91 | 0 | .6, | 2456 | 2301 | 2139 | 2002 | 1893 | 1800 | 1744 | 1724 | 1717 | 1739 | 1785 | 1842 | 1906 | 1996 | 2109 | 2247 | 2425 |
| 1603.52 | 0 | -5 | 264 | 291 | 323 | 362 | 411 | 477 | 567 | 692 | 861 | 1077 | 1333 | 1604 | 1866 | 2118 | 2384 | 2710 | 3156 |
| 158712 | O | -5 | 3883 | 5116 | 6860 | 8035 | 8017 | 7346 | 6287 | 5116 | 4053 | 3195 | 2555 | 2103 | 1802 | 1617 | 1518 | 1471 | 1435 |
| 1569 77 | 1 | ٠5 | 1374 | 1415 | 1473 | 1341 | 1078 | 822 | 622 | 468 | 357 | 278 | 224 | 186 | 161 | 144 | 132 | 122 | 116 |
| 153698 | 1 | -6 | 1087 | 1069 | 1103 | 1163 | 1226 | 1296 | 1398 | 1516 | 1705 | 1920 | 2189 | _2460 | 2678 | 2819 | 2949 | 3195 | 3634 |
| 1504.20 | 1 | .5 | 441 | 569 | 749 | 931 | 1046 | 1119 | 1230 | 1417 | 1678 | 2067 | 2746 | • | | | | | |
| 1483 95 | 0 | -4 | 331 | 41-1 | 543 | 756 | 1122 | 1713 | 2365 | 2467 | 2134 | 1491 | 1013 | 735 | 576 | 463 | 363 | 274 | 209 |
| 1467 56 | 0 | -5 | 1661 | 1371 | 1173 | 1031 | 927 | 855 | 807 | 783 | 778 | 796 | 837 | 904 | 999 | 1127 | 1291 | 1489 | 1713 |
| 1451 17 | 0 | -1 | 196 | 225 | 265 | 337 | 493 | 800 | 1033 | 783 | 504 | 340 | 245 | 192 | 163 | 137 | 109 | 86 | 70 |
| 1433 81 | 1 | -6 | 5156 | 4193 | 3558 | 3047 | 2608 | 2224 | 1892 | 1624 | 1414 | 1271 | 1162 | 1056 | 983 | 941 | 916 | 919 | 940 |
| 1401 03 | i | -6 | 976 | 1047 | 1181 | 1410 | 1766 | 2252 | 2755 | 3022 | 2941 | 2655 | 2335 | 2112 | 2058 | 2214 | 2559 | 2927 | 3016 |
| 1368 24 | 1 | -6 | 2718 | 2224 | 1739 | 1339 | 1042 | 837 | 703 | 615 | 557 | | | | | | | | |

a - The column headed cm⁻¹ contains the wavenumber of the first $k(\tilde{\nu})$ value in the row. The columns headed XE and YE contain the X-exponent and the Y-exponent, respectively, for the row. The columns headed $0, 1, 2, \dots 16$, contain the ordinate values, and the headings give the indices of the ordinate values in the row. In a row which starts with $\tilde{\nu}(0)$, the wavenumber corresponding to the ordinate indexed J is $\tilde{\nu}(J) = \tilde{\nu}(0) - \frac{15798.002}{16384} \cdot J \cdot 2^{XE}$. The $k(\tilde{\nu})$ values in that row are the ordinate value shown times 10^{YE} . Thus the entry indexed 16 in the first row of the table shows that $k = 3400 \times 10^{-7} = 3.40 \times 10^{-4}$ at $\tilde{\nu} = 2008.49 - \frac{15798.002}{16384} \cdot 16 \cdot 2^0 = 1993.06$ cm⁻¹.

JCAMP¹¹ format.

We achieve further reduction in space by multiplying each ordinate value on a line by 10^{-YE} , to create integer ordinate values with the appropriate number of significant figures, and reporting these integer values and the Y-exponent, YE, for the line.

We call the resulting table a Compact Table. It contains the 2437 Y values and

Table for chlorobenzene occupies 2.5 pages. The complete Compact Table for the imaginary refractive index of chlorobenzene is given elsewhere⁷. Table 7.2 presents the small part of it that contains all of the information given in Table 7.1.

In Table 7.2, the first column is labeled "cm⁻¹" and in each row it contains the wavenumber, $\tilde{v}(0)$, of the first ordinate value in the row. This ordinate value is in the column labeled "0". The second column is labeled "XE" for X-exponent, and in each row contains the exponent that is used to calculate the wavenumber spacing between the ordinate values in that row. The third column contains YE, the Y-exponent used to calculate the ordinate values in the row. The remaining column headings, 0, 1, 2, 3, wrough 16, are the indices, J, of the ordinate values in that row. Under these seventeen columns are the seventeen Y values, each presented as an integer of three or four digits with leading zeros omitted.

The ordinate values in a row are obtained from the entries by $Y(J) = (\text{entry})^{Y(J)} = (\text{entry})^{Y(J$

$$\widetilde{v}(J) = \widetilde{v}(0) - \frac{15798.002}{16384} \times J \times 2^{XE} = \widetilde{v}(0) - 0.96423^{\circ} \times J \times 2^{XE}.$$
 (7.1)

The factor 15798.002 / 16384 comes from the use of a He/Ne laser in vacuum, wavenumber 15798.002 cm⁻¹, the fact that fast Fourier transforms yield 2^N spectral points, and a decision that the wavenumber spacing for XE=0 should be near 1 cm⁻¹.

| Table 7.3 - The Compact Table of the molar a | bsorption coefficient of chlorobenzene. a.b. |
|--|--|
|--|--|

| cm i | AT: | YF. | , | 1 | 2 | 3 | 1 | 5 | 5 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 |
|----------|-----|----------|-------------|-------|------|-------|-------------|-------|-------|------|------|-------|-------|------|------|------|-------|------------------|------|
| 21/18 45 | 4 | <u> </u> | 2147 | 2400 | 2623 | 2831 | 2999 | 3119 | 3170 | 3151 | 3095 | 3(;42 | 30/07 | 2998 | 3028 | 3105 | 3244 | 3462 | 3779 |
| 1992 19 | G | 4 | 42/14 | 40.00 | 5156 | 5.46% | 5633 | 5755 | 1904 | 5069 | 6191 | 6241 | 6235 | 6221 | 6253 | 6372 | 5615 | 5998 | 7543 |
| 1974.75 | 1 | 1 | 216 | 11-25 | 1426 | 1721 | 1985 | 2171 | 2258 | 2258 | 2217 | 2186 | 2212 | 2332 | 2574 | 2969 | 3-198 | 4028 | 4386 |
| 1941 96 | 1 | 3 | 43% | 41688 | 3622 | 3053 | 2498 | 2018 | 1636 | 1357 | 1165 | 1020 | 895 | 75-4 | 692 | 618 | 563 | 529 | 503 |
| 190725 | 2 | 3 | 445 | c13 | 456 | 868 | 1264 | 1895 | 2-207 | 2493 | 2356 | 2541 | 3334 | 4277 | 4540 | 3579 | 2175 | 1244 | 768 |
| 1841 68 | 2 | . 3 | 5.2/, | 459 | 466 | 562 | 673 | 750 | 896 | 1081 | 1261 | 1400 | 1557 | 1876 | 2481 | 3223 | 3542 | 3264 | 2691 |
| 1776 11 | 2 | - 3 | 2547 | 1615 | 1138 | 846 | 685 | 505 | 509 | 699 | 928 | 1384 | 2137 | 3021 | 3314 | 2802 | 2022 | 1385 | 1005 |
| 171955 | 2 | 4 | 8276 | 7588 | 7684 | 5.782 | 1827 | 50,00 | | | | | | | | | | | |
| 1689 33 | 1 | -4 | 5063 | 5731 | 6065 | 5318 | 4983 | 4564 | 4205 | 4225 | 4286 | 4687 | 4982 | 5195 | 5376 | 5645 | 6035 | ინ 28 | 7323 |
| 1456 55 | 1 | 3 | 87 K | 1050 | 1315 | 1595 | 1865 | 2042 | 2077 | 1992 | 1851 | 1753 | 1783 | | | | | | |
| 1636 30 | 6 | - 3 | 1769 | 1656 | 1487 | 1340 | 1240 | 1188 | 1189 | 1223 | 1300 | 1431 | 1613 | 1839 | 2078 | 2289 | 2388 | 2405 | 2338 |
| 161991 | O | -3 | 2219 | 2078 | 1930 | 1805 | 1706 | 1621 | 1570 | 1551 | 1544 | 1563 | 1603 | 1653 | 1710 | 1790 | 1890 | 2012 | 2170 |
| 1003.52 | 0 | -2 | 236 | 260 | 288 | 323 | 367 | 425 | 505 | 617 | 767 | 958 | 1185 | 1425 | 1657 | 1879 | 2114 | 2402 | 2804 |
| 158712 | 4) | -2 | 3437 | 4526 | 6065 | 7100 | 7079 | 6483 | 5545 | 4509 | 3570 | 2813 | 2248 | 1849 | 1584 | 1420 | 1332 | 1290 | 1258 |
| 156977 | 1 | -2 | 1203 | 1237 | 1286 | 1170 | 939 | 715 | 541 | 406 | 309 | 240 | 194 | 161 | 139 | 124 | 114 | 105 | 99 |
| 1536 98 | i | -3 | 932 | 915 | 943 | 994 | 1046 | 1104 | 1189 | 1288 | 1447 | 1627 | 1853 | 2080 | 2261 | 2377 | 2484 | 2688 | 3052 |
| 1504.20 | 1 | -2 | 370 | 477 | 627 | 778 | 873 | 933 | 1624 | 1178 | 1394 | 1714 | 2274 | | | | | | |
| 1483 95 | O | -1 | 274 | 3-13 | 449 | 624 | 926 | 1413 | 1950 | 2032 | 1757 | 1227 | 833 | 604 | 473 | 380 | 298 | 224 | 171 |
| 1467 56 | 0 | -2 | 1360 | 1122 | 959 | 842 | 757 | 697 | 658 | 638 | 634 | 648 | 680 | 735 | 812 | 914 | 1047 | 1206 | 1387 |
| 1451 17 | 0 | -2 | 1588 | 1818 | 2145 | 2724 | 3980 | 6456 | 8329 | 6309 | 4057 | 2733 | 1968 | 1539 | 1309 | 1103 | 874 | 687 | 560 |
| 1433 81 | 1 | -3 | 4123 | 3348 | 2838 | 2427 | 2075 | 1766 | 1501 | 1286 | 1118 | 1004 | 917 | 832 | 773 | 739 | 719 | 720 | 736 |
| 1401 03 | 1 | -3 | 763 | 817 | 920 | 1098 | 1373 | 1747 | 2135 | 2339 | 2273 | 2049 | 1799 | 1626 | 1582 | 1699 | 1961 | 2240 | 2305 |
| 1368 24 | ı | -3 | 2074 | 1694 | 1324 | 1017 | 791 | 634 | 532 | 464 | 420 | | | | | | | | |

a - The column headed cm⁻¹ contains the wavenumber of the first $E_{\rm m}(\tilde{\nu})$ value in the row. The columns headed XE and YE contain the X-exponent and the Y-exponent, respectively, for the row. The columns headed $0,1,2,\cdots 16$, contain the ordinate values, and the headings give the indices of the ordinate values in the row. In a row which starts with $\tilde{\nu}(0)$, the wavenumber corresponding to the ordinate indexed J is $\tilde{\nu}(J) = \tilde{\nu}(0) - \frac{15798.002}{16384} \cdot J \cdot 2^{XE}$. The $E_{\rm m}(\tilde{\nu})$ values in that row are the ordinate value shown times 10^{YE} . Thus the entry indexed 16 in the first row of the table shows that $E_{\rm m} = 3779 \times 10^{-4} = 3.779 \times 10^{-1}$ at $\tilde{\nu} = 2008.49 - \frac{15798.002}{16384} \cdot 16 \cdot 2^0 = 1993.06$ cm⁻¹.

b - The units of the E_m values are L mole⁻¹ cm⁻¹.

When the point spacings in two consecutive rows are different, the two rows have different XE values and the spacing between the last point of the first row and the first point of the second row is given by the XE value of the second row.

To observe the procedure for obtaining the \tilde{v} and Y values from the Table 7.2, consider the entry in the first row under the column labeled 4. The wavenumber, \tilde{v} ,

corresponding to that entry is given by:

$$\widetilde{v}(J) = \widetilde{v}(0) - 0.964233 \text{ x } J \text{ x } 2^{XE} = 2008.49 - 0.964233 \text{ x } 4 \text{ x } 2^{0} = 2005.63 \text{ cm}^{-1}.$$
 (7.2)

The k value corresponding to 2005.63 cm⁻¹ is 2683 $\times 10^{-7}$, i.e. k(2005.63) = 0.0002683.

Table 7.3 shows the corresponding table for the decadic molar absorption coefficient which was calculated from $k(\tilde{\nu})$ by the equation

$$E_{\text{m}}(\widetilde{\nu}) = \frac{-\log_{10}[I_{\text{t}}(\widetilde{\nu})/I_{\text{o}}(\widetilde{\nu})]}{C d} = \frac{4\pi \widetilde{\nu}}{2.303C} k(\widetilde{\nu})$$
 (7.3)

where C is the concentration, d is the pathlength and $-\log_{10}[I_1(\widetilde{\nu})/I_0(\widetilde{\nu})]$ is the absorbance. Note that I_1 and I_0 must be fully corrected for losses other than by absorption¹⁰. We follow the practice of analytical spectroscopists, by taking C in mol L^{-1} , d in cm and reporting $E_m(\widetilde{\nu})$ in the units L mole⁻¹ cm⁻¹.

The format of the table for the real refractive index is slightly modified. The common exponent YE is not used because real refractive index values are usually smaller than 9.999. Thus, 5 digits are used and it is implicit that the decimal point comes after the first digit of each entry, e.g. 1.5345 is given as 15345 and 0.8976 is given with a leading zero 08976. In Table 7.4 the values of the real refractive index of chlorobenzene are given for the same wavenumber range as in the previous two tables.

A Fortran program for creating the Compact Table and an input file are given in appendix A. Although the program is given with parameters and features to suit the

Table 7.4 - The Compact Table of the real refractive indices of chlorobenzene.a

| cm 1 | XE | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | _10 | 11 | 12 | 13 | 14 | 15 | 16 |
|-------|------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|--------|-------|--------|-------|--------|-------|
| 29/8 | 49 (| 14978 | 14978 | 14978 | 14977 | 14977 | 14977 | 14977 | 14977 | 14977 | 14976 | 14976 | 14976 | 14975 | 14975 | 14974 | 14974 | 14973 |
| 1992 | 10 0 | 14973 | 14973 | 14972 | 14972 | 14972 | 14972 | 14972 | 14972 | 14971 | 14971 | 14971 | 14970 | 14970 | 14969 | 14968 | 14967 | 14967 |
| 1'474 | 75 1 | 14565 | 1474 | 14963 | 14963 | 14964 | 14966 | 14967 | 14969 | 14969 | 14969 | 14968 | 14967 | 14965 | 14965 | 14966 | 14969 | 14975 |
| 1941 | 96 | 14983 | 14989 | 14994 | 14996 | 14997 | 14996 | 14995 | 14993 | 14991 | 14959 | 14988 | 14986 | 14985 | 14983 | 14981 | 1-4980 | 14978 |
| 1997 | 25 2 | 14975 | 14972 | 14968 | 14965 | 14951 | 14960 | 14963 | 14966 | 14966 | 14963 | 14962 | 14968 | 14982 | 1-1995 | 14997 | 14992 | 14986 |
| 1841 | 68 3 | 14981 | 14977 | 14973 | 14969 | 14967 | 14964 | 14962 | 14960 | 14958 | 14957 | 14955 | 14952 | 14951 | 14954 | 14962 | 14970 | 14975 |
| 1776 | 11 3 | 14975 | 14974 | 14971 | 14967 | 14963 | 14959 | 14954 | 14950 | 14945 | 14940 | 14938 | 14942 | 14952 | 14960 | 14963 | 14960 | 14956 |
| 1719 | 55 3 | 14952 | 14948 | 14946 | 14943 | 14940 | 14936 | | | | | | | | | | | |
| 1689 | 33 | 14934 | 14933 | 14932 | 14931 | 14929 | 14927 | 14925 | 14923 | 14921 | 14919 | 14916 | 14914 | 14912 | 14910 | 14907 | 14905 | 14902 |
| 1656 | 55 | 14898 | 14895 | 14893 | 14891 | 14891 | 14892 | 14893 | 14893 | 14892 | 14889 | 14887 | | | | | | |
| 1636 | 30 (| 14887 | 14887 | 14885 | 14883 | 14880 | 14877 | 14874 | 14871 | 14867 | 14864 | 14861 | 1-4859 | 14858 | 14857 | 14857 | 14858 | 14858 |
| 1619 | 9i (| 14857 | 14855 | 14853 | 14849 | 14846 | 14842 | 14837 | 14833 | 14828 | 14823 | 14817 | 14812 | 14806 | 14799 | 14792 | 14785 | 14776 |
| 1603 | 52 (| 14768 | 14758 | 14747 | 14736 | 14723 | 14708 | 14691 | 14673 | 14655 | 14638 | 14625 | 14615 | 14608 | 14600 | 14588 | 14570 | 14544 |
| 1587 | 12 (| 14512 | 14495 | 14562 | 14743 | 14941 | 15099 | 15208 | 15264 | 15278 | 15265 | 15238 | 15206 | 15174 | 15145 | 15121 | 15103 | 15091 |
| 1569 | 71 | 15071 | 15057 | 15064 | 15080 | 15086 | 15079 | 15067 | 15052 | 15037 | 15022 | 15008 | 14995 | 14983 | 14972 | 14962 | 14952 | 14943 |
| 1536 | 98 | 14934 | 14924 | 14915 | 14906 | 14897 | 14887 | 14878 | 14868 | 14857 | 14847 | 14836 | 14825 | 14814 | 14802 | 14787 | 14770 | 14749 |
| 1504 | 20 | 14726 | 14702 | 14680 | 14663 | 14646 | 14620 | 14581 | 14532 | 14469 | 14383 | 14258 | | | | | | |
| 1483 | 95 (| 14173 | 14070 | 13943 | 13794 | 13663 | 13713 | 14237 | 15138 | 15853 | 16190 | 16153 | 16023 | 15912 | 15834 | 15771 | 15700 | 15624 |
| 1467 | 56 (| 15553 | 15490 | 15436 | 15389 | 15346 | 15308 | 15272 | 15239 | 15207 | 15177 | 15147 | 15118 | 15090 | 15063 | 15036 | 15011 | 14986 |
| 1451 | 17 | 14961 | 14929 | 14885 | 14821 | 14749 | 14809 | 15215 | 15593 | 15656 | 15609 | 15551 | 15497 | 15460 | 15438 | 15416 | 15390 | 15365 |
| 1433 | 81 | 15322 | 15290 | 15266 | 15246 | 15229 | 15214 | 15201 | 15188 | 15177 | 15167 | 15157 | 15148 | 15140 | 15132 | 15125 | 15118 | 15111 |
| 1401 | 03 | 15105 | 15098 | 15092 | 15086 | 15080 | 15077 | 15076 | 15078 | 15080 | 15080 | 15078 | 15075 | 15070 | 15066 | 15063 | 15064 | 15067 |
| 1368 | 24 | 15071 | 15071 | 15069 | 15066 | 15062 | 15058 | 15054 | 15051 | 15047 | | | | | | | | |

a. The column headed cm⁻¹ contains the wavenumber of the first $n(\tilde{\nu})$ value in the row. The column headed XE contains the X-exponent for the row. The columns headed $0,1,2,\cdots 16$, contain the $n(\tilde{\nu})$ values with the decimal point implicitly after the first digit in each value, and the headings give the indices of the $n(\tilde{\nu})$ values in the row. In a row which starts with $\tilde{\nu}(0)$, the wavenumber corresponding to the ordinate indexed J is $\tilde{\nu}(J) = \tilde{\nu}(0) - \frac{15798.002}{16384} \cdot J \cdot 2^{XE}$. Thus the entry indexed 16 in the first row of the table shows that n = 1.4973 at $\tilde{\nu} = 2008.49 - \frac{15798.002}{16384} \cdot 16 \cdot 2^0 = 1993.06$ cm⁻¹, and the entry indexed 4 in the row which starts with 1368.24 cm⁻¹ shows that n = 1.5062 at $\tilde{\nu} = 1360.53$ cm⁻¹.

format of our spectral files 10, it can be easily modified for other file formats.

Before we leave the construction of the Compact Table, a few words about significant figures are appropriate. If ordinate values are known to 1% they should be tabulated with three significant figures. If such numbers are later plotted, however, they frequently cause steps in the graph which were not present in the original spectrum.

The steps arise because information was lost in the truncation to three significant figures, in that the relative accuracy from point to point is much better than the absolute accuracy that can be assigned to each ordinate. (the wavenumbers are taken to be essentially exact, as they are from a Fourier transform spectrometer after proper calibration).

Accordingly, we use a number of significant figures that is acceptable in relation to the ordinate accuracy but yields a plotted spectrum free of steps.

7.3 - Recovery of a spectrum from a Compact Table

The tabulated spectrum consists of ordinate values at wavenumbers that are equally spaced in each region but have different spacings in different regions. To recover the spectrum at a uniform spacing throughout, piecewise cubic spline interpolation 12 is used to interpolate between intensity values at equal wavenumber spacing, $\Delta \tilde{v} = \tilde{v}_1 - \tilde{v}_2$. Specifically the intensity values between two tabulated points, Y_1 at wavenumber \tilde{v}_1 and Y_2 at wavenumber \tilde{v}_2 , are obtained from a polynomial function $P(\tilde{v}) = a_3 \tilde{v}^3 + a_2 \tilde{v}^2 + a_1 \tilde{v} + a_0$ which is made to satisfy the position conditions $P(\tilde{v}_1) = Y_1$, $P(\tilde{v}_2) = Y_2$, and the slope conditions $P(\tilde{v}_1) = S_1$ and $P(\tilde{v}_2) = S_2$. We have chosen the free parameters S_1 , S_2 , as the first derivatives at S_1 and S_2 because they give better interpolation than the second derivatives. The first derivatives were calculated by $P(\tilde{v}_1) = \frac{Y_2 - Y_0}{2\Delta \tilde{v}}$ and $P(\tilde{v}_2) = \frac{Y_3 - Y_1}{2\Delta \tilde{v}}$, where Y_0 and Y_3 are the neighboring tabulated points of Y_1 and Y_2 . Obtaining the 4 coefficients for the cubic polynomial from the values of

 $P(\tilde{v}_1)$, $P(\tilde{v}_2)$, $P'(\tilde{v}_1)$ and $P'(\tilde{v}_2)$ is straightforward, as is the interpolation between points 1 and 2 from these coefficients.

In order to proceed, one needs to recognize 3 problematic areas for the interpolation: 1) the first point in the spectrum, 2) the last point in the spectrum, and 3) the boundaries between different regions.

When \tilde{v}_1 is the first point in the spectrum, Y_0 at \tilde{v}_0 is not known. We assume Y_0 = Y_1 . This is usually the case for absorption index spectra, because the values at the highest wavenumber are close to or equal to zero, and it is also usually true for real refractive index spectra because n is almost constant at the highest wavenumber.

When \tilde{v}_2 is the last point in the spectrum, Y_3 at \tilde{v}_3 is not known. No generally useful assumption is evident. Values beyond the last point can not be assumed to be constant because the end point is usually determined by the limitations of the spectrometer or sample cell, and many samples absorb significantly at low wavenumbers. Therefore we do not interpolate to the last point, but only to the penultimate point in the table. In most cases the wavenumber spacing in the last region is increased one to four times when the table is constructed, so this practice typically reduces the length of the recovered spectrum by only one to four points.

The interpolation between adjacent regions is not straightforward because the spacing in the two regions is different and the two points used to calculate the slope are not equidistant from the central point. For simplicity, Figure 7.1 shows only 35 points in the middle of the original spectrum, and all Y values are shown equal. These 35

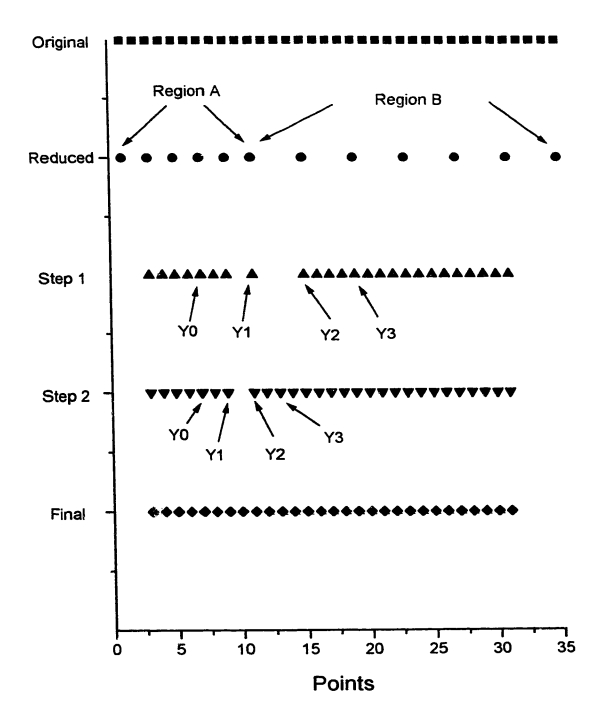


Figure 7.1 - Illustration of the step-wise procedure for interpolating to the original wavenumber spacing at the boundary between regions of different spacings, as described in the text.

points were reduced into two regions, A with 2-point reduction and B with 4-point reduction. Note that the spacing between the last point in region A and the first point in region B is the same as in region B. The interpolation between the two regions A and B is described below. A similar procedure would be followed at the other ends of the two regions to join this 35-point section to the remainder of the original spectrum

The interpolation is first completed in each region from its second point to its next-to-last point. This procedure is straightforward because the first and last points of the region can be used to find the required slopes at the second and next-to-last points, respectively. The result is labeled "Step 1" in Figure 7.1. Then, to interpolate between the two regions, one interpolates the region with the higher reducing factor, region B in the example, to its first point, which is also the last point of region A. As shown in "Step 1" in Fig. 7.1, all values of Y_0 , Y_1 , Y_2 and Y_3 at an equal spacing are known and the interpolation is straightforward between the points denoted by Y_1 and Y_2 . The result is labeled "Step 2" in Figure 7.1. Then, the interpolation of the remaining part of region A can be completed since, again, all values of Y_0 , Y_1 , Y_2 and Y_3 at an equal spacing are known. Note that the interpolation of the more closely spaced region A can not be completed before the interpolation of the less closely spaced region B is completed. The required value labeled Y_3 in "Step 2" is not available before the completion of "Step 2".

The same procedure is followed at all other boundaries to yield the recovered spectrum at the same wavenumbers as the original spectrum, except that it stops at the

last tabulated wavenumber instead of the last wavenumber in the spectrum. The interpolation program and its input file are given in appendix B and can easily be modified to suit any user.

7.4 - The accuracy in the recovered spectrum

Figure 7.2 shows the superimposed original and the recovered absorption index spectra of liquid chlorobenzene⁷. Both spectra are also enlarged to show weak bands. Even with magnification, differences between the original and recovered spectra are not observable. In the lower boxes of Figure 7.3, two bands from this spectrum are expanded, a weak band at high wavenumber and a relatively strong band at low wavenumber. Again, no differences between the original and recovered spectra are visible. To show these differences, the upper boxes of Figure 7.3 show the percent differences between the original and the recovered spectra in these ranges, i.e. the percent inaccuracies of recovery for these bands. Figure 7.4a shows the percent inaccuracies of recovery over the whole spectrum. The magnitudes of the inaccuracies of recovery average 0.2%. Nearly all points in the spectrum are recovered to 1% or better, and the few points that are recovered less accurately are not consecutive.

If the tabulated ordinate values were not truncated to the number of significant figures appropriate to their accuracy, the points in the recovered spectrum would be identical to the points in the original spectrum at the wavenumbers included in the table. To explore the effect of the truncation on the chlorobenzene $k(\widetilde{\nu})$ spectrum, we

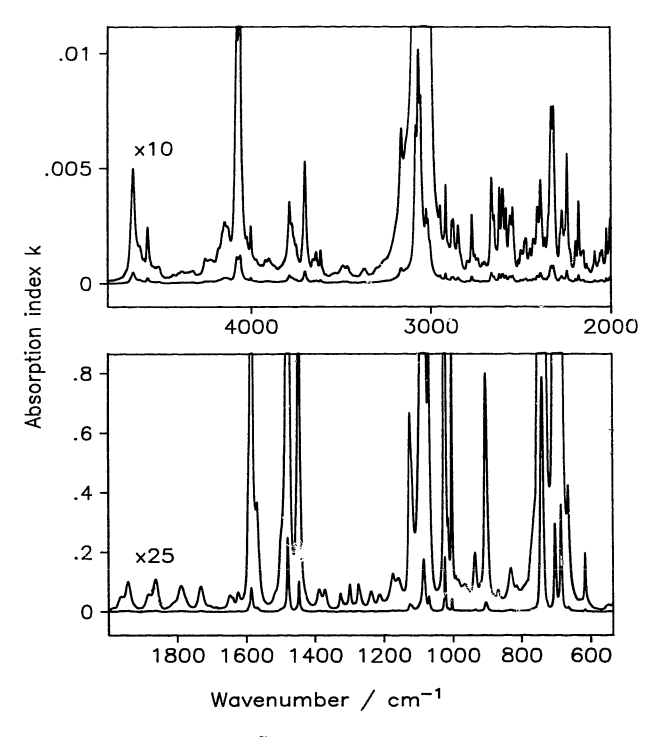


Figure 7.2 - The absorption index, $k(\tilde{\nu})$, spectrum of chlorobenzene at 25°C. In each box, the ordinate scale describes the lower spectrum. For the upper spectra, the ordinate labels must be divided by 10 (upper box) or 25 (lower box). Each curve is the superposition of the original spectrum and that recovered from a Compact Table.

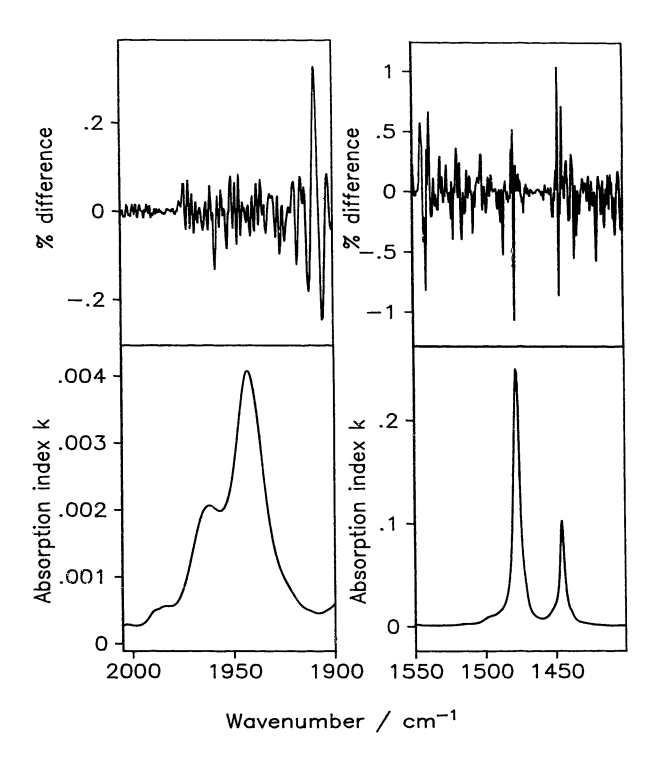


Figure 7.3 - Lower boxes: Two of the bands in Fig. 7.2, with the original spectrum superimposed on that recovered from a Compact Table. Upper boxes: The percent difference between the recovered and original spectra.

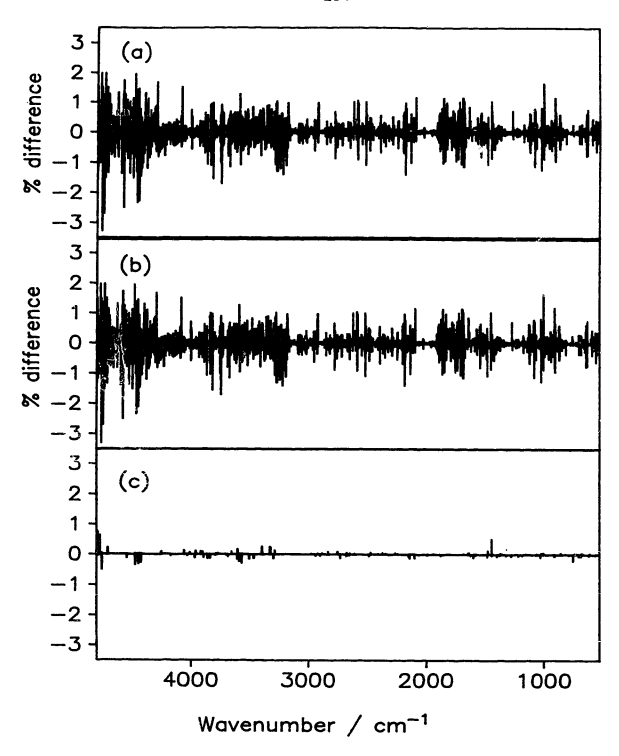


Figure 7.4 - The percent difference between the recovered and original $k(\tilde{\nu})$ spectra when (a) 4 digits or (b) 6 digits were retained in the Compact Table. (c) The difference between the percent differences in the upper two boxes.

constructed Compact Tables with integer ordinate values given to first four and then six significant figures, and recovered the spectrum from each. The accuracies of recovery are shown in Figures 7.4a and 7.4b respectively. The difference between the two is given in Figure 7.4c. In most cases the effect of truncation is less than $\pm 0.1\%$. Again the few disagreements above 0.1% are not in consecutive points.

In addition to the imaginary refractive index spectrum, the real refractive index, $n(\widetilde{v})$, and the molar absorption coefficient, $E_m(\widetilde{v})$, spectra are of common interest. Figure 7.5 shows the recovery of the real refractive index spectrum by two different methods, by interpolation of the tabulated n values (middle box) and by Kramers-Kronig¹³ transformation of the recovered absorption index spectrum (upper box). It is evident that the best method of recovering n values is through the Kramers-Kronig transformation of the recovered k values. However, in both cases the accuracy of recovery is better than $\pm 0.02\%$ at most wavenumbers, and the few larger disagreements are not in consecutive points. Thus, while the Kramers-Kronig transformation of the recovered imaginary refractive index is the best procedure (average accuracy of $\pm 0.005\%$), the $n(\widetilde{v})$ values can be recovered with good accuracy by interpolating the values in a compact $n(\widetilde{v})$ table.

The molar absorption coefficient spectrum, $E_{\rm m}(\widetilde{\nu})$, can be calculated from the imaginary refractive index, as noted earlier. The accuracy of recovery of $E_{\rm m}$ values was tested by two methods. First, the $E_{\rm m}$ spectrum was calculated from the k spectrum, a Compact Table of $E_{\rm m}$ values was created, the $E_{\rm m}$ spectrum was recovered from the table

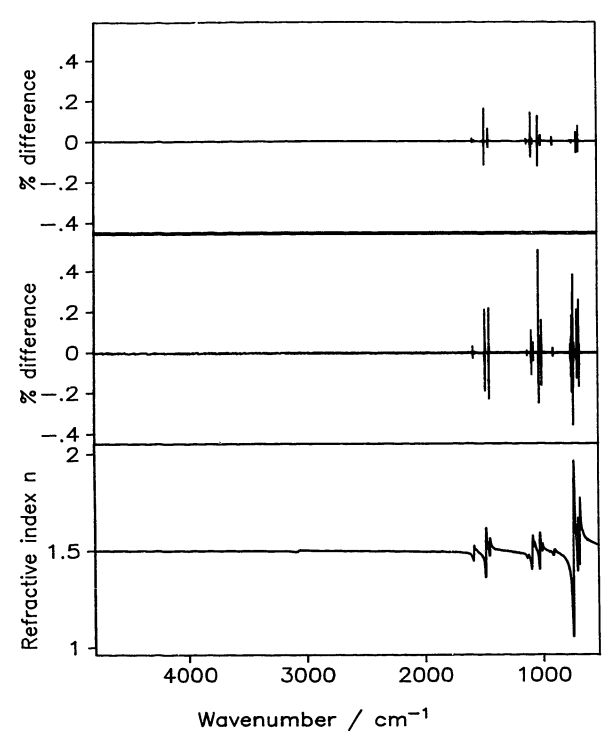


Figure 7.5 - Lower box: The original real refractive index, $n(\tilde{\nu})$, spectrum of chlorobenzene at 25°C superimposed on that recovered from a Compact Table of $n(\tilde{\nu})$ values. Middle box: The percent difference between the recovered and original spectra in the lower box. Upper box: The percent difference between the original $n(\tilde{\nu})$ spectrum and that calculated by Kramers-Kronig transform of the $k(\tilde{\nu})$ spectrum recovered from a Compact Table of $k(\tilde{\nu})$ values.

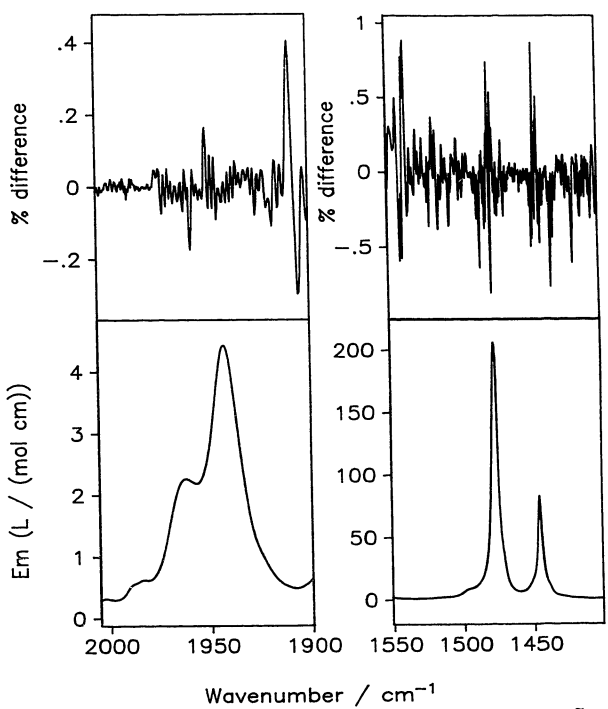


Figure 7.6 - Lower boxes: Two of the bands in the decadic molar absorption coefficient, $E_m(\widetilde{\nu})$, spectrum of chlorobenzene at 25°C. The original spectrum is superimposed on that recovered from a Compact Table of $E_m(\widetilde{\nu})$ values. Upper boxes: The percent difference between the recovered and original spectra.

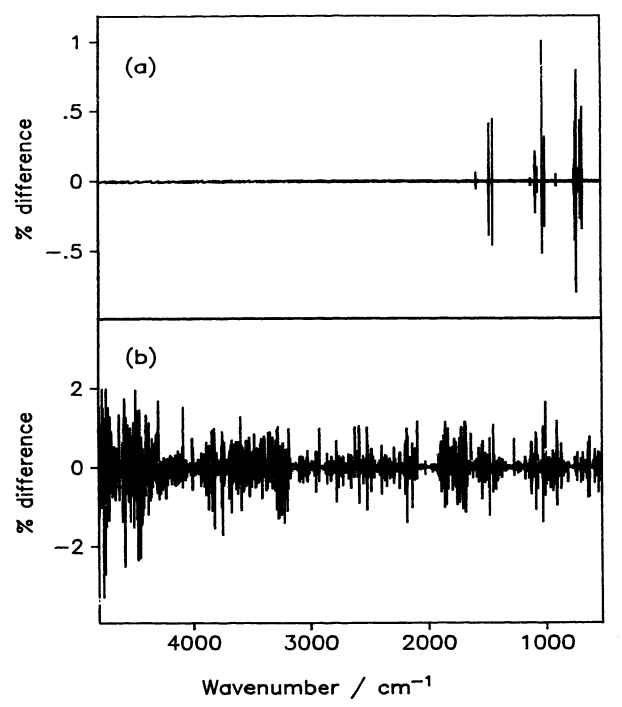


Figure 7.7 - The percent differences between the real (a) and imaginary (b) dielectric constants calculated from the original real and imaginary refractive index spectra and from real and imaginary refractive index spectra recovered from Compact Tables.

by interpolation as described above, and the recovered spectrum was compared with the original. Second, the E_m spectrum was calculated from the original k spectrum and also from the k spectrum recovered from a Compact Table, and the two E_m spectra were compared. The same level of accuracy in the recovered spectrum was obtained from both methods. The lower boxes of Figure 7.6 show two bands of the original molar absorption coefficient spectrum superimposed onto the two bands recovered by the first method. The accuracy of the recovery for these bands is shown in the upper boxes as the percent difference between the original and recovered spectra. The accuracy is expected to be of the same order as the accuracy of the imaginary refractive index, with slight differences because the molar absorption coefficient values are weighted by $\tilde{\nu}$. Comparison of Fig. 7.6 with Fig. 7.2, which shows the same information for the same bands in the imaginary refractive index spectrum, confirms this expectation.

The dielectric constants can be calculated from the refractive indices. The real dielectric constant, $\varepsilon'(\widetilde{v})$, is given by $\varepsilon'(\widetilde{v}) = n^2(\widetilde{v}) - k^2(\widetilde{v})$, while the imaginary part, $\varepsilon''(\widetilde{v})$, also called the dielectric loss, is given by $\varepsilon''(\widetilde{v}) = 2n(\widetilde{v})k(\widetilde{v})$. Figure 7.7 shows the percent differences between the real (a) and imaginary (b) dielectric constants calculated from original and recovered refractive index spectra. For the real dielectric constant, since n > 10k for most of the spectral range, the accuracy of recovery should be that of the n^2 spectrum, i.e. the percent differences should be twice those of the recovered n spectrum. Comparison of Fig. 7.7a and 7.5 shows that this is the case. For the dielectric loss, the recovery should be accurate to the sum of the percent

accuracies of recovery of the real and imaginary refractive indices. Since the percent accuracy of recovery of the k spectrum is much worse than that of the n spectrum, it follows that the accuracy of recovery of the imaginary dielectric constant spectrum (Fig. 7.7b) is essentially that of the k spectrum (Fig. 7.2).

7.5 - Summary

A Compact Table is described to allow numerical reporting of accurate spectral intensity values over the entire infrared spectrum. The table is about one tenth of the size required to report the spectrum in conventional XY format, and allows the intensity values to be recovered with no loss of the experimental accuracy.

The imaginary refractive index and the molar absorption coefficient values can be recovered with average accuracy of about 0.2% and better accuracy than 1% for nearly all points of the spectrum. The real refractive index values can be recovered with average accuracy 0.005%, with most spectral points recovered to better than 0.02%. When the recovered data are used to calculate the dielectric constants, the real dielectric constants have about half of the accuracy of the recovered real refractive index values, and the imaginary dielectric constants have the same degree of accuracy as the imaginary refractive index.

If one has the value of the real refractive index at the highest wavenumber in the spectrum and an accurate program for the Kramers-Kronig transform from $k(\tilde{v})$ to $n(\tilde{v})$,

the $n(\tilde{v})$ spectrum can be recovered from a Compact Table of the absorption index values, $k(\tilde{v})$, to better accuracy than it can be recovered from a Compact Table of $n(\tilde{v})$ values. Thus the intensity properties can be recovered from a single Compact Table of $k(\tilde{v})$ values to at least the accuracies given above. This means, for example, that for liquid methanol a one-page Compact Table contains all of the information needed to recover the complete quantitative intensity information needed to calculate any spectroscopic property between 8000 and 2 cm⁻¹ within the experimental accuracy⁹.

7.6 - Appendix A

7.6.1 - Input file - Comptab.asc

/ no. of spectra, each one to be converted to a table nruns and then for each run / comment head(1) / comment head(2) / input and output filenames filein, fileout / allowed entries: k or n or e spectype / laser wavenumber in cm⁻¹ wl / no. of regions nregion xsreg(i),xfreg(i),factor(i) / starting cm⁻¹, final cm⁻¹, desired spacing in region. one line for each region.

7.6.2 - Summary of variables used in program:

y - a vector containing y values.

xs and xe - starting and ending wavenumbers.

zy(i) - vector zy contains y values after the reduction of the number of points.

z(i,j) - array z contains zy values arranged in rows (i) and columns (j).

nptsreg(i) - number of points in region i.

expo(i) - vector expo contains the Y-exponents.

jz(i,j) - array jz contains values of z(i,j)*10**(-expo(i)) stored as integers.

actspace - the wavenumber spacing in the original file.

nredstep - the reduction factor for the number of points in the region.

newnpts - number of points in the region after reduction.

ind! - number of lines in the region.

ncorect - correction to the index in the original spectral file of the next starting

wavenumber, xsreg(i+1), to ensure that each point is presented only

once and the spacing between xfreg(i) and xsreg(i+1) is that of

factor(i+1).

7.6.3. - Program listings

program Compact Table

implicit real*8(a-h,o-z)

dimension xsreg(35),xfreg(35),factor(35)

reai*4 y(16384),xs,xe,res

dimension z(975,17),zy(16384)

integer*4 nptsreg(35),jz(975,17),expo(975)

integer*2 npts,ier

character*12 filein, fileout

character*192 comm

integer*1 xt,yt

character*1 spectype

character*76 head(2)

open(6,file='comptab.asc')

read(6,*)nruns

do 10 i=1,nruns

read(6,900)head(1)

read(6,900)head(2)

read(6,*)filein,fileout

```
read(6,905)spectype
     read(6,*)wl
     if(spectype.eq.'k') nkflag=0
     if(spectype.eq.'n') nkflag=1
     if(spectype.eq.'e') nkflag=2
     read(6,*)nregion
     do 20 j=1,nregion
20
     read(6,*)xsreg(j),xfreg(j),factor(j)
      open(8, file=fileout)
C
      write table header according to nkflag
      if(nkflag.ne.1) write(8,895)
      if(nkflag.eq.1) write(8,896)
      read data, the following part is for use with files in the .SPC format of Galactic
С
      Industries' SpectraCalc software. modification is needed for a different format of
C
      input data
С
      call readsc(filein,y,npts,xs,xe,xt,yt,res,comm,ier)
      calculate actual spacing
С
      actspace=(xs-xe)/(npts-1)
      start a loop to calculate everything for each region
C
      j=1
      calculate no. of pts in region, the reduction step and finally the new no. of pts in
C
С
      that region
      if((factor(j)/actspace).lt.1) then
       write(*,*)' reduction is impossible, cannot create points'
       stop
       endif
       nredstep=idnint(factor(j)*(wl/16384.d0)/actspace)
 40
       nptsreg(j)=idnint((xsreg(j)-xfreg(j))/actspace+1.0)
       newnpts=idnint((nptsreg(j)-1.0)/nredstep)+1
       xfreg(j)=xsreg(j)-actspace*nredstep*(newnpts-1)
 C
       recalculate nptsreg and newnpts
```

```
nptsreg(j)=idnint((xsreg(j)-xfreg(j))/actspace+1.0)
      newnpts=idnint((nptsreg(j)-1.0)/nredstep)+1
      ntype=nint(dlog(factor(j))/dlog(2.0))
      create a zy array to store the reduced y data
С
      if(j.eq.1)ncorect=nint((xs-xsreg(1))/actspace+1)
      do 50 l=1, newnots
50
      zy(l)=y((l-1)*raredstep+ncorect)
      calculate number of lines in region j
С
      inline=17
      indl=newnpts/inline
      if((indl*inline).ne.newnpts)indl=indl+1
      iflag=0
      create a matrix, z, to store the zy data
C
      do 60 \text{ m}=1,\text{indl}
      do 70 k=1,inline
70
      z(m,k)=zy((m-1)*inline+k)
C
      find 10<sup>n</sup> exponent factor for the line
      if(m.eq.indl) iflag=1
      if(nkflag.ne.1) then
      call findexpo(z,m,expo,iflag,newnpts)
C
      multiply by the exponent factor for the line expo(m)
      do 80 k=1,inline
80
      jz(m,k)=ifix(z(m,k)*10.**(-expo(m))+0.5)
      else
      do 81 k=1,inline
81
      jz(m,k)=ifix(z(m,k)*10000)
      endif
      write line to file
C
      zxline=xsreg(j)-actspace*((m-1)*inline)*nredstep
85
      if(nkflag.ne.1) then
      if(m.ne.indl) write(8,925)zxline,ntype,expo(m),(jz(m,k),k=1,inline)
```

```
if(m.eq.indl) write(8,925)zxline,ntype,expo(m), (jz(m,k),k=1,newnpts-(m-1)*inline)
      else
      if(m.ne.indl) write(8,935)zxline,ntype,(jz(m,k),k=1,inline)
      if(m.eq.indl) write(8,935)zxline,ntype,(jz(m,k),k=1,newnpts-(m-1)*inline)
      endif
60
      continue
      calculate the reduction step for the next region, adjust the starting wavenumber for
С
      that region and go back to statement 40 to redo the whole process for the next
С
      region
С
      j=j+1
      if(i.le.nregion) then
      nredstep=idnint(factor(j)*(wl/16384.d0)/actspace)
      ncorect=ncorect+nptsreg(j-1)+nredstep-1
      xsreg(j)=xfreg(j-1)-nredstep*actspace
      goto 40
      endif
      end of loop to calculate everything for each region
С
10
       continue
895 format(1x,'cm-1',2x,'xe',1x,'ye',3x,'0',4x,'1',4x,'2',4x,'3',4x,'4',4x,'5',4x,'6',4x,'7',4x,
    + '8',4x,'9',3x,'10',3x,'11',3x,'12',3x,'13',3x,'14',3x,'15',3x,'16')
896 format(1x,'cm-1',4x,'xe',4x,'0',5x,'1',5x,'2',5x,'3',5x,'4',5x,'5',5x,'6',5x,'7',5x, '8',5x,
    + '9',4x,'10',4x,'11',4x,'12',4x,'13',4x,'14',4x,'15',4x,'16')
900 format(a76)
905 format(a1)
 925 format(f7.2,1x,i2,1x,i2,17(1x,i4))
 935 format(1x,f7.2,1x,i2,17(1x,i5.5))
       stop
       end
       subroutine findexpo(z,m,expo,iflag,newnpts)
       Suboutine to find the maximum value of z(j,k) in row j. Then to calculate expo(j),
 С
```

```
such that the entries within that line are multiplied by 10**(-expo(j)) when the
С
     table is created.
С
     implicit real*8(a-h,o-z)
     dimension z(975,17)
     integer*4 expo(975)
     Find the largest entry within a row. Use iflag to deal with the last row, which may
С
     contain fewer entries than inline=17.
С
     zmax=0.e0
     if(iflag.ne.1) then
     do 200 i=1,17
200 zmax=amax1(zmax,z(m,i))
      else
      do 210 i=1, new npts-(m-1)*17
210 zmax=amax1(zmax,z(m,i))
      endif
     compute the exponential factor
C
      if(zmax.ge.10000.0) then
      expo(m)=1
      elseif(zmax.ge.1005 3) then
      expo(m)=0
      elseif(zmax.ge.100.0) then
      expo(m)=-1
      elseif(zmax.ge.10.0) then
      expo(m)=-2
      elseif(zmax.ge.1.0) then
      expo(m)=-3
      elseif(zmax.ge.1.d-1) then
      expo(m)=-4
      elseif(zmax.ge.1d-2) then
      expo(m)=-5
      elseif(zmax.ge.1d-3) then
```

```
expo(m)=-6
elseif(zmax.ge.1d-4) then
expo(m)=-7
elseif(zmax.ge.1d-5) then
expo(m)=-7
elseif(zmax.ge.1d-6) then
expo(m)=-7
else
expo(m)=0
endif
return
end
```

7.7 - Appendix B

7.7.1 - Input file - Trecover.asc

```
nruns
                              / no. of tables, each one to be converted to a spectrum
and then for each run
head(1)
                              / comment
head(2)
                              / comment
filein, fileout
                              / input and output filenames
xs,wl,idespace
                              / first wavenumber in Compact Table, laser wavenumber,
                               and the x-exponent, XE, to give the desired spacing in
                               output spectrum (smallest XE in Compact Table)
spectype
                              / allowed values: k or n or e
nregion
                              / number of regions
nptsreg(i),nfactor(i)
                              / number of points in region, reduction factor in the
                               region. one line for each region.
```

7.7.2 - Summary of variables used in program

y - a vector created to contain the ordinate values calculated from the table.

- a vector with the values of the current line in the table. Z - the value of the i th point of the 4 points that are needed for the t (i) interpolation. - the interpolation factor in region j. nfactor(j) nptsreg (j) - the number of points in the table for region j. - the high wavenumber index for the straightforward interpolation in indxhreg(j) region j. indxlreg(j) - the low wavenumber index for the straightforward interpolation in region j. nonhreg(m,k), nonlreg(m,k) Boundary interpolation is defined by the four indices nonhreg(m, 1), nonlreg(m, 1), nonhreg(m,2) and nonlreg(m,2). m=1 to nregion-1 = index of high wavenumber region at the boundary. k=1 if the point indexed is in region m and k=2 if it is in region m+1.

- a dummy vector used in the interpolation (inter1) routine.

7.7.3 - Program listings

idumnon

```
program trecover
implicit real*8(a-h,o-z)
character*12 filein, fileout
dimension y(16384),z(17),t(4)
integer*4 indxhreg(35),indxlreg(35),nonhreg(35,2),nonlreg(35,2), nptsreg(35),
integer*4 nfactor(35), left(975),idumnon(35)
real*4 r(16384)
integer*1 xt,yt
integer*2 npt
character*76 head(2)
character*80 tablhead
character*80 tablhead
character*192 comm
character*1 spectype
data xt /1/, yt /0/, res /0/, ier /0/
open(6,file='trecover.asc')
```

read(6,*)nruns

```
do 10 i=1,nruns
     read(6,900)head(1)
     read(6,900)head(2)
     read(6,*)filein,fileout
     read(6,*)xs,wl,idespace
     read(6,901)spectype
     if(spectype.eq.'k') nkflag=0
     if(spectype.eq.'n') nkflag=1
     if(spectype.eq.'e') nkflag=2
     read(6,*)nregion
     do 20 j=1, nregion
20
     read(6,*)nptsreg(j),nfactor(j)
      calculate total no. of points (npts), indices of straightforward and boundary
С
С
     interpolation regions.
      npts=1+(nptsreg(1)-1)*nfactor(1)
      indxhreg(1)=1
      indxlreg(1)=1+(nptsreg(1)-2)*nfactor(1)
      if(nregion.gt.1) then
      do 30 j=2,nregion
      nonhreg(j-1,1)=indxlreg(j-1)
      nonlreg(j-1,1)=nonlreg(j-1,1)+nfactor(j-1)
      nonhreg(j-1,2)=nonlreg(j-1,1)
      nonlreg(j-1,2)=nonhreg(j-1,2)+nfactor(j)
      indxhreg(j)=npts+nfactor(j)
      npts=npts+nptsreg(j)*nfactor(j)
      indxlreg(j)=npts-nfactor(j)
30
      continue
      endif
      calculate the original end point xe and the final end point xel
C
      xe=xs-(npts-1)*(wl/16384.d0)*2.**(idespace)
```

```
xe1=xs-(indxlreg(nregion)-1)*(wl/16384.d0)*2.**(idespace)
     read table values
С
     open(9,file=filein)
     read(9,910)tablhead
     do 40 k=1, nregion
     calculate no. of lines in region k (lines) and no. of pts in the last line of region k
C
С
     (left)
     lines=nptsreg(k)/17
      do 45 m=1,lines
     left(m)=17
45
     continue
      if(lines*17.ne.nptsreg(k)) then
      lines=lines+1
     left(lines)=nptsreg(k)-(lines-1)*17
      endif
      start reading region by region, one line at a time and create a temporary array, z,
С
      for the line. Then put the values of the line into the correct position in the y array
С
      do 50 k1=1, lines
      if(nkflag.ne.1) read(9,*)wvjunk,njunk,xn,(z(k2),k2=1,left(k1))
      if(nkflag.eq.1) read(9,*)wvjunk,njunk,(z(k2),k2=1,left(k1))
      do 55 k3=1,left(k1)
  if(nkflag.ne.1) y(indxhreg(k)+(k1-1)*nfactor(k)*17+(k3-1)*nfactor(k))=z(k3)*10**xn
  if(nkflag.eq.1) y(indxhreg(k)+(k1-1)*nfactor(k)*17+(k3-1)*nfactor(k))=z(k3)/10000.
55
      continue
50
      continue
40
      continue
      start interpolating in regions that are straightforward.
C
      do 60 l=1, nregion
      if(1.eq.1) t(1)=y(1)
      if(l.ne.1) t(1)=y(nonhreg(l-1,2))
      t(2)=y(indxhreg(1))
```

```
t(3)=y(indxhreg(l)+nfactor(l))
     t(4)=y(indxhreg(l)+2*nfactor(l))
     ind=indxhreg(l)+2*nfactor(l)
     do 70 m=1,nptsreg(1)-2
     call inter1(y,t,l,m,nfactor,indxhreg)
      adjust t
С
      ind=ind+nfactor(1)
      t(1)=t(2)
      t(2)=t(3)
      t(3)=t(4)
      t(4)=y(ind)
70
      continue
60
      continue
      interpolate in boundary regions. In each boundary do the region with bigger
С
      nfactor first.
С
      do 80 11=2,nregion
      if(nfactor(l1-1).gt.nfactor(l1)) then
      t(1)=y(nonhreg(11-1,1)-nfactor(11-1))
       t(2)=y(nonhreg(11-1,1))
       t(3)=y(nonlreg(11-1,1))
       t(4)=y(nonlreg(11-1,1)+nfactor(11-1))
       idumnon(11-1)=nonhreg(11-1,1)
       call inter1(y,t,l1-1,1,nfactor,idumnon)
       t(1)=y(nonhreg(11-1,2)-nfactor(11))
       t(2)=y(nonhreg(11-1,2))
       t(3)=y(nonlreg(11-1,2))
       t(4)=y(nonlreg(11-1,2)+nfactor(11))
       idumnon(11)=nonhreg(11-1,2)
       call inter1(y,t,11,1,nfactor,idumnon)
       else
       t(1)=y(nonhreg(11-1,2)-nfactor(11))
```

```
t(2)=y(nonhreg(11-1,2))
     t(3)=y(nonlreg(11-1,2))
     t(4)=y(nonlreg(11-1,2)+nfactor(11))
     idumnon(11)=nonhreg(11-1,2)
     call inter1(y,t,11,1,nfactor,idumnon)
     t(1)=y(nonhreg(11-1,1)-nfactor(11-1))
     t(2)=y(nonhreg(11-1,1))
     t(3)=y(nonlreg(11-1,1))
     t(4)=y(nonlreg(11-1,1)+nfactor(11-1))
     idumnon(11-1)=nonhreg(11-1,1)
     call inter1(y,t,11-1,1,nfactor,idumnon)
     endif
     continue
     Write output file. This section to statement 10 assumes Galactic Industries .SPC
     ile format, as used by SpectraCalc and GRAMS softwares. Y values are
     converted to single precision and sent to the .SPC write subroutine. Wavenumber
     information is sent by the starting wavenumber, the ending wavenumber and the
     number of points.
     npt=indxlreg(nregion)
     do 90 l=1,indxlreg(nregion)
     r(1)=y(1)
     comm=head(2)
     call writesc(fileout,r,npt,real(xs),real(xe1),xt,yt,res,comm,ier)
     continue
900 format(1x,a76)
901 format(a1)
910 format(a80)
     stop
     end
     subroutine inter1(y,t,l,m,nfactor,indxh)
```

80

С

С

С С

C

90

10

```
implicit real*8(a-h,o-z)
dimension y(16384),t(4)
integer*4 indxh(35), nfactor(35)

c calculate slopes and coefficients
s1=(t(3)-t(1))/2.0d0
s2=(t(4)-t(2))/2.0d0
a=2*t(2)-2*t(3)+s1+s2
b=3*t(3)-3*t(2)-2*s1-s2
do 700 kindx=1,nfactor(1)-1
ratio=dble(kindx)/dble(nfactor(1))
y(indxh(1)+(m-1)*nfactor(1)+kindx)=a*ratio**3+b*ratio**2+s1*ratio+t(2)
700 continue
return
end
```

7.8 - References

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Chapter 8 - Summary

In this work the real and imaginary refractive index spectra of liquid chlorobenzene and toluene were measured across the entire mid-infrared region at 25°C. To evaluate the systematic errors, a comparison was made with spectra measured by other spectroscopists in different laboratories with different instruments. From this comparison, it was estimated that the accuracy is ±0.2% for the real refractive index and ±2-3% for the imaginary refractive index. The real and imaginary refractive indices are fundamental physical quantities. They were used to calculate other measures of infrared absorption intensities of these liquids such as the molar absorption coefficient, the complex dielectric constant and the complex molar polarizability. The molar absorption coefficient and the imaginary dielectric constant and the imaginary molar polarizability have approximately the same accuracy as the imaginary refractive index, i.e. ±2-3%. The real dielectric constant and the real molar polarizability have approximately twice the error of the real refractive index, i.e. ±0.4%.

The optical constants and the molar absorption coefficient spectra of toluene and chlorobenzene measured in this work, together with those of liquid benzene and dichloromethane measured separately, were used to establish secondary infrared intensity standards for liquids. These standards have been published by the International Union of Pure and Applied Chemistry and can be used to calibrate other intensity measurements.

Molecular properties such as the dipole transition moment and the dipole moment derivative with respect to normal coordinate are more directly reflected in the

or the molar absorption coefficient spectrum. These properties are calculated from the integrated intensities of the bands. In order to obtain reliable integrated intensities, the imaginary molar polarizability spectrum had to be separated first into contributions from different transitions. The separation was achieved by curve fitting the imaginary molar polarizability spectra with bands of classical damped harmonic oscillator shape, then calculating the integrated intensities from the parameters of the fitted bands. In this work, an improved curve-fitting program was developed which allows a better fit to be obtained faster. The accuracy of the integrated intensities is estimated at 3-5% for strong bands and 5-10% for weak bands. The results of this work are corrected for liquid dielectric effects and agree usually within a factor of two with literature values for the gas obtained by experimental or by *ab initio* calculation. In these cases the agreement must mean that the vibrations in question have very similar intrinsic intensities in the liquid and gas phases.

The integrated intensities of the 30 fundamentals of liquid chlorobenzene and toluene were compared with each other because the two compounds have the same molecular skeleton with the exception of the chlorine atom and the methyl group. It was for the two liquids that the integrated intensities of many of the fundamentals or of the sum of two fundamentals that result from the splitting of a degenerate fundamental in benzene, are usually within a factor of 2 and so do not change significantly upon substitution.

There are notable exceptions. In toluene the intensity of the CH stretching vibrations is stronger by a factor of 4.6 than that in chlorobenzene. It appears that the chlorine substitution shifts intensity from the CH stretching vibrations to other vibrations. This shift of intensity is compensated in v_{26} , v_7 , v_{10} and v_{11} , in particular in v_7 and v_{10} , for which the integrated intensities in liquid chlorobenzene are significantly greater than those in toluene, by factors of 2.7, 62.1, 55.5 and 6.2, respectively. It is quite possible that the gain in intensity in these vibrations, is associated with the presence of the electronegative chlorine and with the loss of intensity of the CH stretching vibrations.

The procedure used to determine the optical constants of the liquid from transmission measurements is exact but computationally difficult. A simpler, approximate method was developed to obtain the optical constants of the liquids. In this method the apparent absorbance due to reflection losses is calculated by treating the liquid cell as a single slab of window material. This simplifies the computation and the method is useful for nearly all common liquids in cells with alkali halide windows. For all but the strongest absorption bands the approximate method gives imaginary refractive indices that are within ~1% of those obtained from the exact method. Larger deviations occur when the mismatch between the refractive indices of the sample and the window exceed 0.15

Finally, a method for data reduction and presentation is given. The data is reduced and incorporated into a format that we call Compact Table. The Compact

Table format allows a spectrum to be tabulated in about 1/10 of the space required for a traditional table. The format allows direct retrieval of specific values, and also the retrieval, through a recovery program, of the entire spectrum without loss of intensity and line shape information. Other quantities that are calculated from those recovered by the recovery program, are obtained with the same accuracy as if calculated from the original quantities.

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