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AIR FORCE CAMBRIDGE RESEARCH LABORATORIES

L. G. HANSCOM FIELD, BEDFORD, MASSACHUSETTS

## AFCRL Atmospheric Absorption Line Parameters Compilation

R.A. McCLATCHEY W.S. BENEDICT S.A. CLOUGH D.E. BURCH R.F. CALFEE K. FOX L.S. ROTHMAN J.S. GARING

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## AIR FORCE SYSTEMS COMMAND United States Air Force



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OPTICAL PHYSICS LABORATORY

PROJECT 7670

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Abstract

The report describes a compilation of the molecular spectroscopic parameters for a number of infrared-active molecules occurring naturally in the terrestrial atmosphere. The following molecules are included in this compilation: water vapor; carbon dioxide; ozone; nitrous oxide; carbon monoxide; methane; and oxygen. The spectral region covered extends from less than 1  $\mu$ m to the far infrared, and data are presented on more than 100,000 spectral lines. The parameters included in the compilation for each line are: frequency, intensity, half-width, energy of the lower state of the transition, vibrational and rotational identifications of the upper and lower energy states, an isotopic identification, and a molecular identification.

A discussion is provided separately for each molecular species, indicating the sources and accuracy of the data and a general discussion of how the data were obtained.

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## AFCRL Atmospheric Absorption Line Parameters Compilation

#### **1. INTRODUCTION**

About 10 years ago a program was initiated to compile spectroscopic data on individual vibration-rotation lines of water vapor in the 2.7  $\mu$ m region (Gates, et al, 1964).

This work continued resulting in a publication on the 2.05 and 2.7  $\mu$ m bands of carbon dioxide (Calfee and Benedict, 1966) and a third publication on the 1.9 and 6.3  $\mu$ m bands of water vapor (Benedict and Calfee, 1967). Other workers have published similar results on the 15  $\mu$ m bands of CO<sub>2</sub> (Drayson and Young, 1967), the 9.6  $\mu$ m bands of O<sub>3</sub> (Clough and Kneizys, 1965), the CO bands whose fundamental is near 5  $\mu$ m (Kunde, 1967), the CH<sub>4</sub> bands near 3 and 7.5  $\mu$ m (Kyle, 1968) and the unpublished rotational water data calculated by Benedict and Kaplan in 1959 (see Goody, 1964, p. 184).

About 5 years ago an effort was initiated at AFCRL to continue this work with the aim of providing a complete set of data for all vibration-rotation lines of all naturally occurring molecules of significance in the terrestrial atmosphere. With such data at hand, it would be possible to compute the transmittance appropriate for atmospheric paths by first computing the monochromatic transmittance many times in a finely spaced frequency grid and then degrading the results to any appropriate spectral resolution. Up to now the following molecules have been included in this compilation: (1) water vapor; (2) carbon dioxide; (3) ozone; (4) nitrous oxide; (5) carbon monoxide; (6) methane; and (7) oxygen.

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All of these molecules except oxygen are minor constituents of the atmosphere, but nonetheless represent most of the absorption lines in the visible and infrared. Although there is some evidence for decreasing concentration with height of  $N_2O_2$ , CO, and  $CH_4$ , it is probably reasonable for most purposes to assume that all of these gases except  $H_2O_2$ , and  $O_3$  are uniformly mixed by volume in the atmosphere unless other specific information is available. Table 1 provides concentrations and references for these "uniformly mixed gases". Water vapor and ozone are, of course, not uniformly mixed and an appropriate set of models useful in considering the radiation effects of these gases is provided by McClatchey, et al, 1972.

Constituent	ppm by volume	Reference
CO <sub>2</sub>	330	Fink et al, 1964
N <sub>2</sub> Õ	0.28	Birkland and Shaw, 1959
СО	0.075	Shaw, 1968
CH <sub>4</sub>	1.6	Goody, 1964
0 <sub>2</sub>	$2.10 \times 10^{5}$	Valley, 1965

#### Table 1. Concentrations of Uniformly Mixed Gases in Dry Air

#### 2. DESCRIPTION OF COMPILATION

In order to compute the transmittance due to a given spectral line in the atmosphere it is necessary to describe the absorption coefficient as a function of frequency for each line. The four essential line parameters for each line are the resonant frequency,  $\nu_0$  (cm<sup>-1</sup>), the intensity per absorbing molecule, S(cm<sup>-1</sup>/ molecule cm<sup>-2</sup>) the Lorentz line width parameter,  $\alpha_0$  (cm<sup>-1</sup>/atm), and the energy of the lower state, E<sup>u</sup>(cm<sup>-1</sup>). The frequency,  $\nu_0$ , is independent of both temperature and pressure (except for possibly very small pressure effects of less than 0.01 cm<sup>-1</sup>/atm, which have been ignored here). The intensity, S, is pressure-independent, and, as discussed below, its temperature dependence can be calculated from E<sup>u</sup> and  $\nu$ 

The line half-width at half maximum,  $\alpha$ , is by definition proportional to the pressure, p, and its temperature dependence can be estimated as discussed below. The precise line shape is a matter of some uncertainty, but in the derivation

of line parameters from laboratory measurements, it is customary to start from the Lorentz shape (see Goody, 1964) given in Eq. (1).

(1)

$$k(\nu) = \frac{S\alpha/n}{(\nu - \nu_0)^2 + \alpha^2}$$

 $S = \int k(v) dv$ 

2

The validity of Eq. (1) to describe the true line shape is subject to two limitations. The first, which can be precisely estimated and corrected for by the use of the Voigt shape, occurs when  $\alpha_0 P/\alpha_D < 1.0$  where  $\alpha_D$  is the doppler line width which varies with frequency, temperature and molecular mass as given in Eq. (2).

$$\alpha_{\rm D} = \frac{\nu_c}{c} \left( \frac{2 \, {\rm kT}}{m} \right)^{1/2} = 4.298 \, {\rm x} \, 10^{-7} \, \nu_c ({\rm T/M})^{1/2}$$
(2)

where M = molecular weight and here k = Boltzmann's constant and m = mass of a molecule.

For atmospheric molecules and infrared frequencies, modifications of the Lorentz shape begin to be required at pressures below 10 to 100 mb.

The second limitation concerns possible inadequacies of the Lorentz shape, especially in the distant wings of a line  $(|\nu - \nu_0| >> \alpha)$  (see Winters et al, 1964, and Burch et al, 1969) or when the long-range intermolecular forces responsible for collision broadening are dipole-quadrupole, leading to an exponent 1.75 rather than 2.0 for  $(\nu - \nu_0)$ , (Varanasi, 1972). Throughout this compilation we assume the validity of the Lorentz exponent.

The line intensity is temperature dependent through the Boltzmann factor and the partition function as indicated in Eq. (3) (the induced emission term has intentionally been omitted here),

$$S(T) = \frac{S(T_s) Q_v(T_s) Q_r(T_s)}{Q_v(T) Q_r(T)} \exp \left\{ \frac{1.439E''(T-T_s)}{T_s} \right\}$$
(3)

where E" (in cm<sup>-1</sup>) is the energy of the lower state of the transition and where  $Q_v$ and  $Q_r$  are the vibrational and rotational partition functions. The vibrational partition functions for the most abundant isotopes are given in Table 2. Partition functions for the other isotopes are similar. The temperature dependence of the rotational partition function is given by  $(T/T_s)^j$  where j is also provided in Table 2 ( $T_c$  is taken to be 296°K).

It is also necessary to know the temperature variation of  $\alpha$ . In the absence of specific indications discussed under each molecule, the equation  $\alpha$  (T)/ $\alpha$ (T<sub>S</sub>) = (T/T<sub>S</sub>)<sup>-n</sup>, with n = 1/2, corresponding to the assumption of temperature-independent collision diameters, may be made. The validity of the assumption is more uncertain, the larger the dependence of the diameter on the particular rotation-vibration transition, (that is, it is most unrealistic for H<sub>2</sub>O and the low-J transitions of the other molecules). The theory of Tsao and Curnutte (1954) when applied to the determination of line width for H<sub>2</sub>O lines gives a wide variation of n about the mean value of 0.62 (Benedict and Kaplan, 1959). Measurements made with a CO<sub>2</sub> laser (Ely and McCubbin, 1970) indicate a value of n = 1.0 for the P20 line of the 10.4  $\mu$ m CO<sub>2</sub> band.

Molecule	j	Tempera- ture	175	200	225	250	275	296	325
н <sub>2</sub> О	1.5		1.000	1.000	1.000	1.000	1.000	1.000	1.001
CO <sub>2</sub>	1.0		1.0095	1.0192	1.0327	1.0502	1,0719	1.0931	1.1269
O <sub>3</sub>	1.5		1.004	1.007	1.013	1.022	1,033	1.046	1.066
N <sub>2</sub> O	1.0		1.017	1.030	1.048	1.072	1,100	1.127	1.170
co	1.0		1.000	1.000	1.000	1.000	1,000	1.000	1.000
CH <sub>4</sub>	1.5		1.000	1,000	1.001	1.002	1.004	1.007	1.011
0 <sub>2</sub>	1.0		1.000	1.000	1.000	1.000	1.000	1.000	1.001

Table 2. Vibrational Partition Functions

An examination of Eqs. (1) and (3) indicates that it is necessary to know the  $\nu_0$ , S(T<sub>S</sub>),  $\alpha(P_0, T_S)$  and E" value for each line in order to compute a spectrum. The data compilation described here contains these four quantities for each of the more than 100,000 lines between 1  $\mu$ m and the far infrared belonging to the seven molecular species listed in Table 3. Additional identifying information is also supplied for each line as indicated below.

Molecule	Identification No.	Criterion Intensity* Minimum at T=296K	Existing Intensity Minimum at T=296K
H <sub>2</sub> O	1	$3 \times 10^{-27}$	$3 \times 10^{-27}$
CO <sub>2</sub>	2	$2.2 \times 10^{-26}$	$3.7 \times 10^{-27}$
О <sub>3</sub>	3	$3.5 \times 10^{-24}$	$3.5 \times 10^{-24}$
N <sub>2</sub> O	<b>4 .</b>	$3.0 \times 10^{-23}$	$4.0 \times 10^{-23}$
CO	5	$8.3 \times 10^{-23}$	$1.9 \times 10^{-23}$
CH <sub>4</sub>	6	$3.3 \times 10^{-24}$	$3.3 \times 10^{-24}$
0 <sub>2</sub>	7	$3.7 \times 10^{-30}$	$3.7 \times 10^{-30}$

Table 3. Intensity Criteria for Lines Included in Compilation

\*Units are  $cm^{-1}/(molecule-cm^{-2})$ 

In order to establish the "Criterion Intensity Minimum" values given in Table 3, an extreme atmospheric path was considered, assuming the gas concentrations specified in Table 1 and maximum concentrations over the path of  $3 \times 10^{24}$ molecules/cm<sup>2</sup> for water vapor and  $1 \times 10^{20}$  molecules/cm<sup>2</sup> for ozone. This extreme radiation path was the atmospheric path tangent to the earth's surface, and extending from space to space. Using this criterion, lines yielding less than 10 percent absorption at the line center would normally be omitted. Although this absolute line intensity cutoff was established, it has not always been possible to achieve. In some cases it would have been unrealistic to push calculations to this limit when experimental confirmation fell far short. There are two specific areas in which this absolute cutoff has been violated: (1) In regions of very strong absorption, very weak lines above this absolute limit have been neglected; (2) Q-branch lines below this limit have occasionally been included where it is felt that the accumulation of many weak, closely spaced lines would still produce an appreciable absorption under some atmospheric circumstances. In some cases, (for example,  $CO_2$ ), sufficient laboratory measurements and theoretical work were available so that this limit was exceeded throughout the infrared.

In the past, line intensities have been defined in various units, different for each molecular species. It was common to define water vapor concentration in precipitable cm, or  $g/cm^2$ , in the path in question. On the other hand, the amount of  $CO_2$  and the other uniformly mixed gases in a path were often given in cm-atm of gas at STP. In order to unify the units and ultimately to lead to less confusion, we decided to use the more fundamental quantity, molecules/cm<sup>2</sup> as a measure of absorbing gas abundance along the path. The appropriate conversion factors are:

$$1 (cm-atm)_{cmp} = 2.69 \times 10^{19} molecules/cm^2$$

$$1 \text{ g/cm}^2 \text{ of } H_2 \text{O} = 3.34 \times 10^{22} \text{ molecules/cm}^2$$

It was also decided, as indicated in Table 3, to define line (and band) intensities at 296<sup>o</sup>K, the normal room temperature at which most measurements are made. Intensities of all bands are based on the total number of molecules of a given species of all isotopes in their normal abundance, not on the number of the particular isotope responsible for a given band. Isotopic abundance values for all molecules for which data are provided in the compilation are given in Table 4.

Half-widths of lines have been added where available. Details are discussed in the separate sections on individual molecules. In some cases, it is felt that insufficient data exist to warrant the inclusion of a variable half-width. In these cases, a mean, constant value has been inserted for each molecular species and values are given in Table 5.

A shorthand notation was adopted to identify the various isotopic species. It is easy to understand by considering the following examples for  $CO_2$ :  ${}^{16}O^{12}C^{16}O \equiv 626$ ,  ${}^{16}O^{13}C^{16}O \equiv 636$ , and for  $N_2O$ ,  ${}^{14}N^{15}N^{16}O \equiv 456$ , etc. This same type of code is used for the other five molecules.

н <sub>2</sub> о	161 162 181 171	0.99729 0.000300 0.00204 0.000370	Сн <sub>4</sub> Сн <sub>3</sub> D	211 311 212	0.98815 0.01110 0.00060
co <sub>2</sub>	626 636 628 627 638 637 828	0.98414 0.01105 0.00402 0.000730 0.0000452 0.00000820 0.00000812	0 <sub>2</sub>	66 68 67	0.99519 0.00407 0.00074
0 <sub>3</sub>	666 668 686	0.99279 0.00406 0.00203			
N <sub>2</sub> O	446 456 546 448 447	0,99022 0.00368 0,00368 0.00202 0.00037			
со	26 36 28 27	0.98652 0.01107 0.00202 0.000369			

Table 4. Isotopic Abundances

Table 5, Mean Half-width Values

Molecule	Half-width (cm <sup>-1</sup> /atm)	References
$\begin{array}{c} \mathrm{CO}_2\\ \mathrm{O}_3\\ \mathrm{N}_2\mathrm{O}\\ \mathrm{CO}\\ \mathrm{CH}_4\\ \mathrm{O}_2\end{array}$	0,07 0,11 0,08 0.06 0,055 0,060	Yamamoto et al (1969) Lichtenstein et al (1971) Toth (1971) Bouanich and Haeusler (1972) Varanasi (1971) (see Section 6.3) Burch and Gryvnak (1969)

A standard computer format was adopted for card or card-image input and is outlined below, the numbers between vertical lines representing the columns on an IBM card, and the letter-number combinations representing the computer format.\* The first four quantities are:  $\nu =$  frequency in

Withration ID Date Isotone Molecul	
	е
	<del></del>
1-10   11-20   21-20   20-30   36-70   71-73   74-77   78-80	
F10.3 E10.3 F5.3 F10.3 5A6, A5 I3 I4 I3	

\*In the far infrared ( $\nu < 100 \text{ cm}^{-1}$ ) a different format (F10.6) was occasionally chosen for the frequencies where high accuracy microwave measurements are available.

wavenumbers, S = line intensity in cm<sup>-1</sup>/molecule-cm<sup>-2</sup> at 296°K,  $\alpha = half$ -width in cm<sup>-1</sup> atm at 296°K, E'' = energy of the lower state expressed in wavenumbers. The rotation and vibration identification are the most difficult to unify due to the differences in the quantum numbers required to define the upper and lower states for different kinds of molecules. The columns 36 through 70 are indicated below for water and ozone, both triatomic asymmetric molecules, where (') indicates upper state and ('') indicates the lower state.

J', 
$$K_a', K_c'$$
J'',  $K_a'', K_c''$  $v'_1 v'_2 v'_3$  $v_1'' v_2'' v_3''$ 3I31X3I32X3I21X3I21X3I21X3I21X

On the other hand, the quantum numbers specified for  $CO_2$  and  $N_2O$  between columns 36 and 70 are given below, where again (') indicates upper state and ('') indicates lower state, and where r refers to Fermi resonance (see section on  $CO_2$  for more details). The identification system for methane differs from those described here and is described in the section on methane.

 $\frac{v'_{1}v'_{2}\ell'v'_{3}r'}{2x} \frac{v''_{1}v''_{2}\ell''v''_{3}r''}{5x} P(J''+1)$ 

The remaining fields specified above include the entry date of the datum (important primarily for our purposes), the isotopic code as described above, and the molecular identification as given in Table 3.

This work has now proceeded to the point where most of the data consistent with the above-mentioned line intensity limitation are fairly complete in the 1  $\mu$  m to 100  $\mu$ m region. The nature of the remaining uncertainties and omissions is discussed in the sections to follow. The data are frequency ordered on magnetic tape and are contained in records of 321 ten-character words per record. An initial control word indicates the number of words to follow (320 for a full record). Each such record contains 40 card images in the format described above. Thus, it is necessary after reading a record from the tape to decode it according to the format I10, 40 (F10.3, E10.3, F5.3, F10.3, 5A6, A5, I3, I4, I3). In order to aid those using other than Control Data Corporation equipment, Table 6 is provided. The column headed "External BCD Code" indicates the character representation actually used on the tape. If a computer having a different character code is being used, a simple cross-referencing program must be constructed. End of file markers are placed on the tape at the following frequency points: 500, 1000, 2000, 5000, 7500, 10, 000 cm<sup>-1</sup> with a double end of file appearing at the end of the tape.

In Appendix A we have included a listing of a computer program for directly reading the tape on a CDC 6600 computer. (No card decks are available.) For other computer systems the user is left to his own devices. Appendix B provides a computer program and output for generating a synthetic spectrum for a constant pressure path.

	Table 6.	CDC 600
	المتحديث ويتحدث ويتحد الجن محتد ويتحدث ويتحدث ويتحدث	
ource		Console
nguage		Display

Table 6. CDC 6000 Series Fortran Character Codes\*

٦

Source Language	Console Display	External BCD Code	Punch Position in a Hollerith Card Column
Character	Code		var u vorunni
A	01	61	12-1
В	02	62	12-2
C	03	63	12-3
D	04	64	12-4
E	05	65	12-5
F	06	66	12-6
G	07	57 70	12=(
H H	10	710	12-0
		A A	14-7
	12	44	11-1
K .	10	32	11-3
	16	75 44	11-4
IVI N	16	45	11-5
	17	48	11-6
E E	20	47	11-7
â	21	50	11-8
B		51	11-9
S	23	22	0-2
Ť	24	23	0-3
Î Î	25	24	0-4
V V	26	25	0-5
W	27	26	0-6
X	30	27	0-7
Ŷ	31	30	0-8
Z	32	31	0-9
0	33	12	0
1	34	01	
2	35		2
3	36	03	3
4	37	U4	4 E
5	40	UD Ac	C C
10	<b>A</b> 1	07	0 7
	42	1 10	Ŕ
0	70 AA		9
4	45	60	12
	46	40	1 $1$ $1$ $1$ $1$
*	47	54	11-8-4
	50	21	0-1
1 (	51	34	0-8-4
<b>j</b>	52	74	12-8-4
\$	53	53	11-8-3
	54	13	8-3
blank(space)	55	20	space
	56	33	0-8-3
A Contraction of the second	1 57	73	12-8-3

\*Taken from Control Data 6400/6500/6600 Fortran Reference Manual, Publication No. 60174900 Rev. C (1968) Control Data Corporation Interested parties can obtain a copy of the data tape described here by mailing a new, 7-track, 800 BPI certified, 2400 ft long by 1/2 in. wide, magnetic tape to R. A. McClatchey, AFCRL (OPI), L. G. Hanscom Field, Bedford, Ma. 01730.

Due to the large amount of material included on this tape and the likelihood of errors, it is clear that the ultimate test of the accuracy and completeness of the data will be its use by many people in the scientific community. Therefore, we ask the cooperation of all who use these data to keep us informed of any apparent errors or omissions. We would appreciate the receipt of new laboratory data or theoretical work related to improving the data. We will then update this tape as sufficient new or revised data become available.

#### **3. GENERAL REMARKS ON THE DERIVATION OF PARAMETERS**

The four tabulated parameters,  $\nu_0$ , E", S, and  $\alpha$  must of course be derived from experimental observations, subjected to data reduction in the framework of the general theories of molecular spectroscopy. For the basic theory we refer the reader to such textbooks as Herzberg (1950) and Goody (1964). The complexity needed to approach the problem depends both on the type of molecule and the accuracy of the observational data. We here outline the equations and methods used for the general types of molecules, linear triatomic (and diatomic) CO<sub>2</sub>, N<sub>2</sub>O and (CO); nonlinear triatomic, H<sub>2</sub>O and O<sub>3</sub>. Methane, CH<sub>4</sub>, a spherical top, is a special case, as is the diatomic O<sub>2</sub>, where unpaired electrons complicate the rotational structure of the ground and excited electronic states.

The energy states and the transition probabilities between energy states of the molecules are defined primarily by their numerical values as established by experiments and by the indices (quantum numbers) which identify them. In nearly all the cases of interest, mathematical relations of greater or lesser complexity relate the numerical properties to the quantum numbers. We here present the general relations used to generate the tabulated data. The specific data and exceptional cases are discussed later for each molecule.

#### 3.1 Energy Levels and Line Positions

#### 3.1.1 LINEAR MOLECULES

For the triatomic linear molecules  $CO_2$  and  $N_2O$ , the vibrational states are characterized by three quantum numbers, which are zero or positive integers, of pure vibration,  $v_1$ ,  $v_2$ , and  $v_3$  and a fourth number  $l_2 = v_2$ ,  $v_2 - 2$ ... which represents the contribution of the bending mode to the angular rotation. In order to calculate the purely vibrational part of the energy ("the band origins"), it is necessary first to compute an unperturbed energy,

$$G_{v}^{unp}, by; G_{v}^{unp} = \sum_{i} \omega_{i}^{vi} + \sum_{ij} X_{ij} v_{i} v_{j} + g_{22} l^{2} + \sum_{ijk} v_{i} v_{j} v_{k} + \sum_{ijk} v_{i} l^{2} + \dots$$

and then to incorporate the effects of resonance perturbation by combining all close-lying levels with common l and common symmetry in matrices whose diagonal elements are  $G_v^{unp}$ , whose off-diagonal elements are functions of additional molecular constants and the four quantum numbers, and whose eigenvalues are the vibrational energy  $G_v$ . For an excellent discussion and example, the reader is referred to the work on N<sub>o</sub>O by Pliva (1968).

(4)

This perturbation calculation results in the "mixing" of states whose  $G_v^{unp}$  are particularly close, so that the final description of the level by the original four quantum numbers is a poor one. Accordingly, it is useful to add a fifth index, the rank symbol r to label in order of decreasing energy all such mixed states. We have adopted this procedure for CO<sub>2</sub>, where the mixed states are  $(v_1 v_2 l_2 v_3)$ ,  $(v_1 + 1, v_2 - 2, l_2, v_3), \ldots$  etc. The highest value of  $v_1$  and the lowest value of  $v_2$  in each set are retained in the vibrational identification for all levels of the set.

In the present compilation, the above method was used to generate those energy levels which have not been observed; for all observed states the experimental value (averaged from various sources) was used.

The rotational energy of each vibrational state is given by:

$$E_{v,j} = G_{v} + B_{v} \left[ J(J+1) - l^{2} \right] = D_{v} \left[ J(J+1) - l^{2} \right]^{2} + H_{v} \left[ J(J+1) - l^{2} \right]^{3} + \dots, \quad (5)$$

where the constants  $B_v$ ,  $D_v$ ,  $H_v$  for each vibrational state are either determined by observation or calculated from a smaller number of rotational molecular constants. Whenever accurate constants have been observed, these are used; calculated values are reserved for the less important states. The equations for calculations of  $B_v$ and  $D_v$  are similar to those for  $G_v$ , and likewise require modification through the resonance perturbation. Evaluation from data of the small  $H_v$  constants requires highly accurate measurements extending to high J, so that in most cases these are fixed at zero; the non-zero values arise from resonances.

It will be noted in Eq. (4) that the vibrational energy depends on  $l^2$ . When  $l \neq 0$  there are two levels for each  $J \ge l$ , and this degeneracy is removed by rotation. The splitting ("*l*-type doubling") results in two sets of levels, designated <u>c</u> and <u>d</u>, with different effective rotational constants. When l = 1 the splitting is most important, and  $B_c \neq B_d$ ,  $D_c \neq D_d$ , etc.; when l = 2,  $B_c = B_d$  but  $D_c \neq D_d$ , etc.; when l = 3, the constants other than H are the same. Resonances occasionally cause larger deviations.

When the linear molecule has a center of symmetry, as in CO<sub>2</sub> with <sup>16</sup>O at both ends (but not when one oxygen is isotopically different), the paired atoms with zero nuclear spin cause zero statistical weight for rotational levels of a given parity. Thus, only even-J levels exist for the ground vibrational level and for all other levels with  $\ell = 0$  and  $v_3$  even (" $\mathfrak{L}_g^+$  symmetry"); for levels with  $\ell = 0$  and  $v_3$ odd ( $\mathfrak{L}_u^-$ ), only J odd exists; when  $\ell > 0$ , the c- and d- sublevels have different symmetry, so that for  $\ell = 1(\pi v)$  the J = odd levels are c and the J = even levels are d, etc. In the compilation symbols c or d are appended to the rotational quantum number of the lower state only when required, that is for  $\ell \ge 1$  in the molecules without the center of symmetry. For example, R27C means 28c - 27c; Q27C means 27d - 27c.

The line frequencies are determined from the energy states by taking the differences corresponding to all allowed transitions. These depend on the familiar selection rules for the linear molecule;

When  $\Delta l = 0, \Delta J = \pm 1, c \leftrightarrow c, d \leftrightarrow d$ . When  $\Delta l = 1, \Delta J = \pm 1, c \leftrightarrow c, d \leftrightarrow d$ , and  $\Delta J = 0, c \leftrightarrow d$ .

The line positions may thus be conveniently represented for computational purposes as given series in m, where m = J'' + 1 for the R-branch  $(J'' \rightarrow J''+1)$ , m = -J'' for the P-branch  $(J'' \rightarrow J''-1)$ , and m = J'' for the Q-branch  $(J'' \rightarrow J'')$ . A different equation is needed for Q-branch of a given transition than for the P and R branches, because of the differences in the <u>c</u> and <u>d</u> constants. The general equation is

$$\nu (m) = G_{v} + am + bm^{2} + cm^{3} + dm^{4} + em^{5} + fm^{6}, \text{ with } a = (B_{v}' + B_{v}'');$$
  

$$b = (B_{v}' - B_{v}'' - D_{v}' + D_{v}''); c = -2 (D_{v}' + D_{v}''); d = - (D_{v}' - D_{v}'');$$
  

$$e = 3(H_{v}' + H_{v}''); f = (H_{v}' - H_{v}'').$$
(6)

### 3.1.2 NONLINEAR MOLECULES

The nonlinear triatomic molecules  $H_2O$  and  $O_3$  have similar basic structures. The formula for the vibrational energy is identical with Eq. (4), except that the quantum number l and its associated constants do not exist. Vibrational resonances exist; in both  $H_2O$  and  $O_3$ ,  $\omega_1 \sim \omega_3$ , but these are of different symmetry, so that interaction between the band origins and identical rotational states occurs only in (200,002), etc. In addition for  $H_2O$ , and much more closely for the 162 isotope,  $2\omega_2 \approx \omega_1$ , so that the properties of the higher vibrational levels must be computed by taking their resonances into account.

The rotational levels of these three-dimensional rotators, with three different reciprocal moments of inertia  $A_v > B_v > C_v$  are labelled by three quantum numbers, J,  $K_a$ ,  $K_c$ , with  $K_a$  and  $K_c$  assuming all values 0, 1, . . . J, subject to  $K_a + K_c = J$  or J + 1. There are thus 2J + 1 levels of a given J (each again with a rotational statistical weight of 2J + 1); in addition there is a nuclear-spin statistical weight which gives alternate levels, depending on the odd or even parity of  $K_a + K_c + v_3$ , weights of 3 or 1 for H<sub>2</sub>O, 0 or 1 for isotopically symmetrical O<sub>3</sub>. The normal progressions of energy within each J is increasing with increasing K -K; occasionally inversions of a few paired levels may occur in excited vibrational states as a result of rotation-vibration interactions between close-lying states of like over-all symmetry in different vibrational levels. As J increases, there is a tendency for the odd and even levels with  $K_a$  or  $K_c$  close in value to J to approach very closely in energy, so that many apparently single lines are in reality degenerate pairs with unresolvable spacings. The range of  $K_a$ ,  $K_c$  in which these pairings occur, and the general spacing of the rotational levels is governed by the asymmetry parameter,  $\kappa =$ (2B-A-C)/(A-C), which for the ground vibrational states of H<sub>2</sub>O, HDO, O<sub>2</sub> is respectively -0.437, -0.689, and -0.984. As the parameter approaches -1, the energy level formula approaches that of the symmetric top with A > B = C namely,

$$E_{r} = \frac{B+C}{2}J(J+1) + \left(A - \frac{B+C}{2}\right)K_{a}^{2} - D_{J}J^{2}(J+1)^{2} - D_{JK}J(J+1)K^{2} - D_{K}K^{4}$$
(7)

where the D's are centrifugal stretching constants, and where additional terms of powers higher in  $J^2$ ,  $K^2$  are often required. For the asymmetric top molecules with  $K \neq -1$ , the rotational energy cannot be given by a closed formula, but requires the construction and diagonalization of matrices whose diagonal elements are given by Eq. (7) whose off-diagonal elements involve B-C and two additional centrifugal stretching constants of power 4, etc., and whose eigenvalues are the rotational energy. From the eigenvectors of the diagonalization, one obtains effective angular moments about each axis,  $\langle P_a^2 \rangle$ ,  $\langle P_b^2 \rangle$ ,  $\langle P_c^2 \rangle$ , whose sum,  $P^2 = J(J+1)$  and one also obtains other coefficients necessary for the calculation of transition probabilities and other properties of each eigenstate.

The excited vibrational levels of water vapor may differ considerably from the ground state in their values of A, B, and C, and with a relatively high ratio of  $A/\omega_2$  ( $\simeq 0.018$ , as contrasted to 0.00059 for  $B/\omega_2$  in CO<sub>2</sub>), extensive overlapping occurs between rotational levels in nearby vibrational states, making the computation of the higher levels highly uncertain much beyond the limits of observation. Accordingly, for all except the lowest energy levels of the ground state it is preferable to use experimental rather than calculated values when available.

#### 3.2 Line Intensities

• 2

The intensity of any line at frequency  $\nu$  may be expressed in a purely formal way as:

$$S_{m} = \frac{\nu}{\nu_{o}} \cdot S_{v}^{o} \cdot S_{Rot} \cdot F$$
(8)

where  $S_v^o$  is the vibrational intensity of a nonrotating molecule at the vibrational origin,  $\nu_o$ ;  $S_{Rot}$  is the rotational intensity for a rigid nonvibrating molecule and F is a factor that takes into account the fact that both forms of motion are occurring simultaneously. In the rigid case, F = 1, and if  $S_{Rot}$  is normalized so that  $\Sigma_{Rot} S_{Rot} = 1$  and if the band extends over a limited frequency range so that  $\nu/\nu_o \sim 1$ , then  $S_v^o$  as defined by Eq. (8) is identical with the quantity usually denoted by  $S_v$ , the total band intensity,  $\Sigma_m S_m$ .

 $S_{Rot}$  consists of a temperature-independent factor, the rotational line strength,  $L_{R}$ , common to all linear molecules, and a temperature-dependent Boltzmann factor,

If the sum of all transitions from J" is normalized to the statistical weight g = 2J''+1, and if one recalls the definition of the rotational partition function  $Q_r(T) = \Sigma g \exp(-1.439E_R/T)$  (the summation is over all rotational levels of a given v), the normalization condition for all transitions ( $\Sigma S_{Rot} = 1$ ) is fulfilled.

The equations for  $L_{R}$  are as follows:

ΔL	ΔJ	L <sub>R</sub>		
0	0	$l^2(2m + 1)/m(m + 1)$		(9)
0	<u>+1</u>	$( m ^2 - l^2)/ m $		(10)
<u>+1</u>	0	( m +1+l) (m+l) (2m+1)/l	m(m+1)	(11)
+1	<u>+1</u>	( m  + 1 + l) ( m  + l)/ m	and and a second se	(12)

Equation (8) is exact with F = 1 only for a rigid molecule. In the actual molecule, a number of factors may cause deviations, which however will vary regularly along the band. (A very few exceptional situations, "crossing perturbations", may also be recognized.) Depending on the precision of the measurements, the nonrigidity corrections ("F-factors") for the linear molecule, may be expressed as regular functions of the running index, m, as indicated in Eq. (13).

$$F = S_{\text{nonrigid}} / S_{\text{rigid}} = (1 + am + bm^2 \dots).$$
(13)

The first order theoretical correction is  $F = (1 + \zeta m)^2$  or  $a = 2\zeta$ ,  $b = \zeta$ . For nearly all the intensities in the present compilation, a single  $\zeta$  was used for each vibrational band; this is included in the tabulations of band parameters. More

refined measurements and theoretical calculations should, in future revisions of this compilation, permit the use of additional terms in Eq. (13).

For asymmetric top molecules, many more transitions are possible from each rotational level J  $K_a K_c$ , but the same general definitions hold. The values of  $L_{Rot}$  depend on the degree of asymmetry. Tabulations exist (Wacker, 1964) which are useful for orientation purposes. In the present work we either derive  $L_R$  from the rotational constants, for pure-rotation bands and some fundamental bands, or make use of the tables.

The F-factor corrections to the intensities become of considerable significance for most of the vibration-rotation transitions in  $H_2O$ . Various methods are used, based in part on theoretical considerations described later on, and in part on an arbitrary empirical adjustment of calculated values to the best observations.

A few general remarks should be made regarding the temperature dependence of  $S_v^{o}$ .  $S_v^{o}$  is proportional to the product of the relative population of the lower state, given by the vibrational Boltzmann terms exp (-1.439  $G_v$ )/ $Q_v$ , (with  $Q_v \equiv$  $\frac{L}{V}$  g<sub>v</sub> exp (-1.439G<sub>v</sub>) where g<sub>v</sub> is the degeneracy of the level, 1 when l = 0, 2 otherwise), and a temperature independent transition probability,  $u_{yy} = |\int \widetilde{\psi}_{y} u \psi_{y} d\tau|^{2}, \psi$ being the dipole moment function, usually expressed as a Taylor's series expansion in the dimensionless normal coordinates. We rely on measurement to give the total absorption strength in a given spectral region. When the spectral resolution is sufficiently high to distinguish lines of the strongest band in the region, usually the one with the lower level v = 0, from lines of all of the weaker "hot" or isotopic bands which accompany it, no problems arise: The  $S_{tr}^{O}$ of each band, and accordingly the relative  $\mu_{vort}$  is empirically established (along with favorable cases a determination of coefficients of the F-factor, leading to  $\Sigma S_{tr}$ for the region). However, most of the studies of quantitative band intensifies have been made under conditions where only the total intensity is observed and the strengths of the weaker lines, which contribute only a few percent of the total, estimated from theoretical relations.

When all the transitions in the region are of the same type (that is, have identical  $\Delta v_1$ ,  $\Delta v_2$ ,  $\Delta v_3$ ,  $\Delta \ell$ ), a good approximation to the relative transition probability is to use harmonic oscillator wave functions and to assume that the leading term in the expansion of  $\mu$  is formed by the  $\Delta v$ 's: for example if  $\Delta v_1 = 2$ ,  $\Delta v_3 = 1$ , we require that this term be  $\mu_{113} q_1^2 q_3$  with  $\mu_{113} = \partial^3 \mu / \partial q_1^2 \partial q_3$ . Evaluation of the integral for arbitrary values of  $v_1$ ,  $v_2$ ,  $v_3 \rightarrow v_1 + 2$ ,  $v_2$ ,  $v_3 + 1$  then gives the ratio  $\mu_{vv'}^2 / \mu_{o}^2 201 = (v_1 + 2)! (v_3 + 1)! / v_1! v_3!$ . The general formula (for  $\Delta \ell = 0$ ) is  $(v_1 + \Delta v)! (v_2 + \Delta v)! (v_3 + \Delta v)! / v_1! v_2! v_3!$ . When  $\Delta \ell = \pm 1$  similar relations involving integers hold. Use of more realistic wave functions corrected for an-

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harmonicity with lower terms in the dipole expansion lead to very nearly the same ratios. Accordingly, for many transition regions this type of calculation is simple and acceptable.

When, however, as in the case of both  $CO_2$  and  $N_2O$ , the resonance perturbations mix the vibrational wavefunctions, a region includes several bands with different basic transitions. For example, the  $1.6\mu$  region of CO<sub>2</sub> contains four strong  $\Delta l = 0$  bands, composing the resonance quartet 3001r-0, with r=1, 2, 3, 4. If we assume that only the  $\Delta v_1 = 3$ ,  $\Delta v_3 = 1$  transition is involved, the relative transition probability of the four transitions should be given by the squared eigenvectors of the {301, 221, 141, 061} matrix, and the relative transition probability of the four hot transitions from 010 would be the same value (since  $\Delta v_2 = 0$ ) multiplied by the squared eigenvectors of the {311, 231, 151, 071} matrix. These are not identical with the other eigenvectors, and all four hot bands have different transition probabilities. Moreover, the observed ground-state intensities are not proportional to the first set of eigenvectors. Inclusion of a second basic transition integral for  $\Delta v_1 = 2$ ,  $\Delta v_2 = 2$ ,  $\Delta v_3 = 1$ , together with the appropriate eigenvectors can, however, bring the observations in accord with calculation, if the ratio  $\mu_{221}/$  $\mu_{301}$  is of the order of 0.1. The ratio 231-0/221-010 is 3, and accordingly its inclusion increases the over-all -010/0 strength ratio and further shifts the relative probabilities. As one extends the calculations to lower states involving resonating groups with  $v_2 = 2$ , 3, and higher, required in the stronger bands of CO<sub>2</sub> and N<sub>2</sub>O, the situation cannot be predicted in advance. However, calculations similar to those sketched above have been performed; then reliability depends on the accuracy of the input constants, but it is believed that such a computation is useful.

One must also note that in the isotopically modified molecules, the eigenvectors of resonating groups vary widely. Thus, the simple assumption of a constant isotopic abundance ratio for the intensities is very far from correct. It should be adequate however to assume that the relative intensities of the underlying transitions (301-0 and 221-0, in the example cited above) remain isotopically invariant.

It should also be remarked that the eigenvectors are rotationally dependent; the result would be a contribution to the  $m^2$  term in Eq. (9).

#### 4. MOLECULAR SPECIES

#### 4.1 Water Vapor

4.1.1 LINE POSITIONS

The frequencies of the lines of the principal isotope  $({}^{1}H^{16}O^{1}H = Code 161)$  are calculated from a set of energy levels. These were obtained from the best available

data in all spectral regions by a smoothing process which is partly theoretical, partly empirical. The principal sources of data are identified and summarized in Table 7. The energy levels of the ground state,  $v_1v_2v_3 = 000$ , are by far the most extensive and accurate. They are based on a relatively small number (15, at the present writing) of microwave lines whose frequencies are precise to < 0.00001 cm<sup>-1</sup> (0.3 MHz), and several much larger groups of pure rotational lines, whose frequency accuracy may range from 0.001 to 0.002 cm<sup>-1</sup> for isolated lines measured with Michelson-type interferometers to 0.02 to 0.03 cm<sup>-1</sup> for weaker or partially blended lines measured with conventional spectrometers. With the long absorption paths available through the atmosphere, the observed lines extend to quite high energy levels, and by somewhat less accurate measurements with flame sources, to still higher levels, so that pure rotation transitions connect all levels from 0 to more than 4000 cm<sup>-1</sup>. The 000 level is the lower state of the more than 50 vibration-rotation bands observed between 900 and 20,000  $\rm cm^{-1}$ , so that these measurements provide combination differences (common upper level in two or more transitions) for the 000 state that confirm, and for some states greatly improve those of the pure-rotation lines. In particular, the recent laboratory measurements of the 1.9 µm region give low-J energy differences that are selfconsistent to better than 0.001 cm<sup>-1</sup>. A theoretical smoothing of the totality of the observed microwave, pure-rotation, and combination-difference data, by leastsquares fitting to a 29-constant Hamiltonian of the type described for D.O (Benedict et al, 1970) then yields the ground state energy levels used in the compilation. Inasmuch as none of the theoretical calculations have been successful in reproducing all of the available data to their apparent accuracy, the levels adopted for the current compilation are a calculated set for E < 2500 cm<sup>-1</sup>, and for higher energies are determined from observations. It is believed that the accuracy of lines involving these levels is  $\pm 0.005$  cm<sup>-1</sup> for E'' < 1500 cm<sup>-1</sup>;  $\pm 0.02$  cm<sup>-1</sup> for  $1500 \leq E'' \leq 3000$  cm<sup>-1</sup>, and  $\pm 0.05$  cm<sup>-1</sup> up to the tabulated limit. The levels of the other vibration-rotation states are then obtained by averaging the sums of the observed lines and the lower-state levels. When upper-state levels have not been observed, although transitions to them have expected intensities above the lower

limit, estimated values are chosen, either by direct calculations using an appropriate approximate Hamiltonian, or by extrapolation of the observed series of upper-lower rotational energy differences.

For the isotopic forms 181 and 171, the ground-state energies were obtained in the same way, except with much less extensive data (respectively 12 and 9 microwave lines). The dependence on theoretical calculations for the smoothing is more extreme, and the data for levels higher than 1500 cm<sup>-1</sup>, the limit of observation, is quite uncertain. However, since the higher-order constants in the

Dawlen		Range of	Leve	ls	Intensity	Precision		
region	Type of		Rota	tional	Limit	ofV		
cm	Measurement	Vibrational	្វ	Ka	(296 K)	cm <sup>-1</sup>	Ref.	
0-25	L, <u>181,171</u>	0,010	10	6	<-27	.00001	a,b	
0-25	L,162,182	0,010	13	7	<-27	.00001	c,đ	
30-250	<b>L</b>	0	13	7	-23	.005	e	
250-550	L	0	15	11	-24	.03	f	
480-690	LH	0	15	12	-25	.02	q ·	
430-650	F	0,010 etc.	30	15	<-27	.05	ĥ	
700-1100	LH	0,010	19	12	-26	.1	i	
750-1400	A	0,010	18	12	-25	.05	j	
860-1100	<b>A</b>	0,010	18	12	-25	.02	k	
1270-1450	LH	010,020	16	- 9	-27	.05	1	
1200-1700	L,162	010	14	7	-27	.02	m	
1330-1970	L, 181, 171	010	13	6	-26	.01	n	
1840-2500	LH	010,020,001+	.18	10	-26	.03	0,2	
1925-2182	A,U	010,020,001+	28	10	<-27	.01	a	
2390-2970	A,U	100,001, etc.	32	19	<-27	.01	â	
2480-3030	L,162	100,020	18	10	<-27	.005	a	
2900-3500	Α	020,100,001	16	11	-26	.02	r	
2800-3500	F	001,011, etc.	33	13	<-27	.02	S	
2900-4330	L,162	001,100, etc.	14	9	-25	.005	t *	
3340-4030	L,181,171	001,100, etc.	13	7	-26	.01	u	
3940-4300	F	001,011, etc.	33	19	<-27	.02	S	
4032-5090	A,U	001,011,020 +	33	17	<-27	.01	v	
3950-5200	A	001,030,011,etc	18	13	-26	.01	W	
4500-5915	L,162	011,030,110,etc	14	8	-25	.005	t *	
5090-5575	L	011,110	12	7	-24	.005	х	
5540-7000	A at	011,021,120 +	16	9	-26	.01	w	
5550-6720	A,U	011,021,120 +	24	11	<-27	.01	v	
7000-7500	L	101.200	13	8	-24	.05	v	
7400-9000	A	101,002,111 +	16	. 9	-26	.01	Ŵ	
7390-8800	A,U	101,001,111 +	16	9	-25	.01	v	
8300-10000	A	111.012.041	16	8	-25	.03	Z	
9150-9350	A	012,111	10	7	-27	.005		

Table 7. Summary of Principal Data Sources for Water-Vapor Energy Levels

#### References:

DeLucia et al, 1972 a. Steenbeckeliers et al, 1971 b. De Lucia et al, 1971 c. Benedict et al, 1973 đ. Hall and Dowling, 1967 e. Rao et al, 1962 f. Izatt et al, 1969 g. Madden and Benedict, 1956 h. Burch and Gryvnak, 1971 **i.** j. Dionne, 1972 Migeotte et al, 1957 k. Ben-Arych, 1967 1. Williamson et al, 1969 m. Gailar and Dickey, 1960 n. Burch and Gryvnak, 1973 ٥.

- p. Hall, 1972
- q. Benedict et al, 1973
- r. Beer, 1970
- s. Benedict and Sams, 1971;
- t. Pugh, 1972
- u. Fraley et al, 1969
- v. Hall, 1970
- w. Connes et al, 1969
- x. Flaud et al, 1972
- y. Nelson, 1951
- z. Swensson et al, 1970
- aa. Breckenridge and Hall, 1973

Table 7. References for Water Vapor Energy Level Data (Contd)

The data sources are of five types: <u>L</u>, laboratory absorption measurements on water vapor or moist air at room temperature; <u>LH</u>, similar measurements in cells heated to 75-540°C; <u>F</u>, laboratory measurements of emission from oxy-hydrogen or oxy-acetylene flames, yielding  $\sim 10^{19}$ molecules/cm<sup>2</sup> H<sub>2</sub>0 at +2500-3500K; <u>A</u>, measurements of solar radiation through the atmosphere, containing  $10^{22} - 10^{25}$  molecules/cm<sup>2</sup>; <u>U</u>, observations of the ratioed spectrum of sunspot/photosphere, also containing about  $10^{19}$ mol/cm<sup>2</sup> at ~3600K. Isotopic symbols are appended when enriched samples were studied, underlined when a major component, dashed when moderately increased above natural abundance. The range of vibrational and rotational levels is a rough indication of the extent of levels observable down to the intensity limit of S<sup>0</sup> (cm<sup>-1</sup>/mol cm<sup>-2</sup>), corrected to 296K, with the resolving power used. The ground state, (v=0) is implied in all regions.

\* In reference t, the observed contamination of the sample by deuterium to give the HDO abundances varying from 10-200 times normal was not reported. theoretical fit were constrained near their values in  $H_2O-161$ , the tabulated line positions for all pure-rotation lines should be accurate to  $\pm 0.5$  cm<sup>-1</sup>. Upper-state levels for these molecules are likewise available only for the strongest lines, so that the uncertainties of weak lines here might approach  $\pm 1$  cm<sup>-1</sup>.

The asymmetrically substituted HOD (162) molecule has been studied, both in the microwave and infrared regions with thoroughness comparable to 161, so that the frequency data, for the 000,010, 100, 020, 110, and 030 bands should be of the same accuracy as for 161. The 001 and 011 levels may have errors larger by a factor of five.

Quite reliable data can be calculated for the pure-rotation spectra of isotopic forms 182, 172, and 282, which are only of very minimal importance in the atmosphere. Weak lines in vibration-rotation bands of 182 (as yet unobserved, except for a few lines in 100-000) also have been included by estimating constant vibrational shifts from 162.

#### 4.1.2 LINE INTENSITIES

The intensities of pure rotation lines were calculated by a program in which the effects of vibration-rotation interaction on both the rotational eigenfunctions and on the dipole moment were included. The eigenfunctions were obtained from a Hamiltonian with 25 independent constants which gave an excellent fit to the energy levels, and the dipole moment was expressed as a linear expansion in the dimensionless normal coordinates:

$$\mu = \mu_{a} + (\partial \mu / \partial q_{1})q_{1} + (\partial \mu / \partial q_{2})q_{2} + (\partial \mu / \partial q_{3})q_{3} + \dots$$
(14)

The  $\partial \mu / \partial q_i$  coefficients in this expression were obtained from the  $S_v^0$  values of the three fundamental bands by the equation  $S_{vi}^0 = 4.16 \times 10^{-19} \nu (\partial \mu / \partial q_i)^2$ . The algebraic signs of  $\partial \mu / \partial q_i$  were chosen so that the observed vibration-rotation effects on intensities were reproduced in the fundamental bands of  $H_2O$  and verified by noting that these and only these choices also gave a reasonable fit to the band intensities and interaction effects in the isotopic molecules. The value of the permanent moment in the equilibrium configuration,  $\mu_e = 1.847D$  (Clough and Beers, 1973), was derived from Stark Effect measurements on 8 microwave lines in three isotopic water molecules (161, 162, 262), and is in excellent agreement with recent molecular-beam measurements (Dyke and Muenter, 1972). The uncertainty in the calculated intensities should result from inaccuracy of the  $\partial \mu / \partial q_i$  coefficients and neglect of higher terms in the expansion; accordingly, it should be of minor importance for low J lines, and is estimated to be less than 1 percent for E'' < 1000 cm<sup>-1</sup> and less than 10 percent for all lines in the tabulation, with the possible exception of the very low probability lines in such branches as  $R_{53}$ ,  $Q_{55}$ , etc. (Benedict, Classen and Shaw, 1952).

The intensities of vibration-rotation lines are more difficult to calculate. because they differ from those of the rigid asymmetric rotor by three interrelated effects: (1) the centrifugal effects, as used above for pure rotation; (2) the fact that the asymmetry, and hence the rotational eigenfunctions, are vibrationally dependent; and (3) the fact that for all vibrations above 010, the rotational levels of like symmetry and equal J in different vibrational levels are at energies sufficiently close that the vibrational eigenfunctions are rotationally dependent, and in special cases "perturbations" lead to abnormal mixing and intensities. Only the first-named effect has been incorporated in the computations, through the method described by Benedict and Calfee (1967). Table 8 lists the constants used for each band. The footnotes to Table 8 list the sub-branches which are observed to disagree in a regular way from the results of the calculation, and which were adjusted accordingly. The lines that are exceptionally perturbed (effect 3) are identified by the symbol P following K'; for these the total intensity of the two "interacting" lines was divided as required by the calculated vibrational mixing ratios or further adjusted to agree with measurements. No general statement can be made concerning the accuracy of the intensities of individual lines; for the low-J lines of the strongest bands it should be within  $\pm 10$  percent, within a factor of two for other lines of sufficient intensity (>  $10^{-25}$  cm<sup>-1</sup>/mol-cm<sup>-2</sup>) to appear in laboratory and/ or solar spectra, and within an order of magnitude for the weakest lines.

The intensities of the isotopic lines of species 181 and 171 were taken equal to those of 161 multiplied by the abundance factor (except for the cases of strong perturbation where the vibrational mixing ratios are different). The asymmetric isotopic species 162 is widely different, so that a completely independent calculation of its intensities, in both pure rotation and vibration-rotation has been made. The band parameters are included in Table 8.

#### 4.1.3 LINE WIDTHS

Recent high-resolution spectra of air-broadened or  $N_2$ -broadened water vapor lines (Brault, 1972, private communication) have in general confirmed the calculations of Benedict and Kaplan (1959) to  $\pm$  10 percent. A striking discrepancy is, however, observed (Blum et al, 1972) for the lines of highest J which are much narrower than the lower limit of 0.032 cm<sup>-1</sup> atm<sup>-1</sup> which was imposed on the calculated value by choosing a minimum collision diameter equal to the kinetictheory diameter. If the Anderson Theory (Anderson, 1949) is modified by eliminating a distance of closest approach (that is, setting the kinetic theory collision diameter equal to zero) the calculated half-width at half-height of 16<sub>1,16</sub> -15<sub>0,15</sub> is lowered from 0.032 to 0.0098 cm<sup>-1</sup>, still slightly above the observed value (0.0086). The validity of the theory is thus in question for high-J transitions but a revised

÷ .	vo			DAA	5 <mark>0</mark> (296K)	R	Int	eraction	Coeffi	cients
	1	iso	v* v*	Туре	cm <sup>-1</sup> /molcm <sup>-2</sup>	Debyes	α	β	γ	Notes
	1403.489	162	010 000	A	8.0 -022	.066	.0		0	
	1556 005	101		B	1.92 -021	.105	, <b>Ç</b>	'	.0	
	1500.895	101	020 010	B	8.61 -021	.171		1 - A 1 - A		(1)
	1501.279	191	010 000	B	2.12 -020	.121				(1)
	1.774.926	1/1	010 000	B	3.93 -021	.121				(1)
	1594.736	161	010 000	B	1.061-017	.121	.025	.0065	004	(2)
	2062.318	161	100 010	В	8.9 -023	.0167		••••	••••	•
	2161.188	161	001 010	A	4.9 -022	.0394		0108	015	
						n an				
	2709.35	182	100 000	X	1.31 -024	.0437				
	2723.687	162	100 000	A.	6.53 -022	.0437	.0322	.0130	.0077	
n in the Anna Anna				B	2.0 -023	.0076	-1	-,0156	.0	
	2782.014	162	020 000	A	8.1 -023	.0153	.0	0	.0	
			Richard († 1947) Geografie († 1947)	P	9.0 -024	.0051	-,01	-,00150	.0	
	3072.058	161	010 010	ъ	7 00 -027					
	3139.02	181	020 000		1 33 -023	.0121			·	(1)
	3144.96	171	020 000		A 93 -022	.0070		-		(1)
 	3151.631	161	020 000		A.94 -023	.0070				(1)
	3640.245	161	110 010		1 50 -020	.0070		.005	.003	(3)
	3649.690	181	100 000		7 24 -022	.0149				(1)
-	3653.14	171	100 000	P	1 22 -022	.0149			• .	(1)
-	3657.054	161	100 00	R	3 62 -010	.0149	075			(1)
	3808 455				3.02 -029	.0149	.075	.035	.028	(3)
	3707.459	162	001 000	B	3.0 -022	.0255	.C2	00047		
	3736.509	161	011 010	A	3.30 -021	.0708				(1)
÷	3741.571	181	001 000	A	1.60 -020	.0708	· .			(1)
	3748.36	171	001 000	λ.	2.96 -021	.0708				(1)
5 S.	3755.924	161	001 000	A	7.994-018	.0708	.0695	.0310	.0160	(4)
÷. *	4000 054	160	110 000	A	2.0 -023	.0088			.0004	
	4033.334	105	110 000	B	1.5 -024	.0024	(F=	1+0.15Ka	)	.(5)
· · ·	4145.483	162	030 000	A B	2.0 -023 1.5 -024	0	(F=	1-0.15Ka	05m)	(5)
·	4666.720	161	030 000	B	2.0 -022	.00032	.01	.10	.12	
								·		
	5089.539	162	011 000	A	3.0 -023	.0069	02			
n Shara	5180 26	161	120.000	B	b.0 -024	.0031	4			ана алана 1910 г. – С
	5221 20	101	T50 010	8	1.47 -023	.0040				. (1)
ана салана. Стала с	5227 75	101	110 000	: <b>8</b> .	3.66 -023	.0028				(1)
	5234 001	141	110 000	B	D.// -024	.0028				(1)
	5276 774	121	001 010	8	1.83 -020	.0028	.05	015 -	.02	*
	5310.43	101	011 000	а <b>А</b>	7.45 -022	.0286				. (1)
	5320-25	171	013 000	<u>A</u>	1.81 -021	.0203				(1)
		4/4	011 000	A	3.36 -022	.0203				(1)

Table 8. Water Vapor Band Intensity Data and Interaction Coefficients

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		1.1.1				1.	and the second second
		in a sta			and the second		1 A. 19
			e e inte				1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 -
			and the second second	and the second second			
	1. A. 1.	14 C 1			and the second second		
1.14	· · · · · · · · · · · · · · · · · · ·	1 N N 24		8	e studie		and the second of

	ν <sub>0</sub>				S <mark>V</mark> (296K)	9	Int	eractio	n Coeff	lcients
	<b>cm</b> -1	iso	vî çî	Rot Type	cm <sup>-1</sup> /mol cm <sup>-2</sup>	Debyes	α	ß	γ	Notes
	5372.114	162	200 000	A	1.2 -023	.0042				
· ·	6679.21	161	130 010	B	4.32 -024	.00194				
	6755.40	181	120 000	В	7:05 -024	.00112				
	6775.10	161	120 000	В	3.53 -021	.00112				
	6779.08	161	031 010	Å	6.94 -023	.0076		la se pr		
	6844.59	161	021 000	×	1.13 -622	0044				
	6857,32	171	021 000	X	2.09 -023	.0044		· · · · ·		
	6871.512	161	021 000	Å	5.64 -020	.0044			•	
	7186.68	101	200 000	B	1.06 -022	.0042				
e te lege	7201.48	161	200 000	B	5.29 -020	.0042				
	7213.26	161	111 010	X	3.98 -022	.0157				
	7222.68	181	101 000	X	1,49 -021	.0157				
	7235.57	171	101 000	×.	2.76 -022	.0157				Karan Ingelander Maria Maria Maria
-	7249.93	161	101 000	Ŕ	7.47 -019	.0157				e da la cara
	7371.79	161	ó12 <b>010</b>	B	2.17 -024	.00131				
	7417.54	181	002 000	B	1.06 -023	.00131			in a starter	
	7430.54	171	002 000	B	1.96 -024	.00131				
с. с. н . с. н	7445.04	161	002 000	B	5.29 -021	.00131		e stra La tracta	$(s, \xi'_1, \beta_2)$	
								an an a' An an a'		
an an an an a' an Anns an Anns	8238.84	161	041 010	A	5.88 -024	.00204				
	8273.95	161	130 000	R	2.4 -022	.00027	н. 1			
a dia seriesa di series Seriesa di seriesa di se	8341.32	181	<b>631 000</b>	A	7.2 -024	.00102				
•	8356.70	171	031 000	Å	1.33 -024	.00102				
	8373.82	161	031 000	×	3.6 -021	.00102		tin e		
	8734.97	161	121 010	Â	4.10 -023	.0037				en Terrapisco de Secondo
	8761.57	161	210 000	B	3.6 -022	.00031		tan Albanje		
•	8779.75	181	111 000	Å.	9.96 -023	.0037		e de la composición Referencias		
	8792.63	171	111 000	Å	1.85 -023	.0037				
	8807.00	161	111 000	Á	4.98 -020	.0037				
	8966.53	181	012 000	Ď	2.4 -024	.00057				
	9000.13	161	012 000	B	1.2 -021	.00057				
								ent e j	· · · ·	
	0011 59	161	000 000	1 1	A 6 _017	00011				

Table 8. Water Vapor Band Intensity Data and Interaction Coefficients (Contd)

#### Notes to Table 8.

(1) Insufficient data to establish interaction coefficients, these are assumed identical with corresponding band of 161.

(2) Lower line for strong lines (L rigid>1); upper line for others.

- (3) Many lines, especially those with engranced resonance with 001, require special treatment.
- (4) Lower line for strong lines (L rigid>0.5); upper line for others.
- (5) Very close resonance of these two states at  $K_{a}=0$ , all transition municipal affermation to 170-0.

calculation with  $b_{min} = 0$  does reproduce the empirical results better than the original calculation. Accordingly this has been incorporated in the current tabulation. As in previous computations, no vibrational dependence of the width has been included. The widths of the isotopic lines (181 and 171) have been set equal to those of 161; for 162 a new calculation, with  $\mu_e = 1.847$ ,  $q_{air} = 2.62 \times 10^{-26}$  esu and  $b_{min} = 0$  has been made, where  $\mu_e$  is the dipole moment of water,  $q_{air}$  is the effective quadrupole moment of the colliding gas and  $b_{min}$  is the collision diameter.

#### 4.2 Carbon Dioxide

#### 4.2.1 LINE POSITIONS

The line positions and lower-state energies of each of the significant isotopic modifications of  $CO_2$  were calculated for each band by a rotational formula for the linear molecule including terms up to  $D_v J^2 (J + 1)^2$ . For some of the higher states involving Fermi resonances of high rank, an additional term  $H_v J^3 (J + 1)^3$  was required. A very few of the vibrational levels are involved in a rotational perturbation, for which special calculations were needed as discussed below. The constants for each level are summarized in Table 9. For each isotopic species the band constants form a self-consistent set; that is, if a vibrational state appears more than once, as either initial or final state, its value of  $G_v$ ,  $B_v$ ,  $D_v$  (and occasionally  $H_v$ ) is the same. This requirement results in some deviations of the smoothed final line positions from their best observed values, but in general the positions of observed lines, up to J = 40, will be within + 0.01 cm<sup>-1</sup> of the tabulation.

The constants were calculated for each isotopic species separately, using a set of constants similar to those of Chedin and Cihla (1972), for 626, 636, and 628, but the isotopic sets were not consistently readjusted to fit a potential function. However, for isotopic forms for which the data are less extensive, approximate isotopic relations were used.

The data used for the band constants are taken from various sources. The highest precision data are those for the laser transitions 00011-10001 and 00011-10002 in the 626 isotope. The measurements of Gordon and McCubbin (1965, 1966), Oberly et al, (1968), and Drayson (1967) provide the data for interrelations among the lowest states. The most extensive data for the higher states are those from the planet Venus, as observed and summarized by Connes et al, (1969).

The vibrational notation may not be familiar to all users, and accordingly will be explained at this point. Five digits are used for each state, in order,  $v_1v_2 l v_3 r$ . The fifth digit, r, takes the possible values 1, 2, ...  $v_1 + 1$ , and serves to locate the level in the Fermi resonating group of v + 1 levels;  $v_1, v_2^l$ ,  $v_3$ ;  $v_1 - 1(v_2 + 2)^l$ ,  $v_3$ ; ... 0,  $(v_2 + 2v_1)^l$ ,  $v_3$ . The value of r decreases with increasing energy. (In .

Table 9. Vibrational Energy Levels and Molecular Constants for  $\rm CO_2$ 

Energy	G,	B	D	Snergy	Ģ	8	D	Energy	G	8	Ð
Level			Multiply	Level		1997 - C.	Multiply	Lovel	a de la composition d		Multiply -7
	_		by 10				by 10	1.1.1			by 10 '
the second	(cm <sup>-1</sup> )	(cm; )	(cm <sup>-1</sup> )		(cm <sup>-1</sup> )	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )		(cm)	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )
•					<del>2960-171-171-171</del>	tar gerietetete Tartetetet					
00001	100	tope 626		40004	5197.249	0.390092	1.89	41114 c	8081.179	0.387643	1.56
00001	0.00	0.340210	1.331	32203 4	5445.80 <b>3</b>	. 221920	1.535	41113 6	8250.644	.386233	1.355
01101 c	667.379	. 390643	1.355	21113 c	5475.071	.388153	1.61*	41113 6	L	.387813	1.355
01101 d		-, 391 253	1.355	21113 d		. 389423	1.705	41112 6	8425.000	.386817	0.935
10002	1285 412	100401	1 563	40002	5475.565	.390093	0.095*	41112 0	8803 265	.308517	0.965
02201 c	1335.129	. 391682	1.309	05511 cd	5627.254	. 390913	1.44	11132 6		.382744	1.56
02201 d			1.379	21112 ¢	5632.760	. 387018	1,265	03331	d 8863.648	. 383420	1.40
10001	1388.187	. 390188	1.142	21112 d	F770 610	.386128	1.385	11131	8944.146	.381264	1.22
11102 0	1912 470	390736	1 441	2))))) cg	5730.618	389269	1.125	11131 6		. 3818/4	1.18
11102 d		. 391680	1.501	21111 4		. 388613	1.025	20033	9388.990	.38234	1.71
03301 cd	2003.238	. 392420	1,403	10022	5915.209	. 384528	1.57	12232 6	9419.189	.38307	1.37
11101 c	2076,865	. 390416	1.281	02221 c	5958.539	. 385617	1.35	12232	0516 070	20040	1.25
00011	2349.146	391344	1.195	10021	6016.690	383917	1.31	12231	9589.929	. 38247	1.43
		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,						12231	1		1.53
20003	2548.280	, 391183	1.76	30014	6075.983	. 388896	2.052	20031	9631.350	.38125	1.04
12202 C	2585.006	. 391843	1.45	22213 c	6103.670	. 389428	1.29		0007 40	10001	
12202 d	7671 113	100556	1.25	22213 G	6176 693	190485	1.54	21133	2 9987.48	. 38251	1.55
04401 cd	2671.690	. 393162	1.42	41102 c	6179.010	.389642	1.33	21132	10145.43	.38098	1.25
12201 c	2760.735	. 391 5 35	1.44	30013	6227.924	. 386697	1.643	21132	1	. 38258	1.35
12201 d			1.25	22212 c	6288.492	. 388472	1.38	21131	10297.05	.38127	1.10
20001	2797.154	. 390563	0.985	22212 0	6343 664	206461	1.28	21131	1 Tentono	- 38237	1.00
01111.0	2008-010	. 388190	1.349	41101 0	6388.085	. 390290	0.93	00000	0.00	. 390235	1.330
				14411 cd	6398.047	. 390017	1.37				
21103 с	3181.450	.391028	1.63	22211 c	6474.530	. 388662	1.53	01101	c 648.484	. 390598	1.330
21103 d		. 392316	1.75	22211 d	6843 001	207074	1.21	01101	đ	. 391236	1.330
13302 ca 21102 c	3290.009	. 392090	1.32	11122 0	6537.958	384504	1.475	10002	1265,820	. 390920	1.58
21102 d	93991390	.391145	1.37	11122 4		.385722	1.606	02202	1297.269	.391603	1.274
05501 cd	3340.475	. 393908	1.44	11121 c	6679.709	. 384310	1.216	02202	4		1.334
13301 cg	3442.256	.392312	1.35	11121 d	6070 870	- 385127	1.115	10001	1370.067	.389707	1.160
21101 C	1200.390	390401	3.01	00031	0772.319	. 300330	1,341	01102	1896.49	. 391137	1.48
10012	3612.844	387496	1.57	31114 c	6688.274	. 388547	1.735	01102	d .	.392096	1.13
02211 c	3659.277	. 388647	1.383	31114 d		.390246	1.825	03301	cd 1946.343	. 392293	1.34
02211 đ			1,373	31113 c	6863.553	, 386930	1.401	11101	c 2037.093	.390015	1.10
10011	3714.781	. 387051	1.11	11113 a	7023.677	306755	1.405	11101	2283.490	- 390975	1.13
30004	3792.702	. 391760	2.02	31112 4		.388127	1.165				
22203 c	3821.984	. 392359	1.46	20023	7133.819	. 385288	1.75	20003	2507.50	. 391815	1.76
22203 d	-	202424	1.68	12222 c	7166.049	.386017	1.15	12202	¢ 2531.63	. 39220	1.22
14402 cd	1042 402	193434	1.56	31111 0	7263 824	387607	1.05	04401	od 2595.614	. 392987	1.40
22202 c	4007.850	. 391485	1.47	31111 d	1,000,000	.389117	0.85	20002	2645.086	. 389826	1.46
22202 d			1.40	12221 c	7338.249	-385487	1.33	12201	c 2700,25	.39126	1.30
30002	4063.908	. 389606	0.935	12221 4		304403	1.28	12201	d 2760-40	100670	1.34
14401 22201 g	4122.347	. 191030 301710	1.30	01131 g	7602.529	381503	1.315	01111	c 2920.244	. 3050/0	1.325
22201 d			1.13	01131 d		. 382063	1.355	01111	d	.398286	1.325
30001	4225.110	. 391090	0.521		and a straight						1.5
11112 c	4247.713	.38777	1.49	40015	7283.981	. 389550	2.31	21103	c 3127.28	.39155	1.70
11112 0	4314 013	. 388690	1.36	37214 C 19314 A	7407.651	- 783347	1.03	1 3302	a cd 3169.21	. 39308	1.08
11111 c	4390.628	. 387350	) 1.24	40014	7460.530	.387322	1.93	21102	c 3289.71	.39001	1.25
11111 d		. 388230	1.17	32213 c	7505.219	. 388632	1.23	21102	đ	.39120	1.25
00021	4673.327	. 38406	1.319	32213 d			1.34	13301	cd 3361.57	.39188	1.25
21100	A416 160	101.76	1.41	40013	7593.690	. 385562	1.04	21101	C 1433.70	. 39292	1.04
31:104 d	MATD*7.30	. 393076	1.76	32212 4	19393340	L'HUWAGA	1.14	10012	3527.740	. 388030	1.56
31103 c	4591.118	. 18992	5 1.53	40013	7734.452	. 386954	0.59	02211	c 3557.314	. 388666	1.28
31103 d		. 39134	3 1.48	21123 c	7743.700	.305356	1.25	02211	d	20/20/	1.33
3.1102 C	4/53.450	301004	5 1.20	322123 0	7897.571	38968L . 38968L	1.33*	10011	1032.917	. 386/26	1.21
20013	4853.620	. 39818	1.77	32211 4		a la serie de l	0.939**	30001	4145.95	. 39010	1.22*
12212 c	4887.970	. 388951	9 1.33	21122 c	7901.479	. 384008	1.31*	11112	c 4147.234	.38811	1.49*
12212 4	4030 414	1007	1.43	21122 d	7020 040	.385128	1.38**	11112	d	.38915	1.59
31101 6	- 長いいり - 美10	. 109.151	0.995	21121 c	8056.024	. 384404	1.11	11111	c 4287.695	.387654	1.33
04411 cd	4970.909	. 19014	1.42	21121 d		. 38552	1.08	11111	d	. 387977	1.26
20012	4977.830	. 38652	1.35	10032	8192.556	. 38156)	1.56	00021	4543.552	.384360	1.33
12211 c	5061.776	. 38850	1.28	02231 0	8332.880	. 382600	1.30	20012	4740 050	70004	1 010
20011	5099.666	38744	0.921	10031	8293.957	.36080	1.13	12212	c 4770.985	. 38925	1.25
01121 c	5315.730	. 38454	3 1.46					12212	đ		1.27
01121 d		. 38514	9 1.46	.* .** J-c	ievendent i	<b>steractions</b>		04411	cd 4831.99	. 39003	1.34

Energy Level	G	3	D Multiply	Energy Level	G	8	0 Multiply	Energy Level	G	.,	D Aultiply
			by 10 <sup>-7</sup>				by 10 <sup>-7</sup>				by 10 <sup>-7</sup>
	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )		(cm <sup>-1</sup> )	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )		(ca <sup>-1</sup> )	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )
2001.2	1007 2%	0 196947	1.4	21102 d	1201 07	0 36016	1.14		4000.00	0.07/1/	
12211 C	4939.80	. 38831	1.44	13301 cd	3404.93	. 37077	1.28	11112 a	4223.33	. 37700	1.31
12211 d	1707.00		1.40	21101 C	3453.99	. 36881	0.96	11111 c	4367.09	.37600	1.18
20011	4991.350	. 386705	0.94	21101 d		. 37007	0.90	11111 d		. 37686	1.15
01121 c	5168.600	. 384695	1.33	10012	3571.143	. 365284	1.27	00021	4655.205	. 372674	1.20
01121 d		. 182102	1.33	02211 C	3632.52	. 366595	1.19	20013	4821.500	.376411	1.62
21113.0	5357 004	. 388614	1.50	10011	3675 130	. 365524	0.95	20012	4939.350	. 376321	1.21
21113 d	0007.001	. 390108	1.69		0070.100		0.75	21112 a	5593.645	. 375625	1.22
1,312 cd	5397.09	. 38978	1.30	30003	3056.65,	. 367405	1.24	21112 d		.376759	1.22
21112 C	5519.944	. 387038	1.16	30002	3987.610	. 368174	0.91	00031	6945.610	. 369691	1.20
21112 0	5500 17	. 388206	1.30	11112 C	4201.19	. 303000	1.23	00001		6 %	
21111 c	5662.269	. 386837	1.10	03311 cd	4293.35	. 36726	1 19	00001	0.00	. 300100	1.11
21111 d		. 387936	I. w	11111 c	4346.13	. 365716	1.05	01101 c	643.23	. 36857	1.12
				11111 d		. 366 568	1.00	<b>01101</b> d		. 36912	1.12
30014	5951.600	. 389640	2.326	00021	4639.502	. 362372	1.11				
22213 C	5970.949	. 389913	1.57	20013	4791.260	. 365734	1.50	10002	1244.93	. 36820	1.30
30013	6119 618	. 387540	1.33	12212 6	4936.63	.306/0/	1.25	02201 C	1200.00	. 10421	1 11
22212 c	6155.37	. 38852	1.36	20012	4904.850	. 364844	1.11	10001	1342.37	. 36844	0.90
22212 đ			1.30	12211 c	5012.55	. 366735	1.26	00011	2265.973	. 365388	1.11
30012	6241.964	. 385859	1.113	d			1.06	20002 c	2586.43	. 36778	1.04
22211	6326.049	.388103	1.04	20011	5042.570	.366124	0.86	01111 C	299, <b>58</b>	. 36579	1.12
30011	6363 616	387033	1.35	d d	52/7.147	. 302040	1.06	10012	2400.20	. 30033	1.14
11122 C	6374.49,	. 36521	1.46	21113 c	5406.069	. 365893	1.41	02211 C	3529.59	.36671	1.14
<b>11122</b> d		. 38607	1.56	đ		. 366904	1.31	02211 d			1.12
00031	6790.215	.301353	1.33	21112 c	5558.553	.365318	1.10	10011	3567.54	. 36525	1.01
	(552.054	300050	1 ( )		5707.04/	. 366240	1.10	00021	6S06.749	.36260	1.11
31114 C	0552.954	190896	1.03	A 11117	5727.046	. 303940	0.81	20012	4814.570	.364610	1.11
31113 c	6736.6%	. 387368	1.63	10022	5858.022	. 362490	3.357	00001	0.00	. 378656	1.21
31113 d		. 388906	1.58	02221 C	5915.23	. 36376	1.18	01101 c	645.72	. 379036	1.22
31112 c	6692.054	. 386378	1.33	đ			1.15	01101 4		. 379627	1.22
31112 u		. 387706	1.18	10021	5959.5%	. 362605	0.896	10002	1254.93	. 37900	1.40
31111 C	/046.029	. 180828	1.23	30014	5 <b>993.581</b>	. 366337	1.79	02201	1292.90	. 38000	1.22
01131 c	7191.599	. 381803	1 33	30012	6254.592	. 365274	0.80	10001	1355 52	17849	1.22
01131 <b>d</b>		. 382391	1.33	30011	6429.172	. 366616	0.60	00011	2274.33	. 37579	1.21
				00031	6922.210	. 359479	1.10	01111 c	2918.38	. 37617	1.22
40013	7481.510	. 385435	0.33					01111 d		.37675	1.22
40012	7600.130	. 385855	0.63	10032	6120.104	. 359684	1.25	10002	3509.07	. 37641	1.41
10032	7981.1~0	. 3822/3	1.52	10031	9220.363 Tenton	.337030 # 627	0.95	10001	3609.05	.37540 na 828	1.11
10031	Isoto	m 628	1.20	wool	0.00	.378641	1.20s	00001	0.00	. 346799	1.04
00001	0.00	.368164	1.11					01101 c	657.33	.347224	1.04
				01101 c	664.735	. 379065	1.22	01101 đ		. 347736	1.04
01101 c	662.368	. 368588	1.13	01101 d		. 379639	1.22	10002	1230.20	. 34620	
01101 a		. 194158	1.13	10002	1271 075	178718	1.40	02201 C	1312.08	.34812	1.10
10002	1259,430	. 368114	1.26	02201 c	1329.87	. 380061	1.24	10001	1347.22	. 34776	1.04
02201 C	1325.15	. 369520	1.20	02201 4			1.22	00011	5313.97	.34412	1.04
02201 d			1.16	10001	1376.28	. 378806	1.04	01111 c	2959.06	. 34455	1.04
lowl	1365.845	. 368502	0.962					01111 4		. 34504	1.04
	1001 749	36945		11102 C	1916.31	. 37903	1.27	10002	3525.205	. 343840	1.16
11102 C	1701./46	36925	1.26	03301 cd	1995 16	17977	1.32	10001	3639.067	.J440J0 .728	0.907
03301 cd	1988.328	.370182	1.20	11101 c	2062.41	. 37894	1.15	00001	0.00	.35695	1.07
11101 c	2049.346	. 36860 1	1.06	11101 đ		. 37987	1.10	01101 c	659.70	.35737	1.08
11101 d		. 36954	1.01	00011	2340.01	. 375659	1.20	01101 d		.35789	1.00
00011	2332.112	. 365267	1.11					00011	1322.52	.35419	1.07
20002	7600 776	360463	1 4 4	20003	2523.58	. 37924	1.60				
20003 12202 e	2549.425	. 36954	1.25	12202 4		. 300123	1.25				
12202 <b>d</b>			1.27	20002	2641.26	. 37814	1.18				
20002	2614.235	. 367807	1.04	12201 <b>C</b>	2743.66	. 38005	1.20				
04401 <b>cd</b>	2651.975	. 37085	1.22	12201 <b>d</b>			1.2.				
12201 C	2729.264	. 36966	1.30	20001	2776.00	.37927	0.88				
20001	2727 220	. 3690 19	0.77	01111 4	2442.310	. 376664	1.22				
01111 c	2982.106	. 365707	1.13	10012	3590.86	. 375835	1.38				
				02211 C	3645.02	. 377091	1.24				
21103 C	3127.31	. 36855	1.34	"2211 <b>d</b>			1.22				
21103 <b>d</b>		34 157	1.43	10011	3693.64	. 375754	1.06				
13302 cd	3200.15	.3 123	1.28								
ALLVA C	3291.07	· 1001/	****								

Table 9. Vibrational Energy Levels and Molecular Constants for  $CO_2$  (Contd)

this scheme,  $v_2$  is always equal to  $\mathbf{1}$ .) The desirability of labelling the states in Fermi resonance by the same symbols except for  $\mathbf{r}$  is to emphasize the very important role played by the Fermi resonance, which is very nearly exact for most of the groups. It must also be emphasized very strongly that the long-established custom of calling the upper level of the  $(10^{0}0, 02^{0}0)$  resonance at 1388.18 cm<sup>-1</sup>,  $v_1$ , and the lower level at 1285.41,  $2v_2$ , is not only confusing the issue, but is wrong, since the now ten year old work of Amat and Pindert (19651 has shown that the unperturbed position of  $2v_2^0$  lies above that of  $v_1^0$ , in the 626 isotope. This is also true in 627, 628, and 828, but in 636, 637, 638, it lies below. It seems vastly preferable always to label the upper level 10001 and the lower 10002.

#### 4.2.2 LINE INTENSITIES

The intensity data (summarized in Table **10**) are based on quantitative studies of resolved lines where such exist; and where not, on total intensities in a given region. The relations between the strengths of the principal band in a region and its associated weaker "hot" and isotopic neighbors were based on computations relating the transition moments to the vibrational eigenfunctions, taking into account terms up to the quartic in the molecular constants, with particular attention to the degree of Fermi mixing in the states, and an approximate calculation of the "unperturbed" (before Fermi mixing) transition moments. This procedure appears to give fairly good agreement for the ground-state and "first-hot" transitions in such Fermi groups as 2001-0000 and 3001-0000, and seems adequate for estimating strengths of higher unobserved hot bands. However, when experimental data are available (**Burch**, 19701, as for the 12201-11102 band at 828.28 cm<sup>-1</sup>, these were used.

For the isotopic molecules, empirical data were again used where available, as for 628 bands which are forbidden in the symmetric molecules. When the isotopic bands overlap stronger bands of 626, the abundance ratios were **used to** estimate the total strength of **a** group, but the particular Fermi parameters were used to divide the strengths.

The influences of vibration-rotation interactions on line intensities has been **measured** in some cases. As a generalization, it can be said that these are usually negligible for the parallel bands (l' - l'' = 0) but become of considerable importance for all the perpendicular bands (l' - l'' = 0) but become of considerable intensities for all the other bands, Coriolis-type resonances are responsible for inducing a large part of the transition moment in the R- and P-branches, thus "borrowing" intensity from the stronger parallel bands, while leaving the Q-branches unchanged. Accordingly, the latter may remain very weak (or in some cases be unobservable - for example, the 410 group). We have used the approximate equation  $S_m = S_w^0$  (1 t m  $\zeta_w$ )<sup>2</sup> to correct for this interaction, with the values

					-				
		•							•
Center	Level	Lover	'so-	& at 296K	Band	Upper	Lower	Iso-	SV at 296K
Yo			LOFE	-21 -1	Ve	TEAGY	TGAGT	cope	(multiply 4 +22 _1
~1				10 101	·0				10 <b>** mol</b>
(cn=`)				cas°cas°) ∱r	(cm <sup>-+</sup> )				cast cas <sup>−1</sup> ) 5*
					1				
471.415	20003	11101	626	0.0097	681.27	,220,	11101	627	.0082 Q
479.929	13302	12201	626	.0012	681.521	13301	,220,	626	.460
494.537	12202	11101	436	.00079 Q	693.501	11101	10001	626	.77
510.337	2,103	20002	626	.0040	663.870	11101	10001	626	9.05
526.423	1,102	10001	636	.035	669.679	11101	10001	626	148.8
535.903	11102	10001	628	.011 Q	696.623	2220,	21101	626	0.0195
542.1%	21102	20003	626	.00711	699.14	10001	01101	638	. 058
546.275	13302	04401	626	2.723	703.477	10001	01101		9.20
557 .442	14402	05501	626	.00253	703.436	20001	11101	626	0.246
561.097	12202	03301	628	.0058	709.80	10001	01101	637	.0094
564.999	20002	11101	628	.0043	710.765	10011	01111	626	.0202
570.97	13302	03301	626	.0/16	711.545	10001	01101	627	1.26
573.536	13302	04.0,	636	.00115	712.48/	20002	11102	62.	0.013
576.599	11102	0220,	629	.169	713.59	20001	11101	627	.0038
579.605	21102	,220,	626	.0376	720.299	20001	11101	626	4.794
579.95	20002	11101	627	.00072 Q	720.909	10001	01101	626	,953.
581.768	12202	03301	626	1.934	721.563	10001	01101	636	17.9
565.297	12202	03301	636	.034	724.190	20002	1 102	628	0024 0
596.44	11102	02201	627	.0439	732.54	11101	0220,	627	.0474
569.46	21102	12201	636	.00055 Q	733.45	12201	2000,	636	.0021 Q
595.65	20002	12202	636	.908	736.643	20002	11102	626	3.021
596.444	21103	12202	626	.257	739.624	220	0220,	636	0.70
597.062	10002	01101	628	5.19	739.855	21101	,220'	626	.176
597.34,	11102	0220,	626	52.09	741.736	11101	02201	626	79.01
599.028	20003	11102	628 6 7 6	0.0031	748.32	12201	03301	627	0.0025 0
601.70	10002	01101	638	.702	746.5%	20002	04401	636	.0378
607.14	10002	01101	627	1.17	753.055	21102	12202	626	1607
607.27	20003	11102	627	0.021	757.497	,220,	03301	626	3.296
607.993	20002	11101	636	.0111	765.896	13301	04401	634	0.00173 Q
608.828	10012	01111	626	.0175	770.355	,330'	04401	626	.1351
609.1,	20002	13102	637	102	781 872	1 40	05501	636	.136
615.810	20003	11102	626	6.99	789.916	11101	10002	626	.056
617.336	10002	01101	636	20.5	790.536	11101	10002	627	. <b>010</b> Q
616.033	10002	01101	626	1436.	791.060	21102	20003	626	.0548
619.78	21103	20003	636	0.0085	791.452	11101	11102	626	11.23
636 754	01111	20003	636	.0121	926.516	1220.	11102	639	.0009 0
637.56	13302	12202	636	.013	828.265	,220,	11102	624	.201
640.670	11102	10002	636	2.60	929.467	21101	20003	626	.0548
642.319	11102	10002	628	0.78	657.250	13301	12202	624	.0110
643.23	11102	10002	627	0.15	863.15	01111	11101	636	.00495
645.72	01101	00001	637	.642	996.542	02211	12201	626	.0132
646.11	0220,	01101	637	.055	913.423	00011	10001	636	.0614
647.056	11102	10002	626	222	915.584	21101	12202	626	.00195 0
645.484	01101	00001	636	96U. 72 4	917.627	01111	11101	626	.3556
649.074	03301	0220.	636	4.69	932.760	01111	11101		.00135
649.271	. 04401	03301	636	0.266	941.731	10012	20002	626	.01146
649.580	0 05501	04401	636	.01(2	952.310	21101	20003	626	.00064 Q
650.02	12202	11102	627	16.52	963.73	00011	10001	427	.0032
654.870	01111	00011	626	0.885	966.267	00011	10001	639	.0195
655.558	11302	12202	626	0.990	1017.670	00011	10002	636	.0726
657.31	01101	00001	820	. 343	1023.744	01111	11102	636	.00072
657.69	02202	13303	426	.02//	1068 135	00011	10002	627	0.324
659.79	01101	00001	728	.0635	1071.546	01111	11102		.540
<b>661.32</b>	1 3 3 0 1	12201	636	. 0055	1072.682	00011	10002	628	.0285
662.368	01101	00001	628	330.	1074.271	02211	12202	626	.0227
662.782	02201	01101	628	26.2	1060.128	01111	11102		.00243
663.17	01301	0220.	628	1.50	1239.380	11102	01101	629	0.035,
663.547	04401	03301	628	0.078	1244.93	10002	00001	638	.0038
664.735	6 01101	00001	627	62.2	1259.430	10002	00001	628	. 382
665.135	02201	01101	627	4.69	1271.875	10002	00001	627 636	.068
665.49	10110	10001	636	1.58	1365.645	10001	00001	629	. 561
567.379	01101	00001	626	82580.	1376.28	10001	00001	627	.0323
667.750	02201	01101	626	6488.	1386.978	11101	01101	629	.016
668.109	0330,	02201	626	392.4	19/6 321	21102	0220	626	.0003315
666.227	04401	03301	626 626	18.45	1959.02	20003	01101	636	.0001615
668.78	0550.	04401	626	0.976	1960.232	30004	11102	626	.00004 ~.12
649.1	06 06601	05501	626	.077	1965,607	30003	11101	626	.0000712
679.916	12201	11101	629	.060	1880.901	<b>∡000</b> 3	01101	626	.013010

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Table 10. Band Origins and Intensities for CO,

Band Center Vo	Upper Level	Lower Level	Iso- tope	s <sup>9</sup> at 296g (multiply by 10 <sup>-22</sup> -1	Band Center	Upper Level	Lower Level	Iso- tope	S <sup>9</sup> at 296K (multiply by
-1				cm <sup>2</sup> cm <sup>-1</sup> ) f	-9				10 <sup>-22</sup> mol <sup>-1</sup>
(cm <sup>-1</sup> )				· · · ·	(cm <sup>-1</sup> )				can" caa"≛) J
1001 146	12202	01101	676	0.00140	2200 442	11112	11102	620	0.686
1889.514	22203	11102	626	.0012708	2301.041	12211	12201	6%	3.17
1896.490	11102	00001	636	.014906	2301.73	01111	01101	929	0.31
1901.748	11102	10002	628	.0110107	2302.365	10021	10012	6%	0.0473
1905.435	13302	02201	626	.178606	2305.340	20013	20003	626	4.352
1917.627	12202	01101	626	.4226055	2306.717	20012	20002	626 627	2.3%
1932.470	11102	00001	624	4.092062	2307.37	02211	02201	629	10.5
1951.153	21102	10001	626	.0070706	2307.39	00021	100011	629	0.0791
2003.734	20002	01101	626	.001504	2311.675	03311	03301	626	122.9
2004.211	21102	02201	626	.0011804	2311.713	10012	10002	629	7.29
2010.022	11101	00001	6% 636	.0001305	2313.744	11111	11101	626	8.56
2049.346	11101	00001	628	.037203	2313.97	00011	00001	028	3.60
20 <b>65.846</b> 2062.41	11101	01101	628 627	.005203	2335.15 2315.243	11112	11102	627	1.91
2075.380	22202	11102	626	.008504	2317.36	10011	10001	627	0.805
2076.865	11101	00001	626	22.32041	2318.985	10012	01101	627 628	1.29
2094.861	20001	01101	628	0.005504	2322.52	00011	00001	729	1.35
2101.996	20001	01101	636	.014903	2324.146	02211	02201	626	3060.
2112.461	21101	10001	626	.112035	2326.594	10011	10001	624	1103.
2119.119	14401	03301	626	.0156033	2327.432	10002	10002	626	1934.
2120.548	22201	1220.	626 626	.011904	2327.575	01111	01101	627 626	49.7 3330
2129.775	20001	01101	626	1.302046	2336.637	01111	01101	626	76600.
2148.245	30001	11101	626	.0059510	2340.01	00011	00001	627	637.
2165.461	21101	02201	426	.059505	2367.097	10011	10002	6%	0.0131
2170.848	11112	11101	626	.0507	2415.700	10011	10002	628	.0056
2182.507	20012	20001	626	.00092	2429.347	10011	10002	6%	1.059
2205.298	10012	10001	628	.00558	2429.5%	20012	20003	'26	0.00256
2224.647	10012	10001	626 636	1.272	2464.942	21103	01101	626 6 %	.0402
2227.88	13312	13302	636	.00401	2 M o . 7 7	6 20003	00001	62.	.075
2229.724	21113	2'103	636	.00488	2523.56	20003	00001	627 6%	.0060
2236.376	04411	04401	636	.0632	2414.235	20002	00001	629	.186
2238.55	12211	1220,	436	.0384	2618.702	21102	01101	626	.0148
2240.558	20013	20003	636	. 129	2757.229	20002	00001	626	.022
2240.87	20011	20001	636	.0122	2776.00	20001	00001	627	.0023
2242.776	20012	20002	636 638	.0258	4/71.044	21101	01101	020	. 0015
2242.73	02211	02201	636	.149	3125.32,	30004	01101	626	.00024 .40
2245.17	10011	10001	639 638	-0555	3154.605	22203	01101	626 626	.00074 .40
2248.356	01121	01111	636	.0280	3275.113	30003	01101	626	.0102 .02
2248.361	03311	03301	636	1.52	3291.07	21102	00001	629 636	.00017 .14
2250.744	11112	11102	636	1.9'	3339.340	21102	00001	626	.0417 .14
2251.01	02211	02201	637	0.0264	3340.471	22202	01101	626	.0063 .14
2253.53	10011	10002	637	.0096	3399.2%	21113	11101	624	.00413
2254.35	01111	01101	638	3.u	3450.75	13311	03301	636	.0174
2260.045	02211	00011	636	34.6 0.306	3465.433	20013	10001	626	.1786
2261.920	10012	10002	636	20.3	3473.716	12212	02201	636	. 3422
2262.66	10013	10001	636	0.62. 12.3	3492.239	20013	11101	636	.450
2265,973	00011	00001	629	38,8	34%. 3%	10012	00001	63.	.463
2271.763	01111	01101	636 637	616. 7 15	3497.46	4 30002	01101	636	.0 pert 7.314
2274.367	06611	06601	626	0.0068	3500.590	21101	00001	626	.1052 .14
2281.688	22213	22203 04401	626 626	.0369	3504.320	14412	04401	626	.0110
2283.490	00011		636	9602.	3508.07	100'2	00001	637	.0771
2284.286	12211	12201	628 626	0.0115	3509.207 3511.59	21302	11101 01101	62. 626	.0046
2287.205	12212	12263	629	.0262	3517.323	20032	10001	636	.177
2286.352	13311 20012	13301 20002	626 626	.1183	3525.205 3527.703	10012 30014	20001	828 626	.0588
2290, 484	20013	20003	628	. 0180	3527.740	10012	00001	636	94.2
2290.715	13312	21102	626 626	. 3125	3527.757	22212 2001 1	12201	6 %	.0374
2293.621	21113	21102	626	. 3965	3533.942	11122	01111	626	.0353
2296.784	11111	11101	628	.325	3538.822	11112	01101	628	4.449
2299.219	04411	04401	626	4.699	3539.005	20012	11102	629 6%	3.147
2299.262	02221	02211	626	0.076	3543.095	40002	11102	626	.0 pert

			_	-9					•	
Center	Level	Level	tope	(multiply by	Band Center	Opper Level	Lower Level	Leo-	SV at 296K . (multiply by	
vo				10-22 -1-1	Va				-221	
·1.				2 -1 6	-1.				2 –1 i	
(C30))		_		cat" cat") "	(cm <sup>-</sup> )				ດຄື ດຄື)່ຽ	
3549 675	20013	10002	677	0.0334	4907 603			***	6 (0)	
3550.700	30012	20001	626	0.0195	4808.188	40002	01101	626	o.695 O. pert	
3552.B41	12212	02201	626	31.25	4814.570	20012	00001	638	.0134	
3556.749	30013	20002	626	0.0629	4839.737	30013	10001	626	.0744	
3558.595	11112	01101	627	. 668	4853.620	20013	00001	626	90,7	
3566.063	10022	00011	626	.208	4871.46	21112	01101	636 636	.238	
3568.208	20013	10002	626	33.78	4887.970	12212	00001	626	(8.1-08 4)	
3578.670	22213	12202	628	52.2	4896.185	21112	01101	628 628	.0893	
3580.334	11112	01101	626	803.5	4925.010	20011	00001	638	0.00446	
3587.540	10011	00001	638 627	0.703	4928.910	21112	01101	627 676	.00744	
3609.05	10011	00001	637	0.126	4939.350	20012	00001	627	.2306	
3612.844	10012	10001	626 636	10350.	4942.512	30013	10002	626	1.414	
3621.570	20012	10002	636	0.435	4953.363	22212	02201	626	1.042	
3623.454	21112	11102	636	.0433	4959.667	30012	10001	626	0.837	
3632.917	10011	00001	636	160.	4905.381	20012	00001	626	26.56	
3638.067	10011	00001	828	0.0392	4991.35	20001	00001	636	2.12	
3641.717	02211	00001	626	.0240 (1.7-06 Δ)	5013.785	21101 22211	01101	636 636	0.1711	
3655.42	20012	10002	628	.187	5042.57	20011	00001	628	. 227	
3656.805	21112	11102	628 627	.0149	5061.776	12211	100001	626 626	(9.2-08 Δ) .238	
3667.544	10021	00011	626	. 363	5064.680	21111	01101	628	. 260	
3675.130	10011	00001	628 626	47.8	5068.910	20011	00001	627	.0632	
3676.725	20011	10001	628	.164	5114.894	30011	10001	626	0.309	
3676.741 3679 644	30012	20002	626 626	.0915	5123.20	21111	01101	626	10.64	
3683.762	11111	01101	628	3.884	3133.401	*****	02201	040	0.409	
3677.702	21111	11101	628	0.0134	5168.60	01121	00001	636	.00372	
3692.41A	20012	10002	626	42.4	5247.030	10022	01101	626	.0101	
3623.640	10011	00001	627	10.2	5277.07	01121	00001	628	.00149	
3700.290	21112	11102	626	3.549	5291.16	02221	000001	626	.0275	
3702.345	11111	01101	627	0.771	5349.36	10021	01101	626	.00506	
3703.486	30011	20001	626	. 1503	5584.391	00031	10001	626	.00707	
3711.473	20011	10001	626	35.01	5687.166	00031	10002	626	.00751	
3713.714	21111	12201	626	2.816	5858.022	10022	00001	628	.00372	
3714.781	10011	00001	626	15000.	5951.600	30014	00001	636	.00179	
3723.249	20011	10002	636	0.0078	5959.954 5972.52	10021 32214	00001	628 626	.00335	
3726.351	14411	04401	626	.0499	5993.581	30014	00001	628	.00357	
3726.647	12211	02201 03301	626 626	48.5	6020.795 6072.343	31114 40014	01101 10001	626 626	.0465	
3783.14	20011	10002	628	0.0035	6075.983	30014	00001	626	.454	
3799.574	30012	20003	626 626	.0028	6088.21 6100.30	31113	01101 01101	636 628	.00238	
3831.968	30011	20002	626	.00092	6119.618	30013	00001	636	.0290	
3858.109	21111	11102	626 628	.016	6149.760	41114	11102	626	.00179	
					6170.090	32213	02201	626	.01265	
3987.610 4005.948	30002	01101	626	.00818 .14	6175.950	30013	00001	627	.00320	
4416.150	31104	00001	626	.00037 .15	6179.01	41102	00001	626	(1.5-06 77)	
4508.749	40004	01101	626	.00023 .12	6205.503	40013	10001	626	.0127	
4578.090	32203	01101	626	.00018 .20	6227.924	30013	00001	626	4.27	
4591.118	31103	00001	628	.01042	6243.57	31112	01101	636	.00409	
4639.502	00021	00001	628	. 1302	6254.592	30012	00001	628	.01414	
4655.205	22213	02201	627	.0127	6265.170	30012	00001	627	.00275	
4685.780	30014	10002	636	.00186	6308.278	40013	10002	626	.02455	
4687.796	30014	10001	626 638	.00521	6318.17 6346.265	41113 40012	10001	626 626	.0119	
4708.52	21113	01101	636	.0298	6347.854	30012	00001	626	4.27	
4733.50	23313	03301	626	.00670	6356.293	31112	01101	626	0.327	
4748.058	20013	00001	636	.214	6363.616	30011	00001	636	.0127	
4753.450	31102	00001	626	.00298 .10	6388.015	41101	00001	626	(4.8-06 TT)	
4755.705 4768.541	22213	02201	626	.2604	6429.172	30011	00001	628	.00112	
4784.675	2002 3	00011	626	.00149	6466.440	20023	01101	626	.00104	
4786.688 4790.571	31113 30014	11101 m o o 2	626 626	.0119 .1562	6498.67 6503.081	12222	01101 00001	626 626	.00119 .498	
4791.260	20013	00001	628	. 469	6532.653	40011	10001	626	.00130	

Table 10. Band Origins and Intensities for  $\mathbf{CO}_{\mathbf{2}}(\mathbf{Contd})$ 

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ning is being diversities which the surely in Station

	Band	Upper	Lower	I so-	SV At 296K	
	Center	Level	Level	tope	(multiply by	
	~ <b>o</b> ~				10 <sup>-22</sup> m1 <sup>-1</sup>	
	(cm 1)				cant can <sup>-1</sup> ) y	
_						
	6536.445	31111	01101	626	.0476	
	6537.958	11122	00001	626	.0223	
	6562.444	32211	02201	626	.00223	
	6679.709	11121	00001	626	.0263	
				676	01330	Notes to Table IV-2.2
	6745.115	01131	01101	910	.01339	
	0/00.215	00031	00001	476	.1037	Bands deriving all their intensity from J-dependent
	6867 360	11111	03301	626	00112	perturbation are designated o pert, when one near
	6970 704	11132	11101	626	00241	band provides the intensity, $\pi$ , when the listed
	6885 150	01131	11102	628	.00402	intensity is multiplied by $m(m+1)$ , and $\Delta$ , when the
	6897.751	02231	01101	626	.0424	listed intensity is multiplied by $m^2$ (m+1) <sup>2</sup> .
	6905.770	10031	10001	626	.0171	
	6907.144	10032	10001	626	.0290	Q designates bands with intensity below the
	6922.210	00031	00001	628	.0521	criterion limit, whose Q - branches are
	6935.150	01131	01101	626	1.131	significant.
	6945.610	00031	00001	627	0.0112	
	6972.578	00031	00001	626	14.95	·
	7283.981	40015	00001	626	0.00186	
	7460.530	40014	00001	626	.0428	
	7481.51	40013	00001	636	.00112	
	7583.265	41113	01101	626	.00833	
	7593.690	40013	00001	626	.1064	
	7616.620	51102	00001	626	(1.12-06 17)	
	7734.452	40012	00001	626	.02/9	
	7/37.621	41113	01101	626	00148 018	
	7901.479	40001	00001	626	.00145 .018	
	7941 140	10032	00001	626	- 00232	
	8084.060	12232	02201	630	.00193	
	8089.84	10031	00001	616	.00707	
	8103.578	20033	10002	626	.00205	
	8120.104	10032	00001	628	.00201	
	8128.783	20032	10001	676	.00071	
	8135.886	11132	01101	626	.0402	
	8192.556	10032	00001	626	.424	
	8220.363	10031	00001	628	.00201	
	8243.163	20031	10001	626	.00108	
	8254.800	12231	02201	626	.00164	
	8276.767	11131	01101	626	.0461	
	8293.957	10031	00001	626	.614	
	9388.990	20033	00001	626	.00415	
	9478.051	21132	01101	626	.00190	
	9010.970	20032	00001	626	.0233	
	10.11.150	218131		6.76	. (8.79.5	

Table 10. Band Origins and Intensities for CO<sub>2</sub> (Contd)
of  $\zeta^{v}$  included in Table 9. Note that when  $\zeta_{v}$  is larger than 0.01, the total band strength, if defined as the sum of the Q, R- and P-branch lines,  $\underset{m}{\overset{r}{m}} S_{m}$ , may be much larger than  $S_{v}^{o}$ , its value when  $\zeta_{v} = 0$ . The entry in Table 10 is  $S_{v}^{o}$ , not  $\underset{m}{\overset{r}{m}} S_{m}$ .

As a higher-order effect of Coriolis interactions, transitions with l' - l'' = + 2 also become allowed, the effect increasing as  $m^2$ , so that a few such bands are intense enough to appear; here we list the value of  $\mathbf{S_v^o}$  in the formula  $\mathbf{S_m} = \mathbf{S_v^o}(|\mathbf{m}| + \mathbf{m}^2) \exp[(-1.439\mathbf{E''})/\mathbf{T}]/\mathbf{Q_r}$ . There are also a few special cases of near-or crossing-perturbations, where

There are also a few special cases of near-or crossing-perturbations, where a normally **very** weak band "borrows" intensity from a stronger neighbor. A notable example is represented by the bands at 4808.186 cm<sup>-1</sup>, where the 40002-01101 band with zero intensity, resonates with the 21113-01101 band at 4807.692 cm<sup>-1</sup>. The positions and intensities of the individual lines are here calculated using an appropriate Coriolis constant and dividing the strength of 21113 -01101 according to the degree of perturbational mixing.

## 4.2.3 CO, LINE HALF-WIDTHS

The half-widths for  $CO_2$  were assigned by a linear function of the rotational quantum number. The work of Yamamoto, Tanaka and Aoki (1969) gave methods of estimating half-widths for the 15  $\mu$ m and 4.3  $\mu$ m regions, but the differences between their limited comparisons of calculated values and laboratory measurements did not seem to warrant any more complicated procedure in arriving at a half-width dependence upon J for  $CO_2$ .

#### 4.3 Ozone

The band centers and intensities of all bands included in the compilation are given in Table 11.

The pure rotational transitions of ozone have been calculated using the molecular constants of Lichtenstein, Gallagher, and Clough (1971) obtained from microwave measurements. Transitions with upper state J less than 61 and intensities greater than 3.5 x  $10^{-24}$  cm<sup>-1</sup>/mol-cm<sup>2</sup> have been tabulated using a value of  $\mu$ =0.53D for the dipole moment. Lines with K<sub>a</sub> less than 10 should be accurate to 0.001 cm<sup>-1</sup>. Multiplets up to K<sub>a</sub> = 12 (≈100 cm<sup>-1</sup>) have been observed and the calculated contours agree within the experimental accuracy (see Gebbie et al, 1966). At higher values of K<sub>a</sub>, the calculations become increasingly less certain. However, even the highest value of K<sub>a</sub> included in the listing ( that is, K<sub>a</sub> = 20) is substantially less #an the value of K for which the calculation is divergent. The mean line width has been determined to be 0.11 (Lichtenstein et al, 1971).

3and Center	Σ <sub>i</sub> S <sub>i</sub>	Vibrational	Transition	Isotope
cm <sup>-1</sup>	$\frac{\mathrm{cm}^{-1}}{\mathrm{mol/cm}^2}$	Upper State	Lower State	
$\begin{array}{c} 0.\\ 700.930\\ 1007.693\\ 1007.996\\ 1021.096\\ 1027.096\\ 1028.096\\ 1042.096\\ 1042.096\\ 1103.157\\ 2110.790\\ 2785.241\\ 3041.200 \end{array}$	$\begin{array}{c} 4.13 \times 10^{-19} \\ 6.70 \times 10^{-19} \\ 9.49 \times 10^{-21} \\ 2.49 \times 10^{-20} \\ 4.23 \times 10^{-19} \\ 1.62 \times 10^{-19} \\ 5.16 \times 10^{-20} \\ 5.16 \times 10^{-27} \\ 1.29 \times 10^{-17} \\ 3.47 \times 10^{-19} \\ 1.33 \times 10^{-18} \\ 2.32 \times 10^{-20} \\ 1.10 \times 10^{-19} \end{array}$	000 010 101 001 011 002 001 001 100 101 111 003	000 000 100 000 010 001 000 000 000 000	

Table 11. Ozone Transitions Included in Data Compilation

For the  $\nu_2$  fundamental of ozone, upper-state molecular constants have been obtained from microwave data by Tanaka and Morino (1970a). For a discussion of the determination of the band center see Tanaka and Morino (1970b). Transitions up to J = 60 with intensities greater than  $3.5 \times 10^{-23} \text{ cm}^{-1}/\text{mol-cm}^2$  have been tabulated. The dipole moment constant for the  $\nu_2$  fundamental has been taken as  $(\partial \mu / \partial q_2) q_2 = 0.048D$  giving a band intensity of 6.76  $10^{-19} \text{ cm}^{-1}/\text{mol-cm}^2$  based on the measurements of McCaa and Shaw (1968).

The  $\nu_3$  and  $\nu_1$  regions of the ozone spectrum have been studied by Clough and Kneizys (1965, 19661, and Tanaka and Morino (1968). The positions of the  $\nu_1$  and  $\nu_3$  transitions for the main isotope are expected to be accurate to 0.01 cm<sup>-1</sup> for J less than 40 and  $K_a$  less than eight with a degradation of accuracy for higher values of the quantum numbers. In conformity with the results of McCaa and Shaw (1968), the band intensity of  $\nu_3$  has been taken as 130 x  $10^{-19} \text{ cm}^{-1}/\text{mol-cm}^2$ .

Also, more lines have been included than were given in the report of Clough and Kneizys (1965). Goldman, et al (1970) have found that the absorption by large ozone amounts could not be satisfactorily explained without introducing the hot bands and the isotopic bands. Since the energy levels are known for the 101, the 100 and the 010 vibrational levels, transitions for the 101  $\leftarrow$  100 and 101  $\leftarrow$  001 bands may be calculated directly. The line positions for these bands with J less than 30 and  $K_a$  less than six are presumed accurate to +0. 03 cm<sup>-1</sup>. The line positions for the 011  $\leftarrow$  010 hot band have been obtained by applying a calculated shift of -2 1.0 cm<sup>-1</sup> to the  $\nu_3$  transitions of the main isotope. For J less than 30 and  $K_a$  less than six an error of up to 1 cm<sup>-1</sup> may be expected. For the 002  $\leftarrow$  001

hot band, a calculated shift of -15.0 cm<sup>-1</sup> has been applied to the  $\boldsymbol{\nu}_3$  transitions of the main isotope. In this case, the rotational levels of the upper- and **lower-**vibrational states are not accurately described, and these lines must be considered to be useful only for their general contribution to the low-resolution absorption in the region. These **shifts** have been calculated using harmonic frequencies and **an**-harmonic corrections determined from an analysis of the available vibrational data.

For the 668 and 686 isotopes, shifts of -13.1 cm<sup>-1</sup> and -34.1 cm<sup>-1</sup> respectively have been determined from force constant calculations using the structural constants given by Tanaka and Morino (1968). These shifts have similarly been applied to the original  $\boldsymbol{\nu}_3$  line calculations, neglecting the isotopic effect on the rotational constants. Consequently, only the general contribution to low -resolution spectra by these lines is valid. The intensities of the hot bands and the isotopic bands have been determined by applying the appropriate Boltzmann correction, isotopic abundance factor, dipole moment matrix element value, and degeneracy factor to the intensities of the  $\boldsymbol{\nu}_3$  fundamental.

Although the addition of the isotopic bands and the hot bands gives improved agreement between calculated absorption and the observed absorption for large ozone amounts, it appears that an intensity anomaly still exists. In the P branch region (  $\sim$  1000 cm-') the calculated intensity is evidently too low, while in the R branch region the calculated intensity is too high. Until better intensity measure – ments are available, it will be difficult to obtain experimental F factors (see Section 3.2).

The  $(\nu_1 + \nu_3)$  combination band (101  $\leftarrow$  000) has been observed and analyzed using the **rig:d** rotor approximation by Trajmar and McCaa (1964). The value of the band center was found to be 2110.79 cm<sup>-1</sup>. The resolution and calibration accuracy of the observed spectra were not good, so that the line positions up to J = 20 and  $K_a = 4$  are accurate to only 0.3 cm<sup>-1</sup>. The error in line position is expected to be significantly greater for higher quantum numbers. The band intensity for the tabulated lines is 1.19 x 10<sup>-18</sup> cm<sup>-1</sup>.

The  $(\nu_1 \pm \nu_2 \pm \nu_3)$  combination band  $(111 \leftarrow 000)$  at 2785.241 has been studied by Snider and Shaw (1971). The rigid rotor analysis used gives line positions for J less than 25 and K<sub>a</sub> less than 6 accurate to 0.1 cm<sup>-1</sup>. The intensity for this band is 2.46 x 10<sup>-20</sup> cm<sup>-1</sup> and was also obtained from McCaa and Shaw (1968).

Constants for the 3  $\nu_3$  band (003 – 000) have been obtained from a second

order calculation using the results of Tanaka and Morino (1968). The band intensity has been taken as  $1.12 \times 10^{-19} \text{ cm}^{-1}/\text{molecule-cm}^{-2}$  from McCaa and Shaw (1968). The accuracy of the line positions should not be in error by more than  $\pm 5 \text{ cm}^{-1}$ 

#### 4.4 Nitrous Oxide

Nitrous oxide is an asymmetric linear molecule (N-N-O) with three fundamental vibration-rotation bands,  $\nu_1, \nu_2$ , and  $\nu_3$ , at 1284.907, 588.767, and 2223.756 cm<sup>-1</sup>, respectively, for the main isotope. As for  $CO_2, \nu_1 \cong 2\nu_2$  so that Fermi resonance occurs between these two levels, causing shifts of the band centers from the positions otherwise expected. Five different isotopes provide significant absorption. When no reference to a particular isotope is made, the most abundant one, 446, is implied. The abbreviated notations and natural abundances of each isotope used to calculate intensities of the isotopic bands are (446), 99.043 percent; (456), 0.358 percent; (546), 0.358 percent; (448), 0.199 percent; and (447), 0.040 percent. These values differ slightly from those given in Table 4, the difference arising from slight differences in the oxygen and nitrogen natural abundances taken from the 48th edition and the 42nd edition of the Handbook of Physics and Chemistry.\* The 48th edition values were used in the construction of Table 4. These abundance differences are very small, but can be accounted for by modifying the line intensities accordingly.

# 4.4.1 LINE POSITIONS

Table 12 lists the constants used to calculate the line positions. Values of the vibrational energy, G, are relative to the  $00^{\circ}0$  level for each isotope and are probably accurate to a few thousandths of a cm<sup>-1</sup> for the 446 isotope. The uncertainty is much greater for some of the levels of the rarer isotopes. The majority of the constants for the 446 isotope and a few for the other isotopes are from Pliva (1964, 1968a, 1968b). Most of the other values have been derived by Benedict (private communications1 from a variety of data.

Line positions for the P- and R-branches can be calculated by inserting the constants from Table 12 into Eq. (6). The selection rules are:  $AJ = 0, \pm 1; \Delta l = 0, \pm 1; \Delta l = 0, \pm 1; and J = 0 + J = 0$ . The allowed values of J for a given vibrational level must be taken into account in order to determine missing lines. For example, in a  $\phi \leftarrow \Delta$  band (l' = 3 - l'' = 2) the following lines are missing because of rotational energy levels that are not allowed: P1, P2, P3, QO, Q1, Q2, RO and R1.

A few lines of the  $10^{\circ}1, 06^{\circ}0, 06^{2}0$ , and  $10^{\circ}1 - 10^{\circ}0$  bands are shifted from the positions based on the constants in Table 12 because of Fermi interactions. These are discussed in more detail below and summarized in Table 15.

<sup>\*</sup>Published by the Chemical Rubber Co., Cleveland, Ohio

and an	

Energy Level	G	В	D Multiply	M&ply
			by <b>10<sup>-7</sup></b>	by <b>10<sup>-12</sup></b>
	(cm $^{-1}$ )	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )
		Isotope 446		
00 <sup>0</sup> 0	0	0.4190113	1.795	1.17
01 <sup>1°</sup> 0	588.767	0.4191777	1.785	1.17
01 <sup>1d</sup> 0	588.767	0.419969 5	1.785	1.17
02 <sup>0</sup> 0	1168.134	0.4199193	2.445	1.17
02 <sup>20</sup> 0	1177.750	0.4201253	1.165	1.17
02 <sup>2d</sup> 0	1177.750	0.4201253	1.795	1.17
10 <sup>0</sup> 0	1284.907	0.4172563	1.775	1.17
03 <sup>1°</sup> 0	1749.058	0.4196063	2.195	1.17
03 <sup>1d</sup> 0	1749.058	0.4210883	2.195	1.17
03 <sup>3d</sup> 0	1766.958	0.420674	1.805	2.20
03 <sup>3°</sup> 0	1766.958	0.420674	1.805	0.14
11 <sup>10</sup> 0	1880.268	0.4174673	1.765	1.17
11 <sup>1d</sup> 0	1880.268	0.4183803	1.775	1.17
04 <sup>0</sup> 0	2322.570	0.4206113	4.095	16.17
04 <sup>20</sup> 0	(2331.15)	0.4210113	1.350	1.17
04 <sup>2d</sup> 0	(2331.15)	0.4210113	2.50	1.17
12 <sup>0</sup> 0 12 <sup>20</sup> 0 12 <sup>2d</sup> 0 20 <sup>0</sup> 0 00 <sup>0</sup> 1	2461.998 2474.785 2474.785 2563.341 2223.756	0.4181483 0.4187143 0.4187143 0.4224193 0.4155613	2.465 1.210 1.700 1.645 1.795	3.77 1.17 1.17 1.17 1.17 1.17
05 <sup>1C</sup> 0	2897.876		2.085	1.17
05 <sup>1d</sup> 0	2897.876		2.355	1.17
13 <sup>1C</sup> 0	3046.213	0.4177633	2.145	1.17
13 <sup>1d</sup> 0	3046.213	0.4193783	2.165	1.17
13 <sup>3C</sup> 0	3067.749	0.419109	1.805	0.47
13 <sup>3d</sup> 0	; 3067.749	0.419109	1.805	1.87

Table 12. Vibrational Energy Levels and Molecular Constants for  $N_2O$ 

				<u> </u>
Energy	G	В	D Multiply	H Multiply
Dever	i		by 10 <sup>-7</sup>	by lo-:"
	$(cm^{-1})$	$(cm^{-1})$	$(cm^{-1})$	by 10 ,
		<u>Isotope 446</u> (Contd]		
21 <sup>1°</sup> 0	3165.857	0.4158333	1.595	1.17
21 <sup>1d</sup> 0	3165.857	0.4169163	1.595	1.17
01 <sup>1°</sup> 1	2798.290	0.4157723	1.795	1.17
01 <sup>1d</sup> 1	2798.290	0.4165473	1.795	1.17
14 <sup>0</sup> 0	3620.941	0.4187873	3.885	14.17
14 <sup>2°</sup> 0	3631.601	0.4190143	0.375	1.17
14 <sup>2d</sup> 0	3631.601	0.4190143	2.045	1.17
22 <sup>0</sup> 0	3748.252	0.4163273	2.395	2.77
22 <sup>2<sup>c</sup></sup> 0	3766.060	0.4172013	1.200	1.17
22 <sup>2d</sup> 0	3766.060	0.4172013	1.560	1.17
30 <sup>0</sup> 0	3836.373	0.4141473	1.385	1.17
02 <sup>0</sup> 1	3363.974	0.4165443	2.445	1.17
02 <sup>2°</sup> 1	3373.137	0.4167523	1.195	1.17
02 <sup>2d</sup> 1	3373.137	0.4167523	1.795	1.17
10 <sup>0</sup> 1	3480.821	0.4137843	1.745	1.17
23 <sup>10</sup> 0	4335.798	0.4159193	2.045	1.17
23 <sup>1d</sup> 0	4335.798	0.4176813	2.115	1.17
31 <sup>10</sup> 0	4446.379	0.4143703	1.625	1.17
31 <sup>1ª</sup> 0	4446.379	0.4156713	1.365	1.17
03 <sup>1°</sup> 1	3931.258	0.4162253	1.915	1.17
03 <sup>10</sup> 1	3931.258	0.4176843	1.925	1.17
$03^{3^{c}1}$	3948.344	0.417327	1.815	0.14
03 <sup>30</sup> 1	3948.344	0.417327	1.815	2.20
11 <sup>1c</sup> 1	4061.979	0.4140513	1.775	1.17
11101	4061.979	0.4149343	1.735	1.17
40 <sup>0</sup> 0	5105.65	0.4131913	1.795	1.17
320	5026.34	0.4143113	1.795	1.17
2001	4730.828	0.4121163	1.625	1.17
121	4630.164	0.4147633	2.475	4.17
$12^{2^{2}1}$	4642.463	0.4151583	1.315	1.17
12 <sup>2<sup>u</sup>1</sup>	4642.463	0.4151583	1.315	1.17
002	4417.379	0.4120963	1.765	1.17

Table 12. Vibrational Energy Levels and Molecular Constants for  $\rm N_2O\left(Contd\right)$ 

Energy Level	G	В	D Multip <b>ly</b>	H Multiply
	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )	by 10 <sup>-7</sup> (cm <sup>-1</sup> )	by 10 <sup>-12</sup> (cm <sup>-1</sup> )
<u> </u>		Isotope 446 (Contd)		
$21^{1c}1$	5319.175	0.4124313	1.675	1.17
211	5319.175	0.4134703	1.555	1.17
$01^{10}2$ $01^{10}2$	4977.695 4977.695	0.4123583 0.4131183	1.785 1.785	1.17 1.17
-		Isotope 456		-
00 <sup>0</sup> 0	0	0.4189821	1.75	
01 <sup>10</sup> 0 01 <sup>10</sup> 0	575.5 575.5	0.419095 0.419891	1.75 1.75	
10 <sup>0</sup> 0	1280.5	0.41719	1.72	
11 <sup>10</sup> 0 11 <sup>1d</sup> 0	1861.9 1861.9	0.41734 0.41820	1.71 1.70	
20 <sup>0</sup> 0 00 <sup>0</sup> 1	2554.3 2177.659	0.41545 0.41568	1.67 1.75	
01 <sup>1C</sup> 1 01 <sup>1d</sup> 1	2739.63 2739.63	0.415855 0.416605	1.75 1.75	
10 <sup>0</sup> 1	3430.95	0.41387	1.72	
		Isotope 546		
00 <sup>0</sup> 0	0	0.4048564	1.64	_
01 <sup>10</sup> 0 01 <sup>10</sup> 0	585.320 585.320	0.4050304 0.4057724	1.65 1.65	
10 <sup>0</sup> 0	1269.894	0.403269	1.60	
11 <sup>10</sup> 0 11 <sup>10</sup> 0	1863.080 1863.080	0.4034614 0.4043814	1.59 1.57	
20 <sup>0</sup> 0 00 <sup>0</sup> 1	2534.21 2201.604	0.401870 0.401495	1.45 1.65	
01 <sup>1C</sup> 1 01 <sup>1d</sup> 1	2745.709 2745.709	0.4017054 0.4024104	1.65 1.65	
10 <sup>0</sup> 1	3443.659	0.399876	1.60	

Table 12. Vibrational Energy Levels and Molecular Constants for  $\mathbf{N_2O}(\mathbf{Contd})$ 

				_
Energy Level	G	В	D Multiply by <b>10<sup>-7</sup></b>	H Multiply by <b>10<sup>-12</sup></b>
	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )	(cm <sup>-1</sup> )	(cm -1)
		Isotope 448		
00 <sup>0</sup> 0	0	0.395577	1.65	
01 <sup>1C</sup> 0 01 <sup>1d</sup> 0	584.1 584.1	0.39 5749 0.396461	1.65 1.65	
10 <sup>0</sup> 0	1247.9	0.394057	1.55	
11 <sup>10</sup> 0 11 <sup>1d</sup> 0	. <b>1839.8</b> 1839.8	0.39430 0.39513	1.56 1.50	
20 <sup>0</sup> 0 00 <sup>0</sup> 1	2491.3 2218.97	0.39279 0.392317	1.31 1.65	
01 <sup>1C</sup> 1 01 <sup>1d</sup> 1	2786.80 2788.80	0.392549 0.393209		
10 <sup>0</sup> 1	3439.1	0.39078	1.55	
		Isotope 447		
00 <sup>0</sup> 0	0	0.406691	1.72	
01 <sup>1C</sup> 0 01 <sup>1d</sup> 0	586.3 586.3	0.406860 0.407610	1.72 1.72	
10 <sup>0</sup> 0	1265.5	0.405961	1.67	
00 <sup>0</sup> 1	2221.3	0.40334		
01 <sup>1C</sup> 1 01 <sup>1d</sup> 1	2793.55 2793.55	0.403 57 0.40430		

Table 12. Vibrational Energy Levels and Molecular Constants for  $\mathbf{N_2O}\left(\mathbf{Contd}\right)$ 

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#### 4.4.2 LINE INTENSITIES

Each band system contains a main band and associated hot bands that arise from transitions from excited energy levels with the same changes  $\Delta v_1, \Delta v_2$ , and  $\Delta v_3$  in the vibrational quantum numbers as in the main band. The **change**  $\Delta l$  may be either 0 or  $\pm 1$  within the same band system. For example, the 0°01 band is the main band in a system that also contains the hot bands  $01^{1}1 \leftarrow 01^{1}0, 02^{0}1 \leftarrow 02^{0}0,$  $02^{2}1 \leftarrow 02^{2}0, 10^{0}1 \leftarrow 10^{0}0$ , and many others from higher excited energy levels. The relative intensities of the hot bands decrease with increasing value of the lower energy level because of the decrease in its population. Thus, only a few of the hot bands are usually significant. Corresponding bands of the rare isotopic molecules are also part of the system. The main band of the common isotope accounts for 78 percent to 89 percent of the entire system for  $296^{0}K$  samples, depending on the band system.

Table 13 summarizes the normal band systems containing lines with intensities greater than 4 x 10<sup>-23</sup> molecules <sup>-1</sup> cm<sup>2</sup> cm<sup>-1</sup>. Each band system is identified by the main band whose center position is given. The intensities are based on recent measurements of the quantity  $\int (-1/u) \ln T dv$  by Burch et al, (1971a, 1971b, 1971c, 1972). The integration was performed over the entire band system, and u is the N<sub>2</sub>O thickness in molecules cm<sup>-2</sup>. The intensity, S<sub>Q</sub>, of only the Q-branch of the band at 1974.571 cm<sup>-1</sup> is listed because the P and R branches contain no lines with intensities above the cut-off. The Q-branch may be significant although each of its lines is very weak because they occur very close together. Bands containing the 06<sup>o</sup>0 and 06<sup>2</sup>0 levels are not listed because the band systems are very weak. However, as a result of Fermi interaction between a few of the rotational levels, the intensities of the corresponding lines are anomalously high and their positions arc shifted. These lines are discussed in more detail below.

Table 14 lists the intensities of each of the bands with lines above the cutoff intensity, nominally  $4 \times 10^{-23}$  molecules  $^{-1}$  cm<sup>2</sup> cm<sup>-1</sup>. Each line entry in the table corresponds to a given band, and blank lines occur before and after each **band** system. Note that 16 bands are listed for the **00<sup>o</sup>1** band system, whereas only 1 or 2 bands are listed for some of the weaker systems.

Each rotationallevel within a vibrationallevel for which quantum number  $l \neq 0$  is split into two sublevels normally indicated by "c" and "d" (see section 3. 1. 1). A band involving two A levels, or two  $\pi$  levels, or one of each level consists of two sub-bands, one from the c portion of the lower level and one from the d portion. The  $c \leftarrow c$  portion of the first sub-band contains only P- and R-branches, whereas the d  $\cdot$  - c portion contains the Q-branch. Similarly, the d  $\leftarrow$  d and the c  $\leftarrow$  d portions of the second sub-band contain the P t R branches and the Q-branch, respectively. The **intensi-ties**,  $S_{\mu\nu}$  given in Table 14 are the total for both sub-bands.

Center of Main Band (cm <sup>-1</sup> )	Upper Level	Lower Level	Intensity of System Multiply by 10 <sup>-20</sup> Molecules <sup>-1</sup> cm <sup>2</sup> cm <sup>-1</sup>
Main Band (cm <sup>-1</sup> ) 588.767 696.140 938.849 1168.134 1284.907 1634.989 1749.058 1880.268 1974.571 2223.756 2322.624 2461.998 2563.341 2798.290 3363.974 3480.821 3620.941 3748.252 3836.373 4061.979 4335.798 4417.379	Level 0110 10°0 00°1 02°0 10°0 10°0 1110 20°0 00°1 04°0 12°0 20°0 00°1 04°0 12°0 20°0 0111 02°1 10°1 14°0 22°0 30°0 20°0 0110 00°1 00°2 00°2 00°2	Level 00°0 01 <sup>1</sup> 0 10°0 00°0 00°0 01 <sup>1</sup> 0 00°0 00°0 0110 00°0 0	Multiply by 10 $^{-1}$ Molecules $^{-1}$ cm <sup>2</sup> cm <sup>1</sup> 118 $\pm$ 9 0.354 t 0.020 0.254 $\mp$ 0.010 38.5 t-i.5 996 $\pm$ 40 0.278 t 0.02 0.241 $\mp$ 0.02 1.66 $\pm$ 0.08 So = 0.024 $\pm$ 0.002 5710 $\pm$ 250 2.7 t 0.3 33.4 $\pm$ 1.5 135 t-7 9.62-t 0.96 10.6 $\mp$ 0.5 197 t-10 0.56-t 0.02, $-$ 0.05 4.12 t 0.2 8.15 $\pm$ 0.4 0.111 t 0.006 0.1 t 6.1, $-$ 0.07 6.9 t 0.7
4630.164 4730.828 4977.69 5 5026.34 5105.65	$ \begin{array}{r} 1201\\ 2001\\ 0112\\ 3200\\ 4000 \end{array} $	00°0 00°0 00°0 00°0 00°0	0.68 + 0.07 4.4 t-b.4 0.071) t 0.008 0.29 t-b.04 0.29 <del>+</del> 0.03

Table 13. Summary of N<sub>9</sub>O Band Systems

<b>F</b> Sand Center	Upper	Lower	Isotope	S at 296K
	Level	Level	Botobe	$\sqrt{10^{-20}}$
-1		20.00		(Multiply by $10$
(cm )				Molecules cm cm I
588.767 579.367 588.983 580.924 57 1.308 589.208 595.361 585.320 575.5 586.3 584.1	01 <sup>1</sup> 0 0200 0220 0310 0330 0330 1110 0110 011	00 <sup>°</sup> 0 0110 0110 02 <sup>°</sup> 0 0220 02 <sup>°</sup> 0 10 <sup>°</sup> 0 00 <sup>°</sup> 0 00 <sup>°</sup> 0 00 <sup>°</sup> 0	$\begin{array}{c} 446\\ 446\\ 446\\ 446\\ 446\\ 446\\ 446\\ 546\\ 456\\ 447\\ 448\end{array}$	$\begin{array}{c} 98.4 \\ 4.90 \\ 11.26 \\ 0.599 \\ 0.256 \\ 0.958 \\ 0.216 \\ 0.356 \\ 0.356 \\ 0.039 \\ 0.197 \end{array}$
696.140	10 <sup>0</sup> 0	01 <sup>1</sup> 0	446	0.294
938.849	00 <sup>0</sup> 1	10 <sup>0</sup> 0	446	0.223
1168.134 1160.291 1154.436 1153.40	020 0310 0400 0420	0000 010 0200 0220	$\begin{array}{c} 446 \\ 446 \\ 446 \\ 446 \\ 446 \end{array}$	31.7 5.72 0.201 0.228
1177.750	0220	0000	446	See Note A
$\begin{array}{c} 1284.907\\ 1291.501\\ 1293.864\\ 1297.035\\ 1278.434\\ 1297.155\\ \textbf{1300.791}\\ 1285.589\\ 1280.5\\ 1286.4\\ 1269.894\\ 1277.760\\ 1247.9\\ 1255.7\\ 1265.5 \end{array}$	10°0 1110 12°0 1220 20°0 13°0 13°0 13°0 11°0 11°0 11°0 11°0 1	00°0 0110 02°0 0220 10°0 0310 03°0 1110 00°0 01°0 01°0 01°	$\begin{array}{c} 446\\ 446\\ 446\\ 446\\ 446\\ 446\\ 446\\ 446$	$\begin{array}{c} 872.\\ 100.6\\ 3.16\\ 5.92\\ 3.73\\ 0.390\\ 0.364\\ 0.412\\ 3.15\\ 0.354\\ 3.15\\ 0.354\\ 3.15\\ 0.362\\ 1.75\\ 0.200\\ 0.350\end{array}$
1634.969	00 <sup>0</sup> 1	01 <sup>1</sup> 0	446	0.231
1749.058	0310	0000	446	0.200
1880.268 1886.018 1873.231	11 <sup>1</sup> 0 1220 12°1	<b>00<sup>0</sup>0</b> 0110 0110	$\begin{array}{c} 446\\ 446\\ 446\end{array}$	$1.41 \\ 0.156 \\ 0.053$
1974.571	20 <sup>0</sup> 0	0110	446	<b>S<sub>Q</sub> =</b> 0.024
2181.66	06 <sup>0</sup> 0	10 <sup>0</sup> 0	446	See Note B

Table 14. N20 Band Intensities

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Band Center	Upper	Lower	Isotope	S, at <b>296K</b>
ν <sub>c</sub>	Level	Level		(Multiply by 10 <sup>-20</sup>
(cm <sup>-1</sup> )				Molecules $^{-1}$ cm <sup>2</sup> cm <sup>-1</sup> )
2223.756 2209.523 2195.840 2195.387 2195.914 2182.200 2181.386 2181.711 2193.623 2177.659 2164.13 2201.604 2187.389	0001 011 0201 0221 1001 0311 0331 111 0002 0001 0111 011	00 <sup>0</sup> 0 0110 0220 10 <sup>0</sup> 0 0310 0330 11 <sup>1</sup> 0 00 <sup>0</sup> 1 00 <sup>0</sup> 0 01 <sup>0</sup> 0 01 <sup>0</sup> 0	$\begin{array}{c} 446\\ 446\\ 446\\ 446\\ 446\\ 446\\ 446\\ 446$	$5023. \\ 568. \\ 16.8 \\ 31.8 \\ 10.1 \\ 1.958 \\ 1.88 \\ 1.10 \\ 0.201 \\ 18.1 \\ 2.04 \\ 18.1 \\ 2.10$
2218.97 2204.70 2221.3 2207.25	00 <sup>0</sup> 1 01 <sup>1</sup> 1 00 <sup>0</sup> 1 01 <sup>1</sup> 1	0000 0110 0000 0110	448 448 447 447	10.1 1.15 2.01 0.234
2322.624 2309.109	04 <sup>0</sup> 0 05 <sup>1</sup> 0	$00^{0}0$ $01^{1}0$	$\begin{array}{c} 446\\ 446\end{array}$	2.11 0.54
2461.998 2457.446 2452.807 2453.851 2463.345	1200 1310 1400 1420 2200	00 <sup>0</sup> 0 0110 02 <sup>0</sup> 0 0220 10 <sup>0</sup> 0	$\begin{array}{c} 446 \\ 446 \\ 446 \\ 446 \\ 446 \\ 446 \end{array}$	27.6 4.68 0.239 0.321 0.192
2563.341 2577.090 2580.118 2588.310 2551.466 2534.21 2554.3 2491.3	20 <sup>0</sup> 0 21 <sup>1</sup> 0 22 <sup>0</sup> 0 2220 30 <sup>0</sup> 0 20 <sup>0</sup> 0 20 <sup>0</sup> 0 20 <sup>0</sup> 0	0000 0110 0200 0220 1000 0000 0000 0000	$\begin{array}{c} 446 \\ 446 \\ 446 \\ 446 \\ 446 \\ 546 \\ 456 \\ 446 \end{array}$	$120. \\ 12.2 \\ 0.348 \\ 0.648 \\ 0.696 \\ 0.454 \\ 0.384 \\ 0.192$
2474.785	1220	00 <sup>0</sup> 0	446	See Note A
2798.290 2775.207 2784.370 2763.124 2753.508 2770.594	01 <sup>1</sup> 1 0291 021 0311 0311 0331	0000 0110 0110 0200 0220 0220	$\begin{array}{c} 446 \\ 446 \\ 446 \\ 446 \\ 446 \\ 446 \\ 446 \end{array}$	8.06 0.401 0.912 0.025 0.020 0.078
3363.974 3342.491	$02^{01} \\ 03^{1}1$	<b>00<sup>0</sup>0</b> 0110	446 446	8.82 1.57

Table 14. N<sub>2</sub>O Band Intensities (Contd)

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3and Center <sup> <sup> v</sup>c (cm<sup>-1</sup>) </sup>	U <b>pper</b> Level	Lower Level	Isotope	<b>S<sub>v</sub></b> at 296K (Multiply by 10 <sup>-20</sup> Molecules <sup>-1</sup> cm <sup>2</sup> cm <sup>-1</sup> )
3480.821 3473.212 3462.030 3464.713 3445.921 3443.659 3430.95 3439.1	10 <sup>0</sup> 1 1111 12 <sup>0</sup> 1 1221 20 <sup>0</sup> 1 10 <sup>0</sup> 1 10 <sup>0</sup> 1	00°0 0110 02°0 0220 10°0 0000 00°0 00°0	$\begin{array}{r} 446 \\ 446 \\ 446 \\ 446 \\ 446 \\ 546 \\ 456 \\ 448 \end{array}$	$\begin{array}{c} 173.\\ 19.24\\ 0.563\\ 1.064\\ 0.652\\ 0.627\\ 0.627\\ 0.348\end{array}$
<b>3466.54</b> 3474.65	06 <sup>0</sup> 0 06 <sup>2</sup> 0	00 <sup>0</sup> 0 00 <sup>0</sup> 0	$\begin{array}{c} 446\\ 446\end{array}$	See Note B See Note A and B
3748.252 3747.031	$22^{0}_{23}_{0}_{0}$	$0000 0 \\ 0110$	$\begin{array}{c} 446 \\ 446 \end{array}$	3.52 0.512
3620.941	14 <sup>0</sup> 0	00 <sup>0</sup> 0	446	0.492
3836.373 3857.612	<b>30<sup>0</sup>0</b> 3110	<b>00<sup>0</sup>0</b> 0110	$\begin{array}{c} 446 \\ 446 \end{array}$	7.28 0.698
4061.979	11111	00 <sup>0</sup> 0	446	0.092
4335.798	$23^{1}0$	00 <sup>0</sup> 0	446	0.083
4417.379 4388.928	<b>00<sup>0</sup>2</b> 0112	$0000 \\ 0110$	446 446	6.07 0.686
4630.164	12 <sup>0</sup> 1	0000	446	0.597
4730.828 4730.408	<b>20<sup>0</sup>1</b> 2111	<b>00<sup>0</sup>0</b> 0110	$\begin{array}{c} 446\\ 446\end{array}$	3.90 0.410
4977.695	0112	00 <sup>0</sup> 0	446	0.058
5026.34 5105.65	3200 40 <sup>0</sup> 0	0000 <b>00°0</b>	446 446	0.255 0.255

Table 14.  $N_2O$  Band Intensities (Contd)

The intensities of the majority of the bands have been calculated from the experimental values for the entire band systems listed in Table 13. Calculating the intensity **of** an individual band from that of the entire band system involves the population of each energy level, the degeneracy of each level, and the relative transition probabilities.

In some of the band systems, the experimental results indicate that the transition probabilities are the same for all bands within a given system. However, as in the  $12^{0}0$  band system, for example, the transition probabilities are greater for the  $13^{1}0 \leftarrow 01^{1}0$  and  $14^{0}0 \leftarrow 02^{0}0$  bands than for the  $12^{0}0$  band. These differences in transition probabilities were found experimentally by comparing the relative intensities of several isolated lines within the system.

Equations relating the intensities of individual lines to that of the entire band are given in Section 3. These are applicable to the N<sub>2</sub>O bands listed in Table 14 with the exception of two forbidden A  $\leftarrow \Sigma$  bands, 02<sup>2</sup>O and 12<sup>2</sup>O, and those listed in Table 15. The factor ((see Section 3 Eq. (13)) that accounts for the Coriolis vibration-rotation interaction can be assumed to be zero for all bands in Table 14, except for the 03<sup>1</sup>O and 11<sup>1</sup>O bands;  $\zeta$  equals approximately -0.01 and -0.007, respectively, for these bands.

Young (1972) has tabulated the relative intensities of the lines within a given band for which the Coriolis interaction is negligible. A large fraction **of** the intensity of a band system is in the main band; therefore, uncertainties in the relative transition probabilities within a band system do not seriously increase the uncertainty of the main-band intensity calculated on the basis of that of the entire band system. Thus, the percent of uncertainty in the intensity of the main band is approximately the same as that for the entire band system, whereas the uncertainty for the hot bands may be greater.

#### 4.4.3 FORBIDDEN BANDS AND PERTURBED LINES

As a result of Coriolis interactions, transitions with  $l' - l'' = \pm 2$  become allowed. Two  $N_2O$  bands of interest that arise from such "forbidden" transitions are the  $02^20 - 00^{\circ}0$  and  $12^20 - 00^{\circ}0$  bands centered at 1177.750 and 2474.785 cm<sup>-1</sup>, respectively. The relationship between the intensities of the individual lines within a band is different for these bands than'for normal bands. As in the case of CO<sub>2</sub>, the intensities are given by Eq. (15) where  $S_v^{\circ} = 1.36 \times 10^{-5}$  molecules  $^{-1}$  cm<sup>2</sup> cm<sup>-1</sup> for the  $02^{2c}0$  band and  $S_v^{\circ} = 9.5 \times 10^{-6}$  molecules-' cm<sup>2</sup> cm<sup>-1</sup> for the  $12^{2c}0$ band.

A few rotational levels of the  $06^{\circ}0$  vibrational state (centered at J = 46) and the  $06^{\circ}0$  state (centered at J = 28) are perturbed by the corresponding levels of the  $10^{\circ}1$  state to such an extent that the normally very weak lines "borrow" enough

Line	Hand		s <sub>J</sub>
		ν <sub>ο</sub> (cm <sup>-1</sup> )	$\times 10^{-20}$ molecules <sup>-1</sup> cm <sup>2</sup> cm <sup>-1</sup>
R45	$10^{0}1$	1)3508.030	0.207
R45	$06^{0}0$	2)3508.186	0.046
P47	$\begin{array}{c} 10 \\ 0 \\ 06 \\ 0 \end{array}$	113430.230	0.136
P47		<b>2)3430.400</b>	0.031
R27	1 0 0 1	1)3500.040	2.11
R27	<b>06<sup>2</sup>0</b>	2)3499.970	0.0233
P29	1001	113452.330	1.71
P29	<b>06<sup>2</sup>0</b>	<b>2)3452.260</b>	0.0188
R45	1001-1000	<b>3)2226.713</b>	0.0118
P47	10 <sup>0</sup> 1← 1000	312149.246	0.0042

Table 15. Parameters of Perturbed Lines of N20

1) The perturbed line positions of the **10<sup>0</sup>1** band are from observed values by Tidwell et al, (1960).

2) The calculated positions of perturbed lines of the  $06^{\circ}0$  and  $06^{2}0$  bands were determined from known perturbed positions of the  $10^{\circ}1$  lines and Toth's (1971a) values of the line separations.

3) The calculated positions of the perturbed 10°1 ← 10°0 lines were determined from Toth's (1971a) results and the positions of the unperturbed lines calculated on the basis of the constants in Table 12.

4) The perturbed line intensities of the 06°0 and 06<sup>2</sup>0 bands are based on the intensities of the unperturbed lines of the 10°1 band and Toth's (1971a) results. The unperturbed line intensities of the 10°1 and 10°1 ← 10°0 bands are based on the band intensities given. in Table 13 and Young's (1972) relative line intensities.

intensity from the  $10^{\circ}1$  lines to appear, at the same time shifting the positions of the lines related to these rotational levels (**Toth,1971a**). The resulting positions and intensities of the affected lines have been calculated separately and are given in Table 15. The R27 and R29 lines of the  $10^{\circ}1 - 10^{\circ}0$  band are not listed in *the* table because the changes in their intensities and positions are negligible.

# 4.4.4 LINE HALF-WIDTHS

Several workers have measured the half-widths of  $N_2O$  lines broadened by  $N_2$ , which is a good approximation to broadening by air. Toth (1971b) has recently measured the widths of lines in the R branches of the  $10^{\circ}1$  and  $02^{\circ}1$  bands directly from observed spectra with a small correction for **the** effect of **the** spectrometer slitwidth. Margolis (1972) has used the same method to measure the lines in both the P- and R-branches of the  $00^{\circ}2$  band. The results of these two workers are in

good agreement, indicating that there is little, if any, dependence on vibrational mode. Lowder's (1972) recent measurements on the  $00^{\circ}1$  band show approximately the same dependence on J as do the results of Toth and Margolis. although Lowder's values averaged approximately 10 percent less. Oppenheim and Goldman (1971) have measured half -widths indirectly by applying band models to low -resolution spectra of a wide variety of N<sub>2</sub>O samples. Their results are in fair agreement with those of Margolis and Toth. Fraley et al, (1962) reported an average value of 0.05 cm<sup>-1</sup> a t m<sup>-1</sup> for lines in the  $00^{\circ}1$  band; this value is considerably lower than those reported by the workers mentioned above.

We have adopted half-widths of  $N_2$ -broadened lines reported by Toth (1971b) to be used for atmospheric  $N_2O$  lines near 296°K. A few values are listed in the tabulation below; values for other J's can be found by interpolation and for other temperatures by assuming the 'half-widths are inversely proportional to the square root of temperature.

J	o cm <sup>-1</sup> atm <sup>-1</sup>
1	0.0915
4	0.0948
10	0.0892
20	0.0812
30	0.0767
40	0.0748
50	0.0698
60	0.0604

#### 4.5 Carbon Monoxide

For the CO bands, we have depended heavily on the work of Young (1968). Table 16 provides the rotational constants which were used to compute the line positions of the main isotope. The positions of the less abundant isotopes were computed from the appropriate constants for the 26 isotope by applying **the** ratio **of** reduced masses. The accuracy of the line positions in the main isotope is expected to be about + 0.001 **cm<sup>-1</sup>**. **Theaccuracy** of the line positions of the long the other isotopes will be somewhat less, about **+** 0.01 cm<sup>-1</sup>.

The intensities for the fundamental and first two overtone bands were taken from Young and are reproduced in Table 17. The intensities for the isotopic bands were taken to be the same as for the 26 isotope reduced by the relative abundances given in Table 4 and the square of the frequency. The intensity of the fundamental appears to be known very accurately, perhaps within t 2 percent; the first overtone is less accurately known to  $\pm$  10 percent; and the second overtone is known to  $\pm$  4 percent (see Young and **Eachus**, 1966); **Burch** and **Gryvnak** (1967). Half-width values have been added to the compilation following the work of Hunt et'al, **(1968).** 

Symbol	Value
ω <sub>e</sub> χ <sub>e</sub> ω <sub>e</sub> χ <sub>e</sub> Β <sub>e</sub> α <sub>e</sub> γ <sub>e</sub> D <sub>c</sub> β <sub>e</sub>	<b>2169.836</b> 13.295 0.0115 1.931285 0.017535 J.01x10-5 <b>6.12x10<sup>-6</sup></b> 1.0x10-9

Table 16. Rotational Constants Used in the Calculation of CO Line Positions

Table 17. Intensities of CO Bands

	1-о	2-0	3-0
Intensity, S(cm <sup>-1</sup> /molecule-cm <sup>-2</sup> )	9.70x10 <sup>-18</sup>	<b>6.99x10</b> <sup>-20</sup>	<b>4.83x10</b> <sup>-22</sup>

# 4.6 Methane

This molecule is a spherical top, in the rigid rotor approximation. It has tetrahedral  $\{T_d\}$  symmetry in the rest configuration of its nuclei. **Rotation**-vibration interactions are complicated even for its simplest infrared-active bands. Starting from the molecular Hamiltonian and the electric dipole-moment operator, transition energies, selection rules, and intensities may be calculated. However, the techniques and the results are complex, and it is not convenient to summarize them here. Therefore, the user is referred to the original papers on this subject John (1968), Schaffer et al, (1939), Hecht (1960), Moret-Bailly (1961), Herranz (1961), Fox (1962), Dang-Nhu (1969), and Susskind (1972).

For the purpose of the present data compilation, we have established the following conventions in notation. The isotopic species  ${}^{12}CH_4$ ,  ${}^{13}CH_4$ , and  ${}^{12}CH_3D$  are denoted by the digits 211, 311, and 212, respectively, in columns **75** through **77**. The quantum numbers for the initial and final states of a transition are specified in columns 36 through 69 as **follows:** 

except for type E symmetry which has for C the format 1X 1A1 instead of 1A1 111. The initial and final state vibrational and rotational quantum numbers have their usual meanings. To avoid ambiguity, we establish some further conventions for the present data compilation. The selection rule C' - C'' on T<sub>d</sub> symmetry is always **taken** to be  $A_1 - A_2, A_2 - A_1, E - E, F_1 - F_2$ , or  $F_2 - F_1$ . The numbering scheme for the N's

begins with 1, 2, 3 . . . Whenever possible, N increases with increasing groundstate energy; the stronger lines correspond to the **selection** rule, N' - N'' = 0.

This section describes methane spectral data in the 3 to 8.5  $\mu$ m region. The following bands, as summarized in Table 18, have been included:  $\nu$  2,  $\nu$  3,  $\nu$  4,  $\nu$  2+ v<sub>4</sub>, 2v<sub>4</sub>.

Isotope	Band	Center (cm <sup>-1</sup> )	<b>Upper</b> State	Lower State	Intensity (cm <sup>-1</sup> /mole-cm <sup>-2</sup> )
311	ν <sub>4</sub>	1297.88 <sup>a</sup>	00000111	00000000	6.59x10 -20h
211	"4	1305.9138 <sup>b</sup>	00000111	00000000	5.87x10 <sup>-18<sup>i</sup></sup>
211	'2	1533.289'	01100001	00000000	8.91x10 <sup>-20<sup>i</sup></sup>
311	ν <sub>3</sub>	3009.53 <sup>d</sup>	00011001	00000000	1.36x10 <sup>-19h</sup>
211	" 3	3018.9205 <sup>e</sup>	00011001	00000000	$1.21 \times 10^{-171}$
211	2V4	2600 <sup>f</sup>	00000222	00000000	1x10 <sup>-19j</sup>
211	v 2+ v4	2818, <b>2838<sup>g</sup></b>	01100112	00000000	8x10 <sup>-19h</sup>

Table 18. Methane Bands Included in Data Compilation

a. Kyle et **al**, (1970)

b. Michelot and Fox (1973)

c. Dang-Nhu (1968) d. McDowell **(1966)** 

e. Bobin and Fox (1973)

f. Fox (1973)

Benedict (1973)

h. 1/89 times value for corresponding band of 211 isotope
i. Armstrong and Walsh (1960)
j. 1/100 times value for v 3 of 211 isotope
k. 1/15 times value for v 3 of 211 isotope

It is planned to add other weak but significant bands in this spectral region at a later time.

Many line positions, together with their absolute intensities and widths, have been included in the present data compilation, even though the corresponding quantum number assignments are uncertain or unknown at this time. It must be stressed that much of the needed theoretical and experimental research on spectra of methane is still incomplete. Consequently, a large portion of the data compilation should be considered tentative and subject to revision at a later time.

#### 4.6.1 **LINE** POSITIONS

The band  $\nu_3$  at 3.3  $\mu$ m is the strongest infrared-active vibration-rotation fundamental of CH<sub>4</sub>. High-resolution measurements of line positions for <sup>12</sup>CH<sub>4</sub> in the range 2884 to 3141 cm<sup>-1</sup> have been used. See Henryet al, (1970), and Barnes et al, (1972). Quantum number assignments for the allowed lines from 2840 to 3167 c m<sup>-1</sup> were made by **Bobin** and Fox (1973) on the basis of fourth-order perturbation theory. Some forbidden lines were assigned by Barnes et al, (1972) using third order theory. Spectroscopic parameters determined for the ground-state and for  $\nu_3$  of CH<sub>4</sub> are listed in Tables 19 and 20, respectively. The corresponding band of the isotopic species <sup>13</sup>CH<sub>4</sub> has been measured from 2873 to 3123 cm<sup>-1</sup>. The **assignments** of McDowell (1966) with some modifications were used. The corresponding spectroscopic parameters are listed in Table 23.

Table	19.	Spectroscopic	Parameters	for	Ground	State	of	<sup>12</sup> CH
-------	-----	---------------	------------	-----	--------	-------	----	------------------

Parameter	Value (cm <sup>-1</sup> )
B	$5.24059\pm0.00006^{a}$
D <sub>s</sub>	(1.086±0.003)x10 <sup>-4a</sup>
D <sub>t</sub>	(4.403±0.099)x10 <sup>-6b</sup>

a. Barnes et al, (1972)

b. **Ozier** et al, (1970). The error bar is derived from the values given by Barnes et al, (1972) and Husson and Dang **Nhu** (1971).

The band at 7.7  $\mu$ m is the next strongest fundamental of  ${}^{12}$ CH<sub>4</sub>. High resolution measurements in the 1225 to 1393 cm<sup>-1</sup> range made by Botineau (1972) were used to obtain spectral line positions. Quantum-number assignments for the allowed lines, to J=12 for the R and Q branches, and J=13 for the P branch, were made by Michelot and Fox, (1973) on the basis of a fourth order calculation. For J=13 to 15 in the Q branch only, and for some J=13 R-branch lines, quantumnumber assignments were taken from the work of Husson and Poussigue (1971). No forbidden lines of this band have been included in the data compilation yet. The determined spectroscopic parameters are given in Table 21. For  $\nu_4$  of  ${}^{13}$ CH<sub>4</sub>, the measurements of Kyle et al, (1970), extend from 1226 to 1359 cm<sup>-1</sup>, and possibly further. The assignments of Kyle et al, (1970) with some modifications were used; spectroscopic parameters appear in Table 24.

Parameter	Value (cm-')	Parameter	Value (cm")
m n <b>p</b> q s t X V W	3018.9205 9.87157 -4.0622x10 <sup>-2</sup> -2.959x10-4 4.329x10-5 1.942x10-7 -9.11x10 <sup>-8</sup> -4.7921x10-2 3.622x10 <sup>-5</sup>	ghk Lj 2''	4.265x10-3 8.365x10-5 -1.925x10-5 2.509x10-7 5.84x10-8 2.605x10-5 -5.39x10-8 1.49x10-6 7.08x10-8

Table 20. Spectroscopic Parameters\* for  $\nu_3$  of  ${}^{12}CH_4$ 

**\*Bobin** and Fox (1973)

Table 21. Spectroscopic Parameters\* for  $\nu_4$  of  ${}^{12}CH_A$ 

Parameter	Value (cm <sup>-1</sup> )	Parameter	Value (cm-')
m p g s t X V w	1305.9138 5.57 52 -6.4192x10 <sup>-2</sup> 1.91x10-3 J.479x10-6 -1.11x10 <sup>-6</sup> -1.79x10-8 -5.2263x10 <sup>-2</sup> 2.319x10-5	g h k j u z z' z''	$\begin{array}{r} 1.8643 \times 10^{-2} \\ 6.590 \times 10^{-4} \\ 2.88 \times 10^{-6} \\ 1.67 \times 10^{-7} \\ -3.78 \times 10^{-9} \\ 2.077 \times 10^{-5} \\ -1.18 \times 10^{-8} \\ -1.901 \times 10^{-6} \\ 8.28 \times 10^{-8} \end{array}$

\*Michelot and Fox (1973)

The  $\nu_2$  fundamental at 6.5  $\mu$ m is relatively weakly infrared-active, by virtue its **Coriolis** interaction with  $\nu_4$ . Quantum-number assignments are based on the **results** of **Dang** Nhu (1969) who developed a fourth-order theory and applied it to **unpublished** spectra of moderately high resolution. Spectroscopic **parameters** are listed in Table 22.

High-resolution **spectra** of  $v_2 + v_4$  and  $2v_4$  are available in the region of approximately 2450 to 3200 cm<sup>-1</sup> (Plyler et al, 1960; and Hall, 1973). Quantum-number assignments for  $2v_4$  have been made on the basis of a third-order theoretical analysis. A partial analysis of  $v_2 + v_4$  has been **done** in **analogy** with the quantum-number assignments in the work of (Bregier, 1970; and Hilico, 1970), on  $v_2 + v_3$ . Spectroscopic parameters for  $v_2 + v_4$  and  $2v_4$  are given in **Tables 25** and 26 respectively.

Parameter	Value (cm <sup>-1</sup> )
v2 ; B2 B0 d2 d0 z2 f2 f2 e2 e0	1533.289 5.319 5.230 5.7x10-5 2.1x10-5 2.9x10-2 -5.2x10-3 -2.8x10-5 -5.0x10-6

Table 22. Spectroscopic Parameters\* for  $\nu_2$  of  ${}^{12}CH_4$ 

**\*Dang Nhu** (1968)

Parameter	Value (cm <sup>-1</sup> )
ν <sub>0</sub> -2(Βζ <sub>3</sub> )eff	3009.05+0.03
$B_{o}^{+}B_{eff}^{P,Q}$ -2(B $\zeta_{3})_{eff}$	9.963+0.004
$B_{eff}^{P,R}$ - $B_0$	-0.0386 <u>+</u> 0.0008
2(D <sub>0</sub> +D <sub>1</sub> )	$(4.2+0.4) \times 10^{-4}$
D <sub>o</sub> -D <sub>1</sub>	$(2.1\pm0.5)\times10^{-5}$
<b>ö</b> 3	0.046t0.005
${}_{\mathrm{D}}{}^{\mathrm{J}}{}^{\boldsymbol{ au}}$	(4.7+2.8)x10 <sup>-5</sup>
d <sup>J</sup> T	(5.4+1.7)x10 <sup>-5</sup>

Table 23. Spectroscopic Parameters\* for  $\nu_3$  of  $^{13}\text{CH}_4$ 

\*McDowell (1966)

**Table** 24. Spectroscopic Parameters<sup>\*</sup> for  $\boldsymbol{\nu}_4$  of  ${}^{13}CH_4$ 

Parameter	Value (cm <sup>-1</sup> )
ν4 Β' D ξ F4s to44 t134 t224	$1297.88\pm0.02 \\ 5.179\pm0.005 \\ 5.194\pm0.005 \\ (1.16\pm0.02)\times10^{-4} \\ 0.477\pm0.005 \\ (2.26\pm0.02)\times10^{-3} \\ -4.5\times10^{-6} \\ (4.84\pm0.02)\times10^{-4} \\ (-1.46\pm0.01)\times10^{-2} \\ -2.65\times10^{-6} \\ (-1.46\pm0.01)\times10^{-2} \\ (-1.46\pm0.01)\times10^$

\*Kyle et al, (1970)

Table 25. Spectroscopic Parameters\* for  $\boldsymbol{\nu}$  2t  $\boldsymbol{\nu}_4$  of  $^{12}CH_4$ 

Parameter	Value (cm")
<b>v</b> <sub>0</sub> (Q)	2818, 2838

**\*Hiliev** (1970) - Further analysis, including that of the interaction between  $\mathbf{F}_1$  and F2 substates, is necessary in **order** to obtain more complete values of spectroscopic parameters.

Table 26. Spectroscopic Parameters\* for 2  $\nu_4$  of  ${}^{12}CH_4$ 

Parameter	Value (cm-' )
е	2600.29
В	5.175
В	2.520
С	1.0x10-4
44	3.1
у	<b>1.837x10<sup>-2</sup></b>
ћ	6.579x10 <sup>-4</sup>

\*Fox (1962)

#### 4.6.2 LINE INTENSITIES

Although it is recognized that it would be desirable to include in the data compilation all methane lines whose absolute intensity exceeds the cutoff specified in Table 3, it has not been possible to achieve this goal because of present experimental and theoretical limitations.

Absolute intensities for the allowed lines in  $\nu_3$  of  ${}^{12}CH_4$  were calculated, starting from the measured values for R(0); R(1), and R(2). The relative intensities of forbidden lines (Barnes et al, 1972) were used for calculating their absolute intensities. The absolute line intensities for  $\nu_3$  of  ${}^{13}CH_4$  were calculated on the basis of those for  $\nu_3$  of  ${}^{12}CH_4$ , together with the terrestrial  ${}^{12}C/{}^{13}C$  ratio (see Table 4). Calculated and/or measured absolute band intensities are listed in Table 18.

Absolute line intensities for  $\nu_4$  have been calculated on the basis of the measured values for R(0), R(1), and R(2). For <sup>13</sup>CH<sub>4</sub>, absolute intensities were calculated as for  $\nu_3$ . Absolute band intensities are given in Table 18.

Absolute line intensities for  $\nu_2$ , have been calculated on the basis of theoretical relative-line intensities and the experimental absolute band intensity in Table 18.

Absolute -line intensities for  $\nu_2 + \nu_4$  were determined from the solar spectra of Hall (1972), with an assumed  $CH_4$  air mass of approximately  $4 \times 10^{19}$  molecules/cm<sup>2</sup> and an assumed temperature of  $273^{\circ}K$  and a pressure of 0.65 atm.

Absolute line intensities for  $2\nu_4$  were calculated from the relative intensity formula of Fox (1962) together with the absolute band intensity in Table 18. The latter intensity was taken to be approximately 1 /100 of that for  $\nu_3$  of  ${}^{12}CH_4$ .

#### 4.6.3 LINE WIDTHS

The J- and T-dependence of methane line widths is taken from the results of Varanasi et al, (1971, 1972) and Tejwani and Varanasi (1971). The only measured values of air -broadened  $CH_4$  half-widths are for multiples of  $\nu_3$  and  $2\nu_3$  at T= 295°K. For J≈1 to 5,  $\gamma^{\circ} \approx 0.061$  cm<sup>-1</sup> atm<sup>-1</sup>; for J≈15 to 17,  $\gamma^{\circ} \approx 0.048$  cm<sup>-1</sup> atm<sup>-1</sup>. These results agree with values calculated from Y ° for  $CH_4$ -0<sub>2</sub>, assuming N<sub>2</sub> and 0<sub>2</sub> with their partial pressures in air. Measurements and calculations of Yo for  $CH_4$ -H<sub>2</sub> in  $\nu_4$  suggest, but do not prove directly, that Y ° may be independent of vibration-rotation band. For the purpose of the present data compilation, we adopt the value  $\gamma^{\circ}$ =0.055 cm<sup>-1</sup> atm<sup>-1</sup> for all  $CH_4$  lines. There are no results available for the T-dependence of  $\gamma^{\circ}$  for  $CH_4$ -Air or even  $CH_4$ -N<sub>2</sub> or  $CH_4$ -0<sub>2</sub> broadening. From the calculations and measurements for  $CH_4$ -H<sub>2</sub> broadening with T, we assume the usual T<sup>-1/2</sup> dependence. In conclusion, for the present data compilation, we

$$\gamma^{0} = 0.055 \left(\frac{296}{T}\right)^{1/2} \left(\frac{p}{1 \text{ atm}}\right). \tag{16}$$

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### 4.7 Oxygen

The infrared atmospheric bands of oxygen have been interpreted by Van **Vleck** as magnetic dipole transitions between intersystem electronic combinations, a singlet-triplet transition  $\mathbf{a}^{1} \Delta_{g} - \mathbf{X}^{3} \Sigma_{g}$ . The A v=0 (0,0) band at 7882.39 cm<sup>-1</sup> (1.27  $\mu$ m) has an Einstein A coefficient of 1.9x10<sup>-4</sup> sec<sup>-1</sup> Jones and Harrison (1958). The (0,0) band, and the weaker (1,0) band at 9365.89 cm<sup>-1</sup> have been observed in terrestial atmospheric absorption by Herzberg and Herzberg (1947). The (0,1) band at 6325.99 cm<sup>-1</sup> has been observed in the twilight **airglow** emission by Jones and Harrison (1958). In addition to these bands there is an electric quadrupole  $(b_{\Sigma^{+}\sigma^{-a}}^{1}\Delta_{\sigma})$  transition at 1.908  $\mu m$  that has been detected in the emission spectrum of a discharge through 0, and helium (Noxon, 1961). Finally, submillimeterwave rotational transitions (McKnight and Gordy, 1968) and spin reorientation spectra (Zimmerer and Mizushima. 1961) at 60 GHz have been observed in the ground electronic state. The (a-X) system and the microwave spectra are discussed below. The atmospheric "A" band at 7619Å, representing the magnetic dipole transition  $b^{1}\Sigma_{g}^{+} \leftarrow X^{3}\Sigma_{g}^{-}$ , is also included (Wark and Mercer, 1965; Burch and Gryunak, 1967).

With the molecular rotational momentum designated by  $\mathbf{K}$  and the electronic spin vector by  $\mathbf{S}$ , the total angular momentum  $\mathbf{J}$  is given by

# J=K+S.

Thus, for the singlet-delta state J=K, and the energy levels, apart from the vibration, are given by

$$F = B_{v}K(K+1) - D_{v}K^{2}(K+1)^{2}, \qquad (17)$$

where  $\mathbf{B}_{\mathbf{v}}$  is the rotational constant and  $\mathbf{D}_{\mathbf{v}}$  is the first order centrifugal distortion constant for the particular vibrational state. The constants for this state, obtained from measurements of the infrared bands, are listed in Table 27.

The expressions for the rotational energies **of** the rho-type triplet are obtained from a solution of the Hamiltonian (West and Mizushima, 1966):

$$H = B \kappa^{2} + \frac{2}{3} \lambda (3S_{z}^{2} - S^{2}) + \mu \kappa \cdot S, \qquad (18)$$

where A is the spin-spin interaction coupling constant and  $\mu$  is the spin-rotation interaction coupling constant.

1.2

9	
Constant	Value (in cm <sup>-1</sup> )
$B_{0} \\ B_{1} \\ D_{0} \\ \omega_{e} \\ \omega_{e} \\ x_{e} \\ \Delta G \\ \nu_{e} $ 1/2	1.41783 1.4007 <b>4.86x10-6</b> 1509.3 12.9 1483.50 7918.134

Table 27. Constants of the  ${}^{1}\Delta_{g}$  State of  $0_{2}$  (Herzberg and Herzberg, 1947)

Centrifugal stretching effects are approximately corrected for by assuming a K(K+1) dependence for B. A and  $\mu$ , namely:

$$B = B_{v} - D_{v}K(K+1) + H_{v}K^{2}(K+1)^{2},$$
  

$$\lambda = \lambda_{0} - \lambda_{1}K(K+1), \text{ and}$$
  

$$\mu = \mu_{0} - \mu_{1}K(K+1).$$
(19)

For each value of the quantum number at end-over-end rotation, **K**, the triplet energies are given by  $F_1(K)$ ,  $F_2(K)$ , and  $F_3(K)$  where J = K+1, J=K and J=K-1, respectively. Both  $F_1(K)$  and  $F_3(K)$  average about 2 cm<sup>-1</sup> lower than the  $F_2(K)$  component. Values for the constants of the triplet-sigma state, obtained from both infrared and microwave measurements, are given in Table 28.

The band centers are given by

$$G(\mathbf{v}',\mathbf{v}'') = \mathbf{v}_{e} + \omega_{e}^{\prime}(\mathbf{v}'+1/2) - \omega_{e}^{\prime}\mathbf{x}_{e}^{\prime}(\mathbf{v}'+1/2)^{2} + \cdots - \omega_{e}^{\prime\prime}(\mathbf{v}''+1/2) + \omega_{e}^{\prime\prime}\mathbf{x}_{e}^{\prime\prime}(\mathbf{v}''+1/2)^{2} - \omega_{e}^{\prime\prime}\mathbf{y}_{e}^{\prime\prime}(\mathbf{v}''+1/2)^{3} + \omega_{e}^{\prime\prime}\mathbf{z}_{e}^{\prime\prime}(\mathbf{v}''+1/2)^{4} + \cdots, \qquad (20)$$

where the prime and double primes refer to the singlet and triplet states respectively.

The selection rules for magnetic dipole radiation are

 $\mathbf{\Delta J} = \mathbf{0}, \ \mathbf{+} \ \mathbf{1}$ 

and

+ ⊷ +, - ⊷ -, g ⊷ g, u- u.

Constant	Reference	Value (cm <sup>-1</sup> )
$B_{0}^{B_{1}}$ $D_{0}^{O_{1}}$ $H_{0}^{O_{0}}$ $\lambda_{1}$ $\mu_{0}^{U_{1}}$ $\omega_{e} x_{e}$ $\omega_{e} y_{e}$ $\omega_{e} z_{e}$ $\Delta G_{1}/2$	a b b b b c c c c c b b b b b b	1.4376809 1.421979 4.913x10-6 <b>4.825x10<sup>-6</sup></b> <b>3.0x10<sup>-10</sup></b> 1.9847530 -1.950x10-6 <b>-8.42930x10<sup>-3</sup></b> 8.01x10-9 1580.3613 12.0730 <b>5.46x10<sup>-2</sup></b> -1.43x10-3 1556.3856

Table 28. Constants for the $^3\Sigma$	State of <b>0</b> 2
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a. McKnight and Gordy (1968) b. Babcock and Herzberg (1948)

c. West and Mizushima (1966)

The possible branches and the nomenclature used here for the ' $A_{r} - {}^{3}\Sigma_{g}$ ' transition are shown in Table 27. Since the oxygen-16 nuclei obey Bose-Einstein statistics, only the positive rotational levels occur. In the ground state this causes an alternation of lines such that only odd values of the rotational quantum number K are present. For the heteronuclear isotopes of oxygen, however, all rotational levels occur; the  $0^{17}0^{17}$  molecule has alternate rotational levels with nuclear statistical weights of 5 and 7 (Gordy ct al, 1953). In  $0^{16}0^{16}$  (and  $0^{18}0^{18}$ ) there are four branches with  $\Delta K$ =+1, three with  $\Delta K$ =0, and two with  $\Delta K$ =+2, the latter branches being designated by superscripts  $\mathbf{S}$  and  $\mathbf{0}$ . On the magnetic tape the six quantum numbers of the transition, v', J', K', v'', J'', K'', are given in the field of columns 38 through 62. In addition. the nine branches are listed in columns 64 and 65 with the superscript, representing the AK of the transition, preceding the AJ designation. A diagram indicating the transitions involved in the  ${}^{1}\Delta_{g} + {}^{3}\Sigma_{g}$  band is given in Figure 1.

The above description generally applies to the microwave transitions of pure rotation and transitions between levels of the triplet state. In the latter the labelling is given as K+ and K-, designating the transitions  $F_2(K) - F_1(K)$  and  $F_2(K) - F_3(K)$  respectively. Intensities have been calculated (see Townes and Schawlow, 1955). A good summary of the various bands of oxygen molecule can he found in the recent review article by Krupenie, 1972.



Figure 1. Transitions in the  ${}^{1}\Delta_{g} - {}^{3}\Sigma_{g}$  Band of Oxygen

# 5. CONTINUOUS ABSORPTION BY ATMOSPHERIC GASES

Although not part of the data tape of primary concern in this report, a few words should be said about the relatively continuous regions of absorption of particular interest in the atmospheric "windows" near 2500 cm<sup>-1</sup>( $4\mu$ m), from approximately 1250 cm<sup>-1</sup> to 700 cm<sup>-1</sup>( $8\mu$ m - 14  $\mu$ m), and near 450 cm<sup>-1</sup> ( $22 \mu$ m) In other spectral regions the contribution by nearby absorption lines is much greater than that by the continuum absorption, so that for practical purposes the continuum effect can be neglected, although it may be greater than in the windows.

This continuous absorption is caused by one or more of the following three processes: (1) extreme wings of strong collision-broadened absorption lines centered more than 10-20 cm<sup>-1</sup> away; (2) pressure induced absorption resulting from transitions that are forbidden for unperturbed molecules; and (3) the possible existence of the water dimer  $(H_20:H_20)$  in the case of the 8 to 14  $\mu$ m region.

The absorption coefficient due to continuum absorption can be expressed as

$$k = C_s P_s + C_b P_b$$
(22)

where  $C_s$  is the self-broadened coefficient and  $C_b$  is the foreign gas broadening coefficient,  $P_s$  is the partial pressure of species, s, and  $P_b$  is the foreign gas pressure.

Figure 3 gives the spectral dependence of  $C_s$  for water vapor absorption in the 8 to 14  $\mu$ m region for three temperatures (Burch, 1970). The  $C_b$  value has been most reliably measured by McCoy et al, 1969, and is found for nitrogen to be  $C_b^{=}$  0.005 $C_s$  at room temperature.

Figure 2 gives the spectral dependence of  $C_s$  for the water vapor absorption in the region near 4  $\mu$ m for four different temperatures (Burch et al, 1971a). Note that the **T=296<sup>o</sup>K** curve is an extrapolation based on the measurements at higher temperature. These same workers found the ratio  $C_b/C_s$  for nitorgen broadening to be 0.12 ± 0.03.

Figure 4 gives the spectral dependence of the absorption coefficient due to the pressure-induced nitrogen absorption centered near 2330 cm<sup>-1</sup> (Burch et al, 1971a) Measurements have also been made by Shapiro and Gush, 1966, and Farmer and Houghton, 1966. Since the foreign gas broadening in this case results from a gas (oxygen) having a constant mixing ratio in the atmosphere, Eq. (22) reduces to

$$K = Const. \ x \ P \tag{23}$$

Since the nitrogen abundance in the atmosphere is also directly proportional to P, the absorption depends on  $P^2$  and Figure 4 has as ordinate the absorption **coefficien** expressed in the units atm<sup>-2</sup> Km<sup>-1</sup>.

Since line wings as given by the Lorentz shape, Eq. (1), have been found to be in error in **the** extreme wings, an appropriate rule to follow for the truncation of



Figure 2. Normalized Continuum Absorption Coefficient for  ${\bf H_2O}$  at Three Temperatures



Figure 3. Normalized Continuum Absorption Coefficient Between 2400 and 2820 cm  $^{-1}$  for Pure  $\rm H_2O$  at Four Temperatures

line wings and the introduction of continuum absorption coefficients in accordance with Figures 2 through 4 is difficult to state. It is recommended that the user familiarize himself with this problem (see, for example, **Burch** et al, 1969) and in any case the use of the Lorentz shape beyond 20 or 30 wavenumbers of line centers is inappropriate (see also discussion on line shape on p. **3**).



Figure 4. Spectral Plot of Absorption Coefficient for Atmospheric  ${\bf N_2}$  at Four Temperatures

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# Appendix A

Computer Program Listing for Reading Tape

```
PROGRAM RU TP( INPUT. OJTPUT. TAPE2)
    DIMENSION R (400) + TT (15)
     A=0
    IEOF=0
 24 BUFFER IN(2+0) (R(1) +R(340))
    IF (UNIT (2)) 25+30+35
 J5 PRINT 30+TT(1)
 36 FORMAT (* PARITY ERROR AFTER *.F12.3)
    GO TO 24
 3 6 IEOF=IEUF+1
    PRINT +1+IEOF
 3 1 FORMAT (# EOF NO#+15)
      IF(IEDF.GT.7) GO TO 1001
      G 0 TO 4
 25 DECODE (10, 39, R(1)) IRED
 41 FORMAT (F10.3+E10.3+F5.3+F10.3+546+A5+I3+I4+I3)
    I=1
  39 FORMAT(110)
    L=2
     0 0 40 K=1, IRED
    DECODE(80,41,Q(L))(IT(I),1=1,13)
    PRINT 41+(TT(1)+1=1+13)
    L=L+d
     IF(TT(1)+LT+A) PRINT 153+A+TT(1)
      A=TT(1)
153 FORMAT(# O U TOF DRDER +,2F12,3)
  40 CONTINUE
    GO TO 24
1001 CALLEXIT
       s TOP
      END
```

## Appendix **B**

### computer Program for Homogeneous Path Transmittance Calculation

The computer program (Program LBL) provided in this Appendix is intended for use as a check to ensure that a user of the AFCRL Data Tape will be using the data correctly. The spectral region chosen for a sample spectrum (Figure 5) contains spectral lines from four different atmospheric gases. The program was constructed for use with constant pressure and temperature paths only, so the user will be left to his own devices for real atmospheric applications. After computing transmittance at closely spaced monochromatic frequencies, a convolution is performed over a triangular slit function whose half-width can be chosen. Comment cards have been used freely, so no further discussion of this program will be given here.



Figure B1

Synthetic Spectrum Generated using the Data Compilation. Conditions are as follows: pressure = 188 mb, T = 219K, W H<sub>2</sub>O = 2.00 x 10<sup>20</sup>, W<sub>CO2</sub> = 2.05 x 10<sup>21</sup>, W<sub>O3</sub> = 3.23 x 10<sup>18</sup>, W<sub>CO</sub> = 4.67 x 10<sup>17</sup>. Lines belorging to other molecular species contained on tape are not present in this spectral region. Spectral resolution is 0.01 cm<sup>-1</sup>.

PROGRAM LBL(INPJT+0JTPUT+TAPE2) DIMENSION W(7), R(325), GNU(3000), S(3000), A\_PHA(3000), EDP(3000) DIMENSION MOL(3000), CAY1(7), OPD(3000), FNU(1000), TRANS(1000) DIMENSION SUM1(7), CS2(7)

PROGRAM JY R ... MCCLATCHEY.

N O PUNCHED DECKS WILL BEDISTRIBUTED.

THIS PROGRAM GENERATES 4 TRANSMITTANCE SPECTRJY WITH OUTPUT RESULTS P RIN TE D EVE R Y DELV WAVENUMBERS BETWEEN THE INITIAL FREQUENCY. VI, AND T H E FINAL FREQUENCY.V2. CALCULATIONS A Q E PERFORMED FOR A UNIFORM, CONSTANT P R E S S U R E. CONSTANT TEMPERATURE PATH CONTAINING ANY D R ALL O F THE MOLECULAR SPECIES DESCRIBED IN THIS Q E P D Q T IN ARBITRARY AMOUNTS. WOLECULAR ABUNDANCES MUST BE SPECIFIED IN THE UNITS (MOLECULES/CM2). MONOCHROMATIC CALCULATIONS ARE MADE AT FREQUENCY INTERVALS.DV. AND A TRIANGULAR SLIT FUNCTION O FHALF-WIDTH, A, I S CONVOLVED WITH THE MONOCHROMATIC RESULTS.

1EOF=0 DEPTH=0.001 PI=3.14159 SUM=020 IV=1

C C

С

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C C

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**CCCC** C

> C C C C

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C

с С

С

C C R E A D INPUT PARAMETERS (P=PRESSURE), (T=TEMPERATURE), W(1)=H20, W(2)=CO2, M(3)=O3, W(4)=N20, W(5)=CO, W(6)=CH4, W(7)=O2. VIAND V2 ARE FREQUENCY LIMITS FOR WHICH OUTPJT R E SULTS A ? E REQJIRED. DV I S MONOCHROMATIC FREQUENCY INCREMENT. BOUNDIS THE FREQUENCY FROM 4 N YLINE CENTER BEMOND WHICH THELINE WILL BE NEGLECTED. A I STHE HALF-WIDTH OF 4 TRIANGULAR SLIT FUNCTION. DELVIS FREQUENCY IYCREVENT OF CONVOLVED OUTPJT TRANSMITTANCE RESULTS.

READ 77+P+T PRINT 77, P.T READ 81+ (W(M)+M=1+7) PRINT 83 PRINT RL, (W(4),4=1,7) READ 85+ VI+V2+DV+BOUND+A+DELV PRINT 87+ V1+V2+DV+30UND+A+DELV IF (A+2/UV+1.GT. 3000) CALCULATION CANNOT BE DONE IF THERE A R E MORE THAN 3000 LINES READ FROM TAPE I V 4 FREQUENCY RANGE 0 F 2(A+JOUND) CALCULATION CANNOT 3E DONE VBOT=VI-A-dOUND VTOP=V2+A+BOUND MFILE=0 I F (VBOT.GE.500.0) MF ILE=1 I F tVdor.GE.1000.0) MFILE=2 IF tVHoI.GE.2000.0) MFILE=3 I F (VB0F.GE.5000.0) WFILE=4

```
METLE DETERMINES THE NUMBER OF ENDS OF FILE TO BE READ OVER
С
      BEFORE COMING TO THE PERTINENT DATA ON THE TAPE.
С
e
C
      WE ARENOW READY TO READ TAPE.
      I=1
      ILL=1
      BUFFER IN (2+0) (R(1)+R(325))
1
      IF (UNTT(2)) 7,5,3
3
      PRINTHY GNU(I)
      GO TO I
      IEOF=IEUF+1
5
      PRINT 91+ IEOF
      IF (IEOF.GE.7) GO TO 75
      G o TO
      IF (MFILE.GT.IEOF) SO TO 1
7
      DECODE (10,93,R(1) ) IREC
      NT= (8+12EC) -6
      DECODE (10,95,R(NT) )TMAX
      IF (TMAA.LT.VBOT) GO TO 1
      DO 9 K=1+IREC
      DECODE (B0,95.R(L))GNU(I) +S(I) +ALPHA (I) +EDP(I) +IDAT+ISOT+MOL(I)
      し=し+め
      IF (GNU(I).LT.VEDT) GO TO 9
      M=MOL([)
       PATH=S(1)+W(M)/(PI+0.06+P/1013.0)
      IF (PATH-LT-DEPTH) 30 TO 9
       IF (GNU(I).GT.VTOP) GO TO LI
       I=I+1
9
       CONTINUE
       I F (1.GT.2960) 33 T3 11
       GO TO 1
       I1=I
11
       PRINT 97, VHOT, VTOP, GNU( 11), 11
       15=1
       V2P=GNU(I1)-BOUND-A
С
       TAPE HAS BEEN READFOR ALL NECESSARY LINES OR FOR THE MAXIMUMNO.
OF LINES POSSIBLE SUBJECT TO RECYCLING.
С
С
       HALFWIDTHS WILL 3E SUPPLIED BELOW WHEN THEY DO NOT APPEAR
С
       ON TAPE.
С
С
       00 15 [=]LL+I1
       M=MOL(I)
       IF (M.E.).1) GO TO15
       IF (ALPHA(I).GT.0.0) GO TO 13
       IF (M.E3.2) ALPHA(1)=0.07
       IF (4.E2.3) ALP-44(1)=0.11
       IF (M.E3.4) ALPHA(I)=0.08
       IF (M.E2.5) ALPHA(I)=0.06
       IF (M.E3.6) ALPHA(I)=0.055
       IF (M.FQ.7) ALPHA(I)=0.048
IF (ALPHA(I).LT.0.01.0R.ALPHA(I).GT.1.0) ALPHA(I)=0.06
 13
```

74

```
15
      CONTINUE
IS=1
      P0=1013.00
      T0=296.00
      CS1 = (T0 - T) / (T0 + T + 0.5946)
С
С
      ROTATIONAL PARTITION FUNCTION IS DEFINED BELOW
C
      D o 214=1,7
      IF (M.EQ.1) GO TO 17
IF (M.EQ.2) GO TO 19
     , IF (M.EQ.3) GO TO 17
      IF (M.EQ.4) GO TO 19
      IF 14+EQ+5) GO TO19
      IF (M.E2.6) GO TO 17
      If (M.EQ.7) GO TO19
      CS2(4) = ((T0/T) + 1.5)
17
      Go TO 2A
      CS2(4)=F0/T
19
      CONTINUE
CA= ((TO/T) ++0.5) + (P/PO)
51
С
      TEMPERATURE DEPENDENCE OF ALL LINE INTENSITIES COMPUTED HERE.
С
С
      00 23 I=ILL,II
      M=MOL(I)
      S(1)=S(1)+CS2(M)+EX>(-EDP(1)+CS1)
      ALPHA(I)=ALPHA(I)+CA
23
      V=V1-A
      DO 27 M=1,7
25
      CAY1(M)=0.0
27
      SUM1 (M) =0.0
      DETERMINE INDICES (IS AND 16) INDICATING WHICH SPECTRALLINES
C
      ARE TO BE USED IN THE CALCULATION AT FREQUENCY V.
С
С
      Do 33 1=15,11
      IF (V-BOUND-GNU(I)) 29,29,33
IS=I
29
      Go TO 35
      CONTINUE
33
      15=II
      Go TO 49
      D o 39 J=15,11
35
      IF (V+90UND-GNU(J)) 37,37,39
      16=J-1
37
      GO TO 43
      CONTINUE
39
      I6=I1
С
C
      COMPUTE THE OPTICAL DEPTH AND TRANSMITTANCE AT FREQUENCY V.
С
43
      00 45 I=15,16
      M=MOL(I)
```

4

Z=ABS {V=GNU(I)) SUM1 (M) = S(I) + ALPHA(I) / (Z++2+ALPHA(I) ++2) CAY1 (M) = CAY1 (M) + SUM1 (M) CONTINUE 45 CAY=0.0 Do 47 M≠1,7 CAY=CAY+CAY1(M)+W(M)47 OPD(IV) =CAY+0.3183 GO TO 51 OPD(IV)=0.0 49 51 OPD(IV) = EXP(-OPO(IV))IF ((V+QV).6T.V22) 30 TO 53 IF (V.GE.V2+A) 30 TO 53 IF (IV.GE.3000)30 TO 53 IV=IV+1 V=V+DV Go TO ?5 с A T THIS POINT, CYCLE BACK TO STATEMENT 25 AND COMPUTE THE С MONOCHROMATIC TRANSMITTANCE AT V+3V+ETC. С IF STATEMENT 53 IS REACHED . ALL POSSIBLE MONOCHROMATIC TRANSMITTANCE VALUES HAVE BEEN COMPUTED, AND THE SLIT FUNCTION CONVOLUTION WILL! С С NOW BE PERFORMED IN LOOP 57 С С 53 FREQ=VI PRINT 101, IV.V.V2P FINAL=VI+3000.+DV-A-DELV V=V1-A JFNU=1 L=DELV/0V+0.01 IA=1 55 SUM=0+0 Do 57 I=IA+IV SUM=SUM+ (A-ABS (V-FREQ) ) +OPD (1) V=V+DV IF (V-(FREQ+A)) 37,59,59 57 CONTINUE TRANS ( JFNU) = SUM+DV/ (A+A) 59 FNU (JFNJ) =FREQ IF (FRE3.GT.V2) 30 TO 61 IF (FREJ.GT.V2P) GO TO 51 I F (FRE2.GE.FINAL) 30 TO 61 FREQ=FREQ+DELV IF (JFNU.GE.1000) GO TO 61 JENU=JENU+1 IA=IA+L V=FREQ-A SUM=0%0 Go TO 5> С CONVOLVED TRANSMITTANCE RESULTS ARE NOW PRINTED OUT. С С PRINT 103. JENU 51 PRINT 104

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PR.NT 105, (FNU(J) , TRANS(J) , J=1, JFNU) I F (FRED.GE.V2) 30 TO 75 IF (FREQ.GT.V2P) GO TO 57 IF \$JFNU.GE.1000) GD TO 65 IF (FREQ.GE.FINAL) 30 TD 6 3 Go TO 75 V1=FINAL+DELV 53 15=1 Iv=l JFNU=1 V=V1-A GO 1025 IA=IA+L 65 С IF STATEMENT 65 IS REACHED. ADDITIONAL MONOCHROMATIC CALCULATIONS С ARE REGUIRED TO SATISFY THE TOTAL FREQUENCY RANGE OVER WHICH С CONVOLVED RESULTS ARE REQUIRED. С С JFNU=1 V=FREQ-A Go To 52 IV=1 67 С IF STATEMENT 67 IS REACHED, THE DATA FROM THE DATA TAPE WILL BE С REORGANIZED AND THE TAPE WILL BE READ AGAIN. С С JFNU=1 V1=FRED VBOT=VI-A-BOUND D 0 6 7 [ N=1 + 11 IF (GNU(IN).GT.V30T) GO TO 71 CONTINUE 69 IN=I1 71 IJ=IN L=1 DO 73 [=IJ+I1 GNU(L)=GNU(I) S(L) = S(1)ALPHA(L)=ALPHA(I) EDP(L) = EDP(I)MOL(L)=MOL(I) L=L+1 73 I =L ILL=L GO TOL CALL EXIT 75 STOP С 77 FORMAT (E12.5.F7.2) FORMAT (\* PRESSURE =\*,E12.5,\*TEMPERATURE =\*,F7.2) 79 FORMAT (7E10.3) **51** FORMAT (3x,+WATER+,6x,+CO2+,6x,+OZONE+,7X,+N20+,7X,+CO+,8X,+CH4+,7 93 1X,#02#,4X) 85 ' FORMAT (6F10.3)

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37 FORMAT (\* VA =\*+,F10.3,\*V2 =+,F10.3,\*DV=+,F10. 3+\*BOUND \*\*+,F10.3+\*A I=+,F10.3.+DELV =+.F10.3) FORMAT(\* PARITY ERROR ENCOUNTERED AT+,F12.3) FORMAT (\* END OF FILE ENCOUNTERED+.I5) FORMAT (I10) 87 91 93 FORMAT (F10.3.E10.3.F5.3.F10.3.35%, I3.I4.I3) FORMAT (\* VBOT =\*,F12.3.\* VTOP =\*,F12.3.\*GNU =\*,F12.3.\* 11 • 918) 95 97 FORMAT (15,2F10.4) 101 FORMAT (\* JFNU =++I5) FORMAT (5(\* FREQJENCY 103 •)) 104 TRANs. 105 FORMAT 15(F10.30E12.5})

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W. S. Benedict	R. F. <b>Calfee</b>	J. S. Garing					
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		Bedford, M	assachusetts 01730				
S. ABSTRACT							
The report describe	es a compilation	of the molecul	ar spectroscopic parameters				
or a number of infrare	ed-active molecu	les occurring n reincluded in t	aturally in the terrestrial this compilation: water vapor				
u-bon dioxide; ozone;	nitrous oxide; ca	arbon monoxide;	methane; and oxygen. The				
pectral region covered	extends from les 100.000 spectra	al lines. The pa	rameters included in the				
mpilation for each lir	ne are: frequency	y, intensity, hal	f-width, energy of the lower				
nate of the transition, v energy states, an isotop	bic identification,	, and a molecul	ar identification.				
A discussion is pro	wided separately	for each mole	rular enocide indicating the				
Burces and accuracy of	f the data and a	general discuss	ion of how the data were				
otained.							

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Figure 4. Vertical Path to Space From Sea Level for the Midlatitude Winter Model Atmosphere Showing the Separate Contributions to the Total Transmittance