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REMOTE SENSING OF POLLUTANTS

COMPUTERIZED REDUCTION OF LONG PATH ABSORPTION DATA



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REMOTE SENSING OF POLLUTANTS

COMPUTERIZED REDUCTION OF LONG-PATH ABSORPTION DATA

by

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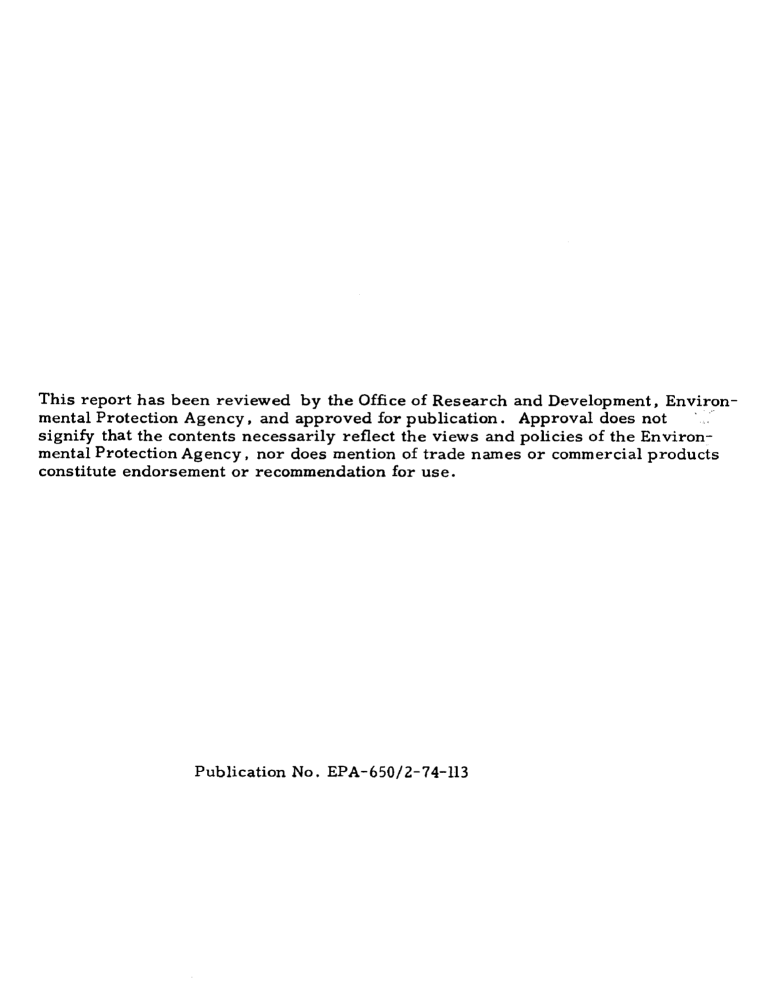
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PREFACE

This report is arranged in a flexible digital format. The sections and subsections are designated by adding a decimal point and a number. For example, 5.1.2 is the second sub-subsection of the first subsection of section 5. References to sections are given without the use of the word section; for example, (7.3) refers the reader to the third subsection of section 7. Equations are invariably and uniquely referred to by a colon notation; that is, (3:7) refers the reader to equation seven of section 3. Figures and tables have a unique reference also; for example, F3.1 and T4.7 are respectively the first figure of section 3 and the seventh table of section 4. Appendixes are indicated by a capital A; that is, A2.1 is the first section of the second appendix. Figures, tables, and equations of appendixes are denoted respectively by FA3.3, TA2.4, and A3:2.

The International System of Units is used throughout except where the traditional unit still holds so strongly that confusion would result from absolute purity. Pollutants for example are often measured in parts per million. Conversion factors are given in A4.

This report is the product of a joint effort by EPA and NOAA. It conforms to EPA format rather than that of NOAA.

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ABSTRACT

Atmospheric gaseous pollutants are very numerous in industrial regions. It is estimated that 25 or more pollutant molecules may be found in the atmosphere in significant quantities. The measurement of the concentration of each gas from the complex spectrum obtained by a long-path infrared spectrophotometer requires the fitting of trial spectra composed from a library of spectra. The fitting procedure adjusts the concentrations of the trial spectra until a "best fit" in a least-squares sense is produced. This report is a description of the physical, mathematical, and calculational principles and procedures for the use of a digital computer program to determine concentrations of atmospheric gases in a path of a few kilometers. Detailed instructions for the computer program and a library of spectra are provided.

1. INTRODUCTION

The Chemistry and Physics Laboratory of the Environmental Protection Agency (EPA) in Raleigh, NC, has been studying the feasibility of various methods of measuring the gaseous pollutants of the earth's atmosphere. One of the most important and promising of these methods is the use of long-path infrared absorption by gases of the atmosphere. This method, useful only for gas concentration measurement and not for measuring the concentration of aerosols, employs a beam of radiation of wide bandwidth from a stable source strong in infrared radiation. The collimated beam passes through a part of the atmosphere to be studied, typically about 1 to 4 km. It may be received at the end of that path or returned by reflectors to a receiver near the transmitter. The receiver contains an infrared spectrometer.

Many methods other than long-path infrared absorption have been considered for the analysis of atmospheric gases. Much effort has been expended to measure accurately the concentrations of pollutants near cities and in the ambient atmosphere far from cities to obtain background concentrations. Other methods include Raman spectroscopy, the use of tunable laser sources, and observation of the ultraviolet electronic spectra of molecules. These methods may either provide the entire analysis or contribute partially. However, in this report, we confine our attention to the use of long-path infrared absorption spectra and attempt to exploit this method as fully as time and resources allow. It is the authors' belief that other methods will prove important in the future, but the long-established science of infrared absorption spectroscopy allows important progress to be made very quickly so that at least preliminary data may be obtained on the pollutants in the atmospheres of cities. It is probable that combinations of several methods will be utilized.

The analysis of the composite infrared spectra prduced by many atmospheric constituents is laborious and difficult. It requires a large computer and a library of spectra. The Wave Propagation Laboratory of the National Oceanic and Atmospheric Administration (NOAA) in Boulder, CO, has worked

for many years in a joint program with EPA to aid in the analysis of such composite spectra. The EPA financed portion of the work has been done under contract number EPA-IAG-077(D) entitled, "Remote Sensing of Pollutants."

This report is specialized in several ways. Firstly, it makes the assumption that the instrument used in obtaining the data is that described in 1.1. This assumption is necessary in order to be specific concerning the parameters used in our computational method. such an assumption, it would be necessary to consider wide ranges of parameters of a long-path spectrophotometer. The complexity caused by this would not clarify the method and would complicate the analysis. is anticipated that this specialization will not hamper the extension of the method to other specific devices. This has been accomplished by letting the input parameters to the computer program which analyzes the complex composite spectra to be at the discretion of the user. Thus, the choice of a different instrument would merely be reflected in different input parameters to the calculation. A second way in which the work has been limited is by the choice of spectra placed in the library. choice has been dictated by the impossibility of obtaining a complete library of spectra with the resources available.

Fortunately, for the immediate application of the method, a relatively small library of spectra is probably quite adequate. The spectra of the gases of importance in the Los Angeles, CA, atmosphere, the Raleigh, NC, atmosphere, and the atmosphere near refineries have been given preference whenever possible. However, the library of spectra has been limited by the unavailability of suitable spectra of some gases. Spectra were obtained from the scientific literature whenever possible. In certain cases, spectra were determined by laboratory experiment when they were not available in the literature. The entries and sources of the spectral library are given in Al.

The major normal atmospheric gases, H_2O , CO_2 , N_2O , and CH_4 , are part of the population. (Oxygen and nitrogen have no significant spectra in the frequency range of interest.) Normal constituents and pollutants are discussed in 2. The listed gases were chosen because of their presence

in the atmosphere and because detailed spectroscopic parameters are available for them, including either line positions, strengths, and line-broadening constants, or high-resolution experimental spectra.

The spectra of molecules have long been used as a definitive identifier, and when the spectrophotometer is properly calibrated, molecular concentrations can also be measured. In many cases, however, the method has been applied to relatively large concentrations in the laboratory, to pure gases, or at most to a mixture of a few constituents. It is true that laboratory spectrometers have been used to obtain low-level concentrations of a few species in the midst of large concentrations of other gases, but this method can only be successfully performed by simple observation when the regions of significant spectral absorption are well separated. Because most organic pollutant molecules have rich infrared spectra, this spectral region may be used for detection and measurement, but the potential for interference between overlapping spectral lines is high.

Thus, the measurement of atmospheric pollutants is complicated by low concentrations and by the simultaneous presence of many gases in temporally and geographically varying amounts. Because low concentrations may be significant for biological well-being, the method of analysis must be sensitive. Table 1.1 (Burriss et al., 1972)* shows typical concentrations of some pollutants in several typical cities. The sensitivity must be sufficient to determine such concentrations and, perhaps, a factor of 10 less. The interference due to aerosols is not considered in this report.

Because hydrocarbons (HC) are a large percentage of the total atmospheric pollutant content of many cities, they need special attention. In Chicago, IL, in 1968, the average HC concentration is 5380 $\mu g/m^3$, compared with CO at 7750 $\mu g/m^3$. The molecules making up the total HC content of the atmosphere for any location are not clearly known because many current analyzes separate only methane and non-methane HC. However, if there is no outstanding single HC and the concentration of HC is distributed among many different molecules, the weak absorption of each will provide

^{*} References are to authors and year of publication.

Table 1.1

ANNUAL AVERAGE CONCENTRATIONS
OF IMPORTANT POLLUTANTS IN CITIES (1968)

CO	HC*	NO ₂	NO	so_2
7750	5380	98	94	340
7000	4220	59	-	57
-	2690	78	67	230
6750	5380	78	54	29
5750	7300	39	40	86
4250	2690	98	54	110
	7750 7000 - 6750 5750	7750 5380 7000 4220 - 2690 6750 5380 5750 7300	7750 5380 98 7000 4220 59 - 2690 78 6750 5380 78 5750 7300 39	7750 5380 98 94 7000 4220 59 - - 2690 78 67 6750 5380 78 54 5750 7300 39 40

^{*}HC = hydrocarbons

Units: µg/m³

(From Burriss et al., 1972)

effectively a small absorption background which can be ignored. This problem can be solved definitively only after a detailed analysis of constituents is obtained for a particular region, and the spectra of these constituents are known.

Thus, the considerations discussed above require the analysis of a complex spectrum of many gases of relatively low amounts. This report is a description of the physical, mathematical, and computational procedures necessary to obtain the concentrations of many gases contained in an atmospheric sample by long-path infrared spectroscopy. The types of gases present in the sample are assumed known; the concentrations are unknown.

1.1 THE PHYSICAL PROBLEM

We consider an atmosphere composed, perhaps inhomogeneously, of normal and pollutant constituents. The pressure is assumed to be one atmosphere $(1.02 \times 10^6 \text{ dynes/cm}^2)$. We assume that the constituents do not

significantly change during the course of an observation. The observation is through a portion of the atmosphere approximately 61 cm in diameter and 1 to 4 km long. (These dimensions arise from the characteristics of a long-path spectrophotometer (see A3).) To focus on a specific problem, we choose the Research Triangle Area of North Carolina and a representative concentration of pollutants for that area will be used when available. For those pollutants whose average concentrations in Raleigh are not known, the known values in other comparable areas will be used. If no reliable values of trace materials can be obtained, they will be assumed to lie in the few parts per billion range. Further information may be found in A1, where TA1.1 and TA1.2 show the lowest concentrations assumed.

Although the method to be described is quite general, for clarity we will use the parameters of a specific system for remote sensing of pollutants (Streiff and Claysmith, 1972). This system, when used in an absorption mode, is a field-type infrared (IR) scanning spectrophotometer with a blackbody light source collimated by a telescope for long-path operation. Some system parameters of interest are given in A3.

The equipment can be used as a receiving spectrophotometer to study emissions from any sufficiently hot source, since it was designed to measure the emission spectra of gases in smokestacks. The methods developed herein are applicable to the study of emissions by substituting a library of emission spectra in place of the library of absorption spectra.

Thus, by means of the long-path spectrophotometer, we obtain the superimposed spectra of the atmospheric constituents in the spectral ranges 3 to 5 μ m (3330-1820 cm⁻¹) and 7 to 13.5 μ m (1430-740 cm⁻¹). The 5 to 7 μ m spectral range is not used because of the very strong absorption of water vapor. It is the purpose of the method described in this report to obtain from the superimposed spectra the concentrations of the molecular species present in the path.

Each of the gases present in the path acts independently of the others according to the Bouguer-Beers law. That is, a small thickness of

gas absorbs a fractional amount, $\frac{dI}{I} = -K'dx$, which implies

$$I = I_0 e^{-K'x}$$
 (1:1)

where I_0 is the initial intensity, K' is the absorption coefficient, and x is the absorber thickness. (Physically, K' is the reciprocal of the distance at which the intensity is 1/e of the initial intensity.) If the first absorber is followed by a second, equal to the first, it absorbs an equal fraction of the radiation incident upon it. If we consider a plane wave impinging on "slabs" of absorbing gases, the effect is the same whether the slabs follow one another or the gases are all placed in one slab. If the gases have absorption coefficients K_1' , K_2' , etc, then the intensity after passage through the absorbers is $I = I_0 e^{-(K_1' + K_2' + \ldots) x}$. The quantities K_1' , K_2' , ... are complicated functions of the frequency (herein designated as v and measured in cm⁻¹) and of the temperature, pressure, and concentrations of the gases. Hence I is also a complicated function of the same variables. The long-path spectrophotometer measures I/I_0 as a function of frequency and, from that data, it is necessary to determine the concentration of the gases.

To see how the concentration enters the problem, we write a K'x as $K\rho x = KW = \sigma N_{o} x$, where N_{o} is the number of absorbing molecules per unit volume, ρ is the density of absorbing gas per unit area of the radiation path, σ is the molecular absorption coefficient, W is the area density of gas, and K is the absorption coefficient. A very wide range of units has been used for these quantities in the literature. The reader should refer to Deutschman and Calfee (1967) and Calfee (1971) for transformations between commonly used sets of units. Table Al.4 lists a summary of these transformations. In this report, W is in units of molecules per square centimeter and I is in units of centimeters squared per molecule.

If the spectrophotometer measures I/I_0 as a function of wavelength after suitable adjustment and calibration, the data available (by taking the natural logarithm of each side of 1:1) is the product $KW = \ln \frac{I}{I_0}$, where KW is a function of frequency. But $KW = \sum_{i=1}^{N} K_i W_i$,

where N is the total number of absorbing gases, K_i is the absorption coefficient, and W_i is the (area) density of the ith gas. If we have the necessary spectral information about the gases, the K_i is known, and we need to determine the concentrations (determinable from W_i) of the gases present. The ROSE* system allows for an I/I output by having an internal blackbody that can be set at the same temperature as the remote blackbody. Their signals fall on the same detector with different chopping frequencies. They are separated and amplified by two phase sensitive locking amplifiers, and then ratioed. The problem that occurs (spurious variation of I/I o) is presumably due to the differences in the optical paths of the two beams. Thus a true I/I o is not provided.

If the instrument only provides relative transmission, I, rather than I/I , it is necessary to scale the data before the calculation of concentrations. The scaling must be done so that the input data to the computer program lies between zero and one, just as does I/I. This can be accomplished by simple scaling only if the response of the spectrophotometer is the same over all frequency ranges. If not, the data must be scaled in sections. In either case, only relative concentrations can be obtained. If absolute concentrations are required, and I/I_0 is not provided by the spectrophotometer, they may be obtained by adding the spectrum of a known amount of calibrating gas. There are several ways of doing this in practice. If, for example, the humidity is known along the path, this could provide an absolute scale. The accuracy would suffer, however, for obtaining low-level concentrations in comparison with the relatively larger concentration of water vapor, and it would be more accurate to choose an independent measurement of a gas of lower concentration. Another method would be to insert a test cell containing a gas of known concentration, adjusted so that its absorption falls in a spectral range unlikely to interfere with the unknown spectra. The concentration should be adjusted so as to fall within the range of most of the unknown gases, by trial and error, if necessary. The material of the cell

^{*} Remote Optical Sensing of Emissions

must be nonabsorbent or accounted for by observing the effect of an empty cell. I_0 can also be obtained by recording a spectrum upwind of an extended source or in a clean environment. The quantity can then be obtained downwind at the same path length and reference blackbody temperature. The resulting I/I_0 will then be an absolute spectrum.

The problem presented by the infrared continuum can under some circumstances be obviated by this method. The infrared continuum, a general increase of all absorption curves over a large part of the spectral range being considered here, occurs when long paths though the atmosphere are used. At present, only empirical methods of dealing with it are known. Over the relatively short path (2-4 km) planned for the spectrophotometer described in this report, the problem will often arise. (See Shapiro and Gush (1966), Burch (1970), and McClatchey et al., (1971).) However, in contamination such as occurs in Los Angeles, the continuum is often a significant contribution to the spectrum.

Determining concentrations is complicated by the inescapably finite signal-to-noise ratio (S/N) of the spectrophotometer and the noise due to atmospheric scintillation. Thus, the signal from the spectrophotometer has a noise component. It is probably not additive, but has been nonlinearly mixed by the system detector. However, a complete noise analysis of the detector is not available, and the character of the noise is dependent on the particular system. Therefore, we will make an assumption, for convenience, that the noise on the signal is additive Gaussian noise, that is, the output of the system is composed of a signal and noise, (I/I_0+N) , where the mean value of N is zero. The bandwidth of the Gaussian noise is determined by the time constant of the circuitry; it has essentially the same spectral bounds as the signal I/I_0 . The precision of the measurement is dependent on the S/N ratio and it, together with the resolution of the ratio digital voltmeter, determine the error. The ratio digital voltmeter of our system has a resolution of 1 part in 10⁴. However, the S/N ratio depends on atmospheric turbulence and is probably never as large as 10^4 (in the absorption mode). Hence the signal-to-noise ratio is usually the limiting factor in the precision of concentration measurement.

Streiff and Claysmith (1972) discuss the signal and noise characteristics of the spectrophotometer system. Their data and a private communication from M. Streiff indicate that practical working ranges of S/N run from 10 to 100. Without further tests of the system under working conditions, a complete noise analysis cannot be performed. To offer guidance on the degradation suffered in precision under finite signal-to-noise ratios, the computer analysis has been performed with an added simulated noise (see section 3.1.2).

In a different sense than the noise of atmospheric fluctuations and the spectrophotometer system discussed above, the presence of water vapor and to some extent that of natural carbon dioxide in the atmosphere constitute important sources of noise or interference. Water vapor concentration varies considerably in time and space, and its very strong absorption spectrum is found through most of the frequency region of interest. It interferes least with other spectra in the region from 2400 to $2600 \, \mathrm{cm}^{-1}$. Unfortunately, only a few gases of interest have observable absorptions there, and thus water vapor must always be one of the unknown gases whose concentration is determined by the procedure described If it is measurable otherwise, it may be desirable to consider removing its effect from the data by preprocessing. When the absolute humidity and temperature of the sample are known, the water vapor spectrum may be computed (1.2) and divided from the absorption spectrum to eliminate its effect. Carbon dioxide is also a naturally variable constituent, but its variation and absorption are less than those of water; and it usually is not an important interferant.

1.2 THE MATHEMATICAL PROBLEM

The transmission spectrum of an atmospheric constituent I/I_0 is a unique function of frequency ν , temperature T, pressure p, and concentration W, that is, $(I/I_0)_i = f_i(\nu, W_i, P, T)$, where the subscript indexes the

specific gas. Figure 1.1 (a-c) shows typical spectra of normal constituents and pollutants, F1.2 shows the normal atmosphere. If the transmission path contains two different gases, the resulting transmission spectrum is obtained by application of the Bouguer-Beers law (1:1). According to the discussion of 1.1, the spectrum due to two gases is obtained by multiplying the relative transmissions, that is,

$$(I/I_0) = (I/I_0)_1 (I/I_0)_2 = e^{-(K_1W_1 + K_2W_2)},$$
 (1:2)

and in general, for N gases,

$$(I/I_o) = \prod_{i=1}^{N} (I/I_o)_i = e^{-\sum_{i=1}^{N} (K_i W_i)},$$
 (1:3)

where (I/I_0) is the transmission of the composite atmosphere.

It should be noted that in whatever part of the frequency range one of the gases has zero (or small) transmission, the composite spectrum (I/I_0) is also zero. In any part of the frequency range where the transmission is one (no absorption) for all gases but one, the spectrum will be identical to that one. Otherwise, the spectrum will be the product of the constituent spectra and will generally present a complex problem to the experimenter attempting to identify the constituents and measure their concentrations. Figure 1.3 shows spectra of two gases, CH_A (methane) and HNO_3 (nitric acid), and their combined spectrum.

The spectra of the gases $(I/I_0)_i$ are functions of ν , T, P, and W. We assume that only the concentrations W_i are unknown, and the compound spectrum recorded as a function of frequency ν is the output of the spectrophotometer system.

In practice, $(I/I_0)_i$ as found in the library is accurate for only those values of P and T for which the spectrum was determined. Departures of $(I/I_0)_i$ from true (I/I_0) , however, are negligible for the range of temperatures and pressures found on the earth's surface.

H₂O

3000

Figure 1.1a
INFRARED SPECTRA OF SELECTED ATMOSPHERIC CONSTITUENTS
(Bands are identified on the overlay)

FREQUENCY (cm-1)

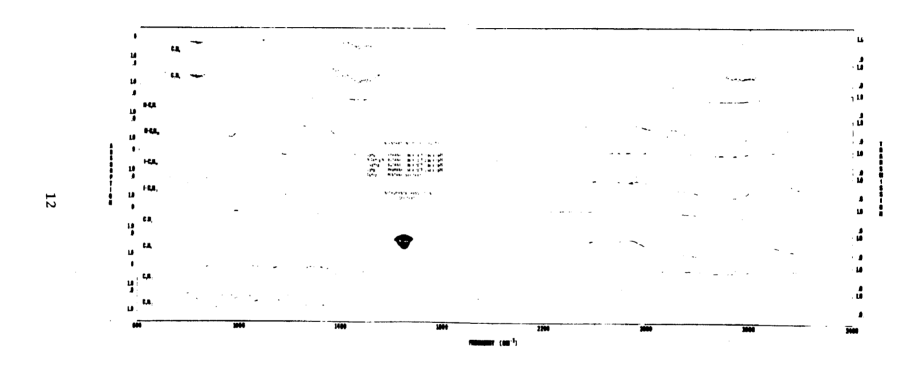


Figure 1.1b
INFRARED SPECTRA OF SELECTED ATMOSPHERIC CONSTITUENTS

13

Figure 1.1c
INFRARED SPECTRA OF SELECTED ATMOSPHERIC CONSTITUENTS

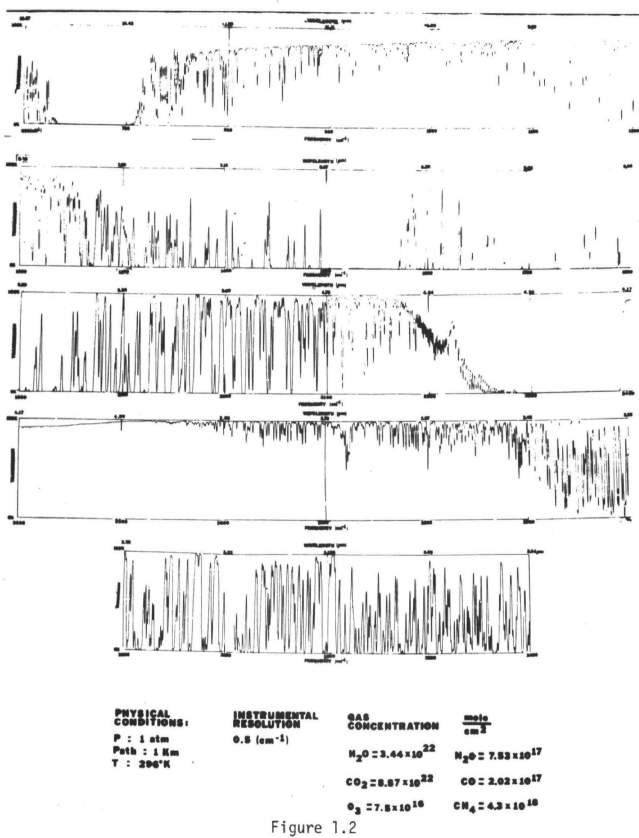


Figure 1.2
INFRARED SPECTRUM OF THE NORMAL ATMOSPHERE

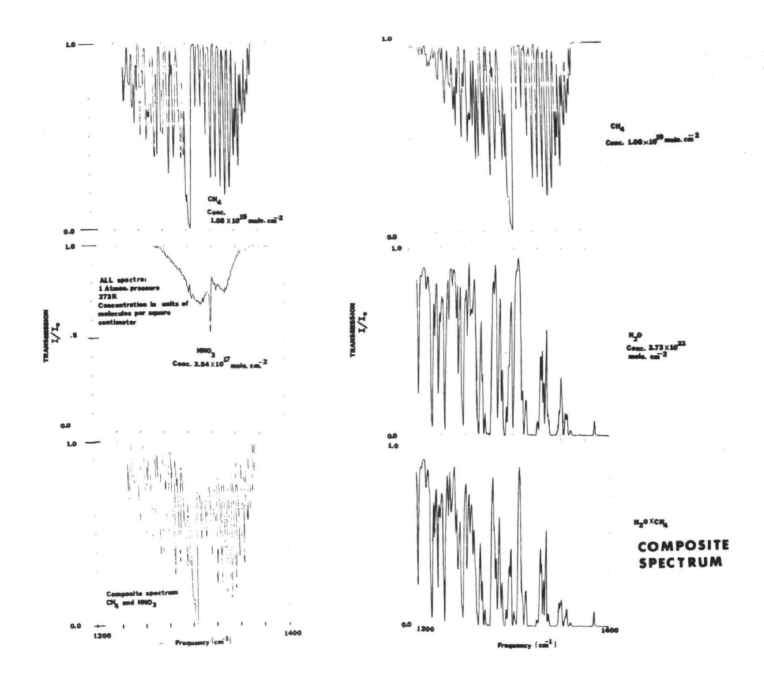


Figure 1.3 A PORTION OF THE TRANSMISSION SPECTRA OF CH $_4$, HNO $_3$, CH $_4$, AND H $_2$ O AND THEIR COMPOSITE SPECTRA

Therefore, in this application p and T are assumed constants, with p = 1 atm(1.02 x 10^6 dynes/cm²) and $T = 296^{\circ}$ K. For special circumstances, accurate $(I/I_o)_i$ may be calculated for any temperature and pressure only for molecules whose line parameters are fully known. This is accomplished using program DEGRADE (Deutschman and Calfee, 1967). For spectra taken experimentally, there is no known transformation to a new set of P and T without a detailed study of the molecule.

We call the output of the spectrophotometer $S(\nu)$. We must choose the concentrations W_i for all the gases which are in the sample path in such a way that the spectrum (I/I_0) of (1:3) is the same as, or as close as possible to, the experimental data $S(\nu)$. Simply put, this is performed by choosing a trial set of W_i , calculating (I/I_0) , and then comparing (I/I_0) with $S(\nu)$. The comparison is made by calculating an error function:

$$E_{j} = \left\{ \frac{1}{n} \sum_{v}^{n}, \left[(I/I_{o}; W_{i}^{j}) - S(v) \right]^{2} \right\}^{\frac{1}{2}}.$$
 1:4

Here the symbol (I/I_o; W_i^j) signifies the composite transmission spectrum, calculated from the spectral library under the assumption that the ith gaseous constituent has the trial concentration W_i^j . Of course (I/I_o; W_i^j) is a function of frequency. The number of terms in the sum is n. The summation over ν is somewhat arbitrary; maximum accuracy will be achieved by using all statistically independent measurements. A good compromise for the number of points was found by trial and error to be 5 per resolution element. (In the calculation described in this report, the data $S(\nu)$ and the spectra of the individual gases are stored at increments of $\Delta \nu = 0.2$ or 0.6 cm⁻¹, depending on the frequency range (see section 2).)

The trial spectra of individual gases are obtained by interpolation and extrapolation from a set of spectra given in the library for specific concentrations which span the range of concentrations expected in the atmospheric sample.

The first estimate of the concentrations, W_{i}^{j} , is generally not correct, of course. After the calculation of the first error, E_{1} , the concentrations are varied in an orderly stepwise manner, calculating successive corresponding errors E_{j} , until the set of concentrations is obtained that minimize E_{j} . These concentrations are then the "best" estimate of the concentrations in the atmospheric path in which the sample data S(v) were obtained.

The calculation is performed by the computer subroutine MINMYZD, developed by Slutz and Winkelman (1964) and applied by Lawrence and Hallenbeck (1965). MINMYZD is described in detail in A2.

This computer routine, although in principle as simple as described above, requires careful design to prevent costly, long computer runs and inaccuracies. The problem is to find the minimum value of a multidimensional function, E, whose variables are the W_i. Care must be taken to avoid local minima in this function and to find the absolute minimum. The subroutine MINMYZD has features which practically assure that result in applications such as herein discussed.

It is not possible and not desirable to discuss here the problems of uniqueness involved in minimization problems such as this one. Instead, the long experience with this program and its successful application to many physical problems gives assurance of its usefulness and accuracy.

Arbitrarily chosen initial concentrations prolong the computer running time and may cause inaccuracy in the final results. Hence initial concentrations are selected for each constituent by computing a "best fit" in a very limited spectral range where the constituent has a prominent spectrum and the interference from other gases is small. These selected concentrations are then employed as initial, approximate choices in the computation over the whole spectral range. When the number of gases is small, these initial values may be sufficient for the experimenter's needs.

It is true that the optimization or best-fit computation here employed is not the only way the concentrations could be obtained from the experimental data. Many other methods exist, under the names of best-fit,

minimization, or inversion methods. No attempt was made to evaluate which of these would be the best method. The MINIMYZD routine was selected because of its long history of successful efficient operation. Methods such as that of Backus and Gilbert (Westwater and Cohen, 1973) offer some advantages of error estimation. Matthes (1973) has devised a method to circumvent problems arising in least-squares fitting which results in negative concentration estimates. This problem may be prevented in the present program by placing a lower bound of zero on any concentration estimate, but it is probably better practice to eliminate such absolute constraints on the MINMYZD subroutine to prevent nonlinear effects on the search process.

2. PRESENT STATUS OF THE LIBRARY OF ABSORPTION SPECTRA

It is essential in the identification of pollutant gases by long-path absorption spectroscopy to possess accurate spectra of the gases expected to occur in the atmosphere under examination. These spectra must be complete in the spectral range covered by the long-path spectrophotometer because 'maverick' gases (i.e., gases present in the sample but not in the spectral library) cause inaccuracy in the calculation of the concentrations. Some examples of such inaccuracies are presented in 3.

The spectra must be known in such a way that they may be determined as a function of temperature, pressure, and concentrations to suit the conditions in the test atmosphere. If, for example, the line position, the line strength, and the broadening parameters of a gas as functions of temperature and pressure are known, the spectra may be computed for any concentration by computer routines (Deutschman and Calfee, 1967, and McClatchey et al., 1973). If these parameters are unknown, but accurately measured experimental spectra are available for concentrations near those of interest, we may extrapolate from them to other concentrations.

To make use of such experimental data, it is necessary that the spectra be obtained at a temperature and pressure near those normally encountered over polluted cities, that resolution is comparable to or better than that of the spectrophotometer used, and that the ordinate of the spectra is calibrated as an absolute transmittance (or, that the line strengths in some way are known).

Whether such spectra are available depends on whether researchers have been interested in these specific molecules. No concerted effort has been made to the authors' knowledge to obtain detailed spectra of all atmospheric constituents including pollutants. However, an effort has been made by a loosely knit group of researchers* to obtain detailed

^{*} The Group on Atmospheric Transmission Studies (GOATS).

spectra on the normal atmospheric constituents. These data are summarized by McClatchey et al., (1973).

In general, however, the search of literature on the spectra of normal and pollutant atmospheric constituents reveals a wasteland punctuated by a few oases of excellent work. Spectroscopists, academic and industrial, have not been supported in obtaining the high-resolution (1 cm⁻¹) infrared spectra with a well calibrated intensity scale at atmospheric pressure needed for concentration measurements in an atmosphere containing many interfering gases.

The chief limitation of the application of this (or any) spectroscopic technique to the detection of atmospheric constituents lies in the ignorance of the spectra of many atmospheric constituents, particularly pollutants. Just as the criminal's fingerprints cannot be used to identify him unless they are in police files, so observation of a gas spectrum only confuses as if it is not in our library. Notably, spectra are unavailable for many hydrocarbons. Also spectra of excited (high temperature) molecules such as found in smoke stacks are not in the literature.

The sources of the spectral library are summarized in Al and detailed descriptions are given there of the parameters of the spectra. Also described are manipulations performed in converting spectra found in the literature into computer compatible form. We summarize in T2.1 the gases available in the spectra library. Spectra for selected concentrations for all these gases are displayed in F1.1(a-c).

Table 2.1 GASES AVAILABLE IN THE SPECTRAL LIBRARY

Gas	Formula	Accuracy of Spectrum
Water	н ₂ 0	1
Carbon dioxide	co ₂	1
Ozone	03	1
Nitrous oxide	N ₂ 0	1
Methane	CH ₄	1
Carbon monoxide	co	1
Hydrochloric acid	HCL	1
Sulfur dioxide	so ₂	2
Nitric acid	HNO ₃	2
Ammonia	NH ₃	2
Nitric oxide	NO	1
Hydrogen sulfide	H ₂ S	2
Ethylene	$C_2^-H_4$	2
Nitrogen dioxide	NO ₂	2
Acetone	CH ₃ CH ₂ HCO	3
Formaldehyde	H ₂ CO	2
Ethane	С ₂ н ₆	2
Normal-butane	$n-C_4H_{10}$	3
Iso-butane	i-C ₄ H ₁₀	3
Propane	C ₃ H ₈	3
Pentane	C ₅ H ₁₂	3

^{*}Accuracy of spectrum: (1) Line information highly trustworthy;

⁽²⁾ Good experimental spectrum available, trustworthy; and

⁽³⁾ Low resolution only, some uncertainty.

3. BEST-FIT CALCULATIONS OF CONCENTRATIONS FROM A SIMULATED COMPLEX SYSTEM

The application of the MINMYZD fitting routine to specific problems requires a controlling program to direct progress from the input of data through the minimization process to the final gas constituent concentrations and error estimates. This program, called EPAGAS, is described in detail in A2.

Using this program, the entire computation was tested by preparing trial spectra, simulating the data obtained from a long atmospheric path spectrophotometer experiment, and then obtaining the concentrations. The tests were made "blind", that is, the programmer was presented with the simulated sample data without knowledge of the concentrations until the computer results were compared with the known composition. In this section, a number of such computations are reviewed. The objectives were to "debug" the program, to test the success of determining the concentrations of gases from their composite spectrum, to determine the accuracy achievable, and to estimate the degradation of accuracy due to finite signal-to-noise ratio in the data. Further, the effects of a lack of a one-to-one correspondence of the library of spectra with the atmospheric constituents are examined.

In these preliminary tests, library spectra of $\rm H_2O$, $\rm N_2O$, $\rm CH_4$, $\rm SO_2$, $\rm HNO_3$, $\rm CO$, $\rm CO_2$, and $\rm O_3$ were employed in various concentrations of combinations to produce a complex spectrum.

3.1 ACCURACY OF CONCENTRATION CALCULATIONS

Tables 3.1 through 3.4 list the trial runs made on the combinations of gases indicated at the top of the table. Each fitting was made over a narrow frequency range to determine the suitability of that range for detection of the gas under investigation and to observe inaccuracies arising when the gas has no absorption in the region.

It should be noted here, and it is emphasized and clarified in A2, that the analysis of complex spectra by the method which is the subject of this report should not be done blindly or automatically.

Table 3.1

CALCULATION OF GAS CONCENTRATIONS
FROM SIMULATED COMPOSITE SPECTRA (SET 1)
(INFINITE SIGNAL-TO-NOISE RATIO)

Code	Range	$W_{\rm g} = 3.73 \times 10^{22}$		$N_2 \cap (4A1)$ $W_g = 1.08 \times 10^{1.8}$		$CH_{4} (5A1)$ $Wg = 1.08 \times 10^{19}$		$SO_2 (20.2)$ Wg = 3.54 x 10^{17}		HNO_3 (30.2) Wg = 3.54 x 10 ¹⁷		Final Transmission Error RMS	
	(cm ⁻¹)	Wc	_ AW	Nc	ΔW	Ψc	ΔW	Wc	ΔW	Wc	∆W	Wtd.	Unwtd.
8888 a 1	230.0-1249.8	3.73x10 ²²	.00	1.09x10 ¹⁸	01	1.08x10 ¹⁹	. ೧೧	17.88x10 ¹⁷	-14.34	0.07x10 ¹⁷	3.47	.03	6.60
b 1:	250.0-1269.8	3.73x10 ²²	.00	1.09x10 ¹⁸		1.08x10 ¹⁹		17.88x10 ¹⁷		0.19×10^{17}	3.44	.06	6.60
c 1	270.0-1289.8	3.73x10 ²²	.00	1.08x10 ¹⁸	.00	1.08x10 ¹⁹	.00	17.88x10 ¹⁷	-14.34	3.55x10 ¹⁷	- 0.01	.03	6.41
d 1	290.0-1309.8	3.73x10 ²²	.00	1.10x10 ¹⁸	02	1.08x10 ¹⁹		0.75x10 ¹⁷		3.45×10^{17}	0.09	.05	1.25
	310.0-1329.8		.00	1.23x10 ¹⁸	-,15	1.06x10 ¹⁹		4.29×10^{17}		3.42x10 ¹⁷	0.12	.03	0.35
f 1:	330.0-1349.8	3.~3x10 ²²	.00	1.21x10 ¹⁸	13	1.08x10 ¹⁹		17	0.03	3.56×10^{17}	- 0.02	.02	0.06
	350.0-1369.8		.00	1.21x10 ¹⁸		1.06x10 ¹⁹	.02	3.32x10 ¹⁷	0.22	4.32×10^{17}	- 0.78	. 03	0.37
	370.0-1389.8		.05	1.21x10 ¹⁸	13	1.48x10 ¹⁹	40	4.88x10 ¹⁷		18.38x10 ¹⁷	-14.84	.03	6.68
	380.2-1400.0		.06	1.21x10 ¹⁸		1.48x1c ¹⁹	40			18.23x10 ¹⁷	-14.69	.03	6.61

Wg = given concentration in simulated sample, molecules $\,\mathrm{cm}^{-2}$

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMY2D to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1).

Wc = calculated concentration, molecules cm⁻²

 $[\]Delta W$ = Wg-Wc (the factor of a power of ten is omitted), molecules cm⁻²

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

Table 3.2

CALCULATION OF GAS CONCENTRATIONS
FROM SIMULATED COMPOSITE SPECTRA (SET 2)

(INFINITE SIGNAL-TO-NOISE RATIO)

	Wavenumber Range	- ` `		CO_2 (2B1) Wg = 1.08 x 10^{21}		O_3 (3B1) Wg = 2.69 x 10 ¹⁷		N_2O (4B1) $Wg = 1.08 \times 10^{1.8}$		CO (6B1) Wg = 1.88 x 10 ¹⁹			Transmission Error	
Code	(cm-1)	Wc_	Δ₩	Wc	ΔW	Wc	∆W.	Wc	Δ₩	Wc	ΔW	Wtd.	Unwtd.	
9999 a	2100.0-2159.4		01	1.08x10 ²¹	.00	2.84x10 ¹⁷	-0.15	1.22x10 ¹⁸	14	1.88x10 ¹⁹	0.00	.03	0.09	
b	2160.0-2219.4		.01	1.05x10 ²¹	.03	5.94x10 ¹⁷	-3.25	1.08x10 ¹⁸	.00	1.88x10 ¹⁹	0.00	. 03	1.45	
c	2220.0-2279.4	3.68x10 ²¹	.05	1.08x10 ²¹	.00	5.94x10 ¹⁷	-3.25	1.08x10 ¹⁸	.00	1.88x10 ¹⁹	0.00	.03	1.45	
ď	2240.4-2299.8	3.69x10 ²¹	.04	1.08x10 ²¹	.00	5.94x10 ¹⁷	-3.25	1.08x10 ¹⁸	.00	2.98x10 ¹⁹	-1.10	.03	1.53	

Wg = given concentration in simulated sample, molecules cm⁻²

Wc = calculated concentration, molecules cm⁻²

ΔW = Wg-Wc (the factor of a power of ten is omitted), molecules cm⁻²

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMYZD to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1).

Table 3.3a
CALCULATION OF GAS CONCENTRATIONS
FROM SIMULATED COMPOSITE SPECTRA (SET 1)
(S/N=10)

	Wavenumber	Wavenumber H_2O (1) Range $Wg = 3.73$				CF1 4 (Wg = 1.08	5A1) x 10 ¹⁹	SO_2 (2) $Wg = 3.54$:	0.2) x 10 ¹⁷	HNO_3 (30.2) Wg = 3.54 x 10^{17}		-	ransmission rror	
Code	(cm-1)	Wc		Wc	ΔW	Wc	<u> </u>	КС	∇R_i	Wc	ΔW	Wtd.	Unwtd.	
S/N = 10														
8 888 a	1230.0-1249.8	3.80x10 ²²	07	0.80×10^{18}	. 28	1.04×10 ¹⁹	. 04	17.88x10 ¹⁷	-14.34	-1.89×10^{17}		2.08	6.86	
ь	1250.0-1269.8	3.74x10 ²²	01	0.82x10 ¹⁸	.26	1.13x10 ¹⁹	05	17.88x10 ¹⁷	-14.34			2.28	6.54	
С	1270.0-1289.8		. 20	1.04x10 ¹⁸	. 04	1.06x10 ¹⁹	.02	17.88×10 ¹⁷	-14.34	4.81x10 ¹⁷	1.27	1.86	6.44	
d	1290.0-1309.8		25	1.03x10 ¹⁸	.05	1.08×10 ¹⁸	.00	0.28x10 ¹⁷	3.26	3.54x10 ¹⁷	0.00	1.50	1.46	
e	1310.0-1329.8		25	0.75x10 ¹⁸	.33	1.02x10 ¹⁹	.06	2.50x10 ¹⁷	1.03			0.76	0.54	
f	1330.0-1349.8	3.72 x10²²	.01	1.21x10 ¹⁸	13	1.11x10 ¹⁹	03	2.75x10 ¹⁷	0.79	3.95x10 ¹⁷		0.80	0.40	
g	1350.0-1369.8	3.75x10 ²²	02	1.21x10 ¹⁸	13	0.99x10 ¹⁹	.09	4.15x10 ¹⁷	- 0.61	4.69x10 ¹⁷		0.28	0.59	
h	1370.0-1389.8	3.71x10 ²²	.02	1.21x10 ¹⁸	13	1.48x10 ¹⁹	40	4.24x10 ¹⁷	- 0.70			0.14	6.55	
i	1380.2-1400.0		.08	1.21x10 ¹⁸	13	1.48x10 ¹⁹	40	5.60x10 ¹⁷	- 2.06	18.22x10 ¹⁷	14.68	0.14	6.63	

Wg = given concentration in simulated sample, molecules cm⁻²

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMYZD to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1.2).

Wc = calculated concentration, molecules cm^{-2}

ΔW = Wg-Wc (the factor of a power of ten is omitted), molecules cm⁻²

Table 3.3b (S/N=30)

	Wavenumber Range	lange $Wg = 3.73 \times 10^{22}$		N_2O (4A1) Wg = 1.08 x $10^{1.6}$		CH 4 (5A1) Wg = 1.08 x 10 ¹⁹		SO_2 (20.2) Wg = 3.54 x 10^{17}		HNO ₃ (30.2) Wg = 3.54 x 10 ¹⁷		Final Transmissic Error RMS	
Code	(cm ⁻¹)	Wc	∑W.	No.	ΔW	We	ΔW	We	ZN	Wc	ΔW	Wtd.	Unwtd.
S/N = 30													
8888 a	1230.0-1249.8	3.76x10 ²²	~.03	0.50×10^{18}	.58	1.08x10 ¹⁹	.00	17.88x10 ¹⁷	-14.34	0.15×10^{17}	3.39	.65	6.59
ъ	1250.0-1269.8		.07	1.23x10 ¹⁸	15	1.07x10 ¹⁹	.01	17.88x10 ¹⁷		-0.69×10^{17}	4.23	.75	6.69
с	1270.0-1289.8	3.68x10 ²²	. 05	1.05x10 ¹⁸	.03	1.05x10 ¹⁹	.03	17.88x10 ¹⁷		4.11x10 ¹⁷	- 0.57	.65	6.42
đ	1290.0-1309.8		.00	1.11x10 ¹⁸	03	1.07x10 ¹⁹	.01	1.00x10 ¹⁷	2.54	3.50x10 ¹⁷	0.04	.46	1.14
e	1310.0-1329.8		02	1.38x10 ¹⁸	30	0.97x10 ¹⁸	.11	4.35x10 ¹⁷	- 0.81	3.40×10^{17}	0.14	.25	0.39
f	1330.0-1349.8		.00	1.21x10 ¹⁸	13	1.05x10 ¹⁹	.03	3.64x10 ¹⁷	- 0.10	3.54x10 ¹⁷	0.00	.26	0.07
g	1350.0-1369.8	3.76x10 ²²	03	1.21x10 ¹⁸	13	1.07x10 ¹⁹	.01	3.06x10 ¹⁷	0.48	4.58x10 ¹⁷	- 1.04	.10	0.52
ħ	1370.0-1389.8		.05	1.21x10 ¹⁸	13	1.48x10 ¹⁹	~.40	4.83x10 ¹⁷	- 1.29	18.20x10 ¹⁷	-14.66	.05	6.58
i	1380.2-1400.0	3.69x10 ²²	. 04	1.21x10 ¹⁸	13	1.48x10 ¹⁹	40	4.54x10 ¹⁷	- 1.00	18.24x10 ¹⁷	-14.70	.06	6.59

Wg = given concentration in simulated sample, molecules cm⁻²

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMYZD to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1.2).

Wc = calculated concentration, molecules ${\rm cm}^{-2}$

ΔW = Wg-Wc (the factor of a power of ten is omitted), molecules cm⁻²

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

Table 3.3c (S/N=100)

	Wavenumber Range	H_2O (1 Wg = 3.73		N ₂ O (4 Wg ≈ 1.08	A1) x 10 ^{1 8}	CH4 ((5A1) x 10 ¹⁹	SO_2 (2) $Wg = 3.54$	0.2) x 10 ¹⁷	HNO_3 (3) Wg = 3.54 x			ransmissior rror
Code	(cm ⁻¹)	Жc	ΔW	We	۷W		ΔW	Wc	Δ₩	Wc	ΔW	Wtd.	Unwtd.
/N = 100 888 a	1230.0-1249.8	3.74x10 ²²	01	1.11x10 ¹⁸	03	1.07x10 ¹⁹	. 01	17.88x10 ¹⁷	-14.34	- 0.03x10 ¹⁷	3.57	.25	6.61
ъ	1250.0-1269.8	3.71×10^{22}	.02	1.13x10 ¹⁸	05	1.07x10 ¹⁹	.01	17.88x10 ¹⁷	-14.34	$0.08 x 10^{17}$	3.46	.22	6.60
c	1270.0-1289.8	$3.73x10^{22}$.00	1.09x10 ¹⁸	~.01	1.09x10 ¹⁹	01	17.88x10 ¹⁷	-14.34	$3.49x10^{17}$	0.05	.20	6.41
đ	1290.0-1309.8		.01	1.09x10 ¹⁸	01	1.08x10 ¹⁹	.00	0.74x10 ¹⁷	2.80	3.50x10 ¹⁷	0.04	.17	1.25
e	1310.0-1329.8	3.72x10 ²²	.01	1.21x10 ¹⁸	13	1.08x10 ¹⁹	.00	4.11x10 ¹⁷	- 0.57	3.50x10 ¹⁷	0.04	.07	0.26
f	1330.0-1349.8	3.74 x 10 ²²	01	1.21x10 ¹⁸	13	1.07×10 ¹⁹	.01	3.57×10^{17}	- 0.03	3.51×10^{17}	0.03	.09	0.06
g	1350.0-1369.8	3.74x10 ²²	01	1.21x10 ¹⁸	13	1.06×10 ¹⁹	.02	3.25x10 ¹⁷	0.29	4.37x10 ¹⁷	0.83	.04	0.40
h	1370.0-1389.8	3.68x10 ²²	.05	1.21x10 ¹⁸	13	1.48x10 ¹⁹	40	4.84x10 ¹⁷	- 1.30	18.37x10 ¹⁷	-14.83	.03	6.66
i	1380.2-1400.0		.06	1.21x10 ¹⁸	13	1.48x10 ¹⁹	40	5.19x10 ¹⁷	- 1.65	18.23x10 ¹⁷	-14.69	.03	6.61

Wg = given concentration in simulated sample, molecules ${\rm cm}^{-2}$

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMYZD to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1.2).

Wc = calculated concentration, molecules cm^{-2}

 $[\]Delta W$ = Wg-Wc (the factor of a power of ten is omitted), molecules cm⁻²

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

Table 3.4

CALCULATION OF GAS CONCENTRATIONS

FROM SIMULATED COMPOSITE SPECTRA (SET 2)

(S/N = 10, 30, 100)

	Wavenumber Range	H ₂ O (1 Wg = 3.73	B1)	CO_2 (2 Wg = 1.08	2B1)	0 ₃ Wg = 2.69	(3B1)	N_2O (4 $Wg = 1.08$	B1)	CO (6	B1)	Eı	ansmissio ror
Code	(cm ⁻¹)	Wc Wc		We	ΔW	Wc	ΔW	Wc 1.00		Wg = 1.88 Wc	<u> Δ</u> ₩	RMS Wtd.	Unwtd.
S/N = 10 9999 a	2100.0-2159.4	3.66x10 ²¹	0.07	1.07x10 ²¹	.01	1.79x10 ¹⁷	0.90	1.34x10 ¹⁸	26	1.91x10 ¹⁹	03	1.09	0.42
ъ	2160.0-2219.4	4.08x10 ²¹	-0.35	0.35x10 ²¹	.73	5.94x10 ¹⁷	-3.25	1.07x10 ¹⁸	. 01	1.88x10 ¹⁹	.00	1.28	1.50
c	2220.0-2279.4	5.10x10 ²¹	-1.37	1.11x10 ²¹	03	5.94x10 ¹⁷	-3.25	1.11x10 ¹⁸	-,03	1.87x10 ¹⁹	.01	2.64	1.58
d	2240.4-2299.8	-0.93x10 ²¹	4.66	1.10x10 ²¹		5.94x10 ¹⁷	-3.25	1.06x10 ¹⁸	.02	1.33x10 ¹⁹	.55	3.54	2.55
S/N = 30							,						····
9999 a	2100.0-2159.4	3.73x10 ²¹	0.00	1.14x10 ²¹	06	2.53x10 ¹⁷	0.16	1.19x10 ¹⁸	11	1.88x10 ¹⁹	.00	0.36	0.09
ь	2160.0-2219.4	3.89x10 ²¹	-0.16	1.27x10 ²¹		5.94x10 ¹⁷	-3.25	1.08x10 ¹⁸	.00	1.88x10 ¹⁹	.00	0.44	1.46
с	2220.0-2279.4	4.72x10 ²¹	-0.99	1.09x10 ²¹		5.94x10 ¹⁷	-3.25	1.09x10 ¹⁸	01	1.89x10 ¹⁹	01	0.86	1.52
4	2240.4-2299.8	2.67x10 ²¹	1.06	1.08x10 ²¹		5.94x10 ¹⁷	-3.25	1.07x10 ¹⁸	.01	1.49x10 ¹⁹	.39	1.19	1.54
S/N = 100				·	- 1 · · · ·							 	
9999 a	2100.0-2159.4	3.72x10 ²¹	0.01	1.07x10 ²¹	.01	3.01x10 ¹⁷	32	1.19x10 ¹⁸	11	1.88x10 ¹⁹	.00	0.10	0.15
ь	2160.0-2219.4		0.05	1.08x10 ²¹	.00	5.94x10 ¹⁷	-3.25	1.08x10 ¹⁸	.00	1.88x10 ¹⁹	.00	0.14	1.45
с	2220.0-2279.4		0.05	1.08x10 ²¹	.00	5.94x10 ¹⁷	-3.25	1.08x10 ¹⁸	.00	1.87x10 ¹⁹	.01	0.26	1.45
đ	2240.4-2299.8		-0.05	1.08x10 ²¹		17	-3.25	1.08x10 ¹⁸	.00	2.59x10 ¹⁹	71	0.31	1.49

Wg = given concentration in simulated sample, molecules cm^{-2}

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMYZD to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1.2).

Wc = calculated concentration, molecules cm⁻²

ΔW = Wg-Wc (the factor of a power of ten is omitted), molecules cm⁻²

To apply directly the method to a sample of unknown composition, utilizing the full library of test spectra, would be an example of how much must be paid (in computer costs) for total ignorance of the sample. Maximum information available on the composition of the unknown sample should be utilized in selecting the library of test spectra. In the examples discussed below (3.1.3), we will see the penalty on the accuracy of the calculation for omitting a spectrum from the library when the gas is in the sample.

Further, we note that suitable choices of frequency regions to investigate the presence of a given gas must be made. Obviously, if at concentration levels of interest, a gas shows no (or weak) absorption, its concentration may not be measured there. In F3.1 (and F1.1a) consider the two spectral regions of absorption by HNO₃, namely those at 880 and 1300 cm⁻¹. The higher frequency band has strong potential interference from H₂O, SO₂, N₂O, CH₄, and all HC. Thus, if HNO₃ is suspected in a sample, the concentration should be calculated in the lower frequency band. Therefore, the analyst should consult the positioning of spectra in F3.1 before beginning his analysis.

Suppose an analyst had unlimited free computer time. Should he not then simplify the exercise of judgment and insert his entire library into the computer and analyze the whole frequency range? But here he will find a requirement for careful judgment also. For, as we will see below, in regions where absorption is effectively zero, the results of the calculation are small but essentially random. Therefore, the analyst must choose from among the calculated concentrations obtained in all regions only those where the absorption is substantial for the gas of interest. Such a weighting of values could be added to the computer program. It is uncertain whether the advantages outweigh the added complexity in the program.

In T3.1 through T3.3, a uniform notation is used. Two separate sets of gases are used. Set 1 consists of $\rm H_2O$, $\rm N_2O$, $\rm CH_4$, $\rm SO_2$, and $\rm HNO_3$, and has the Code 8888. Set 2 consists of $\rm H_2O$, $\rm CO_2$, $\rm O_3$, $\rm N_2O$, and $\rm CO$,

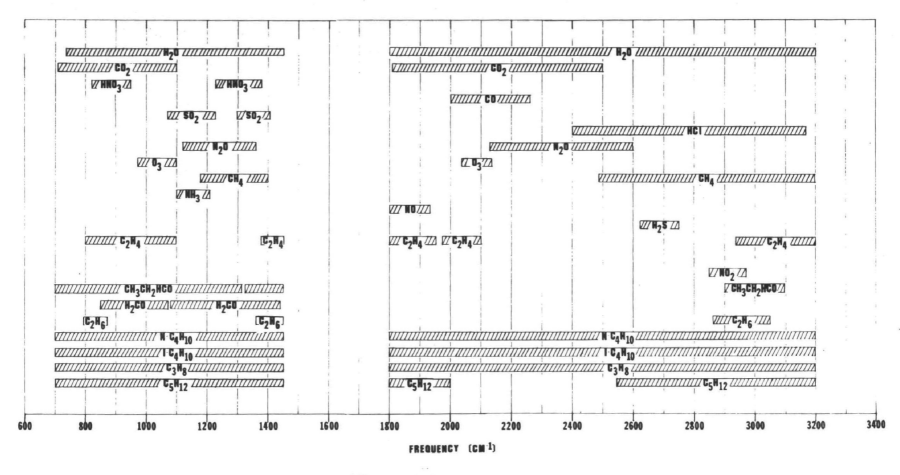


Figure 3.1

PRINCIPAL REGIONS OF ABSORPTION OF SELECTED NORMAL AND POLLUTANT ATMOSPHERIC CONSTITUENTS (The region from 1460 to 1800 cm⁻¹ has strong water vapor absorption)

and has the Code 9999. H₂0 is included in both sets because it is always present in the atmosphere; CO₂, though always present, is not as variable. The other gases were chosen to be representative spectra in the upper and lower frequency ranges (see Fl (a-c) and F3.1). The concentration (molecules cm⁻²) is denoted for each gas by Wg. The concentrations are chosen to match representative values in the Los Angeles area. The frequency range for which the calculation was made is designated by a small letter. The total range was divided into subranges to observe the effect of spectral strength and structure on the accuracy of the computation. The calculated concentration is denoted by Wc. The difference (Wg-Wc)xl0^{-y} is a measure of the accuracy (y is the exponent in Wg). Two measures of rms error are given, the weighted error and the unweighted error. The weighted error is of use to the programmer. Here we will refer to the unweighted error.

The penultimate columns of the tables contain those final transmission errors defined as

$$\begin{bmatrix} \frac{1}{N} & \sum_{k=1}^{N} \left\{ \left[\text{(True Transmission)}_{k} - \text{(Calculated Transmission)}_{k} \right]^{\frac{1}{2}} \right\} ,$$

where CF_k is a "confidence factor" introduced to weight in favor of high concentrations because they produce, generally speaking, less error. (See A2.5.) The transmissions involved are those due to the total sample of all gases. The summation is over each experimental data value of the frequency range given in the tables. It thus is a figure of merit for "goodness of fit" over that range. The unweighted error is obtained by letting all $\operatorname{CF}_k=1$.

The unweighted error presented in the final column is a different sort of measure. It is simply the rms of the fractional errors (Wg-Wc)/Wg, on a line of the table. In the error column, factors of 10 have been omitted.

Tables 3.1 and 3.2 are based on an assumption of infinite signal-to-noise ratio. Tables 3.3 (a-c) and 3.4 illustrate the effect of finite S/N. Tables 3.5 (a, b) and 3.6 illustrate the effect of "maverick" gases. All these cases are discussed in the following sections.

3.1.1 Calculation of Concentrations for Spectra Without Noise

Consider the calculations coded as 8888 (a-i) in T3.1 (Set 1). A simulated sample was prepared, containing concentrations of the gases listed, unknown to the analyst. Using the simulated sample as data, concentrations were computed by the MINMYZD method. In the frequency ranges given in the second column, a concentration of water of Wg = 3.73×10^{22} is relatively strong and the small errors (Δ W) under H_2O are to be expected when, as here, the S/N is assumed infinite. The inaccuracy lies between 0 and 2 percent, the larger error occurring toward the strong "black-out" absorption band of H_2O lying between 1430 and 1800 cm⁻¹. Interference from the other constituents of the sample has little effect on the calculated concentrations of H_2O .

The calculated concentration of N_2^0 from the data is done with an error not exceeding 14 percent. The larger error occurs where the absorption is very small, and such regions should be lightly weighted in a final averaging. By such weighting, we may estimate a maximum error of 5 percent in this case.

Methane (CH₄) has a strong absorption spectrum and very distinctive spectral structure in part of the frequency range covered in T3.1, and the error in the calculated concentration is small except where the absorption is very weak in the range 1370 to 1400 cm⁻¹, being 37 percent. The given concentration of SO_2 , $Wg = 3.54 \times 10^{17}$ molecules cm⁻²,

The given concentration of SO_2 , $Wg = 3.54 \times 10^{17}$ molecules cm⁻², is relatively small. Better results than shown can be obtained by using the stronger absorption spectrum between 1050 and 1200 cm⁻¹ (see F1.1a). Here, the errors, excluding the regions from 1230 to 1310 cm⁻¹, range from 1 to 44 percent. By using the results only from the strongest absorbing region, the errors are quite small, averaging 9 percent.

The last gas considered in this set is HNO3. Its spectrum is strong,

but the concentration is relatively low. Again we see small errors where the absorption is strong, and large errors amounting to 100 percent where the absorption is weak.

The calculations listed under Codes 9999 (a-d) in T3.2 give a similar set of calculations for a different set of gases, namely $\rm H_2O$, $\rm CO_2$, $\rm O_3$, $\rm N_2O$, and CO. The spectra of these gases are shown in F1.1a, except for $\rm O_3$. Its absorption, in the frequency range 2100 to 2300 cm⁻¹, is too weak to show on the scale of the graph; it is listed however in the library of spectra.

In this set of gases, the errors, except for 0_3 , are quite small. The best analysis of 0_3 would be made in the stronger absorption band centered at 1100 cm^{-1} .

It is, of course, not surprising that we may conclude from these computations that errors for a particular gas are minimized by choosing regions of strong absorption of that gas, as free from interference as possible. These results illustrate the earlier discussion that intelligent choices of frequency regions for the computation are necessary.

3.1.2 Effect of Finite Signal-to-Noise Ratio (S/N)

As remarked in section 1.1, the spectrophotometer system has a finite S/N ratio. To estimate the effect of noise on the system, simulated random noise has been added to the data in runs under Codes 8888 (a-i) and 9999 (a-d) as shown in T3.3 (a-c) and T3.4. The S/N ratios of 10, 30, and 100 were simulated. The method of simulation is discussed in A2.

By comparing T3.1 and T3.3, we see that errors (ΔW) are not substantially increased at SN = Ω 0 over the noise-free case. Average errors increase at S/N = 30, and they become considerable in some cases at S/N = 10. Generally, the calculated concentrations are within an order of magnitude of the true sample concentrations (except in the case of SO₂ and HNO₃ in those regions of the spectrum where absorption is weak). Whether these errors are tolerable or not depends on the needs of the experimenter. In many cases, order of magnitude results are all that are needed for preliminary assessments of a polluted atmosphere. Greater

accuracy could be achieved by longer paths and longer integration times, provided that the instrument is stable against long period drift and the sample is stationary.

Even at S/N = 10, the errors may be kept low by choosing regions where the spectrum is strong. It is certainly true, however, that as S/N becomes much less than 10, the accuracy will decrease significantly.

Table 3.4 may be compared with T3.2 for the effect of noise on concentrations calculated in the spectral range from $2100-2300 \text{ cm}^{-1}$. The gases (Set 2) H_2O , CO_2 , O_3 , N_2O , and CO are present in the simulated sample (Codes 9999 (a-d)). Here conclusions parallel those of the above discussion; a signal-to-noise ratio more than 10 is required to keep errors below 20 percent, and S/N = 100 produces negligible errors.

The reader should not be perturbed because a few low S/N cases appear more accurate than those with higher or infinite S/N. This occurs for two reasons: (1) the noise added may accidently provide an occasional better fit; and (2) the repetition of the algorithm seeking the best fit is at the discretion of the operator. In the case of low S/N, the search was allowed to proceed for more iterations than in the less noisy cases in order to provide the best opportunity of overcoming the noise.

3.1.3 Effects of "Maverick" Gases in the Sample

In analysis of a spectrum from a specific geographical location, it is important to have in the library of spectra, utilized in the computer analysis, a spectrum of each of the gases expected in the area. However, the library should be kept as small as possible to minimize computer storage problems. Thus, the library data entered into the computer memory are chosen to match as closely as possible the set of gases that are in the atmosphere under measurement. For a larger than necessary library (i.e., some gases in the library are not in the test atmosphere), the only important penalties are increased computer memory required and the additional costs due to the necessity of breaking the frequency spectrum into separately treated sections, each run separately on the computer, if the limited computer memory requires it.

To test the effect of having gases present in the sample, but not present in the computer library (a "maverick" gas), runs were made on both previously used sets of gases. From each set of calculations, spanning for Set 1, from 1230 to 1400 cm⁻¹ in nine steps, each gas, except H₂O, was successively deleted from the library (T3.5a,b). Similar procedures were followed in the 2100 to 2300 cm⁻¹ range (T3.6).

The gas deleted from the library was included in the simulated data, but was not among those which the computer attempted to fit to the sample data by best choice of concentration. It thus acted as a localized noise or perturbation of the spectrum.

Comparing T3.1 with T3.5 (a,b) and T3.2 with T3.6, we see that appreciable errors may arise by having an insufficient library. The greatest of these appear where the spectrum of the maverick gas overlaps a weak spectrum of a gas whose spectrum is in the library. This result indicates that as a practical measure it is better to add a gas to the library if its presence in the sample is uncertain.

An examination of the spectra of Sets 1 and 2 in F1.1 (a-c) and F3.1 shows that the spectral structure must act as an appreciable noise background where the gas is a maverick. The MINMYZD method, by utilizing goodness of fit over a wide spectral region, partially eliminates this noise effect.

When a gas spectrum is present in the library, but near or at zero concentration in the sample, the calculation occasionally produces negative concentrations, small in absolute value. They could be prevented by a simple change in the program, but it was judged preferable not to affect the convergence process by putting a barrier at zero concentrations. Such negative concentrations should be considered equal to zero. This situation is discussed in A2.5.

Table 3,5a

GAS CONCENTRATIONS FROM SIMULATED COMPOSITE SPECTRA (SET 1)

	Wavenumb er Range	H_2O () $Wg = 3.73$	1A1) × 10 ²²	N ₂ O (4 Wg = 1.08	4A1) × 10 ^{1 8}	CH w (5A1) x 10 ¹⁹	$SU_2 (20)$ $Wg = 3.54 g$	0.2) K 10 ¹⁷	HNO ₃ (30 Wg = 3.54 x).2) 10 ¹ 7		ansmission ror
Code	(cm-1)	Wc	ΔW	Wc	ΔW	Wc	ΔW	Wc	ΔW	Wc	ΔW.	Wtd.	Unwtd.
8888 a,B	1230.0-1249.8	3.78x10 ²²	-0.05			1.10x10 ¹⁹	02	17.88×10 ¹⁷	-14.34	- 0.07x10 ¹⁷	3.61	0.35	7.39
b,B	1250.0-1269.8	4.00x10 ²²	_0,27			1.18x10 ¹⁹	10	17.88x10 ¹⁷	-14.34	24.58x10 ¹⁷ -	21.04	0.74	12.73
c,B	1270.0-1289.8	3.92x10 ²²	_0.19	N ₂ O delete	l from	1.02x10 ¹⁹	06	17.88x10 ¹⁷	-14.34	8.10x10 ¹⁷ -	4.56	1.76	7.52
d,B	1290.0-1309.8	4.04x10 ²²	-0.31	library for	runs	1.34x10 ¹⁹	-,26	- 0.18x10 ¹⁷	3.72	5.81x10 ¹⁷ -	2.27	0.60	2.19
e,B	1310.0-1329.8	3.69x10 ²²	0.04	8888a,B th	rough	1.31x10 ¹⁹	23	0.42x10 ¹⁷	3.12	4.85x10 ¹⁷ -	1.31	0.11	1.70
f,B	1330.0-1349.8	3.73x10 ²²	0.00	8888i,R		1.08x10 ¹⁹	, 00	3.51x10 ¹⁷	0.03	3.56x10 ¹⁷ -	0.02	0.02	0.02
g,B	1350.0-1369.8	3.73x10 ²²	0.00			1.06x10 ¹⁹	.02	3.32x10 ¹⁷	0.22	4.32x10 ¹⁷ -	0.78	0.03	0.41
h,B	1370.0-1389.8	3.68x10 ²²	0.05			1.48x10 ¹⁹	40	4.88x10 ¹⁷	- 1.34	18.38x10 ¹⁷ -	14.84	0.03	7.45
i,B	1380.2-1400.0	3.67x10 ²²	0.06			1.48x10 ¹⁹	40	5.10x10 ¹⁷	- 1.56	18.23x10 ¹⁷ -	14.69	0.03	7.39
\$888 a,B1	1230.0-1249.8	4.66x10 ²²	-0.93	8.51x10 ¹⁸	-7.43			17.88x10 ¹⁷	-14.34	25.61x10 ¹⁷ -	21.77	7.19	13.46
b,B1	1250.0-1269.8	3.3\columbia)22	0.34	4.85x10 ¹⁸	-3.77			17.8\$x10 ¹⁷	-14.34	-17.19x10 ¹⁷	21.44	7.42	13.
c,B1	1270.0-1289.8	2.90x10 ²²	0.83	-0.59x10 ¹⁸	1.67	CH, deleted	from	17.86x10 ¹⁷	-14.34	17	21.51	7.79	12.5
d,Bl	1290.0-1309.8	2.06x10 ²²	1.67	5.52x10 ¹⁸	-4.44	library for		- 1.12x10 ¹⁷	4.66	- 2.27x10 ¹⁷	5.81	10.36	4.42
•,B1	1310.0-1329.8	4.5€€10 ²²	- 0.83	3.71x10 ¹⁸	-2.63	8888a,B1 th	rough	10.34x10 ¹⁷	- 6.80	- 0.27x10 ¹	3.81	0.72	4.15
f,Bl	1330.0-1349.8	4.09x10 ²²	-0.36	1.21x10 ¹⁸	-0.13	8888i,B1		0.32x10 ¹⁷	3.22	9.18x10 ¹⁷ -	5.64	₹.71	3.29
g,B1	1350.0-1369.8	4.33x10 ²²	-0.60	1.21x10 ¹⁸	-0.13			•.96x10 ¹⁷	2.58	24.01x10 ¹⁷ -	20.47	€. 74	10.32
h,Bl	1370.0-1389.8	3.68x10 ²²	0.05	1.21x10 ¹⁸	-0.13			4.88x10 ¹⁷	- 1.34	18.38x10 ¹⁷ -	14.84	6.03	7.48
i,B1	1380.2-1400.0	3.67x10 ²²	0.06	1.21x10 ¹⁸	-0.13			5.10x10 ¹⁷	- 1.56	18.23x10 ¹⁷ -	14.69	0.03	7.39

Wg = given concentration in simulated sample, molecules cm^{-2}

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMYZD to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1.3).

Wc = calculated concentration, molecules cm^{-2}

ΔW = Wg-Wc (the factor of a power of ten is omitted), molecules cm⁻²

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

Table 3.5b

	Wavenumber Range	H_2O (1) $Wg = 3.73$	A1) x 10 ²²	N_2O (4) Wg = 1.08	IA1) × 10 ¹⁸	CH և (Wg = 1.08	(5A1) x 10 ¹⁹	SO_2 (2) Wg = 3.54	0.2) x 10 ¹⁷	HNO_3 (3) $Wg = 3.54$	30.2}		Transmission Error
Code	(cm ⁻¹)	W'c	ΔW	Wc	ΔW	Wc	∆W	Wc Wc	ΔW.	Wc_	Δ₩	Wtd.	Unwtd.
8888 a,B2	1230.0-1249.8	3.73x10 ²²	.00	1.09x10 ¹⁸	-0.01	1.08x10 ¹⁹	.00			0.07×10^{17}	3.47	0.03	1.74
b,B2	1250.0-1269.8	3.73x10 ²²	.00	1.09x10 ¹⁸	-0.01	1.08x10 ¹⁹	.00			0.10×10^{17}	3.44	0.06	1.72
c,B2	1270.0-1289.8	3.73×10^{22}	.00	1.08x10 ¹⁸	0.00	1.08x10 ¹⁹	.00	SO, deleted	from	3.55×10^{17}	- 0.01	0.03	0.01
d,B2	1290.0-1309.8	3,73x10 ²²	.00	1.08x10 ¹⁸	0.00	1.08x10 ¹⁹	.00	library for		3.54×10^{17}	0.00	0.03	0.00
e,B2	1310.0-1329.8	3.64x10 ²²	. 09	0.31x10 ¹⁸	0.77	1.16x10 ¹⁹	08	8888a,B2, t	hrough	5.19×10^{17}	- 1.65	0.11	0.91
f,B2	1330.0-1349.8		32	1.21x10 ¹⁸	-0.13	1.06x10 ¹⁹	. 02	8888i,B2		4.50×10^{17}	- 0.96	0.36	0.51
g,B2		22	35	1.21x10 ¹⁸	-0.13	1.09x10 ¹⁹	01	•		7.82×10 ¹⁷	- 4.28	0.06	2.15
h,B2	1370.0-1389.8	4.04x10 ²²	31	1.21x10 ¹⁸	-0.13	1.48x10 ¹⁹	40			18.25x10 ¹⁷	-14.71	0.02	7.36
i,B2	1380.2-1400.0	4.04x10 ²²	31	1.21x10 ¹⁸	-0.13	1.48x10 ¹⁹	40			18.23×10 ¹⁷	-14.69	0.02	7.35
8888 a,B3	1230.0-1249.8	3.73x10 ²²	.00	1.09x10 ¹⁸	- 0. 01	1.08x10 ¹⁹	.00	17.88x10 ¹⁷	-14.34		·	0.03	7.17
b,B3	1250.0-1269.8		.00	1.09x10 ¹⁸	-0.01	1.09x10 ¹⁹	.00	17.88x10 ¹⁷	-14.34			0.06	7.17
c,B3	1270.0-1289.8	4.05x10 ²²	32	1.32x10 ¹⁸	- 0, 24	1.22x10 ¹⁹	14	17.88x10 ¹⁷	-14.34	HNO, deleted	from	0.72	7.17
d,B3	1290.0-1309.8	3.5Ux10 ²²	.20	3.02x10 ¹⁸	-1.94	0.86x10 ¹⁹	.22	14.24x10 ¹⁷	-10.70	library for	runs	1.76	5.44
e,B3	1310.0-1329.8	4.34x10 ²²	61	2.93x1n ¹⁸	-1.85	0.82×10^{19}	.26	11.80x10 ¹⁷	- 8.26	8888a,B3 thi	ough	0.19	4.25
f,B3	1330.0-1349.8		27	1.21x10 ¹⁸	- 0.13	1.42x10 ¹⁹	34	10.10x10 ¹⁷	- 6.56	8888i,B3		0.81	3.29
g,B3	1350.0-1369.8	3.64x10 ²²	.09	1.21x10 ¹⁸	- 0.13	1.07x10 ¹⁹	.01	6.14x10 ¹⁷	- 2.60			0.04	1.30
h,B3	1370.0-1389.8		. 05	1.21x10 ¹⁸	- ,13	1.48x10 ¹⁹	40	4.86x10 ¹⁷	- 1.32			0.03	0.69
i,B3	1380.2-1400.0		.06	1.21x10 ¹⁸	13	1.48x10 ¹⁹	40	5.10×10^{17}	- 1.56			0.03	0.81

Wg = given concentration in simulated sample, molecules cm⁻²

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMYZD to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1.3).

Wc = calculated concentration, molecules cm⁻²

ΔW = Wg-Wc (the factor of a power of ten is omitted), molecules cm⁻²

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

Table 3.6
GAS CONCENTRATIONS FROM SIMULATED COMPOSITE SPECTRA (SET 2)
(MAVERICK GASES)

	Wavenumber Range	H_2O (1) $Wg = 3.73$	B1)	CO_2 () Wg = 1.08	2B1)	O ₃ (Wg = 2.69	(3B1)	N_2O (4 Wg = 1.08	B1)	CO (6 Wg = 1.88	5B1)		ansmission ror
Code	(cm ⁻¹)	Wc Wc	ΔW	Wc	ΔW	Wc		Wc	^ ^∆W	Wc Wc	ΔW	Wtd.	Unwtd.
9999 a,A	2100.0-2159.4	3.65x10 ²¹	0.08	CO, delete	d from	5.07x10 ¹⁷	- 2.38	0.64x10 ¹⁸	0.44	1.93x10 ¹⁹	- 0.05	0.43	1.21
b,A	2160.0-2219.4	3.73x10 ²¹	0.00	library for		5.94x10 ¹⁷		1.08x10 ¹⁸	0.00	1.88x10 ¹⁹	0.00	0.03	1.63
c.A	2220.0-2279.4	49.64x10 ²¹	-45.91	9999a,A th	rough	5.94x10 ¹⁷		0.93x10 ¹⁸	0.15	1.78x10 ¹⁹	0.10	24.22	23.01
d,A	2240.4-2299.8		- 8.01	-	•	4.40x10 ¹⁷		0.51x10 ¹⁸	0.57	2.98x10 ¹⁹	- 1.10	63.50	4.14
9999 a,Al	2100.0-2159.4	3.68x10 ²¹	0.05	1.29x10 ²¹	- 0.21	O ₃ deleted	from	0.81x10 ¹⁸	0.27	1.91x10 ¹⁹	- 0.03	0.20	0.17
b,Al	2160.0-2219.4	3.72x10 ²¹		1.05x10 ²¹	0.03	-		1.08x10 ¹⁸	0.00	10	0.00	0.03	0.02
c,A1	2220.0-2279.4	3.68x10 ²¹	0.05	1.08x10 ²¹	0.00	9999a,A1 th	rough	1.08x10 ¹⁸	0.00	1.88x10 ¹⁹	0.00	0.03	0.03
d,Al	2240.4-2299.8	3.69x10 ²¹		1.08x10 ²¹	0.00	9999d,A1		1.08x10 ¹⁸	0.00	2,98x10 ¹⁹	- 1.10	0.03	0.55
9999 a,A2	2100.0-2159.4	3.72x10 ²¹	0.01	1.05x10 ²¹	0.03	2.64×10 ¹⁷	0.05	N ₂ O deleted	from	1.89x10 ¹⁹	- 0,01	0.04	0,03
b,A1	2160.0-2219.4	1.58x10 ²¹	2.15	5.76x10 ¹²	- 4.68	5.94x10 ¹⁷	- 3.25	library run		3.88x10 ¹⁹	- 2.00	24.13	3.20
c,A2	2220.0-2279.4	81 24x10 ²¹	-77.51	21	0.28	17	- 3.25	9999a,A2 th	rough	14.11x10 ¹⁹	-12.23	33.67	39.27
d,A2	2240.4-2299.8	11.74x10 ²¹	- 8.01	0.92x10 ²¹	0.16	4.40x10 ¹⁷	- 1.71	9999d,A2		2.98x10 ¹⁹	- 1.10	29.44	4.13
9999 a,A3	2100.0-2159.4	12.55x10 ²¹	- 8.82	11.11x10 ²¹	-10.03	45.48x10 ¹⁷	-42.79	44.14x10 ¹⁸	-43.06	CO deleted	from	16.55	31.08
b,A3	2160.0-2219.4	36.19x10 ²¹		$-5.34x10^{21}$	6.42	5.94x10 ¹⁷	- 3.25	1.79x10 ¹⁸	- 0.71	library for	runs	18.60	16.63
c,A3	2220.0-2279.4	3.59x10 ²¹	0.14	1.08×10^{21}	0.00	5.94x10 ¹⁷	- 3.25	1.14x10 ¹⁸	- 0.06	9999a,A3 th	rough	2.06	1.63
d,A3	2240.4-2299.8	3.61x10 ²¹	0.12	1.08x10 ²¹		5.94x10 ¹⁷	- 3.25	1.08x10 ¹⁸	0.00	5A, beeee		0,03	1.63

Wg = given concentration in simulated sample, molecules cm^{-2}

A composite sample spectrum was obtained by adding the absorption coefficients as a function of frequency and calculating the transmission (See 1.2). The sample spectrum was then analyzed by MINMYZD to obtain the calculated concentrations. The error is the difference between the calculated and given transmission. (See 3.1.3).

Wc = calculated concentration, molecules cm⁻²

ΔW = Wg-Wc (the factor of a power of ten is omitted), molecules cm-2

See 3.1 for definition of the weighted and unweighted final transmission errors (last two columns). Numbers following gas formulae are computer codes.

4. CALCULATION OF THE SELECTED GAS SAMPLE CONCENTRATIONS FROM SPECTROPHOTOMETER DATA

The results presented in 3 were obtained from simulated data samples produced by calculating composite spectra from selected gas spectra for specified concentrations. While these are an adequate test of the computer program, they do not test the overall system of spectrophotometer and computer algorithm. Accordingly, composite spectra were prepared by Dr. Wm. Herget, EPA, Research Triangle Park, NC, on the ROSE* system described in 1.1 and A3. These data were recorded on digital magnetic tape, and this information became the input to the EPAGAS program described in A2. In addition, for comparison, some data were obtained on a Fourier transform spectrometer (FTS).

The data were not obtained from a long atmospheric path because of the difficulty of obtaining, in that case, an independent measure of the gaseous concentrations. Instead, a cell containing approximately measured concentrations of selected gases was placed in the path. The data were normalized, approximately, by removing the cell and running over the same spectral range to determine the spectra of ambient H_2O and CO_2 .

Several problems arose in connection with normalization and wavelength calibration. These are discussed in 1.1 and A2.5, respectively. The result of normalization and wavelength uncertainties in the spectrophotometer output and the added effect of mismatched resolutions of the spectrophotometer and the spectra library result in some small inaccuracies in the calculated concentrations. These problems can all be solved by better calibration methods, and effort is underway to do so. The ROSE spectrophotometer resolution is a function of wavelength, which is characteristic of grating spectrometers. It can be prevented by a non-linear slit drive mechanism or by using a Fourier Transform Spectrometer (FTS). The FTS can be expected to have a linear frequency scale, but the

^{*}Remote Optical Sensing of Emissions.

data may need a shift of frequency to correct for an uncertain zero calibration. The ROSE spectrophotometer data and the data from the FTS correspond well in general. However, in a number of regions, the correspondence is sufficiently poor to cause errors in calculated concentrations. Data with such discrepancies may be seen by comparing F4.1, F4.2, and F4.3 with F4.9 (a-c).

However, in spite of these difficulties, the computational method performs well, and as shown below, the results of the calculation are of good accuracy when precautions are taken to calibrate the data.

Several calculations from experimental data are described in the following sections, utilizing different gases in different frequency ranges.

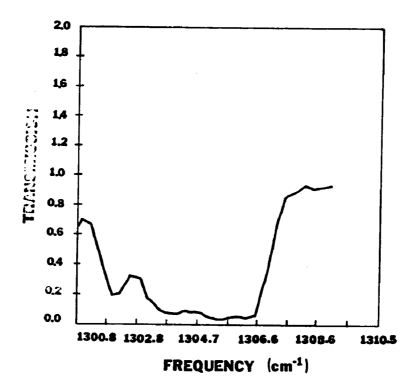


Figure 4.1

SPECTRUM OF SO₂ AND CH₄ OBTAINED BY FTS (1300.8-1310.5 cm⁻¹)

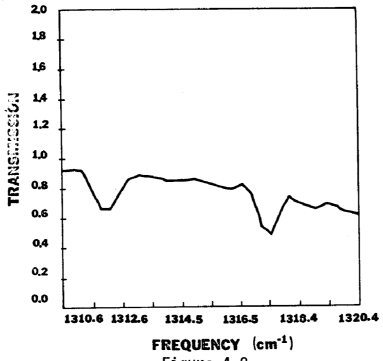


Figure 4.2

SPECTRUM OF SO₂ AND CH₄ OBTAINED BY FTS (1310.6-1320.4 $\rm cm^{-1}$)

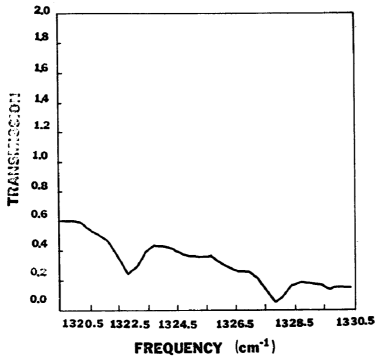


Figure 4.3

SPECTRUM OF SO₂ AND CH₄ OBTAINED BY FTS (1320.5-1330.5 cm⁻¹)

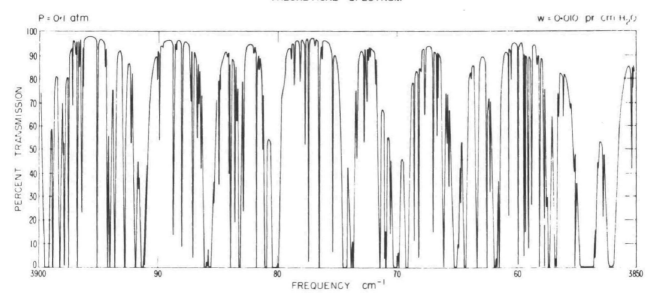
4.1 EFFECT OF MISMATCHED RESOLUTION OF EXPERIMENTAL DATA AND LIBRARY SPECTRA

The resolution of spectroscopic instruments has a large effect on the appearance of molecular spectra. It is well known that instrumental resolution can change the shape of a theoretical spectrum so thoroughly that recognition is very difficult. Appearance of the spectrum is of great importance in any best-fit program because it compares two curves, the one obtained from the spectral library with the one from the data. Figure 4.4 shows an example of a theoretical spectrum degraded by an instrumental bandwidth (Calfee, 1966).

The gross changes of recognizable features seen in F4.4 are not usually a problem in spectral analysis of pollutants because the mismatch in resolution is not as large. However, considerable error in analysis can be introduced by such mismatch. Figure 4.5 is a position of the reconstructed spectrum of carbon monoxide reproduced from data obtained on the ROSE system. The concentration of the sample was 1.5×10^{19} molecules cm⁻² and the spectral slit width was 3 cm⁻¹ according to the accompanying data. Difficulty was encountered in analyzing the spectrum, hence a series of computed spectra were made with various amounts of gas and with different slit widths. Figure 4.6 was computed using a spectral slit width of 3 cm⁻¹ and with 1.5×10^{18} , 7.5×10^{18} , and 1.5×10^{19} molecules cm⁻² to see if a different amount of gas in the cell could be the cause of the discrepancy. This was not the case because none of the computed spectra match the data.

Next, a calculation was made with the same quantities of gas, but with a narrower slit width (2.75 cm^{-1}) . Figure 4.7 shows the result of this calculation. This result approaches closer to the experimental spectrum. Figure 4.8 was made by using 2.3 cm⁻¹ as the spectral slit width. Here the agreement between the computed spectrum and the experimental curve is very good for a concentration of 1.5 x 10^{19} molecules cm⁻². This analysis points up the significance of the spectral slit width in the structure of the spectrum and the best-fit procedure. (The slit width as a function of frequency is shown in A3.)

THEORETICAL SPECTRUM



DEGRADED SPECTRUM

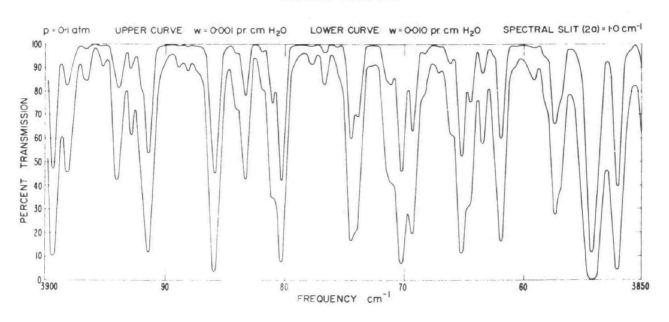
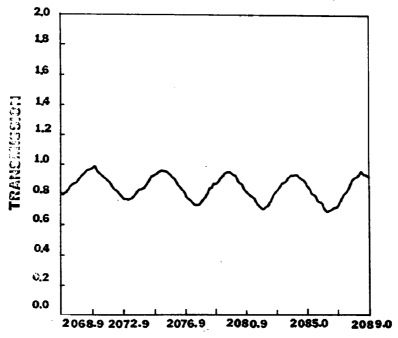


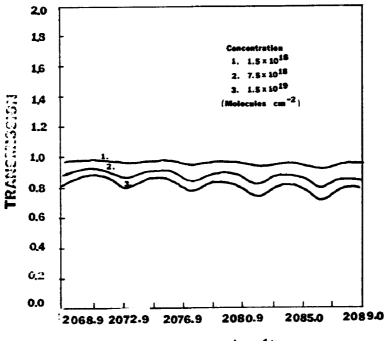
Figure 4.4

COMPARISON OF A THEORETICAL SPECTRUM (HIGH RESOLUTION) OF WATER VAPOR AND THE SPECTRUM OBSERVED THROUGH A SPECTROMETER WITH 1.0 cm $^{-1}$ SLIT WIDTH (IN MANY REGIONS THE COMPARISON IS VERY DIFFICULT)



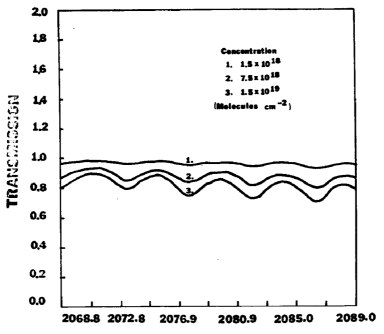
FREQUENCY (cm⁻¹)
Figure 4.5

EXPERIMENTAL SPECTRUM, CO, NOMINAL RESOLUTION 2.5 cm⁻¹ OBTAINED ON ROSE SYSTEM (Compare with F4.6)



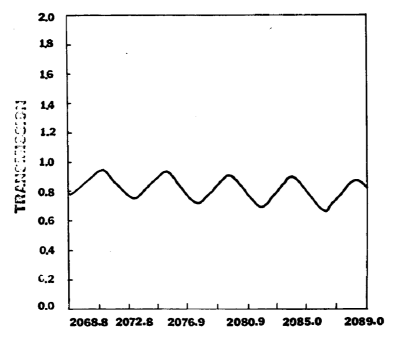
FREQUENCY (cm⁻¹)

Figure 4.6
CALCULATED SPECTRUM, CO,
RESOLUTION 3.0 cm⁻¹
(Compare with F4.5)



FREQUENCY (cm⁻¹)
Figure 4.7

CALCULATED SPECTRUM, CO, RESOLUTION 1.0 cm⁻¹ (Compare with F4.5)



FREQUENCY (cm⁻¹)
Figure 4.8

CALCULATED SPECTRUM, CO, RESOLUTION 2.3 cm⁻¹ (Compare with F4.5) The spectra shown in F4.9 (a-c) were reconstructed from EPA ROSE data of a combination of SO_2 (7.56 x 10^{18} molecules cm⁻²) and CH_4 (1.23 x 10^{19} molecules cm⁻²) with a spectral slit width of 1 cm⁻¹. Figure 4.10 (a-c) shows the computed spectra for the same regions with the same amounts of gas and the same slit width used for the calculations. The most obvious disagreement appears between the curves of F4.9b (experimental) and F4.10b. It would appear that some gas other than SO_2 and CH_4 (or noise) (see 4.4) was in the sample to produce the additional structure. An examination of the entire series shows a general agreement, but again the experimental curves show an indication of a higher resolution than that used for the computed spectra. It would also appear that the use of a lower value of concentration would produce a computed spectra which would come closer to matching the experimental curves. Probably both a lower concentration and a higher resolution would produce a spectrum which would agree with the experimental spectrum.

This discussion illustrates the necessity for an accurate normalization procedure and precision in determining instrumental resolution to avoid ambiguity in spectral analysis. The resolution of spectrometers generally changes over their spectral range. This makes it necessary that either the experimenter or the analyst must be able to compensate for the effect.

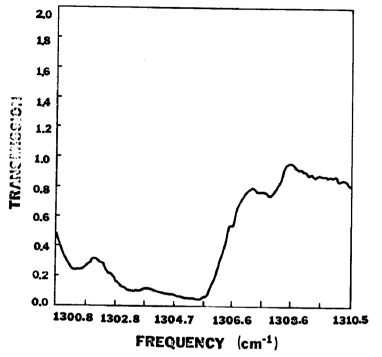
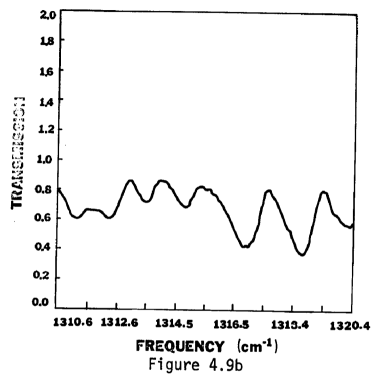
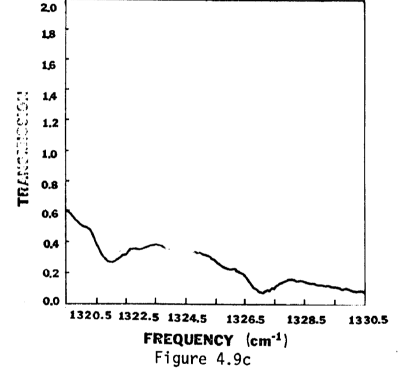


Figure 4.9a

EXPERIMENTAL SPECTRUM OF COMBINATION OF SO2 AND CH₄, NOMINAL SLIT WIDTH 1.0 cm⁻¹ (ROSE SYSTEM) (1300.8-1310.5 cm⁻¹) (Compare with F4.10a,b,c)



EXPERIMENTAL SPECTRUM OF COMBINATION OF SO₂ AND CH₄, NOMINAL SLIT WIDTH 1.0 cm⁻¹ (ROSE SYSTEM) (1310.6-1320.4 cm⁻¹) (Compare with F4.10a,b,c)



EXPERIMENTAL SPECTRUM OF COMBINATION OF SO₂ AND CH₄, NOMINAL SLIT WIDTH 1.0 cm⁻¹ (ROSE SYSTEM) (1320.5-1330.5 cm⁻¹) (Compare with F4.10a,b,c)

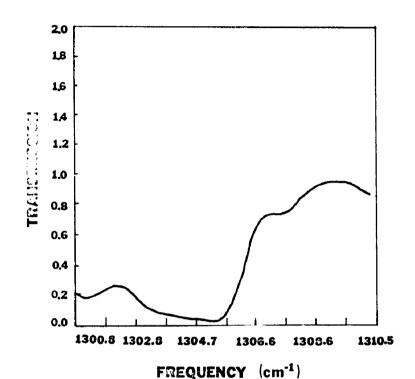


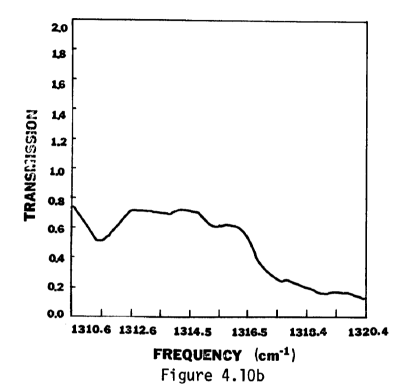
Figure 4.10a

CALCULATED SPECTRUM, SO₂ AND CH₄

(RESOLUTION 1.0 cm⁻¹)

(1300.8-1310.5 cm⁻¹)

(Compare with F4.9a,b,c)



CALCULATED SPECTRUM, SO₂ AND CH₄
(RESOLUTION 1.0 cm⁻¹)
(1310.6-1320.4 cm⁻¹)
(Compare with F4.9a,b,c)

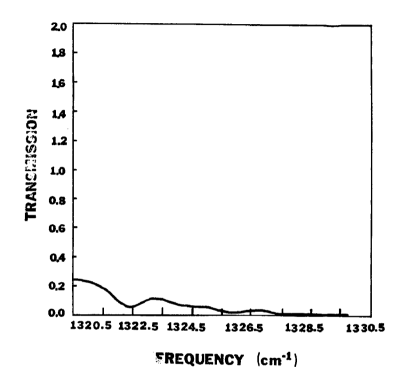


Figure 4.10c

CALCULATED SPECTRUM, SO₂ AND CH₄

(RESOLUTION 1.0 cm⁻¹)

(1320.5-1330.5 cm⁻¹)

(Compare with F4.9a,b,c)

4.2 CALCULATION OF THE CONCENTRATION OF CO

A composite spectrum of H_2O , CO_2 , CO_2 , CO_4 , and SO_2 was prepared on the ROSE spectrophotometer over the spectral region from 1250 to $1330 \, \mathrm{cm}^{-1}$ and in the range 2050 to 2150 cm^{-1} . Because only a relative intensity was obtained, normalization proceeded by obtaining a background run containing only spectra of H₂O and CO₂. The ratio of the two spectra gave the normalized spectrum I/I. The EPAGAS program was then used to calculate the concentration of CO in three spectra regions, 2073-2094. 2094-2114, and 2114-2132 cm⁻¹. T4.1 shows the results of the calculation. The spectral regions are listed in the second column. Several initial trial concentrations were used to determine the mathematical stability of the process. As can be seen for each region, different starting estimates made little change in final calculated concentrations. However, as discussed in 4.1, the resolution used in the spectral library strongly affects the accuracy. Part of the calculations summarized in T4.1 used the nominal spectral library resolution (3.0 cm⁻¹) for this region. However, as pointed out in 4.1, the resolution used in preparing the data was actually closer to 2.3 cm⁻¹. Nine values of concentration were calculated assuming a resolution of 2.3 cm⁻¹. These values average 1.26 x 10^{19} molecules cm^{-2} ; the sample concentration measured chemically was 1.5×10^{19} molecules cm⁻². It is believed that this agreement is excellent in view of the calibration and normalization problems.

Note that the spectral resolution was carefully determined in the region $2073-2094~{\rm cm}^{-1}$ and that the errors are the least there, increasing as the calculation is performed for more distant regions.

By using program DEGRADE (Deutschman and Calfee, 1967) before the fitting routine in EPAGAS, the actual resolution of the sample was determined to be 2.3 cm⁻¹. This choice of resolution was verified by the results given in T4.1.

Table 4.1

CALCULATION OF THE CONCENTRATION OF A SAMPLE OF CO BY EPAGAS TO AN EXPERIMENTAL SPECTRUM AT SEVERAL SPECTRAL REGIONS AND TWO INSTRUMENTAL RESOLUTIONS

	Region	Frequency Correction		CO Trial (Est.) Concentration	CO Calculated Concentration	Final	Averages f Given Resc and Regior Cal.	lution
Code	Spectral	Δν	Resolution	(molecules cm ⁻²)	(molecules cm ⁻²)	Transmissions	· Conc.	Error
la	2073.0-2094.0	0.0	3.0	1.50x10 ¹⁹	7.33×10^{18}	2.51		
b	2073.0-2094.0	0.0	3.0	1.88x10 ¹⁹	7.32x10 ¹⁸	2,51	7.33×10^{18}	2.51
c	2073.0-2094.0	0.0	2.3	5.38x10 ¹⁸	1.18x10 ¹⁹	1.25		
d	2073.0-2094.0	0.0	2.3	1.50x10 ¹⁹	1.18x10 ¹⁹	1.25		
c	2073.0-2094.0	0.0	2.3	1.88x10 ¹⁸	1.18x10 ¹⁹	1.25	1.18x10 ¹⁹	1.25
2a	2094.0-2114.0	0.0	3.0	1.50x10 ¹⁹	9.27x10 ¹⁸	2.24		
b	2094.0-2114.0	0.0	3.0	1.88x10 ¹⁸	$9.29 \text{x} 10^{18}$	2.24	9.28×10^{18}	2.24
c	2094.0-2114.0	0.0	2.3	5.38x10 ¹⁸	1.26x10 ¹⁹	2.18		
d	2094.0-2114.0	0.0	2.3	1.50x10 ¹⁹	1.26x10 ¹⁹	2.18		
e	2094.0-2114.0	0.0	2.3	1.88x10 ¹⁹	1.26x10 ¹⁹	2.18	1.26x10 ¹⁹	2.18
3a	2114.0-2134.0	0.0	3.0	1.50x10 ¹⁹	8.71x10 ¹⁸	1.96		
b	2114.0-2134.0	0.0	3.0	1.88x10 ¹⁹	8.71x10 ¹⁸	1.96	8.71×10^{18}	1.96
c	2114.0-2134.0	0.0	2.3	5.38×10 ¹⁸	1.35x10 ¹⁹	2.19		
d	2114.0-2134.0	0.0	2.3	1.50x10 ¹⁹	1.35x10 ¹⁹	2.19		
e	2114.0-2.34.0	0.0	2.3	1.88x10 ¹⁹	1.35x10 ¹⁹	2.19	1.35x10 ¹⁹	2.19

Average CO Concentration and Error for 3.0 resolution: 8.44x10¹⁸; Error 2.24.

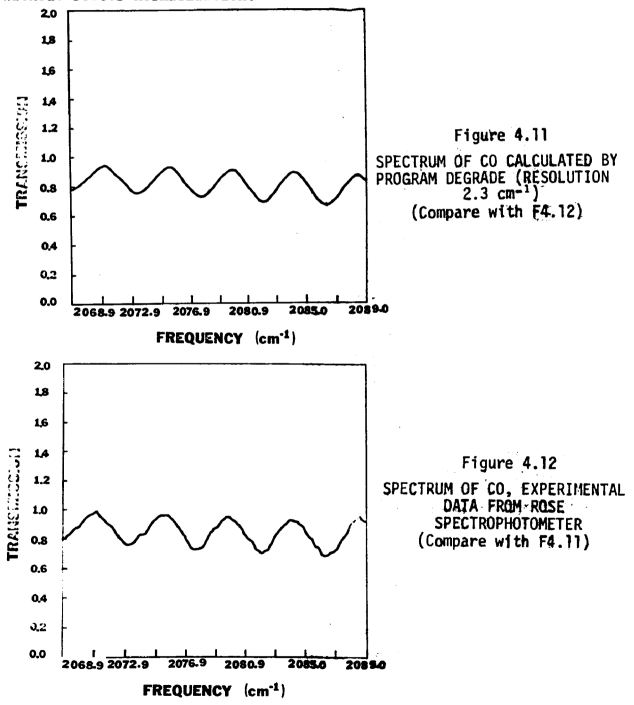
Average CO Concentration and Error for 2.3 resolution: 1.26x10¹⁹; Error 1.87.

Sample Concentration: $1.50 \times 10^{18} \text{ mol cm}^{-2}$ (Obtained by Chemical Analysis).

The calculated concentration with the least error (1.26×10^{19}) compares well with the sample concentration (1.50×10^{18}) .

NOTE: Powers of ten suppressed in errors quoted above. Concentrations are in molecules cm-2.

Figures 4.11 and 4.12 show, respectively, the data produced by the ROSE spectrophotometer and the best fitting calculated spectrum. The slight differences in this case are probably due to noise. Although the shapes of these spectra match quite well, the absolute transmission differences account for the slight discrepancy in calculated and experimental concentration. These absolute transmission differences are unresolvable without better normalization.



The effect of resolution on the goodness of fit in EPAGAS may be seen in F4.13. In the first of these, the dashed curve represents the given experimental data, the full line is the calculated value. The results of the calculation are given in (resolution 3.0 cm⁻¹) T4.1. These curves should be contrasted with the first calculation listed in T4.1 using 2.3 cm⁻¹ (line 7), shown in F4.14. The much better fit of given and calculated data results in a more accurate value of concentration and once again shows the importance of resolution matching.

The efficiency of the program in performing a "best fit" may be seen in F4.14. Notice that the two curves are not sufficiently alike to be fitted exactly. However, the optimization of the adjustable parameters of the calculated spectrum has clearly resulted in an excellent fit in a least-squares sense. An understanding of the change of a typical spectrum with a small change of concentration may be seen in F4.15. Clearly good accuracy and low noise of the instrument data are required for accurate analysis.

TRANSMISSION

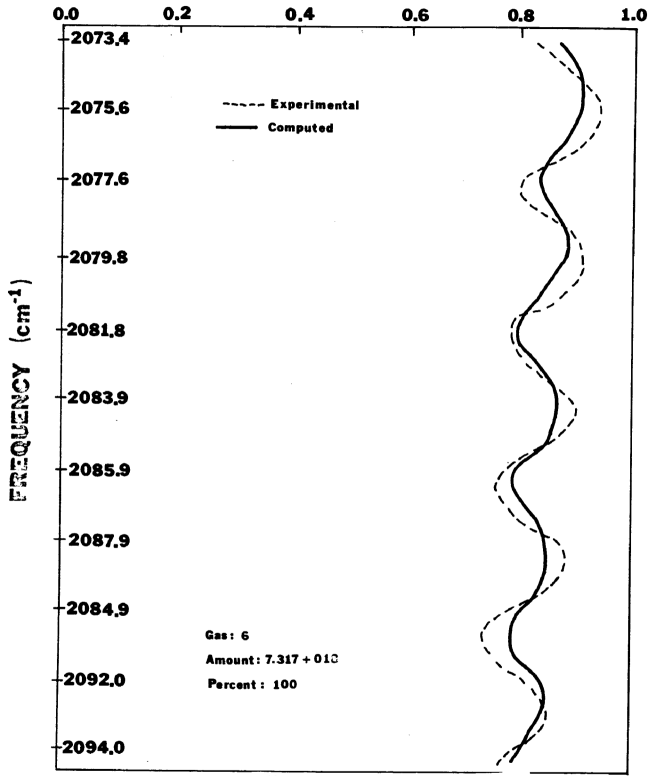


Figure 4.13

COMPUTED BEST-FIT TRANSMISSION OF CO (EPAGAS PROGRAM) RESOLUTION 3.0 \mbox{cm}^{-1}

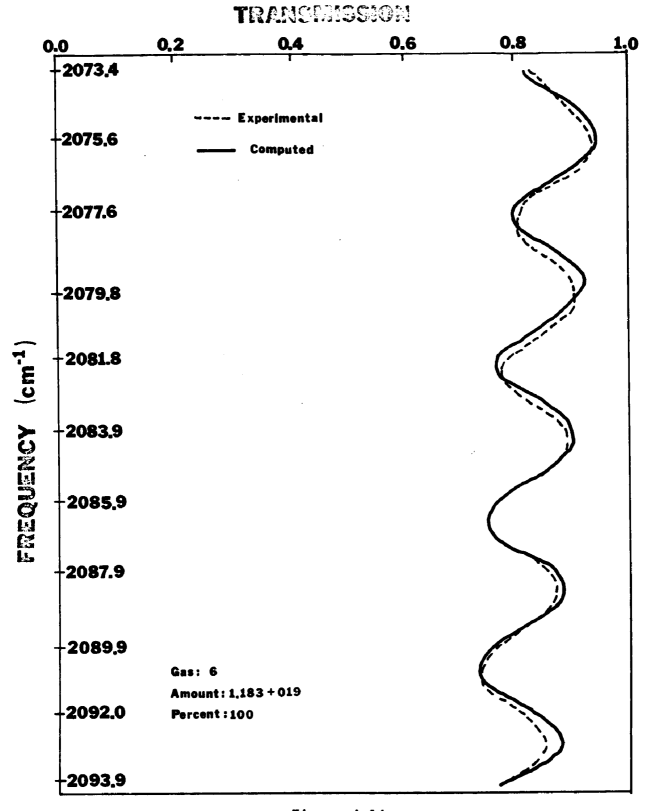


Figure 4.14

COMPUTED BEST-FIT TRANSMISSION OF CO (EPAGAS PROGRAM)

RESOLUTION 2.3 cm⁻¹

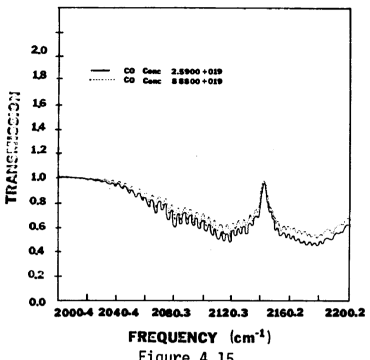


Figure 4.15 IN SPECTRUM O

CHANGE IN SPECTRUM OF CO FOR SMALL CHANGE OF CONCENTRATION (Calculated by Program DEGRADE)

4.3 CALCULATION OF THE CONCENTRATION OF SO₂ IN THE PRESENCE OF CH₄

The gas sample cell used in obtaining data for CO_2 , as discussed in 4.2, contained also SO_2 and CH_4 . These gases have significant spectra in the region from 1150 to 1420 cm⁻¹ (see F1.1 (a-c)). However, the data have good signal-to-noise ratio only in the ranges 1300-1330 cm⁻¹ and 1250-1300 cm⁻¹. We first consider the 1300-1330 cm⁻¹ range. In this range, the spectrophotometer data shows evidence of incomplete elimination of water spectra or noise, hence the data from the Fourier transform spectrometer were utilized. In this range, the spectral structure of CH_4 does not lend itself to fitting uniquely, and we here seek only to find the concentration of SO_2 in the presence of interference due to CH_4 .

Table 4.2 shows the results of concentration calculations by MINMYZD for several different conditions of initial trial concentration and frequency corrections. The stability of the calculated concentration of SO₂, regardless of starting point, is apparent. The frequency corrections listed for the various regions are linear frequency shifts due to an uncertainty of the zero point in the Fourier transform.

As expected, the concentrations calculated for $\mathrm{CH_4}$ here vary widely and are untrustworthy. However, the presence of $\mathrm{CH_4}$ does not radically affect the value of $\mathrm{SO_2}$ concentration computed in any of the spectral regions. The average value of $\mathrm{SO_2}$ concentration molecules cm^{-2} is very close to the chemically determined concentration in spite of the many uncertainties in the data parameters. It is particularly encouraging to note that the concentration of the sample lay outside the span of library concentrations, but the program EPAGAS, by use of the subroutine FINDTRN, was able to extrapolate properly. The library does not include such large concentrations because it is not expected to be so large in field experiments, even in highly polluted geographical regions.

4.4 CALCULATION OF THE CONCENTRATION OF CH

An attempt was made to analyze the spectra for the concentration of $\mathrm{CH_4}$. The results of this calculation were not as successful as the determination of the concentrations of CO and $\mathrm{SO_2}$. The cause of the discrepancy is either noise or unknown interfering spectra (or both) in the data. The problem is illustrated in F4.16. There we see that the optimum fit of $\mathrm{CH_4}$ to the data still allows large errors because structure is found in the data which does not come from the $\mathrm{CH_4}$ molecule.

The source of the structure very likely lies among the following:

- a. The library spectra of CH_4 is incorrect.
- b. There may be small SO_2 lines from an adjacent band not found in the library.
- c. The sample contained an unknown gas.
- d. The spectrum contains instrumental noise.
- e. Water lines are incompletely cancelled.

Table 4.2 CALCULATION OF THE CONCENTRATION OF THE SAMPLE OF SO_2 IN THE PRESENCE OF CH 4 BY APPLICATION OF EPAGAS TO AN EXPERIMENTAL SPECTRUM

	Spectral	Frequency Correction	Concent (molecul	(Est.) rations es cm ⁻²)	(molecul	rations les cm ⁻²)	Final Transmission	Give a	ages for a n Frequency nd Range	
ode	Region	Δν	CH 4	SO₂	CH4	SO ₂	Error	СНц	SO ₂	Error
a*	1300.0-1310.0	0.0	7.56x10 ¹⁸	1.23x10 ¹⁹	5.44x10 ¹⁸	$9.20x10^{19}$	24.89			
ъ	1300.0-1310.0	0.0	1.23x10 ¹⁹	7.56x10 ¹⁸	6.33x10 ¹⁸	8.37x10 ¹⁸	31.77	6.33x10 ¹⁸	8.37x10 ¹⁸	31.77
с*	1300.0-1310.0	-0.9	7.56x10 ¹⁸	$1.23x10^{19}$	7.98x10 ¹⁸	1.47x10 ¹⁹	18.61			
d	1300.0-1310.0	-0.9	1.23x10 ¹⁹	7.56x10 ¹⁸	7.98×10^{18}	9.47x10 ¹⁸	18.67	7.98x10 ¹⁸	9.47x10 ¹⁸	18.67
e	1300.0-1310.0	-1.2	7.56x10 ¹⁸	1.23x10 ¹⁹	7.98x10 ¹⁸	6.77x10 ¹⁸	14.61	•		
f	1300.0-1310.0	-1.2	1.23x10 ¹⁹	7.56x10 ¹⁸	8.00x10 ¹⁸	7.10x10 ¹⁸	14.61	7.99x10 ¹⁸	6.94x10 ¹⁸	14.61
a*	1310.0-1320.0	0.0	7.56x10 ¹⁸	1.23x10 ¹⁹	-6.54x10 ¹⁷	7.77x10 ¹⁸	7.93			
b *	1310.0-1320.0	0.0	1.23x10 ¹⁹	7.56×10^{18}	-6.54×10^{17}	7.77x10 ¹⁸	7.93	-		
С	1310.0-1320.0	-0.9	7.56x10 ¹⁸	1.23x10 ¹⁹	6.68x10 ¹⁸	6.86x10 ¹⁸	4.87			
đ	1310.0-1320.0	-0.9	1.23x10 ¹⁹	7.56x10 ¹⁸	6.75x10 ¹⁸	6.84x10 ¹⁸	4.87	6.72x10 ¹⁸	6.85x10 ¹⁸	4.8
e	1310.0-1320.0	-1.2	7.56x10 ¹⁸	1.23x10 ¹⁹	7.53x10 ¹⁸	6.90x10 ¹⁸	4.89			
f	1310.0-1320.0	-1.2	1.23x10 ¹⁹	7.56x10 ¹⁸	7.53x10 ¹⁸	6.89×10^{18}	4.89	7.53×10^{18}	6.90x10 ¹⁸	4.89
а	1320.0-1330.0	0.0	7.56x10 ¹⁸	1.23x10 ¹⁹	3.15x10 ¹⁸	6.09x10 ¹⁸	9.98			
ь	1320.0-1330.0	0.0	1.23x10 ¹⁹	7.56×10^{18}	3.15x10 ¹⁸	6.09x10 ¹⁸	9.98	3.15x10 ¹⁸	6.09x10 ¹⁸	9.98
С	1320.0-1330.0	-0.9	7.56x10 ¹⁸	1.23x10 ¹⁹	1.89x10 ¹⁹	6.25x10 ¹⁸	6.42			
d	1320.0-1330.0	-0.9	1.23x10 ¹⁹	7.56x10 ¹⁸	2.27x10 ¹⁹	6.23x10 ¹⁸	6.37	2.08x10 ¹⁹	6.24x10 ¹⁸	6.40
e	1320.0-1330.0	-1.2	7.56x10 ¹⁸	1.23x10 ¹⁹	1.99x10 ¹⁹	6.36x10 ¹⁸	5.69			
f	1320.0-1330.0	-1.2	1.23x10 ¹⁹	7.56x10 ¹⁸	2.65x10 ¹⁹	6.37x10 ¹⁸	5.51	2.32x10 ¹⁹	6.37x10 ¹⁸	5.60

Table 4.2 (cont.)

	Spectral			(Est.) rations es cm ⁻²)	Concent	lated rations les cm-2)	Final Transmission	Averages for a Given Frequency and Range			
Code	Region	Δν	CH 4	S0 ₂	CH 4	SO ₂	Error	CH4	SO ₂	Error	
4a	1300.0-1330.0	0.0	7.56x10 ¹⁸	1.23x10 ¹⁹	5.11x10 ¹⁸	6.54x10 ¹⁸	20.05				
Ъ	1300.0-1330.0	0.0	1.23x10 ¹⁹	7.56×10^{18}	5.18x10 ¹⁸	6.56x10 ¹⁸	20.05	5.15x10 ¹⁸	6.55x10 ¹⁸	20.05	
c	1300.0-1330.0	-0.9	7.56x10 ¹⁸	1.23x10 ¹⁹	8.00x10 ¹⁸	6.84x10 ¹⁸	11.49				
d	1300.0-1330.0	-0.9	1.23x10 ¹⁹		8.00×10^{18}	6.84x10 ¹⁸	11.49	8.00x10 ¹⁸	6.84x10 ¹⁸	11.49	
e	1300.0-1330.0	-1.2	7.56x10 ¹⁸	1.23x10 ¹⁹	8.05x10 ¹⁸	7.04x10 ¹⁸	9.22				
_f	1300.0-1330.0	-1.2	1.23x10 ¹⁹	7.56×10^{18}			9.22	8.05x10 ¹⁸	7.04×10^{18}	9.22	
5a*	1310.0-1330.0	0.0	7.56x10 ¹⁸	1.23x10 ¹⁹			9.77				
b*	1310.0-1330.0	0.0	1.23x10 ¹⁹	7.56x10 ¹⁸	1.91x10 ¹⁸		9.77	-			
С	1310.0-1330.0	-0.9	7.56x10 ¹⁸	1.23x10 ¹⁹	9.62x10 ¹⁸		6.34				
d	1310.0-1330.0	-0.9	1.23x10 ¹⁹	7.56x10 ¹⁸	9.62x10 ¹⁸	6.38x10 ¹⁸	6.34	9.62x10 ¹⁸	6.38x10 ¹⁸	6.34	
е	1310.0-1330.0	-1.2	7.56x10 ¹⁸	1.23x10 ¹⁹	1.06x10 ¹⁹		6.05				
f	1310.0-1330.0	-1.2	1.23x10 ¹⁹	7.56x10 ¹⁸	1.08x10 ¹⁹	6.52x10 ¹⁸	6.05	1.07x10 ¹⁹	6.52x10 ¹⁸	6.05	

Average Concentrations	Resolution	CH _A	so,	Error	
for all Regions	0.0	4.88×10^{18}	7.00×10^{18}	20.60	
	-0.9	6.88x10 ¹⁸	7.16x10 ¹⁸	9.55	
	-1.2	5.39x10 ¹⁸	6.75x10 ¹⁸	8.07	
	1	Λ	7		

Sample Concentrations: $CH_4 = 1.23 \times 10^{19} \text{ molecules cm}^{-2}$ (Obtained by Chemical Analysis) $SO_2 = 7.56 \times 10^{18} \text{ molecules cm}^{-2}$

^{*} Not used in Averages

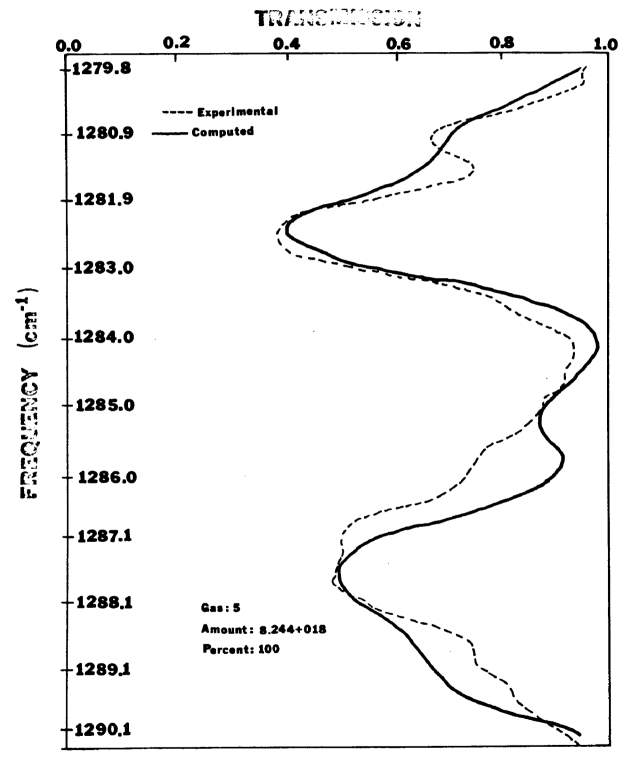


Figure 4.16

COMPUTED BEST-FIT TRANSMISSION OF CH₄
COMPARED WITH THE GIVEN EXPERIMENTAL TRANSMISSION

We comment on these possibilities:

- a. The library spectrum of CH₄ has been checked by many experimenters. It is possible, but considered unlikely, that it is incorrect.
- b. A chemical analysis by EPA of the sample did not indicate any other gases. The analysis method should be reviewed to see if it would reveal other gases than those known to be in the sample.
- c. The spectrum furnished by EPA from the ROSE spectrophotometer clearly has noise in some regions of the spectrum. The section analyzed here appeared less noisy than any other by visual inspection, but if the noise were low frequency drifts, it would not be seen as obvious noise.
- d. An initial suspicion that water absorptions were responsible was discounted by the investigation described below.

In summary, the reason for the discrepancy between the value of 1.23×10^{19} molecules per cm² of CH₄ and the value of 8.22×10^{18} calculated by EPAGAS is not known at this time. Both instrumental uncertainties and possible library inadequacies must be investigated. It should be noted that even with these difficulties, the values differ by a factor of only 1.5.

Table 4.3 shows a summary of calculations done to determine the concentrations of $\mathrm{CH_4}$ and $\mathrm{H_20}$ in the spectrum. The water values are low (for water) and are scattered, when compared in different frequency regions, indicating that the spectra contains very little water absorption. The $\mathrm{CH_4}$ values are, on the other hand, quite consistent. Figure 4.16 shows the impossibility of fitting the given spectrum with a $\mathrm{CH_4}$ -only absorption spectrum. Structure in the given spectrum cannot be matched by any concentration of the $\mathrm{CH_4}$ spectrum.

Table 4.3

CALCULATION OF THE CONCENTRATION OF A SAMPLE OF CH₄ AND H₂O BY APPLICATION OF EPAGAS TO AN EXPERIMENTAL SPECTRUM

	Region	Frequency Correction	Trial Concentr (molecule	ations	Calcul Concentr		Final Transmission
Code	Spectral	Δν	H ₂ O	СН ц	H ₂ O	CH ₄	Error
1a	1250.0-1260.0	-0.6	$3.73x10^{22}$		$-2.17x10^{21}$	7.79x10 ¹⁸	6.07
b	1260.0-1270.0	-0.6	3.73×10^{22}	1.23x10 ¹⁹	6.82×10^{20}	8.46x10 ¹⁸	6.88
С	1270.0-1280.0	-0.6	$3.73x10^{22}$	1.23×10^{19}	$-1.44x10^{20}$	8.67×10^{18}	6.73
đ	1280.0-1290.0	-0.6	$3.73x10^{22}$	$1.23x10^{19}$	$5.04x10^{20}$	7.96x10 ¹⁸	7.14
2a	1250.0-1260.0	-0.9	3.73x10 ²²	1.23x10 ¹⁹			5.41
Ъ	1260.0-1270.0	-0.9	3.73×10^{22}	1.23×10^{19}	7.09×10^{20}	$8.17x10^{18}$	6.14
С	1270.0-1280.0	~0.9	$3.73x10^{22}$	$1.23x10^{19}$	4.42x10 ²⁰	6.85×10^{18}	8.68
d	1280.0-1290.0	-0.9	$3.73x10^{22}$	$1.23x10^{19}$	1.80×10^{20}	7.90x10 ¹⁸	10.20
		Δν		<u> </u>	H_2O -2.82x10 ²⁰	CH ₄ 8.22x10 ¹⁸	Error
Averag	ges for all region	ns -0.6					6.71
and a	given frequency	-0.9			1.04x10 ²⁰	$7.74x10^{18}$	7.61

Correction

5. CONCLUSIONS AND RECOMMENDATIONS

Many geophysical studies require the measurement of constituent concentrations in situations where the identifying signals from many components are mixed in the spectrometer. In remote measurement of aerosols by electromagnetic scatter, in measurement of ocean constituents, in the measurement of pollutants in the water and in the air, the same complex combination of identifying spectra occur. Means of separating and measuring a large number of components of a mixture usually require analysis by a large computer because the data set is quite large.

Not only must the field data be large enough to give sufficient statistically independent information, but the identifiers (spectra) require a large computer memory to avoid breaking the problem into too many pieces.

The calculational algorithms utilized in the program EPAGAS have been shown in this report to determine efficiently and accurately the concentrations of gas from a complex long-path infrared spectrum. Two important requirements are:

- The spectral library must be accurate in wavelength, pressure and temperature broadening parameters, and spectral line or band intensity.
- 2. The spectrophotometer used to obtain the field data must accurately measure the spectral intensity (at least relatively), and the wavelength and the spectral resolution must match the resolution of the spectra in the library.

The first requirement is met (with qualifications detailed in A2) by the spectral library assembled for this report (F1.1(a-c)), but adequate spectral data analysis of complex spectra is rare in the scientific literature, as discussed in A1. Other gases than those considered in this report are important for pollution monitoring. The technological changes in the United States resulting from the fuel shortage and the expected mineral shortage will require new industrial

developments such as oil shale processing. These can be expected to result in new effluents whose spectra will most likely not be well known. Indeed, only a small portion of atmospheric pollutants presently in industrial areas are well enough known spectroscopically.

Thus, it is emphatically recommended that continuing support be given to competent spectroscopists to obtain the basic spectroscopic data needed for air and water pollution. Absorption spectra, Raman spectra, and emission spectra are needed for the various techniques presently in development.

It is further recommended that vigorous efforts be made to obtain field data under controlled conditions to develop experience with the whole system. Initially, the experiments must include, along the path, point samplers to establish the levels of gaseous pollutants for comparison with the spectral analysis. Remote sensing systems must not be considered complete until proven in comparison with more conventional methods.

6. ACKNOWLEDGMENTS

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7. LIST OF SYMBOLS

I Radiation Intensity K' Absorption coefficient (cm ⁻¹) K Absorption coefficient		
K Absorption coefficient (cm² molecules⁻¹) W Area density (or concentration) (cm⁻²) P Density absorbing gas per unit area of radiation (cm⁻³) N Noise power V Spatial frequency (wave- number) (cm⁻¹) T Temperature (°K) P Pressure (dynes-cm² or atm) S(V) Spectrophotometer output E Error function Wg Given concentration (molecules cm⁻²) Wc Computed concentration (molecules cm⁻²) CF Confidence factor λ Wavelength (cm) C Speed of light in vacuum (cm sec⁻¹) f Frequency (sec⁻¹)	I	Radiation Intensity
(cm ² molecules ⁻¹) W Area density (or concentration) (cm ⁻²) P Density absorbing gas per unit area of radiation (cm ⁻³) N Noise power Spatial frequency (wave- number) (cm ⁻¹) T Temperature (°K) P Pressure (dynes-cm ⁻² or atm) S(v) Spectrophotometer output E Error function Wg Given concentration (molecules cm ⁻²) Wc Computed concentration (molecules cm ⁻²) CF Confidence factor A Wavelength (cm) C Speed of light in vacuum (cm sec ⁻¹) f Frequency (sec ⁻¹)	K'	Absorption coefficient (cm ⁻¹)
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E Error function Wg Given concentration	P	Pressure (dynes-cm ⁻² or atm)
Wg Given concentration (molecules cm ⁻²) Wc Computed concentration (molecules cm ⁻²) CF Confidence factor λ Wavelength (cm) C Speed of light in vacuum (cm sec ⁻¹) f Frequency (sec ⁻¹)	S(v)	Spectrophotometer output
(molecules cm ⁻²) Wc Computed concentration (molecules cm ⁻²) CF Confidence factor λ Wavelength (cm) C Speed of light in vacuum (cm sec ⁻¹) f Frequency (sec ⁻¹)	E	Error function
(molecules cm ⁻²) CF Confidence factor Wavelength (cm) Speed of light in vacuum (cm sec ⁻¹) f Frequency (sec ⁻¹)	Wg	. 3
<pre>Wavelength (cm) Speed of light in vacuum</pre>	Wc	
Speed of light in vacuum (cm sec ⁻¹) f Frequency (sec ⁻¹)	CF	
(cm sec ⁻¹) f Frequency (sec ⁻¹)	λ	Wavelength (cm)
	С	-
S/N Signal-to-noise ratio	f	Frequency (sec ⁻¹)
	s/n	Signal-to-noise ratio

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APPENDIX 1

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APPENDIX 1

A1. SOURCES AND CONTENTS OF THE LIBRARY OF SPECTRA

A1.1 MOTIVATION FOR THE LIBRARY

As outlined in the body of the report and explained in detail in A2, EPAGAS is a program which employs a least-squares fitting method to match an experimentally taken spectrum, say of a polluted atmosphere, with an internally generated reference spectrum. Through constructing reference spectra by adjusting the concentrations of gases in its library, EPAGAS fits the data spectrum to a reference spectrum and thus determines the concentration of the gases in the data spectrum.

The closer that the library comes to containing spectra of all the molecules found in the experimental spectra, the more accurate are the results of EPAGAS. (See 3.1.3.) This is the motivation for assembling a large and accurate library of spectra.

Physically, the library consists of a set of condensed computer cards which contain values of the absolute transmittance of each gas at each concentration at each frequency used by EPAGAS-a total of about 256,000 points.

A1.2 SOURCES AND CONTENT

Spectra for the 10 gases shown in F1.1 were assembed by this laboratory* over the past 10 years and are summarized in TA1.1. The spectra of the first seven gases listed in the table (set numbers 1A1 through 7B1) were calculated theoretically using line parameters and self-broadening constants. SO_2 , HNO_3 , and NH_3 spectra were taken experimentally. All assumed room temperature (296°K).

^{*}Wave Propagation Laboratory, ERL/NOAA. These data were assembled from theoretical computations and experimental data from many sources. See McClatchey et al., 1973.

Table Al.1

SET NUMBER	GAS	WAVENUMBER REGION	INCREMENT	OM THE WPL DATA	NUMBER OF POINTS PER CONCENTRATION	CONCENTRATIONS (molecules cm-2)
1 A1	H ₂ O-Water	740.0-1450.0	0.2	1.0	3551	7.46 $\times 10^{21}$, 2.24 $\times 10^{22}$, 3.73 $\times 10^{22}$, 5.22 $\times 10^{22}$, 6.77 $\times 10^{22}$
181	H ₂ O-Water	1800.0-3200.4	0.6	3.0	2335	7.46 $\times 10^{20}$, 2.24 $\times 10^{21}$, 3.73 $\times 10^{21}$, 5.22 $\times 10^{21}$, 6.77 $\times 10^{21}$
2A1	CO2-Carbon Dioxide	710.0-1100.0	0.2	1.0	1951	8.61×10^{20} , 1.08×10^{21} , 1.29×10^{21}
2B1	CO ₂ -Carbon Dioxide	1810.2-2499.0	0.6	3.0	1149	8.61×10^{20} , 1.08×10^{21} , 1.29×10^{21}
3A1	0 ₃ -0zone	970.0-1100.0	0.2	1.0	651	1.08 $\times 10^{17}$, 1.88 $\times 10^{17}$, 2.69 $\times 10^{17}$, 5.38 $\times 10^{17}$,
3B1	0 ₃ -Ozone	2040.0-2139.0	0.6	3.0	166	1.08 ×10 ¹⁷ 1.08 ×10 ¹⁷ , 1.88 ×10 ¹⁷ , 2.69 ×10 ¹⁷ , 1.08 ×10 ¹⁸ 5.38 ×10 ¹⁸ 5.38 ×10 ¹⁷
4A1	N ₂ 0-Nitrous Oxide	1120.0-1360.0	0.2	1.0	1201	5.38×10^{17} , 1.08×10^{18} , 1.88×10^{18}
4B1	N ₂ 0-Nitrous Oxide	2130.6-2599.8	0.6	3.0	783	5.38×10^{17} , 1.08×10^{18} , 1.88×10^{18}
5A1	CH ₄ -Methane	1180.0-1400.0	0.2	1.0	1101	2.69 $\times 10^{18}$, 5.38 $\times 10^{18}$, 1.08 $\times 10^{19}$, 1.88 $\times 10^{19}$, 2.69 $\times 10^{19}$
5B1	CH ₄ -Methane	2488.2-3199.8	0.6	3.0	1187	2.69 $\times 10^{18}$, 5.38 $\times 10^{18}$, 1.08 $\times 10^{19}$, 1.88 $\times 10^{19}$, 2.69 $\times 10^{19}$
6B1	CO-Carbon Monoxide	2000.4-2259.6	0.6	3.0	433	5.38 $\times 10^{18}$, 1.08 $\times 10^{19}$, 1.88 $\times 10^{19}$, 2.69 $\times 10^{19}$, 5.38 $\times 10^{19}$
7B1	HCL-Hydrochloric Acid	2400.6-3169.8	0.6	3.0	1283	2.69 $\times 10^{16}$, 1.08 $\times 10^{17}$, 2.69 $\times 10^{17}$, 1.08 $\times 10^{18}$, 2.69 $\times 10^{18}$
20.1	SO ₂ -Sulfur Dioxide	1074.8-1231.2	0.2	1.0	783	1.06×10^{18} , 3.54×10^{18} , 1.06×10^{19} , 3.54×10^{19}
20.2	SO ₂ -Sulfur Dioxide	1299.2-1409.2	0.2	1.0	551	3.54×10^{16} , 1.06×10^{17} , 3.54×10^{17} , 1.06×10^{18} , 3.54×10^{18}
30.1	HNO ₃ -Nitric Acid	824.8- 951.2	0.2	1.0	633	3.54×10^{17} , 1.06×10^{18} , 3.54×10^{18}
30.2	HNO ₃ -Nitric Acid	1234.8-1383.8	0.2	1.0	746	1.06×10^{17} , 3.54×10^{17} , 1.06×10^{18} , 3.54×10^{18}
40.1	NH ₃ -Ammonia	1112.0-1211.4	0.2	1.0	498	3.54×10^{17} , 1.06×10^{18} , 3.54×10^{18} , 1.06×10^{19}

The theoretical gas spectra were calculated using program DEGRADE (Deutschman and Calfee, 1967) for specific resolutions, temperatures, concentrations, and data increments. In other applications, it is possible to calculate new theoretical spectra for differing sets of these parameters. For the remaining experimental spectra in TA1.1 and all of those in TA1.2, except NO, it is only possible to generate lower resolution spectra of different concentrations using programs LOWRES and MORTRAN. See A1.4 for brief explanations and listings of these and other programs used in processing the library of spectra.

In TA1.1, "increment" refers to the spacing on the wavenumber axis between transmittance values. Wavenumber ν is related to frequency f and wavelength λ by $\nu = \frac{f}{c} = \frac{1}{\lambda}$, where c is the speed of light. Note that in the lower region, 700 to 1450 cm⁻¹, the increments are 0.2 cm⁻¹. In the upper region, 1800 to 3200 cm⁻¹, the increments are 0.6 cm⁻¹ and they are spaced evenly from 1800.0 cm⁻¹. This standardization of spacing is necessitated by the manner in which EPAGAS compares spectra. In the lower region, most spectra in the library were of 1.0 cm⁻¹ resolution, while in the upper region, they were typically of 3.0 cm⁻¹ resolution. Thus the 0.2 and 0.6 cm⁻¹ spacings give five data points per resolution element, providing essentially complete information about the spectra.*

^{*}Trying to compare higher or lower resolution experimental spectra with library spectra will introduce errors, but effects on EPAGAS outputs are presently unknown. It should be noted that most commercial spectrometers change resolution while making a large frequency scan. The adjustments to the program can be made quite simply by changing the input data to correspond to the library requirements.

To increase the library, the literature was searched for usable spectra of 23 molecules. These molecules have been found in polluted atmospheres, but no high resolution spectra had been collected for use in this application. Usable data for only 11 gases, those listed in TA1.2, could be found with the time and resources available. For the remaining 12 gases, either no spectra were available or published information was unusable because no absolute scale of transmittance or absorption was given or because the concentrations could neither be calculated nor estimated with available information. Except in a few special cases (SO₂), it was not possible to obtain spectra experimentally due to the expense and time involved with the procurement and handling of samples and the proper ratioing of data to obtain absolute values of transmittance.

A summary sheet for each of the 11 molecules for which useful information was found follows this discussion of general techniques (A1.3). Note that gas 4 is missing. This gas (SO₃) was deleted, after the coding system had been finalized, when it was determined that the spectra were in error. The important parameters required for the conversion of the published data to a form usable by program EPAGAS are listed in seven columns. First, a photocopy of the region of interest (column 1) was made from the literature and photo-enlarged to a size suitable for scaling on a digitizer. Absorption values for even increments of wavenumber (column 4) were punched onto cards, which in turn were used to produce one card for each frequency together with its corresponding value of transmittance, plus peripheral identifiers such as the resolution (column 2) and concentration (column 3) found in the literature.

It was necessary to produce these cards to manipulate the spectra with existing in-house programs. SPTR, a simple read-punch program, was used for this purpose. As needed, the spectra were then interpolated linearly to finer spacings (column 5) using program INTERP and/or degraded to a lower resolution (column 6) using program LOWRES. The latter program employs a sliding triangular slit function as a low pass filter to smooth high-resolution data into low-resolution data.

Table A1.2

SPECTRA PROCESSED FROM THE SCIENTIFIC LITERATURE

SET NUMBER	GAS	MAVENUMBUR REGION (cm ⁻¹)	INCREMENT (cm-1)	NUMBER OF POINTS PER CONCENTRATION	LOWEST CONCENTRATION (molecules cm-2)
					1.0
1	NO-Nitric Oxide	1805.4 - 1935.0	0.6	217	1.06×10^{18}
2	H ₂ S-Hydrogen Sulfide	2623.8 - 2755.2	0.6	220	1.06×10^{20}
3a	C ₂ H ₄ -Ethylene	800.0 - 1098.0	0.2	1491	1.06×10^{18}
3b	C ₂ H ₄ -Ethylene	1380.0 - 1450.0	0.2	351	3.54×10^{18}
3c	C ₂ H ₄ -Ethylene	1800.0 - 1955.4	0.6	260	3.54×10^{18}
3 d ′	C ₂ H ₄ -Ethylene	1970.4 - 2101.8	0.6	220	3.54×10^{19}
3e	C ₂ H ₄ -Ethylene	2936.4 - 3049.8	0.6	190	3.54×10^{18}
3f	C ₂ H _A -Ethylene	3050.4 - 3200.4	0.6	251	1.06×10^{18}
5	NO ₂ -Nitrogen Dioxide	2853.0 - 2967.0	0.6	1 91	3.54×10^{17}
6a	CH ₃ CH ₂ HCO-Acetone	700.0 - 1149.8	0.2	2250	1.06×10^{18}
6b	CH ₃ CH ₂ HCO-Acetone	1150.0 - 1315.8	0.2	830	3.54×10^{17}
6c	CH ₃ CH ₂ HCO-Acetone	1325.0 - 1450.0	0.2	626	3.54×10^{17}
6d	CH ₂ CH ₂ HCO-Acetone	2900.4 - 3099.0	0.6	332	1.06×10^{18}
7b	H ₂ CO-Formaldehyde	851.0 - 1071.6	0.2	1104	1.06 x 10 ¹⁹
7 c	H ₂ CO-Formaldehyde	1077.0 - 1237.0	0.2		3.54×10^{18}
7 e	H ₂ CO-Formaldehyde	1239.0 - 1439.0	0.2	801 1 001	3.54×10^{18}
7 е 8а	C ₂ H ₆ -Ethane	792.6 - 870.0	0.2		5.54 x 10
8b	C ₂ H ₆ -Ethane	1360.0 - 1450.0		388	3.54×10^{18}
			0.2	451	3.54×10^{18}
8c	C ₂ H ₆ -Ethane	2863.2 - 3050.4	0.6	313	3.54×10^{17}

Table A1.2 (cont.)

SET NUMBER	GAS	WAVEN REGI (cm		INCREMENT (cm ⁻¹)	NUMBER OF POINTS PER CONCENTRATION	LOWEST CONCENTRATION (molecules cm-2)
9a	n-C ₄ H ₁₀ -Normal Butane	700.0 -	1450.0	0.2	3751	3.54×10^{18}
9b	n -C ₄ H ₁₀ -Normal Butane	1800.0 -	3200.4	0.6	2335	3.54×10^{18}
10a	$i - C_A H_{10} - Iso-Butane$	700.0 -	1450.0	0.2	. 3751	1.06 x 10 ¹⁹
10b	i -C ₄ H ₁₀ -Iso-Butane	1800.0 -	3200.4	0.6	2 3 3 5	1.06×10^{19}
11a	C _z H _g -Propane	700.0 -	1450.0	0.2	3751	1.06 x 10 ¹⁹
11b	C _z H _g -Propane	1800.0 -	3200.4	0.6	2335	1.06×10^{19}
12a	C ₅ H ₁₂ -Pentane	700.0 -	1450.0	0.2	3751	3.54×10^{18}
12b	C _S H ₁₂ -Pentane	1800.0 -	1999.8	0.6	334	1.06×10^{19}
12c	C ₅ H ₁₂ -Pentane	2544.0 -	3200.4	0.6	1095	3.54×10^{18}

Note: Insufficient data was found for O_2 , C_6H_6 , SO_3 , HCN, H_2O_2 , C_3H_4O , C_2H_4O , C_4H_8O , C_3H_8 , C_7H_8 , and N_2-N_2 collisional spectra. In most cases, the spectrum was of too low resolution or unknown concentration.

Finally program MORTRAN, using Bouguer-Beer's law, calculated transmittance values for five concentrations from each data point for storage in the library. Any particular problems encountered or special techniques employed for a molecule are summarized in the final paragraph(s) on the individual summary sheets.

In several cases, molecular bands were found that could not be processed, either because absolute transmittance scales were not used by the author, resolution was extremely low, or no value of concentration could be deduced. Such bands were listed as missing if other bands of the same molecule could be processed. It is possible, though unlikely, that bands exist which the literature search did not find. Such bands were neither processed nor listed as missing.

The accuracy of the spectra in the library was controlled primarily by the accuracy of the blowup process. Typically, a 2 x 6 in. journal spectrum was enlarged to 18 x 54 in. and data points were taken every 0.04 in. on the frequency axis of the enlargement. The uncertainty in reproducing the position of a spectrum on the frequency axis was about 0.12 in., or $\frac{1}{2}$ of the data-taking intervals (column 4). The uncertainty in controlling the length of the transmittance axis is $\frac{1}{2}$ 5 percent of full scale. Both of these uncertainties assume the reported spectra were ratioed and calibrated.

The spectra for several gases at concentrations expected in the atmosphere were nearly straight lines. That is, more than 90 percent of the band had transmittance values above 0.97. Such spectra are nearly useless for use in EPAGAS. Thus, the first concentration which caused at least 10 percent of the band to fall under 97 percent transmission, and the next four higher concentrations in the standard sequence were calculated and used in the library. The standard sequence is . . . , 1.06 x 10^n , 3.54 x 10^n , 1.06 x 10^{n+1} , 3.54 x 10^{n+1} , 1.06 x 10^{n+2} , . . . (molecules cm⁻²), where n is an integer. The "lowest concentration" in the last column on table A1.2 refers to the first number selected from this sequence and implies the next four which were used in the library.

Concentrations on TA1.1 and TA1.2 and on the summary sheets are given in molecules per square centimeter for a 1-km path. 1.06×10^{18} molecules cm⁻² corresponds to 3 torr partial pressure at 296° K. For conversion to other units, consult A4 (Calfee, 1971) which lists many of the standard units used in studying gaseous absorption processes and summarizes their interrelationships. See the reference for detailed explanations.

Figures 1.1b and 1.1c display selected spectra for each of the gases processed. The overlays to F1.1a and F1.1b depict band designations. Where disagreement occurs between literature sources, the band was not identified, but the disagreement was noted on the summary sheet. Due to the complicated vibration and rotation modes of the larger molecules in F1.1c, individual bands are not easily identifiable and therefore no overlay is provided.

Figure A1.1 summarizes the state of knowledge in the infrared of absorbing gases found in the atmosphere.

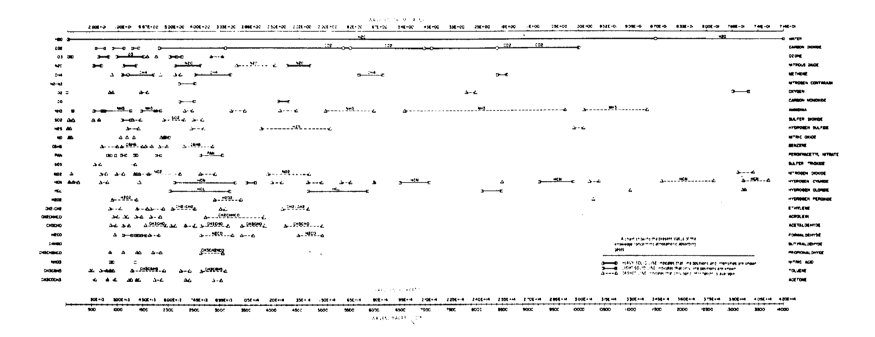


Figure A1.1
INFRARED ABSORBING REGIONS OF ATMOSPHERIC GASES

GAS 1: NO-Nitric Oxide

l Frequency Range (cm ⁻¹)	Original Resolution (cm ⁻¹)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm ⁻¹)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	7 Interpolated Concentrations (mol. cm ⁻²)	Source
1805.4-1935.0	-	-	-	0.6	3.0	1.06x10 ¹⁸ -1.06x10 ²⁰	1,2

The NO spectra were calculated from theoretical line positions (Shaw, 1956), theoretical line strengths (Abels & Shaw, 1966), and experimental half-widths (Abels & Shaw, 1966) using program DEGRADE. Where not reported, positions were computed using formulas presented in Shaw, 1956, and half widths were estimated from adjacent lines. Line parameter cards containing this unreported information are identifiable by a missing last significant figure in the field describing the position and/or half-width.

^{1.} J. Chem. Phys., Vol. 24, #2, p. 399, 1956, Shaw.

^{2.} J. Molec. Spec., Vol. 20, p. 11, 1966, Abels & Shaw.

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GAS 2: H₂S-Hydrogen Sulfide

1 Frequency Range (cm ⁻¹)	Original Resolution (cm-1)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm-1)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	7 Interpolated Concentrations (mol. cm ⁻²)	Source
2623. 8-2755.2	1.2	2.7×10^{20}	0.6	-	-	1.06x10 ²⁰ -1.06x10 ²²	1

1. Phys. Review, Vol. 37, p. 728, 1931, Nielsen & Barker

A strong band, 1100-1400cm⁻¹, is missing. The resolution claimed by Nielsen and Barker seems high. The apparent resolution of their spectrum is close to 3.0cm⁻¹, the resolution required for this region. Therefore, LOWRES was not used in processing this spectrum for the library.

GAS 5: Call,-Ethylene

Prequency Range (cm-1)	Original Resolution (cm ⁻¹)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm-1)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	Interpolated Concentrations (mol. cm ⁻²)	Source
800.0-1098.0	0.8	1.1 x 10 ¹⁹	0.4	0.2	-	1.06x10 ¹⁸ -1.06x10 ²⁰	1
1380.0-1450.0	1.5	3.0×10^{19}	0.2	-	-	$3.54 \times 10^{18} - 3.54 \times 10^{20}$	2
1800.0-1955.4	1.6	1.0×10^{20}	0.6	-	-	$3.54 \times 10^{18} - 3.54 \times 10^{20}$	2
1970.4-2101.8	1.5	2.5×10^{20}	0.6	-	-	$3.54 \times 10^{19} - 5.54 \times 10^{21}$	2
2936.4-3049.8	0.6	3.4×10^{19}	0.6	-	5.0	$3.54 \times 10^{18} - 3.54 \times 10^{20}$	3
3050.4-3200.4	1.5	2.4×10^{19}	0.6	-	-	$1.06 \times 10^{18} - 1.06 \times 10^{20}$	2

No accurate concentrations were reported by Levin and Meyer. The original concentrations given in column 3 were first estimated from Sadler low resolution band strengths and then refined by comparison with neighboring spectra where possible.

For instance, the original spectra in the $2956-3049~\rm cm^{-1}$ and $3050-3200\rm cm^{-1}$ regions were assumed to have been taken at concentrations of $2.7~\rm x~10^{19}~mol-cm^2$ and $3.0~\rm x~10^{19}~mol-cm^2$, respectively. When converted to a concentration of $5.54~\rm x~10^{18}~mol-cm^{-2}$, a noticeable mismatch occurred between the two spectra at $5050\rm cm^{-1}$. To cause the two spectra to coincide at $3050\rm cm^{-1}$, the assumed concentration for the first spectrum had to be increased to $3.4~\rm x~10^{19}~mol-cm^{-2}$ (25.3% increase) and the second lowered to $2.4~\rm x~10^{19}~mol-cm^{-2}$ (19.1% decrease). These refined figures were the ones reported on the summary sheet, and they reflect a 50% uncertainty in the reported concentration.

Once again, the reported resolution seems too high in all the Levin and Meyer spectra. Apparent resolution is about 5.0cm⁻¹ Note that the apparent resolution is satisfactory for the 1800-3200cm⁻¹ region but too low for the 700-1500cm⁻¹ region.

^{1.} J. Chem. Phys., Vol. 40, #8, p. 2096, 1964 Smith & Mills.

^{2.} J. Opt. Soc. Amer., Vol. 16, #3, p. 155, 1928, Levin & Meyer.

^{3.} J. Chem. Phys., Vol. 8, p. 799, 1940, Smith.

GAS 5: NO₂-Nitrogen Dioxide

1 Frequency Range (cm-1)	Original Resolution (cm ⁻¹)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm-1)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	7 Interpolated Concentrations (mol. cm ⁻²)	Source
2853.0-2967.0	0.6	1.1 x 10 ¹⁹	0.6	-	1.0	3.54x10 ¹⁷ -3.54x10 ¹⁹	1

Only about one half of the band is available from the band center to one wing. Another band, centered at $1617 \, \mathrm{cm}^{-1}$ is available, but was omitted from the library due to strong $\mathrm{H}_2\mathrm{O}$ absorption in this region.

^{1.} Private communication from Goldman, University of Denver, 1969.

GAS 6: CH₃CH₂HCO-Acetone

	l Frequency Range (cm ⁻¹)	2 Original Resolution (cm ⁻¹)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm-1)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	7 Interpolated Concentrations (mol. cm ⁻²)	Source
a	700.0-1149.8	(IR^{-7})	8.2 x 10 ¹⁹	1.0	0.2	-	1.06x10 ¹⁸ -1.06x10 ²⁰	1
b	1150.0-1315.8	(IR^{-7})	3.2×10^{18}	1.0	0.2	-	$3.54 \times 10^{17} - 3.54 \times 10^{19}$	1
С	1325.0-1450.0	(IR^{-7})	9.9×10^{18}	1.0	0.2	-	$3.54 \times 10^{17} - 3.54 \times 10^{19}$	1
d	2900.4-3099.0	(IR^{-7})	9.9×10^{18}	1.0	0.6	-	$1.06 \times 10^{18} - 1.06 \times 10^{20}$	1

A band exists from $1680-1825 {\rm cm}^{-1}$ as reported by Cassee and Schachtschneider, but it was omitted due to strong ${\rm H}_2{\rm O}$ absorption in this region.

The resolution is lower than desired (approximately 7cm⁻¹) due to the low resolution of the Beckman IR-7 spectrometer. Since a range of resolutions may be preset on this instrument, one needs to consult the workers to determine the exact resolution of the various runs.

^{1.} J. Chem. Phys., Vol. 44, #1, p. 97, 1966, Cossee & Schachtschneider.

GAS 7: H₂CO-Formaldehyde

	Frequency Range (cm-1)	Original Resolution (cm ⁻¹)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm-1)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	7 Interpolated Concentrations (mol. cm ⁻²)	Source
	851.0- 997.2	0.3	7.1 x 10 ¹⁹	0.2	-	1.0	1.06x10 ¹⁹ -1.06x10 ²¹	1
	9 9 7.4-1071.6	0.3	3.5×10^{19}	0.2	-	1.0	$1.06 \times 10^{19} - 1.06 \times 10^{21}$	1
;	1077.0-1237.0	0.3	4.7×10^{19}	0.2	_	1.0	$3.54 \times 10^{18} - 3.54 \times 10^{20}$	1
	1239.0-1325.0	0.3	2.4×10^{19}	0.2	-	1.0	$3.54 \times 10^{18} - 3.54 \times 10^{20}$	1
	1321.0-1439.0	0.3	5.3×10^{19}	0.2	-	1.0	$3.54 \times 10^{18} - 3.54 \times 10^{20}$	1

Temperatures of the original spectra range from 150°C to 25°C , accounting for small discrepancies which may occur between regions. See Nakagawa & Morino for details.

Note an error on page 90, Figure 1e. The reported cell length is 100cm but should read 10cm.

Nakagawa and Morino report the band head at 1167cm^{-1} to be v_4 , while Herzberg claims it is v_6 .

^{1.} J. Molec. Spec., Vol. 38, p. 90, 1971, Nakagawa & Morino.

GAS 8: C₂H₆-Ethane

	1 Frequency Range (cm ⁻¹)	2 Original Resolution (cm ⁻¹)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm ⁻¹)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	7 Interpolated Concentrations (mol. cm ⁻²)	Source
а	792.6- 870.0	0.6	6.0×10^{19}	0.2	-	-	$3.54 \times 10^{18} - 3.54 \times 10^{20}$	1
b	1360.0-1450.0	1.3	4.0×10^{19}	0.2	-	-	$3.54 \times 10^{18} - 3.54 \times 10^{20}$	
С	2863.2-3050.4	1.4	6.0×10^{19}	0.6	-	-	$3.54 \times 10^{17} - 3.54 \times 10^{19}$	1

No accurate concentrations were reported. The original concentrations given in column 2 were estimated from Sadler low resolution band strengths.

The reported resolution seems too high. The apparent resolution is about $1.0\,\mathrm{cm}^{-1}$ in the $793-870\,\mathrm{cm}^{-1}$ region and about $3.0\,\mathrm{cm}^{-1}$ in the remaining two regions.

^{1.} J. Opt. Soc. Amer., Vol. 16, #3, p. 155, 1928, Levin & Meyer.

GAS 9: C_4H_{10} -N-Butane

-					4 10			
	1	2	3	4	5	6	7	
	Frequency	Original	Original	Increment	Converted	Converted	Interpolated	
	Range	Resolution	Concentration	Data Taken	Increment	Resolution	Concentrations	
	(cm ⁻¹)	(cm ⁻¹)	$(mo1. cm^{-2})$	(cm-1)	(cm ⁻¹)	(cm ⁻¹)	(mol. cm ⁻²)	Source
		_	20	-			18 3	20
a	700.0 -1450.0	-	2.5×10^{20}	5.0	0.2	-	$3.54 \times 10^{18} - 3.54 \times 10^{2}$	1
b	1800.0-3200.4	-	2.5×10^{20}	5.0, 10.0	0.6	-	$3.54 \times 10^{18} - 3.54 \times 10^{2}$	20

The resolution for all regions is unknown, but is approximately $10-15\,\mathrm{cm}^{-1}$. The instrument used was a Perkin-Elmer Model 21.

^{1.} Anal. Chem., Vol. 28, p. 1218, 1956, Pierson, Fletcher, Grantz.

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				GAS 10:	C ₄ H ₁₀ -I-Butane		·	
4	1 Frequency Range (cm-1)	2 Original Resolution (cm ⁻¹)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm-1)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	7 Interpolated Concentrations (mol. cm ⁻²)	Source
а	700.0-1450.0	-	2.5×10^{20}	5.0	0.2	-	1.06x10 ¹⁹ -1.06x10 ²¹	l 1
Ъ	1800.0-3200.4	· -	2.5×10^{20}	5.0, 10.0	0.6	-	$1.06 \times 10^{19} - 1.06 \times 10^{21}$	1 1

1. Anal. Chem., Vol. 28, p. 1218, 1956, Pierson, Fletcher, Grantz.

The resolution for all regions is unknown, but is approximately $10-15\,\mathrm{cm}^{-1}$. The instrument used was a Perkin-Elmer Model 21.

GAS 11: C₃H₈-Propane

	1 Frequency Range (cm- ¹)	2 Original Resolution (cm ⁻¹)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm-1)	5 Converted Increment (cm ⁻¹)	6 Converted Resolution (cm ⁻¹)	7 Interpolated Concentrations (mol. cm ⁻²)	Source
а	700.0-1450.0	-	2.5×10^{20}	5.0	0.2	•	1.06x10 ¹⁹ -1.06x10 ²	1 1
b	1800.0-3200.4	-	2.5×10^{20}	5.0, 10.0	0.6	-	$1.06 \times 10^{19} - 1.06 \times 10^{2}$	1 1

The resolution for all regions is unknown, but is approximately $10-15\,\mathrm{cm}^{-1}$. The instrument used was a Perkin-Elmer Model 21.

^{1.} Anal. Chem., Vol. 28, p. 1218, 1956, Pierson, Fletcher, Grantz.

GAS 12: C₅H₁₂-Pentane

					3 12			
	1 Frequency Range (cm-1)	Original Resolution (cm-1)	3 Original Concentration (mol. cm ⁻²)	4 Increment Data Taken (cm-1)	5 Converted Increment (cm ⁻¹)	Converted Resolution (cm ⁻¹)	Interpolated Concentrations (mol. cm ⁻²)	Source
a	700.0-1450.0) -	9.1 x 10 ¹⁹	5.0	0.2		$3.54 \times 10^{18} - 3.54 \times 10^{2}$	-
b	1800.0-1999.8	3 -	9.1×10^{19}	5.0	0.2	•	$1.06 \times 10^{19} - 1.06 \times 10^{2}$	1
С	2544.0-3200.4	4 -	9.1×10^{19}	10.0	0.6		$3.54 \times 10^{18} - 3.54 \times 10^{2}$	0 1

1. U.S. Naval Research Lab., American Petroleum Institute Research Project 44.

The original spectrum for this molecule was for a <u>liquid</u> sample 0.174mm thick, and therefore it may differ significantly from the gaseous spectrum. Equivalent gaseous concentrations were figured from the density of liquid pentane and the thickness of the sample.

The resolution for all regions is unknown, but is approximately $10-15\,\mathrm{cm}^{-1}$.

A1.3 PROGRAMS USED IN PROCESSING SPECTRA

FORTRAN PROGRAM

CDC 3800

SPTR

M. J. Post

Aug. 17, 1973

Given absorption values A expressed as an integer number between 999 and 0, corresponding to 99.9 and 0.0 percent absorption, SPTR punches one card for each absorption value. On this card are punched wavenumber and the value of transmittance.

Input

 $\underline{\text{Card}} \ A_i - A_n$

Column	<u>Field</u>	Description
1-80	2014	Absorption values A(I)

Output

Card A,

1-10	F10.2	Wavenumber V _i
11-20	F10.3	Transmittance T _i
54-58	F5.2	Resolution RES
59-70	E12.3	Concentration CONC
73-80	A8	Molecule name MOL

Note: 1. An EOF card is required at the end of the data cards.

- 2. RES, CONC, and MOL are entered on source cards, not data cards.
- 3. V_{i} are generated internally by changing source cards.

PROGRAM SPTR

THIS PROGRAM TAKES COMPACTED SPECTRAL ABSORPTION VALUES (FORMAT 2014) AND OUTPUTS CARDS. ONE FREQUENCY AND ONE TRANSMISSION VALUE PER CARD, PLUS PERIPHERAL IDENTIFIERS.

IT IS USEFUL IN CONVERTING DIGITIZED ANALOG SPECTRAL INFORMATION (FROM STRIP CHARTS, FOR EXAMPLE) TO CARDS IN A FORMAT USED IN ASSOCIATED PROGRAMS.

```
PROGRAM SPTR
   DIMENSION A(20)
 1 FORMAT (20F4.4)
2 FORMAT (F10.2.F10.3.33X.F5.2.E12.3.2X.AR)
   V=1149.0
   DV=1.0
   MOL=8H ACETONE
   RFS=0.0
   CONC=3.2E+18
 5 READ 1, (A(I), I=1,20)
   IF (EOF+60)20+10
10 Do 15. K=1.20
   V=V+DV
   T=1.-A(K)
   PRINT 2+V+T+RES+CONC+MOL
15 PUNCH 2.V.T.RES.CONC.MOL
   GO TO 5
20 STOP
   END
```

FORTRAN PROGRAM
CDC 3800

LOWRES

M. J. Post

Aug. 17, 1973

Given a transmission spectrum T(N) of high resolution, LOWRES degrades it to a spectrum TF of lower resolution A.

Used in the computations are the given increment DV and the output increment DELV. A triangular slit function of half-width A is used to compute each new TF.

Input

Card A

9	Column	<u>Field</u>	Description
	1-10	F10.3	Beginning wavenumber
			of given spectrum, V1
	11-20	F10.3	End wavenumber of
			given spectrum, V2
	21-25	F5.2	Increment given
			spectrum, DV
	26-30	F5.2	Lower resolution
			desired, A
	31-35	F5.2	Desired output
			increment, DELV
Card B			
	11-20	F10.3	Given spectrum
			T(N)

Output

Card A

· · · · · · · · · · · · · · · · · · ·		
Column	<u>Field</u>	Description
1–10	F10.3	Wavenumber of output
		transmittance, VI
11-20	F10.5	Transmittance TF_{i} of
		output spectrum
54-5 8	F5.2	Resolution of output
		spectrum RES (=A)
50-70	E12.3	Concentration of output
		spectrum CONC
73-80	A8	Molecular name MOL

NOTE: An EOF card is required at the end of the data cards. DELV must be an exact multiple of DV and A must be an odd multiple of DV.

CONC and MOL are entered on source cards, not data cards.

PROGRAM LOWRES

THIS PROGRAM TAKES A SPECTRUM OF HIGHER RESOLUTION GIVEN IN INCREMENTS OF DV AND DEGRADES IT TO A SPECTRUM OF RESOLUTION A, GIVEN IN INCREMENTS OF DELV, POSITIONED AT THE ORIGINAL FREQUENCIES.

DELV MUST BE AN EXACT MULTIPLE OF DV.

A MUST BE AN ODD MULTIPLE OF DV.

```
DIMENSION T(2000)
  1 FORMAT (2F10.3,3F5.0)
  2 FORMAT (10X, F10.3)
  3 FORMAT (F10.3,F10.5,33X,F5.2,E12.3,2X,A8)
    RES=3.0
    CONC = 1.05E + 19
    MOL=8H
               NO2
    READ 1, V1, V2, DV, A, DELV
    MULT=DELV/DV + .00001
    SLIT=A*2.
    NMX=SLIT/DV-.99999
    SLTFTR=DV/A**2
    VEND=V2-A+DV
    N=1
 20 READ 2,T(N)
    IF (EOF, 60) 30, 25
 25 N=N+1
    GO TO 20
 30 NJ=1
    VI=V1+A-DV
    V=VI+DV
 50 SUMT=0.
    DO 150 NN=NJ,NMX
    TT=(A-ABS(V-VI))*T(NN)
    SUMT=SUMT+TT
150 V=V+DV
    TF=SUMT*SLTFTR
    PRINT 3, VI, TF, RES, CONC, MOL
    PUNCH 3, VI, TF, RES, CONC, MOL
    NJ=NJ+MULT
    NMX=NMX+MULT
    VI=VI+DV*MULT
    IF (VI.GT. VEND)
    V=VI-A+DV
    GO TO 50
    END
```

INTERP

M. J. Post

Aug. 17, 1973

Given a spectrum spaced in increments of DELV, INTERP linearly interpolates it to finer increments DIV and outputs it on cards.

Input

Card A

	Column	<u>Field</u>	Description
	1-5	F5.2	Input spectral
			increment DELV
	6-10	F5.2	Output spectral
			increment DIV
Card B			
	1-10	F10.3	Beginning wavenumber
			of spectrum V1
	11-20	F10.5	First transmittance
			value of spectrum T1
	54-58	F5.3	Resolution of spectrum
			RES
	59-70	E12.3	Concentration of
			spectrum CONC
	73-80	A8	Molecular name MOL
<u>Card</u> C			
	11-20	F10.5	Transmittance values
			T2 _i

Output

$\underline{\mathtt{Card}}\ \mathtt{A_i}$

Column	Field	Description
1-10	F10.3	Wavenumber of
	•	transmittance value,
		vi,
11-20	F10.5	Transmittance value T
54-58	F5.3	Resolution of spectrum
		RES
59-70	E12.3	Concentration of
		spectrum CONC
73-80	A8	Molecular name MOL

```
PROGRAM INTERP
C
      THIS PROGRAM TAKES A SPECTRUM SPACED IN INCREMENTS OF DELV AND LINEARLY
C
C
       INTERPOLATES IT TO FINER INCREMENTS DIV.
С
С
      DFLV MUST BE AN INTEGRAL MULTIPLE OF DIV.
C
    1 FORMAT (2F5.2)
    2 FORMAT (10x+F10+5)
3 FORMAT (F10+3+F10+5+33x+F5+3+F12+3+2X+AB)
       READ 1.DFLV.DIV
       READ 3.V1.T1.RES.CONC.MOL
       N=DELV/DIV + .001
       V[=V1-D]V
   10 READ 2.T2
       IF (FOF+60) 20+12
   12 PART = (T2-T1)/V
      DO 15 K=1.4
      T=T1+PART*(K-1)
      VI=VI+DIV
   PHNCH 3.VI.T.RES.CONC.MOL
15 PPINT 3.VI.T.PES.CONC.MOL
      T1=T2
      30 TO 10
   SO END
```

For a detailed discussion of program DEGRADE, see Deutschman and Calfee, 1967.

```
PROGRAM DEGRADE
      DIMENSION W(5)+F(5)+AB(5)+TT(5)+TF(5)+SUMT(5)+TRAPSUM(5)+FMT1(9)+
     1 FMT2(9) .FMT3(9) .FMT4(9)
      COMMON/1/GNU(3000)+54(3000)+A4 (3000)+CAY(1)/2/T(3000+5)+I5
    I FORMAT (9AB)
    2 FORMAT(F9.2.F12.6.10F10.5)
    3 FORMAT(/)1HOPPESSURE =F9.5.3XAHSLIT A =F9.5.3X13HTFMPERATURE =F9.2
     1//12H ARSURPTANCE + 9×13HCONCENTRATION/)
    4 FORMAT(24H KICKED OUT FROM ST. NO.+214+F10.2)
    5 FOPMAT(* WAVENUMBER WAVELENGTH TRANS 1 ABSORB 1
                                                             TRANS 2 ABSOR
     18 2 TRANS 3 ABSORB 3 TRANS 4 ABSORB 4 TRANS 5 ABSORB 54/1
    6 FORMAT (F8.2.5F10.5)
    7 FORMAT(11H1PRESSURE =F9.5.3X13HTEMPERATURE =F9.2.3X8HSLIT A =F9.5.
     13x4HDV =F7.4+3X6HDELV =F7.4+3X7HBOUND *F7.2/)
    8 FORMAT(F12.4.10X.E11.4)
    9 FORMAT(///33HODATA FOR PLOTTING ARE ON TAPE C-. 14)
   10 FORMAT(I4)
   11 FORMAT(60H)TOO MANY LINE CARDS. EXCEEDS DIMENSION. LAST CARD READ W
     145 =F9.2)
   12 FORMAT(19H117 TOO LARGE. I7 =15.5X9HGNJ(17) =F9.2.5X12HV1-A+BOUND
     1 = F9.2
   13 READ 1. (FMT1(I).I=1.9)
   14 READ 1. (FMT2(1).I=1.9)
   15 RFAD 1. (FMT3(I).I=1.9)
   16 \text{ RFAD } 1 \cdot (\text{FMT4}(I) \cdot I = 1 \cdot 9)
      PRINT1,FMT1,FMT2,FMT3,FMT4
   17 RFAD FMT3+K1+I7+DV+DELV+V1+V2+A+P+NP+ROUND
   18 READ EMT4.TEMP.TEMPO.BX.CX
 1800 READ FMT2 + (W(I) + I=1 + K1)
C
      SFT UP TEMPERATURE CORRECTION CONSTANTS.
C
 1810 \text{ CS1} = (\text{TEMPO-TEMP})/(\text{TEMPO+TEMP+.6950})
 1820 CS2=(TEMPO/TEMP)##BX
 1830 CA=((TEMPO/TEMP)**CX)*P
      CALL Q9 EXUN
      READ IN LINE DATA AND COUNT CARDS
                               VHI=V2+A+BOUND
      VL0=V1-A-BOUND-1.$
   20 I=0
   21 I=I+1
   25 READ
             FHT1 +GN +S+ALF+EPP+MOL
      IF (GN .LT.VLO) GO TO 25
      IF (GN .GT.VHI) GD TO 27
      IF(FOF,60)27,272
  272 IF (MOL.F3.1) GO TO 274
  273 ALF=.1
  274 AA(I)=(ALF*CA)**2
      GVU(I)=GN
 2601 SA(I)=S#ALF#CA#CS2#EXP(-EPP#CS1)
                                                   $ 60 TO 21
   27 11=1-1
   28 FORMAT (2F10.3.15)
      PPINT 28+ GNU(1)+GNU(11)+11
```

```
C CHECK THAT GNU(17) IS NOT GREATER THAN V1-A+BOUND.
   32 XROUND = V1 - A + BOUND
   33 IF (GNU(17) .GT. XBOUND) 34+36
   34 PRINT 12.17.GNU(17).XBNUND
   35 Go TO 1001
   36 PPINT 7.P. TEMP. A.DV. DELV. ROUND
   37 POINT 38. (W(K),K=1.K1)
   38 FORMAT(26X++W1 =+E10.3+5X++W2 =+E10.3+5X+W3 =+E10.3+5X +W4 =+F10.3
     1.5X#W5 =*E10.3/)
   39 PRINT 5
C
       SFT UP CALCULATION COUNTERS.
   40 IS=1
   41 SLIT=2.#A
   46 Iniv=DELV/DV+.01
   47 NODDI=IDIV+1
   48 NODD2 = (SLIT/DV) + 1.00001
   49 \text{ NODD3} = \text{NODD2} - \text{IDIV}
   52 \text{ NJ} = 1
   55 SLTFTR = DV/(A**2)
   98 \text{ AI} = \text{AI}
   69 V = V1 - A
C
       INITIALIZE SUMMATIONS
C
   70 DO 71 K=1+K1
   71 SUMT(K) = TRAPSUM(K) = 0.0
C
      ARSORPTION COEFFICIENT CALCULATION BY SUBROUTINE ARSCOEF
C
C
   79 DO 150 NN = NJ.NODD2
   BO CALL ABSCOEF (V.BOUND.11.2)
C
      TPANSMISSION SUMMATION AND APPLICATION OF VARIABLE PORTION OF SLIT
C
      FACTOR INTEGRAL
  130 Do 138 K=1.Kl
  134 \text{ T(NN+K)} = \text{EXPF(-w(K)*CAY)}
  136 TT(K)=(A-ABSF(V-VI))*T(NN.K)
  138 SUMT (K) = SUMT (K) +TT (K)
  150 V=V+DV
  154 V=VI
C
      TRANSMISSION AND ARSORPTION CALCULATION WITH FINAL SLIT FACTOR
C
С
      APPLICATION
  160 Do 175 K=1.Kl
  161 TF(K)=SUMT(K)#SLTFTR
  162 E(K) = 1.-TF(K)
  163 IF (V-V1) 509+164+166
  164 TRAPSUM(K) = TRAPSUM(K) +0.5*E(K)
  165 Gn TO 174
```

```
166 \text{ TPAPSUM(K)} = \text{TRAPSUM(K)} + \text{E(K)}
C
       OUTPUT STATEMENTS AND FREQUENCY INCREMENTING
  174 ALAM = 1.E+4/V
  175 CONTINUE
  176 PRINTS. V.ALAM. (TF(K).F(K).K=1.K1)
       PUNCH 6.V. (TF(K).E(K).K=1.K1)
  177 GO TO(180+181) - NP
  180 WPITE(2. 6 )V.(TF(K).K=1.K1)
  181 V=V+DELV
  190 Do 191 K=1.Kl
  191 SUMT(K)=0.0
  200 IF(V-V2)201,201,235
C
       SAVING OF ALMEADY CALCULATED TRANSMISSION VALUES FOR NEXT STEP
С
       IN THE CALCULATION
  201 00 205 K=1.K1
  202 DO 204 NN = NODD1+NODD2
  503 KK=NN-IDIA
  204 T(KK+K)=T(NN+K)
  205 CONTINUE
  210 V;=V
  211 V=V-A
  215 DO 219 NN = 1.40DD3
  216 DO 218 K=1.K1
  217 TT(K)=(A-ABSF(V-VI))*T(NN.K)
  218 SUMT(K) = SUMT(K) + TT(K)
  219 V=V+DV
  225 \text{ NJ} = \text{NODD3} + 1
  229 GO TO 79
С
C
       TPAPAZOIDAL RULE CALCULATION OF TOTAL ABSORPTANCE
C
  235 PRINT 3.P.A.TEMP
  236 Do 238 K=1.K1
  237 AB(K) = DELV*(TRAPSUM(K)-0.5*E(K))
  238 PRINT R+AB(K)+W(K)
  239 READ 240.V1.V2
  240 FORMAT (2F10.2)
  241 IF(EDF.60) 250.68
  250 GO TO(251+1001) . NP
  251 END FILE 2
  252 RF*IND 2
  255 Gn TO 1901
C
C
      KICKOUT STATEMENTS
  500 \text{ Nx} = 31
  501 GO TO 1000
  509 \text{ Ny} = 163
 1000 PRINT 4+NX+I1+SNU(II)
 1001 CALL EXIT
      END
```

```
SUBROUTINE ABSCOEF (V-ROUND-11-P)
      COMMON/1/GNU(3000).SA(3000).AA (3000).CAY(1)/2/T(3000.5).15
C
C
      DFTERMINATION OF INDEXING VALUES FOR ABSORPTION COEFFICIENT CALC.
   10 Do 14 I = 15.I1
   11 IF (V-BOUND-GNU(I))12+12+14
   12 15 = 1
   13 Go TO 15
   14 CONTINUE
   15 Do 19 K = 15 \cdot 11
   16 IF (V+BOUND-GNU(K))17.17.19
   17 I6 = K
   18 GO TO 25
   19 CONTINUE
С
      CALCULATION OF ABSORPTION COEFFICIENT WITH LORENTZ LINE SHAPE
C
   25 \text{ CAY1} = \text{CAY2} = 0.0
   30 Dn 46 I = 15 \cdot 16
   32 Y = ABSF(V-GNU(I))
   34 IF (Y-2.) 36.36.42
   36 SUM1=SA(I)/(Y++2+AA(I))
   38 CAY1 = CAY1 + SUM1
   40 GO TO 46
   42 SUM2=SA(1)/Y##2
   44 CAY2 = CAY2 + SU42
   46 CONTINUE
   50 \text{ CAY} = 0.3183^{\circ}(\text{CAY1} \cdot \text{CAY2})
   52 RETURN
      END
          SCOPE
CACLI
[RJN. 2.10000
 (F10.3,E10.3.F5.3.F10.3,43X.12)
 (5E10.3)
 (12.14.2F6.3.2F8.2.F5.2.E10.3.12.F7.2)
```

```
SUBROUTINE ABSCOEF(V, BOUND, 11, P)
       COMMON/1/GNU(3000), SA(3000), AA(3000), CAY(1)/2/T(3000,5), 15
С
       DETERMINATION OF INDEXING VALUES FOR ABSORPTION COEFFICIENT CALC.
C
   10 DO 14 I =15.I1
   11 IF (V-BOUND-GNU(I))12,12,14
   12 	 15 = I
   13 GO TO 15
   14 CONTINUE
   15 DO 19 K = 15,11
   16 IF(V+BOUND-GNU(K))17,17,19
   17 \quad I6 = K
   18 GO TO 25
   19 CONTINUE
С
       CALCULATION OF ABSORPTION COEFFICIENT WITH LORENTZ LINE SHAPE
C
   25 \quad CAY1 = CAY2 = 0.0
   30 DO 46 I = 15, 16
   32 \quad Y = ABSF(V-GNU(I))
   34 	 IF(Y-2.)36,36,42
   36 SUM1=SA(I)/(Y**2+AA(I))
   38 \quad CAY1 = CAY1 + SUM1
   40 GO TO 46
   42 SUM2=SA(I)/Y**2
   44 \quad CAY2 = CAY2 + SUM2
   46 CONTINUE
   50 CAY = 0.3183*(CAY1+CAY2)
   52 RETURN
       END
```

SINGLE-BANK COMPILATION.

FORTRAN PROGRAM CDC 3800

MORTRAN

Dana Gregory
June 29, 1973

Given the original gas concentration, W_o , the corresponding theoretical transmission, $T(W_o, v_j)$, as a function of the wavenumber v_j , and the original gas concentration, W_o , MORTRAN will compute theoretical transmissions, $T1(W_i, v_j)$, for the desired concentrations W_i .

Used in the computation is Beer's Formula,

$$T1(W_i, v_j) = T(W_o, v_j)^{W_i/W_o},$$

where $Tl(W_i v_j)$ is a function of the set of transmission values for each desired concentration.

Input

Card A

	Column	Field	Description
	1-10	F10.2	Beginning wavenumber
	11-20	F10.2	Ending wavenumber
	21-30	F10.2	Wavenumber increment
	31-40	E10.3	Original concentration,
	46-50	15	Woo NW, number of desired concentrations to be used in signal run
Card B	1-10	E10.3	1 st desired
	11-20	E10.3	concentration, W ₁ 2 ^d desired concentration, W ₂

		MORTRAN - (cont.)
Column	<u>Field</u>	Description
•	•	•
•	•	•
71-80	E10.3	i th desired
		concentration, W
$\frac{Card}{c_i}$ C_n		1
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	F10.0	Wavenumber, v _i
11-20	F10.0	Transmission,
		T(W _o ,v _i)
		. 0. j.
Output		
Card A		
1-10	F10.2	Beginning wavenumber
11-20	F10.2	Ending wavenumber
21-30	F10.2	Wavenumber increment
31-40	E10.3	Concentration, W_{i}
		i = 0; i = 1, NW
Card B _i -B _n		
1 1 1 1-5	15	Sequence number
6-10	3PF5.0	Transmission,
		$T(W_i, v_i)$
11-15	3PF5.0	Transmission,
		$T(W_i, v_{j+1})$
•	•	•
- -	•	•
76-80	3PF5.0	Transmission,
	-	$T(W_{i}, v_{j+14})$
		`1' J+14'

MORTRAN - (cont.)

SUBROUTINES USED: None.

MORTRAN: Computes additional transmissions and outputs in packed format.

```
PROCEAT MORTERY
       10005) 11, (5006) V. (6006) T. (11) MS NOTEN SENT
01
       READ SE, BV. F /. TV. W. NN
       IF (EOF,60)850.77
9.2
       FOPMAT(3F10.2.F10.3.5X.IS)
33
       NT=1.000001+(TY-0V)/OV
       NWA= XAPSE (NW)
       IF (NT.LT.5000)74. 100
14
       READ OF, (WM(T), T=1.NWA)
05
       FORMAT (SE10.3)
       30 65 I=1,NT
       READ OF, V(T), T(I), WT
06
       FORMAT(F10.2,510.3,43x,610.3)
       IF (NW.LE. 3)87,7FF
065
       IF (W.EQ.WI) 07.710
0.7
       IF (V(I).FO.RV) 79,08
09
       X=V(I-1)+DV
       (I) V=Y
       IF (APS(Y-X).t (.. 0001) 69,910
09
       CONTINUE
       NCARES=NT/1F
       IF (NOAFDS#1F.NF.NT) 21.22
21
       NCARES=NCARDS+1
22
       I3=1
       IE=IE+14
       PRINT 3001,8V,FV.DV.W
      PUNCH 20.BV.EV.DV.W
      JO 259 K=1+MC499S
      PUNCH 30, K, (T(4), M=18, IE)
      PRINT 3032,K, (T(M), M=19,1E)
      IB=IB+15
      IE=XMINOF(IF+15,NT)
259
      CONTINUE
      00 40 J=1,NWA
      HY(L)MH=TAX3
      00 10 T=1,NT
10
      T1(I)=T(I) ** EYPT
      PUNCH 28, BV, TV, DV, WN (J)
20
      FORMAT(3F10.2,F10.3)
      PRINT 3001, RV.EV.DV.WN(J)
3001
      FORMAT(35X.3F17.2.E17.3)
      IB=1
      IE=18+14
      DO 40 K=1,NCAP95
      PUNCH 30, K, (T1(M), M=IB, IE)
30
      FORMAT(15, (15(3PF5.0)))
      PRINT 3002, K, (*11(L), L=16, I5)
3002
      FORMAT (2X, IE, (15 (3PF5.0)))
      IB=IB+15
      IE=IE+15
      IE=XMINOF(IE, YT)
      CONTINUE
40
      GO TO C1
850
      PRINT 851
851
      FORMAT(12(/), 70x, *SOPRY, THERE ARE NO MORE CAROS*)
```

I

```
CALL EXIT

900 PRINT 901, NT, 3V, EV, 5V

901 FORMAT (20X*ERROR IN CARDS*//7X*NT=*, T5, *FV=*, F10.2, 2X*EV=*, F10.2, B 2X, *DV=*, F10.2)

GALL EXIT

910 PRINT 911, W, VT, Y, X

911 FORMAT (10X*ERROR IN DATA*//10X, 2E10.3/1JX, 2F1J.3)

CALL EXIT

TNO

SINGLE-BANK COMPILATION.
```

APPENDIX 2

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		EPAGAS CALC CLOSER CONST CXMIZ EROR1 EROR2 FINDIT FINDTRN GAUSSL GRAPHS IBINSER MINMYZD PARTIN PLOTD PLOTT PRINT1 PRINT2 PRINT3 READALL READ1 READ2 READ3 READ4 READ6 READ7 READ8 RMSERR THEOTRN		
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APPENDIX 2

A2. PROGRAM EPAGAS DOCUMENTATION ANALYSIS

A2.1 PROGRAM SUMMARY

In this appendix, the main minimization program EPAGAS and all of the subprograms needed by EPAGAS are described. All coding was done in the FORTRAN applicable to the CDC 3800 computer. Each description lists what that particular subprogram does, what input parameters are needed, and what output parameters are produced.

The computer program must be given: (1) a library of theoretical transmissions for each gas constituent which might possibly be present in the measured total transmissions. For each constituent, the theoretical transmissions are known functions of both wavenumber and gas concentration. The library must be large enough to bound the regions of interest in both wavenumber and concentrations to be expected. See THEOTRN description for exact preparation of the library table. Due to possible computer memory limitation, the user may wish to "tailor-make" separate libraries for different experimental situations. However, care must be taken in selecting proper wavenumber spacings and individual concentrations so that double linear interpolation within the transmission table will result in correct transmissions values for wavenumbers and concentrations lying between grid points of the library table.

(2) Next, a set of measured experimental total transmissions TG as a function only of known wavenumber ν must be provided. This spectrum, obtained from the spectrophotometer, will be compared to a trial spectrum TC which is a function of gas constituent, gas concentration W, and wavenumber ν as determined from the library of theoretical transmissions. By adjusting the amounts of W for each gas constituent as a function of wavenumber, the program tries to get TC to agree as best as possible with TG, thus determining the gas composition and the amounts of each gas constituent that were present in the sample producing the total transmission, TG. READ7 and/or READ8 descriptions provide specific information on the preparation details for inputting TG.

- (3) The program must also be provided with a starting set of gas concentrations W for each gas constituent that is suspected to be present in the sample producing the given experimental total transmission TG. Also given is a corresponding set of concentration increments, WI, which are used when the minimization process tries to vary the individual W_i by an amount WI_i to find the best agreement between TG and TC. See the READ6 description for preparation of W and WI.
- (4) Along with the initial values of W and WI, the program also needs error bounds for each of the individual gas constituent concentrations. See EROR1 description for detail error formulas. The errors defined on the gas concentrations are used primarily to help guarantee an acceptable solution by keeping the gas concentrations within physically reasonable bounds. READ2 description provides details for inputting the error parameters.
- (5) Finally, various program controls must be provided. Some of these control parameters are used in directing the minimization process and are listed in detail in the READ3 description. The remaining control parameters are used in directing printing and plotting of intermediate results and in providing certain convergence criterion. Details for these parameters are found in the READ4 description.

A2.1.1 <u>List of Fortran Symbols and Descriptions</u> Single-Valued Variables

<u>Variable</u>	Description
СҒМ	A constant confidence factor to be applied to the experimental transmission data if individual ones
GLB	are not specified. The minimum root-mean-square (rms) error permitted
	before using the minimization routine MINMYZD again.

Variable	Description
IPLOT	A plotting control which, when set to m, permits a graph of the given and currently computed transmissions, as functions of wavenumber, to be plotted after every m th use of MINMYZD.
IPRINT	A printing control which, when set to n, permits printing of intermediate results after every n th use of MINMYZD.
MK	The maximum number of uses of MINMYZD permitted for a particular data set.
N	N = NG + NT. The total number of errors in the error array ERR.
NG	The total number of gas constituents to be found in the library of theoretical transmissions.
NT	The number of experimental transmission values used.
Q	The minimum percent by which each try of MINMYZD must improve the mean square deviation.
RMS	Current root-mean-square (rms) error computed from the error array ERR.
RMSTP	Lowest root-mean-square (rms) error computed from the error array ERR.
RMST	Lowest root-mean-square (rms) error computed from the NG+1 through NG+NT elements of the error array ERR. This rms value corresponds to the errors based on the difference between the given experimental transmission values and the theoretical transmission values currently computed.
RMSW	Lowest root-mean-square (rms) error computed from the 1 through NG elements of the error array ERR. This rms value corresponds to the errors defined on the individual gas concentrations.

Description		
Beginning wavenumber corresponding to the given		
transmission values if they are equally spaced in wavenumber.		
End wavenumber corresponding to the given		
transmission values if they are equally spaced in wavenumber.		
Wavenumber increment corresponding to the given		
transmission values if they are equally spaced in wavenumber.		
Maximum wavenumber value for which the given		
transmission values will range over.		
Minimum wavenumber value for which the given		
transmission values will range over.		
A number used by MINMYZD which tells how much of the		
predicted final step to take.		

Array Variables

	<u>Maximum</u>	
Variable	dimension	Description
CF	150	Confidence factor, $0 \stackrel{<}{-} CF(K)$, $K=1$, NT , corresponding to the NT given transmission values.
ERR	160	Errors based on the individual gas concentrations and the weighted differences between the given and computed transmission values. Elements 1 through NG are the errors corresponding to the gas concentrations, and elements NG+1 through NG+NT are the errors corresponding to the transmission errors.

	Maximum	
Variable	dimension	Description
IDGAS	10	Integer numerical identifiers (I2)
		for each of the gases to be found in
		the gas library of theoretical
		transmissions. IDGAS(K), K=1, NG
		correspond to the K $^{ extsf{th}}$ gas.
IPW	10	One of the error parameters used in
		computing the errors based on the
		individual gas concentrations. This
		parameter is an odd positive integer
		used as a power in the error
		function. IPW controls the rate of
		increase of the error as the current
		gas concentration ventures outside
		the acceptable range for that
		particular concentration. IPW(K),
		$K=1$, NG correspond to the $K^{ ext{th}}$ gas.
KX	10	The first six elements of the array KX
		are input control parameters used by
		the main minimization routine MINMYZD.
		The last four elements of the array are
		output parameters of MINMYZD and
		provide various convergence information.
		<pre>KX(1) is the maximum number of new sets</pre>
		of errors (minimization cycles) to use.
		If KX(1)=0, then one cycle is used. If
		KX(1)<0, then each gas concentration
		increment WI is reduced by 10 percent after
		each complete use of MINMYZD, and
		KX(1) is used for the maximum number
		of cycles to be used in one complete
		use of MINMYZD.

Maximum

Variable

dimension

Description

- KX(2) is the number of cycles to count
 for a bad cycle, that is, one which did
 not result in enough improvement,
 KX(2)-1.
- KX(3) is the percent improvement in tenths in the rms error for the cycle to be counted as a good cycle. Any cycle resulting in less than this causes KX(2) cycles to be counted rather than one cycle, KX(3)-0.
- KX(4) is the limit as to the multiple of a WI increment which may be taken in one full step (ZEP=1). If KX(4)=0, then 100 is used. If KX<0, then no step will be permitted in the -WI direction for the first complete use of MINMYZD. Future uses of MINMYZD uses KX(4).
- KX(5) is the minimum ZEP in 0.001's
 permitted. KX>0.
- KX(6) is the minimum total percent improvement in tenths for ZEP not to be changed. KX(6)>0.
- KX(7): If KX(7)=1, then at least one single step (even if it were a step of WI_i) resulted in an improvement of at least KX(3); otherwise, KX(7)=0.
- KX(8): If KX(8)=1, then there was some
 improvement sometime; otherwise,
 KX(8)=0.

	Maximum	
Variable	dimension	Description
		KX(9): If $KX(9)=-1$, then all WI's
		were too small. Convergence has
		probably been reached. If $KX(9)=0$,
		then the solution matrix was
		singular. If $KX(9)=1$, then the matrix
		was normal.
		KX(10) is the actual total percent
		improvement in tenths.
NU	10	Number of wavenumber values used for
		each gas in the library of theoretical
		transmissions. $NU(K)$ is the number
		of wavenumber values used for the
		K^{th} gas, K=1, NG.
NW	10	Number of concentration values used
		for each gas in the library of
		theoretical transmissions. NW(K) is
		the number of concentrations used for
		the K^{th} gas, $K=1$, NG.
S	5	An array used by READALL and its
		associated reading routines. If $S(K)=0$,
		then the input information was correct.
		If $S(K)\neq 0$, then a reading error occurred.
TC	150	Computed theoretical transmission
		values, $TC(K)$, $K=1$, NT based on the
		NG current gas concentrations being
		used. TC(K) corresponds to the K th
		value. $TC(K) = \frac{NG}{II}$ $TCI(I,K)$.
TCI	10	Individual compu- $^{I=1}$ ted theoretical
101		transmission values TCI(I,K) for a
		given wavenumber U(K), K=1, NT and
		given gas concentrations W(I), I=1, NG.
		0-1 0 1

	Maximum	
<u>Variable</u>	dimension	Description
TG	150	Given experimental transmission values
		TG(K), $K=1$, NT as a function of
		known wavenumber $U(K)$, $K=1$, NT .
TRAN	(5000,5)	A two-dimensional array which contains
		all of the theoretical transmission
		values that make up the gas library.
		The transmission values are a
		function of known wavenumbers,
		known gas constituents, and known
		concentrations for each of the
		specified gas constituents. See
		THEOTRN description for exact
		formulation of the array.
U	150	Known wavenumbers, $U(K)$, $K=1$, NT
		corresponding to the array TG of the
		NT given experimental transmission
		values.
UR	(10,3)	A two-dimensional array UR(I,J)
		containing the wavenumber range and
		spacing used for each of the gas
		constituents in the gas library.
		I corresponds to the I th gas. J=1
		corresponds to the beginning wave-
		number, J=2 corresponds to the end
		wavenumber, and J=3 corresponds to
		the wavenumber increment.
WHR	10	One of the error parameters used in
		computing the errors based on the
		individual gas concentrations.
		This parameter is the half-range of
		permitted variation in concentration for the K th gas constituent and is
		for the K gas constituent and is

	Maximum	
Variable	dimension	Description
		simply the difference between the
		mean concentration WM(K) and the
		lowest concentration of the $K^{ extstyle{th}}$ gas
		constituent.
WIO	10	Original increments to be applied to
		the starting gas concentration WO(K),
		K=1, NG. These are needed by MINMYZD
		when it tries to adjust the original
		concentrations WO to get the best
•		agreement between the given and
		computed transmission values. These
		values may be modified in the
		minimization process, and hence
		become WI(K), K=1, NG.
WI	10	Current incremental values of gas
		concentrations, WI(K), K=1, NG.
WM	10	One of the error parameters used in
		computing the errors based on the
		individual gas concentrations. This
		parameter is the mean concentration
		value of the desired upper and lower
		bounds of the concentrations of the
		$K^{ extsf{th}}$ gas constituent, $K=1$, NG.
WO	10	Original gas concentrations to be used
		in the minimization process. WO(K)
		corresponds to the K th gas constituent,
		K=1, NG.

	<u>Maximum</u>	
Variable	dimension	Description
WR	(10,5)	A two-dimensional array WR(I,J)
		containing the actual gas
		concentrations used in the gas
		library. I corresponds to the I $^{ exttt{th}}$
		gas constituent, and J corresponds to
		the $\mathtt{J}^{ exttt{th}}$ gas concentration of the $\mathtt{I}^{ exttt{th}}$
		gas constituent.
W	10	Current gas concentrations being used
		in the minimization process. $W(K)$,
		K=1, NG corresponds to the K th gas
		constituent.
ww	10	One of the error parameters used in
		computing the errors based on the
		individual gas concentrations. This
		parameter is a "normalization"
		factor and is used to scale the gas
		concentration errors. WW(K), K=1,
		NG are the factor for the K th gas
		constituent.

A.2.1.2 List of Program and Subprogram Descriptions

FORTRAN PROGRAM CDC 3800

EPAGAS
Margot Ackley
July 30, 1973

Program EPAGAS is a computer program which is designed to determine atmospheric composition from measured total transmissions at known wavenumbers. The program utilizes a set of theoretical transmission spectra of selected gas constituents at known values of concentrations and at known wavenumbers, and a set of observed total transmissions at known wavenumbers only. By employing a particular minimization technique, EPAGAS tries to fit the observed transmissions to the theoretical transmissions by varying the concentrations of the individual constituents.

Assuming a certain concentration of each gas constituent in an atmospheric path, a corresponding transmission can be found from a large table of known transmissions as a function of gas constituent, wavenumber, and gas concentration. The total transmission at a specified wavenumber is then computed and compared to the observed transmission value obtained from a spectrophotometer. The difference between the computed and observed transmissions is then minimized by varying the concentrations of the individual gas constituents. The minimization process tries to reduce the rms error of an error function defined on the errors of the given and computed transmissions and on the concentrations. (See A2.3.) The concentration errors are used mainly to help keep the concentrations within such a range as to provide physically realistic solutions.

The program uses various controls to direct its progress in the minimization process. The initial and final gas constituent concentrations, computed transmissions, and errors are always printed together with a plot of the given and computed transmissions as a function of wavenumber. Intermediate results may also be printed and plotted.

For preparation of the theoretical gas library table, see the THEOTRN description. For preparation of the given transmission data and all

EPAGAS - (cont.)

other input needed by the program, see the READALL description. It is imperative that all descriptions pertaining to the program be read and understood.

USES:

CALC	IBINSER	READ1
CLOSER	MINMYZD	READ2
CONST	PARTIN	READ3
CXMIZ	PLOTD	READ4
EROR1	PLOTT	READ6
EROR2	PRINT1	READ7
FINDIT	PRINT2	READ8
FINDTRN	PRINT3	RMSERR
GAUSSL	READALL	THEOTRN
GRAPHS		

FORTRAN SUBROUTINE
CDC 3800
(Internal to EPAGAS)
CALC (W. ERR. RMS)

CALC
Margot Ackley
July 30, 1973

Given the array W of NG current gas concentrations and the selected arguments from the two COMMON blocks, COMMON/100/ and COMMON/101/, CALC computes an error array ERR of length NG + NT, where NT = the number of computed and given transmissions and NG = the number of gas concentrations. CALC also computes the rms error RMS needed by MINMYZD, and the individual rms errors RMSW for the error based only on the concentrations, and RMST for the error based only on the transmissions. (See A2.3.)

The two COMMON blocks utilized are as follows: COMMON/100/TRAN (5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); COMMON/101/NG, NT, RMSTP, RMST, RMSW.

The first NG elements of the array ERR are the errors based on errors from the NG gas concentrations. (See EROR1 description.) The NG + 1 elements through the NG + NT elements of the array ERR are the errors based on errors from the NT given and currently computed transmissions. (See EROR2 description.)

Before computing the individual transmission errors, CALC must first calculate the NT computed transmissions TC based on the current NG gas concentrations W given to CALC. CALC uses subroutine FINDTRN to find the individual computed transmission TCI(i,j) for a given wavenumber U(j) and a given gas concentration W(i). The total transmission for the particular wavenumber U(j) is then the product of all the transmissions for all the gas concentrations W(i) for the particular U(j) where:

$$TC(j) = \prod_{i=1}^{NG} TCI(i,j) .$$

CALC - (cont.)

If the current use of CALC produces an RMS error which is smaller than the previous rms error, then RMSTP = RMS, and the array TC will then contain the computed transmissions producing the lower RMS. If RMS is greater, then the values RMSTP and TC will contain the last set of values resulting in an improved rms error.

For definitions of arguments and variables used, see below descriptions:

EPAGAS: W

READ7(8): NT, U, TG, CF

THEOTRN: NG

USES: EROR1, EROR2, FINDTRN(IBINSER), and RMSERR.

CALC: Computes concentration and transmission errors for MINMYZD.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)

CLOSER (A, X, B)

CLOSER

J. R. Winkelman June 15, 1965

CLOSER is the value of X in the closed region A-X-B, taking on the values of the end points for X outside the range. More exactly, if X-A, the value of CLOSER will be A. If X-B, the value of CLOSER will be B. Otherwise, CLOSER=X.

USES: No subroutines.

CLOSER: Restrict X to a closed region.

FORTRAN SUBROUTINE

CONST .

CDC 3800

Margot Ackley

(Internal to EPAGAS)

July 30, 1973

CONST (S, UMIN, UMAX, ZEP, KX, Q, MK, GLB, CFM, IPRINT, IPLOT)

CONST sets certain constants needed by program EPAGAS and its associated subroutines. These values are as follows:

$$S(1) = S(2) = -1.$$

$$S(3) = S(4) = S(5) = 0.$$

WM(1) = 3.237E22

WHR(1) = 3.437E22

WM(2) = 1.235E20

WHR(2) = 1.235E20

WM(3) = 1.235E20

WHR(3) = 1.235E20

WM(4) = 1.235E20

WHR(4) = 1.235E20

WM(5) = 1.245E20

WHR(5) = 1.245E20

WM(6) = 0.715E20

WHR(6) = 0.715E20

$$WW(1) = WW(2) = WW(3) = WW(4) = WW(5) = WW(6) = 1$$

$$IPW(1) = IPW(2) = IPW(3) = IPW(4) = IPW(5) = IPW(6) = 1$$

UMIN = 1000.

UMAX = 1250.

$$KX(1) = 10$$

$$ZEP = 0.6$$

$$KX(2) = 3$$

$$Q = 10.0$$

$$KX(3) = 50$$

$$GLB = 3.0$$

$$KX(4) = 30$$

$$CFM = 10.0$$

$$KX(5) = 100$$

$$MK = 10$$

KX(6) = 100

IPRINT = 1

$$KX(7) = 0$$

IPLOT = 1

$$KX(8) = KX(9) = KX(10) = 0$$

.

CONST - (cont.)

The definitions of the above arguments are given in the below subroutine descriptions.

S READALL
WM, WHR, WW, IPW READ2
UMIN, UMAX READ1
ZEP, KX READ3
Q, GLB, CFM, MK READ4
IPRINT, IPLOT

CONST utilizes COMMON/100/ where: COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10).

USES: No subroutines.

CONST: Set values for EPAGAS.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)
CXMIZ (KX, G, GLB, MK, RMS)

CXMIZ
Margot Ackley
July 30, 1973

Given the array KX (see READ3 description), the arguments Q and GLB (see READ4 description), MK the remaining uses of MINMYZD to be permitted (see READ4 description), and RMS, the current rms error, CXMIZ checks to see if MINMYZD is to be used again.

If KX(9) = -1, then CXMIZ = 2.0 and returns. If KX(9) = 0, then CXMIZ = 3.0 and returns. If KX(9) = 1, then CXMIZ checks the following: if $MK^{-}0$, if $\frac{Q}{1} > KX(10)$, or if $RMS^{-}GLB$, then CXMIZ = 2.0.

If none of the above occur, then CXMIZ = 1.0 and returns. Where:

CXMIZ = 1, use MINMYZD again,

CXMIZ = 2, do not use MINMYZD again,

CXMIZ = 3, singular matrix, halt computation.

USES: No subroutines.

CXMIZ: Checks for future uses of MINMYZD in EPAGAS.

FORTRAN SUBROUTINE
CDC 3800
(Internal to EPAGAS)
EROR1 (W, ERR)

EROR1
Margot Ackley
July 30, 1973

Given NG values of W, where W(I) is the concentration of the I^{th} gas constituent and corresponding error parameters WM, WHR, WW, and IPW, EROR1 computes the NG errors ERR as defined below:

ERR(I) = WW(I)*
$$\left(\frac{W(I)-WM(I)}{WHR(I)}\right)^{IPW(I)}$$
I = 1, NG

WM, WHR, WW, and IPW are provided through COMMON/100/: COMMON/100/ TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10).

USES: No subroutines.

EROR1: Computes gas constituent errors for EPAGAS.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)

EROR2 (TG, TC, CF)

EROR2
Margot Ackley
July 30, 1973

EROR2 is the error computed from the discrepancy between the given transmission TG and the computed transmission TC, weighted by a confidence factor CF, where

EROR2 = CF*(TG-TC).

USES: No subroutines.

EROR2: Computes transmission errors for EPAGAS.

FORTRAN FUNCTION

CDC 3800

(Internal to EPAGAS)

FINDIT (A, L, M, N)

FINDIT

J. R. Winkelman

December 18, 1962

The value of FINDIT will be that A(L) which would be obtained if the array A was sorted between A(M) and A(N). A(L) has its correct new value, that is, $A(K) \stackrel{<}{-} A(L)$ for $M \stackrel{<}{-} K \stackrel{<}{-} L$, $A(K) \stackrel{>}{-} A(L)$ for $L \stackrel{<}{-} K \stackrel{<}{-} N$. Thus, the array is in better sort on exit from FINDIT. The array A outside of the range M, N is unchanged.

Example: To find the 20 percentile and median of the array from A(10) to A(20),

PMED = FINDIT (A, 15, 10, 20) P20 = FINDIT (A, 12, 10, 14) or FINDIT (A, 12, 10, 20).

USES: PARTIN.

FINDIT: Find the Lth sorted value.

FORTRAN SUBROUTINE
CDC 3800
(Internal to EPAGAS)
FINDTRN (W, UP, TT)

FINDTRN
Margot Ackley
July 30, 1973

Given the arguments W and UP and selected variables from the two COMMON blocks--COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW--FINDTRN finds the corresponding theoretical transmissions TT.

The array W contains the NG current values of gas concentrations, and UP is some given wavenumber within the range of interest. The computed theoretical transmissions are placed in the array TT, which has a maximum length of 10, where TT(I) corresponds to the theoretical transmissions from the gas library as a function of wavenumber UP and concentration W(I) of the Ith gas.

Because the current concentrations W and wavenumber UP may not necessarily correspond to those wavenumbers and concentrations in the gas library table TRAN (see THEOTRN description), FINDTRN uses double linear interpolation within the two-dimensional table TRAN to determine the transmission TT(I) corresponding to W(I) and UP.

Wavenumber UP will never lie outside the range of the table. However, the concentrations W could possibly lie outside the range due to the minimization process which varies only the array W. In the event that the individual W(I) lies above the range of the concentrations of the Ith gas, the highest two concentrations of that gas are used to determine a transmission value TT(I) for W(I) and UP. In the case where W(I) lies below the range of the table, the smallest concentration in the table and a concentration of zero are used in determining the corresponding transmission. The interpolation formula for this case utilizes the fact that, by definition, a transmission value for zero concentration is 1. (See A2.4.) The interpolation process can possibly

FINDTRN - (cont.)

result in transmissions outside the range 0 to 1, but the structure of the error functions tends to discourage selections of those concentrations W which make this occur.

USES: IBINSER.

FINDTRN: Finds theoretical transmissions as a function of wavenumber and concentrations for EPAGAS.

FORTRAN SUBROUTINE
CDC 3800
(Internal to EPAGAS)
GAUSSL (A, NRD, NR, NC, D)

GAUSSL L. David Lewis

January 20, 1970

Given an augmented matrix A, its row dimension NRD (from the DIMENSION statement defining A), the number of rows NR (equals the order of the matrix), and the number of columns NC (equals NR plus the number of vectors);

<u>Find D</u>, the determinant of the matrix and the solution vectors of the indicated linear system. The given matrix is destroyed, and the given vectors are replaced by the solution vectors.

Restrictions: A must be stored columnwise;

NC-NR, NR-NRD. If NC = NR, only D is computed.

Method: Gauss elimination with pivotal condensation.

Error conditions: D = 0, the matrix was singular; the augmented matrix has been destroyed.

USES: No subroutines.

GAUSSL: Determinant of a matrix by Gauss elimination and solution.

FORTRAN SUBROUTINE
CDC 3800
(Internal to EPAGAS)
GRAPHS (A, B, Y, NGC, NXC, NUC, NOC, NBC, NQ)

GRAPHS
Margot Ackley
July 30, 1973

Edits an array NQ of 101 alphanumeric characters (R1 format) to allow computer plotting of functions.

For A<B, A is the minimum ordinate associated with NQ(1) and B is the maximum ordinate associated with NQ(101). Y is the value to be plotted. If $A \le Y \le B$, the quantity D = 1 + 100*(Y-A)/(B-A) is computed and rounded to the nearest integer I. Then NQ is examined. If NQ(1) contains the blank character NBC or a decimal point (.), then the graph character NGC is stored there; if NQ(I) does not contain NBC or a decimal point, then the intersection character NXC is stored in NQ(I). If Y is outside the range A,B, then I is computed as though Y were in the range A + n*(B-A), B + n*(B-A) for appropriate n. Then if NW(I) \neq NBC, NXC is stored in NQ(I); otherwise if Y<A, the underflow character NUC is stored in NQ(I), and if Y<B, the overflow character NOC is stored in NQ(I). If a character is zero, NQ will be left unchanged. If B<A, the computation proceeds as though A and B were interchanged and I was replaced by 101-I.

In addition, if the value of D before rounding differed from an integer by more than 3/8, NGC (or NXC, NUC, or $N\overline{OC}$ as appropriate) will be stored in NQ(I+1) or NQ(I-1) so that the two adjacent characters will, when plotted, straddle the proper position. If the value of D before rounding differed from an integer by an amount between 1/8 and 3/8, a decimal point (.) will be stored in NQ(I+1) or NQ(I-1) so that it, along with the graph character NGC stored in NQ(I), will indicate the direction in which the plotted character should have been moved by a slight amount. A decimal point will be stored, however, only if it replaced a blank character NBC.

GRAPHS - (cont.)

In use, to plot n functions, GRAPHS would be called n times, with the n function and scale values, then a line would be printed. A suitable print statement might be: WRITE OUTPUT TAPE 61, 99, X, (NQ(I), I=1, 101), 99 FORMAT (1HbE16.9, 2X, 101RI), where X is the abscissa value or perhaps one of the function values.

Restrictions: A must not equal B; zero (0) cannot appear as a plotted character.

USES: No subroutines.

GRAPHS: Plot graphs on the line printer.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)
IBINSER (X, N, P)

IBINSER
Margot Ackley
July 30, 1973

Given an array X of length N containing monotonically increasing numbers and given a number P, IBINSER performs a binary search in array X to find the integer value I for which $X(I)^{-}P>X(I+1)$. IBINSER is then set equal to I, that is, $1^{-}IBINSER^{-}N$. If N=2, IBINSER=1.

USES: No subroutines.

IBINSER: Binary search of a monotonically increasing array.

MINMYZD

CDC 3800

R. Gregg Merrill
(J. R. Winkelman)

(Internal to EPAGAS)

March 20, 1970

MINMYZD (W, WI, NG, CALC, ERR, N, ZEP, KX, PRNT, RMS, XTM)

This subroutine is used to minimize a function of several variables when the true value of the function is known at several points. In other words, it fits an arbitrary function of NG variables to N known points. For example, many positions of a satellite over a few days may be known and the elements (period, major axis, etc.) are desired which will give an orbit for which the rms distance from the given position is a minimum.

Because the convergence procedures and flexibility are much greater than would be required on "nice functions," a suggested set of control factors KX will be given so that the average programmer does not need to become familiar with all the options. (See summary.)

In detail, the subroutine starts with a given set of NG-10 parameters W and an increment WI for each W. Within the WI array, a zero implies that the corresponding W is not to be changed. Also given are the N-160 errors ERR (an error at each of the given points where these are of no concern to MINMYZD) and an rms error RMS where RMS may be obtained by RMS = RMSERR(ERR,N). NG may be 1, but N must be $\stackrel{>}{\sim}$ NG.

MINMYZD makes a step in W_i of WI_i and looks at the resultant errors obtained by calling CALC(W,ERR,RMS). If the RMS for this new set is smaller, the slopes of the individual errors with respect to WI_i are calculated and stored. If larger, a step of WI_i is made in the other direction and slopes are calculated. If neither RMS is smaller, WI_i is halved for future use, as the step size is probably too large. If no errors changed at all, WI_i is set to zero (normally because WI was halved out of existence and is no longer important to the minimization process).

After all W's have been varied, the matrix

$$(\partial ERR_{i}/\partial W_{j})*(\Delta W_{j}) = -(ERR_{i})$$
 (1)

is solved for the changes ΔW_i in W. The new parameters are then $W_i = W_i + ZEP * \Delta W_i$. If the important $\partial ERR_i / \partial W_j$ is nearly constant, ZEP should be 1. As a minimum is approached, ZEP must be made smaller. This is done internally under control of the control values KX.

After the new ERR's and RMS have been calculated, PRNT is checked. If PRNT is zero, no internal printing is done. If PRNT \neq 0, a line is printed specifying the ZEP used and the RMS corresponding to that ZEP. It also prints a ZEP and RMS which would have theoretically minimized the curve (slopes are known for ZEP = 0, and the theoretical value RMS p for any ZEP can be compared against the actual RMS evaluated at the given ZEP to predict the second derivatives and the corresponding minimum).

If at any time a new RMS is less than the old one, the W and ERR arrays are replaced by the better values and the necessary adjustments made to KX (7 and 8).

The KX array contains six input control parameters and four output parameters. These control the action of MINMYZD. None of the controls KX(1-6) are altered by the program.

- KX(1) The total number of cycles to take, even if conditions are still good.
- KX(2) The number of cycles to count for a cycle which does not cause enough improvement.
- KX(3) The acceptable improvement in 0.1 percent (105 is 10.5 percent).
 Any cycle resulting in less than this causes KX(2) cycles to be counted rather than one cycle.

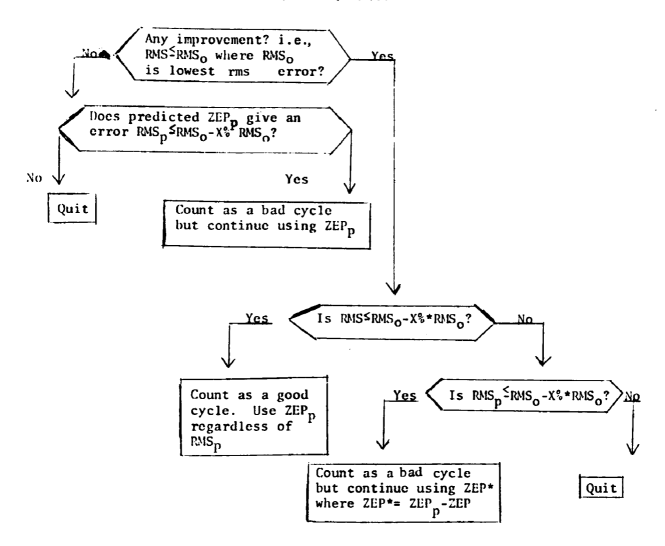
MINMYZD - (cont.)

- KD(4) The limit as to the multiple of a WI increment which may be taken in one full step (ZEP=1). A zero will be used as 100. A small slope might predict a step of 10⁶ which is clearly too large. If KX(4) is negative, no step is made in the -WI direction. It is normally faster to have KX(4)<0 for the first few uses of MINMYZD.
- KX(5) Minimum ZEP to be used in 0.001 (10 is 0.010). If ZEP is allowed to go too small (alas, the predictions are not always good) the step does not go anywhere. If ZEP minimum is too large, ZEP cannot be made small enough to account for the nonlinearity in ERR for final convergence.
- KX(6) If good improvements are made, it is natural for ZEP to become smaller as MINMYZD moves away from the region in which the derivatives were calculated. If the total improvement KX(10) is greater than KX(6) in 0.1 percent, then the ZEP at exit will be the same as the ZEP on entry, that is, things are going well so stick to the same ZEP. Otherwise, exit with the last predicted ZEP_D.
- KX(7) through KX(10) are used to convey certain information to the user.
- KX(7) If 1, at least one single step (even if it were a setup of WI_i) resulted in an improvement of at least KX(3); otherwise, it is zero.
- KX(8) If 2, there was some improvement sometime; otherwise, zero.
- KX(9) If -1, the WI's are all too small; the system has probably converged. If 0, the matrix was singular; this is not expected to occur. +1 is normal.
- KX(10) Total improvement in 0.1 percent relative to the RMS input.

 750 would mean the new RMS was 4 of the old RMS.

After PRNT is checked, the improvement is checked against KX(3). If sufficient, one is subtracted from the cycle count set up from KX(1).

A ZEP gives an error RMS: (X = 0.1(KX(3)))



MINMYZD - (cont.)

If the count is -1, the new ERR is put into equation (1) and the system continues from there using the predicted ZEP_p. Any time the cycle count reaches zero, an exit is made.

If the improvement is not good enough, KX(2) is subtracted from the cycle count. If the count $\stackrel{<}{\sim}0$ or the predicted RMS is also not good enough, an exit is made. Otherwise, a calculation is made using a ZEP based on the old ZEP and the predicted ZEP, hoping for an improvement. If no improvement is made but the predicted RMS is good enough, then calculations are made using the predicted ZEP, otherwise, an exit is made. (See flow chart for termination of MINMYZD.)

Helpful hints: If KX(2) is one less than KX(1), then, after a "good" cycle, one bad cycle will end it. If a "bad" cycle occurs immediately, then one more try with the adjusted ZEP will usually give good results. On complicated functions (almost anything will work on simple ones), it may be necessary to set KX(5) higher from 100 to 300 for a few steps if ZEP is being predicted too small. The XTM matrix, currently (160,11), is the partial derivatives matrix XTM(I,K) = $\partial ERR_1/\partial W_k$, I=1, N, K=1, NG, where K is the Kth non-zero WI, that is, there are no zero columns.

If one or more rows of XTM completely dominate, it may result in a singular matrix. This condition is checked, and the matrix is weighted if necessary. The weights are stored in the column (KK) following the normal information. This number XTM(I,KK) will normally be 1. If it is less than 1, then the Ith row has been multiplied by XTM(I,KK) and appears smaller than when the row was actually calculated.

SUMMARY:

Given: Arrays W and WI, NG words long where W contains the parameters, WI contains the suggested increments, and a zero WI means that the corresponding W is not to be varied; a subroutine CALC, which given W, calculates an error array ERR (positive and/or negative), N long, and the rms error RMS; a variable ZEP; and values for PRNT, KX(1), KX(2), KX(3), KX(4), KX(5), and KX(6).

MINMYZD exits with improved values of W; WI may be slightly altered; the array ERR of new errors associated with W; RMS correspond to ERR; ZEP may be a better value (must be smaller toward the end); KX(7)=1, at least one single step exceeded 0.1 percent improvement (from KX(3)) or else is zero; KX(8)=1 some improvement occurred sometime or else is zero; KX(9)=1), all is well, =0 unexpected, =-1 all WI's reduced to zero; you are done, congratulations! KX(10) the relative improvement for this one step of MINMYZD is 0.1 percent.

Normally KX(9) should be tested, and then test KX(7,8,or 10) or RMS to determine whether to call MINMYZD again or to quit as good enough. Note that all values needed for input are proper as returned from MINMYZD, so only a call is necessary, no further work.

<u>USES</u>: CALC, CLOSER, internal MINMYZD (FINDIT (PARTIN)), and GAUSSL. MINMYZD: To minimize an arbitrary function of several variables.

PARTIN

CDC 3800

J. R. Winkelman

(Internal to EPAGAS)

December 18, 1962

PARTIN (K, L, M, N, I, J)

Given the array K, a location in it K(L), and the limits of the area of concern K(M) to K(N), PARTIN shuffles the array until the following inequalities hold:

$$K(IJ) \stackrel{<}{-} K(L)$$
 for $M \stackrel{<}{-} IJ \stackrel{<}{-} J$
 $K(IJ) \stackrel{>}{-} K(L)$ for $I \stackrel{<}{-} IJ \stackrel{<}{-} N$

$$K(IJ) \stackrel{>}{-} K(L)$$
 for $I \stackrel{<}{-} IJ \stackrel{<}{-} N$

where M-J-I-N. I and J are output.

PARTIN uses fixed-point arithmetic internally so that it handles fixed and floating arrays without prejudice.

USES: No subroutines.

PARTIN: Partition (partial sort) an array.

PLOTD

CDC 3800

Margot Ackley

(Internal to EPAGAS)

July 30, 1973

PLOTD (W, IDGAS)

Given the above arguments and selected arguments from the two COMMON blocks, COMMON/100/ and COMMON/101/, PLOTD plots an on-line graph of the difference between the given transmissions TG and final computed transmissions TC versus the wavenumber U as a function of the final set of gas concentrations W. Scale for the transmission differences is set from -0.10 to +0.10.

The two COMMON blocks utilized are as follows: COMMON/100/TRAN (5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

For definitions of arguments and variables used, see below descriptions:

EPAGAS:

W

THEOTRN:

IDGAS, NG

READ7(8):

NT, U, TG

CALC:

TC

PLOTD starts a new page.

USES: GRAPHS

PLOTD: Plots difference of given and final computed transmissions versus

wavenumber.

PLOTT

CDC 3800

Margot Ackley

(Internal to EPAGAS)

July 30, 1973

PLOTT (W, IDGAS, NM)

Given the above arguments and selected arguments from the two COMMON blocks, COMMON/100/ and COMMON/101/, PLOTT plots an on-line graph of the given transmissions TG and corresponding computed transmissions TC versus the wavenumber U as a function of the current values of the gas concentrations W. Scale for transmissions is set from 0 to 1.

The two COMMON blocks utilized are as follows: COMMON/100/TRAN (5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

For definitions of arguments and variables used, see below descriptions:

EPAGAS:

W

THEOTRN:

IDGAS, NG

READ7(8):

NT, U, TG

CALC:

TC

If NM=0, a title line stating that the plot is before use of MINMYZD is printed; if NM>0, a title line stating that the plot is after NX uses of MINMYZD is printed; and if NM<0, a title line stating that the plot is after final use of MINMYZD is printed.

PLOTT starts a new page.

USES: GRAPHS

PLOTT: Plot curves of given and currently computed transmission versus

wavenumber.

PRINT1

CDC 3800

Margot Ackley

(Internal to EPAGAS)

July 30, 1973

PRINT1 (Q, GLB, CFM, MK, IPRINT, IPLOT, UMIN, UMAX, ZEP, KX, IDGAS, W, WI)

Given all of the above arguments and selected arguments from the two COMMON blocks, COMMON/100/ and COMMON/101/, PRINT1 prints all the input data for program EPAGAS.

The two COMMON blocks are as follows: COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

For definitions of arguments and variables printed, see below descriptions:

READ4:

Q, GLB, CFM, MK, IPRINT, IPLOT

READ1:

UMIN, UMAX

READ3:

ZEP, KX

THEOTRN:

IDGAS, NG

READALL:

W, WI

READ2:

WM, WHR, WW, IPW

READ7(8):

NT, U, CF, TG

PRINT1 starts a new page.

USES: No subroutines.

PRINT1: Prints all input for EPAGAS.

PRINT2

CDC 3800

Margot Ackley

(Internal to EPAGAS)

July 30, 1973

PRINT2 (NM, Q, GLB, MK, ZEP, KX, RMS, IDGAS, WO, W, WI, ERR)

Given the above arguments and selected arguments from the two COMMON blocks, COMMON/100/ and COMMON/101/, PRINT2 prints the intermediate data and errors before the first use of MINMYZD and after any successive uses of MINMYZD.

The two COMMON blocks are as follows: COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

For definitions of arguments and variables printed, see below descriptions:

READ4:

Q, GLB, MK

READ3:

ZEP, KX

CALC:

RMS

THEOTRN:

IDGAS, NG

READ6:

WO

READALL:

W, WI

CALC:

ERR, RMST, RMSW, TC

READ2:

WM, WHR, WW, IPW

READ7(8):

NT, U, CF, TG

If NM=0, a title line stating that the data is before use of MINMYZD is printed. If NM>0, a title line stating that the data is after NM uses of MINMYZD is printed.

PRINT2 starts a new page.

USES: No subroutines.

PRINT2: Prints intermediate data and errors for EPAGAS.

PRINT3

CDC 3800

Margot Ackley

(Internal to EPAGAS)

July 30, 1973

PRINT3 (IDGAS, WO, W, ERR, RMS)

Given the above arguments and selected arguments from the two COMMON blocks, COMMON/100/ and COMMON/101/, PRINT3 prints the final data and errors.

The two COMMON blocks are as follows: COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/100/NG, NT, RMSTP, RMSW.

For definitions of the arguments and variables printed, see below descriptions:

THEOTRN:

IDGAS, NG

READ6:

WO

READALL:

W

CALC:

ERR, RMS, RMST, RMSW, TC

READ2:

WM, WHR, WW, IPW

READ7(8):

NT, U, CF, TG

PRINT3 starts a new page.

USES: No subroutines.

PRINT3: Prints final data and errors for EPAGAS.

FORTRAN SUBROUTINE

CDC 3800

Margot Ackley

(Internal to EPAGAS)

READALL (S, UMIN, UMAX, ZEP, KX, Q, GLB, MK, CFM, IPRINT, IPLOT, W, WI,

WO, WIO)

Given the array S of length 5 (see CONST description), READALL reads in and checks input data to be used by EPAGAS. Each of the seven different types of data must be preceded by the appropriate header card containing the card type number in column 1.

	Error array	
Card type	element	FORTRAN reading subprogram
. 1	S(3)	READ1 (UMIN, UMAX)
2	S(4)	READ2 (FAKEARG)
3	none	READ3 (ZEP, KX)
4	none	READ4 (Q, GLB, CFM, MK, IPRINT,
		IPLOT)
6	S(1)	READ6 (WO, WIO)
7	S(2)	READ7 (UMIN, UMAX, CFM)
8	S(2)	READ8 (UMIN, UMAX, CFM)

S(K) = 0 indicates no error in reading and $S(K) \neq 0$ indicates that an error has occurred. If a card type 5 is read in, S(5) = -1 and indicates an error. See above subprogram descriptions for preparation of data. If 1, 2, 3, 4 type data is not read in, then those values given by CONST will be used in computation. Type 6 data must initially be read in, and type 7 or 8 (but not both) must also be initially read in. As many sets of data as desired may be processed. However, a type 9 card must follow every set of data (9 in column 1). This card will cause checking

of data and begins computation if no reading errors occur. READALL provides printed error analysis for any errors.

Arrays W and WI are set equal to the arrays WO and WIO. These are the gas concentrations W with corresponding increments WI, and the original concentrations WO with corresponding original increments WIO. If no concentrations are read in after the initial set, then each new set of data will be processed, starting with the original set of concentrations WO and increments of WIO. If one wishes to use the final computed set of concentrations of the last data set as the starting concentrations of the next data set, this may be accomplished by using READ6 with the appropriate header card, followed by a card with a 2 in column 80. This last set of concentrations and increments will then become the original concentrations and increments for all future sets of data.

After the initial set of data, sets of data need only to contain those values which one wants to vary. The unchanged values which will be used in processing the new data set will be those which had last been read in.

READALL utilizes two blocks of COMMON: COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

<u>USES</u>: READ1, READ2, READ3, READ4, READ6, READ7, and READ8.

READALL: Reads in and checks data for EPAGAS.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)

Margot Ackley
July 30, 1973

READ1

READ1 (UMIN, UMAX)

READ1 reads in the minimum wavenumber value UMIN and the maximum wavenumber value UMAX for which the given transmissions will range over.

If UMIN-0 and/or UMIN-UMAX, then an error has orrcurred and READ1 returns equal to -1. If no error occurs, then READ1 = 0.

CARD FORMATS:

CONTROL CARD-read by READALL

Col.	<u>Field</u>	Description
1	Integer	Card type - use 1
	DATA CARD-read by READ1	
1-10	Floating	UMIN>0
11-20	Floating	UMZX>UMIN

USES: No subroutines.

READ1: Reads in wavenumber range for EPAGAS.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)
READ2 (FAKEARG)

READ2
Margot Ackley
July 30, 1973

Given NG, the number of gas constituents, READ2 reads in the necessary gas concentration error parameters to be used by EROR1. NG values each of WM, WHR, WW, and IPW are read in.

If WW(K)<0, WM(K)<0, WHR(K)-0, and/or IPW(K)-0 for K=1, NG, then a reading error has occurred and READ2 returns equal to -1. If no error occurs, then READ2=0. See EROR1 description for detailed use of WW, WM, WHR, and IPW. FAKEARG is a fake argument and has no real use in the function. It is set to zero.

READ2 utilizes two blocks of COMMON: COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

CARD FORMATS:

CONTROL CARD-read by READALL

Col.	<u>Field</u>	Description
1	Integer	Card type - use 2
	DATA CARD-read by READ 2	
1-13	E13.6	WM(K) ^{>} 0
21-33	E13.6	WHR(K)>0 \cdot K=1, NG
41-53	E13.6	WW(K) > 0
69-70	Integer	IPW(K)>0

<u>USES</u>: No subroutines.

READ2: Reads in gas concentration error parameters.

FORTRAN SUBROUTINE	
CDC 3800	
(Internal to EPAGAS)	
READ3 (ZEP, KX)	

READ3 Margot Ackley

June 30, 1973

READ3 reads in the necessary controls needed for subroutine MINMYZD which are ZEP and the first six elements of the array KX.

CARD FORMATS:

CONTROL CARD-read by READALL

Col.	Field	Description
1	Integer	Card type - use 3
	DATA CA	RD-read by READ3
1-10	Floating	ZEP (0 $<$ ZEP $<$ 1), a number which tells how much of the predicted final step to take.
11-20	Integer	KX(1), the maximum number of new sets of errors (cycles) to use. Must be a whole number. If set to zero, then one cycle is used. If negative, then each increment value used in MINMYZD is decreased by 10 percent after each complete use of MINMYZD, and KX(1) is used for the number of cycles.
21-30	Integer	$KX(2)$, the number of times to count a bad cycle, $KX(2)^{>}$ 1.

Col.	<u>Field</u>	Description
31-40	Integer	KX(3), the percent improvement in
		tenths in the rms error to be
		accepted as a good cycle. Any cycle
	•	resulting in less than this causes
		KX(2) cycles to be counted rather
		than one cycle, $KX(3)^{>}0$.
41-50	Integer	KX(4), the limit as to the multiple
		of a WI increment which may be taken
		in one full step (ZEP=1). If set to
		zero, then 100 is used. If KX(4) is
		negative, then no step will be
		permitted in the -WI direction for the
		first complete try of MINMYZD.
		Future tries of MINMYZD, use KX(4).
51-60	Integer	<pre>KX(5), minimum ZEP in 0.001's,</pre>
		KX(5)>0.
61-70	Integer	KX(6), total percent improvement
		in tenths, KX(6)>0.

USES: No subroutines.

READ3: Reads controls for MINMYZD.

FORTRAN SUBROUTINE
CDC 3800
(Internal to EPAGAS)
READ4 (Q, GLB, CFM, MK, IPRINT, IPLOT)

READ4
Margot Ackley
July 30, 1973

READ4 reads in the necessary program controls for EPAGAS: Q is the number of percent by which each complete try of MINMYZD must improve the mean square deviation when using CALC, GLB is the minimum rms error permitted before using MINMYZD again, MK is the maximum number of uses of MINMYZD permitted, and CFM is a constant confidence factor to be applied to the given experimental transmission data if individual ones are not provided.

IPRINT=n permits printing of intermediate results using PRINT2 after every nth use of MINMYZD, and IPLOT=m permits a graph of the given and computed transmissions as functions of wavenumber to be plotted after every mth use of MINMYZD. If either IPRINT and/or IPLOT is zero, then no intermediate printing and/or plotting will occur.

CARD FORMATS:

CONTROL CARD-read by READALL

Col.	Field	Description
1	Integer	Card type - use 4
	DATA CARD	-read by READ4
1-10	Floating	Q> 0
11-20	Floating	GLB ² 0
21-30	Floating	CFM>0
31-40	Integer	MK>0
41-50	Integer	IPRINT [≥] 0
51-60	Integer	IPLOT ² 0

READ4 - (cont.)

USES: No subroutines.

READ4: Read program controls for EPAGAS.

FORTRAN FUNCTION	READ6
CDC 3800	Margot Ackley
(Internal to EPAGAS)	July 30, 1973
READ6 (WO, WIO)	

Given NG, the number of gas constituents, READ6 reads in NG original gas concentrations WO(K) and their corresponding increments WIO(K), K=1, NG. If the first data card read in contains a 2 in column 80 instead of WO(1) and WIO(1), then the original start concentrations and increments will be set to the final concentrations and increments of the previous data run and READ6 is set to -1. If not, then READ6 reads in the rest of WO and WIO values and returns equal to zero.

READ6 utilizes one COMMON block where: COMMON/101/NG, NT, RMSTP, RMST, RMSW.

CARD FORMATS:

CONTROL CARD-read by READALL

Col.	<u>Field</u>	Description	<u>n</u>
1	Integer	Card type - use 6	
	DATA CA	RD-read by READ6	
1-13	E13.6	WO(K)	
21-33	E13.6	WIO(K) K=1, NG	

USES: No subroutines.

READ6: Reads in starting gas concentrations and corresponding increments.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)
READ7 (UMIN, UMAX, CFM)

READ7
Margot Ackley
July 30, 1973

Given UMIN, UMAX, and CFM, where UMIN and UMAX are, respectively, the minimum and maximum wavenumbers for the corresponding given transmissions and CFM is a constant confidence factor to be used for each individual transmission value if separate factors are not provided; READ7 reads in individual transmissions TG(K) and corresponding confidence factors CF(K). The transmissions are assumed to be equally spaced in wavenumber starting at UB, stepping in UI increments, and ending at UE. NT number of pairs of TG(K) and CF(K) are read in where NT=1 + (UE-UB)/UT. The first data card (a.) read in gives the values UB, UE, and UI. The following data cards (b.) contain TG and CF.

If any of the following conditions occur: UB<UMIN, UE>UMAX, UI-0, NT>150, or TG(K)<0, TG(K)>1, CF(K)<0, K=1, NT, then an error has occurred and READ7 returns equal to -1. Otherwise, if no error occurs, then READ7=0 upon return. If CF(K)=0, then CF(K) is set to CFM for all appropriate K=1, NT.

Before return, the array U(K), K=1, NT is filled where U(K)=UB+(K-1)*UI.

READ7 utilizes two blocks of COMMON: COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

CARD FORMATS:

CONTROL CARD-read by READALL

Col.	Field	Description
1	Integer	Card type - use 7
		DATA CARD (a.) - read by READ7
1-10 11-20 21-30	Floating Floating Floating	UB UB-UMIN UE UE-UMAX UI UI>O
		DATA CARD (b.) - read by READ7
1-5 6-10 11-15 16-20	F5.4 F5.5 F5.4 F5.5	TG(j) CF(j) TG(j+1) CF(j+1) j=1, 9, 17, <nt< td=""></nt<>
21-25 26-30 31-35 36-40	F5.4 F5.5 F5.4 F5.5	TG(j+2) CF(j+2) TG(j+3) CF(j+3)
•		· · · · · · · · · · · · · · · · · · ·
71-75	F5.4	TG(j+7)
76-80	F5.5	CF(j+7)

<u>USES</u>: No subroutines.

<u>READ7</u>: Reads in given transmissions and corresponding confidence factors for EPAGAS.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)
READ8 (UMIN, UMAX, CFM)

READ8
Margot Ackley
July 30, 1973

Given UMIN, UMAX, and CFM, where UNIM and UMAX are, respectively, the minimum and maximum wavenumbers for the corresponding given transmissions and CFM is a constant confidence factor to be used for each individual transmission value if separate factors are not provided, READ8 reads in the individual wavenumber values U(K), the corresponding given transmission TG(K), and its confidence factor CF(K), K=1, NT. The first data card (a.) read in gives the value of NT, the number of given transmissions to be read. The following cards (b.) contain TG, U, and CF.

If any of the following conditions U(K)<UMIN, U(K)>UMAX, TG(K)<-0.2, or TG(K)>1.01, or CF(K)<0, K=1, NT, or NT>150 occur, then an error has been encountered and READ8 returns equal to -1. Otherwise, no error occurred and READ returns equal to 0. If CF(K)=0, then CF(K) is set to CFM for all appropriate K=1, NT.

READ8 utilizes two blocks of COMMON: COMMON/101/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

CARD FORMATS:

CONTROL CARD-read by READALL

Col.	<u>Field</u>	Description
1	Integer	Card type - use 8

READ8 - (cont.)

	<u>D.</u>	ATA CARD	(a.) - read by	READ8
Col.	Field		Descri	ption
1-3	Integer	NT	0 <nt<sup><150</nt<sup>	
	<u>D.</u>	ATA CARD	(b.) - read by	READ8
1-10	Floating	TG(K)		
11-20	Floating	U(K)	K=1, NT	
21-30	Floating	CF(K)		

USES: No subroutines.

READ8: Reads in given transmission and corresponding wavenumbers and confidence factors for EPAGAS.

FORTRAN FUNCTION

CDC 3800

(Internal to EPAGAS)

RMSERR (A, N)

RMSERR

J. R. Winkelman

June 15, 1965

Given an array A of length N, RMSERR is the root mean square of the array A where

RMSERR =
$$\left(\frac{1}{N} \sum_{i=1}^{N} A_i^2\right)^{\frac{1}{2}}$$

USES: No subroutines.

RMSERR: Root mean square of an array.

FORTRAN FUNCTION
CDC 3800
(Internal to EPAGAS)
THEOTRN (IDGAS)

THEOTRN

Margot Ackley

July 30, 1973

THEOTRN reads in the gas library of theoretical transmissions for a given number of specified gases as a function of both gas concentration and wavenumber. If any errors occur in reading or certain variables are outside permitted ranges, THEOTRN returns immediately equal to one. If all reading is correct, then THEOTRN returns equal to zero.

Before the actual gas library is read in, THEOTRN first reads in a set of data cards that specify which gases are in the library, which concentrations for each gas appear, and which wavenumber region is used for each gas.

Variables appearing on the data cards with their corresponding dimensions and restrictions, if applicable, are given below:

NG: Total number of different gases to be found in

the library, 0 < NG-10.

IDGAS: Array of maximum length 10 which contains a

numerical (I2) identifier for each of the gases, where IDGAS(I) = identifier for the Ith gas, I=1,

NG.

NU: Array of maximum length 10 which contains the

number of wavenumber values used for each gas,

where NU(I) = number of wavenumber values for the

Ith gas, I=1, NG.

NW: Array of maximum length 10 which contains the number

of concentrations used for each gas, where NW(I) = number of concentrations for the Ith gas, I=1, NG;

 $0 < NW(I)^{<} 5$.

THEOTRN - (cont.)

UR:

A two-dimensional array UR(I,J) dimensioned (10,3). UR contains the wavenumber range and spacing used for each of the gases. I corresponds to the Ith gas. J=1 corresponds to the beginning wavenumber, J=2 corresponds to the end wavenumber, and J=3 corresponds to the wavenumber increment. Note that 1. + (UR(I,2) - UR(I,1))/UR(I,3) should be equal to NU(I).

WR:

A two-dimensional array WR(I,J) dimensioned (10,5). WR contains the actual concentrations used for each gas. I corresponds to the Ith gas. J corresponds to the Jth concentration for the particular Ith gas and must be given in strictly increasing order where 1.E-20 < WR(I,1) < WR(I,2) < ... < WR(I,J) < ... < WR(I,NW(I)).

Note that the number of wavenumber values and the wavenumber range and spacing are the same for all concentrations for a given gas.

DATA CARD FORMATS:

	ar	_	
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Col.	<u>Field</u>	Description	
1-4	Integer	NG	
Card 2.			
2-4	Integer	IDGAS(1)	
6-8	Integer	IDGAS (2)	
•	•	•	
•	•	•	
•	•	•	

<u>Col.</u>	<u>Field</u>	Description	
•	•	IDGAS (NG)	
Card 3.			
1-4	Integer	NU(1)	
5-8	Integer	NU(2)	
•	•	•	
•	•	•	
•	•	•	
•	•	NU (NG)	
Card 4.			
1-4	Integer	NW(1)	
5-8	Integer	NW(2)	
•	•	•	
•	•	•	
•	•	•	
•	•	NW (NG)	
Card 5.	 		—
1-8	Floating	UR(1,1)	
9-16	Floating	UR(1,2)	
17-24	Floating	UR(1,3)	
25-32	Floating	UR(2,1)	
33-40	Floating	UR(2,2)	
41-48	Floating	UR(2,3)	
49-56	Floating	UR(3,1)	
57-64	Floating	UR(3,2)	
65-72	Floating	UR(3,3)	
73-80		B1ank	
Use s	uccessive cards up	to a maximum of four.	

Card 5. +J, 1 - J - 13

Col.	<u>Field</u>	Description	
1-10	E10.3	WR(I,1)	
11-20	E10.3	WR(I,2) I=1, NG	
•	•	•	
•	•	•	
•	•	•	
•	•	WR(I,NW(I))	

Use successive NG cards, one for each gas.

THEOTRN next reads in the actual gas library.

LIBRARY CARD FORMAT:

Col.	<u>Field</u>	Description
1-2	Integer	NAME
3-4	Integer	ICON
5-8	Integer	NN
9-12	F4.3	T(J)
13-16	F4.3	T(J+1)
17-20	F4.3	T(J+2)
21-24	F4.3	T(J+3)
•	•	
77-80	F4.3	T(J+17)

The gas with identification IDGAS(1) must come first, then the gas with IDGAS(2) is next, etc. Gas with identification IDGAS(NG) comes last. NAME on each gas library card corresponds to IDGAS of the particular gas being read in. Within a particular gas, the concentrations W must be increasing. The first concentration corresponds to ICON=1, second concentration corresponds to ICON=2, and last concentration corresponds to

THEOTRN - (cont.)

ICON=NW(I) for the Ith gas. Within a given gas and a given concentration, the cards are serialized by NN where NN=1 for the first card, NN=2 for the second card, etc. The maximum value for NN within a set is NU(I)/18 for the Ith gas. T(J) corresponds to the theoretical transmission for the particular Ith gas and its particular concentration. T(1) is the transmission corresponding to first wavenumber UR(I,1), T(2) is the transmission corresponding to the wavenumber UR(K,1) + UR(I,3), and the last transmission, T(NU), corresponds to the last wavenumber UR(I,2) for the Ith gas. Within a given gas and particular concentration, all cards have 18 values of transmission except for the last card which may have less.

THEOTRN does extensive checking of the card order when reading in the library. If the cards are out of order, an error has occurred and THEOTRN terminates equal to one.

As the individual transmission T is read in, it is stored in the large transmission array TRAN (I,J) which has been dimensioned (5000,5). J corresponds to the concentration number ICON, where 1 - J - 5 for all gases. I=1 corresponds to the transmission of the first gas, first wavenumber. I=NU(1) corresponds to the transmission of the first gas, last wavenumber. I=NU(1) + 1 corresponds to the transmission of the second gas, first wavenumber. I = NU(1) + NU(2) corresponds to the transmission of the second gas, last wavenumber. I = $\sum_{i=1}^{NG} NU(i)$ last wavenumber. Unless all gases have the same number of concentrations, the actual amount of storage utilized will not be rectangular.

THEOTRN utilizes two COMMON blocks COMMON/100/ and COMMON/101/ where COMMON/100/TRAN(5000,5), NU(10), NW(10), UR(10,3), WR(10,5), TG(150), U(150), CF(150), TC(150), WM(10), WHR(10), WW(10), IPW(10); and COMMON/101/NG, NT, RMSTP, RMST, RMSW.

USES: No subroutines.

THEOTRN: Reads in gas library of theoretical transmissions for EPAGAS.

A2.2 SIMULATION OF GAUSSIAN NOISE

To study the effects of noise in the transmission data, a method of obtaining random Gaussian noise was developed. Many large-scale computers (CDC 3800 in particular) provide a library routine which furnishes the user with a sequence of uniformly distributed random numbers ranging from 0 to 1. Given two such numbers A_i and B_i , a Gaussian random variable X_i can be obtained from a transformation provided by Box and Muller (1958), where

$$X_i = (-2 \log_e A_i)^{\frac{1}{2}} \cos (2\Pi B_i).$$

Three different signal-to-noise ratios 10, 30, and 100 were incorporated in this report. Letting SNR be the signal-to-noise ratio desired, then SNR is defined as

$$SNR = \frac{\sigma_d}{\sigma_n} ,$$

where σ_d is the standard deviation of the data and σ_n is the standard deviation of the Gaussian noise for the set of data to be analyzed. The noise N_i used to modify a particular transmission value T_i is defined as

$$N_i = \sigma_n X_i$$
,

where the noise over the set has mean zero and standard deviation σ_n .

The new transmission values ${\rm TN}_{\dot{1}}$ incorporating the noise are then defined as

$$TN_i = T_i + N_i$$
.

A2.3 CONSTITUENT AND TRANSMISSION ERROR FORMULATION

To direct its progress, the minimization process uses a total rms error E as a measure of the "goodness of fit." By varying the amounts of each of the gas constituents, the process tries to minimize the total rms error E which is both a function of the rms error E_T from the individual transmission errors and the rms error E_W from the errors of the individual constituent amounts. (See EROR2 and EROR1 descriptions, respectively.)

For the benefit of the user, all three rms errors E, E_T , and E_W are standard printed variables. In this program, the total rms error E and the constituent rms error E_W are calculated. From these two, the transmission rms error E_T can then be calculated as follows.

Let NG = the number of gas constituents,

NT = the number of transmission values,

A = the individual errors from the gas constituents, i=1, NG,

 A_i = the individual errors from the transmissions, i=NG+1, NG+NT,

N = NG + NT.

Then,

$$E = \left[\frac{1}{N} * \sum_{i=1}^{N} A_i^2\right]$$

$$E_{W} = \left[\frac{1}{NG} * \sum_{i=1}^{NG} A_{i}^{2}\right]^{\frac{1}{2}}$$

But

$$E^2 = \frac{1}{N} * \sum_{i=1}^{NG} A_i^2 + \frac{1}{N} * \sum_{i=NG+1}^{NG+NT=N} A_i^2$$

$$E^2 = \frac{NG}{N} * \frac{1}{NG} * \sum_{i=1}^{NG} A_i^2 + \frac{NT}{N} * \frac{1}{NT} * \sum_{i=NG+1}^{N} A_i^2$$

and

$$E_{\mathbf{T}}^2 = \frac{1}{NT} * \sum_{i=NG+1}^{N} A_i^2$$

Therefore,

$$E^2 = \frac{NG}{N} * E_W^2 + \frac{NT}{N} * E_T^2$$

$$E_{T} = \left[\frac{N * E^{2} - NG * E_{W}^{2}}{NT}\right]^{\frac{1}{2}}$$

A2.4 INTERPOLATION FORMULAS USED IN TRANSMISSION LIBRARY

Given a wavenumber ν and a gas concentration W for a specified constituent, a corresponding transmission $T(\nu,W)$ can be found from the transmission library. Because the library contains transmissions at discrete values of wavenumber and concentration, a form of double linear interpolation must be utilized.

Assuming the ν and W lie within the region of the transmission table, that is,

$$W_{\min} \stackrel{<}{-} W_1 \stackrel{<}{-} W \stackrel{<}{-} W_2 \stackrel{<}{-} W_{\max}$$

$$v_{\min} < v_1 < v_2 < v_2 < v_{\max},$$

then by interpolating first in the W direction, we have,

$$\frac{T(v_1, W) - T(v_1, W_1)}{W - W_1} = \frac{T(v_1, W_2) - T(v_1, W_1)}{W_2 - W_1}$$
(A2:1)

and

$$\frac{T(v_2, W) - T(v_2, W_1)}{W - W_1} = \frac{T(v_2, W_2) - T(v_2, W_1)}{W_2 - W_1}$$

Therefore,

$$T(v_1, W) = \frac{W - W_1}{W_2 - W_1} * \left[T(v_1, W_2) - T(v_1, W_1) \right] + T(v_1, W_1),$$

$$T(v_2, W) = \frac{W - W_1}{W_2 - W_2} * \left[T(v_2, W_2) - T(v_2, W_1) \right] + T(v_2, W_1). \quad (A2:2)$$

and

Interpolating next in the v direction, we have

$$\frac{T(v,W) - T(v_1,W)}{v - v_1} = \frac{T(v_2,W) - T(v_1,W)}{v_2 - v_1}$$
(A2:3)

Therefore,

$$T(v,W) = \frac{v - v_1}{v_2 - v_1} * \left[T(v_2,W) - T(v_1,W) \right] + T(v_1,W). \tag{A2:4}$$

By substituting the expressions for $T(v_1,W)$ and $T(v_2,W)$ from (A2:2) into equation (A2:4), we have the following interpolation formula

$$T(v,W) = \frac{v - v_1}{v_2 - v_1} * \left[\frac{W - W_1}{W_2 - W_1} * \left(T(v_2, W_2) - T(v_2, W_1) + T(v_1, W_1) \right) \right] - T(v_1, W_2) + T(v_2, W_1) - T(v_1, W_1) + T(v_1, W_2) - T(v_1, W_2) - T(v_1, W_2) - T(v_1, W_2) + T(v_1, W_2) - T(v_1, W_2) - T(v_1, W_2) + T(v_1, W_2) - T$$

Because of the structure of the program, the wavenumber v will always lie within the region of the transmission table. However, the concentration W for a particular gas constituent could possibly lie outside the region due to the minimization process which varies only the W's. In the event W is outside the region, we have two possible cases:

$$(1) \qquad \mathbf{W}_{\max} < \mathbf{W}$$

For the first case, let

$$W_{\text{max}-1} = W_1 < W_{\text{max}} = W_2 < W.$$

Then,

$$\frac{T(v_{1},W) - T(v_{1},W_{1})}{W-W_{1}} = \frac{T(v_{1},W_{2}) - T(v_{1},W_{1})}{W_{2}-W_{1}}$$

$$\frac{T(v_{2},W) - T(v_{2},W_{1})}{W-W_{1}} = \frac{T(v_{2},W_{2}) - T(v_{2},W_{1})}{W_{2}-W_{1}},$$
(A2:6)

and

$$\frac{T(v,W) - T(v_1,W)}{v - v_1} = \frac{T(v_2,W) - T(v_1,W)}{v_2 - v_1}.$$
 (A2:7)

By referring to (A2:1) and (A2:3), we can then use (A2:5) for the final formula with $W_1 = W_{max-1}$ and $W_2 = W_{max}$. For the second case, where

$$W_1 = 0 - W < W_{min} = W_2$$

or

$$W_1 = 0 < W_{\min} = W_2$$
,

we may utilize the fact that

$$T(v,0) = T(v,W_1) \equiv 1 \text{ for all } v.$$
 (A2:8)

Applying (A2:8) to (A2:5), we have

$$T(v, w) = \frac{v - v_1}{v_2 - v_1} * \left[\frac{w - 0}{w_2 - 0} * \left(T(v_2, w_2) - 1 + 1 - T(v_1, w_2) \right) + 1 - 1 \right] + \frac{w - 0}{w_2 - 0} * \left[T(v_1, w_2) - 1 \right] + 1.$$
(A2:9)

$$T(v, W) = 1 + \frac{W}{W_2} * \left[\frac{v - v_1}{v_2 - v_1} * \left(T(v_2, W_2) - T(v_1, W_2) \right) + T(v_1, W_2) - 1 \right]$$
(A2:10)

where

$$W_2 = W_{\min}$$

In summary, given v and W to find T(v,W), where $v_{min} = v_1 = v_2 = v_{max}$: For $w_{min} = w_1 = w_2 = w_{max}$,

$$T(v,W) = \frac{v - v_1}{v_2 - v_1} * \left[\frac{W - W_1}{W_2 - W_1} * \left(T(v_2, W_2) - T(v_2, W_1) + T(v_1, W_1) - T(v_1, W_2) \right) + T(v_2, W_1) - T(v_1, W_1) \right] + \frac{W - W_1}{W_2 - W_1} * \left[T(v_1, W_2) - T(v_1, W_1) \right] + T(v_1, W_1).$$

For $W_{max} < W$,

$$T(v,W) = \frac{v-v_1}{v_2-v_1} * \left[\frac{W-W_{max}-1}{W_{max}-W_{max}-1} * \left(T(v_2,W_{max}) - T(v_2-W_{max-1}) + T(v_1,W_{max-1}) - T(v_1,W_{max}) \right) + T(v_2,W_{max-1}) - T(v_1,W_{max-1}) \right] + \frac{W-W_{max}-1}{W_{max}-W_{max-1}} * \left[T(v_1,W_{max}) - T(v_1,W_{max-1}) \right] + T(v_1,W_{max-1}) .$$

For W < W_{min},

$$T(v,W) = 1 + \frac{W}{W_{\min}} * \left[\frac{v - v_1}{v_2 - v_1} * \left(T(v_2, W_{\min}) - T(v_1, W_{\min}) \right) + T(v_1, W_{\min}) - 1 \right]$$

A2.5 PROGRAM EXECUTION TECHNIQUES

Although program EPAGAS is a totally self-contained program, its proper utilization requires art as well as science if successful results are desired. Before attempting a run, the user should thoroughly acquaint himself with all the program and subprogram descriptions. The following categories require careful checking before the program is run and are a trouble-shooting checklist in case of improper results.

A2.5.1 Theoretical Gas Library Spectra

The user cannot hope to have good results if the library of spectra does not contain all of the gas constituents which are (or are suspected) to occur in the experimental data. Also the wavenumber range for the gas constituents may not be sufficient for the range of the experimental data. If this occurs, an error message will be printed. The immediate solution to this problem is either to increase the wavenumber range of the library spectra or to analyze a smaller region of the experimental data.

The range of the concentrations of a particular gas constituent is another item that should be checked. The library should have a range of concentrations that covers the expected concentration of a particular gas constituent. If, during the process of trying to find a solution, EPAGAS tries to use a concentration outside the region of given concentrations, a method of linear extrapolation is used to derive a particular gas concentration. For concentrations less than 10% of the total range of the region lying beyond the outer limits of the library, there generally will not be problems. However, concentrations greater than 10% could cause the program to give erroneous results.

The spacing of the library spectra in both wavenumber and concentration is also very important. The program uses linear interpolation in both wavenumber and concentration to determine transmissions for a given wavenumber and a trial concentration. (See FINDTRN decription.) If the grid spacing is not fine enough to give reasonably correct results with linear interpolation, the program may not be able to give satisfactory results. The fineness of the library is

often limited by available computer memory and by the total number of gas constituents present. In the case where the user has experimental data containing many gas constituents, it might be necessary to limit the wavenumber region to a fairly narrow band or to pick regions where only a few gas constituents are prominent.

The program has extensive built-in checks to make sure that the library spectra and associated wavenumber and concentration information are properly read in. If any of the library cards are out of order or if inconsistencies occur, an error message is printed and further execution is terminated. In the event this happens, carefully check the THEOTRN description for proper input of the library spectra.

A2.5.2 Experimental Transmission Data

Unsuccessful results can often be caused by experimental data which have an intolerable noise level. Signal-to-noise ratio, S/N, should be on the order of 100 or better. For S/N<100, the user should select wavenumber regions that have only a single gas constituent or regions which have little interference from other gases.

Before utilizing EPAGAS, the user may find that it is necessary to do some "preprocessing" of the data. Due to a variety of reasons, instrumental drift in particular, the data may have to be corrected either linearly or nonlinearly for both wavenumber and/or transmission. Absolute transmission values of I/I_O without noise should range from 0 to 1. Past experience in the use of program EPAGAS has shown that wavenumber drift is particularly detrimental to obtaining successful results. Wavenumber and intensity calibrations can be obtained from careful comparison with known spectra.

Another very important item to check is the instrumental resolution. Resolution should be the same or very nearly the same as the resolution used in producing the library spectra. It is recommended that instrumental resolution should deviate from that of the library by no more than 10 percent. Also the resolution should not fluctuate as a function of wavenumber which can be a major problem for some types of spectrometers unless it is corrected.

The temperature and pressure of the experimental data should also correspond as close as possible to those used in producing the library spectra.

In order that the experimental data is well defined, spacing in wavenumber should be sufficiently fine. If the spacing is too coarse, important spectral line features may be lost. If this is the case, a suitable solution would almost be impossible to find. Wavenumber spacings of at least 1/5 of the resolution are strongly recommended.

Two other important aspects of the experimental data should also be taken into consideration. The slit function of the instrument should closely approximate that of the library. The digital resolution in the conversion of the analog I/I information to digital form has a definite effect on the final accuracy of determinations of gas concentrations. The higher the digital resolution, the more accurately EPAGAS can determine the concentrations.

A2.5.3 Minimization or "Goodness of Fit" Controls

Often less than desirable results are obtained because EPAGAS has not been permitted to "work hard enough" at finding a satisfactory solution. A thorough understanding of the descriptions of MINMYZD, READ3, READ4, and CXMIZ would be extremely helpful (along with some experience in using EPAGAS) in remedying this problem. MINMYZD description gives the main minimization procedures and presents a discussion of the convergence controls that direct its progress toward a solution. READ3 description provides information on the controls, used by MINMYZD, which are labelled ZEP and array KX. Particular attention should be paid to the first six elements of the array KX. The description of READ4 gives a discussion of the variables Q, GLB, and MK which are used in conjunction with the 9th and 10th elements of the array KX by CXMIZ to determine if another try by MINMYZD to find a solution should be made. It is recommended that when first using the program on a new set of experimental data, the controls should be set as loosely as computer costs will permit.

A2.5.4 Error Parameters

Proper formulation of the errors defined on the individual gas concentrations is most helpful for successful utilization of EPAGAS. The user though must keep in mind that the term "error" is used rather loosely in connection with the gas concentrations. These errors are mainly a way in which to guarantee that the minimization process does not try to select individual gas concentrations which are known to be physically unsuitable. The errors (or better, restraints) on the individual gas concentrations are of the general form,

$$ERR(I) = WW(I) * \left(\frac{W(I) - WM(I)}{WHR(I)}\right)^{IPW(I)},$$

where I = ith gas constituent. W is the current trial concentration being used by the minimization process, and all the other variables are input error parameters. (See EROR1 description.) WM is considered to be the mean acceptable concentration. For most cases (unless additional knowledge exists of the amount of the gas constituent present), WM is simply the mean concentration value based on the highest and lowest library concentrations of the particular gas constituent. WHR is the half range of permitted variation of the particular concentration and is simply the difference between the highest library concentration and the mean concentration. As long as W remains within the concentration range of the library, the absolute value of in a fairly small error. If the $\left|\frac{W-WM}{WHR}\right|$ will be between 0 and 1, resulting minimization process tries to select W's outside of the library region, the error will become rapidly larger as W gets farther and farther away from the acceptable region. The variable IPW serves the purpose of determining just how fast the errors will grow as W moves farther and farther away. IPW should always be an odd, positive number to preserve the "direction" of the error. The larger IPW is, the faster the error will increase once W has entered the unacceptable regions. From past experience, IPW=3 usually is a good starting value. WW is a weighting or normalization factor and is used to keep individual

gas constituent errors in line with each other and also in proportion with the individual transmission errors. Sometimes several runs have to be made before a proper determination of the WW's can be made; however, unity is always good for an initial try.

It is generally best to keep the gas constituent error parameters fairly loose; however, if prior knowledge of the concentration of a particular gas constituent in the experimental data exists, then one may wish to narrow the acceptable region around the suspected concentration. This would be done by setting WM equal to the suspected concentration, by considerably decreasing WHR, by increasing IPW up to 9 or 11, and by setting WW to 10 or 20 times the values of WW used for the other gas constituents.

The errors, depending on the computed and given transmission values, are of the general form,

$$ERR(I) = CF(I)*(TG(I)-TC(I)),$$

where I = ith value corresponding to the ith wavenumber. TG is the given experimental transmission, TC is the currently computed transmission based on the current gas concentrations as determined from the minimization process, and CF is a confidence or weighting factor. The role of CF is twofold. If some regions of the experimental data are more reliable than others, then the corresponding CF's should be larger than those applied to transmissions in the more uncertain regions. This has an effect on the minimization process so as to let it somewhat "ignore" the less reliable data and not to spend its efforts seeking some erroneous solution to very noisy or bad data. Also, the CF's play a similar role for the transmission errors that the WW's do for the gas concentration errors. should be taken to select the CF's so that the transmission errors are proportional to the concentration error. The user must always keep in mind that the minimization process does not involve the individual gas concentration or transmission errors, but rather tries to minimize the composite rms error of all the errors. Hence, it is vitally important

that the effects of the gas concentration errors and the transmission errors on the rms error are properly balanced.

A2.5.5 Miscellaneous Execution Techniques

To commence its minimization process, program EPAGAS needs to be furnished with a set of beginning gas constituent concentrations. If no prior knowledge of the concentrations exists, either the mean or the center concentration is always a good choice. As mentioned in the previous section, if the concentration of a particular gas constituent is approximately known, then this value should be the one selected for the beginning gas concentration. Also, in this case, the gas error parameters should be tightened for this particular gas constituent. From prior experience, improper selection of beginning trial gas concentrations has generally had no disastrous effects on the final solution except to increase computation time. In most cases, a unique solution is found regardless of the starting point. However, in some rare cases, a change in the starting positions or a change in the order of the gas constituents can sometimes be helpful in seeking a suitable solution. The size of the initial increments by which the individual concentrations are varied by the minimization process must also be carefully selected. increments are too large, the first change in a particular gas concentration may completely "jump" over the actual true concentration. On the other hand, if the increments are too small, a proper solution may not be reached because applying the increment to the original gas concentration results in an insignificant change. When in doubt, use an increment on the larger size because the minimization process will decrease the size of the increment, but will never increase it.

Program EPAGAS is equipped to furnish intermediate printing of current gas concentrations, of currently computed transmissions, and of corresponding errors, along with a graph showing the given and currently computed transmissions as a function of wavenumber. Careful studying of these intermediate results can often furnish the user with a better feeling of how certain changes in the individual gas concentrations affect the overall rms error.

The intermediate printing also provides the user with the current values of selected elements of the KX array, KX(7) through KX(10). These values also can give the user an insight into the success of each cycle or try of the minimization process. (See MINMYZD description.)

Last, but not least, there are some sets of experimental data which do not lend themselves properly to the minimization technique of determining atmospheric gas constituents and their corresponding concentrations from their superimposed infrared spectra. This may occur whenever the data is too noisy or too imprecise, or whenever the scales (I/I_0 or v) must be corrected nonlinearly. Without good data, good results cannot be achieved, but if the nonlinear corrections are known, they may be applied in the computer program.

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APPENDIX 3

A3. CHARACTERISTICS OF THE ROSE LONG-PATH SPECTROPHOTOMETER SYSTEM

Design range (transmission) 0.4 to 4.0 km

(0.25 to 2.5 mi.)

Spectral regions 3 to 5.5 μ (1820 to 3330 cm⁻¹ and

7 to 13.5 μ (740 to 1430 cm⁻¹)

Resolution to 0.01 u

Scan time 2-min. minimum (either spectral region),

130-min. maximum

(either spectral region)

Display wavenumber, I, I/I (see section 1.1)

Source and Reference Blackbodies

Operating temperature 1800°K
Temperature stability +4°C

Emissivity 0.99 - 0.01

Aperture $12.7 \times 6.3 \text{ mm} (0.50 \times 0.25 \text{ in.})$

Telescopes

Focal length 304.8 cm (120 in.)

Clear aperture 61.0 cm (24 in.)

Field of view 0.25 deg. (-0.125 deg.)

Blur circle diameter $150~\mu$

Primary f/no 2.0

Secondary magnification 2.5

Source Modulator

Operating frequency 570 Hz

Reference Chopper

Frequency 330 Hz
Stability +0.3 Hz

Monochromator

Mfgr. and model Perkin Elmer (Norwalk, CT)

Model 210B

Type Littrow grating with linear wavenumber

drive

Gratings 3 to 5.5 μ ; 240 1/mm 20 3400 cm⁻¹

 (2.94μ)

7 to 13.5 μ ; 101 1/mm 22°2' 1333 cm⁻¹

 (7.5μ)

masked to 15.0 cm² net area (nominal)

Slits width, 0 to 2.0 mm micrometer-

controlled; height, 0 to 12.0 mm

micrometer-controlled

Wavenumber drive 3/32 to 6 rpm in steps of 2x

Min. scan time approximately 2 minutes

Detector

Mfgr. and model Santa Barbara Research Center

(Goleta, CA) 8679^{-1}

Type Mercury-doped Germanium

Operating temp. 26° K (58 psia H₂)

Size $0.2 \times 2.0 \text{ mm} (0.008 \times 0.08 \text{ in.})$

Field of view 90

D* (λm , 630) 0.84 x 10^{10} m Hz¹2/W

Detector Cooler

Mfgr. and model Cryogenic Technology Inc.

(Waltham, MA) Model 20 with

3833-005 temperature controller

Type closed-cycle He refrigerator,

consisting of compressor and cold

head connected by flexible hoses

Operating temp. range approximately 10° to 28°K

Operating temperature 26°K (58 psia H₂)

Cooldown time approximately 15 min.

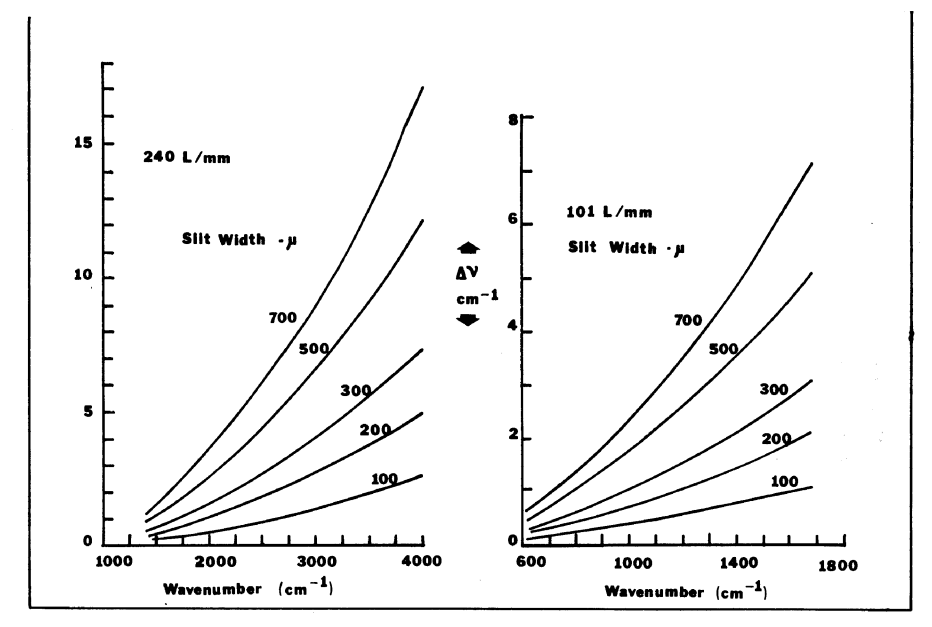


Figure A3.1
RESOLUTION VS. WAVENUMBER FOR VARIOUS SLIT WIDTH SETTINGS
FOR ROSE SPECTROPHOTOMETER

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APPENDIX 4

A4. CONVENIENT CONVERSION FACTORS

 Θ = temperature deg. K,

M = molecular weight

A = Avogadro's number = 6.0225×10^{23}

S = Line intensity

K = Absorption coefficient

U = Optical path

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16. ABSTRACT

Atmospheric gaseous pollutants are very numerous in industrial regions. It is estimated that 25 or more pollutant molecules may be found in the atmosphere in significant quantities. The measurement of the concentration of each gas from the complex spectrum obtained by a long-path infrared spectrophotometer requires the fitting of trial spectra composed from a library of spectra. The fitting procedure adjusts the concentrations of the trial spectra until a "best fit" in a least-squares sense is produced. This report is a description of the physical, mathematical, and calculational principles and procedures for the use of a digital computer program to determine concentrations of atmospheric gases in a path of a few kilometers. Detailed instructions for the computer program and a library of spectra are provided.

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