FT-IR OPEN-PATH MONITORING GUIDANCE DOCUMENT
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by
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Notice

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Preface

A need exists for a document describing remote sensing over an open path with a Fourier transform infrared (FT-IR) spectrometer. This monitoring technique is developing rapidly, and some guidance should be given to assist the newer users. There is also a need to coordinate other developmental efforts and to attempt to standardize certain perfunctory tasks. Several attempts have been made to provide such a document, but they all seem to lack the guidance that users of this technique require. For example, none answer the question of how one actually obtains background data. Also, none present a clear description of how the system functions.

There is some difficulty in producing a document that answers all of the questions most people have about the field use of the long-path, open-path FT-IR monitoring technique. The reason is that many of the questions do not have ready answers. To be sure, the Michelson interferometer is well understood and characterized, but, for example, the final procedure for handling the effects of interferences caused by water vapor may take several years to fully understand and develop. Chapter 2 is primarily a description of the optics that make up an open-path FT-IR system. While writing the remaining chapters, both authors came to realize that the field investigator should have a thorough understanding of how the system functions. Therefore, we tried to cover the most important concepts of the system in some technical depth. In referring to long-path, open-path monitoring, we mean measurements taken along a total path that is at most 1 km long and is open to the atmosphere.

With these items in mind, the reader should not view this document as the final chapter concerning the use of the FT-IR technique. Rather, it should be considered as an interim product. As new developments occur, this document can be updated so that it will continue to be a useful guide. Because of this intended use, other experimenters in IR open-path monitoring are asked to contribute to the document as they make the significant progress that is sure to come.
Preface to the Second Edition

During the 18 or so months since the publication of the first edition of this document we have had many comments from the readers. We have also had the opportunity to acquire a large data set at the EPA facility in Research Triangle Park, North Carolina.

We have made several additions in the present edition that we believe will make the document more helpful to the users of long-path FT-IR monitoring techniques. We have added a new chapter that deals specifically with the single-beam spectrum and the measurements that are important for quality assurance purposes. There is also a new chapter that describes the production of a water vapor spectrum that can be used with the analysis software. The chapter on resolution has been expanded to incorporate our findings over the past year. The chapter on quality assurance has also been expanded in that it now contains a portion of the data that we have acquired. We hope the data will provide some guidance to the users of this technique. We have added some further definitions to the glossary of terms chapter and have updated the bibliography so that it now contains more than 250 entries.

Although we have tried to make this document fairly general and not instrument specific, there are instances where this has not been possible. Our experience is with an FT-IR system that operates at 1-cm⁻¹ resolution and uses a specific data analysis package. The data contained in this document and some of the statements reflect this fact.

The authors wish to emphasize that this document is meant to (1) serve as a primer to new users and others interested in learning about the fundamental aspects of long-path, open-path FT-IR monitoring and (2) give current users of FT-IR monitors guidance regarding general operating procedures. This document is not intended to be a standard operating procedure, nor is it meant to imply that the FT-IR technique is an EPA-approved method for ambient air analysis. The development of an EPA Toxic Organics (TO) method using open-path FT-IR systems is forthcoming and will be in a different format from this document. We anticipate that the guidance document will continue to be revised as new developments are made in the field of open-path FT-IR monitoring.

GMR
JWC
# Contents

Notice ................................................................. ii  
Preface ........................................................................ v  
Preface to Second Edition ........................................... vii  
Figures ........................................................................ ix  
Tables ........................................................................... xi  
Acknowledgement ........................................................... xii

1 Introduction ................................................................. 1-1  
   1.1 Overview of Document ........................................ 1-1  
   1.2 References .......................................................... 1-2

2 The Fourier Transform Spectrometer ................................. 2-1  
   2.1 Introduction and Overview ..................................... 2-1  
   2.2 The Michelson Interferometer ................................. 2-2  
      2.2.1 Interference ................................................ 2-3  
      2.2.2 Resolution ............................................... 2-6  
      2.2.3 Throughput .............................................. 2-10  
      2.2.4 The Detector ............................................. 2-10  
      2.2.5 The IR Source ........................................... 2-12  
   2.3 Transfer Optics, Telescopes, and Beam-Return Optics ... 2-12  
      2.3.1 Bistatic System ......................................... 2-15  
      2.3.2 Monostatic System ..................................... 2-16  
   2.4 The Electronics ................................................... 2-17  
   2.5 The Computer .................................................... 2-18  
   2.6 The Data Output .................................................. 2-19  
      2.6.1 Beer's Law ................................................ 2-19  
      2.6.2 The Interferogram .................................... 2-20  
         2.6.2.1 Truncation ........................................ 2-21  
         2.6.2.2 Phase Shift ....................................... 2-21  
      2.6.3 The Transform ........................................... 2-22  
      2.6.4 The Single-Beam Spectrum ......................... 2-22  
      2.6.5 Data Analysis ........................................... 2-23  
         2.6.5.1 Generation of the Absorption Spectrum ... 2-23  
         2.6.5.2 Generation of the Reference Spectrum ... 2-23
6.2.1 The Longest Path ........................................ 6-3
6.2.2 Shortest Path Requirements ............................ 6-4
6.2.3 Short Path Versus Long Path ......................... 6-4
6.2.4 Prevailing Winds .................................. 6-6
6.2.5 Slant Path Versus Horizontal Path ............... 6-7
6.3 Changing the Path .................................. 6-7
6.4 Ancillary Measurements ................................. 6-8
6.5 A Specific Case ........................................ 6-8
6.6 References .............................................. 6-11

7 Resolution Considerations in Long-Path, Open-Path FT-IR Spectrometry ........................................ 7-1
7.1 Introduction and Overview .............................. 7-1
7.2 Definition of Resolution ................................ 7-3
7.3 Trading Rules in FT-IR Spectrometry ............... 7-4
7.4 Example Spectra of CO₂ and Water Vapor .......... 7-6
  7.4.1 Resolution Effects ................................ 7-6
    7.4.1.1 Laboratory Measurements .................... 7-8
    7.4.1.2 Long-Path Measurements ..................... 7-11
  7.4.2 Zero-Filling Effects ............................... 7-12
  7.4.3 Apodization Effects ............................... 7-15
7.5 Effect of Resolution on Quantitative Analyses ...... 7-18
  7.5.1 Studies from the Literature ....................... 7-18
  7.5.2 Case Study: The Effect of Resolution and Related Parameters on the CLS Analysis of Multicomponent Mixtures ........................................ 7-21
    7.5.2.1 Mixtures of CO and ¹³CO ...................... 7-22
    7.5.2.2 Mixtures of Acetone, Methylene Chloride, and Ethanol 7-23
      7.5.2.2.1 Effect of the Number of Data Points on the CLS Analysis 7-24
      7.5.2.2.2 Effect of S/N Ratio on the CLS Analysis 7-26
    7.5.2.3 Mixtures of Methylene Chloride and Nitrous Oxide 7-28
    7.5.2.4 Conclusions and Recommendations Based on Case Study 7-29

7.6 General Conclusions and Recommendations .......... 7-30
7.7 Guidance for Selecting Resolution and Related Parameters ........................................ 7-31
7.8 References .............................................. 7-34

8 Quality Assurance and Quality Control .................. 8-1
8.1 Introduction and Overview ........................... 8-1
8.2 Project Plan Categories ................................ 8-2
8.2.1 Category Definitions ........................................ 8-3
8.2.2 Category I Points to be Addressed ...................... 8-3
  8.2.2.1 Project Description .................................... 8-4
  8.2.2.2 Project Organization and Responsibilities .......... 8-4
  8.2.2.3 QA Objectives .......................................... 8-5
  8.2.2.4 Site Selection and Sampling Procedures ............ 8-6
  8.2.2.5 Sample Custody ......................................... 8-6
  8.2.2.6 Calibration Procedures and Frequency ............... 8-6
  8.2.2.7 Analytical Procedures .................................. 8-7
  8.2.2.8 Data Reduction, Validation, and Reporting .......... 8-7
  8.2.2.9 Internal Quality Control Checks ...................... 8-8
  8.2.2.10 Performance and System Audits ...................... 8-8
  8.2.2.11 Preventive Maintenance ............................... 8-8
  8.2.2.12 Calculation of Data Quality Indicators ............. 8-9
  8.2.2.13 Corrective Action ..................................... 8-9
  8.2.2.14 Quality Control Reports to Management .......... .... 8-9
  8.2.2.15 References ............................................. 8-9
  8.2.2.16 Other Items ............................................. 8-9

8.3 Case Study: QA Data Collected over Two and One-Half Months
  at a Semipermanent Field Site ................................. 8-10

8.4 Recommendations of Tests to Be Included in a QA Program
  for FT-IR Long-Path Monitors ................................ 8-16
  8.4.1 Noise Measurements ...................................... 8-17
  8.4.2 Stability of Instrument .................................. 8-17
  8.4.3 Accuracy and Precision .................................. 8-18
  8.4.4 Completeness and Representativeness of Data ........ 8-20
  8.4.5 Comparability of the Data ............................... 8-20
  8.4.6 Ancillary Measurements ................................... 8-21
  8.4.7 Documentation ............................................ 8-21

8.6 References ................................................... 8-21

9 Glossary of Terms for FT-IR Open-Path Remote Sensing ...... 9-1
  9.1 Introduction and Overview .................................. 9-1
  9.2 Terms ....................................................... 9-1
  9.3 References .................................................. 9-7

10 Bibliography .................................................. 10-1
  10.1 Introduction and Overview ................................ 10-1
  10.2 Publications ............................................... 10-1
## Figures

<table>
<thead>
<tr>
<th>Number</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-1</td>
<td>A Schematic of the Simplest Form of a Michelson Interferometer</td>
<td>2-4</td>
</tr>
<tr>
<td>2-2</td>
<td>Schematic of Interference Created by Division of Amplitude</td>
<td>2-5</td>
</tr>
<tr>
<td>2-3</td>
<td>Center Burst Increasing as the Wave Number Range Expands</td>
<td>2-7</td>
</tr>
<tr>
<td>2-4</td>
<td>Interferograms for a Range of 3500 cm(^{-1})</td>
<td>2-8</td>
</tr>
<tr>
<td>2-5</td>
<td>Interferogram of Two Cosine Waves as a Function of (\Delta T)</td>
<td>2-9</td>
</tr>
<tr>
<td>2-6</td>
<td>The Bistatic Configuration</td>
<td>2-13</td>
</tr>
<tr>
<td>2-7</td>
<td>The Monostatic Configuration</td>
<td>2-14</td>
</tr>
<tr>
<td>2-8</td>
<td>Data Reduction Flow Chart</td>
<td>2-24</td>
</tr>
<tr>
<td>3-1</td>
<td>Single-Beam Spectrum Along a 414-m Path</td>
<td>3-3</td>
</tr>
<tr>
<td>3-2</td>
<td>Single-Beam Spectrum Recorded at a 20-m Total Path Length</td>
<td>3-4</td>
</tr>
<tr>
<td></td>
<td>Indicating Nonlinear Operation</td>
<td></td>
</tr>
<tr>
<td>3-3</td>
<td>Region Between 1000 and 1025 cm(^{-1})</td>
<td>3-5</td>
</tr>
<tr>
<td>3-4</td>
<td>Subtraction of Spectra for the Determination of Line Shifts and</td>
<td>3-8</td>
</tr>
<tr>
<td></td>
<td>Resolution Changes</td>
<td></td>
</tr>
<tr>
<td>3-5</td>
<td>Effect of Stray Light</td>
<td>3-9</td>
</tr>
<tr>
<td>3-6</td>
<td>The RMS Baseline Noise Measured Between 980 and 1020 cm(^{-1}),</td>
<td>3-11</td>
</tr>
<tr>
<td></td>
<td>2480 and 2520 cm(^{-1}), and 4380 and 4420 cm(^{-1})</td>
<td></td>
</tr>
<tr>
<td>4-1</td>
<td>Synthetic (I_0) Spectrum</td>
<td>4-3</td>
</tr>
<tr>
<td>4-2</td>
<td>A Possible Configuration for (I_0) Spectrum Acquisition</td>
<td>4-4</td>
</tr>
<tr>
<td>4-3</td>
<td>Procedure for Acquiring a Short-Path Background Spectrum</td>
<td>4-6</td>
</tr>
<tr>
<td>5-1</td>
<td>The Portion of a Single-Beam Spectrum over Which Methane Absorbs</td>
<td>5-5</td>
</tr>
<tr>
<td>5-2</td>
<td>Methane Region with Synthetic Background Spectrum Superimposed</td>
<td>5-5</td>
</tr>
<tr>
<td>5-3</td>
<td>Methane Reference Spectrum and the Calculated Absorption Spectrum</td>
<td>5-7</td>
</tr>
<tr>
<td>5-4</td>
<td>Water Vapor Spectrum Made for the Methane Absorption Region</td>
<td>5-7</td>
</tr>
<tr>
<td>5-5</td>
<td>Atmospheric Ozone Absorption Spectrum and Ozone Reference Spectrum</td>
<td>5-8</td>
</tr>
<tr>
<td>5-6</td>
<td>Ozone Measured at Research Triangle Park During June</td>
<td>5-9</td>
</tr>
<tr>
<td>6-1</td>
<td>Sulfur Hexafluoride Reference Spectrum</td>
<td>6-6</td>
</tr>
<tr>
<td>6-2</td>
<td>Aerial Photograph of a Superfund Site</td>
<td>6-9</td>
</tr>
<tr>
<td>7-1</td>
<td>Single-Beam IR Spectra of CO(_2) Measured at 0.25-, 0.50-, 1.0-,</td>
<td>7-7</td>
</tr>
<tr>
<td></td>
<td>and 2.0-cm(^{-1}) Resolution with No Apodization and No Additional Zero Filling</td>
<td></td>
</tr>
<tr>
<td>7-2</td>
<td>Single-Beam IR Spectra of Water Vapor Measured at 0.25-, 0.50-,</td>
<td>7-8</td>
</tr>
<tr>
<td></td>
<td>1.0-, and 2.0-cm(^{-1}) Resolution with No Apodization and No</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Additional Zero Filling</td>
<td></td>
</tr>
<tr>
<td>7-3</td>
<td>Single-Beam IR Spectra of Water Vapor Measured at 2-, 1-, and</td>
<td>7-9</td>
</tr>
<tr>
<td></td>
<td>0.5-cm(^{-1}) Resolution over a 150-m Path</td>
<td></td>
</tr>
<tr>
<td>7-4</td>
<td>IR Spectra of Water</td>
<td>7-13</td>
</tr>
</tbody>
</table>
Absorbance Spectra of CO\textsubscript{2} Measured at 0.25-cm\textsuperscript{-1} Resolution with a Zero-Filling Factor of 1, 0.5-cm\textsuperscript{-1} Resolution with No Zero-Filling, and 0.5-cm\textsuperscript{-1} with a Zero-Filling Factor of 2 ........................................ 7-14
 Absorbance Spectra of Water Vapor Measured at 0.25-cm\textsuperscript{-1} Resolution with a Zero-Filling Factor of 1, 0.5-cm\textsuperscript{-1} Resolution with a Zero-Filling Factor of 2, and 1-cm\textsuperscript{-1} Resolution with a Zero-Filling Factor of 4 ............. 7-15
 Absorbance Spectra of CO Measured at a Nominal 0.125-cm\textsuperscript{-1} Resolution with No, Triangular, Happ-Genzel, and Norton-Beer-Medium Apodization Functions .............................................. 7-16
 Absorbance Spectra of Water Vapor Measured at 0.5-cm\textsuperscript{-1} Resolution with a Zero-Filling Factor of 2 and with No, Triangular, Happ-Genzel, and Norton-Beer-Medium Apodization Functions .............................................. 7-17
 Reference 0.25-cm\textsuperscript{-1} Spectra of \textsuperscript{13}CO and CO and Spectra of Synthetic Mixtures of 150 ppm CO and 100 ppm \textsuperscript{13}CO Measured at 0.25-, 0.5-, 1.0-, and 2.0-cm\textsuperscript{-1} Resolution .............................................. 7-23
 Concentration Calculated from CLS Analysis vs. Known Concentration for \textsuperscript{13}CO/CO Mixtures Measured at 2-cm\textsuperscript{-1} Resolution .............................................. 7-24
 Reference 0.25-cm\textsuperscript{-1} Spectra of Acetone, Methylene Chloride, and Ethanol and Spectra of Synthetic Mixtures of 100 ppm Acetone, 100 ppm Methylene Chloride, and 500 ppm Ethanol Measured at 1.0-, 2.0-, and 4.0-cm\textsuperscript{-1} Resolution .............................................. 7-25
 Spectra of Synthetic Mixtures of 100 ppm Acetone, 100 ppm Methylene Chloride, and 500 ppm Ethanol Measured at 1-cm\textsuperscript{-1} Resolution with 0, 1, 5, 10, and 25% Noise Added .............................................. 7-27
 Reference 0.25-cm\textsuperscript{-1} Spectra of N\textsubscript{2}O and Methylene Chloride and Spectra of Synthetic Mixtures of 50 ppm N\textsubscript{2}O and 100 ppm Methylene Chloride Measured at 0.25-, 0.5-, and 1.0-cm\textsuperscript{-1} Resolution .............................................. 7-28
 Concentration Calculated from CLS Analysis vs. Known Concentration for N\textsubscript{2}O/Methylene Chloride Mixtures Measured at 0.25-cm\textsuperscript{-1} Resolution .............................................. 7-29
 Return Signal Magnitude of the FT-IR Monitor Measured Daily at 0700 and 1200 .............................................. 8-11
 The RMS Baseline Noise Measured Between 980 and 1020 cm\textsuperscript{-1}, 2480 and 2520 cm\textsuperscript{-1}, and 4380 and 4420 cm\textsuperscript{-1} .............................................. 8-12
 Repeatability of the Position of the Water Vapor Singlet at 1014.2 cm\textsuperscript{-1} Measured on November 10, 1993, December 22, 1993, and January 4, 1994 .............................................. 8-13
 Measurement of Ambient Methane Concentration and Single Beam Intensity at 987 cm\textsuperscript{-1} on November 17 and 18, 1993 .............................................. 8-14
 Peak Area of 2998.8-cm\textsuperscript{-1} Absorption Band of CH\textsubscript{4} and the 1014.2-cm\textsuperscript{-1} Absorption Band of Water Vapor Measured on November 17-18, 1993 .............................................. 8-16
<table>
<thead>
<tr>
<th>Number</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>6-1</td>
<td>Minimum Usable Path Lengths</td>
<td>6-5</td>
</tr>
<tr>
<td>7-1</td>
<td>Resolution Test Data</td>
<td>7-7</td>
</tr>
<tr>
<td>7-2</td>
<td>Optimal Wave Number Region and Minimum Resolution</td>
<td>7-19</td>
</tr>
<tr>
<td>7-3</td>
<td>Effect of the Number of Data Points on the CLS Analysis</td>
<td>7-25</td>
</tr>
<tr>
<td>7-4</td>
<td>The Effect of Zero Filling on the CLS Analysis</td>
<td>7-26</td>
</tr>
<tr>
<td>7-5</td>
<td>Effect of Noise on the CLS Analysis</td>
<td>7-27</td>
</tr>
</tbody>
</table>
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Chapter 1
Introduction

The Michelson interferometer has had a remarkable history in that new uses for the device have been found for more than 100 years. One use of the interferometer that has experienced rapid growth since the mid-1960s is as the main optical component of Fourier transform infrared (FT-IR) spectrometers. Although there have been several applications of FT-IR spectrometers to unique and difficult problems, the majority of FT-IR systems have been used to make qualitative measurements under controlled conditions in the laboratory. More than 20 years ago, some efforts were made to use the instrument for making quantitative measurements of atmospheric gaseous pollutants over extended open paths (Hanst 1970; Herget and Brasher 1979). Although these efforts were largely successful, they were overlooked by the great majority of people engaged in environmental monitoring. During the 1980s there was steady but slow progress in development of the technique. In the late 1980s, a revival of the technique occurred, initiated in part during a meeting of the Chemical Manufacturer’s Association in Houston (Russwurm and McClenny 1990; Levine et al. 1991; McClenny et al. 1991), and today there is a large amount of developmental activity taking place. (See the bibliography in Chapter 10.)

This document describes the components of FT-IR monitors and is intended to provide guidance for the FT-IR operator in field monitoring applications. It is a point of reference for further development and evaluation of FT-IR open-path monitors as field instruments.

1.1 Overview of Document

A brief discussion of the FT-IR open-path monitor and its function is given in Chapter 2, along with a more in-depth description of the various components of the sensor. Chapter 3 includes the preliminary procedures for setting up the FT-IR instrumentation for monitoring. Chapter 4 is a discussion of background spectra, and Chapter 5 is a discussion of water vapor spectra. Chapter 6 presents guidance on how to set up the monitoring instruments within the physical constraints of a site. Chapter 7 presents experimental data that illustrate the effect of resolution and related parameters on the spectral data. Chapter 8 contains quality control and quality assurance guidelines, incorporating portions of an approved quality assurance plan, and includes selected QA data we collected over a recent one-year period. Chapter 9 is a glossary of terms, and Chapter 10 is a general bibliography of work that addresses FT-IR monitoring and the principles of FT-IR spectrometry.

Each chapter begins with a summary highlighting the primary contents of the chapter. This is followed by an introduction and overview of the chapter.
1.2 References


Chapter 2
The Fourier Transform Spectrometer

SUMMARY

The major topics discussed in this chapter are the following:

- The basic principles of FT-IR spectrometers
  - Resolution and throughput
  - Detectors and sources
  - Electronics and computer requirements

- The fundamental aspects of the interferogram, the Fourier transform, and single-beam spectra

- The optics used in long-path, open-path FT-IR monitors
  - Transfer optics, telescopes, and beam return optics
  - Monostatic and bistatic configurations

- Beer's law and data analysis procedures

2.1 Introduction and Overview

This chapter describes the components of a complete FT-IR monitoring system, which include the following: the FT-IR spectrometer, the transmitting and receiving optics, the electronics, the computer, and the data output. The discussions in this chapter are based on the general configurations of instruments that are commercially available at the time of this writing. There are currently other manufacturers with instruments in the design or developmental stages.

However, familiarity with the basic principles of FT-IR spectrometry is required if proper operational choices are to be made under varying field conditions. And, the better the operator understands the functions of the instrument, the more likely it is that reliable data will be produced. This chapter includes a description of long-path, open-path FT-IR monitors and an in-depth discussion of the various components of FT-IR spectrometers. The integral components of an FT-IR monitoring system, which include the interferometer, detector, IR source, transfer and beam-return optics, electronics, and computers, are described. The fundamental processes of FT-IR spectrometry, including the interference phenomenon, generation of the interferogram, optical throughput, resolution, and the Fourier transform, are
explained. A brief discussion of Beer's law and its application to the data analysis is provided. In addition to providing quantitative results, the relationships that are explained by Beer's law are important when estimating detection limits and determining optimum path lengths.

The heart of an FT-IR system is the interferometer. Most, but not all, commercial instruments use the Michelson interferometer. A detailed description of the Michelson interferometer is provided in Section 2.2. The trace of the output of the interferometer is referred to as an interferogram. The interferogram is the actual data produced by an FT-IR spectrometer and contains all of the information about the spectrum. However, the information contained in the interferogram is not in a form that is readily recognizable to most spectroscopists. To change the data into a form that is more easily interpretable, the raw data are converted into a spectrum (a plot of intensity versus wave number) by performing a Fourier transform on the interferogram. A computer system with the appropriate software packages is used to apply this and all other necessary mathematical functions to the data. Although the execution of these calculations is virtually invisible to the operator, a basic understanding of the principles involved is necessary to ensure that the optimum parameters are used to collect and process the FT-IR data.

All quantitative data analysis in long-path, open-path FT-IR spectrometry is based on Beer's law. Beer's law states that for a constant path length, the IR energy traversing an absorbing medium diminishes exponentially with concentration. Mathematically, this is written as

\[ I(v) = I_0(v)e^{-\alpha(v)CL} \]

where \( I_0(v) \) is the intensity of the incident beam, \( \alpha(v) \) is the optical absorption coefficient of the absorbing material (e.g., target gas) as a function of wave number \( v \), \( C \) is the concentration of the target gas, and \( L \) is the path length.

Two primary configurations, monostatic and bistatic, are used to transmit the IR beam along the path, as described in Section 2.3. The monostatic system has both the IR source and the detector at one end of the path and a retroreflector at the other. The retroreflector returns the beam either along or collinear to the original path, which doubles the effective path length and thus the measured absorbance of the target gas. The bistatic system has the detector at one end of the path and the source at the other. This configuration minimizes the optical components that are required for open-path monitoring. However, in the bistatic system, the IR beam is limited to a single pass along the path. Both types of configurations are currently in use for environmental monitoring.

2.2 The Michelson Interferometer

The primary optical component in an FT-IR instrument is a Michelson interferometer. It is not generally necessary
to have a fundamental understanding of how the interferometer functions to obtain reliable data with an FT-IR instrument. However, familiarity with some of the aspects of the interferometer is required if proper operational choices are to be made under varying field conditions. To that purpose, a brief discussion of the optics of the FT-IR instrument is included in this subsection. The following major topics are discussed: interference (Section 2.2.1), resolution (Section 2.2.2), throughput (Section 2.2.3), the detector (Section 2.2.4), and the IR source (Section 2.2.5).

A variety of devices have been used over the last 200 years to study interference phenomena. These devices are conveniently classified by the amount of four primary attributes that they exhibit: monochromatism, fringe localization, fringe production by division of wave front or by division of amplitude, and double or multiple beams. The interferometric device that today bears his name was first introduced by A. A. Michelson in 1881 (Michelson 1881). It is the most famous of a group of interferometers that produce interference fringes by the division of amplitude. Four years after Michelson introduced the interferometer, it was shown that the Fourier transform of the interferogram was the original spectrum or intensity as a function of wavelength. The Michelson interferometer has been used to define and measure the standard meter, to measure the angular separation of binary stars, and to provide the experimental data for one of the four cornerstones of relativity theory. During recent times, the Michelson interferometer has been used successfully to measure the concentrations of various chemicals that absorb energy in the IR portion of the electromagnetic spectrum. (See Chapter 10, Bibliography.) It is currently being developed as an instrument to make similar measurements over extended open paths, and it is in this context that the interferometer is discussed here.

A schematic of the simplest form of a Michelson interferometer is shown in Figure 2-1. It consists of a beam splitter and two mirrors, one of them movable. The figure also shows an arrangement for the light source and the detector. For the most accurate use, the two mirrors must be kept perpendicular to one another. One of the two mirrors moves along the optic axis. During this motion the perpendicularity cannot change. This requirement can represent a stringent limitation for the mechanisms involved with the motion. The light incident on the beam splitter should be collimated, because uncollimated light gives rise to poor resolution.

2.2.1 Interference

This section is presented for completeness and because there seems to be some confusion as to how the interferogram arises. It is somewhat mathematically rigorous and can be omitted without jeopardizing the ability of the operator to obtain reliable FT-IR data.

Interference is the underlying physical phenomenon that allows a Fourier transform
Figure 2-1. A Schematic of the Simplest Form of a Michelson Interferometer.

An instrument to obtain spectrometric data. The interference phenomenon cannot be physically explained by the simple addition of the intensities of two or more optical beams. The amplitudes of the individual interfering beams must be added according to the principle of superposition, and the total intensity must be calculated from that result. Interference phenomena are linear in amplitude. The principle of linear superposition, which is operating here, follows directly from Maxwell's equations and the fact that these equations are linear differential equations. To arrive at the basic equation that describes how the Michelson works, consider the arrangement of Figure 2-2. A monochromatic electromagnetic plane wave is incident on a device at A that divides its amplitude into two components. After the division, the individual beams traverse a medium along different paths and are somehow recombined at a point P in space. On arrival at point P, the two beams, which need not be collinear, have the following amplitudes.

\[ A_1 = A_0 e^{i(\omega t - 2\pi n T_1 / \lambda)} \]
\[ A_2 = A'_0 e^{i(\omega t - 2\pi n T_2 / \lambda)} \]

The two \( A_0 \) terms are the amplitudes of the individual beams, the \( \omega \) is the angular frequency of the radiation, \( n \) is the index of refraction of the medium, and the two \( T \) terms are the physical path lengths that each
beam has traversed. The product $nT$ is called the optical path length that the beam has traveled. At point $P$, where the two beams are recombined, the total amplitude is the sum of these terms. The intensity is then given by the product of this sum and its complex conjugate. Thus the intensity at point $P$ is given by Equation 2-1.

$$I(P) = A_0 A_0^* + A_0 A_0^* + A_0 A_0^* e^{-i[2\pi(\tau_2 - \tau_1)/2]}$$

$$+ A_0 A_0^* e^{i[2\pi(\tau_2 - \tau_1)/2]}$$

(Eq. 2-1)

The first two terms are the intensities of the original two beams, and the last two terms are called the interference terms. When the amplitudes $A_0$ and $A_0^*$ are equal, they can be combined.

By using the relation $2\cos x = e^x + e^{-x}$, the intensity at point $P$ is given by Equation 2-2.

$$I(P) = 2I_0 \left[1 + \cos \left(2\pi \left(\frac{T_2 - T_1}{\lambda}\right)\right)\right]$$

(Eq. 2-2)

Here $I_0$ is the intensity of either beam. Thus as the difference of the path length, $T_2 - T_1$, changes, the intensity at point $P$ can vary from $0$ to $4I_0$. The fact that $I$ can be $4I_0$ does not violate the conservation of energy law. There is no physical requirement that the intensity at every point in space be $2I_0$. The requirement is that the interference term averaged over space must be zero (Rossi 1957).

When a plane monochromatic wave is incident on the beam splitter of the Michelson interferometer, the amplitude ideally is evenly
divided along each leg. At any position of the moving mirror, the detector output is proportional to the integral of the intensities over wavelength, and this recording is called the interferogram. From Equation 2-2, it is seen that at zero path \( (T_2 - T_1 = 0) \) difference, the cosine term is 1 for all wavelengths. Thus for all wavelengths, the intensity is \( 4I_0 \), and the output of the detector is large compared to any other mirror position. This is quite noticeable in the interferogram and is commonly called the center burst.

This center burst does not appear when the radiation is monochromatic. Figure 2-3 shows how the center burst builds as the wave number range is expanded to include more wavelengths. The interferograms in this figure were calculated from Equation 2-1 in the following way. All the wave numbers have the same intensity and add incoherently. The wave number value was stepped in increments of 0.1 cm\(^{-1}\). The retardation (actually, the term \( T_2 - T_1 \)) was taken in increments of the wavelength of a He-Ne laser. At each position of the mirror the proper phase for each wave number was used to calculate the intensity, and then the intensities were added. The interferograms were actually calculated for 6000 incremental movements of the mirror; however, only a portion of the data is shown for clarity. The interferograms in Figure 2-3 are for (A) a 2-cm\(^{-1}\) range, (B) a 50-cm\(^{-1}\) range, (C) a 500-cm\(^{-1}\) range, and (D) a 3500-cm\(^{-1}\) range. The two interferograms shown in Figure 2-4 are for a range of 3500 cm\(^{-1}\), but curve B has a 1500 K blackbody radiation curve superimposed on it, and it appears quite similar to the interferogram actually recorded by the FT-IR spectrometers.

Equation 2-2 shows that as the mirror moves, the path difference causes a modulation of the intensity at each wavelength. The modulation can be used to advantage in open-path FT-IR monitors. For example, if the IR beam traverses the interferometer before it is sent along the open path, any background radiation entering the system from the surroundings is not modulated and will not be processed by the electronics. However, a portion of this unmodulated light will still be incident on the detector and in extreme situations could cause the detector to become saturated. Therefore, it is prudent to avoid setting the instrument up along a path that includes bright (hot) IR sources.

2.2.2 Resolution

The resolution of an instrument determines how close two absorption features can be and still be separated enough for analysis. There are several criteria for this instrument parameter, but the one most often used for the FT-IR instrument is described below. Equation 2-2 shows that all wavelengths have a maximum and are in phase with one another at zero path difference. The most common definition of resolution for the FT-IR spectrometer states that two absorbing features centered at wavelengths \( \lambda_1 \) and \( \lambda_2 \) will be resolved if the mirror moves at least to the point where these two wavelengths are again in phase. To
Figure 2-3. Center Burst Increasing as the Wave Number Range Expands.
Figure 2-4. Interferograms for a Range of 3500 cm\(^{-1}\). Interferogram B has a 1500 K blackbody radiation spectrum superimposed on it.
Figure 2-5. Interferogram of Two Cosine Waves vs. $\Delta T$. The wavelengths differ by 10 cm. The minimum occurs when the two waves are 180° out of phase.

To determine when this occurs, the following example may be considered. If only two spectral features situated at $\lambda_1$ and $\lambda_2$ make up the spectrum, then the interferogram is made up of two spectra, each described by Equation 2-2. The result of adding these two spectra is shown in Figure 2-5 and is given by Equation 2-3.

$$I(P) = 4I_0 \left[ 1 - \cos \left( \frac{\pi \Delta T}{\lambda_1} \left( \frac{1}{\lambda_1} - \frac{1}{\lambda_2} \right) \right) \cos \left( \frac{\pi \Delta T}{\lambda_2 + \lambda_1} \left( \frac{1}{\lambda_1} + \frac{1}{\lambda_2} \right) \right) \right]$$

(Eq. 2-3)

The second cosine term produces a high-frequency signal that is modulated by a low-frequency signal described by the first cosine term. It is the first term that is of interest when determining the resolution of the system. The signal is a maximum when the argument of this cosine term is $2N\pi$, where $N = 0, 1, 2, \ldots$. Thus, the first time that the two wavelengths are in phase after the center burst is when $N = 1$, so that

$$\pi \Delta T (1/\lambda_2 - 1/\lambda_1) = 2\pi$$
This implies that

\[ \Delta T = 2/(1/\lambda_2 - 1/\lambda_1) \]

However, the term in the denominator is the difference in the wave numbers of the absorption peaks, so that \( \Delta T = 2/\Delta v \). Thus, if the operator desires a resolution of 0.5 cm\(^{-1}\), the optical path difference must be 4 cm. Because the beam traverses the path in the interferometer twice, the actual motion of the mirror must be only 2 cm. It should be noted here that this is an idealized result. The fact that the interferogram is first truncated and then apodized changes this result somewhat (Marshall and Verdun 1990; Beer 1992).

The question of what resolution should be used for a specific data collection task is not addressed in this section. It is discussed in more depth in Chapter 7. The answers to the resolution questions are specific to the gases to be monitored and the effects of water vapor in their regions of absorbance. At the present time, each monitoring situation must be considered separately.

### 2.2.3 Throughput

The throughput of an optical system is defined as the product of the area of an aperture \( A \) and the solid angle \( \Omega \) of the light beam at that aperture. This quantity is theoretically a constant throughout the system, so that once it is defined for an aperture it is known for all apertures. For small angles, the solid angle of the beam can be shown to be equivalent to the product \( \pi \theta^2 \), where \( \theta \) is the half angle of the field of view of the instrument. It can be shown that the throughput is related to the f\# of the system by recognizing that \( \theta = 1/(2f\#) \), so that the throughput is equal to \( A \pi (1/(4f\#))^2 \). With FT-IR instruments, the selection of the system f\# is generally a compromise. An important consideration is the solid angle of the beam as it traverses the interferometer. A portion of the beam traversing the interferometer at a large angle will travel over a longer path through the interferometer, and a beam traversing at a smaller angle will travel over a shorter path. This angular dispersion tends to degrade the resolution of the instrument, because energy at the same wavelength appears to the interferometer as though it covers a range of wavelengths. Smaller f\#s are at first attractive because they indicate that a smaller aperture can be used. However, small f\#s imply large solid angles and therefore a loss in resolution. The manufacturers of these instruments have taken this into account in the instrument design, but nevertheless the aperture size is fixed, and once a specific instrument is purchased, there is little, if anything, the operator can do to change the throughput.

### 2.2.4 The Detector

The detector in most FT-IR instruments used for monitoring atmospheric pollutant gases is a semiconductor device made of mercury, cadmium, and telluride, commonly called an MCT detector. There are three modes of operation for this device, as a photovoltaic device, as a photoelectromagnetic device, and as a photoconductive device. The MCT photoconductive detector
is the one most often used in the FT-IR instrument. This device converts a beam of photons to an electrical current that can be measured. In addition to the spectral region that the detector responds to, the two most important parameters of the detector are the noise equivalent power (NEP) and the sensitivity of the detector in terms of a quantity called $D^*$ (pronounced "Dee Star"). For an MCT photoconductive device, the spectral response ranges from 2 to 20 μm, or from 500 to 5000 cm$^{-1}$. The NEP is given in terms of W/(Hz)$^{1/2}$. For the detectors used in FT-IR instruments, this parameter has a value of about $5 \times 10^{-12}$. The user should be aware that this parameter represents a measure of the inherent noise in the detector and that small numbers are better than large numbers. The $D^*$ is a measure of the sensitivity of the detector and has units of cm(Hz)$^{1/2}$/W. This number is actually defined as the ratio of the square root of the detector area to the NEP, or $D^* = \sqrt{A/\text{NEP}}$. For the MCT photoconductive detectors, this number is about $5 \times 10^{10}$ at 10 μm. Here, larger numbers are better. However, there is little that the operator can do about the magnitude of these parameters once the instrument is purchased. But if the detector has to be replaced, some acceptance criteria for the NEP and the $D^*$ should be specified.

There are, in general, two types of MCT detectors available, wide band and narrow band. Each has somewhat different characteristics. Wide band MCT detectors cut off at around 500 cm$^{-1}$, whereas the narrow band MCT detectors cut off near 600 cm$^{-1}$. For long-path measurements the region below 722 cm$^{-1}$ is nearly opaque because of absorption in the atmosphere, so a wide band detector does not offer any real advantages. Also, the $D^*$ for the narrow band detector is 5 to 10 times higher than that for the wide band detector. There is also an indium antimonide (In-Sb) detector that can be used in the higher wave number region with advantage. One particular application for using this type of detector is the measurement of HF.

An important requirement for the detectors in FT-IR monitoring systems is that they must be cooled to operate properly. Liquid nitrogen temperatures (77 K) allow optimum operation of these detectors. Currently, two techniques are used to cool the detector. The first is to place the detector in a Dewar that uses liquid nitrogen as a refrigerant. For this mode, a supply of liquid nitrogen must be available for use in the field, and the operator must fill the Dewar periodically. This has not been a major problem in the past, as the liquid nitrogen requirement is only a few pints per day of operation. The second technique to achieve cold temperatures is with a cryogenic cooler, such as a Stirling engine or a closed-cycle helium refrigeration system. In the Stirling engine, the heat is exchanged through a wall from the enclosure to cool the gas. Currently, the major problems with this cooling device is the mean time between failures is too short, and these coolers seem to add noise to the spectra. One-half year of continuous operation is about the maximum that can be expected. If unattended operation is a necessity, the Stirling engine is
one choice. Other options for unattended operation are the use of larger detector Dewars with longer hold times or devices that automatically refill the detector Dewar.

2.2.5 The IR Source

All IR sources that are available today for use with the FT-IR monitor are heated elements that are open to the atmosphere. They are resistive devices that radiate approximately as black body radiators. These devices operate at a color temperature from 1200 to 1500 K. The Wien displacement law states that the product of the wavelength of maximum power output and the temperature of the source, $\lambda_{\text{max}} T$, is a constant equal to 0.2987 cm K$^{-1}$. This indicates that there is an inverse relation between peak wavelength and source temperature. Planck’s radiation law shows that there is more energy at all wavelengths for hotter sources. The ideal would be a source that is at about 3000 K, so it would have a peak at about 1.1 μm. The materials that are necessary to make such a source have not been available until now.

Perhaps the most detrimental characteristic of the available sources is that they are large compared to the focal length of the collimating optics. From geometrical optics, it is clearly seen that the beam can never be better collimated than the angle that the source subtends at the collimating optics (lens or mirror). Thus, all available FT-IR spectrometers have beam divergences that are too large for the rest of the optics. This means that retroreflectors or receiving optics are overfilled, and much of the initially available energy is lost. Ultimately, this divergence restricts the path length that can be used to advantage. Perhaps as further developments occur, a small hot source will be developed that will minimize this difficulty.

2.3 Transfer Optics, Telescopes, and Beam-Return Optics

There are two primary geometrical configurations available for transmitting the IR beam along the path. One is a bistatic system (Figure 2-6); the other a monostatic system (Figure 2-7). The monostatic system has both the IR source and the detector at the same end of the path, whereas the bistatic system has the detector at one end of the path and the source at the other. In the bistatic system, the optical path length is equal to the physical path length, whereas in the monostatic configuration, the optical path length is twice the physical path length. In this document, we always refer to the optical path length. The reflecting optics for a bistatic FT-IR monitoring system are relatively straightforward (see Section 2.3.1), whereas the optics in a monostatic system may include one or more telescopes, an additional beam splitter, and return-beam optics. The possible configurations for a monostatic system are described in Section 2.3.2.

There are several types of telescopes that could be used to transmit and collect the IR beam. Those in current use are the Cassegrain and the Newtonian telescopes. They are optically equivalent, and the only difference is the placement of the diagonal mirror that removes the beam from the
Figure 2-6. The Bistatic Configuration.
Figure 2-7. The Monostatic Configuration.
telescope. In the Cassegrain, the beam exits from the end of the telescope, and for the Newtonian it exits from the side. Optically, the number of reflections is the same, and other things being equal, the reflection losses are also essentially the same. The geometrical configuration for the whole instrument is slightly different for these two designs. In one, the overall package enlarges vertically, whereas in the other it grows longitudinally. These are minor points as far as instrument operation is concerned.

In most cases, the beam is expanded before it is sent along the path, and in principle there are essentially no size limits. The physical quantity that is of interest to the operator is how much of the IR beam is absorbed by the gases of interest. This is, in general, very small (about 1 part in a 1000). This fact is not changed by expanding the beam. In general, beam expansion allows more energy to be transmitted along the path, thereby increasing the overall signal-to-noise ratio (S/N). There are, of course, practical limitations to the size of the optics that can be accommodated.

As a rule, the optics in either system are reflecting optics rather than refracting optics to avoid transmission losses. Once the IR energy has been collected, the optics must reduce the size of the beam so that it can pass through the system to the detector without vignetting and also to set the solid angle of the beam so that the resolution remains acceptable.

2.3.1 Bistatic System

The bistatic configuration minimizes the optical components that are required for open-path monitoring. At the source end of the path there must be some method for collimating the beam. This can easily be done with a mirror shaped as a parabola or one of the other conic sections. At the receiving end of the path, a collector, similar in design to the collimator, may be used to transfer the beam to the interferometer and the detector. In commercially available instruments, the diameter of the collector generally is the same as that of the transmitter, although there is no optical necessity for this choice.

There are two configurations that can be used for bistatic systems. One configuration places the IR source, interferometer, and transmitting optics at one end of the path and the receiving optics and detector at the other end (Figure 2-6A). The advantage of this configuration is that the IR beam is modulated along the path, which enables the unmodulated background radiation to be rejected by the system's electronics.

The other configuration places the IR source and transmitting optics at one end of the path and the receiving optics, interferometer, and detector at the other end of the path (Figure 2-6B). This is the more common configuration of bistatic systems in current use. The main drawback to this configuration is that the IR source is not modulated before it is transmitted along the path. Therefore, the system has no way to
distinguish between the active IR source and the background IR radiation.

Because the IR radiation makes only a single pass through the optics, less of the radiation is lost in bistatic systems. Also, this makes the systems more amenable to passive measurements, such as emission measurements or absorption measurements with a natural background hot source.

Ambient monitoring in confined areas or rough terrain may pose certain logistics problems with a bistatic system. One is that this mode of operation requires two power sources, one at each end of the path. Another is that there is only one pass through the absorbers. The absorbance for most gases of interest is very small, and for short paths, as those encountered with plumes, a single pass through the gas may be insufficient.

2.3.2 Monostatic System

There are currently two techniques in use for returning the beam along the optical path when the monostatic mode is used. One is to set up an arrangement of mirrors that translates the beam slightly for its return path, and another is to place a retroreflector array at the end of the path. These two configurations are optically equivalent.

In the first configuration, an optical system is placed at the end of the path that translates the IR beam slightly so that it does not fold back on itself (Figure 2-7A). The receiving end then has a second telescope slightly removed from the transmitter with the detector at the primary focus. This technique circumvents a possible objection to the second monostatic configuration.

The second configuration for the monostatic monitoring mode uses the same telescope for the transmitting and the receiving optics and uses a retroreflector array at the end of the beam (Figure 2-7B). A retroreflector is an arrangement of mirrors that reflects the beam so that the incident and reflected directions of propagation are collinear but opposite to one another. It is made of three reflecting surfaces that are mutually orthogonal, such as the floor and two adjacent walls of a room. It is often assumed that after reflection the beam returns along the same path over which it was transmitted. That is true only in a gross sense because if the beam is small compared to the reflecting surfaces, a measurable translation takes place, and there is always an inversion and a reversion of the beam. In order to transmit and receive with the same optics, a beam splitter must be placed in the optical path. An objection to this configuration is that the IR energy must traverse this beam splitter twice, once on the transmitting end and once on the receiving end. The most effective beam splitter transmits 50% of the light and rejects the other 50%. Thus, in two passes, the transmission is only 25% of the original beam. Because this loss affects the S/N, it may be a significant drawback of this configuration of the monostatic mode.
Because of the design of the orthogonal mirror retroreflector, it is quite insensitive to small motions such as those caused by a wind. Retroreflector arrays are also very easy to align with the transmitting-receiving telescope, and a few degrees of misalignment will pose no problem to the operator. This does not seem to be the case with the spherical mirror configuration. Although this arrangement is also easy to align, it seems somewhat more sensitive to a small error.

2.4 The Electronics

An in-depth discussion of the electronics of the FT-IR system is beyond the scope of this document. However, some points that are of interest to the operator are covered. The interferogram is in the form of intensity versus position of the moving mirror. As the mirror moves, the detector measures a varying intensity, and this signal is first amplified and then sent to an analog-to-digital (A/D) converter for digitization. There are two important features that pertain to this digitization process.

The first concerns the dynamic range of the signal, which can actually be too large. There must be enough resolution in the A/D converter so that the least significant bit can always be reserved for recording the noise in the system. If this not the case, the spectrum derived by performing the Fourier transform on the interferogram will be distorted. For this reason, most commercially available instruments today use a 16-bit A/D converter. This means that if the noise is recorded on the least significant bit, the highest signal that can be recorded is a factor of $2^{16}$ above the noise. This is not really as high as it seems. For example, if the noise is about 1 $\mu$V, the largest signal that can be recorded is about 0.5 V.

The second feature concerns the amount of data that has to be recorded in digital form so that the original waveform can be reproduced. There is some relation between the rate at which a signal varies and the number of sampling pulses that are needed to reproduce it exactly. The sampling theorem from modern communication theory, sometimes called the Nyquist theorem, states that at least $2/f_m$ equally spaced samples are needed each second to reproduce the waveform without distortion. Here $f_m$ is the maximum frequency component that is contained in the original waveform.

As the mirror of the Michelson interferometer moves, each wavelength is modulated at a frequency that is related to the velocity of the mirror and the particular wavelength. Since the light must traverse the distance from the beam splitter to the mirror twice (down and back) in any one cycle, 2 times the actual velocity can be used for the determination of the frequency range to which the electronics must respond. If the mirror moves at a speed of 1 cm/s and the wavelength range is from 2.5 $\mu$m to 20 $\mu$m, then the frequency range is from 8000 Hz to 1000 Hz. Thus, according to the Nyquist theorem, the digital sampling rate must be at least 16000 equally spaced samples per second.
There is also a need for measuring the position of the mirror and for signaling to the electronics when to record data. This is done by using an He–Ne laser. The laser beam is sent through the interferometer and modulated in the same way as all other wavelengths are. The amplifier output is capacitively coupled to the rest of the electronics so that the He–Ne interferogram is an ac signal with negative and positive parts. The zero crossings of this ac signal are sensed, and the instrument records a data point at the zero crossings. Electronically, zero crossings are easier to detect than maxima or minima because of the sign change.

2.5 The Computer

The final requirement for an FT-IR system is a computer. This discussion is meant to be a discussion of the minimum requirements only.

The data storage requirements can be fairly large if the interferogram is stored. This requires about 100 kilobytes for each interferogram recorded at 1-cm⁻¹ resolution, and some means for archiving the data must be available. For use with a generic, commercially available software package, the minimum requirements for a computer are that it must use a 80286 microprocessor and have 640 kilobytes of RAM, a monochrome screen, and enough capacity to store whatever the user thinks is a minimum block of data. If the same computer has to be used for data collection and data analysis, then, in addition, there must be enough space to store a set of library spectra. The set that is currently commercially available contains about 150 compounds and uses about 4 megabytes of memory. If surveillance work must be done, then a computer faster than a 80286 model must be used. For ordinary field work at Superfund sites, the ideal system seems to be either a single computer with the ability to operate in both foreground and background modes or two computers—one to control the instrument and to record the data and the other, a much more capable machine, to be used for data analysis.

In addition to the computer hardware, a software package is required that will control the FT-IR system and record either the interferogram (preferably) or the single-beam spectrum (Section 2.7.4) produced by the Fourier transform performed on the interferogram. There is currently one generic software package that can be configured for any of the commercially available FT-IR systems. Other FT-IR systems use proprietary software for data collection. In addition to data collection, the software should also provide some means for data analysis. This is discussed further in Section 2.6.5.

2.6 The Data Output

This section contains a discussion of the interferogram generated by the FT-IR system, the Fourier transform that is applied to the interferogram, and the single-beam spectrum into which the interferogram is transformed. The data reduction process, starting with the interferogram and ending with the unknown gas concentration, is
described. A discussion of Beer's law of how energy diminishes as it traverses an absorbing medium is presented first.

2.6.1 Beer's Law

The fundamental physical law that is quoted as the basis for all FT-IR data analysis is Beer's law (Beer 1852). It states that, for a constant path length, the intensity of the incident light energy traversing an absorbing medium diminishes exponentially with concentration. Mathematically, this is written as

\[ I(v) = I_0(v)e^{-a(v)CL} \]

(Eq. 2-4)

where \( I_0(v) \) is the intensity of the incident spectrum, \( a \) is the optical absorption coefficient of the gas and is a function of the wave number \( v \), \( C \) is the concentration of the gas, and \( L \) is the path length. There are many possible sets of units for these quantities that are variously used by the workers in FT-IR open-path monitoring. Whatever the set chosen, it must be noted that the product \( aCL \) must be a unitless quantity. Thus, if the absorption coefficient has units of \( (\text{cm} \cdot \text{atm})^{-1} \), the concentration must be in atmospheres and the path length must be in centimeters. One primary difficulty that confronts the user of FT-IR open-path monitors is determining the quantity \( I_0 \). This is discussed in detail in Chapter 4, Background Spectra.

The mathematical functional form of Beer's law explains many physical phenomena. These include atmospheric pressure as a function of altitude, thermal expansion of metal rods, radioactive decay, and the electrical discharge of capacitors, to name but a few. Although there is no physical basis for doing so, in the field of optical spectroscopy, this functional form is often stated by using logarithm to the base 10. The available analysis software also uses logarithms to the base 10. To understand how the change is to be made, consider the following.

The fundamental formula is \( Y = a^X \). The problem is to solve this equation for the quantity \( X \), which is the power that \( a \) must be raised to obtain \( Y \). To solve this, the concept of logarithms is introduced so that \( \log_a(Y) = X \log_a(a) \). This is read as "the logarithm of \( Y \) to the base \( a \) equals \( X \) times the logarithm of \( a \) to the base \( a \)." By definition, \( \log_a(a) = 1 \) so that

\[ X = \log_a(Y) \]

Logarithms can be determined by using any number for the base, but only two are commonly used in the physical sciences. They are the base 10 and the base \( e \). The number \( e \) is defined as the limit of \((1 + 1/N)^N\) as \( N \) goes to infinity. The number \( e \) occurs naturally in mathematics, and particularly it occurs naturally in equations like Beer's law. The question is then how must Beer's law be written to account for the change to base 10.

The logarithmic form of the equation \( Y = a^X \) can be written in any other base \( b \) such that

\[ \log_b(Y) = \frac{X}{\log_b(a)} \]
\[ \log_b(Y) = X \log_b(a) \]

Solving this for \( X \) gives

\[ X = \frac{\log_b(Y)}{\log_b(a)} \]

Substituting this \( X \) into the original equation gives

\[ \log_b(Y) = \log_b(Y) \log_b(a) \]

In terms of the bases \( e \) and 10, this is written as

\[ \log_{10}(Y) = \log_{10}(e) \log_e(Y) \]

Historically, natural logarithms are written by using the prefix \( \ln \), while logarithms to the base 10 are written with the conventional \( \log \) as a designator. Logarithms to all other bases also use the convention \( \log \) as nomenclature, but then the actual base is specified as a subscript.

If the power of 10 is used, Beer’s law is written as follows.

\[ I(\lambda) = I_0(\lambda) 10^{-20(\lambda)CCL} \]

The mathematics given here is transparent to the operator and of little significance throughout the remainder of this document. However, it should be noted that when absorbances are given as numerical values, the logarithm to the base 10 has been used.

Although most FT-IR workers cite and discuss Beer’s law, it is not directly used. The absorption coefficient is generally not known. One implication of Beer’s law that is used is the concept of reciprocity. That is, if the concentration diminishes by a factor of 2 but the path length increases by a factor of 2, the measurement will yield the same results. This is not always true, and it is generally accepted that if the quantity \( aCCL \) becomes larger than about 0.1, the concept of reciprocity is no longer valid.

Beer’s law has been restated so that it includes many applications, and, as restated, the law has assumed several other names, as the Lambert-Beer law or the Bouguer-Lambert-Beer law, but these other names are not correctly used. Beer wrote the law for a purpose other than the way it is used today. When Beer published his original work in 1852, he was conducting experiments designed to measure the absorption of various materials that were then being used in the field of photometry. His entire endeavor was directed toward investigating the effects of the thickness of a material. He therefore did not write the law in terms of concentration, nor is there any evidence that he considered the effects of a changing concentration. The law as used today, with concentration as an explicit term in the exponent, seems to have first been published by B. Walter almost 40 years after Beer’s original work (Walter 1889).

### 2.6.2 The Interferogram

The primary data produced by an FT-IR instrument is the interferogram, and it is the piece of data that should be recorded. However, the mechanics of the FT-IR
instrument itself can alter the appearance of the interferogram, and this influence must be accounted for during data analysis. Two of these effects, truncation and phase shift, are discussed below.

2.6.2.1 Truncation

As discussed earlier, the interferogram is the intensity measured by the detector as a function of the position of the moving mirror. It contains all the information about the spectrum that is familiar to most operators. In actual operation, the mirror in the interferometer moves, at most, a few centimeters and stops and then returns to its original position. This finite movement truncates the interferogram at each end. It can be shown that this truncation actually limits the sharpness of the absorbing features that are of interest to the experimentalist. This is analogous to creating a square wave from a series of sine and cosine functions. There it is seen that the higher frequency components sharpen the edges of the square wave. The situation with the FT-IR data is identical. The information at the ends of the interferogram is really information about the high-frequency components. A simple truncation (stopping the mirror motion after a certain distance) behaves mathematically as though the interferogram were multiplied by what is called a boxcar function. That is, the interferogram is multiplied by a function that is 1 in a region from mirror position 1 to mirror position 2 and is zero elsewhere. The effect of this multiplication is to broaden the spectral line features. Truncation also causes a phenomenon called ringing in the wings of the spectral features. That is, the truncation of the interferogram adds oscillations into the wings of the spectral features. These unwanted features can be removed by applying an apodization function to the interferogram prior to the Fourier transform. There are several apodization functions that can be applied to the data, but an in-depth description is beyond the scope of this chapter. They are addressed briefly in Chapter 7. The point is that the operator should be aware of this effect and that some choices can be made during data analysis that will affect the shape and intensity of the spectral features.

2.6.2.2 Phase Shift

A second instrumental effect on the data that occurs is a shifting in the relative phase of the wavelength, which is caused by the optics and the electronics. There are two components for each of the wavelengths whose intensities are added to make up the interferogram. They are the magnitude of the intensity and the relative phase. The optics and the electronics cause slight phase shifts as the signals are processed, and these are frequency dependent. The shifts are normally accounted for when the Fourier transform is done, but no record of them is saved. Therefore, if the interferogram is not recorded, it cannot be retrieved by simply performing an inverse transform on the spectrum itself. It is primarily for this reason that the interferogram should be saved, even though it is somewhat more costly in disk space.
2.6.3 The Transform

The transform is performed on the interferogram by machine and therefore is done numerically. There are many algorithms to accomplish this, and which of these is used in the software provided with the commercially available instruments is not known. The mathematical basis for the transform is described below.

The complex motion of items such as vibrating strings or drumheads or other periodically varying quantities can always be described by a sum of sine and cosine terms known as a Fourier series. The frequencies in these terms are called the fundamental frequencies at which the item or quantity can vibrate. The actual motion is then a linear combination of these fundamental frequencies. When this summing of terms is done, it is said that a harmonic analysis has been performed on the original vibratory motion. A study of such analyses shows that there are related pairs of variables such as time and frequency or position and momentum. In an analogous manner, functions can be analyzed, but here the more general Fourier transform must be used. The Fourier series is used to describe a periodic function as an infinite sum of sine and cosine terms whose frequencies are multiples of some fundamental. The transform allows the analysis of nonperiodic functions as an integral (also a summation) over a continuous range of frequencies. In one of its forms that relates time and frequency, the Fourier transform \( F(\omega) \), a function of frequency, is related to \( G(t) \), a function of time, as is shown in Equation 2-5. In the present situation, the function \( G(t) \) is the interferogram produced by the system, and \( F(\omega) \) is called the single-beam spectrum. The \( i \) in the term \( G(t) \) is a dummy variable, but in this case it is really the position of the moving mirror from the center burst position.

\[
F(\omega) = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{\infty} G(t) e^{-i\omega t} dt
\]

(Eq. 2-5)

2.6.4 The Single-Beam Spectrum

Some of the literature in this field refers to the single-beam spectrum as the inverse transform of the interferogram because the instrument takes the transform of the incoming signal to start with. Mathematically, this merely changes the sign in the exponent of Equation 2-5, which means a phase shift in the sine and cosine terms. The term single-beam spectrum is a historical holdover from the time when spectroscopists used a double-beam instrument and determined the transmission directly from a ratiometer in the electronics of the instrument. The present-day FT-IR systems do not use a double-beam system, but some of the terminology remains.

The single-beam spectrum contains all the information about the absorbing species of interest. But most workers do not use this spectrum for any direct data analysis. In some systems, it is this spectrum that has been stored on disk. At 1-cm\(^{-1}\) resolution this spectrum takes about 35 kilobytes of memory, which is much less than the
interferogram (100 kilobytes). However, storage capacity of the computers available today makes this issue a minuscule one.

2.6.5 Data Analysis

The data analysis includes generating an absorption spectrum from the raw interferogram data, developing or obtaining the appropriate reference spectra, and then applying the chosen analytical method to determine the concentration of the target gases. The analytical methods and the procedures for generating an absorption spectrum from the interferogram and reference spectra of the target compounds are discussed below.

2.6.5.1 Generation of the Absorption Spectrum

As shown in Figure 2-8, the data analysis generally starts with the recording of the interferogram. Although in some cases the interferogram is analyzed directly to determine the concentration of the target gas, more commonly, the interferogram is automatically converted into a single-beam spectrum through the numeric process that is called a fast Fourier transform. A single-beam spectrum is generated and recorded for each sampling period. We call this spectrum the analytical, or field, spectrum. A background spectrum is generated by one of the methods described in Chapter 4. Then a transmission spectrum is obtained by dividing the field spectrum by the background. The absorption spectrum is obtained by taking the negative logarithm of the transmission spectrum. The absorption spectrum is used for all further data analysis.

2.6.5.2 Generation of the Reference Spectrum

A reference spectrum is usually generated by using a high concentration of gas in a relatively short cell. The cell is usually at least 1 m long, although multipass cells with longer path lengths are also used. A pure sample of gas mixed with an inert gas, such as nitrogen, is used. The concentration of gas used to generate the reference spectrum should yield a range of absorbance values that match as closely as possible those expected to be found in atmospheric measurements. The system can use a flowing stream of gas, but the total pressure should be around 1 atm. The process of producing a reference spectrum is then the same as outlined above.

The production of reference spectra is an exacting undertaking and requires great attention to the experimental details. It is not likely that most users of the FT-IR technique will prepare their own reference spectra. Reference spectra are currently available commercially. There is, however, no independent organization responsible for validating the accuracy of these spectral libraries.

2.6.5.3 Analytical Methods

After the reference spectra of the target gases are obtained, the appropriate wave number region for analysis must be
Analytical Spectrum
Generated from Each Sampling Period

Background Spectrum
Generated as Described in Chapter 4

Transmission Spectrum = \frac{\text{Analytical Spectrum}}{\text{Background Spectrum}}

Absorption Spectrum = -\log \text{[transmission]}

The absorption spectrum is used for all further data analysis.

Figure 2-8. Data Reduction Flow Chart.
selected. The selection should be based on an examination of reference spectra and the type of analytical method chosen. Two issues must be addressed to make this selection. Ideally, the gas should have a high absorption coefficient in the selected region, and the region should be free of absorption bands from interfering species. If interfering species are present they must be identified and accounted for in the analysis methods.

Once an appropriate wave number region is selected, data analysis can proceed. The concentration of the unknown gas can be determined in three general ways, as described below: the comparison method, scaled subtraction, and multicomponent analysis techniques. Each method uses a reference spectrum of the gas being investigated.

2.6.5.3.1 Comparison Technique

One method of determining the concentration is to measure the absorbance at a particular wave number and compare it with the absorbance of the reference spectrum at the same wave number. Then, if reciprocity holds (as implied by Beer’s law), the concentration is obtained as follows. The absorbance \( A \) is the product of \( \alpha \), the optical absorption coefficient, \( C \), the concentration of the gas and \( L \), the path length. Thus \( A = \alpha CL \) and the unknown concentration can be found from the following expression.

\[
\frac{A_{\text{ref}}}{A_{\text{unk}}} = \frac{C_{\text{ref}} L_{\text{ref}}}{C_{\text{unk}} L_{\text{unk}}}
\]

(Eq. 2-6)

Solving for the unknown concentration gives the following.

\[
C_{\text{unk}} = C_{\text{ref}} L_{\text{ref}} A_{\text{unk}} / L_{\text{unk}} A_{\text{ref}}
\]

(Eq. 2-7)

This concentration has the same units as the units of the reference concentration, which is prepared as described in Section 2.6.5.2.

2.6.5.3.2 Scaled Subtraction Technique

The scaled subtraction technique is similar in principle to the comparison technique. This technique is particularly useful if there are spectral features due to interfering species that overlap with those of the target compound. However, for scaled subtraction to be successful, either the target compound or the interfering species should have at least one unique absorption band. High-resolution data can be used to an advantage with this technique.

The scaled subtraction can be done as follows. Most software packages allow two spectra to be subtracted interactively. In this case the reference spectrum should be subtracted from the analytical spectrum until the absorption maximum of the band of interest is zero. Once the subtraction is completed the software reports a scaling factor. This factor can be multiplied by the concentration used to generate the reference spectrum to obtain the concentration of the target gas in the analytical spectrum. There is some operator skill involved in subtracting spectra interactively; therefore, some practice
in using this technique is recommended before the actual field spectra are measured.

2.6.5.3.3 Multicomponent Analysis Techniques

Multicomponent analysis techniques can be used to advantage when there are several target compounds to be analyzed for and there are several interfering species present. This is the case often encountered in open-path FT-IR monitoring, so some type of multicomponent analysis technique is generally the preferred method of analysis. There are several techniques that are used to perform multicomponent analyses of IR spectra. Multicomponent analysis techniques encompass a discipline unto themselves, and a complete discussion of the various techniques is beyond the scope of this document. The reader is referred to an excellent review by Haaland (1990). The most common multicomponent analysis method used in open-path FT-IR monitoring is based on a classical least squares (CLS) fitting algorithm. This is discussed below.

The CLS technique performs a linear regression by using the unknown and the reference spectra over a wave number region. The slope calculated in the regression is then used as a multiplier of the reference concentration to obtain the unknown. The ratio of the path lengths must also be accounted for. Thus if the slope is found to be 1 and the ratio of the reference path length to the path length used for the measurement is 1/10, then the unknown concentration is 1/10 of the reference concentration.

The process of using the linear regression is more suitable than either the comparison technique or the scaled subtraction technique because the shape of one spectrum is compared with the shape of the reference spectrum. If the correlation coefficient is also calculated, it gives a measure of this comparison. There is one problem with this technique that has generally been overlooked. That is that the range of selected wave numbers over which to perform the calculation can be too small. This choice of wave numbers should be made so as to encompass the maximum range of absorbance. To obtain meaningful statistical values for these quantities, the total noise should make up less than 5% of the range of the absorbance values.

2.7 References


Chapter 3
Initial Instrument Operation

SUMMARY

This chapter offers guidance and recommendations with respect to the initial tests that should be performed on the FT-IR system to verify that the instrument is set up to operate properly. Specific areas that are addressed include the following.

- The characteristic features of the single-beam spectrum
- The distance at which the detector is saturated and operating in a nonlinear fashion
- The return signal intensity as a function of distance
- The uniformity of the IR beam intensity
- The contribution of stray light to the total return signal
- The determination of the system noise
- The effect of water vapor concentration on the return signal intensity

3.1 Introduction and Overview

The assumption made for the discussion in this chapter is that the manufacturer has set up the FT-IR and it is running according to his specifications. Initially, the setup procedure for each field study should be the same, although certain procedural differences are dictated by specific data requirements.

Before putting the instrument into continuous monitoring service, the operator should conduct some initial tests and determine the following.

- The distance at which the detector is saturated and operating in a nonlinear fashion (Section 3.4)
- The return signal as a function of distance (Section 3.5)
- The stray light inside the instrument (Section 3.6)
- The uniformity of the beam intensity (Section 3.7)

The operator should become quite familiar with the single-beam intensity profile and with the gross features of the single-beam spectrum. Beyond that, the operator must start several control charts that will provide
long-term information about the return intensity and the noise levels. Some ancillary items such as water vapor concentration, ambient temperature, and ambient pressure should also be recorded.

This chapter includes a discussion of each of these items and also addresses the items that must be recorded so that adequate information concerning the long-term stability of the FT-IR can be obtained. In addition, some data are presented that have been recorded by FT-IR instruments in the past.

The tests outlined in this chapter should be performed before data are recorded with the FT-IR monitor. A failure to do so could result in the acquisition of erroneous data that could lead to inaccurate concentration measurements. Many of the tests involving the initial instrument setup are similar to those proposed for use in the routine quality assurance procedures presented in Chapter 8 of this document.

3.2 The Single-Beam Spectrum

Figure 3-1 shows a single-beam spectrum taken with 1-cm\(^{-1}\) resolution. The total scan time was 5 min and the total path length was 414 m. The vapor pressure of water during this measurement was 12 torr. There are several features in the spectrum that should be noted. First, the regions 1415 to 1815 cm\(^{-1}\) and 3547 to 3900 cm\(^{-1}\) are where the infrared energy in the beam is totally absorbed by water vapor. The operator will notice that, for a given path length, the width of the region for complete absorption varies as the amount water vapor in the atmosphere changes from one day to the next. Also the center of the region should become somewhat transparent as the path is made shorter.

The strong absorption in the 2234- to 2389-cm\(^{-1}\) region is due to carbon dioxide, and the atmosphere in this wave number region should remain opaque at all times, even when the instrument is used to monitor over short paths. The opaque regions should be flat, and they represent the baseline of the spectrum. Any deviation from zero in these regions indicates that something is wrong with the instrument operation. However, Figure 3-1 shows these regions to be elevated. This is due to stray light in the instrument, and these regions are marked "S" in the figure.

The operator should pay particular attention to the spectrum in the region around 600-700 cm\(^{-1}\). The spectrum below this wave number region should be flat and at the baseline. If the spectrum has an elevated baseline below the detector cutoff, in this example the 650-cm\(^{-1}\) region, the instrument may be operating in a nonlinear manner. If this is the case, the operator will see what seems like a dip appear as the retroreflector or source is brought closer to the FT-IR. An example of this is given in Figure 3-2 for a single-beam spectrum recorded at a 20-m path length.

When the path is sufficiently long (200 m and 10 torr H\(_2\)O) or the water vapor concentration is large, an absorption band should be noticeable at 2720 cm\(^{-1}\). This
Figure 3-1. Single-Beam Spectrum Along a 414-m Path. S indicates stray light.

peak is the Q-branch of deuterated water (HD0), and it is also possible to observe the P and the R branches.

The spectral region around 3000 cm\(^{-1}\) is also strongly absorbed by water vapor, although it is not opaque. The absorption features of methane are in this region. This is also the region of the C-H stretching frequencies. The atmosphere from 3500 to 3900 cm\(^{-1}\) is opaque, again because of water vapor. There is still some sensitivity and therefore an elevated signal return above 4200 cm\(^{-1}\), and this is the region where hydrogen fluoride is absorbing.

The return beam intensity at approximately 987, 2500, and 4400 cm\(^{-1}\) should be recorded so as to form a basic set of data about the instrument's operation. Along with this, the operator should record the path length. The total return signal is dependent on the path length and the amount of water vapor in the atmosphere. When
using the single-beam spectrum to gauge how well the instrument is functioning, the operator should try to select regions that are not greatly impacted by water vapor.

Figure 3-3 shows the region between 1000 and 1025 cm\(^{-1}\) enlarged and plotted in absorbance. The operator should notice that there are water vapor lines at 1010, 1014, and 1017 cm\(^{-1}\). These lines will be in every spectrum as long as the product of the water vapor concentration and the path length is large enough. The lines at 1010 and 1017 are actually doublets and cannot be resolved at 1-cm\(^{-1}\) resolution. The line at 1014.2 cm\(^{-1}\) is a singlet and can be used as a check for wave number shifts and resolution. A procedure for doing each of these is given in the subsections below.
Both wave number shifts and resolution changes indicate that something has changed in the instrument geometry, and if these occur they should be discussed with the instrument manufacturer. A subtle, apparent wave number shift can be observed if the atmospheric absorption line used for a shift determination is an unresolved doublet. In this case, if one line becomes more intense with respect to the other, the envelope peak will appear to be shifted.

3.2.1 Wave Number Shift

To determine whether wave number shifts have occurred, the operator should have an absorption spectrum that contains the water vapor line at 1014.2 cm\(^{-1}\) and one for which there is no shift present. The HITRAN database for water vapor is used in this document as a guide (University of South Florida 1993), and it positions this water vapor line at 1014.2 cm\(^{-1}\).

For any particular instrument, the line assignment may be slightly different (\(\pm 0.2 \text{ cm}^{-1}\)) because of the instrument's geometry, but it should not shift in time. However that may be, it is the responsibility of the operator to determine precisely where the water line is and whether shifts occur with time. The operator may also choose to determine wave number shifts by using the HDO lines in the 2720-cm\(^{-1}\) region. This measurement is somewhat more sensitive to shifts in the higher wave number (shorter wavelength) region.

In principle, any absorption or single beam spectrum can be used as a guide to determine wave number shifts. There are two methods available for determining shifts. The first is simply to compare the positions of
the peaks of the two spectra on the computer monitor. The second is to subtract the second spectrum from the first and study the result. The second spectrum should be normalized to the first by a simple multiplication before the subtraction is done, or the subtraction can be done interactively. After subtraction, wave number shifts will result in a curve that appears to be the first derivative of the line shape. The wave number where zero amplitude occurs will be shifted from the original peak wave number by an amount that is proportional to the wave number shift. This is shown schematically in Figure 3-4A.

3.2.2 Change in Resolution

The other possible change that can occur in the spectra as time passes is a resolution change in the instrument. If a change in the resolution has occurred but there is no peak shift, the result will appear to have the shape of an 'M' or a 'W', depending on which spectrum has been recorded with the largest amount of water vapor. This is shown schematically in Figure 3-4B. If there are no changes in the line, then the result of subtraction will be random noise.

3.3 Distance to Saturation

One of the early pieces of information to obtain with an FT-IR monitor is the path length at which the detector is saturated. As discussed in Section 3.2, this is easily noticed by a negative dip in the single-beam return in the 650-750-cm\(^{-1}\) region below the detector cutoff. As the retroreflector or the light source is brought closer to the detector, this dip will appear. This response is opposite to the response that is due to an absorption feature in the spectrum. This distance is important because it represents the minimum path length over which it is possible to operate without making changes to the instrument. In the monostatic case, it is possible to rotate the retroreflector to lower the return intensity. If necessary, it is possible to lower the intensity of the FT-IR instruments by simply using a fine wire mesh
screen to cover the aperture. A plastic screen should not be used because plastics have absorption features in the infrared.

To measure this distance, simply move the light source or the retroreflector away from the transmitting telescope until the negative dip just disappears from the single-beam spectrum. This distance should be recorded as the minimum working distance available without making instrument changes.

3.4 Return Intensity as a Function of Distance

Some attempt is made to collimate the infrared beam before it is transmitted along the path. It is, however, impossible to completely collimate the beam because of the size of the light source. Therefore, most beams either are always diverging as they traverse the path or become diverging at some point along the path. Once the beam is bigger than the retroreflector or the receiving telescope, the return signal should diminish as the square of the distance. That is, beyond some distance the signal is reduced by a factor of 4 when the path length is doubled. The reason that this return signal versus distance determination is necessary is twofold. The first is that the commercially available instruments that use the monostatic configuration both have a stray light signal when the telescope is blocked. The return beam should never be allowed to approach this signal. The second is that at some distance the system noise will become an appreciable part of the signal, and this represents the maximum usable distance.

To determine the return signal as a function of distance, the operator must start with the retroreflector or the light source at the minimum working distance as determined above. Then, the operator should move the light source or the retroreflector back by some distance and record the signal. This process should be continued until the signal level reaches the noise level or just levels off. It should be noted that the leveling-off effect can also be caused by the return signal's reaching the stray light level. These data should then be plotted and used for quality assurance/quality control purposes.

3.5 Determination of the Stray Light Signal

The stray light in the instrument can be measured without regard to the distance to the light source or the retroreflector. It is not expected that instruments using the bistatic configuration will have any measurable stray light, but a one-time check is appropriate. To measure the stray light, the operator must block the receiving telescope while the signal is being recorded. It is important to use an appropriate blocking material to do this. No surface that can reflect any of the infrared energy back to the instrument can be used, nor any material that is transparent. The best blocking material is a piece of black cloth such as is used in the construction of a photographic film changing bag. For systems that transmit the beam through the interferometer before transmitting it along the path, the beam can simply be slewed away from the retroreflector. This return signal should be recorded and then plotted on the graph as return signal versus distance.
discussed above. A record of the stray light spectrum should be made and compared to the single-beam spectrum recorded at the selected working distance.

Stray light inside the instrument can also be caused by strong sources of IR energy that are in the field of view of the instrument. For example, it is possible to have the sun in the instrument’s field of view during sunrise and sunset. This will probably give rise to an unwanted signal that actually comes from reflections inside the instrument.

The stray light actually causes an error in the determination of the gas concentrations and must be subtracted from the data spectra before processing. Thus it has to be recorded at every monitoring session and periodically in the case of a permanent installation. The effect of stray light on photometric accuracy is illustrated by the absorption feature shown in Figure 3-5.

The absorption line in Figure 3-5A has a transmission of 0.5 at the peak. The baseline (100% transmission) has unity amplitude.

![Graph A](image)

![Graph B](image)

**Figure 3-5.** Effect of Stray Light. A. Spectral feature without stray light. B. Spectral feature with stray light.
Now suppose that stray light exists in the instrument in the amount of 10% of the original return signal. This means that the baseline goes to an amplitude of 1.1, but the absorption feature goes to an amplitude of 0.6, as shown in Figure 3-5B. Thus the new transmission of the absorption is 0.6/1.1, and this is not equal to 0.5 but to 0.5454, or is in error by about 9.2%.

Mathematically, this is just verifying the fact that \( A/B \neq (A + C)/(B + C) \). It might be questioned whether the effect of this stray light is offset by the effect of the air in the interferometer enclosure, which most likely contains the pollutant gas also.

However, the transmission due to the gas in the cell is increased from that along the path in the ratio of the path lengths. Thus, for our data taken at Research Triangle Park, NC, along a 414-m path, the ratio of the path lengths is at most 1/414, and therefore the additional absorption is negligible. A more in-depth analysis of the problems introduced by stray light indicates that actual line shape distortions may take place. It is also quite likely that the simple subtraction of stray light as suggested above will not remove all the error incurred. Therefore, all efforts should be made to at least minimize the amount of stray light reaching the detector.

3.6 Determination of the Random Noise of the System

The random noise of the system is determined from an absorption spectrum made from two single-beam spectra taken sequentially. These spectra are to be taken under the same operating conditions as will be used for the acquisition of data spectra. That is, the same acquisition time and path length should be used for the noise determination as will be used for data acquisition. There should be no time allowed to elapse between the acquisition of the two spectra. This determination will be somewhat dependent on the water vapor concentration in the atmosphere, so the water vapor concentration should also be determined.

The use of the word noise suggests a random signal that is primarily produced by the system electronics. When measurements are taken in the open air, this may not exactly be the case. If the period of data acquisition for the two spectra is long (because of a large number of scans, for example), then atmospheric effects may contribute to the measurement. For that reason, the wave number regions that are used for the determination should be carefully chosen. Three regions are suggested below, but the low wave number region may not be suitable in all situations because of the presence of gases other than water vapor.

The actual wave number range over which the noise should be calculated will vary with the resolution used. Statistically, it can be shown that about 98 data points is an optimum number (Mark and Workman, 1991). For 1-cm\(^{-1}\) resolution, this means that data should be taken over a 50-cm\(^{-1}\) region.

Once the two spectra have been acquired, an absorption spectrum should be
made by using one of the two spectra as a reference or background spectrum. Which one is used for the background is not important. The noise is then determined in the three regions 958-1008 cm\(^{-1}\), 2480-2530 cm\(^{-1}\), and 4380-4430 cm\(^{-1}\).

There are several ways that are described in statistics texts to determine the noise. We will specify the use of the root mean square (RMS) deviation as the appropriate measure of the noise. The first step is to perform a linear regression over the wave number region and determine the slope and the intercept of the line. At each wave number, the next step is then to calculate the difference between the calculated line and the actual ordinate value. The squares of these differences are then used to calculate the RMS deviation as is described below.

To calculate the slope and the intercept from a linear regression, use the following expressions.

\[
\text{NUMERATOR} = N \sum X_i Y_i - \sum X_i \sum Y_i
\]

\[
\text{DENOMINATOR} = N \sum (X_i)^2 - \sum X_i \sum X_i
\]

\[
\text{SLOPE} = \frac{\text{NUMERATOR}}{\text{DENOMINATOR}}
\]

\[
\text{INTERCEPT} = \frac{1}{N} (\sum Y_i - \text{SLOPE} \times \sum X_i)
\]

In this case the \(X_i\) values are the wave numbers and the \(Y_i\) values are the absorbance values at the particular wave numbers.

The differences are then found as \(D_i = Y_i - \gamma_i\), where now the \(Y_i\) values are calculated from the line by using the expression \(Y_i = \text{slope} \times X_i - \text{intercept}\).

Once this is accomplished the RMS deviation is determined with the following expression.

\[
\text{RMSDEV} = \left( \frac{\sum D_i^2}{N-2} \right)^{1/2}
\]

Mathematical representations of the RMS deviation vary in what the denominator is. Quite often the denominator is written simply as \(N\). The term \(N-2\) is used here because the slope and the intercept are calculated from the data. This reduces the degrees of freedom by 2 and hence the \(N-2\) (see, e.g., Mark and Workman 1991).

Some results of these measurements are shown in Figure 3-6. The data shown in the figure were taken over the region 980-1020 cm\(^{-1}\) in order to include the water vapor peaks at 1014 cm\(^{-1}\). To reduce the effect of water vapor to a minimum it is possible to create two spectra by using an artificial background, subtract the water vapor of one from the other, and then make the noise determination.
The return-beam intensity is a function of the absorption due to water vapor in the atmosphere. It is therefore a function not only of the path length but also the water vapor concentration in the atmosphere. It must be clear to the operator that relative humidity is not important in this case and has no relevance to the FT-IR data, and it is actually the water vapor partial pressure in torr that must be used. However, over a period of one day, the water vapor concentration may not change very much, so acquiring a set of data over a range of water vapor concentrations will take some time. Some thought must also be given to the problem of measuring the water vapor concentration when the temperature is below freezing. The Smithsonian psychrometric tables give data to an ambient temperature of
5 °F, but it is not clear that a simple sling psychrometer should be used to make the wet and dry bulb measurements.

The measurements for water vapor can be made at any place along the path. The operator should note, however, that some investigators feel that the concentration of water vapor along the path actually changes. We have made some measurements with a sling psychrometer and have not seen any appreciable changes along the path. We have seen rather subtle changes in the absorption due to water vapor from the spectra themselves. This is most easily noticed if the subtraction technique is used. For example, we acquired a set of data taken at 1-min intervals. After the spectra were converted to absorption spectra, the first spectrum was subtracted from the others in order. The residual water in the spectra indicated that minor but noticeable changes take place minute by minute.

3.8 References

University of South Florida. 1993. USF HITRAN-PC. University of South Florida, Tampa, FL

Chapter 4  
Background Spectra

SUMMARY

The topics and specific points of emphasis discussed in this chapter include the following.

- The generation of the transmission spectrum and the absorption spectrum
- The need for a background spectrum and the difficulties in obtaining one that is adequate
- The methods for generating a suitable background spectrum
  - Synthetic backgrounds
  - Upwind backgrounds
  - Short-path backgrounds
  - Averaged backgrounds
- Specific field guidance for measuring the background spectrum

4.1 Introduction and Overview

In current use, long-path, open path FT-IR data are obtained from single-beam measurements. That is, there is no reference, or background, spectrum taken simultaneously with the sample spectrum to null the spectral features due to the characteristics of the source, beam splitter, detector, and interfering species in the atmosphere. To remove these background spectral features, the single-beam field spectrum is divided by a single-beam background spectrum, or \( I_0 \) spectrum. This operation (illustrated in Figure 2-8) generates a transmission spectrum. According to Beer's law, the absorption spectrum is then calculated by taking the negative logarithm of the transmission spectrum. It is the absorption spectrum that is used for data analysis.

Ideally, the background spectrum is collected under the same experimental conditions as those for the sample spectrum, but without the target gas or gases present. However, in the field it is not possible to obtain the \( I_0 \) spectrum directly because the target gas cannot be easily removed from the atmosphere. This chapter presents a discussion of the problems associated with obtaining the \( I_0 \) spectrum and the methods that are used to generate a background spectrum.

There are currently four methods for obtaining \( I_0 \): synthetic, upwind, short-path, and averaged background spectra. Synthetic
background spectra can be generated by selecting data points along a single-beam spectrum and then calculating the high-order polynomial function that best fits the selected points or by repeatedly subtracting the spectral features due to interfering species from a single-beam spectrum. Both of these methods can introduce distortions or spurious features in the actual intensity profile. Therefore, much care must be taken when generating synthetic background spectra.

For short-term monitoring efforts, the FT-IR path is generally chosen to be perpendicular to the wind field. If the area of the emission source is relatively small, a background spectrum can be acquired along a path that is upwind from the source. However, it is difficult to make this type of measurement frequently if the FT-IR system has to be moved from one side of the source area to another. Also, errors may be introduced in the measurements if the water vapor in the atmosphere changes significantly between the times that the background spectrum and the sample spectra are acquired.

Another option for obtaining the $I_0$ spectrum is to bring the retroreflector or external IR source close to the receiving telescope. This effectively eliminates the absorption caused by the target gases and records a true instrument background. One problem with this method is that the detector can be saturated at short paths because too much IR radiation is incident on the detector element.

When the experimental conditions are fairly constant over a measurement period, it is possible to average several backgrounds that have been taken over this time. This average $I_0$ can then be used for the entire data set for that period. However, most of the time, the experimental conditions are not constant enough to perform this type of long-term averaging. For example, the concentrations of water vapor, $CO_2$, and gases emanating from other sources are constantly changing.

The change in water vapor concentration must be considered the biggest potential source of error in the background measurement. An accurate record of the partial pressure of water vapor should be maintained. These data should be taken continuously, or at least every two hours during quiet days and every half hour during times when fronts are passing.

Acquisition of the $I_0$ spectrum represents one of the more difficult tasks in FT-IR long-path, open-path monitoring. Currently, there is not a universal method for obtaining a satisfactory background spectrum. The method chosen to obtain $I_0$ must be determined on a site-by-site basis. This chapter includes a rationale for the use of an $I_0$ spectrum, as well as advice on the appropriate techniques for generating it. These techniques are each discussed below.
4.2 Synthetic Background Spectra

The software that is supplied with the commercially available instruments has a routine that allows a synthetic spectrum to be generated. This is accomplished by selecting data points along some original single-beam spectra and then calculating a high-order polynomial function that best fits the selected points. Thus, once a single-beam spectrum is produced, it can be used to generate a synthetic $I_0$. Synthetic $I_0$ spectra can be made that cover only selected wave number regions, or they can be made to cover the entire wave number region that the FT-IR uses. An example synthetic spectrum is shown in Figure 4-1.

Some care must be used when synthetic $I_0$ spectra are generated so that distortions are not introduced into the intensity function. For this reason, when data points are selected, they should never be selected at the peaks or even within an absorbing feature. The final curve that is produced must be a smooth function without artificial dips and peaks and must follow the baseline of the single-beam spectrum from which it is made.

![Figure 4-1. Synthetic $I_0$ Spectrum. The peak at 1110 cm$^{-1}$ has intentionally been left in as a fiducial point.](image)

4-3
It is essentially impossible to construct a synthetic spectrum in the wave number regions where water vapor and carbon dioxide absorb strongly. The individual lines are overlapping so that it is very difficult to judge where the background curve should be set, and in much of the region there is almost no energy being returned to the detector. Even at the shortest path length possible, there are still portions of the spectrum that are completely absorbed by the water vapor and the carbon dioxide. Because of this strong absorption, these regions of the spectrum are not used in the data analysis.

4.3 Upwind Background Spectra

For short-term monitoring efforts, the path is generally set out so as to be perpendicular to the wind field. If need be, the operator can change the orientation of the path so that this geometry is maintained. If the area of the source is relatively small and its upwind side is accessible, an upwind $I_0$ spectrum can be acquired. A usable background spectrum can also be acquired by taking data along the side of source. (See Figure 4-2.) As long as the wind is not blowing across the source area and transporting the emissions across the path used for $I_0$, these spectra should be satisfactory.

Another technique for acquiring an upwind background spectrum is to wait until the wind shifts so that the path is along an upwind side of the source. This works well for isolated sources, but if there are other places emitting chemicals, then this method can lead to errors in identifying compounds and in quantifying just what is coming from the source under study.

![Figure 4-2. A Possible Configuration for $I_0$ Spectrum Acquisition.](image)

There are some advantages to taking true upwind background spectra this way. First, it is likely that sources are not isolated and the chemical species of interest are emanating from several places in the area. The compounds entering the area being investigated are thus included in the upwind background spectrum. If the configuration can be set up so the side of the source area can be used, a second retroreflector or IR source can be used, and the $I_0$ spectrum can be taken frequently and without having to transport the entire system from one place to another.

There are also some disadvantages to this procedure. It is difficult to take upwind and downwind spectra frequently if the system has to be moved from one side of the source area to another. Generally, this type of spectrum is taken once at the beginning of
the monitoring period and once at the end (each day).

4.4 Short-Path Background Spectra

Another possible technique for obtaining the $I_0$ spectrum is to bring the retroreflector or external IR source close to the receiving telescope. This effectively eliminates the absorption caused by the compounds of interest and records a true background. This background is called the short-path $I_0$. If two retroreflectors are available, this task is fairly easy to perform. The FT-IR monitor can be pointed first to one retroreflector and then the other quite easily with some regularity.

Figure 4-3 presents a procedure for acquiring a short-path background spectrum. The recommendation for the frequency of repeating this procedure is that a background should be considered valid for no longer than one day or until there is a major change in the operational parameters. Although a short-path background spectrum may be valid for an extended period, it should be revalidated on a daily basis. Necessary checks for a short-path background spectrum include the following.

- Comparison of the curvature of the baseline in the short-path spectrum with the curvature of the baseline in the field spectra
- Inspection of the spectrum for wavelength shifts and resolution changes, as discussed in Section 3.2

One difficulty with this procedure is deciding on an appropriate distance for placing the retroreflector or external IR source. One difficulty is that the detector can easily be saturated when the path length is too short. However, if the reflector is placed far enough away to overcome this problem, then the absorption may become large enough to be detrimental. Thus, this distance must be determined for each instrument at least once. When the detector is saturated, the signal seems to drop below zero at the low-wave-number end of the single-beam spectrum, and the response of the detector is nonlinear. The retroreflector or external IR source must be far enough away so that this dip in the signal strength disappears. One way to accomplish this is to reduce the light intensity by rotating the retroreflector or using wire mesh screens to attenuate the signal. (See Section 3.3.)

A second difficulty in monostatic systems is that the retroreflector will subtend different angles when it is at different distances. The retroreflector may be the actual optical field stop of the instrument, and changing the distance can cause distortions in the spectrum. When the distance is increased, the retroreflector subtends smaller angles, and the instrument uses different cones of light. This problem can be overcome by placing a field stop in the instrument that uses a smaller field of view than the smallest anticipated from the retroreflector. However, that is a job for the manufacturer because stops cannot be placed just anywhere in the optical train without causing other problems.
ACQUIRING A SHORT-PATH BACKGROUND SPECTRUM

1. Calculate the required path length for each gas that is being measured.
   a. Calculate the absorption coefficient of the gas in the wave number region that is being used for
      analysis. From the reference spectrum, obtain the product of the concentration and the path
      length used to generate the reference data. Then measure the absorbance of the peak that is
      being used for analysis. The absorption coefficient is given by the following.
      \[
      \alpha = \frac{A}{CL}
      \]
   b. Estimate the maximum anticipated concentration of the target gas. Express this in the same
      units as the concentration of the reference has been expressed (generally, ppm).
   c. Set the absorbance to the number that has been experimentally determined from the RMS noise
      as an estimate of the lowest possible detection limit for the FT-IR system, for example, \(10^{-6}\),
      and use the expression
      \[
      L = \frac{A}{\alpha C}
      \]
      to find the path length to be used for the background spectrum.
   d. A sample calculation with the toluene peak at 1031.6 cm\(^{-1}\) is given below.
      \[
      \alpha = \frac{A}{CL}
      \]
      Read \(A\) at 1031.6 as 0.0203.
      The \(CL\) product = 496 ppm-m
      \[
      \alpha = \frac{0.0203}{496} = 4.1 \times 10^{-5}
      \]
      Estimate the maximum concentration to be 0.050 ppm.
      Calculate \(L\):
      \[
      L = \frac{A}{\alpha C} = 10^{-4}/(0.000041 \times 0.050)
      \]
      or
      \[
      L = 48.8 \text{ m}
      \]

2. Repeat Step 1 for each target gas and set up the retroreflector or light source at the minimum
   calculated distance.

3. Take a spectrum at the same resolution as will be used to take data. So that the noise in the data
   spectrum will be the predominant source of noise, take the background spectrum for at least 4 times
   the number of scans used for the data spectrum. (This judgment should be based on the time
   involved with taking a spectrum.)

4. Use this spectrum for the background spectrum.

Figure 4-3. Procedure for Acquiring a Short-Path Background Spectrum.
4.5 Averaged Background Spectra

When the experimental conditions are fairly constant over a measurement period, it is possible to average several background spectra that have been taken over this time. This average $I_0$ can then be used for the entire data set for that period. Because all the individual spectra making up the average should have the same noise and there should be no other errors, the final error in this average background should be smaller by a factor of $\sqrt{N}$. Here, $N$ is the number of spectra in the average.

However, most of the time the experimental conditions are not constant enough to perform the averaging. The water vapor concentration is changing most of the time, and so is the concentration of carbon dioxide. If other sources are in the area, the concentrations of the gases emanating from them are not likely to be constant. If any of these gases are also being monitored, the use of an average $I_0$ will not give true absorption spectra for the entire monitoring period.

Currently, no published data seem to be available that have been taken over periods of more than a few weeks. Therefore it is not known whether $I_0$ spectra can be taken over extended periods, sorted according to the similarity of conditions, and then averaged. If this could be done, it might be possible to acquire a set of $I_0$ spectra that are universal. The experimenter could use these as he uses the library of pure compound spectra. Thus, with 10 torr of water vapor and a path length of 200 m, background VII might be used, etc.

Acquisition of the $I_0$ spectrum represents one of the more difficult tasks associated with using an FT-IR spectrometer. Most of the data that have been published until now has come from short-term monitoring programs. The $I_0$ spectra that have been used have been taken at various times during the days of operation. These spectra have then been used for the next series of atmospheric spectra to be analyzed until the next background is taken. Although shifts in the wind direction have been used to determine the need for a new background, changes in water vapor concentration have not. Little information has been published on the stability of the instrument itself or on the effect of taking a new background after some instability develops. There is some evidence that has been derived from bench-top laboratory work that indicates the $I_0$ should be taken as often as every other atmospheric spectrum. There is also some evidence that a single $I_0$ can be used over a long period of time with no detrimental effects. Neither of these observations has been corroborated by any in-depth study of the background spectra.

The work performed at Research Triangle Park, NC, indicates that a synthetic background ($I_0$) spectrum can be used for extended periods with some care. After one year of operation, the $I_0$ spectrum had to be changed four times. Three times were caused by instrument component changes, and only one time did the atmospheric conditions require a change in $I_0$. That time the water vapor concentration changed from 23 torr to 9 torr in a 15-min period.
Thus, much work is yet to be done in establishing the appropriate manner and frequency for acquiring the \( I_0 \) data.

4.6 Why Use a Background

The primary data that is produced by an FT-IR is the interferogram. It contains all the information that is required to obtain the concentrations of the gases that the experimenter wants. However, the information in the interferogram is in a form that is somewhat cumbersome to most people that are familiar with spectroscopy. In addition, the primary physical law that governs the analysis of the data is Beer's law, and this is defined in terms of the more conventional spectra that are familiar to most people. Thus, with few exceptions, the interferogram is converted to a single-beam spectrum via a Fourier transform and divided by a background, or \( I_0 \), spectrum to get a transmission spectrum. This is then converted to an absorption spectrum by finding the negative logarithm of the transmission spectrum. At this point, the absorption spectrum is compared with the absorption spectrum (after it has undergone the same mathematical processing) of a pure gas to obtain the concentration.

All this mathematical processing is performed by computer and therefore is done numerically with whatever algorithms were available at the time the analysis software was written. A relevant question is whether all the processing is necessary, or if it is being done merely because it has always been done that way.

It is likely that most workers would essentially refuse to work with the interferogram directly because spectra have historically been used. However, this does not seem to be true of the single-beam spectrum. The single-beam spectrum also contains all the information about the concentrations, and it looks like a normal spectrum. The water vapor and the carbon dioxide absorption peaks are readily discernable, and any absorption due to other gases should also be discernable. Thus, for identification purposes, using the single-beam spectrum should not be a problem.

The problem seems to arise when quantitation is required. If the reference spectrum were also available in terms of a single-beam spectrum, a direct comparison could be made between the data spectrum and the reference spectrum, but only if the intensity levels of each were known on some absolute basis. Beer's law gives no hint of how the data are to be analyzed in the absence of an \( I_0 \) spectrum. It is true that the single-beam spectrum is recorded with some intensity level for the ordinate. But unless it is put on an absolute basis, the single-beam spectrum alone is not a sufficient piece of information to determine the transmission through the atmosphere.

Currently, there does not seem to be a satisfactory way to use the single-beam spectrum alone for the final analysis.
4.7 General Advice About Background Spectra

All of the currently used methods for generating a background spectrum are fraught with difficulties. No one method is generally accepted as the best method for acquiring a background. It is crucial for the operator to be aware of the importance of this spectrum and of two criteria for a valid background.

1. The background cannot contain any absorption features due to the target gas or gases.

2. The background spectrum must be valid for the time period over which it is used.

Although there seems to be agreement that the first point above is a requirement, no such consensus has been reached for the second. The time periods over which single backgrounds are used by various workers vary from a day to several months. However, one point has become clear. Whenever any optical component (light source, mirror, window, etc.) is changed in the instrument, a new background must be acquired.

There are few guidelines as to what represents a valid background spectrum for the production of accurate data. One point is that the curvature of the baseline (maximum intensity) must be quite close to the curvature of the baseline of the field spectra.

There are two primary choices for the background that is to be used with a specific data set. If the absorbing peaks are narrow, as they are for methane or hydrogen fluoride, it is possible to construct a synthetic background for the analysis. But for broad absorbing features like those exhibited by acetone, this is difficult. With broad features, even small changes in the curvature of the baseline can produce large errors. In general, the operator is advised to use a synthetic background whenever possible. Taking a short-path background should be considered only when the absorption feature of the target gas is very broad. A final reason to consider the use of a synthetic background is that it is essentially the only background that allows actual atmospheric concentrations to be determined.
Chapter 5
Water Vapor Spectra

SUMMARY

Specific topics that are addressed in this chapter are the following.

- Selection of spectra that can be used for generating a water vapor reference spectrum
- Creation of the water vapor reference spectrum itself
- Subtraction from the water vapor spectrum all absorption features of CO₂, N₂O, CH₄, CO, and the so-called pollutant gases
- Example water vapor spectra for methane and ozone, selected because they present different problems to the operator

5.1 Introduction and Overview

Water vapor absorption lines are present in all regions of the mid-IR wavelength region, as was shown in Figure 3-1. The water vapor spectrum interferes with the spectrum of almost every volatile organic compound in the atmosphere. Because of this, the absorption features of water vapor have to be accounted for during the analysis of field spectra.

Some amount of the water vapor absorption is accounted for if there is water vapor absorption in the background spectrum, as described in the previous chapter. However, when a synthetic background is used, all the water will still be in the field spectrum, and some residual amount will be there when other backgrounds are used. It is possible to account for the water vapor by considering it as an interfering species in the analysis package. The software commercially available for performing a classical least squares analysis allows the operator to choose interfering gas species that are present in the wave number region of interest. To do this, however, a water vapor spectrum must be available. Although there are water vapor spectra available commercially, they are not suitable for use in this application because of line shape differences and other small instrumental effects, as well as insufficient path length.

This chapter explains how to use field spectra on site to produce a spectrum of pure water that can subsequently be used in the analysis of field spectra. Examples of water vapor spectra that can be used for the analysis of methane and ozone are discussed, as these gases represent, perhaps, the
extreme challenges that the operator will encounter.

5.2 Water Vapor Spectra Considerations

Any single-beam spectrum that exhibits a sufficient amount of water vapor absorption in the wave number region of interest can be used for the production of a water vapor reference spectrum. Spectra taken at short path lengths or during very dry periods may not be satisfactory. At Research Triangle Park, NC, we have seen the water vapor partial pressure change from a low of less than 1 torr in the winter to a high of 28 torr during the summer. Changes in the water vapor concentration of this magnitude, along with any instrument changes, may require that a new water vapor spectrum be produced. It is the responsibility of the operator to determine when the water vapor spectrum has to be remade, and no hard and fast rules on the frequency for creating a new spectrum are presently available. If the error bars of the analysis increase from one data set to another, a first step in determining the cause is to compare the water vapor reference spectrum with the water vapor in the field spectra.

The primary concern for the production of a water vapor spectrum is that the final result must not contain any of the target gas. If the water vapor spectrum does contain even a small amount of a target gas, the analysis will be in error by that amount. The ease with which the absorption features of the target gas can be removed from the water vapor reference spectrum is dependent on many instrumental factors, and the process can be quite time-consuming. The removal of the target gas absorption is done by subtraction and in some instances requires great attention to detail.

5.3 General Process for the Production of a Water Vapor Spectrum

The general procedure that is to be followed to produce a water vapor reference spectrum is given below.

1. Select two single-beam spectra that will be combined into a water vapor absorption spectrum.

2. Use one of the spectra to create a synthetic spectrum that is to be used as the background.

3. Create the absorption spectrum that is to be used as the water vapor reference spectrum.

4. Subtract any absorption features that are known to be present from the target gas.

5. Analyze a number of the field spectra for the target gas, using the water vapor reference spectrum as an interfering species.

6. Determine if any of the target gas absorption remains in the water vapor spectrum.

7. Repeat Steps 4-6 until you are sure that there is no absorbance due to the target gases remaining.
A word of caution is necessary here. If analysis is to be done for more than one gas, the synthetic background in Step 2 should be created for all the gases of interest at the same time. Otherwise, when the absorption spectrum is created, some or all of the water vapor absorbance will disappear. If this occurs the process has to be started over.

5.3.1 Selection of Spectra

The spectra selected in Step 1 that are to be used to produce a water vapor reference spectrum should be taken at the same resolution as the field spectra. The water vapor spectrum should have a signal-to-noise ratio that is the same as or better than the field spectra. If possible, the water vapor concentration for these two spectra should be representative of the water vapor concentration in the field spectra to be analyzed. Curvatures and any other special features of the baselines of these spectra should be the same as those of the field spectra baselines over the wave number regions of interest. If both of these spectra are closely spaced in time, they should contain approximately the same amount of the target gas. The operator should select spectra at times when a minimum of the target gas is expected to be present. However, the operator should be aware that the target gas will be present in both spectra and should consider the ramifications of that fact.

5.3.2 Generation of Synthetic Background

Either of the two spectra selected above (Step 1) can be used to create a synthetic background that is then used to create an absorption spectrum. The synthetic background must be created over the same wave number region as will be used for the final analysis. The wave number region can be larger than that used for analysis, but it cannot be smaller. If the two spectra selected above are closely spaced in time they will probably both contain approximately the same amount of the target gas. This will certainly be the case with methane and nitrous oxide regardless of the time selected, but any set of spectra for ozone may have widely varying absorption due to ozone. Again, which one of the spectra is used for the synthetic background generation is arbitrary. Once the synthetic background has been prepared, it should be stored with an appropriate name.

5.3.3 Generation of the Absorption Spectrum

Use the remaining spectrum (after Step 2) as the data spectrum and the newly created synthetic background to create an absorption spectrum (Step 3). This is done exactly as all the other absorption spectra are created. (See Figure 2-8.) At this point it is not likely that the baseline of the absorption spectrum is at zero absorbance. To make the baseline zero, measure the height of the baseline above zero and simply subtract that amount from the spectrum. This completes the process of creating an absorption
spectrum, and it should be saved with an appropriate name.

5.3.4 Subtraction of the Target Gas

This step (Step 4) is really an iterative process. The newly created water vapor spectrum must be used in the analysis of other spectra. The results of these analyses must be examined in an attempt to determine whether any of the target gas remains. If some remains, it must be subtracted from the water vapor spectrum. The process must then be repeated until the operator is sure that no target gas absorbance remains.

In order to analyze the water vapor spectrum, the operator must process several data spectra and use the newly created water vapor spectrum as an interfering species spectrum. The field spectra to be used should be chosen so that the target gas is a minimum. If the analysis shows the target gas to go through zero and actually become negative, then the water vapor spectrum still contains an absorbance due to the target gas. The negative concentration should be subtracted from the water vapor reference spectrum. Some gas concentrations will not go to zero at any time but will reach a minimum. This minimum can be set to zero in some instances, but it should at least be set to the minimum that the gas is known to achieve from other sources.

The actual subtraction can be done in two ways. The first way is to use the reference spectrum of the target gas and create a spectrum that represents the amount of the target gas to be subtracted. This is done by multiplying the reference spectrum by an appropriate factor. This factor can be calculated by using the concentration path length product used in the reference gas and the path length that was used for data acquisition.

The second way is to do the subtraction interactively with the software. In this way, the operator can see the results of the subtraction directly and has a little more control of the process. Both procedures require some practice, and the operator must be aware that his first attempt may not be satisfactory.

5.4 Methane and Ozone Examples

Figure 5-1 shows the portion of a single-beam spectrum over which methane absorbs. The methane concentrations at Research Triangle Park are generally measured at about 2.5 ppm. We have seen methane concentrations as high as about 6 ppm in this area. The spectrum in Figure 5-1 actually contains water vapor and methane, although the methane is not very noticeable. Figure 5-2 has superimposed on it the synthetic background that will be used to manufacture an absorption spectrum. The synthetic background has been raised slightly above the single beam spectrum for clarity.

Figure 5-3 shows the absorption spectrum that has been made from the two spectra shown in Figure 5-2. Also shown in this figure is the methane reference spectrum. The task is now to subtract out the methane
Figure 5-1. The Portion of a Single-Beam Spectrum Over Which Methane Absorbs.

Figure 5-2. Methane Region with Synthetic Background Spectrum Superimposed.
that is present in this absorption spectrum. There are two absorption peaks in the methane spectrum that are virtually not impacted by water vapor. They are at 2916.7 and 2926.8 cm$^{-1}$. The methane concentration can be calculated from the peak height of these two peaks. The concentration path length product of the reference spectrum is 81 ppm-m. The field spectrum was taken at a path length of 414 m, so the reference spectrum represents a concentration of 81/414 = 196 ppb. The measured peak height of the 2926.8-cm$^{-1}$ line is 0.00771 absorbance units for the reference spectrum and 0.1069 absorbance units for the field spectrum. Thus if the reference spectrum is multiplied by the factor 0.1069/0.00771 = 13.9 it will represent the same amount of methane as is in the field spectrum. That is to say that the absorbance in the data spectrum is indicative of 196 ppb $\times$ 13.9 = 2.7 ppm of methane. After multiplication, the reference can be subtracted from the field spectrum, and the methane should be removed from the water vapor spectrum.

Figure 5-4 shows the spectrum that has the methane removed and is now usable as a water vapor spectrum in the region of the methane absorbance. The operator must note, however, that many gases other than methane absorb in this region and, if they are still in the spectrum, they will cause errors in the analysis.

Removal of ozone from the water vapor spectrum is much more difficult, primarily because the absorption feature is broad. Figure 5-5 shows an atmospheric ozone absorption spectrum and the ozone reference spectrum. The slight elevation of the absorption spectrum in the vicinity of 1050 cm$^{-1}$ is indicative of an absorbance due to ozone with a concentration of about 100 ppb. A significant problem with ozone occurs when the synthetic background is made. Here the absorption spectrum is so broad that at least one point in the ozone spectrum region must be chosen for the baseline. It is easiest to think that the point in the center of the spectrum where the absorbance dips almost to the baseline should be used. However, the ozone absorbance does not go to zero at that wave number. So, a very careful estimate for where that point should be placed needs to be made. A difficulty arises that is associated with the MCT detector, for which there is a curvature of the baseline in this wave number region. This means that the baseline cannot be made by connecting a line between two points along the curve.

Once the synthetic background has been made and the operator has an absorption spectrum, the ozone still has to be subtracted from it. There is almost no possibility that the ozone concentration will actually go to zero at any location but it should go through a minimum. For areas such as Research Triangle Park, where the atmospheric ozone is produced locally and not transported into the area from elsewhere, that minimum occurs at about 6:00 in the morning.
Figure 5-3. Methane Reference Spectrum and the Calculated Absorption Spectrum.

Figure 5-4. Water Vapor Spectrum Made for the Methane Absorption Region.
A plot of several days of ozone concentration taken at Research Triangle Park during the month of June is shown in Figure 5-6. The negative values indicate that there is a significant amount of ozone in the water vapor spectrum. The question is just how much to subtract from the spectrum, because the ozone concentration does not go to zero. In this area, we are fortunate because there are other instruments that make measurements of ozone, and they can be used to determine the ozone minimum. Ozone is a criteria pollutant and is also monitored by the individual states. These data are generally available as hourly averages and may be useful to the operator who is trying to subtract the correct amount of ozone from the water vapor reference spectrum.

The operator must be aware that gases other than methane and ozone must be subtracted from the water vapor reference spectrum, even though they are all not covered here. Certainly the gases CO₂, N₂O, and CO must be subtracted from the water vapor reference. When data are taken at industrial sites, any gas that is to be monitored or used as an interfering species must be subtracted from the water vapor reference spectrum.
Figure 5-6. Ozone Measured at Research Triangle Park During June.
Chapter 6
Siting

SUMMARY

The topics and specific points of emphasis discussed in this chapter include the following:

- Siting considerations for long-term and short-term monitoring efforts
- Factors to consider when selecting the path
  - Short path versus long path
  - Prevailing winds
  - Slant path versus horizontal path
- What conditions warrant changing the path
- What ancillary measurements are required
- Calculation of the minimum path length required to detect specific concentrations of selected target gases
- An example of a specific monitoring site

6.1 Introduction and Overview

There are two kinds of monitoring programs for which siting needs to be discussed. One is a long-term effort with the instrument placed in a more or less permanent position. The second is a short-term program designed to take data at a site for a period from a few days to a few weeks. Each of these situations, while similar, requires somewhat different thinking to actually site the instrument. The short-term program is more flexible in that the path configuration can be based on the meteorological conditions at the time of the monitoring program. Long-term monitoring programs must be designed to allow for changes in the direction of the path as dictated by changing meteorological conditions, or useful data might be lost. Siting considerations for both situations are described in this chapter. Criteria for selecting the path, changing the path, and choosing the ancillary measurements to make at a monitoring site are also discussed.

There is little information in the literature pertaining to the long-term monitoring program. Even for the short-term program, the parameters that were considered in selecting the actual direction and length of the path have been discussed in only a cursory manner. A typical statement is "The path was set up based on a knowledge of the prevailing winds." But
what the wind field actually was during the observation period or where the prevailing wind data were taken is almost never presented. There is a similar lack of information concerning the selection of the path length and the partial pressure of water vapor in the atmosphere. For that matter, there is almost no discussion in the literature about how the length of the path is selected.

There are some sophisticated methods being studied by various groups that use real-time meteorological data for making decisions about the path. It seems that these techniques are more suited for permanent monitoring installations and not for short-term programs such as monitoring at small waste sites. Under any circumstances these methods have not been adequately tested, and an evaluation of these methods is considered to be beyond the scope of this document at the present time.

Other ongoing relevant work is being documented by the U.S. Environmental Protection Agency, which is preparing a set of changes to Part 58 of Chapter 1 of Title 40 of the Code of Federal Regulations (40CFR58) that will define the appropriate ambient air monitoring criteria for open-path (long-path) monitors (U.S. Environmental Protection Agency, 1994). These amendments have not been finalized or approved, and they specifically address the monitoring of the gases called the criteria pollutants. But the proposed amendments are significant in that they describe just how the path is to be chosen in terms of obstructions, height above the ground, and changes in path height. They also describe the appropriate positioning of the path in relation to buildings, stacks, and roadways.

Several factors must be considered when selecting the path. These factors include (1) instrumental parameters, such as the signal-to-noise ratio (S/N) of the system and the divergence of the IR beam; (2) the characteristics of the target gases, such as concentrations and absorption coefficients; (3) the presence and concentrations of interfering species, such as water vapor and CO₂; (4) meteorological data, such as wind direction and speed; and (5) physical constraints, such as the area of the emission source, the extent of the plume, and the availability of suitable sites to accommodate the instrument and peripherals.

Example calculations using Beer's law are given in this chapter to illustrate the minimum path length required to measure a specific concentration of a target gas. For example, the minimum path length required to measure ammonia at a concentration of 10 ppb would be approximately 21 m, assuming a minimum detection limit of 3 x 10⁻⁴ absorbance units, no interfering species, and a uniform concentration throughout the path. In contrast, the minimum path length required to measure 10 ppb of chlorobenzene would be 230 m. In general, the length of the path must be chosen to be the minimum length that will allow the measurement to be made with a meaningful statistical accuracy.

An example using a specific Superfund site is given. Procedures are described for selecting a usable path for a short-term intensive study.
6.2 Selecting the Path

To select the length and the position of the path, the investigator must have some understanding of the ramifications of these choices. The immediate questions concern (1) the effect that the path has on the data that is produced and (2) the procedure that the operator follows for selecting a path.

Preliminary answers to those questions are found by referring to Beer’s law. But the complete answers are more complicated. They include scattering and absorption by aerosols, the effects of water vapor and carbon dioxide on the S/N; and spectral interferences. When measuring plumes of finite extent, a path longer than the width of the plume is actually detrimental.

The proposed amendments to 40 CFR 58 describe the following considerations for selecting the path.

- At least 80% of the path must be between 3 and 15 m above the ground.

- At least 90% of the path must be at least 1 m vertically or horizontally away from walls, etc.

- If the path has to be near a building, then it must be on the windward side of the building.

- Buildings or other obstructions may possibly scavenge the gases of interest. At least 90% of the path must have unrestricted airflow and be located away from obstructions so that it is removed by at least twice the height that the obstacle protrudes above the path.

- At least 90% of the path must be at least 20 m from the drip lines of trees.

- When monitoring is done for ozone, 90% of the path must be at least 10 m from a road that carries fewer than 10,000 cars a day. This criterion changes to 250 m for heavily traveled roads (> 110,000 cars per day).

There are other proposed changes to 40 CFR Part 58 that are applicable to the use of FT-IR open-path monitors, but they are for concerns other than siting. The interested reader should obtain a copy of 40 CFR Part 58 from the Office of Federal Register, National Archives and Records Administration, Washington, DC. It is also available in most public libraries.

6.2.1 The Longest Path

It is possible to determine the maximum usable path in several ways. One is to use the noise equivalent power of the detector as the minimum signal that can be recorded. Another is to use a minimum S/N that the operator is willing to accept. Then if either the noise equivalent power (NEP) signal or the S/N is known at one distance and the energy falls off as the inverse square of the distance, the maximum possible path can be calculated. Any attempt to actually do this calculation results in the conclusion that the maximum path is essentially infinite.
In a practical sense the absorption due to water vapor will limit the usable path long before the theoretical limit calculated above can be reached. The water vapor line at 1014.2 cm\(^{-1}\) has an absorbance of 0.01 at a total path length of about 30 m when the water vapor partial pressure is 10 torr. If an absorbance of 1 is considered the maximum allowable for this line, then the maximum usable total path is about 3 km.

6.2.2 Shortest Path Requirements

The shortest path for various gases can be calculated from the absorbance measured in the reference spectra, a knowledge of the minimum measurable absorbance, and the assumption that reciprocity holds. To make this calculation, the operator must have chosen the wave number region that will be used for analysis and obtained the absorbance of the gas from the reference spectrum over that region. The operator must also choose a minimum concentration that is to be measured. Then, by using the minimum detectable absorbance, the minimum path can be calculated as follows.

1. Measure the absorbance at the appropriate wave number for the target gas from the reference spectrum. Record the concentration path length product at which this spectrum was taken.

2. Calculate the absorption coefficient \( \alpha \) for this gas by using the following formula.

\[
\alpha = \frac{A_r}{C_r L_r}
\]

where \( A \) is the absorbance and \( CL \) is the concentration-path length product. The subscript \( r \) refers to the reference spectrum.

3. Assume a minimum concentration that will be measured, and set the minimum detectable absorbance at 3 times the RMS baseline noise as measured under normal operating conditions, for example, \( 3 \times 10^{-4} \).

4. Calculate the minimum usable path \( (L_m) \) from

\[
L_m = \frac{A_m}{\alpha C_m}
\]

where \( A_m \) is the minimum absorbance \( (3 \times 10^{-4}) \) and \( C_m \) is the minimum concentration assumed in Step 3, and \( \alpha \) is the absorption coefficient calculated in Step 2.

The results of the above calculations for four different gases are given in Table 6-1.

6.2.3 Short Path Versus Long Path

As shown in the previous section, the selection of the path length begins by calculating the minimum usable length from Beer's law. If a retroreflector is used, the physical path can be half the optical path determined above. This is advantageous when plumes of finite size are being measured because the path length may be chosen close to the physical extent of the plume.
## Table 6-1. Minimum Usable Path Lengths*

<table>
<thead>
<tr>
<th>Gas</th>
<th>Wave Number (cm(^{-1}))</th>
<th>Absorbance (A_r)</th>
<th>Concentration (C \times L_r) (ppm-m)</th>
<th>Absorption Coefficient (\alpha)</th>
<th>Minimum Concentration (C_m) (ppb)</th>
<th>Minimum Usable Path Length (L_m) (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(p)-dichlorobenzene</td>
<td>822</td>
<td>0.085</td>
<td>500</td>
<td>(1.7 \times 10^4)</td>
<td>10</td>
<td>176</td>
</tr>
<tr>
<td>Chlorobenzene</td>
<td>1025</td>
<td>0.0227</td>
<td>170</td>
<td>(1.34 \times 10^4)</td>
<td>10</td>
<td>223</td>
</tr>
<tr>
<td>Toluene</td>
<td>1031</td>
<td>0.0203</td>
<td>496</td>
<td>(4.09 \times 10^{-5})</td>
<td>10</td>
<td>734</td>
</tr>
<tr>
<td>Benzene</td>
<td>1038</td>
<td>0.0027</td>
<td>27</td>
<td>(1.0 \times 10^{-4})</td>
<td>10</td>
<td>300</td>
</tr>
</tbody>
</table>

*\(A_r\) = absorbance of the reference spectrum, \(C \times L_r\) = concentration path length product at which the reference spectrum was taken, \(\alpha\) = absorption coefficient, \(C_m\) = minimum concentration, \(L_m\) = minimum usable path length.

The length of the path must be chosen to be the minimum length that will allow the measurement to be made with a meaningful statistical accuracy. For the calculations above, this distance was determined by using a minimum absorbance of \(3 \times 10^4\), or about 3 times the best detection limit that is achievable at the present time. For homogeneously distributed gases, the path can be made longer with some advantage. But for plumes of finite extent, making the path longer than the plume is wide would be a detriment and should not be done. This is because the measurement actually determines the path average concentration, and if a portion of the path has zero concentration, there is a dilution effect. Another reason for choosing a path that is as short as possible is that the effects of spectral interferences will be minimized.

A different example can be described as follows. There are times when a release of a tracer gas such as \(SF_6\) is desirable. The question is how much must be in the path if it is to be detected. From Figure 6-1, it is seen that the absorbance of \(SF_6\) at 947 cm\(^{-1}\) is 1.56. The concentration path length is 66 ppm \(\cdot\) m. Thus for a working detection limit of \(3 \times 10^4\) absorbance units and a path length of 50 m, the minimum average concentration in the path must be \(C = \frac{A}{\alpha L}\). The absorption coefficient \(\alpha\) is obtained from \(\alpha = \frac{A}{C L} = 1.56/66\). Thus \(C = (3 \times 10^4 \times 66)/(1.56 \times 50) = 0.25\) ppb. It is clearly seen in this example that because of its large absorption coefficient, not much \(SF_6\) is required for detection.

There is no distance with any of the available instruments that will reduce the absorption due to water vapor and carbon dioxide below the detection limits. In fact, in the wave number region of strongest absorbance for these gases the atmosphere is generally totally opaque. That is, there is so much light being absorbed that none returns to the detector. These regions are not usable for data analysis with the FT-IR systems.
For long-term monitoring programs with permanent installations, the only real option is to place retroreflectors or light sources (depending on the instrument configuration) at various distances and switch from one to the other periodically or on some predetermined schedule. A scanning system is available with some versions of open-path FT-IR monitors that facilitates this. Currently, almost no work has been done to define various lengths for various conditions. Thus, this chore must be individually repeated for each monitoring program.

6.2.4 Prevailing Winds

When using the FT-IR long-path technique, the operator depends on the wind to deliver the gases being emitted by a source to the infrared beam. Knowledge of the prevailing winds is important when setting up the path for long-term monitoring programs, but may be much less important for short-term programs. Most operators of open-path monitors have been concerned with short-term programs and know that the wind almost never comes from where the prevailing wind rose predicts. The short-term program usually demands that the operator be
prepared to change the path configuration when the wind changes. For either the long-term or the short-term program, the ideal situation is to have more than one retroreflector or light source. This allows the path direction and length to be changed as the requirements of the program dictate without having to transport the instrument itself.

When emission rates need to be calculated from data taken with an FT-IR instrument, the wind direction and speed must be known. The direction of the path with respect to the wind must also be known. A knowledge of the historical prevailing winds is of little use for this task. When emission rates are required, the wind field at the path must be measured directly.

6.2.5 Slant Path Versus Horizontal Path

Path orientation is important because the wind is the primary mode of transportation of the gases being monitored. Wind speed and direction can change dramatically over small regions when measured close to the ground. This is true not only because of the changing terrain but also because the motion of the air (a wind) must at least approach zero at the surface. There is some indication that the concentration contours of gases become very complex with altitude, at least in part because of turbulence. There are no data in the FT-IR literature that describe the variation of concentration with altitude. Because of these uncertainties, a comparison of the use of a slant path and a horizontal path cannot be made.

6.3 Changing the Path

The beginning of this chapter included a discussion of some of the ramifications of path selection. The question here is when should the path length or direction be changed? Obviously, if the plume from a point source or an area source is being monitored and the wind changes direction, the path should be changed. Changing the path, however, should be done in accordance with some plan. Items that need to be covered in the plan include the conditions that make a change necessary, as described above. They should also consider the ramifications of the change, both the advantages and the disadvantages. For example, if the concentrations of gases crossing a fence line are being monitored, there is little point in changing the direction of the path.

Change in the length of the path should be considered only for purposes of taking a background spectrum or when spectral interferences from compounds like water vapor become so strong that the absorption due to the target compounds is overwhelmed. Whether any accurate monitoring can be done under that condition has not been studied. Certainly, it makes no sense to reduce the path length to the point where the target compounds cannot be monitored.
6.4 Ancillary Measurements

There are several reasons why some ancillary measurements must be made when taking data with an FT-IR open-path sensor. One is the requirement to take data that can be used for quality control and quality assurance purposes. (See Chapter 8 for a discussion of quality control and quality assurance procedures.) Another is that many programs will require ancillary data such as wind speed and direction. Also, for the foreseeable future, the amount of water vapor in the atmosphere should be monitored because too many unanswered questions about water vapor exist. By far, water vapor represents the strongest spectral interference, and unless it is measured separately, problems may arise when the data are analyzed. It should be noted that a measurement of relative humidity is not satisfactory for this work. The actual partial pressure of water vapor must be found, and if relative humidity is measured, then the temperature must also be measured. The ambient pressure should also be recorded. At any one monitoring location operators can expect to experience a small change in ambient atmospheric pressure. In some cases, the data may have to be corrected for these changes. However, when acquiring data in places of high altitude, such as Denver, CO, a substantial change in pressure can be expected when compared to sea level. The operator must determine whether his experiment demands that these changes be accounted for in the data.

Guidance for selecting and setting up the instruments for making meteorological measurements can be found in a government handbook (U.S. Environmental Protection Agency 1989). Although this document does not directly address the long-path measurements, it does present useful information about meteorological instrumentation and measurements. For the long-path situation, the only measurements that should be obtained are probably those at or along the path itself.

6.5 A Specific Case

The literature offers very little information about procedures for selecting the path at any given site. The task is again divided into two parts: (1) selecting a path or set of paths for long-term monitoring at a fixed installation and (2) selecting a usable path for short-term intensive studies. For this document, we will discuss the latter case only and do so by presenting a real case.

Figure 6-2 is an aerial photograph of a Superfund site undergoing remediation. The active region of the site is at the middle left of the photograph. Two large repository pits can be seen, one in the top middle of the photograph and the other in the top right. The former pit has been filled and capped with dirt while the latter is open and in the process of being lined. To gauge the size of the site, note that the vehicles immediately to the left of the FT-IR monitor are D8 bulldozers.
Figure 6-2. Aerial Photograph of a Superfund Site Undergoing Remediation.
The site lies between two ridges (not shown), and the prevailing winds blow along the valley through the site from upper left to lower right. The first pit rises sharply from the active area for about 50 ft, and then the terrain falls off about 20 ft to the second pit. The road in front of the FT-IR monitor rises sharply in front of the second pit. The surrounding terrain is forest, with the trees rising about 40 ft above the ground level. Permission had been obtained to make measurements with the FT-IR monitor with the proviso that there would be no interference with the ongoing remediation. The FT-IR operators were not given access to the active area, which was defined as starting at the buildings in the foreground and extending to the forested region to the left of and behind the capped pit. The remediation operation entailed digging soil from the active area, repacking it in metal drums, and moving it to the lined pit areas. Fluids that were encountered in the active area or in old drums were brought to the settling tanks seen in the middle of the photograph. The predominant chemical in the site was the herbicide Dicamba. Dicamba has a very low vapor pressure, but two by-products were thought to be present. They were benzonitrile and benzaldehyde, and the goal of the FT-IR study was specifically to measure these two compounds. Funding had been allocated for one week of field work. Although the proposed amendments to 40 CFR Part 58 were not available at the time of this study, they were almost exactly followed.

Benzonitrile has a single usable absorption band at 757 cm\(^{-1}\). The absorbance of this band is 0.0346 when the concentration path length product is 186 ppm-m. Repeating the calculation described above for the absorption coefficient gives \( \alpha = \frac{0.0346}{186} = 1.9 \times 10^{-4} \). It was thought that benzonitrile would have a concentration of about 10 ppb. At the time of this study, the FT-IR instrument had a minimum detection limit of about 1 x 10\(^{-3}\) absorbance units. The minimum usable path is calculated as follows.

\[
I_m = \frac{1 \times 10^{-3}}{(1.9 \times 10^{-4} \times 10 \times 10^{-3})} = 525 \text{ m}
\]

(Eq. 6-1)

The problem is somewhat more complicated than this because there is a weak interfering carbon dioxide peak at 757 cm\(^{-1}\). The task was then to site the instrument so that the retroreflector could be placed 250 m away while adhering to the constraints imposed by the remediation process. To obtain electrical power without the use of a generator, the only logical place to put the FT-IR monitor was at the main entrance to the site, as it is shown in the Figure 6-2 photograph. From there, only two possible paths of 250 m were available. They are shown in the photograph at the RR positions. The path that extends from the FT-IR monitor to the right RR position in the photograph was selected as the primary path because this would encompass the entire plume coming from the active area according to the prevailing winds. The secondary path, extending from the FT-IR monitor to the bottom of the capped repository pit (at the left RR position), was not really satisfactory
because it rose too high above the ground level and went directly over the settling tanks. A path that is too high above the active area would leave the possibility that the plume might go under the beam. If the beam were to go directly over the settling tanks, flux calculations would be virtually impossible.

As an aside, during the week-long field program the remediation process was halted because of a problem with the lining of the second pit. Also, during this time, the wind never blew more than 0.5 mph, and its direction was almost always from the bottom to the top of the photograph, contrary to expectation.

6.6 References


Chapter 7
Resolution Considerations in FT-IR Long-Path, Open-Path Spectrometry

SUMMARY
The topics and specific points of emphasis discussed in this chapter include the following.
- The definition of resolution in FT-IR spectrometry
- The trading rules between resolution, the signal-to-noise (S/N) ratio, and measurement time
- Example spectra of atmospheric constituents and selected VOCs that illustrate the following effects:
  - Effect of resolution on peak shape and intensity
  - Effect of apodization and zero filling on peak shape and intensity
- A discussion of the effect of resolution on quantitative analysis
- A case study illustrating the effects of resolution, zero filling, and baseline noise on the CLS analysis of multicomponent mixtures

7.1 Introduction and Overview

One important issue regarding the use of long-path, open-path FT-IR systems for monitoring hazardous air pollutants is the appropriate spectral resolution to be used during data acquisition. A resolution should be chosen to maximize the ability to resolve spectral overlap while maintaining a balance between the S/N, analysis time, and data storage requirements. Several factors must be considered when determining the optimum resolution for measuring the IR spectra of atmospheric constituents along an open long path. These factors include (1) the ability to distinguish between the spectral features of target analytes and those of ambient interfering species in the atmosphere, such as water vapor and CO₂; (2) the trade-offs between resolution, IR peak absorbance, and S/N; and (3) practical considerations, such as measurement time, computational time to process the interferogram, and the size of the interferogram file for data storage. The use of an inadequate instrumental resolution can distort the true absorption spectrum, affect the quantitative relationship between absorbance and concentration, and diminish the ability to resolve spectral overlap. Resolutions ranging from 0.25 to 2 cm⁻¹ have been suggested for use in FT-IR monitoring,
but there currently is no consensus as to what resolution is generally applicable.

This chapter describes the fundamental aspects of resolution in FT-IR spectrometry and illustrates the effects of resolution and related instrumental parameters on the measured spectrum. The trading rules that determine the balance between resolution and S/N are discussed. Test spectra were obtained in the laboratory and along an open path to illustrate the effects of resolution, apodization, and zero filling on the IR spectra of CO₂ and water vapor, common atmospheric species that can interfere with analytical measurements, and selected gases and VOCs. Studies from the literature that address resolution requirements in long-path FT-IR monitoring are discussed. A case study illustrating the effects of resolution, zero filling, and baseline noise on the CLS analysis of multicomponent mixtures is also presented.

In FT-IR spectrometry, the minimum separation in wave numbers (cm⁻¹) of two spectral features that can be just resolved is inversely related to the maximum optical path difference in centimeters of the two mirrors employed in the Michelson interferometer. If the desired resolution is increased by a factor of 2, for example, from 1- to 0.5-cm⁻¹ resolution, the moving mirror in the interferometer must travel twice as far. In practical terms this means that the scan time will be approximately twice as long, the interferogram file will be approximately twice the size for data storage, and the time required to process the interferogram will be longer for the higher resolution measurement.

The instrumental resolution also affects the S/N. In general, if the size of the aperture, or Jacquinot stop, in the interferometer is held constant, the baseline noise in an FT-IR spectrum is directly proportional to the resolution of the interferometer for measurements made in equal times. For example, changing the resolution from 1- to 0.5-cm⁻¹ increases the noise level by a factor of 2 for equal measurement times. Therefore, to obtain the same baseline noise level for the 0.5-cm⁻¹ spectrum as was measured for the 1-cm⁻¹ spectrum, the measurement time would have to be quadrupled (because the S/N is proportional to the square root of the measurement time).

Resolution may also affect the peak absorbance of the bands being measured. For narrow and weak spectral features, the peak absorbance will approximately double on halving the resolution. In this case, the S/N would be the same for the spectra acquired at the higher and lower resolution settings, provided the measurement time was equal. For weak, broad spectral features whose peak absorbance does not change as a function of resolution, the lower resolution measurement is preferable. For strongly absorbing bands, whether they are broad or narrow, calibration curves of absorbance values measured at different resolutions and plotted versus concentration must be developed to ascertain the optimum resolution to be used.

In general, the minimum limit of detection (LOD) should be found for measurements made at the lowest possible
resolution that adequately resolves the spectral features of the analyte from those of interfering species. The use of an inadequate resolution can distort the true absorption spectrum, affect the quantitative relationship between absorbance and concentration, and diminish the ability to resolve spectral overlap. Conversely, the use of a higher resolution than is required can result in a poorer S/N and an unnecessary increase in measurement time, processing time, and data storage requirements.

There is currently no consensus as to what resolution and related parameters are generally applicable in long-path, open-path FT-IR monitoring. Most likely, the optimum resolution will need to be determined on a case-by-case basis, depending on the spectral characteristics of the target compounds and their concentration, the path length, and the presence of interfering species. In field measurements, a qualified judgement must be made taking into account these factors in addition to the practical considerations discussed above.

7.2 Definition of Resolution

An understanding of the resolution requirements in FT-IR long-path, open-path monitoring requires an understanding of the basic principles involved in generating an interferogram and the operations performed on the interferogram prior to converting it to a spectrum. The following discussion is an attempt to describe these basic principles in a way that will be of general use to analysts in FT-IR monitoring. For a more rigorous treatment of the fundamentals in FT-IR spectrometry, the reader is referred to the definitive text by Griffiths and de Haseth (1986) or several other excellent references (Horlick 1968; Bell 1972; Herres and Gronholz 1984) and Section 2.4.2 of this document.

As shown in Section 2.4.2, the minimum separation in wave numbers of two spectral features that can be resolved is inversely related to the maximum optical path difference, in centimeters, of the two interferometer mirrors employed in the Michelson interferometer. The closer the separation of the two spectral features, the greater the optical path difference must be before the spectral features can be resolved.

In terms of the measured spectrum, resolution can be defined as the minimum separation that two spectral features can have and still be distinguished from one another. A commonly used requirement for two spectral features to be considered resolved is the Raleigh criterion. This criterion states that two bands that have identical intensity, band shape, and peak width are resolved when the minimum of one band falls on the maximum of the other. When this is the case, there is a dip corresponding to approximately 20% of the absorption maxima between the two overlapping spectral features. It should be noted that this criterion is valid only for a sinc² instrument line shape, such as that found in a dispersive spectrometer or an FT-IR instrument using triangular apodization.
The actual spectral resolution in the frequency domain that can be obtained by an interferometer is also affected by the truncation of the interferogram and the application of various apodization functions. The apodization functions can increase the bandwidth and also change the line shape. Apodization is discussed further in Section 7.3.3.

7.3 Trading Rules in FT-IR Spectrometry

The quantitative relationships between the S/N, resolution, and measurement time in FT-IR spectrometry are referred to as "trading rules". The factors that affect the S/N and dictate the trading rules are expressed in Equation 7-1, which gives the S/N of a spectrum measured with a rapid-scanning Michelson interferometer. (The derivation of Equation 7-1 is given by Griffiths and de Haseth [1986].)

\[
\frac{S}{N} = \frac{U_c(T) \theta \Delta \nu^{1/2} \xi D^*}{(A_d)^{1/2}}
\]

(Eq. 7-1)

where \( U_c(T) \) = the spectral energy density at wave number \( \nu \) from a blackbody source at a temperature \( T \)

\( \theta \) = the optical throughput of the spectrometric system

\( \Delta \nu \) = is the resolution of the interferometer

\( t \) = is the measurement time in seconds

\( \xi \) = the efficiency of the interferometer

\( D^* \) = the specific detectivity, a measure of the sensitivity of the detector

\( A_d \) = the area of the detector element

As shown in Equation 7-1, the S/N of a spectrum is proportional to the square root of the measurement time \( (t^{1/2}) \). For measurements made with a rapid scanning interferometer operating at a constant mirror velocity at a given resolution, as would most likely be the case in FT-IR monitoring applications, the S/N increases with the square root of the number of scans being averaged.

The relationship between the S/N and resolution is not as straightforward as implied in Equation 7-1. If the physical parameters of the spectrometric system, such as the measurement time, optical throughput, and the interferometer efficiency, are assumed to be constant for measurements made at both high and low resolution, the S/N will be halved on doubling the maximum retardation of the interferometer \( (\Delta \nu_{\text{max}}) \) or halving the resolution \( (\Delta \nu/2) \). Because the S/N is proportional to the square root of the measurement time, the measurement time required to maintain the original baseline noise level must be increased by a factor of 4 each time \( \Delta \nu_{\text{max}} \) is doubled, or \( \Delta \nu \) is halved, for measurements made at a constant optical throughput.
The optical throughput does not necessarily remain constant throughout the range of resolutions that could be used to measure atmospheric gases. In low-resolution measurements, a large optical throughput is allowed for the interferometer, and the throughput is limited by the area of the detector element or the detector foreoptics. Most commercial low-resolution FT-IR spectrometers operate with a constant throughput for all resolution settings.

Instruments capable of high-resolution measurements are equipped with adjustable or interchangeable aperture (Jacquinot) stops installed in the source optics that reduce the solid angle of the beam passing through the interferometer. Spectra collected at high resolutions are generally measured with a variable throughput, which decreases as the spectral resolution increases.

In high-resolution measurements made under variable throughput conditions, the throughput is halved as \( \Delta_{\text{max}} \) is doubled. This results in an additional decrease in the S/N by one-half, which requires increasing the number of co-averaged scans by another factor of 4 to obtain the original S/N. Thus, for high-resolution FT-IR spectrometers operating under variable throughput conditions, the total measurement time is increased by a factor of 16 when \( \Delta_{\text{max}} \) is doubled.

The above discussions apply only to the effect of resolution on the baseline noise level. Resolution may also affect the peak absorbance of the bands being measured. For a weak and narrow spectral feature whose full width at half height (FWHH) is much less than the instrumental resolution, the peak absorbance will approximately double on doubling \( \Delta_{\text{max}} \). Assuming this band was measured under constant-throughput conditions, its S/N would be the same for measurements taken at the higher and lower resolution settings, provided the measurement times are equal. However, the degree of overlap by nearby spectral features will be reduced when the measurement is taken at a higher resolution. Therefore, in this case the higher resolution measurement is preferred.

For weak, broad spectral features whose peak absorbance does not change as a function of resolution, the lower resolution measurement is preferable when the optical throughput is constant. For strongly absorbing bands, whether they are broad or narrow, calibration curves of absorbance values measured at different resolutions and plotted versus concentration must be developed to ascertain the optimum resolution to be used.

In general, the minimum LOD should be found for measurements made at the lowest possible resolution that adequately resolves the spectral features of the analyte from those of interfering species. Increasing the resolution beyond this point degrades the S/N. In FT-IR monitoring, the optimum resolution will be determined by the band widths of the absorption lines in the spectra of the target compounds, the presence of interfering species, and the S/N of the
system. The optimum resolution will most likely vary with respect to specific analytes and measurement conditions.

7.4 Example Spectra of CO₂ and Water Vapor

Water vapor and CO₂ have IR absorption bands with theoretical bandwidths as narrow as 0.1 cm⁻¹, according to the USF HITRAN-PC database (University of South Florida 1993). To fully characterize the IR spectra of these compounds, which have absorption bands that may overlap with those of target compounds, an FT-IR spectrometer capable of high-resolution measurements must be employed.

A series of experiments was conducted on a benchtop FT-IR spectrometer in the laboratory and with a transportable FT-IR monitor in the field to illustrate the effects of resolution on the IR spectra of CO₂ and water vapor. Three separate series of experiments were performed. In the first set of experiments, single-beam sample and background spectra were collected at various instrumental resolution settings with the benchtop spectrometer purged with nitrogen. These spectra were used to generate the data given below in Section 7.4.1.1 (Table 7-1). In the second set of experiments, a background interferogram was collected at 0.125 cm⁻¹ with the spectrometer purged, and a sample interferogram was collected with the sample compartment open to the laboratory air. These interferograms were used to generate lower resolution spectra by reducing the number of data points used for the Fourier transform. These spectra are shown in Figures 7-1 and 7-2. In the third set of experiments, single-beam spectra were collected along an open path of 150 m at 0.5-, 1.0-, and 2.0-cm⁻¹ resolution with a transportable FT-IR monitor. These spectra are shown in Figure 7-3.

All laboratory spectra were collected on a benchtop FT-IR spectrometer, which has a nominal instrumental resolution selectable to 0.125 cm⁻¹. The data system uses two 68000 data processors and contains 2 megabytes of RAM. The long-path spectra of water vapor were obtained on a portable FT-IR spectrometer with resolutions selectable to a nominal 0.5 cm⁻¹. Data acquisition and manipulations were carried out by using a commercial software package on a 486/33-MHz personal computer with 8 megabytes of RAM.

7.4.1 Resolution Effects

The effects of resolution on the IR spectra of CO₂ and water vapor obtained from laboratory measurements and long-path measurements are addressed in this section. The laboratory measurements illustrate the relationship between resolution and scan time, data processing time, data storage requirements, spectral definition, and S/N. The long-path measurements are used to characterize the water vapor spectrum in a region in which it interferes with the analysis of an example target compound, toluene.
**TABLE 7-1. RESOLUTION TEST DATA**

<table>
<thead>
<tr>
<th>Resolution (cm(^{-1}))</th>
<th>Fourier Transform Points</th>
<th>File Size (bytes)</th>
<th>Scan Time (s)*</th>
<th>Process Time (s)</th>
<th>RMS Noise 2150-2100 cm(^{-1}) (10(^{-3}) Abs)**</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25</td>
<td>131072</td>
<td>300268</td>
<td>194</td>
<td>249</td>
<td>1.5115</td>
</tr>
<tr>
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<td>65536</td>
<td>168268</td>
<td>112</td>
<td>119</td>
<td>0.9007</td>
</tr>
<tr>
<td>1.0</td>
<td>32768</td>
<td>100267</td>
<td>69</td>
<td>60</td>
<td>0.4504</td>
</tr>
<tr>
<td>2.0</td>
<td>16384</td>
<td>67803</td>
<td>49</td>
<td>32</td>
<td>0.2347</td>
</tr>
<tr>
<td>4.0</td>
<td>8192</td>
<td>35034</td>
<td>28</td>
<td>16</td>
<td>0.1102</td>
</tr>
<tr>
<td>8.0</td>
<td>4096</td>
<td>18650</td>
<td>18</td>
<td>10</td>
<td>0.0590</td>
</tr>
<tr>
<td>16.0</td>
<td>2048</td>
<td>10266</td>
<td>13</td>
<td>8</td>
<td>0.0225</td>
</tr>
</tbody>
</table>

* Time to collect 100 scans.

** RMS noise for 1-min measurement times.

---

**Figure 7-1.** Single-Beam IR Spectra of CO\(_2\) Measured at (A) 0.25-cm\(^{-1}\), (B) 0.50-cm\(^{-1}\), (C) 1.0-cm\(^{-1}\), and (D) 2.0-cm\(^{-1}\) Resolution with No Apodization and No Additional Zero Filling.

7-7
Figure 7-2. Single-Beam IR Spectra of Water Vapor Measured at (A) 0.25-cm\(^{-1}\), (B) 0.50-cm\(^{-1}\), (C) 1.0-cm\(^{-1}\), and (D) 2.0-cm\(^{-1}\) Resolution with No Apodization and No Additional Zero Filling.

7.4.1.1 Laboratory Measurements

Single-beam spectra of the \(p\)- and \(r\)-branch of \(\text{CO}_2\) were recorded at resolutions ranging from 0.25 to 2 cm\(^{-1}\) as shown in Figure 7-1. No apodization or additional zero filling was applied to the interferograms prior to performing the Fourier transform. When plotted on the scale from 2400 to 2280 cm\(^{-1}\), the spectral features of the \(\text{CO}_2\) bands appear to be defined equally well at 0.25- and 0.5-cm\(^{-1}\) resolution, although slight differences were observed in the FWHH measurements. In the spectrum measured at 1-cm\(^{-1}\) resolution, there is a more noticeable degradation of the rotational fine structure. This structure is completely lost in the spectrum measured at 2-cm\(^{-1}\) resolution, and the \(r\)-branch appears as a broad continuum. The absorption bands that make up the
rotational fine structure of CO$_2$ have bandwidths of approximately 0.2 cm$^{-1}$, according to the USF HITRAN-PC database (University of South Florida 1993). Thus, these bands are not fully resolved, even at 0.25-cm$^{-1}$ resolution, and a resolution of 0.125 cm$^{-1}$ is required to fully characterize these bands.

Similar results were observed in spectra of water vapor measured at resolutions of 0.25, 0.5, 1, and 2 cm$^{-1}$ with no apodization or additional zero filling (Figure 7-2). The single-beam spectrum of water vapor between 3720 and 3620 cm$^{-1}$ exhibits several isolated, sharp features as well as overlapping features that are nearly baseline resolved in the spectrum measured at 0.25-cm$^{-1}$ resolution. The spectrum measured

![Graph showing spectra of water vapor at different resolutions](image)

**Figure 7-3.** Single-Beam IR Spectra of Water Vapor Measured at (A) 2-cm$^{-1}$, (B) 1-cm$^{-1}$, and (C) 0.5-cm$^{-1}$ Resolution over a 150-m Path.
at 0.5-cm\(^{-1}\) resolution exhibits a slight degradation in spectral definition as compared to the 0.25-cm\(^{-1}\) spectrum, although the general characteristics of the bands are retained. The overall band structure is still present in the spectrum measured at 1-cm\(^{-1}\) resolution; however, the distinction between some of the closely spaced, weaker bands is lost. In the spectrum measured at 2-cm\(^{-1}\) resolution, there is no longer any evidence of the spectral definition exhibited in the previous spectra, and the bands are significantly broader.

The effect of resolution on scan time, data processing time, data storage requirements, and baseline noise levels was also determined. For these tests, single-beam sample and background spectra were measured independently at each resolution setting, while holding the Jacquinot stop constant. The number of scans for each spectrum was 100. The RMS noise levels were calculated from spectra with a 1-min measurement time. The results from this test are presented in Table 7-1.

As shown in Table 7-1, the time required to collect 100 scans increases significantly upon acquiring data at higher resolutions. This is because the moving mirror in the interferometer must travel a greater distance as \(\Delta_{\text{max}}\) is increased for higher resolution scans. On average, the total scan time for 100 scans for this particular instrument increased by a factor of 1.6 each time \(\Delta_{\text{max}}\) was doubled, indicating the poor duty cycle efficiency.

The time required to process the interferogram also increases as \(\Delta_{\text{max}}\) (and as the number of data points collected for the interferogram) increases. In these examples, the processing time is that required to perform the Fourier transform and related operations on both the background and sample interferograms and to calculate the absorption spectrum. The time required to process the interferogram increased on average 1.8 times each time \(\Delta_{\text{max}}\) was doubled. It should be noted that these data were processed on a relatively old data system using a 68000 processor chip. With newer and faster computers the time required to perform the Fourier transform is not as much of a factor. For example, on a 486/33MHz machine with 8 megabytes of RAM, the time required to perform the Fourier transform and plot a single-beam spectrum is 1.65, 3.23, and 7.74 s for 2-, 1-, and 0.5-cm\(^{-1}\)-resolution interferograms, respectively. However, if time resolution is an important parameter in a specific FT-IR monitoring application, then the combination of scan time and processing time should be considered, or the interferograms should be stored for post-run processing.

The amount of disk space required for data storage increases almost by a factor of 2 each time the resolution is increased by a factor of 2. For 3.5-in floppy disks with 1.4 megabytes of storage capacity, this means that, for example, 14 interferograms collected at 1-cm\(^{-1}\) resolution could be stored on one disk, whereas only eight 0.5-cm\(^{-1}\) interferograms could be stored on one disk at
a time. The newer data acquisition software packages and data stations make more efficient use of disk space. For example, 21 interferograms collected at 1-cm⁻¹ resolution on a newer system could be stored on one 3.5-in. floppy disk. If large amounts of data are expected to be collected, such as might be the case in routine FT-IR monitoring studies, data storage requirements could be an important consideration.

The RMS noise measured between 2200 and 2100 cm⁻¹ increases as Δ_max increases. The data in Table 7-1 were taken from absorption spectra created from background and sample spectra collected over a 1-min scan time with a constant aperture at each resolution setting. These data follow closely the twofold increases in baseline noise expected each time the resolution is increased by a factor of 2. It should be noted that only the baseline noise level was measured in this experiment. Resolution may also affect the peak absorbance of the bands being measured. For example, for a weak spectral feature whose FWHH is much less than the instrumental resolution, the peak absorbance will approximately double on increasing the resolution by a factor of 2. Therefore, the S/N would be the same for measurements taken at the higher and lower resolution settings, provided the measurement times are equal. For broad bands, the peak absorbance will not be affected by changes in resolution, and the lower resolution measurement would be preferred.

7.4.1.2 Long-Path Measurements

The laboratory measurements described above illustrate some of the trade-offs encountered between resolution and other experimental parameters. However, the path length used in those studies was insufficient to detect many of the water vapor bands that interfere with bands of target pollutants, such as toluene. Russwurm (1992) has addressed the limitations that the presence of overlapping water vapor bands impose on the ability to detect and quantify toluene by long-path FT-IR spectrometry. With a CLS analysis of the toluene band at 1031 cm⁻¹, the detection limit for toluene in the presence of 10.5 torr of water vapor was estimated to be approximately 1 ppm. In this spectral region over a path length of 420 m, the absorbance due to water vapor was found to be strong compared to that of toluene. These data clearly indicate that to optimize the detection limits of FT-IR monitors for difficult target compounds, such as toluene, the water vapor spectrum must be well characterized.

Single-beam spectra measured at 2-, 1-, and 0.5-cm⁻¹ over a 150-m path length are shown in Figure 7-3. The interferograms were processed with triangular apodization and no additional zero filling. The spectra are plotted over the wave number region used to quantify toluene. As expected, the FWHHs of the water vapor absorption bands between 1000 and 1060 cm⁻¹ are narrower in the 0.5-cm⁻¹ resolution spectrum as compared to the 1- and 2-cm⁻¹ resolution spectra. In addition, spectral features are resolved in the
0.5-cm⁻¹ spectrum that appear as a single band in the other two spectra. For example, the band at 1010 cm⁻¹ in the 1-cm⁻¹ spectrum is resolved into a doublet at 1010 and 1010.7 cm⁻¹ in the 0.5-cm⁻¹ spectrum. Also, bands appearing at 1028.3 and 1029.5 cm⁻¹ are much better resolved in the 0.5-cm⁻¹ spectrum. In fact, they are not resolved at all in the 2-cm⁻¹ spectrum.

To completely resolve all of the overlapping bands in the spectrum of water vapor over this wave number region, the spectrum must be recorded at 0.125-cm⁻¹ resolution. The theoretical spectrum of water vapor from the USF HITRAN-PC database is shown in Figure 7-4A.

A 0.125-cm⁻¹ resolution spectrum of water vapor recorded on the ROSE system (Herget 1992) is shown in Figure 7-4B. In these spectra the band at 1018 cm⁻¹ can be resolved into two components, and the bands at 1010 and 1010.7 cm⁻¹ are completely baseline resolved. Whether or not measuring ambient spectra at 0.125-cm⁻¹ resolution improves the results of quantitative analyses for difficult target compounds, such as toluene, has not yet been fully investigated.

7.4.2 Zero-Filling Effects

When the interferogram contains frequencies that do not coincide with the frequency sample points, the spectrum resembles a "picket fence" (Herres and Gronholz 1984). An example of this effect is shown in Figure 7-5 in the spectrum of CO₂ measured on a benchtop FT-IR instrument. In this example, the spectrum of CO₂ measured at 0.25-cm⁻¹ resolution with no apodization and a zero-filling factor of 1 (Figure 7-5A) exhibits excellent peak shape. However, in the spectrum measured at 0.5 cm⁻¹ with no additional zero filling (Figure 7-5B), the peaks of several absorption bands are squared off. This effect can be overcome by adding zeros to the end of the interferogram before the Fourier transform is performed. This operation is referred to as zero filling. Zero filling increases the number of points per wave number in the spectrum, and, in effect, interpolates the spectrum. Normally, some multiple (e.g., 2, 4, etc.) of the original number of data points is added to the interferogram. This improves the photometric accuracy of the FT-IR spectrum and increases the digital resolution. As shown in Figure 7-5C, zero filling the interferogram measured at 0.5 cm⁻¹ by an additional factor of 2 eliminated the picket fence effect.

It should be noted that zero filling improves only the digital resolution, and not the resolution of the FT-IR spectrum. An example of this is illustrated in Figure 7-6 for spectra of water vapor measured at 0.25-, 0.5-, and 1-cm⁻¹ resolution. The spectrum measured at 0.25-cm⁻¹ resolution was zero filled by a factor of 1, the 0.5-cm⁻¹ spectrum was zero filled by a factor of 2, and the 1-cm⁻¹ spectrum by a factor of 4. In this case, each of the interferograms contained the same number of data points after being zero filled. Even with additional zero filling, the 0.5-cm⁻¹ spectrum does not match the spectral definition of the spectrum obtained at 0.25-cm⁻¹ resolution. (Compare Spectra A
Figure 7-4. IR Spectra of Water. (A) 10 torr of water vapor over a 300-m path, from the USF HITRAN-PC database (University of South Florida 1993). (B) Water vapor recorded at 0.125-cm\(^{-1}\) resolution of the ROSE system (reproduced with permission from W.F. Herget).
Figure 7-5. Absorption Spectra of CO₂ Measured at (A) 0.25 cm⁻¹ with a Zero-Filling Factor of 1, (B) 0.5 cm⁻¹ with No Zero Filling, and (C) 0.5 cm⁻¹ with a Zero-Filling Factor of 2.

and B in Figure 7-6.) The loss of spectral features is more dramatic in the zero-filled 1-cm⁻¹ spectrum. For example, shoulders at 3905 and 3884 cm⁻¹ that are detectable in the 0.25- and 0.5-cm⁻¹ spectra were not observed in the 1-cm⁻¹ spectrum. Also, side lobes appear in the 1-cm⁻¹ spectrum that was zero filled by a factor of 4 (Figure 7-6, Spectrum C). These side lobes are also present, but are not as severe, in the 0.5-cm⁻¹ spectrum zero filled by a factor of 2 (Spectrum B in Figure 7-6).

The picket fence effect is less extreme if the spectral components are broad enough to be spread over several sampling positions. As a rule of thumb, the original interferogram size should be doubled by zero filling by an additional factor of 2. When a CLS analysis of the spectral data is performed, in general it has been found that one order of zero filling (which is 2 times the original number of data points used in the Fourier transform) yields a factor of 2 lower error than that with no additional zero filling. An example of this is
given in Section 7.5.2.2.1. It should be noted that zero filling does increase the file size and the time required for data processing.

7.4.3 Apodization Effects

As shown in Chapter 2, the Fourier transform integral has infinite limits for the optical path difference. Thus, to measure the true spectrum of the source, the interferometer must scan infinite distances. However, because the mirror can move only a finite distance, the exact reconstruction of the spectrum is impossible. The finite movement of the interferometer mirror truncates, or cuts off, the interferogram. This, in effect, multiplies the interferogram by a boxcar truncation function. This function may cause the appearance of side lobes on both sides of the absorption band. The corrective procedure for eliminating these side lobes is called apodization. Apodization is
performed by multiplying the measured interferogram by a mathematical function. Typical apodization functions include triangular, Happ-Genzel, and Norton-Beer functions. An example of the effect of these apodization functions on the FT-IR spectrum of CO is shown in Figure 7-7.

Apodization affects the effective spectral resolution, the apparent peak absorbance, and the noise of any FT-IR spectrum. The apparent absorbance of narrow bands will be most affected by the choice of apodization function. In general, the bands in a spectrum computed with no apodization will be more intense than bands in the spectrum of the same sample computed from the same interferogram after applying an apodization function.

Apodization also degrades resolution slightly. An example of this is illustrated by the spectra of water vapor in Figure 7-8. In this case, subtle differences are observed, for

![Apodization Functions](image)

**Figure 7-7.** Absorption Spectra of CO Measured at a Nominal 0.125-cm\(^{-1}\) Resolution with (A) No, (B) Triangular, (C) Happ-Genzel, and (D) Norton-Beer-Medium Apodization Functions.
example, at 3948 and 3947 cm\(^{-1}\) and 3924.4 cm\(^{-1}\), in the spectrum generated with no apodization (Figure 7-8A) and the spectra generated by using the three types of apodization functions.

In general, to obtain the optimum S/N for spectra of small molecules with resolvable fine structure, the use of no apodization is preferable if side lobes from neighboring intense lines do not present an interference. If side lobes are present and interfere with either qualitative or quantitative analyses, apodization becomes necessary. For broad absorption bands, the measured absorbance is about the same in apodized and unapodized spectra. Overall, the greatest noise suppression will be obtained with the strongest apodization function, but the spectral resolution and band intensities will be greatest for weaker apodization functions (Griffiths and de Haseth 1986). The optimum apodization function has yet to be determined for general use in long-path FT-IR monitoring.

![Figure 7-8](image)

**Figure 7-8.** Absorption Spectra of Water Vapor Measured at 0.5-cm\(^{-1}\) Resolution with an Additional Zero-Filling Factor of 2 and with (A) No, (B) Triangular, (C) Happ-Genzel, and (D) Norton-Beer-Medium Apodization Functions.
Triangular and Happ-Genzel apodization functions are commonly used in OP/FT-IR monitoring, although Griffiths et al. (1995) have indicated that a Norton-Beer medium function actually gives a better representation of the true absorbance. In all cases, however, the same parameters should be used to collect the field spectra that were used to record the reference spectra. The choice of apodization function may be limited by this requirement. If spectra from a commercial or user-generated library are to be the reference spectra for quantitative analysis, then the parameters that were used to generate those reference spectra should be used to collect the field spectra. Otherwise, errors in the concentration measurement will occur.

7.5 Effect of Resolution on Quantitative Analyses

The determination of analyte concentrations by FT-IR spectrometry depends on the linear relationship between IR absorbance and concentration as described by Beer’s law. This linear relationship is observed only when the true peak absorbance of the analytical band(s) is less than a value of 0.7 and the spectrum is measured at a resolution that is higher than the FWHH of the band. If the FWHH of the band is narrower than the instrumental resolution, the measured spectrum is actually a convolution of the instrument line shape and true band shape. As a result, the measured absorbance will be only approximately linear with concentration. The higher the resolution for the spectral features of the IR band chosen for quantification, the better the approximation. The apodization function also has an effect on linearity. This section describes studies from the literature that have addressed the effects of resolution and related parameters on quantitative analyses.

7.5.1 Studies from the Literature

Strang et al. (1989) designed and evaluated an FT-IR system for monitoring toxic emissions from semiconductor manufacturing processes. This system was used to analyze part-per-billion levels of organic vapors and metal hydrides such as arsine, phosphine, and diborane in simulated workplace environments. The optimal wave number region for quantification and the effects of resolution and spectral overlap on the accuracy of quantitative results were studied. Spectral measurements were taken at resolutions ranging from 0.5 to 8 cm\(^{-1}\) to determine the optimum balance between (1) analysis time, (2) data storage space, (3) S/N, (4) accuracy of quantitative analyses using a CLS program, and (5) the ability to differentiate compounds with overlapping spectra. It should be noted that these data were acquired over a 20.25-m path in a multipass cell. Therefore, findings from this study may not be applicable to long, open-path measurements. Discussion of this study is included in this text because of the interest in workplace monitoring. Also, the methodology used to determine the optimum resolution for the short-path measurements can also be applied to longer path measurements.
The authors specified the following four issues that must be resolved for a CLS analysis at a given resolution to be acceptable.

1. Whether the CLS result varies by more than 50% of the theoretical value
2. Whether false positives or false negatives develop as a result of degraded resolution
3. Whether the amount of error in the measurement will cause potentially toxic concentrations of the target analyte in air to be measured incorrectly
4. Whether the detection limit obtained with the CLS program changes as a function of resolution

Using these criteria, the authors determined the minimum allowable resolution for the target compounds. The results are summarized below in Table 7-2.

In this study, the higher resolutions required for the metal hydrides arsine, diborane, and phosphine were a result of spectral overlap with other target analytes and with interfering atmospheric compounds. For example, phosphine overlaps with CO$_2$ and arsine overlaps with water vapor. The authors also determined the effects of decreased resolution on the accuracy of the quantitative results. In the case of diborane, only the 0.5-cm$^{-1}$ resolution measurements exhibited a linear relationship for all concentrations. Measurements taken at 2- and 4-cm$^{-1}$ resolution deviated from linearity as the concentration decreased.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Wave Number Region (cm$^{-1}$)</th>
<th>Minimum Resolution (cm$^{-1}$)</th>
</tr>
</thead>
<tbody>
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<td>Acetone</td>
<td>1287-1167</td>
<td>8</td>
</tr>
<tr>
<td>Arsine</td>
<td>2132-2106</td>
<td>2</td>
</tr>
<tr>
<td>Diborane</td>
<td>2522-2515</td>
<td>0.5</td>
</tr>
<tr>
<td>o-Dichlorobenzene</td>
<td>1060-1002</td>
<td>8</td>
</tr>
<tr>
<td>2-Ethoxyethanol</td>
<td>1192-984</td>
<td>8</td>
</tr>
<tr>
<td>Freon 11</td>
<td>876-813</td>
<td>8</td>
</tr>
<tr>
<td>Freon 13B1</td>
<td>1137-1031</td>
<td>8</td>
</tr>
<tr>
<td>Freon 22</td>
<td>1193-1063</td>
<td>8</td>
</tr>
<tr>
<td>Nitrogen trifluoride</td>
<td>960-833</td>
<td>8</td>
</tr>
<tr>
<td>Phosphine</td>
<td>2440-2390</td>
<td>4</td>
</tr>
<tr>
<td>Sulfur hexafluoride</td>
<td>965-915</td>
<td>8</td>
</tr>
</tbody>
</table>

*From Strang et al. (1989). These data are applicable to a 20,25-m path.
The effect of resolution on the ability to quantify overlapping compounds by the CLS analysis was investigated by using mixtures of Freons 11, 13B1, and 22. These compounds have relatively broad spectral features that overlap. The mixtures were analyzed at 0.5-, 2-, 4-, and 8-cm\(^{-1}\) resolution. Each of the individual compounds could be quantified accurately at each resolution in a 1:1:1 mixture at concentrations of 10, 1, and 0.1 ppm.

Strang and Levine (1989) have also determined the LODs for the same target compounds in the previous study as a function of resolution. For most compounds, there was very little difference in the LODs estimated at resolutions of 0.5, 2, 4, and 8 cm\(^{-1}\). However, for diborane and phosphine the LOD was difficult or impossible to measure at 8 cm\(^{-1}\) resolution. In the case of diborane, the peak selected for quantification had a FWHM of 7 cm\(^{-1}\). At 8-cm\(^{-1}\) resolution there are only two data points every 8 cm\(^{-1}\), so a peak 7 cm\(^{-1}\) in width is not defined well enough to be quantified by the CLS program. For phosphine, the peak shape is severely degraded at 8 cm\(^{-1}\). Although the CLS program could quantify the peak, the LOD was significantly higher (0.7 ppm-v/v) for spectra measured at 8 cm\(^{-1}\) resolution as compared to those measured at 0.5 cm\(^{-1}\) resolution (0.07 ppm-v/v). This example also illustrates one of the advantages of the CLS program over single peak absorbance measurements in quantitative analysis. The single peak absorbance measurement is difficult to make for broad bands, whereas the CLS program uses multiple data points over the entire spectral range of the broad band.

Spellicy et al. (1991) addressed several issues regarding spectroscopic remote sensing with respect to the Clean Air Act. One issue that was addressed was the optimum resolution for remote sensing FT-IR applications. The authors presented theoretical calculations describing the relationship between absorbance and concentration for a single Lorentzian line with a half-width of 0.1 cm\(^{-1}\) measured at resolutions from 0.01 to 0.1 cm\(^{-1}\). Under these conditions, linearity was observed only at a highest resolution case and at the lowest concentrations. The deviation from linearity most likely would be observed in small molecules such as HCl, CO, CO\(_2\), and H\(_2\)O, which have sharp spectral features. For larger compounds, such as heavy hydrocarbons that exhibit broader IR bands, the linear relationship between absorbance and concentration is more likely to be followed.

More recently, Marshall et al. (1994) conducted a laboratory study to determine the effect of resolution on the multicomponent analysis of VOCs with a CLS program. When they analyzed for target VOCs, such as acetone, chloroform, toluene, methanol, 1,1,1-trichloroethane, methyl ethyl ketone, carbon tetrachloride, and the xylene isomers, over a short path, resolutions lower than 4 cm\(^{-1}\) had an adverse effect on the multicomponent analysis. Resolutions of 1 to 2 cm\(^{-1}\) were found to be adequate for these target compounds when the CLS program was used.
Griffiths et al. (1993) reported the advantages and disadvantages of using low-resolution measurements in long-path FT-IR monitoring. Among the advantages cited were the smaller size and greater portability of the instrument, an improved S/N, and a lower cost. The disadvantages included a greater difficulty in visualizing the IR bands of the target compounds and potential deviations from Beer’s law. A test case of measuring the xylene isomers at resolutions of 2, 4, 8, and 16 cm\(^{-1}\) was presented. By using a partial least squares (PLS) program, good quantitative results were obtained at the relatively low resolution measurements. These results also indicated that the PLS program might be better than the CLS program for distinguishing and quantifying target compounds with overlapping features.

Bittner et al. (1994) have reported on high-resolution FT-IR measurements of VOCs at a variety of monitoring sites. By recording spectra at 0.125-cm\(^{-1}\) resolution, detection limits for benzene of 0.5 ppm-m were achieved at path lengths between 60 and 100 m at a fuel storage area. The high-resolution measurements allowed the narrow benzene band at 674 cm\(^{-1}\) to be separated from the strong CO\(_2\) absorption bands in that spectral region.

\subsection{5.2 Case Study: The Effect of Resolution and Related Parameters on the CLS Analysis of Multicomponent Mixtures}

A study using laboratory-generated spectral mixtures that have overlapping features was conducted to investigate the effect of instrumental resolution and related parameters on the CLS analysis results (Childers and Thompson 1994). The study was designed to simulate conditions that might be encountered in long-path measurements. The results from three separate cases are discussed.

1. Analytes with narrow bands that overlap, such as those for CO and \(^{13}\)C-labeled CO

2. Analytes with broad bands that overlap, such as those for acetone, methylene chloride, and ethanol

3. Analytes with narrow and broad bands that overlap, such as those for nitrous oxide and methylene chloride.

The effect of the number of data points and the noise level on the CLS analysis is also illustrated. Because the mixtures analyzed in this study were created from a linear combination of reference spectra, the effect of resolution on the relationship between concentration and absorbance is not addressed.

The spectra were collected on a research-grade, benchtop FT-IR spectrometer equipped with an MCT detector. A gas cell, 50 mm long and 32 mm in diameter, was used to obtain reference spectra of CO, \(^{13}\)CO, acetone, methylene chloride, ethanol, and nitrous oxide at room temperature and atmospheric pressure. The reference spectra were acquired at resolution settings of 0.125 and 1 cm\(^{-1}\). The original interferograms were processed by using the appropriate number of
data points to yield spectra with nominal resolutions ranging from 0.25 to 8 cm\(^{-1}\). No additional zero filling was used on the 0.25-cm\(^{-1}\) spectra because of memory limitations in the data system. The 0.5-cm\(^{-1}\) spectra were zero filled by an additional factor of 2, and the 1.0-cm\(^{-1}\) spectra were zero filled by an additional factor of 4. As a result, the 0.25-, 0.5-, and 1.0-cm\(^{-1}\)-resolution spectra had the same number of data points, that is 131,072. The 2-, 4-, and 8-cm\(^{-1}\) spectra were generated from interferograms that contained 16,384, 8192, and 4096 data points, respectively. No additional zero filling was performed on these interferograms. A triangular apodization function was applied to each interferogram prior to performing the fast Fourier transform.

The concentrations of the individual analytes in the gas cell were determined by comparing the maximum absorbance values of the 0.5-cm\(^{-1}\) spectra to those of reference spectra in a commercial spectral library. The absorbance values of the reference spectra were then normalized to values corresponding to a concentration of 100 ppm. These spectra were added mathematically to produce synthetic mixtures with varying concentrations of each analyte. Synthetic noise corresponding to an amplitude of 1, 5, 10, and 25% of the most intense peak in each spectrum was added to the mixtures. Mixtures with 10% noise added were then analyzed by using a CLS algorithm.

### 7.5.2.1 Mixtures of CO and \(^{13}\)CO

Synthetic mixtures of CO and \(^{13}\)CO were generated by adding reference spectra of CO corresponding to concentrations of 150, 300, 450, and 600 ppm to reference spectra of 100 ppm \(^{13}\)CO. The 0.25-cm\(^{-1}\) reference spectra and the spectra of the synthetic mixtures corresponding to 150 ppm of CO and 100 ppm of \(^{13}\)CO recorded at 0.25, 0.5, 1.0, and 2.0 cm\(^{-1}\) resolution are shown in Figure 7-9. CO and \(^{13}\)CO have several bands that overlap or nearly overlap in the spectral region between 2118 and 2137 cm\(^{-1}\). For example, overlapping bands at 2123.6 and 2124.2 cm\(^{-1}\) and at 2131.0 and 2131.6 cm\(^{-1}\) are nearly baseline resolved in the 0.25 cm\(^{-1}\) spectrum. These bands can still be distinguished at a resolution of 0.5 cm\(^{-1}\), but appear as only one band in the 1.0- and 2.0-cm\(^{-1}\) spectra. Even though these bands are not resolved at 1.0 and 2.0 cm\(^{-1}\) resolution, the CLS analysis accurately determined the concentration of CO in the mixture when it was analyzed for both CO and \(^{13}\)CO. Plots of the calculated concentration versus the known concentration of CO were linear over the range of 0 to 600 ppm of CO in the presence of 100 ppm \(^{13}\)CO. When the mixtures were analyzed for only CO, a positive bias and an increase in the magnitude of the errors in the measurements were observed. Although the bias was relatively constant, the error increased as the resolution decreased from 0.25 to 1.0 cm\(^{-1}\). In the case
Figure 7-9. Reference 0.25-cm⁻¹ Spectra of (A) \(^{13}\)CO and (B) CO and Spectra of Synthetic Mixtures of 150 ppm CO and 100 ppm \(^{13}\)CO Measured at (C) 0.25⁻¹, (D) 0.5⁻¹, (E) 1.0⁻¹, and (F) 2.0-cm⁻¹ Resolution.

Of the 2-cm⁻¹ measurements, CO could not be detected at 150 ppm in the mixture if \(^{13}\)CO was excluded from the analysis. (See Figure 7-10.)

7.5.2.2 Mixtures of Acetone, Methylene Chloride, and Ethanol

Synthetic mixtures of acetone, methylene chloride, and ethanol were
Figure 7-10. Concentration Calculated from CLS Analysis vs. Known Concentration for $^{13}$CO/CO Mixtures Measured at 2-cm$^{-1}$ Resolution. The (■) represents a value obtained during analysis for both $^{13}$CO and CO, and the (♦) represents a value obtained during analysis for CO only.

generated by adding reference spectra of ethanol corresponding to concentrations of 125, 250, 375, and 500 ppm to reference spectra of 100 ppm each of acetone and methylene chloride. The 0.25-cm$^{-1}$ reference spectra and the spectra of the synthetic mixtures corresponding to 500 ppm of ethanol and 100 ppm each of acetone and methylene chloride recorded at 1.0-, 2.0-, and 4.0-cm$^{-1}$ resolution are shown in Figure 7-11. In this case, the spectrum of each analyte could be adequately measured at 1.0-cm$^{-1}$ resolution. At 4-cm$^{-1}$ resolution, the Q-branches of these compounds were no longer detected. However, this did not diminish the ability to quantify ethanol in these mixtures with the CLS algorithm. Plots of the calculated concentration versus the known concentration of ethanol were linear over the entire range from 0 to 500 ppm ethanol in the presence of 100 ppm of acetone and 100 ppm of methylene chloride for each resolution setting. Although the concentration of ethanol could be determined in the low-resolution measurements, the errors in the CLS analysis increased with decreasing spectral resolution. The high-resolution spectra contain more data points per wave number than do the low-resolution measurements. To determine if this contributed to the increase in the magnitude of the errors, the effect of the number of data points on the CLS analysis was investigated. The effect of S/N on CLS was also investigated.

7.5.2.2.1 Effect of the Number of Data Points on the CLS Analysis

The average error in the CLS analysis for ethanol in the acetone, methylene chloride, ethanol synthetic mixtures for each resolution setting is shown in Table 7-3.

For the 0.25-, 0.5-, and 1.0-cm$^{-1}$ measurements, in which additional zero filling was used to keep the number of data points the same, the average error was relatively constant at approximately 9%. However, for the 2.0-, 4.0-, and 8.0 cm$^{-1}$ measurements, in which no additional zero filling was used, the average error increased with a decrease in the number of data points. (Note that the 0.25-, 0.5-, and 1.0-cm$^{-1}$ spectra were generated
Figure 7-11. Reference 0.25-cm⁻¹ Spectra of (A) Acetone, (B) Methylene Chloride, and (C) Ethanol and Spectra of Synthetic Mixtures of 100 ppm Acetone, 100 ppm Methylene Chloride, and 500 ppm Ethanol Measured at (D) 1.0-, (E) 2.0-, and (F) 4.0-cm⁻¹ Resolution.

<table>
<thead>
<tr>
<th>Resolution (cm⁻¹)</th>
<th>Additional Zero Filling</th>
<th>Data Points</th>
<th>Average Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25</td>
<td>None</td>
<td>131072</td>
<td>9.13</td>
</tr>
<tr>
<td>0.5</td>
<td>2 ×</td>
<td>131072</td>
<td>8.93</td>
</tr>
<tr>
<td>1.0</td>
<td>4 ×</td>
<td>131072</td>
<td>9.55</td>
</tr>
<tr>
<td>2.0</td>
<td>None</td>
<td>16384</td>
<td>24.54</td>
</tr>
<tr>
<td>4.0</td>
<td>None</td>
<td>8192</td>
<td>32.73</td>
</tr>
<tr>
<td>8.0</td>
<td>None</td>
<td>4096</td>
<td>42.27</td>
</tr>
</tbody>
</table>
from an original interferogram collected at 0.125-cm\(^{-1}\) resolution, whereas the 2.0-, 4.0-, and 8.0-cm\(^{-1}\) spectra were generated from an original interferogram collected at 1.0-cm\(^{-1}\) resolution.)

To show that the increase in error is related to the number of data points per wave number, and is not necessarily a direct result of degrading the spectral resolution, spectral mixtures of acetone, methylene chloride, and ethanol obtained at 1-cm\(^{-1}\) resolution were processed by using no additional zero filling, an additional zero filling factor of 2, and an additional zero filling factor of 4. This resulted in interferograms having 32,768, 65,536, and 131,072 data points, respectively. These results were also compared to those obtained for spectra measured at 2 cm\(^{-1}\), which were generated from interferograms containing 16,384 data points.

As can be seen in Table 7-4, the accuracy of the measurements was not affected by the number of data points per spectral element. However, the magnitude of the error in the measurements was related to the number of interferogram data points used to generate the spectra. On average, the error in the CLS analysis decreased by a factor of 1.4 each time the number of data points used to process the interferogram was doubled.

7.5.2.2 Effect of S/N on the CLS Analysis

Synthetic noise was added to 1-cm\(^{-1}\) spectral mixtures containing 100 ppm acetone, 100 ppm methylene chloride, and 0 to 500 ppm ethanol at levels corresponding to 1, 5, 10, and 25% of the maximum absorbance value in each spectrum.

**TABLE 7-4. THE EFFECT OF ZERO FILLING ON THE CLS ANALYSIS**

<table>
<thead>
<tr>
<th>Conc. (ppm)</th>
<th>1-cm(^{-1}) Resolution</th>
<th>1-cm(^{-1}) Resolution</th>
<th>1-cm(^{-1}) Resolution</th>
<th>2-cm(^{-1}) Resolution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>4 × Zero Fill</td>
<td>2 × Zero Fill</td>
<td>No Zero Fill</td>
<td>No Zero Fill</td>
</tr>
<tr>
<td>0</td>
<td>Below MDL</td>
<td>Below MDL</td>
<td>Below MDL</td>
<td>Below MDL</td>
</tr>
<tr>
<td>125</td>
<td>121.40 (8.49)</td>
<td>119.65 (11.89)</td>
<td>124.53 (17.24)</td>
<td>119.85 (22.47)</td>
</tr>
<tr>
<td>250</td>
<td>248.33 (9.35)</td>
<td>248.87 (12.87)</td>
<td>252.07 (18.15)</td>
<td>252.18 (22.53)</td>
</tr>
<tr>
<td>375</td>
<td>379.85 (9.58)</td>
<td>368.40 (13.83)</td>
<td>380.78 (17.94)</td>
<td>373.87 (24.27)</td>
</tr>
<tr>
<td>500</td>
<td>500.81 (10.29)</td>
<td>509.94 (14.73)</td>
<td>492.42 (20.83)</td>
<td>509.24 (28.90)</td>
</tr>
</tbody>
</table>
Spectral mixtures containing 100 ppm acetone, 100 ppm methylene chloride, and 500 ppm ethanol at each noise level are shown in Figure 7-12. In these mixtures, the average error in the CLS analysis was found to be directly proportional to the percentage of noise added to the spectrum. (See Table 7-5.)

<table>
<thead>
<tr>
<th>% Noise</th>
<th>Average Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.07</td>
</tr>
<tr>
<td>1</td>
<td>0.96</td>
</tr>
<tr>
<td>5</td>
<td>4.78</td>
</tr>
<tr>
<td>10</td>
<td>9.55</td>
</tr>
<tr>
<td>25</td>
<td>23.89</td>
</tr>
</tbody>
</table>

Figure 7-12. Spectra of Synthetic Mixtures of 100 ppm Acetone, 100 ppm Methylene Chloride, and 500 ppm Ethanol Measured at 1-cm⁻¹ Resolution with (A) 0, (B) 1, (C) 5, (D) 10, and (E) 25% Noise Added.
7.5.2.3 Mixture of Methylene Chloride and Nitrous Oxide

Synthetic mixtures of methylene chloride and nitrous oxide were generated by adding reference spectra of nitrous oxide corresponding to concentrations of 12.5, 25, 37.5, 50, 75, and 100 ppm to reference spectra of 100 ppm methylene chloride. The 0.25-cm\(^{-1}\) reference spectra and the spectra of the synthetic mixtures corresponding to 50 ppm of nitrous oxide with both 0 and 100 ppm of methylene chloride recorded at 0.25-, 0.5-, and 1.0-cm\(^{-1}\) resolution are shown in Figure 7-13.

The spectrum of nitrous oxide exhibits sharp bands that are resolved at 0.25 cm\(^{-1}\), but are not as well resolved at 0.5-cm\(^{-1}\) resolution. These bands become a broad continuum in the 1.0-cm\(^{-1}\) spectrum. At first glance, one would expect the CLS analysis to perform better for the 0.25-cm\(^{-1}\) spectra, in which the N\(_2\)O bands are fully resolved. However, this is not the case in these mixtures. When analyses are performed for

![Figure 7-13](image_url)

**Figure 7-13.** Reference 0.25-cm\(^{-1}\) Spectra of (A) N\(_2\)O and (B) Methylene Chloride and Spectra of Synthetic Mixtures of 50 ppm N\(_2\)O and 100 ppm Methylene Chloride Measured at (C) 0.25-, (D) 0.5-, and (E) 1.0-cm\(^{-1}\) Resolution.
N$_2$O over the entire band envelope from 1231 to 1329 cm$^{-1}$, N$_2$O is not detected at concentrations less than 75 ppm in the 0.25-cm$^{-1}$ spectra. (See Figure 7-14.) However, when the mixture was analyzed for N$_2$O by using the methylene chloride region from 1243 to 1292 cm$^{-1}$, N$_2$O was accurately quantified with a high precision. Similar results were obtained for the 0.5-cm$^{-1}$ resolution spectra. In contrast, the mixtures recorded at 1.0-cm$^{-1}$ resolution could be analyzed successfully over both regions. At 2-cm$^{-1}$ resolution, the CLS analysis performed best over the N$_2$O region. Apparently, in multicomponent mixtures, the CLS algorithm does not perform well in regions where one or more components exhibit only baseline noise. This effect seems to be amplified in higher resolution measurements of compounds with sharp spectral features, such as N$_2$O. These results indicate that for spectra with sharp features the CLS should be performed over a narrow range that contains absorbing features.

7.5.2.4 Conclusions and Recommendations Based on Case Study

The following conclusions regarding resolution requirements in long-path FT-IR monitoring can be drawn from this simulated study using well-characterized spectral data sets.

In spectra with overlapping sharp features, the CLS algorithm can accurately quantify target analytes, even when the bands used for analysis are not fully resolved. However, a failure to identify all of the overlapping components in a mixture can result in a bias and an increase in the error in the CLS analysis. Thus, the real value in performing higher resolution measurements might be to facilitate identifying the species present to be included in the CLS analysis set.

Figure 7-14. Concentration Calculated from CLS Analysis vs. Known Concentration for N$_2$O/Methylene Chloride Mixtures Measured at 0.25-cm$^{-1}$ Resolution. The (■) represents a value obtained during analysis over the methylene chloride region, and the (♦) represents a value obtained during analysis over the N$_2$O region.

In the case of spectra with overlapping broad features, such as those found in acetone, methylene chloride, and ethanol, the accuracy of the CLS analysis is not affected by the resolution setting. However, the
magnitude of the errors in the CLS analysis is related to the number of data points per wave number in the analyte spectrum. Therefore, the errors in the CLS analysis will increase with decreasing resolution unless additional zero filling, or some other means, is used to increase the number of data points in the spectrum. However, the use of zero filling or interpolation to indiscriminately increase the number of data points in the spectrum is not recommended, because interpolated data points do not contain independent information. In these mixtures, the errors in the CLS analyses were also found to increase proportionally with increases in the noise level.

The results from the CLS analysis of spectral mixtures with overlapping broad and sharp bands, as was the case with methylene chloride and nitrous oxide, were not as straightforward to interpret. When analyzing for nitrous oxide in spectral regions where methylene chloride did not exhibit any absorption bands, the CLS algorithm performed better at lower resolutions. In regions where the two compounds exhibited overlapping spectral features, comparable results were obtained for measurements taken at 0.25-, 0.5-, and 1.0-cm⁻¹ resolution.

In summary, resolution requirements will vary for different target compounds and sampling conditions. In field measurements these requirements will depend on several factors, such as path length, concentration of the target compounds, and the presence of interfering species. Although the simulated studies described here do not provide a definitive answer regarding the resolution question, similar studies using target analytes and possible interfering species should be performed prior to field studies to establish guidelines for data acquisition and analysis.

7.6 General Conclusions and Recommendations

As stated in the introduction of this chapter, there is currently no consensus as to what resolution is generally applicable in FT-IR long-path, open-path monitoring. A spectral resolution of 0.125 cm⁻¹ is required to fully characterize the spectra of atmospheric CO₂ and water vapor. Spectra taken along a 150-m path show that there are significant differences in the water vapor spectra measured at nominal resolutions of 2, 1, and 0.5 cm⁻¹. The effect of these differences on the computer-assisted quantitative analyses for target pollutants has not, however, been fully examined for long-path FT-IR measurements. In previous studies, Strang et al. (1989) have shown that for several organic vapors a resolution of 8 cm⁻¹ is sufficient to obtain quantitative results over a short path if a CLS program is used. In contrast, Spellicy et al. (1991) have presented theoretical results that suggest that the FT-IR spectra of small molecules with very fine spectral features will obey Beer’s law at only high resolution (0.01 cm⁻¹) and at very low concentrations. Recently, Marshall et al. (1994) and Griffiths et al. (1995) have indicated that 1- to 4-cm⁻¹ resolution, or possibly a lower resolution, is adequate for measuring certain VOCs using CLS and PLS multicomponent analysis programs.
Clearly, there is much fundamental research that must be done to "resolve" the resolution question. Experiments similar to those done by Strang et al. (1989), Strang and Levine (1989), Marshall et al. (1993), and Griffiths et al. (1993) should be conducted over a long, open path for the hazardous air pollutants stipulated under the Clean Air Act Amendments of 1990. Measurements of these compounds should be taken at different resolutions, concentrations, and path lengths to determine the optimum experimental conditions for obtaining the best S/N and detection limits. Most likely, the optimum resolution will be different for the various compounds. However, at least minimum resolution requirements could be determined.

Although the question of what resolution should be used in FT-IR long-path, open-path monitoring has not been answered, the reader should have an appreciation for the factors related to resolution that affect spectral measurements. Instrument manufacturers and software vendors have made great strides in simplifying the use of FT-IR instruments. Most FT-IR software is menu driven and some instruments can be operated at the push of a button. Although these developments facilitate the collection of FT-IR data, they also allow data to be collected without a knowledge of the principles behind the measurement. Analysts working in this field must be aware of the effects of different instrumental parameters on the measured spectrum. Grasselli et al. (1982) have published criteria for presenting spectra from computerized IR instruments, with an emphasis on FT-IR measurements.

The authors established recommendations and guidelines for reporting experimental conditions, instrumental parameters, and other pertinent information describing the acquisition of FT-IR spectra. These guidelines should be followed when reporting FT-IR data.

7.7 Guidance for Selecting Resolution and Related Parameters

In this section, general criteria and guidelines are suggested for choosing the optimum resolution for acquiring spectral data. The choice of resolution and related parameters, such as apodization and zero filling, to be used for data collection will be determined by several factors. As stated before, there is no consensus as to what the optimum parameters should be. The parameters need to be optimized for the specific experiments planned, taking into consideration the goals of the monitoring study. The following guidelines should be taken into account when choosing the optimum instrumental parameters.

1. Consider the bandwidths of the absorption features used to analyze for specific target compounds. If the absorption bands of the target compounds are relatively broad, there may be no need to acquire high-resolution spectra. When this is the case, no additional information will be gained, and the measurements will have a poorer S/N and will require longer data collection, computational times, and larger data storage space. The analyst must be aware, however, that the spectral features of atmospheric constituents
such as CO₂, H₂O, and CH₄ can be completely resolved only at a resolution of 0.125 cm⁻¹. Because these compounds are in every long-path spectrum and often overlap with the target analyte, access to high-resolution data may be required to develop the analysis method. There is some thought that the real advantage of high-resolution spectral data is the ability to visualize the spectral features and to identify interfering species. This information can then be used in developing the analysis method.

2. Determine if interfering species are present. If the comparison method or scaled subtraction is used for quantitative analysis, the resolution should be sufficient to separate spectral features of the target compounds from those of interfering species. For example, in the case of toluene the absorption band used for analysis at 1031 cm⁻¹ is relatively broad. At first, this would indicate that a low-resolution measurement would be sufficient. However, this band overlaps with bands due to atmospheric water vapor and CO₂. Therefore, a higher resolution measurement is required to separate the toluene from those of interfering species.

3. Acquire reference spectra of the target compounds. If the specific target compounds are known prior to beginning the monitoring study, reference spectra of the compounds of interest should be recorded at various resolutions. This can be accomplished by collecting a reference spectrum at the highest resolution setting on the instrument, and then processing the data by using the required number of Fourier transform data points for each desired resolution. When this method is used, only one spectrum has to be collected. By comparing the spectra processed at different resolutions, the operator can determine the lowest resolution measurement that still resolves the spectral features of interest. This resolution setting should be used as a starting point for future measurements. If this is not possible, the operator should consult reference libraries to help determine the optimum resolution setting required to characterize the target analyte.

4. Develop calibration curves of the target compounds. If an inadequate resolution is used, the relationship between absorbance and concentration will not be linear. This relationship is also affected by the apodization function. Calibration curves covering the concentration range of the target compounds expected in the ambient measurements must be developed at different resolutions and with the use of different apodization functions to determine the optimum settings. If the compound of interest does not respond linearly with respect to concentration, a correction curve will need to be fitted to the data and used in the quantitative analysis package.

5. Use the same parameters to collect field spectra as were used to record the reference spectra. If spectra from a commercial or user-generated library are to be the reference spectra for quantitative analysis, then the parameters that were used to generate those reference spectra should be used to collect the field spectra. Otherwise, errors in the measurement will occur.
6. **Determine that the instrument is producing data at the specified resolution.**

The following factors should be considered here: (a) that the operator has selected the proper parameters, and (b) that the instrument is operating to the manufacturer's specifications and that the manufacturer's specifications are a true indication of the capabilities of the instrument.

a. Most software packages allow the resolution to be selected from a menu. The software then automatically sets the proper parameters to collect data at the selected resolution. Therefore, there is very little opportunity for operator error. In older versions of software that are not menu driven, but instead require entering line commands, many of the parameters affecting resolution, such as the number of data points used for the Fourier transform, must be entered manually. In this case, the operator must know, and enter correctly, all of the proper parameters, and there is a greater chance of error.

b. If the instrument is not producing data of the selected resolution, it is also possible that the instrument is malfunctioning or that the manufacturer overstated the capabilities of the instrument. The following procedure can be used to determine if the instrument is producing data at the specified resolution. There is a cluster of absorption bands between 1008 and 1020 cm\(^{-1}\) due to water vapor that can be used to verify the resolution of the FT-IR monitor. There is a doublet centered at 1010.5 cm\(^{-1}\), a single band at 1014.2 cm\(^{-1}\), and a pair of bands at 1017.5 and 1018 cm\(^{-1}\). The singlet at 1014.2 cm\(^{-1}\) has a theoretical bandwidth of approximately 0.3 cm\(^{-1}\) (USF HITRAN-PC [University of South Florida 1993]). This band is a good reference band for determining the actual resolution measured by a medium- or low-resolution spectrometer. For example, if this band is measured at an instrument setting of 0.5 cm\(^{-1}\), the FWHH should be 0.5 cm\(^{-1}\). If measured with an instrument capable of achieving a higher resolution, for example 0.25 cm\(^{-1}\), the FWHH should be the theoretical value of 0.3 cm\(^{-1}\). The doublet centered at 1010.5 cm\(^{-1}\) is just resolved at 0.5-cm\(^{-1}\) resolution, but is not resolved at 1-cm\(^{-1}\) resolution. The theoretical bandwidth of each of these two absorption bands is approximately 0.1 cm\(^{-1}\) (USF HITRAN-PC), which makes them good reference peaks for instruments capable of measuring at a resolution of 0.125 cm\(^{-1}\). The two peaks at 1017.5 and 1018 cm\(^{-1}\) are resolved at 0.125 cm\(^{-1}\), but not at lower resolutions. If the instrument is not performing to specifications, there is most likely an alignment problem with the interferometer or source optics. Unless the operator is trained to perform this alignment, a representative from the manufacturer must service the instrument.

The bands centered at 1014.2 cm\(^{-1}\) are a good test of resolution, but they are not as sensitive to misalignment in the
interferometer. Other bands that may also be used are the 2169-cm$^{-1}$ band of CO and the HDO doublet centered at approximately 2720 cm$^{-1}$. These bands at shorter wavelength (higher wave number) are more sensitive to interferometer misalignment and can also be used to determine the stability of the interferometer.

These are general guidelines to be used when choosing instrumental parameters to collect data. In reality, the user’s choice of parameters that can be actually used may be limited by either the specifications of the spectrometer or by the software. For example, one software package supplied with FT-IR long-path, open-path systems allows only triangular apodization with no additional zero filling for processing the interferogram with the menu-driven commands. These parameters cannot be changed unless the user has the capability of editing the software code. As the resolution requirements of long-path, open-path FT-IR monitors become better defined, the manufacturers will most likely produce instruments and software to meet those needs.

7.8 References


University of South Florida. 1993. USF HITRAN-PC, University of South Florida, Tampa, FL.
Chapter 8
Quality Assurance and Quality Control

SUMMARY

The topics and specific points of emphasis discussed in this chapter include the following:

- The need for a quality assurance (QA) project plan
- QA project plan categories and relevant EPA documents
- A general format for a 16-point QA project plan
- A discussion of specific quality assurance and quality control (QA/QC) issues related to FT-IR long-path, open-path monitoring
- Portions of an approved QA project plan
- A case study presenting QA data collected over a two and one-half month period
- Recommendations of procedures to be included in a QA program

8.1 Introduction and Overview

For open-path FT-IR spectrometry to become an accepted method for environmental monitoring, QA procedures must be established. While QA issues have been addressed (Kricks et al. 1992; Russwurm 1992a,b; Weber et al. 1992; Kagann et al. 1994), there is currently no consensus regarding the proper QA procedures required to validate open path FT-IR data. In fact, there is very little information in the literature that addresses the quality of the data generated by FT-IR long-path, open-path systems. For example, when error bars are given, they are often merely stated, and no discussion of how they were derived is supplied. As the FT-IR long-path technique begins to be used for routine monitoring, this approach will not be satisfactory, and a more extensive QA plan must be developed.

The development of and adherence to a QA project plan requires the operator to consider exactly how the data generated by an FT-IR long-path, open-path monitoring program will be obtained, processed, interpreted, and used. When implemented properly, the QA plan will alert the operator if the instrument is not functioning properly or is generating erroneous data, and it will contain recommendations for the corrective action to be taken. The various levels of QA project plan designs apply to programs ranging from research and development
programs to routine monitoring programs that must produce legally defensible data. This chapter covers some of the points that will have to be addressed for any QA program.

The general points that must be addressed for any QA program are given in this chapter. These points are drawn from specific documents that address QA requirements for data obtained for the U.S. Environmental Protection Agency. Points of emphasis discussed in these documents include the following: project description, organization, and responsibilities; QA objectives; site selection and sampling procedures; sample and data custody; calibration procedures and frequency; analytical procedures; data reduction, validation, and reporting; internal quality control checks; performance and systems audits; preventive maintenance; calculation of data quality indicators; corrective action; quality control reports to management; and references.

These, and other, items are addressed with respect to FT-IR long-path, open-path monitoring in this chapter. In addition, portions of the QA project plan for a 1989 Superfund Innovative Technology Evaluation study in Delaware are presented as an example for use in FT-IR long-path, open-path monitoring. Also, a case study involving the acquisition of QA data over a two and one-half month period is presented in Section 8.3, and recommendations for procedures to be included in a QA program are given in Section 8.4.

8.2 Project Plan Categories

The U.S. Environmental Protection Agency has published a document that defines four different categories of QA project plans, as described in Section 8.2.1. The program with the least requirements is a research and development program (Category IV), whereas a program that produces data by routinely monitoring the atmosphere (Category I) has the most. When considering programs that are specifically designed to obtain data for the U.S. Environmental Protection Agency, it is convenient to refer to two documents that describe QA project plans: (1) Preparing Perfect Project Plans: A Pocket Guide for the Preparation of Quality Assurance Project Plans and (2) Preparation Aids for the Development of RREL Quality Assurance Plans (U.S. Environmental Protection Agency 1989, 1991). Actually, there are four parts to the second document, one for each of the four categories, and each has its own document number.

Some of the features of the items that need to be addressed in the Category I project plans are covered in Sections 8.2.1 and 8.2.2. It is intended that these sections be a paraphrasing of the two documents listed above. People who have to deal with these issues must obtain a copy of the documents and follow the guidance given there. To give some specificity to the various points of a QA project plan, the QA project plan used for a recent program conducted by EPA is used as an example. The program entailed taking data with an FT-IR open-path monitor in an industrial complex and comparing that data to
data obtained by a canister technique according to method TO-14 (Russwurm and McClenny 1990). Funding was allocated for 10 days of measurements with the FT-IR instrument in the field.

8.2.1 Category Definitions

- Category I projects are those that are designed so that their results can be used directly, without additional support for compliance or other litigation. As such, they must be able to withstand legal challenge and therefore have the most rigorous and detailed requirements. These projects are critical to the goals of the U.S. Environmental Protection Agency.

- Category II projects are those whose data complement other projects. When combined with the output from other projects, these data can be used for rule making or policy making.

- Category III projects are those producing data that allow the evaluation and selection of basic options for use in feasibility studies.

- Category IV projects are those that are associated with research and development projects. The results are used to assess the basic or underlying assumptions or suppositions of other work. Because of the nature of these projects, they have the minimum number of items that need to be addressed in a QA program.

These categories are intended to be fairly general and broad, and any project must fit into one of the four. The number of items that must be addressed for each of the categories is 16, 13, 12, and 6 for Categories I through IV, respectively. The project described above for acquisition of data for comparison purposes and referred to as an example in Sections 8.2.2.1 through 8.2.2.16 was considered to be a research program, and therefore it fell into Category IV.

8.2.2 Category I Points to Be Addressed

The 16 items that must be addressed for the QA project plan in this category are listed below.

1. Project description
2. Project organization and responsibilities
3. QA objectives
4. Site selection and sampling procedures
5. Sample custody
6. Calibration procedures and frequency
7. Analytical procedures
8. Data reduction, validation, and reporting
9. Internal quality control checks
10. Performance and systems audits
11. Preventive maintenance
12. Calculation of data quality indicators
13. Corrective action
14. Quality control reports to management
15. References
16. Other items

These 16 items are discussed briefly below.
8.2.2.1 Project Description

The most important feature of the project description is that a person who is unfamiliar with the project, but is familiar with the technology, must be able to understand this section.

For the example project, the following items were included in this section.

- An in-depth discussion of the comparison program. The primary aspect of this was to relate in detail how the data would be taken. The FT-IR instrument is a long-path monitor, and the canister technique is a point monitor. Thus, it was decided to transport the canister along the path while the FT-IR monitor was acquiring data. Topics that had to be included in the QA project plan were the number of traversals along the path and the number of scans the FT-IR spectrometer would make.

- A brief description of the FT-IR technique and the canister technique. The techniques were described, and appropriate documents for each technique were referenced.

8.2.2.2 Project Organization and Responsibilities

This section must describe the relationships among all of the people connected with the project, including the QA manager, and give their responsibilities. It should be noted that, somewhere in the organization conducting the program, there should be an autonomous QA representative. Most organizations have this function removed from the technical staff and under the jurisdiction of an administrative manager. The primary function of this person is to act as final arbiter for any disputes about the QA aspects of the program. Therefore, he should not be administratively connected with the technical program.

For the project being outlined as an example, the management and administrative staff consisted of U.S. Environmental Protection Agency staff and staff from two contractors. The following descriptions were therefore included in this section.

- All the principal people involved and their duties
- The relationship of one person to the other concerning decision-making responsibilities
- The lines of communication among the project personnel

All lines of communication among the various project personnel, including any subcontractors, must be described. If no subcontractors are used, it is sufficient to simply state that fact.

8.2.2.3 QA Objectives

This section of the QA project plan must cover items such as detection limits,
precision, accuracy, completeness of the data, representativeness of the data, and comparability of the data. There must be a discussion of the impact of not meeting these objectives and how these indicators will affect the legal defensibility of the data.

Inadequately addressing these items is probably the most frequent cause of the rejection of a QA project plan.

Three items detection limits, precision, and accuracy must be numerically defined as QA objectives. Completeness of the data defines what percentage of the total number of possible data points that are available under the sampling schedule are expected to be captured. Representativeness of the data implies how well the acquired data account for the variability of the real situation. Representativeness of the data concerns itself with sampling. Comparability of the data indicates how well the data can be compared to that taken with other instrumentation.

A portion of the example field program (Russwurm and McClenny 1990) was designed to determine how to best make measurements of the detection limits, the accuracy, and the precision. Therefore, no specific definitions were required. At the present time there are no generally accepted definitions for these quantities for FT-IR open-path monitors.

For the completeness of the data, the following procedure was used. A 10-h working day was assumed, and 2 h were allocated for setup and warm-up time of the instrument. The data-taking period was assumed to be 0.5 h for each spectrum. Therefore, it was anticipated that 16 spectra a day would be taken. During the study, it was noted that the concentration of the gases being monitored was changing at a much higher rate than could be accommodated by the 0.5 h allocated to each data spectrum. Therefore, the sampling schedule was altered to account for the change. It should be noted that this is certainly an appropriate response to changing conditions. The QA plan is a guide for subsequent operations and not something that is unchangeable. All changes must be recorded, and the rationale for the change must be presented.

A large section of the plan concerned itself with the representativeness and comparability of the data. The reason that the canisters had to be transported along the path was precisely to satisfy the requirements of these two items.

8.2.2.4 Site Selection and Sampling Procedures

In addition to a physical description of the site, the rationale for selecting any individual site must be presented. This discussion must address how the site will allow the data objectives to be met.

The sampling portion of this section must address the scientific and regulatory (if any) objectives that the sampling protocol allows. It must also address how any calibration samples are to be obtained and
delivered to the system. In the case of the FT-IR system, if a gas sample cell is to be used, then its preparation and use must be addressed in this section.

Site selection and sampling procedures played an important role in the example study. Several months before the field portion of the example study, a search for suitable sites was started. The general area of the study had been selected, but the individual sites and FT-IR paths needed to be chosen. Therefore, this section of the example QA plan contained the following descriptions for the site selection process and sampling procedures.

- Background information on site selection. During a number of trips to the general area, various land owners were contacted, and several sites were selected as usable for the study.

- Predominant requirements for the site. Proximity to the source, an unobstructed path length of up to 300 m, a path that could be used to transport the canister while the FT-IR monitor was taking data, and a site that was safe for the operators.

- Procedure for using a QA gas cell. A description of using a short cell filled with a high concentration of gas for calibration purposes was described here. The use of this cell was also intended for determining the precision of the instrument.

8.2.2.5 Sample Custody

This section must present complete sample custody procedures and personnel responsibilities in handling samples. Because the FT-IR data are stored on disk, all procedures for ensuring the integrity of the data on the disk and the legal defensibility of those procedures must be addressed.

For the example program, no sample custody procedures were required because it was a research program.

8.2.2.6 Calibration Procedures and Frequency

The FT-IR open-path monitor is not calibrated in the classical sense. That is, a sample of known concentration is not presented to the instrument for measurement. During FT-IR sampling, the absorbance values in all the spectra obtained at various wave numbers for the specific gases are always compared to the absorbance values of the reference spectra. These reference spectra are made with pure samples of the gas. Production of reference spectra is a formidable task, and few laboratories are equipped for such an undertaking. Because of this, only a limited number of spectral libraries that can be used for quantitative analysis exist.

Although measurements of the precision can be made by using a short cell filled with a pure sample of the gas, this is not routinely done at the present time. There
is currently no agreed-upon procedure for the use of such a cell.

The project being used as an example did not require that the system be calibrated. However, a written procedure for using a short cell to make precision measurements was part of the QA plan.

8.2.2.7 Analytical Procedures

For the most part, unless other techniques are to be used along with the FT-IR system, there is nothing to address in this section. The analytical procedures that are to be used to determine the concentrations of any short-cell gas mixtures would have to be addressed here.

In the QA project plan being used as an example, the procedures for preparing the short cell and the gas mixtures were given here. The cells are generally filled with a pure gas sample and then backfilled with nitrogen so that the total pressure is 1 atm. All the apparatus used for this procedure was described in this section.

This section also contained a description of the procedure to be used for the analysis of the canister samples. This was a brief description, but it referenced the TO-14 procedure manual. The entire QA procedure for that portion of the effort did not have to be presented in the plan, but it had to be referenced. If one had not been available, then it would have had to be written and given in this section.

8.2.2.8 Data Reduction, Validation, and Reporting

The data reduction procedures must be discussed in this section. This includes the least-squares regression analysis, or other multicomponent analysis method, if it is to be used. All statistical methodology that is used as an aid to data interpretation must be described here. This section can also include sample calculations. All procedures to be used for flagging the data and removing outliers from the data set must be stated in this section. The flow of data and the procedures that will be used to transfer the data from where it is generated to the end user must also be described.

The QA project plan being used for discussion purposes included an analysis of the spectral data with a classical least squares algorithm. The actual process of recording the interferogram, performing the Fourier transform, and the least-squares analysis was not discussed in detail. The published papers that describe these techniques were referenced. The process of selecting the wave number range for analysis was discussed, and the wave number regions that were used were listed in this section of the QA project plan.

The TO-14 procedure for data reduction also was referenced, but the actual details were not presented in the plan.

The major portion of this section dealt with the comparison of the data which was to be done by regression techniques. At
various stages in the program, two canister samples were taken simultaneously, and the comparison procedure for these data was discussed.

Validation of the data was described as the process of reviewing the study logbooks to make sure that no unwanted contamination of the samples had occurred. For example, one sample was invalidated because an automobile had followed the person transporting the canister along the path.

8.2.2.9 Internal Quality Control Checks

This section is designed to determine what internal QC checks are to be made. It must also cover why these checks are necessary and how they will help to achieve the data quality objectives. For example, this section would describe the use of control charts that show the peak-to-peak readings of the single-beam spectrum at various path lengths and with varying amounts of water vapor.

At the time of the example study, there were no requirements for internal QC measurements written into the plan. A portion of the study was designed as an attempt to develop procedures for making these measurements.

8.2.2.10 Performance and System Audits

System audits are generally done before any data are taken. They are designed to answer questions about the proposed procedures and sampling protocols to be used in the program. The performance audits are designed to determine whether the instrument is operating as it was described in the other sections of the study’s QA project plan. Both of these audits are generally done by people who are not associated with the daily operation of the instrument.

There were no provisions for either system or performance audits for the study being used as an example.

Although it is easy to envision the system audit for the FT-IR instrument, the performance audit of the instrument would be much more difficult to do at this time. This is usually done by using the instrument to measure a known concentration of gas and determining the response. There has been no systematic development of a procedure for doing this for the FT-IR systems to date. Currently, not all commercially available systems have provisions for putting a short cell in the beam.

8.2.2.11 Preventive Maintenance

This section requires a description of the preventive maintenance procedures and the schedule on which they will be performed. The time involved in performing preventive maintenance will affect the total amount of data that can be taken, and this must be accounted for in the description of the completeness requirements.

In the example QA project plan, no preventive maintenance schedule was given
because the study was a short-term intensive program. Over the 10-day field program, no maintenance of the instrument was required.

8.2.2.12 Calculation of Data Quality Indicators

This section must contain detailed procedures that are to be used for determining the data quality. It must include all the statistical routines that are to be used. Items that must be addressed are precision, accuracy, outliers, etc.

Currently, no generally accepted definitions for precision and accuracy are available for the FT-IR open-path technique.

8.2.2.13 Corrective Action

The corrective action portion of the QA project plan is a set of contingency plans that try to address "what if" questions. These corrective action plans serve as a check for the general tendency to want to perform quick fixes of equipment to get as much data as possible. This is certainly the case with short-term field programs, but it is generally better to take the instrument off line and repair or adjust it properly. The corrective action plans should describe how this is to be done and specifically what criteria will be used to make the judgement as to when to discontinue data collection and shut down the instrument for repair. This section was not a requirement for the example QA project plan.

8.2.2.14 Quality Control Reports to Management

This section must state what reports will be transmitted to whom and when they will be transmitted. There should be a description of the contents of each report, and all the QA/QC data that must be included in each report should be stated in this section of the QA project plan.

The QA project plan for the example study did not address this requirement because there was no consistent QA data generated in the program.

8.2.2.15 References

When references related to the present program are available, they should be included in the QA project plan. For example, as mentioned in Sections 8.2.2.1 and 8.2.2.7, appropriate documents were referenced for the canister technique used in the example QA project plan.

8.2.2.16 Other Items

The QA project plan contains information that the principal investigator will need at various points of the program. Because of this, it is somewhat personalized, and this section can include any other items that are considered important to the program.

There were no other items included in the QA project plan for the example study.
8.3 Case Study: QA Data Collected over Two and One-Half Months at a Semipermanent Field Site

The study described here was designed to evaluate the stability of a long-path FT-IR system and to determine the precision and accuracy of the concentration measurements (Thompson et al. 1994). The following criteria were used to assess the stability of the instrument: electronic noise, the magnitude of the return signal, the RMS baseline noise, and the repeatability of the position and full width at half height (FWHH) of selected absorption bands. Ambient concentrations of CH₄, N₂O, and CO were measured to test the use of these data for determining the precision and accuracy of the FT-IR open-path monitor. Measurements were made daily over two and one-half months, from November 1993 to mid-January 1994.

Spectral data were acquired by using a monostatic FT-IR monitor. Each spectrum consisted of 64 co-averaged scans recorded at a nominal 1-cm⁻¹ resolution. Triangular apodization was used. The collection of each spectrum required approximately 5 minutes. A spectrum was taken every 15 minutes. Single-beam spectra were typically acquired over a 7- to 8-h time period. Absorption spectra were created by ratioing the single-beam spectra to a synthetic background spectrum generated from a 2048-scan single-beam spectrum recorded over the 414-m path. This background spectrum was recorded at the beginning of the experiment and was used throughout the study. The data were analyzed by using a CLS software package and reference spectra from a commercial library.

The site is located near I-40, one of the main traffic arteries for the Research Triangle Park, NC, area. The instrument was kept in a climate-controlled shed, which is heated during the winter months. The total path length was 414 m, and it extended over an open, grassy field and a small parking area with very limited traffic. The beam path rose from about 1.8 m to 12.8 m above the ground as it was directed from the FT-IR spectrometer to the retroreflector array, which was mounted on a tower.

The instrumental electronic noise was measured each morning before the detector was cooled with liquid nitrogen. This signal typically ranged between 600 and 620 counts with the instrument in the single-beam mode. Shortly after the detector was cooled, the instrument was aligned and the maximum return signal was recorded. The return signal was recorded again (without realignment) around noon to check the stability of the signal. On clear days the single-beam return signal ranged from 10,500 to 13,500 counts. (See Figure 8-1.) Certain atmospheric conditions caused the return signal to vary from day to day. For example, the return signal dropped by 20-30% during fog. On some mornings, when the humidity was close to or below the dew point, condensation or ice formed on the retroreflector, resulting in a lower return signal in the early morning measurement. As the condensation evaporated, an increase in return signal...
counts was measured. To remedy the problem of condensation, a heat lamp was mounted on the tower and directed at the retroreflector. After the heat lamp was installed on December 10, the noon and early morning return signals were nearly the same. The use of the heat lamp did not cause an increase in noise or detected IR signal.

The RMS baseline noise measured over 26 days is illustrated in Figure 8-2. The baseline noise was determined by collecting two back-to-back, 64-scan, co-added spectra. One spectrum was ratioed against the other to obtain an absorption spectrum. The RMS noise (in absorbance units) was calculated over three spectral regions: 980-1020, 2480-2520, and 4380-4420 cm⁻¹. During operations when condensation did not form on the retroreflector, the baseline noise was on average approximately 2 × 10⁻⁴ for the 980-1020-cm⁻¹ region, 2.5 × 10⁻⁴ for the 2480-2520-cm⁻¹ region, and 9 × 10⁻⁴ for the 4380-4420-cm⁻¹ region. In these measurements, the 980-1020-cm⁻¹ region included water vapor bands. For a true measurement of instrument performance that is not influenced by temporal changes in water vapor concentration, it is recommended that the region from 968 to 1008 cm⁻¹ be used. During measurement periods when condensation formed on the retroreflector, the baseline noise for these regions increased to 9.7 × 10⁻⁴, 5.5 × 10⁻⁴, and 2.9 × 10⁻³, respectively.
The wave number stability of the instrument was determined by monitoring the peak position and the FWHH of the water vapor singlet at 1014.2 cm\(^{-1}\). Band positions typical of data collected at the beginning, in the middle, and near the end of the study are depicted in Figure 8-3. No shift in the frequency was observed during this time period. Also, no shifts were observed in the 1-cm\(^{-1}\) spectra collected under a variety of weather conditions, including rain, freezing rain, sleet, snow, and low (single-digit) temperatures. To determine if the water vapor singlet at 1014.2 cm\(^{-1}\) broadened, a spectrum collected at the beginning of the experiment was subtracted from a spectrum in the middle and end of the experiment. No broadening was evident during the middle of the experiment; however, a slight broadening for some of the spectra at the end of the experiment was observed. The FWHH of the water vapor singlet in spectra taken during different atmospheric conditions was also examined. When a clear day spectrum was
subtracted from any of these spectra, no broadening was evident. It should be noted that short wavelengths (higher wave numbers) will be more sensitive to spectral shifts and changes in resolution. The HDO doublet centered at 2720 cm⁻¹, the CO band at 2169 cm⁻¹, or other water vapor bands in the higher wave number region can also be used to test for shifts and changes in resolution.

The feasibility of using ambient gases for accuracy and precision measurements was also investigated. Ambient concentrations of N₂O, CH₄, and CO were measured on a daily basis. Each morning between 0715 and 0930 the concentrations of these gases increased, then steadily decreased during the remainder of the day. However, concentrations of N₂O and CH₄ remained constant, approximately 250 ppb and 1.7 ppm, respectively. To determine whether the increases in concentration during the first 3 h of operation were due to an instrument effect or to the proximity of the site near a major highway, data were collected continuously for 36 h.

Figure 8-3. Repeatability of the Position of the Water Vapor Singlet at 1014.2 cm⁻¹ Measured on (A) November 10, 1993, (B) December 22, 1993, and (C) January 4, 1994.
Data were collected from November 17 at 0730 until 1730 on November 18. The CH₄ concentration data exhibited scatter during an early morning fog episode and decreased steadily during the day. (See Figure 8-4A.) A step in the CH₄ concentration measurement was observed when the liquid nitrogen in the detector was depleted at approximately 2345 on November 17. The CH₄ concentration value was 1.70 ppm just before the liquid nitrogen was depleted, increased to 1.9 ppm after liquid nitrogen was refurbished, and remained 10% higher compared to the previous levels. The concentration data for CO showed a similar, stepped increase.

The CH₄ concentration data exhibited irregular behavior during a 6-h period shortly after the detector Dewar was refilled with liquid nitrogen. To determine if the instrument was operating properly during this time period, the single-beam intensity at 987 cm⁻¹ was measured from archived spectra. The single-beam spectra had a lower intensity during the fog episode, then leveled off until the detector ran out of coolant. (See Figure 8-4B.) After this sudden drop, the

![Graph showing concentration levels over time](image_url)

Figure 8-4. Measurement of (A) Ambient Methane Concentration and (B) Single-Beam Intensity at 987 cm⁻¹ on November 17 and 18, 1993.
single-beam intensity returned to its original reading and remained relatively constant throughout the remainder of the experiment. This indicates that the instrument was working properly during the episode of high measured CH₄ levels.

One other observation during this time period concerned the effect of water vapor concentration on CLS analysis for CH₄. On November 17 a cold front moved through the area in the late evening, and the water vapor pressure dropped rapidly. Because the water vapor spectrum is used as an interfering species in the CLS concentration analysis for CH₄, the sudden change in water vapor pressure could have had an effect on the CH₄ concentration measurements. The relative concentration of water vapor along the path was determined by measuring the peak area of the absorption band at 1014.2 cm⁻¹. Likewise, the relative concentration of CH₄ was determined by measuring the peak area of the absorption band at 2998.8 cm⁻¹. This peak was chosen because the water vapor bands do not interfere with it. However, it was later brought to our attention that this CH₄ band actually overlapped nearly exactly with a water vapor band (W.F. Herget, ETG, personal communication). This might explain the dip in the peak area measured at 2998.8 cm⁻¹ during the time that the water vapor concentration was decreasing rapidly. Therefore, this band is not a good choice for this type of data analysis; the CH₄ bands at 2916.8 and 2927 cm⁻¹ do not overlap with water vapor and should be used instead. As shown in Figure 8-5, the relative water vapor concentration decreased rapidly when the front moved through the area. The relative CH₄ concentration increased during this period. Similar trends were observed between plots of the CH₄ peak area and CH₄ concentrations determined by the CLS concentration analysis. This indicates that the change in water vapor concentration did not greatly affect the CLS analysis for CH₄, and the fluctuations in the CH₄ concentrations were real.

The data collected in this study indicate that for this particular FT-IR monitor the return signal and baseline noise are repeatable and are instrumentally stable over extended periods, but are subject to variations due to weather conditions. The peak positions and the FWHHs of the water vapor singlet at 1014.2 cm⁻¹ were repeatable from day to day and were not affected by rain, freezing rain, snow, or single-digit temperatures. The variability in the concentrations of CH₄ limits its use in instrument stability studies and for accuracy and precision determinations. A step in the concentration measurements associated with the depletion and refurbishment of detector coolant is not yet understood. Future experiments with a QA cell and surrogate standards are planned to further investigate this effect.
8.4 Recommendations for Tests to be Included in a QA Program for FT-IR Long-Path Monitors

We are currently evaluating and developing procedures to determine the quality of data taken with FT-IR monitors. The following is an outline of criteria that we have used in a preliminary QA program to verify the performance of an FT-IR long-path system. Development of a QA plan for FT-IR monitoring should include, but not necessarily be limited to, these types of measurements. These tests are designed to determine that the instrument is operating properly and producing good data. Some of these issues
were discussed in Chapter 3 for the initial verification of instrument performance, but can be used for routine QA procedures as well. Other criteria for development of a QA plan, such as siting criteria or data chain-of-custody, should be addressed as warranted, but are not discussed here. These procedures were developed for a research and development program, but factors relevant to routine monitoring programs were also taken into consideration. Bear in mind that two separate issues must be addressed in a QA plan. One is whether or not the instrument is working properly. The other issue is if the method used for quantitative analysis is producing the correct results.

8.4.1 Noise Measurements

Measurements of two types of noise can be routinely taken, instrumental electronic noise and random baseline noise. Electronic noise is recorded before the detector Dewar is filled with liquid nitrogen. This small signal is indicative of the electronic noise of the system with no detector signal. This should remain relatively constant and typically contributes less than 0.25% of the total return signal. If some electrical component of the system is producing spurious noise, it will become apparent from this measurement.

Random baseline noise is measured by recording back-to-back spectra after the detector has been filled with liquid nitrogen. One spectrum is then ratioed to the other and the absorption spectrum is calculated. The result is a spectrum of the random system noise. The RMS or peak-to-peak noise in absorbance units can then be calculated from these spectra. These spectra should be acquired by using the instrumental parameters to be used during the analytical measurements. The baseline noise measurements should be taken in a spectral region that is devoid of absorption due to water vapor or other atmospheric gases. If not, changes in water vapor concentration over the measurement time will influence the magnitude of the noise calculations. Noise measurements can also be taken over the spectral region chosen for analysis of a target gas to give an estimate of detection limits.

For the baseline noise measurement it is best to record these two spectra back to back, as passage of time between the two spectra might also include changes in atmospheric conditions or concentrations of species in the path. Spectra taken at longer time intervals during the study can be ratioed in this manner to determine baseline stability or systematic noise.

8.4.2 Stability of Instrument

Several aspects related to the stability of the instruments can be measured. One is the repeatability of the noise measurements described above. These noise measurements should be taken daily and recorded on a control chart to alert the operator of any gross changes or trends in the deterioration of the baseline noise.

Another measurement to be taken daily, or several times during the day, is the return intensity. This can be measured either
as the single-beam intensity at a selected wave number region or the magnitude of the interferogram. At this time, the position of the zero peak difference of the interferogram should also be recorded, and the single-beam spectrum should be examined for evidence of system nonlinearity (see Chapter 3, Section 3.4). The single-beam intensity should be measured in different wave number regions to determine if the source characteristics have changed or the interferometer alignment has been altered. The high wave number (short wavelength) portion of the spectrum will be most sensitive to interferometer misalignment and will show a decrease in intensity relative to the other wave number regions if changes have occurred.

In addition to being dependent on the instrument's performance, the return intensity also depends on atmospheric conditions. For example, fog has a deleterious effect on the return signal. Thus, the atmospheric conditions must be noted when the measurement is taken. As with the noise measurements, the return intensity should also be plotted daily on a control chart. A decrease in return intensity could be related to a drop in the source intensity, misalignment of the external optics, misalignment of the interferometer optics, deterioration of the system optics, or a loss in the detector Dewar hold time.

The positions of selected absorption bands should also be recorded. In long-path measurements the water vapor singlet at 1014.2 cm⁻¹ works well for this purpose. The FWHH of this band can also be measured to determine the repeatability of the resolution of the system. The subtraction technique described by Russwurm (1992b) can be used to detect small frequency shifts or subtle changes in the instrument resolution. If the resolution of the system is deteriorating, and the band becomes broader than a spectrum recorded previously, the subtraction result will appear as an "M". Shifts in frequency will produce a derivative shape in the subtraction result. Note that the FWHH measurement and the spectral subtractions should be done on an absorption spectrum, and not on a single-beam spectrum or transmission spectrum.

As mentioned previously, bands in higher wave number regions actually may work better for this test, as they will be more sensitive to spectral shifts and changes in resolution. In addition to the water vapor bands centered at 1014.2 cm⁻¹, the HDO bands centered at 2920 cm⁻¹ can be used for this test.

All of the above measurements should be recorded on at least a daily basis and compared to existing data to establish that the instrument is performing properly.

8.4.3 Accuracy and Precision

The determination of accuracy and precision are not as straightforward as the tests used to determine if the instrument is performing properly. The concentrations of ambient gases, such as CH₄ or N₂O, can be used to a certain extent for these purposes.
This approach has the advantage that certain gases are always present in open-path spectra and no changes have to be made to the instrumental configuration to measure these gases. If the ambient concentrations of these gases are assumed to be constant, precision measurements can be made. For example, we have measured the concentration of N_2O continuously over a five-day period in the late spring to be within ±3.5% of the mean value. On the other hand, as discussed in Section 8.3, we have seen CH_4 concentrations increase by a factor of 2.5 in a short time during measurements taken in the late fall. Therefore, care must be taken to account for possible emission sources if ambient gases are used for QA purposes. Ambient gases can also be used to test for frequency shifts or changes in resolution as described above. The use of ambient gases for QA purposes can be used to estimate the precision of the measurement, but this approach does not really address the accuracy of the method.

The alternative approach to using ambient gases for QA data is to insert a short gas cell that contains surrogate standards into the beam. This has the disadvantage of an attenuated IR beam due to the transmitting and reflecting properties of the windows used in the cell. Thus, the performance of the instrument is somewhat degraded. However, this approach does have the advantage of having a known quantity of target gas in the path. Assuming this quantity is accurately known and it remains constant, accuracy and precision measurements can be made with a short cell technique. To date, however, no universally applicable technique using a short cell has been developed.

As discussed in Chapter 3, stray light in the monostatic configuration and background blackbody radiation in the bistatic configuration can affect the quantitative results. The signal due to stray light or background radiation must be subtracted from the sample spectrum prior to quantitative analysis. The stray light in the spectrometer of a monostatic instrument can be measured by blocking the return beam with some type of opaque and nonreflective material. In our experience the signal due to stray light has been relatively constant, provided no components of the system are changed. The background blackbody radiation of a bistatic system can be measured with the IR source off. This response will vary for different sites and can also change throughout the day. Therefore, this signal must be recorded on a more frequent basis. It is not obvious at this point, however, that simple subtraction adequately compensates for the effects of stray light or background radiation linearly over a range of absorption values. Thus, the accuracy of the system might be affected.

In addition to determining the accuracy and precision of the instrument, the accuracy and precision of the method used for quantitative analysis must also be determined. In most cases, an automated software package, such as one that uses the CLS algorithm, is used to determine the concentrations of the target gases. These procedures can be checked manually by
comparing the sample spectra to spectra of reference gases with a known concentration. Interactive subtraction procedures that yield a scaling factor for the reference spectrum can be used to check the concentration measured by the CLS software. In addition, the reference spectra can be scaled to the desired concentration and then added to the sample spectrum. When the composite spectrum is then analyzed, the measured concentration should reflect the amount of reference gas added. Care must be taken in choosing the spectral regions used to analyze for each target compound. For example, the optimum region for analysis does not always encompass the entire absorption envelope. Possible interfering species must also be accounted for in the analysis method.

The operator should also be aware that any time a concentration spike appears that cannot be immediately attributed to a known source, the actual spectral data should be examined to verify the presence of the compound in question and its concentration. This can be done by first subtracting the appropriate absorption spectra of any interfering species from the sample spectrum. The signature and absorbance values of the resultant spectrum should then be compared to the reference spectrum for proper features and intensities. Also, any time that the concentrations of the target compounds seem to correlate with changes in water vapor concentration, the spectra of the target compounds should be examined to verify that the changes in concentration are real. If the concentrations of the target compounds exhibit either positive or negative inflections with respect to changes in water vapor concentration, the analysis method should be altered to alleviate the problem.

Ultimately, if the instrument is operating properly and a suitable analysis method is developed, the accuracy of the FT-IR technique will be determined by the accuracy of the reference spectra. To date, no way of validating or certifying these reference spectra exists.

8.4.4 Completeness and Representativeness of Data

These requirements will vary with specific monitoring applications. Care must be taken to ensure that data points are acquired frequently enough to account for the variability of the target gas concentration. Failure to do this will make it difficult to discern between real changes in the target gas concentration and possible variability in the FT-IR measurements.

8.4.5 Comparability of the Data

If possible, the FT-IR data should be initially compared to an established method. This can be difficult because the FT-IR produces a path-averaged concentration, whereas most established methods use some type of point monitor. As discussed in Section 8.2.2, some of the FT-IR data have been compared to the canister method. In a current study, we are comparing ozone measurements recorded with the FT-IR monitor to those taken with personal sampling devices and to hourly averages from
the state ozone monitoring program. Although not exact, these comparisons can give the operator an idea if the FT-IR measurements are within generally accepted values. If not, corrective action should be taken.

8.4.6 Ancillary Measurements

The type of ancillary measurements required will vary, depending on the type of study being conducted. For any long-path, open-path measurements, the ambient temperature, water vapor concentration, ambient pressure, and wind velocity should be recorded. The operator should also be aware of the effect of changes in altitude on ambient pressure. If the instrument is housed in an enclosed environment, the temperature of that environment should also be recorded. We have also found it useful to record the temperature inside the spectrometer itself, especially in cold weather situations.

8.4.7 Documentation

As with any analytical methodology, a log of instrument usage, downtime, and repairs, as well as notes regarding unusual observations, should be maintained. These notes can prove invaluable for analyzing data that appear to be abnormal. Records should be kept that are appropriate for the type of study being conducted. For example, requirements for a research and development project may be different from those required for legally defensible data.

8.5 References


Chapter 9
Glossary of Terms for
FT-IR Open-Path Remote Sensing

9.1 Introduction and Overview

This chapter contains a glossary of terms for remote sensing, with an emphasis on those terms relevant to FT-IR long-path, open-path monitoring. When possible, definitions of terms have been drawn from authoritative texts or manuscripts in the fields of remote sensing, air pollution monitoring, spectroscopy, optics, and analytical chemistry. In some cases, general definitions have been augmented or streamlined to be more specific to long-path, open-path monitoring applications. These definitions were intended to remain scientifically rigorous and still be generally applicable to a variety of FT-IR open-path remote-sensing issues.

9.2 Terms

Absorbance: The negative logarithm of the transmission, \( A = \ln(I/I_0) \), where \( I \) is the transmitted intensity of the light and \( I_0 \) is the incident intensity. Generally, the logarithm to the base 10 is used, although the quantity \( I \) really diminishes exponentially with \( A \) (Pfeiffer and Liebhafsky 1951). If the term "fractional transmission" is used for the ratio \( I/I_0 \), then the implication is that the instrument’s slit function (see "Instrument function") is accounted for (Penner 1959).

Active system: A system that radiates energy to the surrounding environment (for example, a radio transmitter).

Apodization: A mathematical transformation carried out on data received from an interferometer to alter the instrument’s response function. There are various types of transformation; the most common are boxcar, triangular, Happ-Genzel, and Beer-Norton functions.

Average concentration: For FT-IR or differential absorption spectroscopy systems, this quantity is the result of dividing the integrated concentration (the quantity that is measured) by the path length used for the measurement. It has units of parts per million, parts per billion, micrograms per cubic meter, etc. (McClenny and Russwurm 1978).

Background spectrum: 1. With all other conditions being equal, that spectrum taken in the absence of the particular absorbing species of interest. 2. Strictly, that radiant intensity incident on the front plane of the absorbing medium. 3. A spectrum obtained from the ambient black body radiation entering the system. This background must be considered in FT-IR systems, in which the IR beam is not modulated before it is transmitted along the path. For FT-IR systems that do not use a separate source of infrared energy, the background is the source of infrared energy.

Band pass filter: A filtering device that allows the transmission of only a specific band of
energies. It can be wide band or narrow band.

**Bandwidth:** The width of a spectral feature as recorded by a spectroscopic instrument. This width is listed as the full width at the half maximum of the feature or as the half width at the half maximum of the spectral feature. This is also referred to as the line width (Lengyel 1971).

**Beer’s law:** Beer’s law states that the intensity of a monochromatic plane wave incident on an absorbing medium of constant thickness diminishes exponentially with the number of absorbers in the beam. Strictly speaking, Beer’s law holds only if the following conditions are met: (1) perfectly monochromatic radiation, (2) no scattering, (3) a beam that is strictly collimated, (4) negligible pressure-broadening effects (Pfeiffer and Liebhafsky 1951; Lothian 1963). For an excellent discussion of the derivation of Beer’s law, see Penner (1959).

**Bistatic system:** A system in which the receiver is some distance from the transmitter. This term is actually taken from the field of radar technology. For remote sensing, this implies that the light source and the detector are separated and are at the ends of the monitoring path.

**Broad band system:** Any system that admits a broad range of energies into its signal-processing section. Alternatively, a system that has a flat response to a large range of energies.

**Closed path:** The optical path over which the beam travels in a sensor that is entirely enclosed. This is the case when White cells or Harriot cells are used with the system.

**Cooler:** A device into which the detector is placed for maintaining it at a low temperature in an IR system. At a low temperature, the detector provides the high sensitivity that is required for the IR system. The two primary types of coolers are a liquid nitrogen Dewar and a closed-cycle Stirling cycle refrigerator.

**COSPEC:** An acronym for "correlation spectrometer." This is a misnomer for the type of instrument implied. It is really an instrument that uses a diffraction grating either in the active, bistatic or the passive mode. This instrument is more correctly included in the class of instruments using differential absorption spectroscopy techniques. It is used primarily to measure sulfur dioxide and nitrogen oxide.

**DIAL:** An acronym for "differential absorption lidar." This system uses two pulsed frequencies from the same laser or from different lasers to measure the concentration of gas over a path. The two laser lines are at different positions within the absorption feature. The difference of the amount of light backscattered at these two wavelengths is the quantity used for the measurement.

**DOAS:** An acronym for differential optical absorption spectroscopy. A technique whereby, in principle, any known difference in absorbance is used to determine the concentration of a gas. Generally, the
absorption difference is taken between the spectral line center and the wing.

**Electromagnetic spectrum:** The total of all possible frequencies of electromagnetic radiation. Different sources may emit over different frequency regions. All electromagnetic waves travel at the same speed in free space (Halliday and Resnick 1974).

**Fingerprint region:** The region of the absorption spectrum of a molecule that essentially allows its unequivocal identification. This region covers the wave number range from 650 to 1300 cm\(^{-1}\) (Willard et al. 1974).

**Flux:** The number or mass of particles or molecules that pass through a given unit area of surface per unit of time (Calvert 1990).

**Fourier transform:** A mathematical transform that allows an aperiodic function to be expressed as an integral sum over a continuous range of frequencies (Champeney 1973). The Fourier transform of the interferogram produced by the Michelson interferometer in an FT-IR is the intensity as a function of frequency.

**FT-IR:** An abbreviation for "Fourier transform infrared." A spectroscopic instrument using the infrared portion of the electromagnetic spectrum. The working component of this system is a Michelson interferometer. To obtain the absorption spectrum as a function of frequency, a Fourier transform of the output of the interferometer must be performed. For a brief overview of the FT-IR, see the publication by Nicolet (Nicolet Analytical Instruments 1986). For an in-depth description of the FT-IR, see Griffiths and deHaseth (1986).

**GASPEC:** An acronym for "gas filter correlation spectrometer." The earliest of these devices was described by Luft in 1943, and they have been used in various configurations ever since. The primary feature of these devices is a pair of gas cells. One cell contains a carefully selected quantity of the target gas and the other a gas that is spectroscopically inactive. The difference in spectral transmittance of the two cells is an indicator of the concentration of the target gas in the atmosphere (Ward and Zwick 1975).

**Infrared spectrum:** That portion of the electromagnetic spectrum that spans the region from about 10 cm\(^{-1}\) to about 12,500 cm\(^{-1}\). It is divided (Willard et al. 1974) into (1) the near-infrared region (from 12,500 to 4000 cm\(^{-1}\)), (2) the mid-infrared region (from 4000 to 650 cm\(^{-1}\)), and (3) the far-infrared region (from 650 to 10 cm\(^{-1}\)).

**Instrument function:** The function superimposed on the absorption line shape by the instrument. This is sometimes referred to as the slit function, a term taken from instruments that use slits to obtain resolution.

**Intensity:** The radiant power per unit solid angle. When the term "spectral intensity" is used, the units are watts per steradian per nanometer. In most spectroscopic literature,
the term "intensity" is used to describe the power in a collimated beam of light in terms of power per unit area per unit wavelength. However, in the general literature, this definition is more often used for the term "irradiance," or "normal irradiance" (Calvert 1990; Stone 1963).

Interference: The physical effects of superimposing two or more light waves. The principle of superposition states that the total amplitude of the electromagnetic disturbance at a point is the vector sum of the individual electromagnetic components incident there. For a two-component system of collinear beams of the same amplitude, the mathematical description of the result of addition is given by \( I(p) = 2I_0(1 + \cos(\Delta)) \), where \( I_0 \) is the intensity of either beam, and \( \Delta \) is the phase difference of the two components. The cosine term is called the "interference term" (Halliday and Resnick 1974; Stone 1963). See also "Spectral Interference."

Interferogram: The effects of interference that are detected and recorded by an interferometer; the output of an FT-IR and the primary data that is collected and stored (Stone 1963; Griffiths and deHaseth 1986).

Interferometer: Any of several kinds of instruments used to produce interference effects. The Michelson interferometer used in FT-IR instruments is the most famous of a class of interferometers that produce interference by the division of an amplitude (Tolansky 1962).

Irradiance: Radiant power per unit projected area of a specified surface. This has units of watts per square centimeter. The term "spectral irradiance" is used to describe the irradiance as a function of wavelength. It has units of watts per square centimeter per nanometer (Calvert 1990).

Laser: An acronym for the term "light amplification by stimulated emission of radiation". A source of light that is highly coherent, both spatially and temporally (Lengyel 1971).

LIDAR: An acronym for the term "light detection and ranging" (Calvert 1990). A technique for (1) detecting the presence of gases and aerosols by measuring the backscattered portion of a laser beam and (2) determining the range of these gases and aerosols by electronically gating the detected signal and performing a calculation based on the speed of light (about 30 cm/ns).

Light: Strictly, light is defined as that portion of the electromagnetic spectrum that causes the sensation of vision. It extends from about 25,000 cm\(^{-1}\) to about 14,300 cm\(^{-1}\) (Halliday and Resnick 1974).

Light scattering: The redirection of light waves due to interaction with molecules or aerosols. If the size of the body causing the scattering is small compared to the wavelength of the incident radiation, the scattering is termed "Raleigh scattering." If the size is large compared to the wavelength, the scattering is termed "Mie scattering." Scattering by molecules is generally Raleigh
scattering, and scattering by aerosols is Mie scattering. As light travels through a medium, two physical processes diminish the intensity in the forward direction — scattering and absorption. The sum of these two effects is called extinction (van de Hulst 1981).

**Long-path monitoring:** This is a monitoring technique that uses an extended, open path. LIDAR systems can make measurements over a path length of a few kilometers. DOAS systems can make measurements of ozone over a path of up to 2 km. The FT-IR systems customarily use paths with length less than 1 km.

**Minimum detection limit:** The minimum concentration of a compound that can be detected by an instrument with a given statistical probability. Usually the detection limit is given as 3 times the standard deviation of the noise in the system. In this case, the minimum concentration can be detected with a probability of 99.7% (Calvert 1990; Long and Winefordner 1983).

**Monitoring path:** The actual path in space over which the pollutant concentration is measured and averaged.

**Monitoring path length:** The length of the monitoring path in the atmosphere over which the average pollutant concentration measurement is determined (U.S. Environmental Protection Agency 1994).

**Monostatic system:** A system with the source and the receiver at the same end of the path. For FT-IR systems and for DOAS systems, the beam is generally returned by a retroreflector. For LIDAR systems, the backscattered portion of the laser beam is measured directly.

**Open-ended system:** A system in which the remote sensor uses light reflected from targets of opportunity (walls, trees, etc.) or skylight as a source.

**Open-path analyzer:** An automated analytical instrument that is used for a method of measuring the average atmospheric pollutant concentration in situ along one or more monitoring paths that are 5 m or more in length (U.S. Environmental Protection Agency 1994).

**Open-path monitoring:** Remote sensing over a path that is completely open to the atmosphere. Thus, the concentration of a particular gas in the beam path can be changed by winds and diffusion. The open path is the most frequently used in remote sensing.

**Optical remote sensing:** A generic term used to describe any of a number of optical measurement techniques that measure some quantity or constituent of the atmosphere. These techniques include DIAL, DOAS, FT-IR, GASPEC, LIDAR, etc. One thing that is common to all the techniques is that no sample must be collected.

**Parts per million meters:** The unit term for the quantity that is measured by many remote sensors. It is the unit associated with the quantity path-integrated concentration. It
is a possible unit of choice for reporting data from remote sensors because it is independent of the path length.

**Passive system:** Any system that does not radiate energy to its surroundings.

**Path-averaged concentration:** The result of dividing the path-integrated concentration by the path length. Path-averaged concentration gives the average value of the concentration along the path (McClenny and Russwurm 1978).

**Path-Integrated concentration:** The quantity that is measured by a remote sensor over a long path. It has units of concentration times length.

**Plume:** The gaseous and aerosol effluents emitted from a chimney or other source and the volume of space they occupy. The shape of a plume and the concentration of pollutants within it are very sensitive to meteorological conditions (Calvert 1990).

**Point analyzer:** An automated analytical method that measures pollutant concentration in an ambient air sample extracted from the atmosphere at a specific inlet probe point (proposed changes [U.S. Environmental Protection Agency 1994]).

**Probe:** The actual inlet where an air sample is extracted from the atmosphere for delivery to a sampler or point analyzer (proposed changes [U.S. Environmental Protection Agency 1994]).

**Radiometry:** The measurement of various quantities such as intensity associated with radiant energy (Calvert 1990). This is in contrast to the term photometry, which assumes the spectral sensitivity of the human eye as the detector (Walsh 1965).

**Real-time system:** Any monitoring system that acquires and records data at a rate that is comparable to the rate at which the concentration is changing.

**Reference spectra:** Spectra of the absorbance versus wave number for a pure sample of a set of gases. The spectra are obtained under controlled conditions of pressure and temperature and with known concentrations. For most instruments, the pure sample is pressure-broadened with nitrogen so that the spectra are representative of atmospherically broadened lines. These spectra are used for obtaining the unknown concentrations of gases in ambient air samples.

**Resolution:** The minimum separation that two spectral features can have and still, in some manner, be distinguished from one another. A commonly used requirement for two spectral features to be considered just resolved is the Raleigh criterion. This states that two features are just resolved when the maximum intensity of one falls at the first minimum of the other (Jenkins and White 1950; Tolansky 1962). This definition of resolution and the Raleigh criterion are also valid for the FT-IR, although there is another definition in common use for this technique. This definition states that the minimum
separation in wave numbers of two spectral features that can be resolved is the reciprocal of the maximum optical path difference (in centimeters) of the two interferometer mirrors employed (Griffiths and deHaseth 1986; Nicolet 1986).

Retrorreflector: The CIE (Commission Internationale de l'Eclairage) defines retroreflection as "radiation returned in directions close to the direction from which it came, this property being maintained over wide variations of the direction of the incident radiation." Retrorreflector devices come in a variety of forms and have many uses. The one commonly described by workers in remote sensing uses total internal reflection from three mutually perpendicular surfaces. This kind of retrorreflector is usually called a corner cube or prismatic retrorreflector (Rennison 1980).

Slit function: See "Instrument function."

Source: The device that supplies the electromagnetic energy for the various instruments used to measure atmospheric gases. These generally are a Nernst glower or globar for the infrared region or a xenon arc lamp for the ultraviolet region. There are several lasers that also are used as sources for DIAL and LiDAR instruments.

Spectral intensity: See "Intensity."

Spectral interference: When the absorbance features from two or more gases cover the same wave number regions, the gases are said to exhibit spectral interference. Water vapor produces the strongest spectral interference for infrared spectroscopic instruments that take atmospheric data.

Synthetic background: A spectrum that is made from a field spectrum by choosing points along the baseline and connecting them with a high-order polynomial or short, straight lines. The synthetic background is then used to find the absorbance spectrum.

Truncation: The act of stopping a process before it is complete. In FT-IR spectrometers, the theoretically infinite scale of the interferogram is truncated by the finite movement of the interferometer mirror.

Unistatic system: A system that has the source and the receiver at the same place; now more commonly referred to as a monostatic system.

Wave number: The number of waves per centimeter. This term has units of reciprocal centimeters (cm⁻¹).

9.3 References


Chapter 10
Bibliography

10.1 Introduction and Overview

This chapter contains a bibliography of journal articles, books, and conference proceedings that address FT-IR long-path monitoring, as well as general references that present discussions of the basic principles of FT-IR spectrometry. This is not a reference section for literature cited in the text of this guidance document, but is a general bibliography. References in this document are listed at the end of the chapter in which they are cited.

This bibliography is by no means exhaustive, but is intended to give a broad overview of the available literature in the FT-IR long-path discipline. During the initial phase of this overview, a computer-assisted on-line literature search was conducted for citations listed in Chemical Abstracts through September 29, 1992. A subsequent on-line search was conducted for citations listed in Chemical Abstracts from 1992 through July 22, 1994. These literature searches focused on, but were not limited to, current publications regarding FT-IR long-path, open-path monitoring, in keeping with the emphasis of this document. Other citations were gleaned from the reference sections of articles on file. Several early citations are also given to provide an important, historical perspective into long-path IR monitoring in environmental analysis. Only articles that are on file in our laboratory are listed in the bibliography. This bibliography will continue to be updated as revisions to this document are made.

Several of the more recent citations are from conference proceedings, which may limit their general availability. They are included to give the reader access to current research in the field that has in many cases not yet appeared in the peer-reviewed literature. Also, inclusion of the conference proceedings provides the reader with pertinent information regarding conferences and meetings that typically address FT-IR long path or remote-sensing issues.

10.2 Publications


Air & Waste Management Association, Pittsburgh, PA, pp. 529-540.


Scotto, R.L., B.C. Sanders, R.H. Kagann, and O.A. Simpson. 1992. Validation of a Gaussian Plume Dispersion Model Based on Data from the Kansas Open-Path FTIR Intercomparison Study. Proceedings of the
85th Annual Meeting and Exhibition, Air & Waste Management Association, Pittsburgh, PA, pp. 92-73.04.


University of South Florida. 1993. USF HITRAN-PC. University of South Florida, Tampa, FL.

to Environmental and Industrial Safety Problems, Air & Waste Management Association, Pittsburgh, PA, pp. 44-54.


