1. Introduction

With the recent developments in Fourier transform infrared (FTIR) spectrometers it is becoming more feasible to place these instruments in field environments [1]. As a result, there has been enormous increase in the use of FTIR techniques for a variety of qualitative and quantitative chemical measurements. These methods offer the possibility of fully automated real-time quantitation of many analytes; therefore FTIR has great potential as an analytical tool. Recently, the U.S. Environmental Protection Agency (U.S.EPA) has developed protocol methods for emissions monitoring using both extractive and open-path FTIR measurements. Depending upon the analyte, the experimental conditions and the analyte matrix, approximately 100 of the hazardous air pollutants (HAPs) listed in the 1990 U.S.EPA Clean Air Act amendment (CAAA) can be measured. The National Institute of Standards and Technology (NIST) has initiated a program to provide quality-assured infrared absorption coefficient data based on NIST prepared primary gas standards. Currently, absorption coefficient data has been acquired for approximately 20 of the HAPs. For each compound, the absorption coefficient spectrum was calculated using nine transmittance spectra at 0.12 cm\(^{-1}\) resolution and the Beer’s law relationship. The uncertainties in the absorption coefficient data were estimated from the linear regressions of the transmittance data and considerations of other error sources such as the nonlinear detector response. For absorption coefficient values greater than \(1 \times 10^4 \text{ mmol/mol}^{-1} \text{ m}^{-1}\), the average relative expanded uncertainty is 2.2%. This quantitative infrared database is currently an ongoing project at NIST. Additional spectra will be added to the database as they are acquired. Our current plans include continued data acquisition of the compounds listed in the CAAA, as well as the compounds that contribute to global warming and ozone depletion.

Key words: air pollutants; database; gas standards; infrared spectrometer.

Accepted: December 3, 1998

Available online: http://www.nist.gov/jres
sources including the quantitative libraries by Infrared Analysis [8],¹ MIDAC Corporation [9], and Sprouse Scientific [10]. Comparisons of reference spectra from the available quantitative collections shows that the agreement of reported intensities is frequently ± 10 % or worse. Impurity bands present in reference spectra can also interfere with the interpretation of field results.

The National Institute of Standards and Technology (NIST) has initiated a program to develop a quality-assured quantitative database of infrared spectra based on NIST prepared primary gas standards. Data acquisition is currently focused on the hazardous air pollutant species listed in the CAAA [4]. Since the database is designed to facilitate ground-based open-path FTIR measurements, the data were acquired with samples at room temperature and pressure broadened with nitrogen to one atmosphere. Currently, absorption coefficient data are available for approximately 20 HAPs on a U.S. EPA priority list. The data are stored in the standard JCAMP-DX format [11] to enable universal access to the data. Unapodized interferograms were acquired at 0.12 cm⁻¹ resolution and have been processed to generate data at a number of different resolutions and apodizations, providing the users with data that closely match their experimental parameters. At each wavenumber in the spectrum the absorption coefficient $a$ is given as defined by the Beer-Lambert equation:

$$I_n(\nu) = I_0(\nu)10^{-a(n)c_l}$$

where $I_n(\nu)$ and $I_0(\nu)$ are the transmitted and incident light intensities, $c$ denotes the concentration of absorbing species, and $l$ is the path length. A digital signature accompanies each data file, allowing users to ensure the integrity and source of the data file and traceability to NIST.

This quantitative infrared database is an ongoing project at NIST. Additional spectra will be added to the database as they are acquired. Our current plans include continued data acquisition of the compounds listed in the CAAA (Appendix A) [4], as well as the compounds that contribute to global warming and ozone depletion.

## 2. Materials and Methods

### 2.1 Materials

The volatile organic compounds (VOCs) used to prepare these gas standards were obtained from commercial suppliers with the highest purity available, in most cases the stated purity was 99.9 %. Purity analyses were performed on the VOCs using gas chromatography with mass selective detection, differential scanning calorimetry, and Karl Fischer coulometric methods. Generally, the compounds were found to be 99.9 % pure by gas chromatography and differential scanning calorimetry. The Karl Fischer titrations measured significant amounts of water in a number of the samples. Table 1 lists the samples in three categories; compounds that had a mass fraction of water greater than 0.1 %, compounds that had a mass fraction of water less than 0.1 %, and compounds that were not measured by the Karl Fischer method. These results were included in the gravimetric values.

Ultra-high-purity nitrogen (99.9995 %) was used as the balance gas. The primary gas standards were prepared in aluminum cylinders having an internal volume of 6 L and equipped with brass valves. The cylinders were pre-cleaned by a commercial supplier in a manner that minimizes contamination by trace hydrocarbons and halocarbons and then treated to deactivate the internal walls.

### 2.2 Gravimetric Standards Preparation

The procedure to prepare µmol/mol (commonly referred to as part-per-million) level gravimetric gas standards of VOCs in nitrogen has been described in detail previously [12]. An evacuated, preweighed cylinder is fitted with the appropriate CGA-350 fitting equipped with a septum. A pure organic liquid is introduced into the cylinder with a septum. A pure organic liquid is introduced into a gas tight syringe. The syringe containing the analyte is weighed on a microbalance with a capacity of 100 g, an uncertainty on the order of 5 µg. The fitting on the cylinder is heated with a heat gun to approximately 80 °C. Then the syringe needle is inserted into the septum while the cylinder valve is opened. If all the liquid is not immediately pulled into the evacuated cylinder,
the syringe is heated gently. The syringe is then removed and weighed immediately and the weight of the organic material in the cylinder is determined by the difference in the syringe weights.

Next, ultra-high-purity nitrogen is added to the cylinder to a precalculated pressure, and the cylinder is weighed. The amount-of-substance fraction (commonly called mole fraction) of the VOCs is calculated from the weight of the pure VOCs and the weight of the nitrogen placed in the cylinder. The cylinders are weighed on a top-loading balance with a maximum capacity of 32 kg, and an uncertainty on the order of 0.1 g. All balances were calibrated with NIST-traceable weights. The final mole fractions of the gravimetric standards prepared for this work range from 1 \( \mu \text{mol/mol} \) to 1000 \( \mu \text{mol/mol} \), with the upper limit dependent on the vapor pressure of the individual compound. The standard concentrations were chosen based on the infrared band strengths. For standards up to 50 \( \mu \text{mol/mol} \), the expanded uncertainty (coverage factor of \( k = 2 \) and thus a two standard deviation estimate, representing a 95% confidence interval) of the gravimetric values is 0.5% based on the uncertainties from the weighing procedures. For standards ranging from 50 \( \mu \text{mol/mol} \) to 1000 \( \mu \text{mol/mol} \), the expanded uncertainty in the gravimetric values is 0.2%. Finally, the gas standards are analyzed using a gas chromatograph (GC) equipped with a flame-ionization detector. The data are fitted to a quadratic equation to verify the gravimetric procedure. The GC results confirm the gravimetric values to < 1.0%.

2.3 Data Acquisition

FTIR spectra at 0.12 cm\(^{-1}\) resolution were acquired using a liquid-nitrogen cooled, mercury-cadmium-telluride (HgCdTe) detector with the optical bench under vacuum. The primary gas standards were flowed continuously at 1 L/min at atmospheric pressure through a multipass absorption cell with a total volume of 7.5 L and a maximum path length of approximately 20 m. The mirror spacing was measured with the cell disassembled. The total path length for one pass, including the additional length at the cell entrance and exit, measured 1.356 m with a standard uncertainty of approximately 0.001 m. All other path lengths were derived from the mirror spacing [13]. The accuracy of this measurement was confirmed by comparing band intensities from laser studies [14] with the integrated band intensity of the \( n + \nu_3 \) band of SO\(_2\) obtained using this FTIR spectrometer [15]. The ambient temperature and pressure were monitored periodically throughout the measurements and referenced to a NIST-calibrated thermometer and capacitance manometer.

All background spectra were taken with ultra-high-purity nitrogen flowing through the cell. To test the stability of the sample concentration, several short scans were recorded first. Once it was verified that the ratio of consecutive scans showed no drift in the absorbance, a longer scan was recorded to obtain a signal-to-noise of 1000 or better. For the benzene-in-nitrogen mixtures, the transmittance was reproducible to ± 0.5% within 15 minutes.

Generally, the data were acquired from three different gravimetric concentrations at three different path lengths to generate a total of nine spectra. This procedure was chosen to provide a large dynamic range of data, so that both the strong and weak bands could be observed. Figure 1 shows the dynamic range of the transmission spectra acquired for the benzene samples. The final absorption coefficients were calculated from the nine transmission spectra and have been corrected to 296 K and 1.013 \( \times 10^5 \) Pa (760.0 Torr) using the ideal gas law.

2.4 Wavenumber Calibration Using Water Vapor

The most convenient method for calibrating the wavenumber scale of the instrument is to use selected water vapor lines in both the (1200–1900) cm\(^{-1}\) and (3500–4000) cm\(^{-1}\) regions of the spectrum. These lines are always found in the spectrum and have been measured and tabulated by Toth [16], using the FT instrument at Kitt Peak, to a standard uncertainty of better than 0.0005 cm\(^{-1}\). Water vapor spectra were obtained with 1.3 kPa (10 Torr) of ambient air in the multipass cell to minimize pressure-broadening effects. The water vapor peak positions were identified using the boxcar apodization function and second derivative peak search routine. The wavenumber shifts of selected water lines measured on our spectrometer compared to those of the Kitt Peak measurements are shown in Fig. 2. As expected, a small but significant shift is found which is linear with wavenumber. By fitting the ratio of the measured wavenumbers to the calibration wavenumbers, a correction factor was derived. When corrected wavenumbers are subtracted from those values given by Toth, the root-mean-square deviation obtained for 158 lines is 0.0042 cm\(^{-1}\) which is a good indication of the standard uncertainty of the frequency in our measurements. To maintain consistent wavenumber accuracy throughout this work, the wavenumber calibration is checked periodically and any time after the interferometer has been adjusted.

2.5 Data Processing

In all cases, unapodized interferograms were truncated to yield spectra at nominal resolutions of
Fig. 1. Transmission spectra of benzene-in-nitrogen samples at three different concentrations and path length combinations.

Fig. 2. Wavenumber calibration using selected water vapor lines.
The resolution $\Delta \nu$ is given by the relationship $\Delta \nu = 1/L$, where $L$ is the maximum retardation of the interferometer [17]. Then the interferograms were transformed using the Mertz phase correction, a zerofilling factor of two, a boxcar apodization function, and a detector nonlinear correction routine using the software package supplied with the instrument [18]. Pointby-point absorption coefficients were calculated from nine transmission spectra using the known concentrations and path lengths. Figure 3 shows a Beer’s law plot for benzene at three representative wavelengths with the absorbance, $A = -\log_{10}(T)$ where $T$ is the transmittance. The uncertainty in the absorbance was derived from the uncertainty in the transmittance based on the relationship $\Delta A \sim (1/T) \Delta T$. The data were modeled with a linear regression and a weighting factor given by $(1/\Delta A)^2 = (T/\Delta T)^2$. For transmittance values less than 0.02, the transmittance was set to $1 \times 10^{-3}$. Generally, correlation coefficients $r^2 = 0.9997$ were obtained from the linear regressions, confirming the Beer’s law behavior of this system.

Since many potential users collect data at different resolutions and apodization functions, an effort was made to provide the absorption coefficient data that would closely match the users’ acquisition parameters. Deresolve [19], a program designed to degrade the resolution of high-resolution absorbance reference spectra, was used to generate the lower resolution data. Deresolve generates a transmittance spectrum from an absorbance spectrum then calculates the inverse Fourier-transform. The resulting interferogram is truncated and convoluted with a specified apodization function. Finally, the interferogram is transformed and converted back into an absorbance spectrum. It is anticipated that this program will accompany this database in future releases.

An absorbance spectrum for a given concentration and path length can be calculated from the tabulated absorption coefficient data by multiplying the absorption coefficient data by the desired mole fraction in units of $\mu$mol/mol and by the desired path length in meters. It is important to emphasize that the absorbance spectrum, calculated as described above, will only be accurate at low absorbances where the absorbance is linear. The absorbance levels where non-linearities become an important factor depend on the resolution and apodization function of the spectrum as well as the natural width of the absorption feature.

Fig. 3. Plot of the absorbance versus concentration multiplied by path length for selected lines along with the list squares fit to the data. The largest uncertainty in the absorbance is $(1.0 \times 10^{-3})$ absorbance units.
3. Artifacts

3.1 Residual H₂O, CO, and CO₂

Careful examination of the individual spectra indicated that residual water, carbon monoxide, and carbon dioxide features are present in the absorbance spectra. This indicates that there are different levels of H₂O, CO, and CO₂ in the background spectra compared to the sample spectra. Peak heights of H₂O, CO, and CO₂ lines were measured and compared to line intensities tabulated in the HITRAN database [7]. All of the CO and CO₂ peak heights correspond to less than ±2 μmol/mol of CO and CO₂, where the negative value represents higher mole fractions of the contaminant in the background spectrum. Trace amounts of CO and CO₂ in the ultra-high-purity nitrogen are likely to be the largest sources of the CO and CO₂. The amount of water in the absorbance spectra generally varied from −2 μmol/mol to 25 μmol/mol. By comparing the water levels for all of the spectra, it was clear that the water levels depended on the compound as well as the concentration of the primary gas standards. This suggested that moisture in the pure VOCs is the largest source of water. This hypothesis was confirmed by Karl Fischer coulometric measurements that quantitated the moisture in the original VOCs. In fact, the mass fractions of water in the pure VOCs were in excellent agreement with the mass fraction of water in the gas standards calculated from the FTIR spectra.

3.2 Electronic Noise

During the course of this work, it was noted that consistent noise spikes occurred in the spectra at 1974.8 cm⁻¹, 2962.2 cm⁻¹, and 3949.5 cm⁻¹. These artifacts also appeared in spectra taken when no light reached the detector. An effort was made to eliminate these artifacts, but was not successful. Since these features are clearly artifacts, the noise spikes were eliminated from the spectra by replacing the data in a 0.4 cm⁻¹ region centered about the spike with a line. The parameters for the line were obtained by using a linear function to model the data on either edge of these features.

3.3 Baseline Drift

An examination of all the transmission spectra indicates that there was a higher probability for the 3150 cm⁻¹ to 3400 cm⁻¹ region of the baseline to drift more than other regions. This artifact has been attributed to changes in the detector conditions during the course of the data acquisition. An effort was made to minimize these errors by taking background spectra at the beginning and end of a data run. The background spectrum that most closely matched the sample spectrum was used to calculate the transmission spectrum.

4. Uncertainties

The actual statistical or Type A [20] uncertainties in these measurements are represented by the uncertainties obtained from the linear regressions of the data. ORTHO [21], a Fortran subroutine that performs a least squares fit for a set of linear equations or power series was used to obtain the point-by-point absorption coefficients, α, along with the associated uncertainty in the absorption coefficients, uα. Figure 4a shows the absorption coefficient data calculated for ethylene and Fig. 4b shows uα as a function of wavenumber. It is clear from Fig. 4b that there are significantly larger uncertainties in the absorption coefficient data in the regions of the spectrum where H₂O, CO, and CO₂ absorb due to significant variations in the levels of these species in the background and sample spectra. Because it is difficult to completely remove these features from the spectra, the absorption coefficients are not certified in regions of the spectra where H₂O [(1325–1900) cm⁻¹ and (3550–3950) cm⁻¹], CO [(2050–2225) cm⁻¹], and CO₂ [(2295–2385) cm⁻¹] absorb.

Figure 5 shows uα as a function of the associated a, demonstrating that the uncertainty in the absorption coefficient can be approximated by a linear function of a, with uα = ma + b. Regions where H₂O, CO, and CO₂ absorb were not included in this analysis. Table 2 lists the slope m and intercept b parameters which can be used to approximate uα for each compound contained in the database along with the mean and the standard deviation of the mean of the m and b parameters. These results indicate that the b parameter for each compound can be replaced by the mean value for b.

The results in Table 2 also show that the relative Type A uncertainty, which can be approximated by m for a > 1 × 10⁻⁴ (μmol/mol)⁻¹ m⁻², is significantly larger for three compounds: benzene, bromomethane, and ethyl tert-butyl ether. The uncertainties reported for the gravimetric standards in Sec 2.2 are based only on the uncertainties in the weighing procedures. Additional factors can affect the final gravimetric concentration [22] and may be responsible for the uncertainties observed for benzene, bromomethane, and ethyl tert-butyl ether. For example, a compound may react with the cylinder walls. Ten additional benzene samples were intercompared with FTIR spectrometry and the integrated band absorbances for two of the standards used for the database were significantly different compared to the other benzene samples. An effort is currently underway to improve the benzene results.
The nonlinear response of HgCdTe detectors has been documented [17]. This will add an additional non-statistical Type B [20] relative uncertainty to the measurements, which has been included in the uncertainty analysis. An estimate of the standard uncertainty is 1.0 % of the absorption coefficient and was obtained by comparing integrated band intensities for measurements made with a deuterated triglycine sulfate (DTGS) detector to measurements with a HgCdTe detector. The HgCdTe measurements were processed using the nonlinear correction routine supplied with the instrument [18].

Additional experimental variables contribute to the overall uncertainty of the absorption coefficients. Type B estimates of the relative standard uncertainties for the cell path length, pressure, temperature and FTIR stability are listed in Table 4. Estimates of the Type B relative uncertainty in the sample concentration where approximately a factor of ten lower than the sample to sample variability. Since gravimetric standards were measured for each compound, uncertainties in the absorption coefficients due to the sample concentrations are folded into the evaluation of the Type A uncertainties. An additional Type B uncertainty was included for samples, which showed the presence of water in the FTIR spectra, but were not tested for water content by the Karl Fischer method. The magnitude of this uncertainty component was estimated by the amount of water measured in the FTIR spectra.

The Type B relative uncertainties were combined by the equation:

$$u_{\text{Brel}} = \sqrt{u_1^2 + u_{\text{press}}^2 + u_{\text{temp}}^2 + u_{\text{FTIR}}^2 + u_{\text{NL}}^2 + u_{\text{water}}^2},$$

(2)

where the relative standard uncertainties in the cell path length, pressure, temperature, FTIR stability, detector nonlinearities, and sample water content are denoted by $u_1$, $u_{\text{press}}$, $u_{\text{temp}}$, $u_{\text{FTIR}}$, $u_{\text{NL}}$, and $u_{\text{water}}$ respectively. The uncertainty attributed to the detector nonlinearities clearly dominate the Type B relative uncertainties. The combined Type B uncertainties are listed in Table 2.
Fig. 5. The uncertainty in the absorption coefficient $u_a$ plotted as a function of the associated absorption coefficient $a$ along with a linear approximation of $u_a \approx ma + b$.

The expanded uncertainty is defined as $U = ku_c$ with the standard uncertainty $u_c$ determined from the experimental Type A and Type B standard uncertainties and the coverage factor $k = 2$. The Type A and Type B uncertainties were combined as:

$$U = 2(u_a^2 + (u_{\text{type B}} a)^2)^{1/2} = 2((ma + b)^2 + (u_{\text{type B}} a)^2)^{1/2}.$$  

(3)

$U$ can be simplified to:

$$U \approx 2(Ba^2 + Ca + D)^{1/2}.$$  

(4)

where the coefficients $B$, $C$, and $D$ are listed in Table 3. The value of the absorption coefficient at a given wavelength is asserted to lie in the interval defined by $(a \pm U)$ with a level of confidence of approximately 95%. In the relationship between $U$ and $a$, $D$ represents the uncertainty in the baseline. As the absorption coefficient gets larger, the $Ba^2$ term dominates. For values of $a$ greater than $1 \times 10^{-4}$ (μmol/mol)$^{-1}$ m$^{-1}$, the relative expanded uncertainty can be expressed as $U_{rel} \approx 2B^{1/2}$ as listed in Table 3.

5. Summary

In response to the growing interest in quantitative gas measurements using FTIR spectrometry, NIST has initiated a program to develop a quality-assured quantitative database of infrared spectra based on NIST prepared primary gas standards. The database currently has absorption coefficient data for twenty-one compounds that are listed in the 1990 USEPA Clean Air Act Amendment. For each compound, the absorption coefficient spectrum was calculated using nine transmittance spectra at 0.12 cm$^{-1}$ resolution and the Beer’s law relationship. The uncertainties in the absorption coefficient data were estimated from the linear regressions of the absorbance data and considerations of other error sources such as the nonlinear detector response.
Table 2. Type A and Type B standard uncertainties for each compound

| Compound    | Type A $\sigma_a = ma + b$ | Type B $u_{\text{total}}$ relative uncertainty |
|-------------|----------------------------|-----------------------------------------------|
| Slope       | Intercept                  |                                               |
| Benzene     | $1.3 \times 10^{-2}$       | $9.9 \times 10^{-8}$                         | 0.010 |
| Ethylene    | $2.6 \times 10^{-3}$       | $8.8 \times 10^{-8}$                         | 0.010 |
| Acetone     | $5.2 \times 10^{-3}$       | $4.9 \times 10^{-8}$                         | 0.010 |
| Ethanol     | $8.8 \times 10^{-4}$       | $2.8 \times 10^{-7}$                         | 0.010 |
| Methanol    | $2.0 \times 10^{-3}$       | $2.7 \times 10^{-7}$                         | 0.010 |
| 2-Propanol  | $2.0 \times 10^{-3}$       | $7.6 \times 10^{-8}$                         | 0.010 |
| Ethyl acetate | $1.9 \times 10^{-3}$     | $2.7 \times 10^{-7}$                         | 0.010 |
| $n$-Butanol | $7.9 \times 10^{-4}$       | $5.7 \times 10^{-7}$                         | 0.010 |
| Bromomethane | $1.0 \times 10^{-2}$     | $1.2 \times 10^{-3}$                         | 0.010 |
| Acetonitrile | $9.5 \times 10^{-4}$      | $7.3 \times 10^{-4}$                         | 0.010 |
| Ethylene oxide | $3.5 \times 10^{-3}$     | $1.6 \times 10^{-3}$                         | 0.010 |
| Propylene oxide | $3.0 \times 10^{-3}$     | $1.7 \times 10^{-3}$                         | 0.010 |
| Vinyl acetate | $2.3 \times 10^{-4}$     | $1.1 \times 10^{-3}$                         | 0.010 |
| Toluene     | $6.7 \times 10^{-3}$       | $2.1 \times 10^{-7}$                         | 0.010 |
| Ethyl acrylate | $9.0 \times 10^{-4}$     | $1.8 \times 10^{-7}$                         | 0.010 |
| Methyl ethyl ketone | $2.4 \times 10^{-3}$   | $8.4 \times 10^{-8}$                         | 0.010 |
| Ethyl tert–butyl ether | $2.3 \times 10^{-3}$ | $2.5 \times 10^{-7}$                         | 0.010 |
| Mean        | $3.7 \times 10^{-3}$       | $1.7 \times 10^{-3}$                         | 3.3 % |
| Standard deviation of the mean | $1.7 \times 10^{-4}$ | $6.3 \times 10^{-8}$                         | 2.1 % |

Table 3. Final uncertainty coefficients for each compound where the expanded uncertainty is expressed by, $U = 2(Ba^2 + Ca + D)^{1/2}$. For values of $a > 1 \times 10^{-4}$, the relative expanded uncertainty can be simplified to $U_{\text{rel}} = 2B^{1/2}$

| Compound | $B$ | $C$ | $D$ | $a > 1 \times 10^{-4}$ | Relative expanded uncertainty for $a > 1 \times 10^{-4}$ |
|----------|-----|-----|-----|----------------------|----------------------------------------------------------|
| Benzene  | $2.6 \times 10^{-4}$ | $4.2 \times 10^{-9}$ | $2.7 \times 10^{-14}$ | 3.3 % |
| Ethylene | $1.1 \times 10^{-4}$ | $8.5 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.1 % |
| Acetone  | $1.3 \times 10^{-4}$ | $1.7 \times 10^{-9}$ | $2.7 \times 10^{-14}$ | 2.3 % |
| Ethanol  | $1.0 \times 10^{-4}$ | $2.9 \times 10^{-9}$ | $2.7 \times 10^{-14}$ | 2.0 % |
| Methanol | $1.0 \times 10^{-4}$ | $6.6 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.0 % |
| 2-Propanol | $1.0 \times 10^{-4}$ | $6.5 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.0 % |
| Ethyl acetate | $1.0 \times 10^{-4}$ | $6.3 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.0 % |
| $n$–Butanol | $1.0 \times 10^{-4}$ | $2.6 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.0 % |
| Bromomethane | $2.0 \times 10^{-4}$ | $3.3 \times 10^{-9}$ | $2.7 \times 10^{-14}$ | 2.8 % |
| Acetonitrile | $1.0 \times 10^{-4}$ | $3.1 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.0 % |
| Ethylene oxide | $1.1 \times 10^{-4}$ | $1.2 \times 10^{-9}$ | $2.7 \times 10^{-14}$ | 2.1 % |
| Propylene oxide | $1.1 \times 10^{-4}$ | $9.8 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.1 % |
| Vinyl acetate | $1.1 \times 10^{-4}$ | $8.6 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.1 % |
| Toluene | $1.9 \times 10^{-4}$ | $3.0 \times 10^{-9}$ | $2.7 \times 10^{-14}$ | 2.0 % |
| Ethyl acrylate | $1.6 \times 10^{-4}$ | $1.1 \times 10^{-9}$ | $2.7 \times 10^{-14}$ | 2.5 % |
| Methyl tert–butyl ether | $1.6 \times 10^{-4}$ | $7.7 \times 10^{-10}$ | $2.7 \times 10^{-14}$ | 2.1 % |
| Sulfur dioxide | $1.1 \times 10^{-4}$ | $2.7 \times 10^{-4}$ | $2.7 \times 10^{-14}$ | 2.1 % |
For absorption coefficient values greater than $1 \times 10^{-4}$ ($\mu$mol/mol)$^{-1}$ m$^{-1}$, the average relative expanded uncertainty is 2.2%. Plots of the absorption coefficient data for the compounds currently in the NIST Quantitative Infrared Database are shown in Appendix B. Data are shown for spectra at 0.125 cm$^{-1}$ resolution with 3-term Blackman-Harris apodization. The data, at a number of resolutions and apodization functions, is available on compact disc in JCAMP-DX format with a viewer program. A digital signature accompanies each file, allowing users to ensure the integrity and source of the data file and traceability to NIST. Updates to the database are available over the internet.

6. Appendix A. Compounds in the 1990 U.S.EPA Clean Air Act Amendment

Table A1. List of the compounds in the 1990 U.S.EPA Clean Air Act amendment. Approximately 100 of these compounds have the vapor pressure required to prepare primary gas standards. The Chemical Abstracts Service Registry number is denoted by CAS No.

| CAS No. | Chemical name | CAS No. | Chemical name |
|---------|---------------|---------|---------------|
| 75070   | Acetaldehyde  | 67663   | Chloroform    |
| 60355   | Acetamide     | 107302  | Chloromethyl methyl ether |
| 75058   | Acetonitrile  | 126998  | Chloroprene   |
| 98862   | Acetophenone  | 1319773 | Cresols       |
| 53963   | 2-Acetylamidinothiophene | 95487 | $o$-Cresol |
| 107028  | Acrolein      | 108394  | $m$-Cresol    |
| 79061   | Acrylamide    | 106445  | $p$-Cresol    |
| 79107   | Acrylic acid  | 98828   | Cumene        |
| 107131  | Acrylonitrile | 94757   | 2,4-D, salts and esters |
| 107051  | Allyl chloride| 3547044 | DDE           |
| 92671   | 4-Aminobiphenyl| 334883 | Diazomethane |
| 62533   | Aniline       | 132649  | Dibenzofurans |
| 90040   | $o$-Anisidine | 96128   | 1,2-Dibromo-3-chloropropane |
| 1332214 | Asbestos      | 84742   | Dibutylphthalate |
| 71432   | Benzene       | 106467  | 1,4-Dichlorobenzene(p) |
| 92875   | Benzidine     | 91941   | 3,3-Dichlorobenzidine |
| 98077   | Benzocthloride| 111444  | Dichloretidyle ether |
| 100447  | Benzyl chloride| 542756 | 1,3-Dichloropropane |
| 92524   | Biphenyl      | 62737   | Dichlorvos    |
| 117817  | Bis(ethylyl)phthalate | 111422 | Diethanolamine |
| 542881  | Bis(chloromethyl) ether | 121697 | N,N-Diethylaniline |
| 75252   | Bromoform     | 66475   | Diethyl sulfate |
| 106990  | 1,3-Butadiene | 119904  | 3,3-Dimethoxybenzidine |
| 156627  | Calcium cyanamide | 60117 | Dimethyl aminooazobenzene |
| 105602  | Caprolactam   | 119937  | 3,3-Dimethyl benzidine |
| 133062  | Captan        | 79447   | Dimethyl carbamoyl chloride |
| 63252   | Carbazyl      | 68122   | Dimethyl formamide |
| 75150   | Carbon disulfide| 57147  | 1,1-Dimethyl hydrazine |
| 56235   | Carbon tetrachloride| 131113 | Dimethyl phthalate |
| 463581  | Carbonyl sulfide| 77781  | Dimethyl sulfate |
| 120809  | Catechol      | 534521  | 4,6-Dinitro-o-cresol |
| 133904  | Chloramben    | 51285   | 2,4-Dinitrophenol |
| 57749   | Chloradene    | 121142  | 2,4-Dinitrotoluene |
| 7782505 | Chlorine      | 13911   | 1,4-Dioxane |
| 79118   | Chloroaetic acid| 122667 | 1,2-Diphenyldihydrazine |
| 532274  | 2-Chloroaethylenophenone | 106898 | Epichlorohydrin |
| 106906  | Chlorobenzene | 106887  | 1,2-Epoxybutane |
| 510156  | Chlorobenzilate| 140885  | Ethyl acrylate |
| CAS No. | Chemical name          | CAS No. | Chemical name                  |
|---------|------------------------|---------|--------------------------------|
| 100414  | Ethyl benzene          | 62759   | N-Nitrosodimethylamine         |
| 51796   | Ethyl carbamate        | 59892   | N-Nitrosomorpholine            |
| 75003   | Ethyl chloride         | 56382   | Parathion                      |
| 106934  | Ethylene dibromide     | 32688   | Pentachloronitrobenzene        |
| 107062  | Ethylene dichloride    | 87865   | Pentachlorophenol              |
| 107211  | Ethylene glycol        | 108952  | Phenol                         |
| 151564  | Ethylene imine         | 106503  |  \( p \)-Phenylenediamine     |
| 75218   | Ethylene oxide         | 75445   | Phosgene                       |
| 96457   | Ethylene thiourea      | 803512  | Phosphine                      |
| 75343   | Ethyldiene dichloride  | 723140  | Phosphorus                     |
| 50000   | Formaldehyde           | 85449   | Phthalic anhydride             |
| 76448   | Heptachlor             | 133636  | Polychlorinated biphenyl       |
| 118741  | Hexachlorobenzene      | 1120714 | 1,3-Propane sulfone            |
| 87683   | Hexachlorobutadiene    | 57578   | Beta-Propiolactone             |
| 77474   | Hexachlorocyclopentadiene | 123386 | Propionaldehyde                |
| 67721   | Hexachloroethane       | 114261  | Propoxur                       |
| 822060  | Hexamethylene-1,6-diisocyanate | 78875 | Propylene dichloride          |
| 680319  | Hexamethylphosphoramide| 75569   | Propylene oxide                |
| 110543  | Hexane                 | 75558   | 1,2-Propylenimine              |
| 302012  | Hydrazine              | 91225   | Quinoline                      |
| 7647010 | Hydrochloric acid      | 106514  | Quinone                        |
| 7664393 | Hydrogen fluoride      | 100425  | Styrene                        |
| 123319  | Hydroquinone           | 96093   | Styrene oxide                  |
| 78591   | Isophorone             | 1746016 | 2,3,7,8-Tetrachlorodibenzo \( p \)-dioxin |
| 58899   | Lindane                | 79345   | 1,1,2,2-Tetrachloroethane      |
| 108316  | Maleic anhydride       | 127184  | Tetrachloroethylene            |
| 67561   | Methanol               | 7500450 | Titanium tetrachloride         |
| 72435   | Methoxychlor           | 108883  | Toluene                        |
| 74839   | Methyl bromide         | 95807   | 2,4-Toluene diamine            |
| 74873   | Methyl chloride        | 584849  | 2,4-Toluene diisocyanate       |
| 71556   | Methyl chloroform      | 95534   | \( o \)-Toluidine              |
| 78933   | Methyl ethyl ketone    | 800135  | Toxaphene                      |
| 60344   | Methyl hydrazine       | 120821  | 1,2,4-Trichlorobenzene         |
| 74884   | Methyl iodide          | 79005   | 1,1,2-Trichloroethane          |
| 108101  | Methyl isobutyl ketone | 79016 | Trichloroethylene              |
| 624839  | Methyl isocyanate      | 95954   | 2,4,5-Trichlorophenol          |
| 80626   | Methyl methacrylate    | 88062   | 2,4,6-Trichlorophenol          |
| 1634044 | Methyl \( tert \) butyl ether | 121448 | Triethylamine                  |
| 101144  | 4,4-Methylene bis(2-chloroaniline) | 1582098 | Trifluralin                    |
| 75092   | Methylene chloride     | 540841  | 2,2,4-Trimethylpentane         |
| 101688  | Methylene diphenyl     | 108054  | Vinyl acetate                  |
| 101779  | 4,4-Methyleneedianiline| 593602 | Vinyl bromide                  |
| 91203   | Naphthalene            | 75014   | Vinyl chloride                 |
| 98953   | Nitrobenzene           | 75554   | Vinylindene chloride           |
| 92933   | 4-Nitrobenzenephenyl   | 1330207 | Xylenes                        |
| 100027  | 4-Nitrofenol           | 95476   | \( o \)-xylene                 |
| 79469   | 2-Nitropropane         | 108383  | \( m \)-xylene                 |
| 684935  | N-Nitro-N-methylurea   | 106423  | \( p \)-xylene                  |
7. Appendix B. Absorption Coefficient Data

The following figures, Fig. B1 through B21, show the absorption coefficient data for compounds currently in the NIST Quantitative Infrared Database, SRD 79. In all cases the data represent spectra at 0.125 cm\(^{-1}\) resolution with 3-term Blackman-Harris apodization. In all cases, the absorbance was defined as \(-\log_{10}(I/I_0)\).

**Fig. B1.** Benzene.

**Fig. B2.** Ethylene.
Fig. 3. Acetone.

Fig. B4. Ethanol.
Fig. B5. Methanol.

Fig. B6. 2-Propanol.
Fig. B7. Ethyl acetate.

Fig. B8. n-Butanol.
Fig. B9. Bromomethane.

Fig. B10. Acetonitrile.
Fig. B11. Ethylene oxide.

Fig. B12. Propylene oxide.
Fig. B13. Methyl ethyl ketone.

Fig. B14. Ethyl tert-butyl ether.
Fig. B15. 1,3-Butadiene.

Fig. B16. Acrylonitrile.
Fig. B17. Vinyl acetate.

Fig. B18. Toluene.
Fig. B19. Ethyl acrylate.

Fig. B20. Methyl tert-butyl ether.
Acknowledgments

The authors wish to thank S. J. Wetzel, D. vanVlack, and C. L. Calhoun for their help with the FTIR data acquisition and analysis and C. Mack and S. Margolis for the analysis of the pure VOCs. Additionally, the authors would like to thank J. C. Travis, B. Phillips, G. Russwurm, W. Bell, P. Woods, P. Hanst, W. Herget, R. Kagann, R. Richardson, and P. Griffiths for many helpful discussions regarding this work.

9. References

[1] C. Henry, Analytical Chemistry News & Features, April 1, 1998, p. 273 A.
[2] (Proposed) Test Method 320 Measurement of Vapor Phase Organic and Inorganic Emissions by Extractive Fourier Transform Infrared Spectroscopy, Federal Register, Part 63, Appendix A.
[3] Compendium Method TO-16, Long-Path Open-Path Fourier Transform Infrared Monitoring of Atmospheric Gases, EPA Contract 68-D5-0049, NERL, U.S.EPA, Research Triangle Park, NC.
[4] Federal Clean Air Act, Title II, Public Law 101-549.
[5] R. L. Richardson, Jr. and P. R. Griffths, Appl. Spectros. 52, 143–153 (1998).
[6] Quantitative Infrared Vapor Phase Spectra, Contract No. 68D90055 (U.S. Environmental Protection Agency, Emission Measurement Branch, Research Triangle Park, North Carolina).
[7] Rothman L. S., Rinsland C. P., Goldman A, Massie S. T., Edwards D. P., Fland J.-M., Perrin, A., Camy-Peyret C., Dana V., Mandin J.-Y., Schroeder J, McCann A., Gamache R. R., Waterson R. B., Yoshino K., Chance K. V., Jucks, K. W., Brown, L. R., Nemtchinov V., Varanasi, P. 1998 The HITRAN Molecular Spectroscopic Database and HAWKS (HITRAN Atmospheric Workstation): 1996 Edition Journal of Quantitative Spectroscopy and Radiative Transfer 60 (in press). See also Phillips Laboratory/Geophysics Directorate, Hanscom AFB, MA; http://www. HITRAN. com.
[8] P. L. Hanst and S. T. Hanst, Infrared Spectra for Quantitative Analysis of Gases Infrared Analysis, Inc., Anaheim, CA (1996).
[9] Gas Phase Infrared Spectral Standards, MIDAC Corporation, Irvine, CA 92614.
[10] J. F. Sprouse, Quantitative Infrared Spectral Database, Sprouse Scientific Systems, Charlotte, North Carolina (1991).
[11] R. S. McDonald and P. A. Wilks, Jr. Appl. Spectros. 42, 151–162 (1988).
[12] P. M. Chu, G. C. Rhoderick, W. J. Lafferty, and F. R. Guenther, Proceedings of the Air & Waste Management Association, 89th Annual Meeting and Exhibition, Nashville, Tennessee (1996). See also, W. P Schmidt and H. L Rook, Anal. Chem. 55, 290–294 (1983); G. C. Rhoderick and W. L. Zielinski, Anal. Chem. 60, 2454–2466 (1988); G. C. Rhoderick Fresenius, J. Anal. Chem. 341, 521–531 (1991).
[13] R. L. Sams, private communication.
[14] W. J. Lafferty and A. S. Pine, G. Hilpert, R. L. Sams, and J.-M. Flaud, J. Mol. Spectrosc. 176, 280–286 (1996).
[15] W. J. Lafferty, R. L. Sams, and J. M. Flaud, 88th Annual Meeting of the Air and Waste Management Association, San Antonio, Texas (1995).
[16] R. A. Toth, J. Opt. Soc. Am. B 8, 2236–2255 (1991). R. A. Toth, J. Opt. Soc. Am. B, 10, 2006–2029 (1993). [17] P. R. Griffiths, J. A. deHaseth, Fourier Transform Infrared Spectrometry, John Wiley & Sons, Inc., New York (1986).
[18] OPUS 2.2, Bruker Instruments, Billerica, MA 01821.
[19] Deresolve, W. J. Phillips and Arnold Engineering Development Center, TN 37389-4300. See also see http://www.epa.gov/ttn/emc/fhir/deresolv.html.
[20] Guide to the Expression of Uncertainty in Measurement, ISBN 92-67-10188-9, 1st Ed. ISO, Geneva, Switzerland, (1993); see also B. N. Taylor and C. E. Kuyatt, Guidelines for Evaluating and Expressing the Uncertainty of NIST Measurement Results, NIST Technical Note 1297, U. S. Government Printing Office, Washington, DC (1994).
[21] T. H. Wampler, J. Res. Natl. Bur. Stand. (U.S.) 73B, 59–80 (1969) and references therein.
[22] W. R. Miller and G. C. Rhoderick, Fresenius J. Anal. Chem. 351, 221–229 (1995).

About the authors: P. M. Chu, F. R. Guenther, and G. C. Rhoderick are Research Chemists in the Analytical Chemistry Division of the Chemical Sciences and Technology Laboratory. W. J. Lafferty is a Scientist Emeritus in the Optical Technology Division of the Physics Laboratory. The National Institute of Standards and Technology is an agency of the Technology Administration, U. S. Department of Commerce.