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Widely-tunable and sensitive optical sensor for multi-species detection in the mid-IR

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Abstract

Pulsed cavity ringdown spectroscopy (CRDS) technique was used to develop a novel widely-tunable laser-based sensor for sensitive measurements of ethylene, propene, 1-butene and allene in the mid-IR. The use of an external-cavity quantum cascade laser (EC-QCL) enabled the sensor to cover a wide wavelength range from 10 to 11.1 µm (900 – 1000 cm⁻¹) to detect multiple gases relevant to combustion and environment. The sensor operation was validated in a room-temperature static cell using well-characterized absorption lines of carbon dioxide near 938.69 cm⁻¹ and 974.62 cm⁻¹. Detection limits for ethylene, propene, 1-butene, and allene were measured to be 17, 134, 754 and 378 ppb, respectively, at 296 K and 760 Torr for a single-pass path-length of 70 cm with averaging time of 4 ms. The excellent sensitivity of the optical sensor enabled it to measure the aforementioned gases at levels smaller than 1% of their recommended exposure limits. To the best of our knowledge, this is one of the first successful applications of the pulsed CRDS technique to measure trace levels of multiple gases in the 10 – 11 µm wavelength region.

Keywords: Multi-species detection, cavity ringdown spectroscopy, air pollution monitoring, combustion diagnostics

1. Introduction

The monumental advancements in laser technology over the past few decades have made laser-based sensors ubiquitous in a wide array of applications including noninvasive medical diagnostics [1–3], environmental pollution monitoring [4,5], industrial process control [6,7], detection of explosives [8,9] and several others. Laser sensors have gained attention primarily due to their ability to carry out real-time, non-intrusive measurements of species with high sensitivity and selectivity. Recent improvements in laser technology such as the advent of external cavity quantum cascade lasers (EC-QCLs) with wide tuning...
ranges of the order of 100 cm\(^{-1}\) enable spectroscopic measurements of broadband absorbers as well as multiple species with a single laser. The ability of an EC-QCL based sensor to measure multiple species owing to its extensive tuning range is an enormous advantage when compared with competing diode laser-based sensors which can usually measure only a single species because of their narrow operating ranges. A widely-tunable laser when coupled with a mirror cavity that retains high reflectivity over a broad wavelength range enables trace detection of multiple species and has applications in environmental pollution monitoring, industrial gas sensing and combustion. In this vein, we have developed and validated a novel laser-based sensor that uses a widely tunable EC-QCL coupled to a moderately high finesse mirror cavity to obtain high sensitivity over a wide wavelength range from 10 to 11.11 µm (1000 – 900 cm\(^{-1}\)). We demonstrate the high sensitivity of the sensor over a wide tuning range by measuring trace amounts of ethylene, propene, allene, and 1-butene, individually and in a multi-gas mixture.

Ethylene, propene, allene, and 1-butene play a crucial role in the oxidation and pyrolysis of large hydrocarbons, either as intermediates [10–12] or as precursors to a host of radical intermediates [13]. Ethylene and propene also play a significant role in the formation and growth of polycyclic aromatic hydrocarbons which are primary precursors of soot particles. Detection of trace amounts of volatile organic compounds such as ethylene and propene is vital in the study of air quality. Industrial emissions, waste incineration plants and vehicle exhaust fumes are the major sources of ethylene and propene in the atmosphere. Ethylene and propene can rapidly undergo oxidation reactions producing ground-level ozone [14,15], which has adverse effects on crops and human health [16–19]. Moreover, the aforementioned target molecules are widely used as feedstock or intermediates in chemical and petroleum industries and thereby a single diagnostic that can measure all four gases in trace amounts will be invaluable in their leak detection.

Ethylene has previously been measured in trace concentrations using a variety of sensitive absorption techniques. Photoacoustic spectroscopy has been extensively used for monitoring trace ethylene in a number of studies. Altuzar et al. [20] developed a CO\(_2\) gas laser based photoacoustic spectrometer to measure ethylene concentrations in air samples from different locations in Mexico city. Weidmann et al. [5] developed an ethylene sensor based on direct absorption in a multi-pass Herriot cell to monitor ethylene from vehicle exhaust and in air samples from a busy urban tunnel. They reported a noise equivalent sensitivity of 30 ppb for ethylene with 80 seconds of total acquisition time. Wahl et al. [21] developed an ultra-sensitive ethylene sensor based on cw-cavity ringdown spectroscopy technique to measure trace ethylene levels in a fruit packing facility. They reported detection limit of 2 ppb with 4.4 minutes of measurement time. Manne et al. [22,23] performed a proof-of-concept study to measure ethylene using four
strategies, namely pulsed cavity ringdown spectroscopy, cavity-enhanced spectroscopy [22], intrapulse and interpulse spectroscopic techniques with a multi-pass cell [23]. They reported detection limits in the range of 5 ppb to 130 ppb with 5 to 10 seconds of integration time.

Recently, Chrystie et al. [15] developed a laser-based sensor to non-intrusively measure propene concentrations in a chemical reactor (shock tube). In general, there has been little work on the development of laser-based sensors for the detection of propene, allene or 1-butene. Detection of propene, allene, and 1-butene has traditionally been carried out using mass spectrometry [12] or gas chromatography techniques [13,24]; both of which require sampling using physical probes. The narrow tuning range of previously used laser sources generally limited the optical sensors to the detection of a single molecule and this probably made laser sensors unappealing as compared to the gas sampling techniques. The wide tuning range and high sensitivity of the current sensor, coupled with the traditional advantages of laser-based sensors such as non-intrusiveness, portability, robustness and rapid response, make it better suited for environmental monitoring and industrial gas sensing applications.

In this work, we have implemented pulsed cavity ringdown spectroscopy (pulsed CRDS) with ringdown mirrors of 99.5% reflectivity. The wide tuning range of the EC-QCL coupled with the relatively flat reflectivity curve of the ringdown mirrors used here allowed us to develop a sensor with a broad wavelength operating range. In what follows, we present a detailed description of the design of our sensor. We then describe the validation of the sensor using CO₂ gas and describe sensor optimization. We then demonstrate the sensor to measure small amounts of ethylene, propene, allene and 1-butene. Finally, we show the capability of the sensor to measure trace amounts of gases in a multi-gas mixture. To our knowledge, this is the first implementation of pulsed CRDS with EC-QCL to achieve a highly sensitive and widely tunable sensor in the mid-IR (900-1000 cm⁻¹). Additionally, this work represents the first laser-based measurements of trace amounts of propene, allene, and 1-butene.

2. Theory of cavity ringdown spectroscopy

Cavity ringdown spectroscopy (CRDS) was introduced almost three decades back, in 1988, by O’Keefe and Deacon [25] and since then has been extensively used for sensitive detection of gas phase [26–30] and liquid phase samples [31,32]. Cavity ringdown may be implemented with pulsed or continuous-wave lasers to develop highly sensitive gas sensors. A brief description of CRDS implemented with pulsed lasers is given here. In pulsed CRDS, the light pulse from a laser is admitted into a stable optical cavity configuration and undergoes numerous passes. When the laser pulse ends, the light inside the cavity undergoes an exponential
decay. The exponential decay rate is dependent on the reflectivity of mirrors forming the cavity, the length of the cavity, and absorption losses inside the cavity. The time constant of the exponential decay or the ringdown time is the time taken for the light intensity to fall to 1/e of its initial value. It is given as:

$$\tau = L / (c[(1 - R) + \alpha L])$$  \hspace{1cm} (1)$$

In the absence of any absorbing molecules inside the cavity, the ringdown time is given as:

$$\tau_0 = L / (c[(1 - R)])$$  \hspace{1cm} (2)$$

In the above equations, \(L\) is the total length of the cavity, \(c\) is the speed of light, \(R\) is the reflectivity of the mirrors forming the cavity and \(\alpha\) is the absorption coefficient in cm\(^{-1}\). The absorption coefficient can be calculated from Eq. (1) and Eq. (2) as:

$$\alpha = \frac{1}{c} \left( \frac{1}{\tau} - \frac{1}{\tau_0} \right)$$  \hspace{1cm} (3)$$

The absorption coefficient, \(\alpha\), is a function of the total pressure of the gas, \(P\), the mole fraction of the absorbing species, \(X\), the linestrength, \(S\), and the lineshape \(\phi_\nu\):

$$\alpha = S(T)PX\phi_\nu(P, T)$$  \hspace{1cm} (4)$$

Equation (4) enables the determination of the mole fraction of the absorbing species when spectroscopic parameters (\(S\) and \(\phi_\nu\)) are known.

### 3. Experimental apparatus

The experimental setup for the designed sensor is shown in Fig.1. We used a pulsed EC-QCL (Daylight Solutions 11100-UT), tunable over 9.53-12.95 \(\mu\)m (775-1020 cm\(^{-1}\)), with a maximum average power of 15 mW. Laser pulse widths can be varied between 40 to 500 ns and repetition frequencies as high as 100 kHz are possible. Single mode emission at the wavelength of interest was achieved by adjusting the laser head temperature and diffraction grating orientation via the laser controller. Single mode emission was ensured by sending the laser beam to a spectrum analyzer (Bristol 721). The IR beam emitted by EC-QCL was co-aligned with a 10 mW He-Ne laser (632 nm) to facilitate the cavity alignment process. The IR beam (beam diameter \(= 2\) mm) emerging from the EC-QCL head was directed into an optical cavity formed by two plano-concave mirrors (II-VI Infrared) with 1 m radius of curvature, 1" diameter, and a manufacturer specified reflectivity of 99.5% at 10.6 \(\mu\)m (943.39 cm\(^{-1}\)). The ringdown mirrors were placed 70 cm apart to
form a stable optical cavity configuration. A plano-convex lens (ZnSe, 500 mm focal length) was positioned between the EC-QCL head and the ringdown cavity to improve the mode-matching between the laser beam and the TEM$_{00}$ mode of the cavity. A concave mirror (Aluminum, 100 mm focal length) stationed at the exit of the optical cavity was used to focus the IR beam on a thermoelectrically-cooled photovoltaic detector (Vigo PVMI 3TE-10.6) with a bandwidth of 10 MHz. The detector signal was recorded with a digital oscilloscope (Tektronix DPO 3014) having a sampling rate of 2.5 GS/s.

Although the EC-QCL can emit IR radiation over a wide wavelength range (9.53-12.95 µm), its tuning range is not mode-hop free. Therefore, laser head temperature and grating orientation had to be iteratively tuned to avoid mode-hops. However, the strategy to tune temperature and diffraction grating was not successful for all desired wavelengths. Despite this limitation, the wide tunability of the EC-QCL proved to be an enormous advantage in our quest to develop a sensor capable of detecting multiple species in trace amounts. The output wavelength of the EC-QCL was quite reproducible from one day to next and the drift in the emitted wavelength was negligible over long periods (> 8 hours) of continuous operation.

![Experimental setup used for pulsed CRDS measurements in the static cell. BS: Beam Splitter, MMO: Mode matching optic.](image)

Figure 1. Experimental setup used for pulsed CRDS measurements in the static cell. BS: Beam Splitter, MMO: Mode matching optic.
To make the best use of the energy available from the pulsed laser, the laser pulse width was varied to identify its optimum value. The frequency chirp of the laser or the laser linewidth increases from 0.032 cm$^{-1}$ at 80 ns pulse width to 0.08 cm$^{-1}$ at 120 ns pulse width, as measured in an earlier study [30]. It is desirable to minimize frequency chirp as it can lead to artificial broadening of narrow spectral lines. However, high detector signal is also a necessity. Therefore, to minimize frequency chirp while ensuring high SNR of the detector signal, laser pulse width of 100 ns having a frequency chirp of 0.064 cm$^{-1}$ was used throughout this work.

The gas cell, with the two ringdown mirrors at its two ends, was connected to a gas tank, a vacuum pump, and two calibrated pressure transducers (MKS 627D capacitance manometers with 1000 and 10000 Torr full-scale pressure range and accuracy of 0.12%). High purity propene (99.95%), allene (99.5%), 1-butene (99.95%) gases, dilute mixture of ethylene (1051 ppm ethylene in argon), dilute mixture of carbon dioxide (1.05% carbon dioxide in nitrogen), ultra-pure nitrogen (99.999%), and Ultra-pure argon (99.999%) were used for the experiments. The diluted mixtures of individual gases were prepared manometrically using a gas handling system. The pressure in the gas handling system was monitored by two pressure transducers (MKS 627D capacitance manometers with 100 and 1000 Torr full scale range). In the majority of cases, multiple dilution steps were required to achieve the desired concentration of target gases.

4. Sensor validation and optimization

The sensor was validated using carbon dioxide as a test molecule because its spectroscopic parameters (linestrength, lineshape) are very well-known [33]. Carbon dioxide has relatively strong absorption lines in the 930 – 985 cm$^{-1}$ wavelength region and the ringdown mirrors have their maximum reflectivity near 944 cm$^{-1}$. However, ethylene has a very strong vibrational band ($\nu_7$) centered near 949.533 cm$^{-1}$ and even trace amount of ethylene, present as an impurity, can cause appreciable interference. Therefore, simulations were performed using the HITRAN database [33] to identify absorption lines of CO$_2$ where the interference from ethylene is minimal. Subsequently, two absorption lines of CO$_2$ centered near 938.69 cm$^{-1}$ ($(1\ 0\ 0\ 01) \rightarrow (0\ 0\ 0\ 11)$, P26) and 974.62 cm$^{-1}$ ($(1\ 0\ 0\ 01) \rightarrow (0\ 0\ 0\ 11)$, R18), having negligible interference from ethylene, were selected to demonstrate the wide tuning range of this sensor.

With a view to minimizing the effect of artificial broadening due to the frequency chirp of the laser, absorption measurements of CO$_2$ were performed at a relatively high pressure of 1200 Torr. Figure 2 shows a comparison between the simulated and measured absorption coefficients for the absorption lines centered near 938.69 cm$^{-1}$ and 974.62 cm$^{-1}$. Each measured data point was obtained by using Eq. (3) after determining
the ringdown time for pure N\textsubscript{2} (τ\textsubscript{0}) and for a 1.05% CO\textsubscript{2}/N\textsubscript{2} mixture (τ). The measured absorption coefficients (red square) agree very well with the simulated absorption coefficients (black line) in both wavelength regions, thereby validating the pulsed-CRDS setup.

![Absorption Coefficient vs Wavenumber](image)

Figure 2. Comparison of measured absorption coefficients (red squares) with HITRAN simulations [33]. Mixture: 1.05% CO\textsubscript{2}/N\textsubscript{2}; T = 296 K; P = 1200 Torr.

Ringdown times were measured in an optical cell filled with ultra-pure Argon (760 Torr pressure) across the entire wavelength range of the sensor (900 – 1000 cm\textsuperscript{-1}) in steps of ~ 10 cm\textsuperscript{-1}. Figure 3 shows the measured ringdown time and the calculated mirror reflectivity as a function of wavenumber. The mirror reflectivity at 950 cm\textsuperscript{-1} was found to be 99.45% which compares well with the manufacturer quoted value of 99.5%. The ringdown time and mirror reflectivity are largest at the center (950 cm\textsuperscript{-1}) of the mirror curve and drop off on either side. However, the variation is not too significant and the sensor can be used to make species measurements over the entire target wavelength range of 900 – 1000 cm\textsuperscript{-1}.
With a view to determine the minimum detection limit and optimum averaging time, Allan Variance was calculated at each wavelength of interest. The Allan Variance was computed using the decay rate (inverse of ringdown time) and was scaled by the speed of light to make it equivalent to absorption coefficient [34]. The static cell was filled with ultra-pure argon to a pressure of 760 Torr, and the detector signal was recorded at each wavenumber step for 4000 ring-down events which corresponds to a time period of 40 ms (laser repetition rate is 100 kHz). In the current work, Allan Variance was calculated using the new algorithm proposed by Huang and Lehmann [34] as it leads to significant reduction in noise and more accurate determination of the optimum averaging time when compared with the traditional algorithm. The Allan Variance drops very steeply initially and reaches minima near an averaging time of ~ 4 ms (corresponding to 400 ringdown events) for almost all the wavenumbers, as shown in Fig.4. Huang and Lehmann[34] had obtained a similar optimal averaging size of 500 ringdown events in their work; however, their corresponding optimal averaging time was 72 s because of the significantly larger time-gap between successive ringdown events. The high repetition rate of the laser (100 kHz) used in our work enables us to achieve millisecond time resolution even after averaging 400 events and this feature is valuable for carrying out real-time measurements of species in transient environments. For each ringdown measurement reported here, the extracted ringdown times were averaged over 400 ringdown events. For the averaging time of 4 ms, the Allan Variance is less than 1.9 x 10^{-7} cm^{-1} across the entire wavelength range of the sensor. It is to be noted that the averaging time of 4 ms is required to measure the absorbance at a specific wavelength and
is not the time resolution of the diagnostic to measure the concentration of a given species. Measuring the
concentration of a given species in a mixture necessitates the measurement of $\tau$ and $\tau_0$ at one or more
wavelengths, thereby increasing the time required to determine the concentration of a species. In the current
diagnostic, since the laser has to be tuned manually from one wavelength to the next, it is not possible to
assign a time resolution. However, for the case of the concentration being extracted from absorbance
measurement at a single wavelength, the concentration measurements will have a time resolution of 4 ms.

Figure 4. Allan Variance as a function of averaging time for different wavenumbers. The gray vertical line indicates the
location of minimum Allan variance.

The high sensitivity of the sensor over a wide tuning range of 100 cm$^{-1}$ makes it an ideal diagnostic to
measure trace concentrations of multiple gases. To demonstrate the capability of the sensor for multi-species
detection, four hydrocarbon molecules (ethylene, propene, allene and 1-butene) with strong absorption
bands in the operating range of the sensor were selected (Fig.5). Due to the frequency chirp of the laser,
estimated to be 0.064 cm$^{-1}$ for a pulse width of 100 ns, the sensor is best suited to detect molecules with
broad spectral features as selected here. Figure 6 shows simulated absorption coefficients for trace amounts
of the four molecules at a nominal temperature of 296 K and pressure of 760 Torr. The simulated absorption
coefficients shown here were obtained by scaling the absorbance measurement reported by Pacific
Northwest National Laboratory (PNNL) database [35] assuming a path length of 1 cm. At the given
experimental conditions, the peak absorption coefficients for the ppm-level concentration of all gases are
larger than 3.7 x 10$^{-5}$ cm$^{-1}$ which is about two orders of magnitude higher than the sensor detection limit of
1.9 x 10$^{-3}$ cm$^{-1}$, thereby enabling the sensor to measure trace amounts of these hydrocarbon gases. The
absorption peaks for all molecules are very well separated except the peaks of propene and 1-butene which
lie within 0.9 cm$^{-1}$ of each other. Ethylene has strong absorption band ($\nu_7$) centered near 949.41 cm$^{-1}$ that
can be attributed to the CH\textsubscript{2} wagging, while propene has strong absorption band (v\textsubscript{19}) centered near 912.62 cm\textsuperscript{-1} due to the CH\textsubscript{2} twisting and wagging [36]. Strong peaks of 1-Butene are observed near 911.71 cm\textsuperscript{-1} which are attributed to the CH\textsubscript{2} out-of-plane bending mode [37]. Allene absorption near 901.8 cm\textsuperscript{-1} is assigned to one of the weaker lines of the v\textsubscript{4} vibrational band centered at 865 cm\textsuperscript{-1} which is attributed to the CH\textsubscript{2} twisting [38]. Absorption measurements for all four species were carried out near the peaks of their absorption coefficients.

Figure 5. Simulated absorption coefficients of selected target gases. Mole fraction = 1 ppm; T = 296 K; P = 760 Torr.
5. Results and discussion

5.1 Trace Gas Mixtures

The capability of the sensor to measure trace concentrations was initially demonstrated by measuring small amounts of ethylene, propene, allene and 1-butene individually. Measured absorption coefficients were compared with the PNNL database [35] as it is the most reliable source for the absorption cross-sections of these molecules. All measurements reported here were performed at a nominal temperature of 296 K and pressure of 760 Torr; measured data are shown in Fig. 7. As discussed earlier, the mode-hop of the laser does not permit absorption coefficient measurements at constant wavelength intervals. The measured absorption coefficients for all four hydrocarbons agree very well with the PNNL database which confirms the sensor accuracy and sensitivity to trace amounts of target species.
Figure 7. Comparison of measured absorption coefficients (symbols) with the PNNL Database (lines) [35]. T = 296 K; P = 760 Torr.

Representative raw detector signals for the ringdown events of all four target molecules and single-exponential fits to the experimental data are shown in Fig. 8. The single-exponential function fits very well to the measured detector signals for all gases used in this study and over a range of concentrations. The ringdown event for the 5.6 ppm ethylene/argon mixture represents the largest absorption coefficient measured in this study, and the high quality of single-exponential fit for the small ringdown time of 113 ns demonstrates that the frequency chirp of the laser does not cause multi-exponential decays or other non-idealities at the experimental conditions used in this study (296 K and 760 Torr).
5.2 Detection Limits

Detection limits for individual gases were obtained by successively diluting gas mixtures in argon bath gas and measuring the absorption coefficient at each dilution step till the measured absorption coefficient reached an asymptotic value, as shown in Fig. 9 for propene. Similar trends were obtained for the other 3 molecules and the results are listed in Table 1. The detection limits for all gases follow the same trend as observed in Allan Variance analysis (Fig. 4) with allene measured near 902.4 cm\(^{-1}\) having the largest value of the detection limit, and ethylene measured near 950.08 cm\(^{-1}\) having the lowest detection limit. However, the detection limits calculated using Allan Variance analysis are an order of magnitude smaller than the values obtained using mixture dilution method. Mixture dilution method leads to larger values for detection.
limits because it is affected by uncertainties in the ringdown times for both the absorber gas and the bath gas (argon), whereas the Allan Variance analysis only considers the uncertainties in the ringdown times of the bath gas. Therefore, the mixture dilution method provides more realistic values of the detection limits and has been subsequently used to report the minimum detection limit for each molecule at a nominal temperature of 296 K and pressure of 760 Torr (Table 1). The detection limits in terms of ppb were calculated at the peak absorption coefficient for each molecule within the operating wavelength range of the sensor and were scaled from the PNNL database for each molecule. The detection limits presented here provide a very good estimate of the uncertainty in the absorption measurements reported in this study.

![Absorption Coefficient vs Dilution Steps](image)

**Figure 9.** Absorption coefficient as a function of mixture (propene / Ar) dilution. T = 296 K; P = 760 Torr.

**Table 1. Detection limits determined using the mixture dilution method.**

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Wavenumber [cm⁻¹]</th>
<th>Detection Limit [cm⁻¹]</th>
<th>Detection Limit [ppb]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allene</td>
<td>902.4</td>
<td>6 x 10⁻⁶</td>
<td>754</td>
</tr>
<tr>
<td>1-Butene</td>
<td>912.71</td>
<td>2.8 x 10⁻⁶</td>
<td>378</td>
</tr>
<tr>
<td>Propene</td>
<td>912.11</td>
<td>2.6 x 10⁻⁶</td>
<td>134</td>
</tr>
<tr>
<td>Ethylene</td>
<td>950.08</td>
<td>7.6 x 10⁻⁷</td>
<td>17</td>
</tr>
</tbody>
</table>
Here, we discuss the detection limit of our sensor with other competing methods. Assuming an optimistic value of detection limit of 1% absorbance using direct absorption spectroscopy [39], the minimum detection limit for 70 cm static cell will be $1.43 \times 10^5 \text{ cm}^{-1}$, which translates to 2.38 to 18.8 times lower sensitivity compared to the pulsed CRDS sensor described in the current study. Since the pulsed CRDS sensor covers the wavenumber region containing highly strong absorption bands for all molecules of interest (Fig. 5), the current sensor will be more sensitive compared to laser sensors in other wavelength regions, such as in the near-IR as long as mirrors with similar reflectivity are utilized. It is pertinent to point out that higher reflectivity mirrors are easily available in the near-IR region, which when coupled with a cw laser to develop cw-cavity ringdown spectroscopy based sensor may achieve higher detection limits [21]. However, the spectral features of hydrocarbons are not as well separated in the near-IR region and interference from CO$_2$/H$_2$O tend to more problematic.

Fourier Transform Infrared (FTIR) spectrometer studies have typically been carried out to measure absorption cross-section of various absorption bands and use integrated absorbance [36–38, 40], though single-point absorbance measurements as used in the current study are also possible. FTIR spectrometers have few advantages over laser-based sensors notably their ability to cover wide wavelength ranges, typically from near IR to mid IR. However, FTIR measurements typically require a lot of averaging which limits their time resolution, and FTIR spectrometers tend to be bulky thereby making these unfeasible for field measurements as compared to the compact laser-based sensors. Although the QCL system used in the current work had some mode-hop issues, improvements in QCL technology are happening at a rapid pace and the new systems are expected to sweep through large wavelength regions rapidly without mode-hops.

### 5.3 Multi-gas Mixtures

The ability of the sensor to detect multiple species simultaneously was demonstrated by applying the sensor to a multi-gas mixture containing trace amounts of the target gases. The multi-gas mixture composition was selected to contain gases at 1% of their 8-hour time-weighted-average (TWA) exposure limits. The TWA exposure limits for ethylene, propene, and allene are 200 ppm [41], 500 ppm [41], and 1000 ppm [42], respectively. Therefore, a mixture containing 2.2 ppm ethylene, 5.2 ppm propene, and 10 ppm allene was prepared manometrically in argon bath gas. Multiple dilution steps were used to prepare such a mixture. Figure 10 shows the comparison of measured absorption coefficients with those calculated using the PNNL database. The total calculated (PNNL) absorption coefficient of the mixture is shown in gray triangles, while the measured total absorption coefficient is marked with red hollow circles. The total absorption coefficient was measured at 4 specific wavenumbers – three near the peaks of the three species and one near an off-line ethylene frequency. The concentration of ethylene may be calculated using differential absorption strategy since the contribution of allene and propene to the total absorbance at the on-peak and off-peak frequencies of ethylene are almost identical and can thus be subtracted. Using this methodology, the
ethylene concentration was calculated to be 2.3 ppm, which is very close to the manometric value of 2.2 ppm. The total absorption coefficient measured at the two other wavenumbers (901.55 cm\(^{-1}\) and 911.983 cm\(^{-1}\)) may be written as:

\[
\alpha_x x_e + \alpha_p x_p + \alpha_a x_a = \text{Total absorption coefficient} \tag{5}
\]

where \(\alpha\) is the absorption coefficient and \(x\) is the mole fraction. The subscripts \(e, p,\) and \(a\) represent ethylene, propene, and allene, respectively. Writing Equation (5) for both wavenumbers (901.55 cm\(^{-1}\) and 911.983 cm\(^{-1}\)), and substituting in the absorption coefficients for all gases and mole fraction of ethylene, results in a system of two linear equations containing two unknowns, the mole fraction of propene \((x_p)\) and the mole fraction of allene \((x_a)\). The equations are shown below for 901.55 cm\(^{-1}\) and 911.983 cm\(^{-1}\), respectively:

\[
1.45 x_p + 2.8 x_a = 38.12 \\
10.4 x_p + 1.16 x_a = 65.52
\]

On solving the above set of linear equations, we obtain \(x_p\) to be 5.1 ppm and \(x_a\) to be 11 ppm. The mole fraction of propene is within 2% of the expected value of 5.2 ppm, whereas the mole fraction of allene is 10% higher than the expected value of 10 ppm. The overestimation of allene is possibly due to the contribution of impurities (other hydrocarbons such as butene in the allene/argon mixture) to the total measured absorbance near 901.55 cm\(^{-1}\). Mode-hopping of the EC-QCL near the allene peak did not permit measurements near the allene peak (901.85 cm\(^{-1}\)). These measurements demonstrate the capability of the sensor to accurately measure trace amounts of hydrocarbon gases in a multi-gas mixture, which makes this sensor an attractive candidate for environmental pollution monitoring and gas sensing in petrochemical industries.
Additionally, to demonstrate the potential of this sensor to measure trace species in the atmosphere, ethylene concentrations were measured in air samples collected from our laboratory and from an automobile parking garage. Due to the potential interference from carbon dioxide at 949.47 cm\(^{-1}\), these ethylene measurements were carried out at a nearby frequency of 949.835 cm\(^{-1}\). The sample collected from the automobile garage had significantly larger concentration of ethylene (415 ppb) compared to the sample collected from laboratory air (55 ppb). The elevated levels of ethylene in the garage air sample are attributed to the unburnt hydrocarbon exhaust from automobiles and lie within the previously reported values of 17 to 800 ppb measured in urban areas [43].

The pulsed CRDS diagnostic developed in this study is able to achieve sub-ppm detection limits for ethylene, propene, allene and 1-butene. Further improvements in absorption sensitivity are possible by utilizing higher reflectivity mirrors in conjunction with higher power lasers [30]. A laser source that does not suffer from mode-hops will also improve the overall operation of the sensor.
Conclusions

A novel widely-tunable sensor based on pulsed cavity ringdown spectroscopy has been successfully developed and used for performing sensitive absorption measurements of ethylene, propene, allene and 1-butene in the mid-IR. The high sensitivity of the laser-based sensor enabled detection limits of 17 ppb, 134 ppb, 754 ppb and 378 ppb for ethylene, propene, allene and 1-butene, respectively at a nominal temperature of 296 K and pressure of 760 Torr. Trace amounts of ethylene, propene, allene and 1-butene were measured individually as well as simultaneously in a multi-gas mixture containing gases at 1% of their TWA exposure limits. To the best of our knowledge, this work represents one of the first successful implementations of the pulsed CRDS technique to measure trace amounts of multiple gases in the mid-IR (900 – 1000 cm^{-1}) region. Furthermore, this is the first laser-based sensor capable of measuring trace amounts of propene, allene, and 1-butene.

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References


[34] Huang H and Lehmann K K 2010 Long-term stability in continuous wave cavity ringdown spectroscopy experiments Appl. Opt. 49 1378


[38] Es-Sebbar E, Jolly A, Benilan Y and Farooq A 2014 Quantitative mid-infrared spectra of allene and propyne from room to high temperatures J. Mol. Spectrosc. 305 10–6

[39] Ren W, Davidson D F and Hanson R K 2012 IR laser absorption diagnostic for C2H4 in shock tube kinetics
studies Int. J. Chem. Kinet. 44 423–32

[40] Arefae M, Es-sebbar E and Farooq A 2014 Absorption cross-section measurements of methane, ethane, ethylene and methanol at high temperatures J. Mol. Spectrosc. 303 8–14

[41] Airgas Inc. 2015 Material Safety Data Sheet Flammable Gas Mixture: 1-Butene / Cis-2-Butene / Ethane / Ethylene / Isobutane / Isobutylene (Isobutene) / Isopentane / Methane / N-Butane / N-Pentane / Propane / Propylene / Trans-2-Butene

[42] Airgas Inc. 2009 Material Safety Data Sheet Flammable Gas Mixture: 1,2-Propadiene 0.1-10% / Methane 90-99%