Quartz-enhanced photoacoustic sensor for ethylene detection implementing optimized custom tuning fork-based spectrophone

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Abstract: The design and realization of two highly sensitive and easily interchangeable spectrophones based on custom quartz tuning forks, with a rectangular (S1) or T-shaped (S2) prongs geometry, is reported. The two spectrophones have been implemented in a QEPAS sensor for ethylene detection, employing a DFB-QCL emitting at 10.337 μm with an optical power of 74.2 mW. A comparison between their performances showed a signal-to-noise ratio 3.4 times higher when implementing the S2 spectrophone. For the S2-based sensor, a linear dependence of the QEPAS signal on ethylene concentration was demonstrated in the 5 ppm –100 ppm range. For a 10 s lock-in integration time, an ethylene minimum detection limit of 10 ppb was calculated.

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1. Introduction

Trace gas detection finds applications in a wide variety of fields [1]: monitoring of ammonia, nitric oxide, volatile organic compounds (VOCs), ethylene and acetone in human breath to assist the early detection of diseases, such as asthma, cancer and diabetes [2]; the detection of greenhouse and polluting gases for environmental monitoring [3]; hydrocarbon detection for the prediction of production outputs and estimate reserves in the petrochemical industry [4]. Non-optical techniques for trace gas detection, such as gas chromatography, gas spectrometry and electrochemical sensing can be costly, bulky, slow or can suffer from hysteresis. Conversely, laser optical absorption based techniques are less costly, offering long-term stability, high compactness and sensitivity and can operate in real-time [5,6]. Laser absorption spectroscopy has been widely employed in the mid-infrared (mid-IR) spectral range, where the strongest absorption lines for several gas species are located. These absorption features are typically related to molecular ro-vibrational transitions. Photoacoustic spectroscopy (PAS) has been demonstrated as one of the most robust and sensitive trace gas optical detection techniques, capable of performing measurements at sub-parts-per-trillion (ppt) concentration levels [6–8]. PAS is based on the detection of sound waves generated by gas absorption of modulated optical radiation using a sensitive microphone and a resonant acoustic cell. In 2002, an alternative approach was proposed, exploiting a commercial high-quality factor quartz tuning fork (QTF) instead of microphone and resonant cell, named quartz-enhanced...
photoacoustic spectroscopy (QEPAS) [9]. The use of spectrophones, comprising quartz tuning forks acoustically coupled with micro-resonator (mR) tubes to enhance the signal-to-noise ratio, allowed a significant reduction of the detection module volume and immunity to environmental noise, while guaranteeing a high detection sensitivity and selectivity [10,11]. In 2015, a first generation of custom quartz tuning forks with rectangular prong geometry was designed, realized and tested, allowing a study on the influence of the dimensions of a quartz tuning fork on the main physical parameters controlling the QEPAS sensing performance, namely the resonance frequency, the quality factor and the electrical resistance [12]. Based on this study, in 2018 a second generation of custom quartz tuning forks was designed to investigate different prong geometries, such as the prong spacing up to 1.5 mm or prongs with a T-shaped longitudinal cross section or rectangular grooves carved on the prongs' surfaces [13].

The possibility to customize the quartz tuning forks for QEPAS sensing applications by varying the prong geometry and size made QEPAS extremely versatile. Depending on the specific target gas, sensing application or laser source, QEPAS sensors can be designed, implementing an appropriately selected quartz tuning fork. For example, the performance of a QEPAS sensor can be improved by employing a quartz tuning fork with a resonance frequency of few kHz, when detecting gas species characterized by slow non-radiative relaxation rates, such as CO, CO₂ or NO [14,15]. If implementing quartz tuning forks with both fundamental and first overtone flexural mode resonance frequencies lower than 40 kHz, two approaches can be employed: the double antinode excited QEPAS (DAE-QEPAS), where the two antinodes of the quartz tuning fork first overtone mode are simultaneously excited by one laser source [16] to improve the single-gas detection sensitivity, and the dual-gas QEPAS, where two different lasers excite independently both flexural modes of the quartz tuning fork to simultaneously detect two gas species [17]. Large prong spacing (≥800 μm) custom quartz tuning forks permit alignment tolerances to be relaxed and the development of QEPAS sensors employing no-commercial or no-conventional laser sources. These were the cases of: i) the extension of QEPAS to the THz spectral range, where the laser beam diameters are comparable with the prong spacing (300 μm) of the standard quartz tuning fork [18]; ii) the implementation of a high power (<1 W) fiber-amplified near-IR laser source [19]; iii) the exploitation of the tunability range provided by a monolithic array of 32 distributed feedback-QCLs for QEPAS sensing of broadband absorbers [20].

In this work, starting from the analysis of the geometrical and physical parameters of the available two generations of custom quartz tuning forks, two spectrophones with similar sizes were realized. In this way, we were able to design a QEPAS setup including a single acoustic detection module capable to accommodate two different spectrophones. Therefore, a quick switch between the two spectrophones was possible, requiring only a slight alignment after replacement. The performance of the two spectrophones were tested to detect ethylene (C₂H₄) in the mid-infrared spectral range. Ethylene detection is fundamental for its rising demand in the petrochemical industry as well as for air quality monitoring, fruit storage and transportation environment monitoring in food industry, and for inflammatory diseases diagnosis in medicine [21–25]. Depending on the application, the required ethylene detection sensitivity can vary from 1 ppm down to 10 ppb: climacteric fruits ripening and resulting early epinasty are triggered by concentrations in the 0.025-1 ppm range [26]; emissions from industrial facilities are regulated at less than 1 ppm [27]; a threshold of 10 ppb in air is considered a pollutant for plants [28] and traces of ethylene in the range of 10 ppb in breath are indicative of lipid peroxidation in lung epithelium [29]. In 2009, a near-infrared QEPAS-based sensor for ethylene detection was demonstrated, exhibiting a sensitivity of 0.5 ppm for 70 s integration time [30]. A mid-infrared QEPAS-based sensor was demonstrated in 2016, achieving a minimum detection limit (MDL) of 50 ppb in 70 s targeting an absorption line falling at 10.53 μm [31]. Recently, this MDL was lowered to 8 ppb at an integration time of 90 s, operating the sensor at pressures lower than 200 torr [32]. In this work, we demonstrate
that the accurate selection of the employed spectrophone with a specific quartz tuning fork geometry allows the realization of a QEPAS sensor with a minimum detection limit of lower than 10 ppb achievable in shorter integration times, which is suitable for all the above listed applications.

2. Spectrophones design and electrical characterization

The performance of two spectrophones were compared employing tuning forks from the two generations of custom quartz tuning forks realized so far. The two quartz tuning forks were selected in order to allow realizing two interchangeable spectrophones of the same size and provide a QEPAS signal as large as possible. Hence, both spectrophones can be accommodated and easily interchanged within the same acoustic detection module, consisting of a vacuum-tight gas cell equipped with two ZnSe windows antireflection (AR) coated in the 7-12 μm spectral range and gas-in and gas-out connectors. All 2nd generation custom quartz tuning forks have a total height of 14.1 mm, as reported in Table 1, with a prong length of 9 mm. Among them, QTF-S08-T, having T-shaped cross section-prongs, exhibited the largest QEPAS signal [13]. The QTF-S08-T acoustically coupled with two 12.4 mm-long tubes with internal diameter of 1.59 mm provided an increase of the signal-to-noise ratio (SNR) by a factor of 60 with respect to the bare tuning fork, representing a record for mid-IR QEPAS sensing [13]. Hence, this spectrophone was selected for this work and will be named hereafter as S2.

In Table 1 the total height, the resonance frequency and quality factor of the fundamental and first overtone modes of the 1st generation of custom quartz tuning forks described in [33,34] are summarized and compared with QTF-S08-T, as reported in [13]. The surrounding air pressure at which the resonance parameters were measured are also reported.

Table 1. Total height (H), fundamental and first overtone flexural mode resonance frequency (f₀, f₁), and quality factor (Q₀, Q₁) of second- and first-generation QTFs and operation surrounding air pressure, as reported in [13] and [33,34], respectively.
quartz tuning forks and mR tubes comprising the two spectrophones S₁ and S₂, as shown in Fig. 1(a), are summarized.

<table>
<thead>
<tr>
<th>Spectrophone</th>
<th>Lprong (mm)</th>
<th>Tprong (mm)</th>
<th>sprong (mm)</th>
<th>Ltube (mm)</th>
<th>IDtube (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S₁</td>
<td>11.0</td>
<td>0.5</td>
<td>0.6</td>
<td>4.0</td>
<td>0.84</td>
</tr>
<tr>
<td>S₂</td>
<td>9.4</td>
<td>2</td>
<td>0.8</td>
<td>12.4</td>
<td>1.59</td>
</tr>
</tbody>
</table>

A sketch of the two spectrophones is shown in Fig. 1(a), where the S₁ mR tubes are positioned 5.8 mm below the top of the prong, close to the position of the second antinode of the QTF#3 first overtone mode, spaced 50 μm from the quartz tuning fork surface, while the S₂ tubes are placed at a distance of 200 μm from the QTF-S08-T surface, 2 mm below the top of the prong, as already discussed in [33]. The proper selection of the two quartz tuning forks allowed the realization of two interchangeable spectrophones to be hosted in the same acoustic detection module, depicted in Fig. 1(b), allowing the laser beam to be focused between the two quartz tuning forks prongs and through the related mRs with slight vertical adjustments. A cylindrical cavity having a height of 24 mm and a diameter of 13 mm is drilled inside the acoustic detection module in order to host S₁ or S₂, alternately.

The electrical properties of the described spectrophones were studied by using an experimental setup depicted in Fig. 1(b).

The resonance properties of the two spectrophones were analysed by applying a sinusoidal voltage excitation with an amplitude of 170 mV. The quartz tuning fork current signal was transduced in a voltage signal by a custom-designed low-noise transimpedance amplifier employing a 10 MΩ feedback resistor and demodulated by a lock-in amplifier. The electrical responses of S₁ and S₂ at 120 Torr when the quartz tuning forks vibrate at the first overtone and fundamental flexural mode, respectively, are reported in Fig. 2.
For each spectrophone, the quartz tuning fork resonance frequency and quality factor were determined by fitting the acquired resonance spectra with Lorentzian curves. The Lorentzian fit of the S1 curve exhibited a maximum at 21466.12 Hz, with a full width high maximum of 1.02 Hz, leading to a Q-factor of 20880, whereas, for S2 a resonance frequency of 12462.02 Hz with a Q factor of 12530 was measured.

3. QEPAS measurements

The two spectrophones were implemented in the same QEPAS-based sensor, whose architecture is depicted in Fig. 3.

A distributed-feedback quantum cascade laser emitting at ~10.337 μm (Thorlabs, Inc., model QD10337HHLAS) was used as the laser source exciting the ethylene molecules within the acoustic detection module. With an injected current of 441.8 mA, the optical power measured with a pyroelectric power meter at the HHL output was 74.2 mW. The laser beam was focused into the acoustic detection module, fixed on a five-axis stage for alignment purposes, by using a 75 mm focal length ZnSe lens with a 7-12 μm AR coating. Such a focal length was chosen to achieve a tradeoff between the need for a small-diameter beam focused between the QTF#3 prongs of S1 (S_p = 0.6 mm) and a small numerical aperture of the same
beam passing through the 24.8 mm-long S\textsubscript{2} dual-tube micro-resonator. A \textasciitilde{}2 mm diameter pinhole was placed between the lens and the acoustic detection module to cut the beam tails that could hit the mR tubes and/or the quartz tuning fork prongs, avoiding in this way an increase of a non-zero background that would worsen the sensor ultimate detection sensitivity. The far field spatial intensity distribution of the laser beam in the focal plane of the focusing lens was acquired using a pyroelectric camera (pixel size 100\textmu m x 100\textmu m). The use of a pinhole results in a focused beam diameter of 0.29 mm, well below the prong spacing of the employed quartz tuning forks (0.6 mm for S\textsubscript{1} and 0.8 mm for S\textsubscript{2}). A \textasciitilde{}7\% optical power reduction was measured, with an optical power focused inside the acoustic detection module of P = 61.6 mW. QEPAS measurements were performed using the wavelength modulation and dual-frequency detection method: a sinusoidal dither matching half of the quartz tuning fork resonance frequency of the employed spectrophone was applied to the QCL current driver and the transduced quartz tuning fork signal was demodulated by the lock-in amplifier at the quartz tuning fork resonance frequency. The lock-in integration time was set at 100 ms. The demodulated signal was thus digitized and stored on a personal computer by means of a data acquisition board, with the card sampling time set at 3 times the lock-in integration time.

The first measurements were performed using spectrophone S\textsubscript{1}. With the best alignment conditions, 97\% of the laser power was measured to pass through the micro-resonator tubes and between the prongs. Preliminary measurements were conducted to identify the strongest absorption line falling within the tunability range of the QCL and the gas pressure and laser modulation amplitude providing the highest SNR. A wide-range spectral scan was performed while a certified concentration of 100 part-per-million (ppm) of C\textsubscript{2}H\textsubscript{4}:N\textsubscript{2} passed through the system with a controlled rate of 30 sccm. Furthermore, at a laser operating temperature of 15\degree C, the laser injection current was swept by adding a voltage ramp to the sinusoidal dither, with 2.1 V amplitude and 3 mHz frequency. The gas pressure and laser modulation amplitude voltage were varied in order to obtain the largest QEPAS signal, while maintaining the noise low. The optimal values resulted 120 Torr and 23 mV, respectively. In Fig. 4, a comparison between the absorption cross section simulated by using the HITRAN database [37] and the scan obtained by employing the spectrophone S\textsubscript{1} is shown.

![Fig. 4. 100 ppm C\textsubscript{2}H\textsubscript{4}:N\textsubscript{2} absorption spectrum simulated by using HITRAN database (a) and QEPAS spectral scan obtained at the same concentration over the QCL tunability range acquired by employing the spectrophone S\textsubscript{1} (b).](image-url)

The comparison between the two graphs in Fig. 4 shows a very good correspondence between the simulated data and the acquired QEPAS spectral scan. In the investigated spectral range, the line located at 966.38 cm\textsuperscript{-1} is the strongest C\textsubscript{2}H\textsubscript{4} absorption feature with a linestrength of 2.21\texttimes{}10\textsuperscript{-20} cm/mol. A narrower spectral scan around the largest intensity absorption peak was recorded by applying a ramp with an amplitude of 300 mV and a frequency of 10 mHz to the QCL current driver. The obtained QEPAS spectrum is shown in Fig. 5 (green line). The spectrophone S\textsubscript{1} was subsequently replaced with S\textsubscript{2}. After realigning
the system, by slightly adjusting the laser beam vertical position, a 97.9% power transmission through the acoustic detection module was measured, slightly higher than with S1, justified by the larger QTF-S08-T prong spacing (800 µm) and the tubes’ internal diameter. The spectral scan of the 966.38 cm⁻¹ absorption peak for 100 ppm of C₂H₄:N₂ acquired by employing S₂ is shown in Fig. 5 (orange line), obtained by using the same operating conditions as for S₁.

![Graph showing QEPAS signal of 100 ppm C₂H₄:N₂ acquired when S₁ (orange solid line) or S₂ (green solid line) are implemented in the system. Both spectra were acquired at 100 ms lock-in integration time.](image)

The ethylene peak signal, measured by implementing the spectrophone S₁ in the acoustic detection module, was 175.8 mV, with a 1σ noise of 0.175 mV, leading to a signal-to-noise ratio (SNR) of ~1000 and an SNR = 1 minimum detection limit of ~100 part-per-billion (ppb). When implementing the spectrophone S₂, a signal of 615.8 mV, with a 1σ noise of 0.180 mV, was obtained. The similar noise values measured with the two spectrophones, as well as the similar values of the optical power transmitted through the spectrophones, confirmed that the accurate selection of the implemented quartz tuning forks, together with the proper selection of the lens focal length, allows the assembly of two interchangeable spectrophones that can be set in the same acoustic detection module. Although showing a comparable noise, the spectrophone S₂ provides an SNR of 3420, ~3.4 times higher than the one obtained by employing the spectrophone S₁. Despite that S₁ exhibited a higher quality factor, lowering the resonance frequency from 21.4 kHz (S₁) down to 12.4 kHz in S₂ together with the record sound amplification obtained for S₂ [13] lead to an improvement of the sensor performance when implementing S₂. An ethylene MDL of ~30 ppb was achieved. Then, a S₂-based QEPAS sensor calibration was performed. Figure 6(a) shows the QEPAS spectral scans measured for 40 ppm, 30 ppm, 20 ppm, 10 ppm and 5 ppm of C₂H₄:N₂ obtained by diluting ethylene with pure N₂. The signal acquired in pure N₂ is also shown for comparison. The peak values were extracted from each spectral scan in Figs. 5 and 6(a) and plotted as a function of the ethylene concentration in Fig. 6(b).
Fig. 6. QEPAS spectral scans measured for (a) 40 ppm, 30 ppm, 20 ppm, 10 ppm and 5 ppm C2H4:N2 and pure N2 and peak values (b) measured for each ethylene concentration (blue squares) with the corresponding best linear fit (red line).

By linearly fitting the peak values as a function of the ethylene concentration, a calibration curve was obtained. The linear fit yields a slope of 6.12 mV/ppm, with an intercept of 4 mV, comparable with the 1.2 mV background noise level measured in pure N2. The calculated R-squared value is equal to 0.999.

The 1σ noise can be lowered by further averaging the signal over longer times [38]. An Allan-Werle deviation analysis was performed with the aim of estimating the 1σ noise (and thus the achievable minimum detection limit) as a function of the lock-in integration time. The obtained Allan-Werle deviation plot is reported in Fig. 7.

Fig. 7. Allan-Werle deviation plot of the QEPAS signal in ppb units as a function of the lock-in integration time. For a 10 s integration time, a minimum detection limit of ~10 ppb was achieved (red dashed line).

For a lock-in integration time of 10 seconds a minimum detection limit as low as 10 ppb was achieved. This detection sensitivity improved to 7 ppb by increasing the integration up to 30 second. For longer integration times, drift effects emerge, induced by the mechanical instability mainly caused by the vacuum pump, and the sensor performance deteriorates. A future improvement of the sensor long term stability can be obtained by employing low-vibration pump. However, the achieved sensitivity level using the S2-based QEPAS sensor fulfills the requirements for ethylene detection in the food industry, pollution monitoring and breath sensing.
4. Conclusions

In this work we designed and realized two interchangeable custom quartz tuning fork-based spectrophones, namely S\textsubscript{1} and S\textsubscript{2}, capable of high QEPAS sensing performance. The implemented quartz tuning forks were selected among the two different custom quartz tuning forks generations realized so far, in order to assemble two spectrophones having equivalent sizes. The performances of S\textsubscript{1} and S\textsubscript{2} were tested in a QEPAS-based sensor for ethylene detection, designed to easily allow switching of the two spectrophones. At 100 ppm C\textsubscript{2}H\textsubscript{4}:N\textsubscript{2} concentration and 100 ms integration time, S\textsubscript{2} provided a QEPAS signal ~3.4 times higher than that measured using S\textsubscript{1}. This result shows that an appropriate design of the quartz tuning fork as well as its acoustic coupling with resonator tubes can efficiently improve the ultimate performance of a QEPAS sensor. The performance of the ethylene S\textsubscript{2}-based QEPAS sensor was investigated in detail, by calibrating the sensing system and determining its ultimate detection sensitivity. For a lock-in integration time of 10 s, a minimum detection limit of 10 ppb was estimated, making the sensor suitable for the detection sensitivity required in breath sensing, air quality control and fruit transportation control.

Funding

Welch Foundation (C0568); National Natural Science Foundation of China (NSFC) (61622503, 61575113, 61805132); Ministry of Education of the People’s Republic of China, 111 project (D18001).

Acknowledgments

The authors from Dipartimento Interateneo di Fisica di Bari acknowledge financial support from THORLABS GmbH, within the joint-research laboratory PolySense.

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