Terahertz quartz enhanced photo-acoustic sensor

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A quartz enhanced photo-acoustic sensor employing a single-mode quantum cascade laser emitting at 3.93 Terahertz (THz) is reported. A custom tuning fork with a mm spatial separation between the prongs allows the focusing of the THz laser beam between them, while preventing the prongs illumination. A methanol transition with line-strength of 4.28 × 10−21 cm has been selected as target spectroscopic line. At a laser optical power of ~ 40 µW, we reach a sensitivity of 7 parts per million in 4s integration time, corresponding to a 1σ normalized noise-equivalent absorption of 2 × 10−10 cm−1 W/Hz1/2. © 2013 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4812438]

Recent technological innovation in photonics and nanotechnology is now enabling Terahertz (THz) research to be applied in an increasingly wide variety of applications, such as information and communications technology, medical sciences, global environmental monitoring, homeland security, quality and process controls. Most of the above applications involve the use of THz spectroscopic systems. Explosives, narcotics, and toxic gases (i.e., carbon monoxide, hydrogen cyanide, hydrogen chloride and methanol) have indeed distinct spectral “fingerprints” and strong absorption bands across the THz range.

To address the above application requirements, suitable sensing systems equipped with high power, widely tunable sources, and high speed, high sensitivity detectors have to be developed. To date, only a few systems have been successfully demonstrated to operate across the far-infrared. Photomixing-based sensors in the 0.5–1.5 THz range have been recently proposed1 for detection of HCN, CO, H2CO, narcotics, and toxic gases (i.e., carbon monoxide, hydrogen cyanide, hydrogen chloride and methanol) have indeed distinct spectral “fingerprints” and strong absorption bands across the THz range.

The described systems suffer from the low power levels of the employed THz sources and/or the low sensitivity of the detection units. THz quantum cascade lasers (QCLs) can offer a significant improvement in terms of compactness and sensitivity levels, stemming from the single-mode high spectral purity emission,4 the relatively broad tunability range (up to 10% of the central frequency),5 the high continuous wave (CW) output power levels (up to 138 mW),6 and the relatively good compactness provided by Stirling cryocooler systems. THz QCLs have indeed recently demonstrated interesting performance in high resolution molecular spectrometries, in both direct absorption7–10 or wavelength modulation spectroscopy,11 showing a minimum detectable absorption xmin in the 10−6 cm−1 Hz−1/2 range, mostly thanks to the employed sensitive cryogenic detectors. Improving such detection sensitivities requires either switching to novel low noise equivalent power (NEP) room temperature nanodetectors,12,13 or moving to alternative high performance spectroscopic techniques.

Quartz-enhanced photo-acoustic spectroscopy (QEPAS) showed high sensitivity, fast time-response and high compactness in the near-IR and mid-IR spectral ranges. The distinguishing feature of QEPAS is the use of a low loss quartz tuning fork (QTF) for detection of the optically generated sound.14 Very efficient mid-IR QCL-based QEPAS sensors have been recently demonstrated for trace detection of several chemical gas species,4–17 with a record normalized noise-equivalent absorption (NNEA) sensitivity of 2.7 × 10−10 W cm−1 Hz−1/2 (σ) for SF6,18 corresponding to an xmin of 1.5 × 10−8 cm−1 Hz−1/2 and a detection limit of 50 parts per trillion (ppt) in 1 s. One of the main advantages of the photoacoustic spectroscopy techniques is that no optical detection is required.19 Thus, the extension of the QEPAS technique in the THz range would allow to avoid the use of low-noise but expensive, bulky, and cryogenic bolometers.

Standard QTFs have resonance frequencies of ~32.7 kHz and are characterized by a very small sensitive volume between its prongs (~0.3 × 0.3 × 3 mm3). In QEPAS experiments, it is critical to avoid laser illumination of the QTF, since the radiation blocked by the QTF prongs results in an undesirable non-zero background. This background can be several times larger than the thermal noise level of QEPAS and carries a shifting fringe-like interference pattern, which strongly limit the sensor detection sensitivity.18,20 The reduced space (300 µm) between the QTF prongs, comparable with the wavelength of THz sources, so far has represented the main limitation preventing the use of QEPAS system in THz range. Larger size QTFs are, therefore, mandatory to operate in the THz range.

In this paper, we report on the development of a THz QEPAS sensor employing a 3.93 THz QCL and custom-made...
QTF. Standard photolithographic techniques were used to etch the custom QTF, starting from z-cut quartz wafer. Chromium/gold contacts were deposited on both sides of the QTF. The overall QTF dimension was 3.3 cm × 0.4 cm × 0.8 cm; each prong was 2.0 cm long and 2.5 mm wide. The gap between the prongs was ~1 mm. The QTF first flexion resonance falls at $f_0 \sim 4246$ Hz, in agreement with previous studies reported for a similar QTF. At atmospheric pressure we measured a Q factor of 9930. For our QEPAS THz sensor demonstration we selected methanol as target gas molecule. Methanol is widely used as a solvent, detergent, or even denaturant additive for industrial ethanol, and its ingestion can be fatal due to its toxication.

A scheme of the employed experimental system is shown in Fig. 1. A single-mode 250 μm wide, 1.5 mm long bound-to-continuum Fabry-Perot QCL fabricated in a single plasmon configuration and emitting at 3.93 THz (76.3 μm) (Ref. 22) was mounted on the cold finger of a continuous-flow cryostat configuration and emitting at 3.93 THz (76.3 μm). The THz beam was transmitted through the prongs without hitting the QTF. Standard photolithographic techniques were used to etch the custom QTF, starting from z-cut quartz wafer. Chromium/gold contacts were deposited on both sides of the QTF. The gap between the prongs was ~1 mm. The QTF first flexion resonance falls at $f_0 \sim 4246$ Hz, in agreement with previous studies reported for a similar QTF. At atmospheric pressure we measured a Q factor of 9930. For our QEPAS THz sensor demonstration we selected methanol as target gas molecule. Methanol is widely used as a solvent, detergent, or even denaturant additive for industrial ethanol, and its ingestion can be fatal due to its toxication.

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To test our sensor, we selected methanol. In the laser emission range methanol exhibits a rotational absorption line falling at 131.054 cm$^{-1}$ ($v = 1, K = 6, J = 11 \leftarrow (1, 5, 10)$) with line-strength $S = 4.28 \times 10^{-21}$ cm/molecule, about two orders of magnitude stronger than that of the nearby methanol absorption lines. Thus, we performed direct absorption spectroscopic measurements (using a 14 cm-long cell filled with pure methanol at 2 Torr) and, as expected, we observed a main absorption line. The line-strength extracted from absorption measurements was in good agreement with the expected one. So in our QEPAS experiments we exploit the absorption measurement to fine tune the laser frequency on the selected methanol absorption line at 131.054 cm$^{-1}$.

Gas mixtures with different methanol concentrations have been obtained by diluting methanol vapors, collected from a reservoir held at the vapor pressure (~120 Torr at 300 K), with pressurized N$_2$. For measurements at low concentrations, we used a certified 100 ppm methanol/N$_2$ gas mixture. Preliminary measurements were performed to determine the best operating conditions in terms of QEPAS signal-to-noise, as a function of gas pressure, laser current modulation depth and resonance frequency. Note that the possibility to use a lower resonance frequency with respect to standard QTF, partially relaxes the gas excess energy relaxation time requirements and allows to work at low pressures, taking advantages of the corresponding larger Q factors. The best operating conditions have been observed at 10 Torr pressure and 600 mV peak-to-peak modulation voltage. Under these conditions, the physical parameters of the QTF, using N$_2$ as gas carrier, were $Q = 76300$, $f_0 = 4246.73$ Hz, and $R = 6.5$ MΩ. From these data, we extracted a QTF thermal noise of 0.12 μV, about one order of magnitude smaller than that typically observed for standard QTFs.

A representative spectral scan of 0.75% methanol in N$_2$ gas mixture is shown in Fig. 2(a). With a pressure broadening coefficient of 10 MHz/Torr (7.4 MHz/millibar) (Ref. 26) the expected line-width is ~100 MHz (half width at half maximum, HWHM), with a negligible Doppler contribution (4.5 MHz HWHM). We used this estimate to convert the horizontal scale from time to frequency span (MHz). Note that the laser linewidth is much lower than the spectral line width of the methanol absorption feature. On the same energy...
span, in Fig. 2(b) are reported the line-strengths of the main methanol transitions (vertical bars), as tabulated in the Jet Propulsion Laboratory (JPL) database. The noise fluctuations are ~±25 μV. (b) Line strengths of the main methanol transitions (vertical bars), as reported in the JPL database, falling in the energy range corresponding to the frequency span of Fig. 2(a).

Stepwise concentration measurements were performed to verify the linearity of the QEPAS signal as a function of the methanol concentration. The system was operated in the locked mode, i.e., with the QCL frequency set to the center of the absorption line. Under this condition an optical laser power of 40 μW is focused on the QTF. In Fig. 3, the mean value of the QEPAS peak signal is plotted for different methanol concentrations from 6.5% down to 0.01% (100 ppm). The associated error bars take into account both the QEPAS signal fluctuations (standard deviation) and the uncertainty on the methanol concentration (uncertainty on the reading of the pressure gauge of 0.2 Torr). The experimental data show the expected linear dependence of the QEPAS signal from the methanol concentration.

In order to determine the best achievable sensitivity of the QEPAS sensor we performed an Allan variance analysis measuring and averaging the QEPAS signal at zero methanol concentration (pure N₂). The obtained Allan deviation in ppm is shown in Fig. 4. For a 4 sec averaging time (i.e., 0.04 Hz bandwidth), we achieve a detection sensitivity of 7 ppm, corresponding to a minimum absorption coefficient \( \alpha_{\text{min}} = 9.5 \times 10^{-7} \text{ cm}^{-1} \) (laser power of \( \sim 40 \mu \text{W} \)). The calculated NNEA is \( 2 \times 10^{-10} \text{ cm}^{-1} \text{W/Hz}^{1/2} \) (1σ), comparable with the best result obtained in the mid-IR and at least one order of magnitude better than those obtainable with room-temperature pyroelectric detectors and in strong competition with the sensitivities achieved with the most sensitive cryogenic bolometers.

In conclusion, we extend the possibility to employ the QEPAS technique in the THz spectral range, using a THz QCL source and a custom QTF. The simple apparatus architecture and the independence of the detection unit from the laser frequency make the proposed sensor highly versatile in the spectral domain actually accessible to THz QCLs. In addition, the possibility to employ high power CW (\( \sim 100 \text{ mW} \)) THz QCLs for the detection of strong THz absorbing molecules like OH or HF, having line-strengths of the order of \( 10^{-18} \text{ cm} \), would allow QEPAS detection sensitivity in the few parts per trillion concentration range. Furthermore, the possibility to implement compact QEPAS.
THz systems, with no optical detector needed, based on portable closed-cycle Stirling cryo-cooler opens the way to the use of THz sensor systems for in-situ security and environmental monitoring.

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