Dual-Gas Quartz-Enhanced Photoacoustic Sensor for Simultaneous Detection of Methane/Nitrous Oxide and Water Vapor

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ABSTRACT: The development of a dual-gas quartz-enhanced photoacoustic (QEPAS) sensor capable of simultaneous detection of water vapor and alternatively methane or nitrous oxide is reported. A diode laser and a quantum cascade laser (QCL) excited independently and simultaneously both the fundamental and the first overtone flexural mode of the quartz tuning fork (QTF), respectively. The diode laser targeted a water absorption line located at 7181.16 cm⁻¹ (1.392 μm), while the QCL emission wavelength is centered at 7.71 μm and was tuned to target two strong absorption lines of methane and nitrous oxide, located at 1297.47 and 1297.05 cm⁻¹, respectively. Two sets of microresonator tubes were positioned, respectively, at the antinode points of the fundamental and the first overtone flexural modes of the QTF to enhance the QEPAS signal-to-noise ratio. Detection limits of 18 ppb for methane, 5 ppb for nitrous oxide and 20 ppm for water vapor have been achieved at a lock-in integration time of 100 ms.

Gas spectroscopic techniques based on optical absorption have been widely demonstrated as fast, highly sensitive, and selective tools to detect and monitor one or more gas species in a mixture.¹,² Once a low-noise, narrow-bandwidth laser source and a strong, interference-free absorption line are selected, several sensing approaches can be employed.¹ Multipass cells allow an increase of the optical path length up to 1 order of magnitude.³ High-finesse optical cavities exploit optical field enhancement when operating in a steady-state regime, allowing an increase of the intracavity optical power by more than 2 orders of magnitude.⁴ Both approaches require fast, low-noise, ultrasensitive optical detectors to acquire the weak optical field exiting the multipass cell or the cavity. Moreover, these techniques suffer from mechanical instability and complex optical alignment procedures and are strongly wavelength-dependent, hence limiting their optical coupling with multiple sources. Photoacoustic (PAS) spectroscopy is also based on an optical absorption process but differs in the way the absorption signal is detected. In PAS sensors, the gas is confined inside an acoustic cell and the exciting laser light is modulated at one of the cell acoustic resonance frequencies. The modulated absorption and non-radiative relaxation occurring in the gas generate pressure waves, that is, sound. These waves are then detected by using low-noise, highly sensitive microphones.⁵ Quartz-enhanced photoacoustic spectroscopy (QEPAS) exploits an alternative sound detection approach, by substituting the resonant cell and microphone with a quartz tuning fork (QTF), acting as a sharply resonant acoustic transducer.⁶ The QTF can be coupled with a pair of microresonator tubes in order to confine and amplify the sound waves. In on-beam QEPAS, microresonator tubes are aligned on both sides of the QTF, close to one of the antinode points of a QTF in-plane flexural resonance mode.⁷ The laser beam is focused through the tubes and between the prongs and is modulated at the resonance frequency (or one of its subharmonics) of the selected QTF vibrational mode. The pressure wave deflects the prongs and an electrical signal proportional to the stress field is generated exploiting the quartz piezoelectricity. QEPAS sensors can be very compact, do not require an optical detector and are wavelength independent. Indeed, the QTF response depends only on the sound waves intensity and not on the wavelength of the exciting laser used, which makes it suitable for coupling multiple sources emitting at different
wavelengths for multi-gas detection. Many gas-sensing applications require simultaneous detection and quantification of different components in a gas mixture, such as detection of isotopes concentration ratio, environmental monitoring of greenhouse gases, monitoring of different biomarkers in human breath and study of molecular relaxation dynamics. Different approaches have been explored to achieve multigas detection. One possibility is to exploit a single laser source with a wavelength tuning range covering absorption lines of the different gas targets. Optically coupling an external cavity with Fabry–Perot quantum cascade lasers (QCLs), a tunability range of more than 100 cm⁻¹ can be achieved. The QCLs emission wavelength can be finely tuned by a rotating grating to target different gas species. However, external cavity QCLs are bulky, suffering from power fluctuations and the emission wavelength stability is limited by mechanical instability of the grating. Another approach to widely tune a QCL is to integrate a heater to the laser chip. A QCL with an integrated-heater operating in the 1276–1284 cm⁻¹ region was implemented to detect methane and acetylene for combustion applications. Although the high time resolution achieved (20–40 μs), this approach can be applied only to the case in which the absorption lines of the different gases are separated by few cm⁻¹. An alternative approach employs multiple laser sources, one for each gas to be detected, switching between them and activating one laser at one time. A photoacoustic sensor was developed with five lasers sequentially operated along a single optical interaction path using a scanning galvanometer as an optical multiplexer with the capability of switching between two lasers in less than 1 s. QEPAS is perfectly suitable for multigas sensing applications requiring either a wide tuning range of the exciting laser source or the employment of multiple sources due to the wavelength independence responsivity of the QTF. In ref 14, a single intercascade laser was tuned to target absorption features of methane, ethane and propane, whereas in ref 15, the emission wavelength of a distributed feedback-QCL (DFB-QCL) was tuned to detect carbon monoxide and nitrous oxide. In ref 16, two lasers matching the absorption lines of methane and ammonia, respectively, were alternatively focused through the QEPAS detection module.

In all the cases reviewed so far there is a finite time interval between the detection of several gas species, which corresponds to the time needed to tune the laser emission wavelength between different absorption features and/or to switch between multiple laser sources. Hence, these approaches can be referred to as quasi-simultaneous dual-gas detection. Differently, in ref 17, a photoacoustic sensor was developed with three lasers in combination with three independent resonators along three optical paths. Sub part-per-million detection limits for CH₄, H₂O, and HCl were demonstrated, but the cross-talking among the resonators did not allow a real-simultaneous detection of the gases. A multigas QEPAS sensor based on three bare QTFs with different response frequencies for trace gas detection was reported in ref. [18]. Three near-infrared laser diodes were used to simultaneously monitor water vapor, methane and acetylene in the parts per million range.

A straightforward way to guarantee simultaneous detection is developing a wavelength-modulation division multiplexing (WMDM) scheme. Different laser sources are independently modulated, one for each gas species to be detected and then combined using a single detector. The recorded signal is then demodulated at the different modulation frequencies. Optical sensors with WMDM configuration based on multipass absorption have been reported in the literature. The development of custom QTFs opened the way to exploit QEPAS technique in WMDM configuration. If the QTF frequency of the in-plane flexural mode is decreased to a few kHz, the frequency of the in-plane first overtone mode, which is ~6.2 times higher than the fundamental one, becomes accessible for QEPAS operation. A proof-of-concept of simultaneous dual-gas QEPAS detection exploiting a custom QTF was reported in ref 23, where two DFB diode lasers operating in the near-IR region were used as the excitation sources for the fundamental and the first overtone mode of a bare QTF and simultaneous detection of water vapor and acetylene was demonstrated.

In this work, a WMDM configuration dual-gas QEPAS sensor operating in two different spectral regions, near-IR and mid-IR, is reported. A DFB-QCL emitting at 7.7 μm allowed methane (CH₄) or nitrous oxide (N₂O) detection, while a diode laser emitting at 1.392 μm was exploited for simultaneous water vapor monitoring. In contrast to the approach proposed in ref 18, which precludes the use of a microresonator system, the dual gas QEPAS sensor presented here employed a custom QTF acoustically coupled with two microresonator dual-tube systems to enhance both the fundamental and the first overtone QEPAS signal-to-noise ratio, thereby lowering the detection limits down to concentrations in the part per billion range.

### EXPERIMENTAL SECTION

**Dual-gas spectrophone design.** Simultaneous dual-gas detection with QEPAS technique can be obtained only by exciting at the same time two different flexural vibrational modes of a QTF, that is, the fundamental and the first overtone mode. The related resonance frequencies can be chosen at the design phase by properly selecting the prongs length and thickness and making both modes suitable for QEPAS operation. In ref 24, three different QTF designs were proposed and tested, having both the fundamental and the first overtone mode frequencies lower than 40 kHz. Among them, we selected the QTF labeled as QTF#1 since it showed the highest quality factor value and consequently the best QEPAS performance, for both fundamental and first overtone modes. QTF#1 has two rectangular prongs with length of 17 mm, thickness of 1 mm and a quartz crystal width of 0.25 mm. These QTF geometrical parameters leads to a fundamental frequency as low as 2.88 kHz, while the first overtone mode resonates at ~17.78 kHz. The spacing between the two prongs is 0.7 mm. The fundamental mode has an antinode point on the top of the QTF while the first overtone mode has two antinode points, one coincident to that of the fundamental mode and the other one close to the middle of the prong (at 9 mm from the QTF#1 top). In this work, two pairs of microresonator tubes were employed to simultaneously enhance the fundamental and the overtone mode vibrations. One pair was located at 2 mm from the top of the prongs for the fundamental mode and the other pair at 9.5 mm from the top of the prongs for the first overtone mode, as discussed in ref 24. The length and internal diameter of tubes must be accurately selected to optimize the QEPAS performance of the spectrophone. The length of the tubes is correlated to the sound wavelength by the relation $\lambda = \frac{v}{f}$, where $v$ is the sound speed and $f$ is the QTF resonance frequency. The optimal
tube length was experimentally found to be between $\lambda/4$ and $\lambda/2$. For the first overtone mode, the corresponding length range is $4.8 \text{ mm} < l < 9.6 \text{ mm}$, and $l = 8.5 \text{ mm}$ was selected. For the fundamental mode, the optimal length falls between 30 mm and 60 mm. However, such a length would require a bulky acoustic detection module to accommodate the spectrophone and thus complicate the optical alignment operations. For these reasons, tubes with a length of 9.5 mm have been employed, which are 1 mm longer than the tubes used for the first overtone mode in order to maintain the spectrophone size as compact as possible. A low QEPAS signal enhancement is thus expected for the fundamental mode. Therefore, this mode was chosen to detect the gas target with the highest concentration. For both antinodes, tubes having an ID = 1.36 mm were mounted, which was proved to be the optimum internal diameter in terms of the signal-to-noise enhancement factor with respect to the bare QTF. The gap between tubes and the QTF surface was set to 150 $\mu$m. A schematic of the spectrophone is shown in Figure 1.

The vibrational properties of the QTF coupled with the microresonator tubes, namely the resonance frequency and the quality factor of both flexural modes were measured by electrically exciting the QTF. A sinusoidal voltage signal excites the QTF resulting in a piezoelectric charge displacement on its prongs via an inverse piezoelectric effect. Charges are collected by electrodes deposited on the QTF surface and the QTF current signal is converted into an output voltage by means of a trans-impedance preamplifier. Finally, the output voltage is demodulated by a lock-in amplifier at the same frequency of the excitation signal. The resonance curves for the fundamental and first overtone mode of a bare QTF were fitted using a Lorentzian function to extract the resonance frequencies and the full width at half-maximum (fwhm) values. The quality factor is calculated as $Q = f/\text{fwhm}$. In Figure 2, the QTF resonance curves of the fundamental and first overtone modes measured at 200 Torr are shown. The fundamental mode exhibits a resonance frequency of $f_0 = 2871.05 \text{ Hz}$ and a quality factor of 6680, while the overtone resonance mode occurs at $f_1 = 17747.70 \text{ Hz}$ with a quality factor of 17070.

The first overtone mode exhibits a higher quality factor with respect to the fundamental one as already demonstrated in ref 24. The fundamental mode will be used to monitor the water vapor concentration in air (whose concentration is expected to be in the percent-range), while the first overtone mode will be exploited to detect CH$_4$ and N$_2$O trace gas concentration. The distance between the fundamental antinode point and the second antinode of the first overtone mode is $\sim 8 \text{ mm}$, allowing an easy alignment of the two laser beams through the microresonator tubes.

**Architecture of dual-gas QEPAS sensor.** The dual-gas spectrophone is enclosed in a gas cell, forming the acoustic detection module (ADM). The gas cell is equipped with a ZnSe window antireflection (AR)-coated in the range of 7−12 $\mu$m on the front side and a N-BK7 window AR-coated in the 1−1.7 $\mu$m range on the back side. Two connectors are used as a gas inlet and a gas outlet. The ADM was implemented in the QEPAS sensor setup depicted in Figure 3. Two lasers were used as excitation sources. A DFB-QCL emitting at 7.73 $\mu$m was used to alternately target methane and nitrous oxide absorption peaks exciting the first overtone QTF mode. The beam was then focused through the microresonator tubes located at the lower antinode point of the overtone mode by means of a 7−12 $\mu$m AR-coated Zn−Se lens having a focal length of 75 mm. A pinhole was used as a spatial filter to cut the laser beam tails and prevent that the incident light hits the QTF prongs or the microresonator tubes and generates a background photothermal noise. A diode laser emitting at 1.392 $\mu$m was used to target water vapor. A 1.05−1.7 $\mu$m AR-coated N-BK7 lens having a focal length of 75 mm was used to focus the diode laser beam between the QTF prongs and the tubes located at to the antinode point of the fundamental mode. The beam waists of the QCL and the diode laser measured on the QTF plane have a diameter of 0.28 mm and 0.24 mm, respectively, well below the prong spacing of the QTF (0.7 mm). The portion of optical power transmitted through the microresonator tubes and between the prongs was 97.5% and 99.2% for the QCL and the diode laser, respectively. QEPAS signals were detected using the wavelength modulation technique with 2f-detection and the two lasers were independently modulated.

A sinusoidal dither was applied to the current driving the diode laser at the half of the fundamental mode frequency $f_0/2 = 1435.52 \text{ Hz}$, while the current injected into the QCL was modulated at the half of the first overtone mode frequency, $f_1/2 = 8873.85 \text{ Hz}$. The QTF signal was converted into a voltage signal using a trans-impedance preamplifier with a gain factor of 30; and then was simultaneously demodulated at the fundamental and first overtone frequencies using two lock-in amplifiers. The integration time was set at 100 ms for both lock-in amplifiers. The demodulated signals were recorded and

![Figure 1](image1.png)

**Figure 1.** Schematic of the spectrophone implemented. The QTF is coupled with two pairs of microresonator tubes aligned perpendicularly to the QTF. The first pair of tubes (mR$_1$), each one 9.5 mm long, is located at the fundamental antinode point near the QTF top, the second pair (mR$_2$), each tube 8.5 mm long, is positioned 9.5 mm below the prongs top, at the second antinode point position of the QTF first overtone mode; the preamplifier (PA) board is shown on the bottom.

![Figure 2](image2.png)

**Figure 2.** Spectrophone resonance curves (blue squares) and related Lorentzian fits (red solid curves) of the fundamental and the first overtone modes at 200 Torr in air.
analyzed on a personal computer by using a multichannel data acquisition card from National Instruments. A gas line system was assembled to manage flow rates, pressures and compositions of the gas mixtures to be analyzed. One Y-ended gas line branch was connected to the gas cylinders containing certified concentrations of gas targets: 10 000 ppm of CH₄ in N₂ and 10 000 ppm of N₂O in N₂. The other gas line branch was connected to a valve allowing laboratory air to enter the line. Since the water vapor concentration in the laboratory environment can vary, a Nafton humidifier was inserted just before the ADM (not shown in Figure 3) to keep the water concentration in the ADM fixed to 1.6%. The pressure inside the ADM was fixed at 200 Torr and the flow set at 25 sccm by means of a pressure controller, a valve system and a vacuum diaphragm pump.

**RESULTS AND DISCUSSION**

**Simultaneous QEPAS Detection of Methane and Water.** As a first step, the QEPAS sensor was tested for detection of methane and water vapor in a gas mixture, independently. The QCL and diode laser were alternatively switched on to excite the targeted CH₄ and H₂O absorption lines, respectively. A preliminary study showed that the CH₄ QEPAS signal is maximized at a gas pressure of 200 Torr. The HITRAN database was used to simulate the absorption cross-section of pure methane at 200 Torr in the QCL. (a) Hitran simulation of the pure methane absorption cross-section at 200 Torr; (b) QEPAS spectral scan of 1000 ppm of CH₄ at 200 Torr measured by demodulating the QTF signal at the overtone mode frequency; (c) Hitran simulation of the pure water vapor absorption cross-section at 200 Torr; (d) QEPAS spectral scan of 1.6% of H₂O at 200 Torr measured by demodulating the QTF signal at the fundamental mode frequency.
The simulation result is depicted in Figure 4(a). The strongest absorption peak of methane falls at 1297.47 cm\(^{-1}\) with a line strength of \(3.9 \times 10^{-20}\) cm/molecule. No visible water lines fall in the investigated QCL emission spectral range. Figure 4(b) shows the corresponding QEPAS spectral scan acquired for a mixture of 1000 ppm of CH\(_4\) with 1.6% of water vapor at 200 Torr. The QCL temperature was set to 12.5 °C and the injected current was scanned in the range 207–257 mA applying a 2 mHz ramp to the current driver. The QEPAS signal was demodulated at \(f_1\) for methane detection. The amplitude of the QEPAS signal related to the selected line (targeted at a laser current of \(I_{\text{QCL}} = 243.7\) mA) was optimized by varying the laser modulation depth. The highest CH\(_4\) signal of 146 mV was observed at a modulation depth of 35 mV. The other features composing the QEPAS spectrum of Figure 4(b), like the ones peaked at 224.8 mA and 231.8 mA, correspond to methane absorption lines with weaker line strengths as confirmed by a HITRAN database simulation. The QCL was then turned off and the diode laser was used to excite the water vapor absorption line. Figure 4(c) shows the simulated absorption cross-section of pure water at 200 Torr in the 7180–7185 cm\(^{-1}\) range, obtained using the HITRAN database. Within the current dynamic range of the diode, the strongest absorption peak of H\(_2\)O falls at 7181.16 cm\(^{-1}\) with a line strength of \(1.4 \times 10^{-20}\) cm/molecule. Figure 4(d) shows the QEPAS spectral scan for a mixture of 1000 ppm of CH\(_4\) with 1.6% of water vapor, at 200 Torr.

These measurements were obtained by setting the temperature of the laser diode to 25 °C and scanning the laser current in the range 50–115 mA, by applying a 2 mHz ramp to the current driver. The diode modulation amplitude was set to 12 mV and the QEPAS signal was demodulated at \(f_0\). The selected H\(_2\)O line shows a peak value of 0.68 mV. The QEPAS absorption features at 80.7 mA, 89.5 mA and 108.6 mA correspond to water vapor absorption lines with weaker line strengths. The optical power focused inside the ADM was 128.0 mW at the QCL current set to target the selected CH\(_4\) absorption line and 6.2 mW for the diode laser current set to hit the strongest H\(_2\)O line absorption line.

It is mandatory to verify that the QEPAS signal demodulated at \(f_0\) is not influenced by the first overtone mode vibration and that the QEPAS signal demodulated at \(f_1\) is not affected by the fundamental mode vibration. Therefore, a preliminary test was conducted to investigate the QTF response while simultaneously exciting the fundamental and first overtone vibrational modes. Figure 5(a) shows the QTF signal demodulated at \(f_0\) when only the first overtone mode is excited and the QTF signal demodulated at \(f_1\) when both the fundamental and first overtone modes are excited (blue circles). The strongest absorption peak of methane falls at 1297.47 cm\(^{-1}\) with a line strength of \(3.9 \times 10^{-20}\) cm/molecule. No visible water lines fall in the investigated QCL emission spectral range. Figure 5(b) shows the corresponding QEPAS spectral scan acquired for a mixture of 1000 ppm of CH\(_4\) with 1.6% of water vapor at 200 Torr. The QCL temperature was set to 12.5 °C and the injected current was scanned in the range 207–257 mA applying a 2 mHz ramp to the current driver. The QEPAS signal was demodulated at \(f_1\) for methane detection. The amplitude of the QEPAS signal related to the selected line (targeted at a laser current of \(I_{\text{QCL}} = 243.7\) mA) was optimized by varying the laser modulation depth. The highest CH\(_4\) signal of 146 mV was observed at a modulation depth of 35 mV. The other features composing the QEPAS spectrum of Figure 4(b), like the ones peaked at 224.8 mA and 231.8 mA, correspond to methane absorption lines with weaker line strengths as confirmed by a HITRAN database simulation. The QCL was then turned off and the diode laser was used to excite the water vapor absorption line. Figure 4(c) shows the simulated absorption cross-section of pure water at 200 Torr in the 7180–7185 cm\(^{-1}\) range, obtained using the HITRAN database. Within the current dynamic range of the diode, the strongest absorption peak of H\(_2\)O falls at 7181.16 cm\(^{-1}\) with a line strength of \(1.4 \times 10^{-20}\) cm/molecule. Figure 4(d) shows the QEPAS spectral scan for a mixture of 1000 ppm of CH\(_4\) with 1.6% of water vapor, at 200 Torr.
vibrational modes are simultaneously excited by the two laser sources (blue circles).

The QCL and the diode laser currents were tuned to detect the selected CH₄ and H₂O absorption lines falling at 1297.47 and 7181.16 cm⁻¹, respectively, in a mixture containing 600 ppm of CH₄ and 1.6% of water vapor. Similarly, Figure 5(b) shows the QTF signal demodulated at f₀ when only the fundamental vibrational mode is excited using the diode laser (green squares) and when both the lasers operate (blue circles). The perfect overlap between the curves in both figures indicates that there is no cross-talking between the QTF fundamental mode and first overtone mode operations, even in the presence of the two microresonator tubes placed at the antinode points. This allows the excitation of the two vibrational modes simultaneously, acquisition of the QTF signal and independent demodulation at the two resonance frequencies to identify the contribution due to each gas species and retrieve both gas concentrations. The calibration of the sensor for methane detection was performed acquiring spectral scans of mixtures of CH₄ in laboratory air with different gas target concentrations, ranging from 50 to 1000 ppm. In all generated mixtures, the water vapor concentration was fixed by using the Nafion humidifier. The sensor was used in the dual-gas detection mode in order to measure the CH₄ concentration and simultaneously monitor the water vapor level. Figure 6(a) shows the QEPAS signal acquired for methane at seven different concentrations and the correlated water QEPAS signal (see Figure 6(b)).

The CH₄ peak values were extracted for each spectral scan and plotted as a function of the concentration in Figure 6(c). A linear fit of the experimental data points was performed to obtain the sensor calibration curve for methane. The squared-R value of 0.999 confirms the linearity of the QEPAS signal with the methane concentration, with a slope of 0.146 mV/ppm and an intercept of 0.85 mV. For each dilution, the water vapor level has been monitored by scanning the H₂O absorption line with the diode laser. The H₂O QEPAS peak signals extracted from Figure 6(b) show a peak value fluctuation <0.25% value, demonstrating the capability of the Nafion humidifier to efficiently fix the water concentration in the gas line. For CH₄, a 1σ-noise level of 2.7 μV was estimated at 100 ms lock-in integration time corresponding to a minimum detection limit (MDL) of 18 part-per-billion (ppb), while for H₂O with a 100 ms integration time we obtained a 1σ noise of 1.0 μV corresponding to an MDL of 20 ppm.

**Simultaneous QEPAS Detection of Nitrous Oxide and Water.** The realized QEPAS sensor allows also the simultaneous detection of water vapor and nitrous oxide, since the QCL emission spectral range covers absorption lines of N₂O. With the QCL temperature set to 12.5 °C and the QCL current scanned in the range of 152–287 mA, several N₂O absorption lines can be targeted. The highest intensity line falls at 1297.05 cm⁻¹ with a line strength of 1.7×10⁻¹⁹ cm/molecule. The QCL modulation amplitude and the gas pressure maximizing the N₂O QEPAS signal were found to be 130 mV and 200 Torr, respectively. For water vapor detection, the diode laser working conditions remain the same as before. Figure 7(a) shows the QTF signal demodulated at f₁ for a mixture of 400 ppm of N₂O and 1.6% H₂O in air. The selected N₂O absorption line is excited at a QCL current of I_QCL = 271.1 mA and shows a peak value of V = 227 mV. The sensor was calibrated for N₂O detection in the range of 300–800 ppm starting from a 10 000-ppm certified concentration of N₂O:

![Image](https://example.com/figure7.png)

**Figure 7.** (a) Nitrous oxide spectral scan detected at f₁ exciting the first overtone vibrational mode of the QTF with the quantum cascade laser for a mixture of 400 ppm of nitrous oxide and 1.6% of water in air at 200 Torr. (b) Peak values for each N₂O concentration (square symbols) and the corresponding best linear fit (red line).

The QEPAS peak value scales linearly with N₂O concentration in the selected range with a slope of 0.60 mV/ppm, an intercept of 0.51 mV and a squared-R value of 0.997, as shown in Figure 7(b). A 1σ-noise level of 2.7 μV at an integration time of 100 ms corresponds to an MDL of 5 ppb for N₂O detection. Furthermore, an analysis of the stability of the dual-gas sensor was performed by acquiring the QEPAS signal of N₂O and H₂O vapor for 20 min. This analysis is usually performed by fixing the laser current to correspond to the gas absorption peak, while the gas keeps flowing in the gas line. To avoid QEPAS signal fluctuations due to the possible laser sources instabilities and drifts, we applied two fast voltage ramps to the QCL and diode laser in order to scan the selected N₂O and H₂O absorption lines, respectively. The results are shown in Figure 8. The peak-to-peak amplitude V_ripp and frequency f_ripp of the ramps applied to the diode laser and to the QCL were 15 mV and 30mHz, and 60 mV and 50 mHz, respectively. The f_ripp values were selected to provide a fast absorption line scan and avoid down-sampling reconstruction.

As clearly visible in Figure 8, both the H₂O and N₂O peak signal values remain nearly constant with fluctuations lower than 0.9%, demonstrating the time stability of the dual gas QEPAS sensor.

### CONCLUSIONS

In this work, we reported on the realization of a dual-gas QEPAS system operating in the near- and mid-IR, capable of simultaneous detection of two different gas species in a mixture. The sensor was based on a frequency-modulated multiplexing scheme. A custom QTF allowed the simultaneous
excitation of the fundamental and the first overtone flexural modes using a near-IR diode laser and a mid-IR DFB-QCL as excitation sources. The feasibility of the simultaneous detection of water vapor/methane and water vapor/nitrous oxide was demonstrated. The sensor was calibrated for the detection of methane and nitrous oxide with a constant concentration of water vapor fixed at 1.6%, showing a linear response of the QEPAS signals with the gas target concentration. The minimum detection limits achieved were 18 ppb for CH₄, 5 ppb for N₂O and 20 ppm for water vapor, at a lock-in integration time of 100 ms. The developed QEPAS sensor is suitable for real-time simultaneous detection of two gas species in ambient air, and in particular to detect one gas species (methane or nitrous oxide) and monitor the water vapor in the mixture. This is particularly important for QEPAS sensing, since when operated with slow relaxing gas, such as NO, CO or CO₂, and CH₄ water vapor acts as promoter of the sensing, since when operated with slow relaxing gas, such as vapor in the mixture. This is particularly important for QEPAS species (methane or nitrous oxide) and monitor the water gas species in ambient air, and in particular to detect one gas

**Figure 8.** Simultaneous QEPAS spectral scans for a mixture of 600 ppm of nitrous oxide in humidified air. (a) H₂O vapor signal detected at f₁ applying a narrow ramp to the diode laser current centered on the water vapor line at 7181.16 cm⁻¹. (b) N₂O signal detected at f₂ applying a narrow ramp to the QCL current centered on the N₂O absorption line at 1297.05 cm⁻¹.

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The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

**Notes**

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