Compact all-fiber quartz-enhanced photoacoustic spectroscopy sensor with a 30.72 kHz quartz tuning fork and spatially resolved trace gas detection

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An ultra compact all-fiber quartz-enhanced photoacoustic spectroscopy (QEPAS) sensor using quartz tuning fork (QTF) with a low resonance frequency of 30.72 kHz was demonstrated. Such a sensor architecture has the advantages of easier optical alignment, lower insertion loss, lower cost, and more compact compared with a conventional QEPAS sensor using discrete optical components for laser delivery and coupling to the QTF. A fiber beam splitter and three QTFs were employed to perform multi-point detection and demonstrated the potential of spatially resolved measurements.

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Laser based trace gas sensing methods are widely used in many applications, such as atmospheric chemistry,1 industrial process control and combustion diagnostics,2 as well as medical diagnostics and the life sciences,3 due to their high detection sensitivity and selectivity. Among these methods, photoacoustic spectroscopy (PAS) is one of the most effective trace gas detection technologies. PAS employs a broadband microphone for acoustic wave detection and is characterized by a low cost and robust architecture. However, most microphone-based PAS cells have a low resonance frequency, which makes PAS cells more sensitive to environmental and sample gas flow noise. Moreover, the size of a typical photoacoustic cell is relatively large.3

Quartz-enhanced photoacoustic spectroscopy (QEPAS) technique5 is a significant improvement of the microphone-based PAS method. QEPAS uses low cost, commercially available millimeter sized piezoelectric quartz tuning forks (QTFs) as acoustic wave transducers which possesses high selectivity and immunity to ambient acoustic noise.5–8 In QEPAS technology, the acoustic energy is accumulated in the sharply resonant QTF instead of a large photoacoustic cell as in conventional PAS. Therefore, the limitation of gas cell size no longer exists and the cell volume can be reduced, and even the cell can be optional.

In QEPAS, usually an optical fiber collimator and focusing lens are used for laser delivery and coupling.9–11 An all-fiber configuration has the advantages of easier optical alignment, lower insertion loss, reduction in sensor size, and lower cost. To-date, QEPAS has not been applied to multipoint detection and spatially resolved trace gas concentration measurements.

In this work, a single mode fiber was used for laser delivery to construct an ultra compact all-fiber QEPAS sensor system. In addition, a fiber beam splitter (FBS) was employed to split the laser beam into three branches, which resulted in three point QTF based gas detection and demonstrated multi-point detection and spatially resolved trace gas concentration measurements.

Commercially available QTFs with a resonant frequency \(f_0\) of \(\sim 32.76\) kHz are usually used in QEPAS sensors. The QEPAS signal amplitude is inversely proportional to the QTF resonance frequency. A QTF with a smaller \(f_0\) will result in a longer effective integration time, which increases the QEPAS signal. In this paper, a QTF with \(f_0\) of 30.72 kHz was used as an acoustic wave transducer. The length, width, and thickness of the QTF prongs and the gap spacing between the two prongs are listed in Table I.

The experimental sensor system is shown in Fig. 1. Water vapor (H\(_2\)O) was chosen as the target analyte. A 1.395 \(\mu\)m continuous wave, distributed feedback (CW-DFB) fiber-coupled diode laser was employed as the laser excitation source. The diode laser output was divided in a 1:1:1 ratio by means of a single mode FBS for the three all-fiber QEPAS sensor system with three QTFs. The single mode fiber employed in the sensing platform has a length of 3 m, a core diameter of 8.2 \(\mu\)m, and an insertion loss of \(<0.3\) dB. The fiber end was placed on an adjustable base which can be varied in the x, y, and z directions. The fiber output was placed close to the QTF prongs without an optical focusing lens. The distance between the laser output and the QTF prongs was 300 \(\mu\)m, and the beam diameter at the center of the QTF was \(\sim 100\) \(\mu\)m. Modulation of the laser current was implemented by applying a sinusoidal dither to the direct current ramp of the diode laser at half of the QTF resonance frequency \((f = f_0/2)\). The piezoelectric signal generated by the QTF was detected by a low noise transimpedance amplifier with a 10 M\(\Omega\) feedback resistor and converted into a voltage. Subsequently, this voltage was transferred to a

<table>
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<th>TABLE I. Parameters of geometries for QTF.</th>
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<td>QTF with (f_0) (kHz)</td>
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<td>30.72</td>
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lock-in amplifier for a measurement of the $2f$ component generated by the QTF.

Laboratory air was employed as the target analyte which contained 0.46% H$_2$O as determined by means of a tunable diode laser absorption spectroscopy (TDLAS) method. The H$_2$O absorption line located at 7168.4 cm\(^{-1}\) was selected for laser excitation and the laser power for each sensor branch was $\sim$4.83 mW. The all-fiber QEPAS sensor performance using the three QTFs with $f_0$ of 30.72 kHz was evaluated. The impact of $\Delta Y$ (see Fig. 1 inset) between the laser beam and the top of QTF prongs on QEPAS signal level was investigated, and the experimental results are shown in Fig. 2. The modulation depth of laser wavelength was set to 0.39 cm\(^{-1}\). The $2f$ QEPAS signal amplitude increased rapidly with $\Delta Y$ when it was $\leq$0.6 mm. The peak region of the $2f$ signal amplitude occurred in the range of $\Delta Y$ from 0.6 mm to 1 mm. With a further increase of $\Delta Y$, the signal amplitude decreased due to the more challenging QTF prong vibration when the acoustic wave source is at the bottom of the QTF prongs. At $\Delta Y$ of 0.8 mm, the $2f$ QEPAS signal amplitude was 163.2 mV, 166.6 mV, and 171.3 mV for QTF1, QTF2, and QTF3, respectively. The small differences of signal response among the three sensors mainly resulted from small differences of the three QTFs such as the Q-factor. In the following experiments, the optimum value of $\Delta Y$ of 0.8 mm was chosen to achieve the maximum QEPAS signal amplitude.

The laser wavelength modulation depth was optimized in order to improve the $2f$ QEPAS signal amplitude. The dependence of the all-fiber QEPAS signal amplitude as a function of the laser wavelength modulation depth is depicted in Fig. 3. The all-fiber QEPAS signal amplitude increased with the modulation depth, but when the modulation depth was $>$0.59 cm\(^{-1}\), no further significant change was observed. Therefore, a modulation depth of 0.59 cm\(^{-1}\) was found to be optimum.

The measured $2f$ QEPAS signal and noise at a modulation depth of 0.59 cm\(^{-1}\), and $\Delta Y$ of 0.8 mm for the three QTFs is shown in Fig. 4. The signal amplitude was 222.9 mV, 229.2 mV, and 233.2 mV for QTF1, QTF2, and QTF3, respectively. The QEPAS based sensor noise was determined as a standard deviation from the signal far from the targeted absorption line. The $1\sigma$ noise level was 4.81 $\mu$V, 4.76 $\mu$V, and 4.87 $\mu$V for QTF1, QTF2, and QTF3, respectively. The three all-fiber QEPAS sensors have almost the same signal and noise response. The signal-to-noise ratios (SNRs) calculated from the results measured were 46.3, 48.2, and 47.9. This resulted in a minimum detection limit (MDL) for H$_2$O of 99.4 ppm, 95.4 ppm, and
Three QTFs were used for H$_2$O detection simultaneously to split the laser beam into three sub sensor systems, and lower insertion loss, and lower cost. FBS was employed for multi point detection. The MDL of 99.4 ppm, 95.4 ppm, and 96 ppm was achieved for QTF1, QTF2, and QTF3, respectively. The corresponding calculated normalized noise equivalent absorption (NNEA) were $1.11 \times 10^{-7}$ cm$^{-1}$ W/Hz, $1.06 \times 10^{-7}$ cm$^{-1}$ W/Hz, and $1.07 \times 10^{-7}$ cm$^{-1}$ W/Hz, respectively.

H$_2$O detection at three different locations was performed to demonstrate the potential of spatially resolved trace gas concentration measurements. Locations A, B, and C were in three different cavities. The H$_2$O concentrations of the three different locations were controlled by changing the temperature and relative humidity. The measured results are shown in Fig. 5. The signal amplitude was 223 mV, 97 mV, and 61.1 mV for QTF1, QTF2, and QTF3, respectively. The H$_2$O concentration deduced from the measured QEPAS results were 0.46%, 0.20%, and 0.13% for location A, B, and C, respectively. This was also confirmed by means of the QEPAS based sensor system can be further improved by using an acoustic resonator.

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FIG. 5. H$_2$O detection for three different locations using the all-fiber QEPAS sensor.