Calibration-free mid-infrared exhaled breath sensor based on BF-QEPAS for real-time ammonia measurements at ppb level

Biao Li, Chaofan Feng, Hongpeng Wu, Suotang Jia, Lei Dong

A B S T R A C T

Breath analysis, compared with blood analysis, can enable real-time non-invasive diagnostics through quantification of exhaled biomarkers. Here, we demonstrate, for the first time, a calibration-free mid-infrared (MIR) exhaled breath sensor for real-time ammonia measurements at ppb-level, which can be used for the non-invasive diagnosis of liver and kidney diseases. The exhaled breath sensor employs a 10.359-μm MIR quantum cascade laser (QCL) to target a strong NH₃ absorption line and adopts the beat-frequency quartz-enhanced photoacoustic spectroscopy technique (BF-QEPAS) to remove the requirements of calibration process and wavelength locking in the conventional QEPAS technique. By studying the adsorption-desorption effect and optimizing the modulation depth and modulation frequency of the sensor system, a detection limit of 9.5 ppb is achieved at an integration time of 3 ms. The ammonia content exhaled by 8 healthy volunteers is recorded and the real-time measurement results are analyzed. Compared with conventional QEPAS sensors, the proposed BF-QEPAS-based sensor offers higher sensitivity, faster response time, as well as no need for calibration and wavelength locking.

1. Introduction

Variations in ammonia (NH₃) levels in human body are related to the dysfunction of the liver and kidneys since ammonia is mainly excreted from human body through the metabolic processes of the liver and kidneys. When diseases occur in the liver and kidneys, metabolic disorders cause higher ammonia levels in the body [1–3]. However, current medical measurement regarding ammonia levels in human body relies on blood analysis that is an invasive diagnostic with the risk of infection. In fact, ammonia can enter alveoli via passive diffusion and then is exhaled. The breath analysis for exhaled ammonia can be used for the identification and monitoring of procedures for clinical diagnostics due to the fact that the correlation between breath ammonia and breath ammonia has been verified [4].

At present, trace ammonia detection is demanded in various fields and a variety of detection methods were reported. In 2016, Andreas et al. [5] completed the detection of ammonia based on the chemo-resistive method. In 90% relative humidity, the detection sensitivity reaches 400 ppb. In 2021, Shao et al. [6] realized the detection of ammonia based on multi-walled carbon nanotube technology. The detection sensitivity reached 200 ppb, and the response time and recovery time were 17 s and 18 s, respectively. In 2014, Yao et al. [7] reported an ammonia sensor based on graphene/microfiber hybrid waveguide technology with a resolution of 300 ppb. Although all these methods enable the ammonia measurements, there is an urgent need for developing new ways to accomplish the detection of respiratory ammonia in humans owing to the requirements of extremely high resolution, small sample volume, rapid response time, and long calibration interval in respiratory analysis for clinical diagnostics.

In recent years, with the development of photoacoustic technology, more and more photoacoustic gas sensing systems were reported, among which the trace-gas sensor based on quartz-enhanced photoacoustic spectroscopy (QEPAS) is a suitable candidate for respiratory analysis as it is featured as better immunity to environmental acoustic noise and stronger capability to analyze trace gas samples. Instead of conventional microphones, the QEPAS based sensor employs a quartz tuning fork (QTF) as a resonant acoustic transducer, which ‘listen’ to a weak acoustic wave generated when optical radiation interacts with a trace gas [8,9]. So far, the QEPAS technology has been widely used in environmental monitoring, industrial process control, as well as security and defense [10–26]. In 2011, Lewicki et al. realized real-time measurement of human exhaled ammonia based on QEPAS, in which a 10.34-μm
distributed feedback quantum cascade laser (DFB-QCL) was employed using a 3f wavelength locking technique. The sensor system achieved a high detection sensitivity of 6 ppb [27]. But the inherent high-Q characteristic of the QTF makes it susceptible to the environmental parameters, including temperature, pressure, humidity as well as flow and requires a long time to accumulate weak acoustic energy, which results in a frequent calibration requirement and a slow response time. Furthermore, the use of the 3f wavelength locking technique locking the laser wavelength onto the absorption line made the system complex and fragile.

In 2017, Wu et al. [28] put forward beat-frequency quartz-enhanced photoacoustic spectroscopy (BF-QEPAS), which offers unique advantages over the conventional QEPAS in terms of response time and calibration interval. The basic concept of BF-QEPAS requires that the laser modulation frequency is detuned from the QTF resonant frequency in such a way that when the laser wavelength has a quick scan through the targeted absorption line, a beat frequency signal between two frequencies can be generated, based on which the resonance frequency and Q-factor of the QTF, as well as the trace gas concentration, can be rapidly obtained simultaneously. Hence, BF-QEPAS avoids the calibration process and wavelength-locking requirement and permits real-time monitoring of a targeted trace gas. In this work, for the first time, we develop a calibration-free mid-infrared (MIR) exhaled breath sensor for real-time ammonia measurements at ppb level using BF-QEPAS and a 10.359 μm mid-IR QCL, removing the technical defects of the conventional QEPAS breath sensors, in which the frequency and Q-factor of the QTF requires to be calibrated frequently. Compared with previous work [30,31], our MIR exhaled breath sensor benefits from BF-QEPAS, offering higher sensitivity, faster response time, and no calibration requirement.

2. Selection of absorption line and light source for ammonia detection

A strong targeted absorption line is particularly important for the detection of human exhaled ammonia since exhaled ammonia levels are between 0 and 1500 ppb in general for a healthy person, which requires a high detection sensitivity. According to the HITRAN database, the simulated ammonia absorption intensity between 1 μm and 20 μm is shown in Fig. 1. The NH3 molecule has four fundamental vibrational modes (v1, v2, v3, v4), in which the v2 band and other hot bands (2v2-v3) near 10 μm (1000 cm⁻¹) obviously have the strongest absorption. In such a wavelength region, the main spectral interference comes from H2O and CO2 which are the important components of exhaled air. Spectral simulations for exhaled breath between 965.0 cm⁻¹ to 966 cm⁻¹ at 130 Torr were performed using the HITRAN database with H2O (5%), CO2 (5%), and NH3 (10 ppb), as shown in the inset of Fig. 1. The use of low pressure is to avoid the spectral overlaps from the pressure broadening. One free from H2O interference potential NH3 absorption line can be detected at 965.35 cm⁻¹. Although a weak CO2 line overlaps with the NH3 line, we assess that there will not have a significant spectral interference from the weak CO2 line, due to the fact that the linewidths of the two lines are very different so that an appropriate modulation depth matching the NH3 linewidth can just pick out the NH3 line, and on the other hand the concentration level of NH3 in exhaled breath is much greater than 10 ppb. Therefore, the absorption line located at 965.35 cm⁻¹ was determined as the targeted line.

A continuous-wave (CW) DFB-QCL emitting at a center wavelength of 10.359 μm was employed as the light source (Ningbo Healthy Photo Technology, QC-Qube 200831-AC712), to target the selected ammonia absorption line. The output wavelength and the power of the QCL as a function of the driving current were recorded using a wavelength meter and a power meter, respectively, as shown in Fig. 2. The wavelength tuning range covers from 964.955 cm⁻¹ to 966.873 cm⁻¹. The corresponding current and temperature tuning rates with regards to wavelength are calculated to be 0.009 cm⁻¹/mA and 0.0377 cm⁻¹/°C, respectively. The horizontal dash grey line in Fig. 2 marks the position of the selection absorption line. Since a higher optical power can induce a stronger photoacoustic signal, the laser power should be set as high as possible. Considering the limitation of the maximum laser current at 550 mA according to the laser datasheet, a laser power of 48 mW was selected, corresponding to the driving current and temperature of 532 mA and 30 °C, respectively.

3. Design of sensor system

The schematic diagram of the sensor system is shown in Fig. 3(a). The sensor system consists of three parts, a respiratory sampling system, a photoacoustic sensing unit, and a control and data processing unit. The respiratory sampling system was designed to collect the exhaled breath and to regulate the gas pressure and flow, providing a suitable measurement environment for photoacoustic detection. After a human subject breathed into the mouthpiece, the content of CO2 and the airway pressure were monitored by a CO2 mainstream module (Zhejiang National Medical, C500) and a mini pressure meter (CFsensor, XGZP6847A), respectively. Two needle valves were installed upstream and downstream of the photoacoustic sensing unit, which worked together with a diaphragm pump providing negative pressure, to set the pressure and gas flow rate of the photoacoustic sensing unit. A pinhole-
sized vent was set after the CO\textsubscript{2} mainstream module to discharge the excess exhalation. A one-way valve was installed in front of the upstream valve, to quickly flush the photoacoustic sensing unit and thus to reset zero signal when stopping blowing.

The photoacoustic sensing unit is the core part of the sensor system, which used the BF-QEPAS technique and employed a small-sized piezoelectric QTF as an acoustic transducer. The QTF has a high resonance frequency of 15 Hz (32,768 Hz) in a vacuum, resulting in immunity to low-frequency environmental noise for the photoacoustic measurement. The sound and vibration of the diaphragm pump does not affect QTF performance. Two acoustic micro-resonators with the lengths of 4.0 mm and the inner diameters of 0.8 mm were placed at two sides of the QTF, respectively, forming a spectrophone with an on-beam QEPAS configuration. Such a configuration can improve the signal-to-noise ratio up to 30 times [9]. The spectrophone was enclosed into a gas cell with two ZnSe windows, whose inlet and outlet were directed to the respiratory sampling system. The beam of the used DBF-QCL was focused by a ZnSe plane convex lens with a focal length of 25 mm and then passed through the acoustic micro resonators and the QTF in the gas cell. A low noise trans-impedance pre-amplifier with a feedback resistor of 10 M\textOmega was used to amplify the weak electrical signal from the QTF.

In the control and data processing unit, a laser driver (Healthy Photon QC750-Touch\textsuperscript{TM}) was employed, providing the operating current of the laser, and controlling its temperature. A commercial two-channel waveforms generator circuit board (Juntek MHSS200A) produced a rapid scanning wave for the generation of the beat frequency signal and a sinusoidal wave for the wavelength modulation, both of which were added and then sent to the laser driver. The electrical signal from the QTF was demodulated by a lock-in amplifier board (FEMTO, LIV-BVD-150-H), then was collected by a 16-bit data acquisition card (DAQvntech, USB_HRF4626), and was finally processed by the LabVIEW program that ran on an upper computer. Two prototype photos of this sensor system regarding its appearance and internal structure are shown in Fig. 3(b).

Since ammonia is considered a polar molecule, it tends to stick to metal surfaces, delaying the response time and degrading the measurement accuracy. It is, therefore, important to understand the adsorption mechanism of ammonia on surfaces. The adsorption-desorption process under dynamic equilibrium can be expressed as [29]:

$$\frac{V}{\tau} \frac{\partial c_a}{\partial t} + c_a \cdot \Phi = c^0_a \cdot \Phi - \frac{\partial c_b}{\partial t} \cdot S$$  

(1)

where \(V\) is the volume of the container [cm\textsuperscript{3}], \(c^0_a\) is the ammonia concentration [cm\textsuperscript{-3}] at the measurement point, \(\Phi\) is the gas velocity [cm\textsuperscript{3}/s] in the gas path, \(c_b\) is the ammonia concentration at the gas path inlet, \(\partial c_b/\partial t\) represents the effective adsorption rate [cm\textsuperscript{3}/s] of ammonia molecules at quasi-steady state, \(S\) is the surface area of the gas cell [cm\textsuperscript{2}]. When \(c_b \cdot \Phi = \Phi \cdot S\), the inflow rate of ammonia molecules is much greater than the adsorption rate and thus the viscous effect of the gas can be negligible. There are three ways to meet the above condition. First, since the ammonia concentration levels in exhaled breath are unpredictable, a higher gas flow rate \(\Phi\) and a smaller surface area \(S\) are helpful to reduce the adsorption effect. Secondly, a higher temperature makes the ammonia molecules move quickly and reduces \(\partial c_b/\partial t\). Thirdly a non-metallic material such as Teflon can provide a small \(\partial c_b/\partial t\).

Based on the discussion about the adsorption-desorption effect, the piping in the respiratory sampling system was made of Teflon and was heated to 42 °C by means of a heating belt. In such a temperature, the adsorption effect of ammonia was reduced, the exhaled water vapor did not condense and the laser temperature can be easily controlled. The surface area of the gas cell is only 15 cm\textsuperscript{2} thanks to the capability of QEPAS to analyze trace-gas samples. The gas flow rate was set to 60 sccm (standard cubic centimeter per minute), which is the maximum value that the respiratory sampling system can provide due to the capacity limit of the diaphragm pump operating at 130 Torr.

4. Self-calibration process

Compared with the conventional QEPAS, the BF-QEPAS has the advantage of being able to self-calibrate the QTF resonant frequency \(f_0\) and quality factor \(Q\). According to the beat frequency theory, \(f_0\) can be expressed as [28]:

$$|f_0 - f| = \frac{1}{\Delta t} = \Delta f$$  

(2)

where \(f\) is the modulation frequency of the laser, \(\Delta t\) is the time difference between two adjacent peaks or valleys of the beat frequency signal and \(\Delta f\) is the difference between \(f_0\) and \(f\). Since \(f\) is known and \(\Delta t\) can be obtained from the beat frequency signal, \(f_0\) can be derived from Eq. (2). According to the classical oscillator theory, the quality factor can be written as:

$$Q = \frac{\pi f_0}{\Delta f}$$  

(3)

where \(\tau\) is the response time of QTF which can be obtained from the decay time of the beat frequency signal. Thus, \(Q\) can also be calculated.

To prove the feasibility of this self-calibration approach, a certified 1-ppm NH\textsubscript{3}/N\textsubscript{2} gas mixture was filled into the sensor. A right-trapezoid-like waveform with a period of 330 ms was applied as the scanning waveform, corresponding to the laser current range of 525–538 mA and the wavelength range of 965.45 cm\textsuperscript{-1} - 965.3 cm\textsuperscript{-1}. A 32.7 kHz sinusoidal waveform with a modulation depth of 0.047 cm\textsuperscript{-1} was used as the modulation signal. The integration time and the filter slope of the lock-in amplifier were set to 3 ms and 12 dB [28], respectively, corresponding to a detection bandwidth of 83.33 Hz to provide a sufficient response bandwidth and to maintain efficient background noise suppression. The beat frequency signal was demodulated at the 1 Hz mode. The measured beat frequency signal is shown in Fig. 4(a). The fluctuations of the beat frequency signal between 0 s and 0.05 s were from the neighboring ammonia line at 965.5 cm\textsuperscript{-1}. The time difference \(\Delta t\) between two adjacent peaks was measured to be 0.01887 s. The decay time \(\tau\) was obtained to be 0.0377983 s through an exponential
fitting. Hence \( f_0 \) and \( Q \) can be calculated to be 32,753 Hz and 3889, respectively, according to Equ. (2)-(3). A traditional electric excitation method that must interrupt the process of gas concentration measurement was used to obtain the accurate \( f_0 \) and \( Q \). The method gave \( f_0 = 32,754 \) Hz and \( Q = 3891 \). The good consistency between the results from the two methods verified the feasibility and reliability of the self-calibration approach. To maximize the detection sensitivity of the sensor system, the amplitude of the first maximum peak of the beat frequency signals was chosen as the signal value. A LabVIEW program was written and run in the upper computer to analyze the acquired beat frequency signal in real-time and self-calibrate the sensor system using the obtained \( f_0 \) and \( Q \).

5. Experimental results and discussions

5.1. Modulation depth and frequency optimization

In conventional QEPAS-based sensors, the 1\( f \) signal amplitude is usually larger than the 2\( f \)'s. The reason why the 2\( f \) signal is usually adopted in the conventional QEPAS-based sensors is because the maximum value of the 2\( f \) signal corresponds to the peak of the measured absorption line. However, in BF-QEPAS, there is no difference between the waveforms of the 1\( f \) and 2\( f \) signals since an oscillating beat-frequency signal is measured. Thus, the 1\( f \) signal with a larger amplitude should be used to improve the detection sensitivity. The modulation depth was first optimized in the case of the 1-ppm NH\(_3\):N\(_2\) gas mixture filled into the gas cell. The amplitude of the first maximum peak of the beat frequency signals was acquired and depicted in Fig. 5 as a function of the modulation depth. The amplitude linearly increases together with the modulation depth increasing in the range of 0.004–0.025 cm\(^{-1}\). Then the amplitude reaches the maximum value when the modulation depth is equal to 0.047 cm\(^{-1}\), which is the optimal modulation depth.

The frequency difference \( \Delta f \) between \( f_0 \) and \( f \) is a very important parameter since the different \( \Delta f \) determines the different detection sensitivity. The amplitude of the first maximum peak of the beat frequency signal was recorded as a function of the modulation frequency \( f \) at the optimal modulation depth of 0.047 cm\(^{-1}\), as shown in Fig. 5 as a function of the modulation depth. The amplitude linearly increases together with the modulation depth increasing in the range of 32,704 Hz and 32,804 Hz, corresponding to a \( \Delta f \) of 50 Hz. Since \( f_0 \) is obtained in each measurement of the beat frequency signals, the new modulation frequency \( f \) can be calculated by Eq. (2), ensuring the consistency of the detection sensitivity once the QTF resonant frequency \( f_0 \) drifts.

5.2. Evaluation of sensor linearity

The linearity of the BF-QEPAS-based sensor system was assessed. The different concentration levels of NH\(_3\), 50 ppb, 100 ppb, 200 ppb, 400 ppb, 600 ppb and 1 ppm, were generated by diluting the certificated 1 ppm NH\(_3\):N\(_2\) gas mixture using the gas dilution system and were directed into the sensing system in turn. 165 points were measured continuously for each concentration level with a data updating rate of 330 ms/point. High purity N\(_2\) was introduced into the sensor for zero baseline measurement. Fig. 7(a) shows the experimental results. The amplitude of the beat frequency signals for 50 ppb NH\(_3\) was 0.32 mV, while the standard deviation \( \sigma \) of the noise signal was 0.06089 mV, resulting in a signal-to-noise ratio of 5.25. The corresponding detection
limit and normalized noise equivalent absorption (NNEA) are 9.5 ppb and $1.42 \times 10^{-9}$ cm$^{-1}$ W/Hz$^{1/2}$, respectively. An average of the signal values was then performed for each concentration level, and the relationship as a function of concentration value was recorded in Fig. 7 (b). The R-squared value between the measured and nominal concentrations was 0.999, confirming the good linear response of the sensor system to the NH$_3$ concentration.

6. Real-time ammonia measurement in exhaled human breath

Real-time ammonia measurement of exhaled human breath based on the BF-QEPAS sensor system was demonstrated. Eight healthy volunteers were asked to relax and complete the gentle exhalation in a normal posture, as shown in Fig. 8 (a). The subject does not need to make special preparatory movements before exhalation and try to empty the lungs of as much gas as possible during exhalation, making the measurements more accurate. The pressure values of the airway were displayed on the upper computer screen using a graduated bar so that the subjects can control their exhalation strength within the pressure range of 8–12 Torr according to the real-time pressure measurement. Such an exhalation control is to enable the subjects to have a continuous and stable blow and thus to maintain the stable pressure and flow rate of the sensor system.

The measurement result of one typical exhalation is presented in Fig. 8(b). The black line plus diamond represents the values of the airway pressure, which should be a horizontal line during an exhalation owing to the active pressure control. The blue line plus triangle represents the subject’s exhaled CO$_2$ content, which is a real-time measurement of the reference profile for exhaled ammonia content. A healthy human exhales CO$_2$ at 4–6% at a time. And the CO$_2$ content should gradually rise with the blowing time, which can reflect the complete process of one exhalation. When any of the two curves are abnormal, it implies that a complete breathing process is not collected and the NH$_3$ data for this exhalation should be discarded. The red curve represents the NH$_3$ concentration values, which can be divided into four phases. Phase I is the rising process of the NH$_3$ signal. According to the airway pressure curve, the exhalation begins at 2 s, while the NH$_3$ concentration starts to rise at 3 s and ends at 10 s. The delay between the start of blowing and the maximum signal is partly because of the adsorption effect and partly because the exhaled breath needs to pass through a certain length of the tube to enter the gas cell. Subsequently, the concentration signal arrives in and stays a plateau, which is Phase II. However, this ammonia concentration does not come from the alveoli, but from the fermentation of residual food in the oral cavity since each subject has all kinds of food every day, and hence cannot be used as a valid test result. In Phase III, with the NH$_3$ in the oral cavity evacuated, the NH$_3$ concentration drops and then stays at a lower plateau, as shown in the cyan rectangle of Fig. 8(b). The measured value at this time, corresponding to 290 ppb, reflects the true concentration level of NH$_3$ in the alveoli and can be used as a clinical reference. At the time of 32 s, the exhalation process enters Phase IV where the exhalation stops as the airway pressure drops to zero. The zeroing process of the NH$_3$ signal takes ~9 s (32–41 s) due to the adsorption effect, while the CO$_2$ signal quickly returns to zero. The measurement results of the eight healthy subjects are presented in Fig. 8 (a), having a concentration distribution in the range of 150–640 ppb, which are below the safety threshold of 1500 ppb. The result shows that even the healthy subjects have a large concentration differences between individuals.

7. Conclusions

An exhaled breath sensor for real-time ammonia measurements at ppb level was developed based on the BF-QEPAS technique. This sensor combined the main characteristics of QEPAS with the new benefits of BF-QEPAS, thus providing higher sensitivity, faster response time,
Deployment of the sensor system in hospitals for field tests of the patients with liver and kidney disease to evaluate its performance will be conducted in the coming future.

CRediT authorship contribution statement

Biao Li: Writing – original draft, Investigation. Chaofan Feng: Software, Validation, Data curation. Hongpeng Wu: Visualization, Project administration. Suotang Jia: Methodology. Lei Dong: Conceptualization, Supervision, Writing – review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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References


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