Introduction

Alcohol misuse remains a major health issue and a dominant cause of serious traffic accidents in almost all societies. Many informative and persuasive initiatives have been undertaken. Their importance is beyond dispute, but a holistic approach including clever management, adequate resources, and sustainability, is required for them to be maximally effective [1]. The availability of adequate methods and devices to determine breath alcohol on a massive scale is also needed.

Breath alcohol analysis methodology has developed steadily during recent decades. The physiological rationale for replacing blood analysis with breath analysis is now widely recognized [2,3] and some countries have even introduced breath alcohol concentration (BrAC) to drunk driving legislation [4].

Breath alcohol sensor technologies based on electrochemical devices and infrared spectroscopy have evolved and facilitated the development of products for collecting evidence and screening [5]. Industrial standards have also been established for evidential instruments [6], alcohol interlocks [7], and alcometers for screening [8] and for public use [9].

There is a growing demand for more user-friendly and less obtrusive techniques and devices that do not sacrifice accuracy and reliability. The DADSS (driver alcohol detection system for safety) program is one prominent example [10,11]. It is aiming at unobtrusive operation, while maintaining requirements on accuracy which are matching or even surpassing those of evidential instruments [6].

Keywords: Breath alcohol analysis; Infrared spectroscopy; Accuracy; Unobtrusiveness

Abstract

The study objective was to evaluate a novel method and technology for unobtrusive determination of breath alcohol in relation to current industrial accuracy standards. The methodology uses carbon dioxide as a tracer gas detected by sensor technology based on infrared spectroscopy. Part one of the investigation was to analyse the performance of hand-held prototype devices and included tests of resolution, unit-to-unit variation during calibration, response to alcohol containing gas pulses created with a wet gas simulator, and cross sensitivity to other substances. In part two of the study, 30 human participants provided 1465 breath tests in both unobtrusive and obtrusive use modes. The results of both parts of the study indicate that the prototype devices exceeded present industrial accuracy requirements. The proposed methodology and technology eliminate the previous contradiction between unobtrusiveness and high accuracy.

In earlier publications, we have demonstrated methods for contactless determination of BrAC [12,13] in screening applications where sobriety is expected to be the norm. The physiological rationale of using a tracer gas, e.g., CO₂, for contactless determination was examined [14], and the usefulness of this technique in patients with reduced consciousness was demonstrated [15]. Recently, unobtrusive BrAC determination [16] was demonstrated and progress concerning measurement accuracy was reported [17].

In this paper, the seemingly contradictory demands for both unobtrusiveness and high accuracy are addressed. New experimental results from tests using both artificial and human subjects are provided. The interpretation of these results is discussed in a wider context, including an issue recently raised by Grubb et al. [18] in this journal, in which the use of CO₂ as a tracer gas was supposed to “introduce methodological flaws”.

Methods and Materials

In this section we address the methodology used in the present investigation, in relation to industrial standards, the adapted technology, and breath alcohol determination in general. The distinction between technical and physiological error sources is outlined, with special attention to the need for unobtrusive measurement.

Industrial Standards

In Table 1, the current industrial standard accuracy requirements are summarized. The technical accuracy of breath alcohol analysers is commonly expressed as a combination of an offset error expressed in mg/L and a calibration error expressed in the percentage of the reading. In alcohol interlocks, the allowed error close to the concentration limit is also specified as a maximum allowed error using a function test with an artificial gas pulse generator. The cross-sensitivity to common endogenic or exogenic substances is specified as the ratio between the maximum allowed reading and the reading at the same concentration of ethyl alcohol (EtOH).

In many applications, breath alcohol analysers are used to classify
whether a specific concentration limit has been exceeded or not. For example, most European countries have a legal concentration limit of 0.25 mg/L for drunk driving; which approximately corresponds to a blood concentration of 0.05%. In the classifier situation, the accuracy of a breath alcohol sensor can be described by its ability to discriminate between true positive and negative outcomes, and the corresponding false positive and negative outcomes.

**Technology, Prototypes, and Modes of Operation**

The prototypes in the present investigation were based on infrared spectroscopy employing a White cell [19] with a 1.2 m optical path for EtOH and a 30 mm optical path for CO₂. The cell includes emitters and detectors tuned to the absorption peaks of EtOH at a wavelength of 9.5 µm, and CO₂ at 4.26 µm. Signal conditioning, including algorithms for determining BrAC, is performed by on-board electronic circuitry. Photographs of the measuring cell and the hand-held prototype device are shown in Figure 1. Further details of the technology and engineering solutions have been published elsewhere [13,17].

Figure 1 illustrates one of the operational modes of the hand-held breath analyser in its testing environment inside a vehicle. The breath is delivered with a short distance between the device and the test subject’s mouth. In this ‘short-range’ mode, the breath will be diluted by a factor of 1.5-2.5 with ambient air. In the present investigation, two additional modes were included. The first was a ‘long-range’ mode from a distance of approximately 15 cm, for which the dilution may be a factor of 5-10. The other investigational operational mode is delivered with a short distance between the device and the test subject’s mouth. In this ‘short-range’ mode, the breath will be diluted by a factor of 1.5-2.5 with ambient air. In the present investigation, two additional modes were included. The first was a ‘long-range’ mode from a distance of approximately 15 cm, for which the dilution may be a factor of 5-10. The other investigational operational mode was to attach a mouthpiece to the inlet to ensure an undiluted breath.

**Resolution, Calibration, and Function Tests**

Breath analyser resolution is determined by noise behaviour. The prototypes were tested by analysing the EtOH channel output, recording the output signal, and comparing it to the signal obtained from standard concentrations of various substances, including algorithms for determining BrAC, is performed by on-board electronic circuitry. Photographs of the measuring cell and the hand-held prototype device are shown in Figure 1. Further details of the technology and engineering solutions have been published elsewhere [13,17].

Table 1: Technical accuracy requirements according to current industry standards.

<table>
<thead>
<tr>
<th>Standard</th>
<th>Offset error (mg/L)</th>
<th>Calibration error (% of reading)</th>
<th>Function test error</th>
<th>Cross-sensitivity (% of EtOH reading)</th>
</tr>
</thead>
<tbody>
<tr>
<td>EN16280:2012 General public [9]</td>
<td>0.04</td>
<td>±20%</td>
<td>Unspecified</td>
<td>Unspecified</td>
</tr>
<tr>
<td>EN15964:2011 Screening [8]</td>
<td>±0.02</td>
<td>±10%</td>
<td>Unspecified</td>
<td>Acetone 8% Carbon monoxide 20% Methane 25%</td>
</tr>
<tr>
<td>EN50436-1.2:2013 Alcohol interlocks [7]</td>
<td>0.02</td>
<td>±15%</td>
<td>±0.05 mg/L</td>
<td>Acetone 40% Carbon monoxide 100% Methane 67% Methanol 200% Isopropanol 100% +8 more substances</td>
</tr>
<tr>
<td>OIML R 126:2012 Evidential instruments [6]</td>
<td>0.02</td>
<td>±5%</td>
<td>Unspecified</td>
<td>Acetone 20% Carbon monoxide 50% Methanol 100% Isopropanol 100%</td>
</tr>
</tbody>
</table>

The cross-sensitivity of each prototype device was tested by exposing it to controlled concentrations of various substances, including algorithms for determining BrAC, is performed by on-board electronic circuitry. Photographs of the measuring cell and the hand-held prototype device are shown in Figure 1. Further details of the technology and engineering solutions have been published elsewhere [13,17].

The equipment used for the function test (Figure 2, right) includes bottled gas with precisely controlled EtOH and CO₂ concentrations. Gas pulses from the bottles are passed through an aqueous solution, corresponding to the specific EtOH concentration, where it becomes saturated with water vapour corresponding to a human breath.

The functional tests were performed with N=97 of 0.1 mg/L gas pulses on four units, using the output of an evidential instrument (Evidenzer; Nanopuls AB, Uppsala, Sweden) as a reference. Thus, each breath test could be quantified with respect to its deviation from the reference value. A histogram of the statistical distribution provided evidence of both systematic and random error.

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**Unobtrusive Measurement and Physiological Error Sources**

In accordance with Zaouk et al. [10] and Ferguson et al. [11], breath alcohol determination is said to be unobtrusive if it does not interfere with the normal activities of a sober test subject. A breath may be unobtrusively detected by a sensor located at a distance of 10-20 cm from the test subject’s mouth, but it will be diluted with ambient air by a factor of as much as 5-10. This corresponds to the long-range operational mode used in this investigation.

When used in the short- and long-range operational modes, the BrAC value was calculated using the equation (1) [13,14]:

\[
\text{BrAC} = \frac{\text{EtOH}_{\text{meas}} \times D = \text{EtOH}_{\text{meas}} \times (\text{CO}_2_{\text{alv}} - \text{CO}_2_{\text{background}}) / (\text{CO}_2_{\text{meas}} - \text{CO}_2_{\text{background}})}
\]

Where EtOH_{meas}, CO₂_{meas}, CO₂_{alv}, and CO₂_{background} represent measured values, and CO₂_{alv} represents the alveolar CO₂ concentration, which was set at 4.8 vol% in this investigation [22]. D is the dilution factor. The background CO₂ concentration, CO₂_{background} is typically 0.04-0.06%
Human Subjects Test

Out of 30 test subjects, 19 were male and 11 were female, and all were between 19 and 70 years old. Male subjects were given 0.6 g of EtOH per kilogram of body mass and the corresponding amount given to female subjects was 0.55 g of EtOH per kilogram of body mass. Alcohol was consumed in less than 15 minutes and provided an intoxication level of approximately 0.4mg/L. BrAC determinations were then performed every 20 minutes during the EtOH elimination phase. Each measurement set consisted of one test in a reference instrument, one test into the prototype fitted with a mouthpiece, one breath test towards the prototype at a distance of approximately 3 cm, and one test towards the prototype at a distance of approximately 15 cm. In the latter two tests, CO₂ was used to account for the dilution of the breath sample. In total, 1465 tests were performed with the prototype and statistically analysed. The study results thus include technical, physiological, and behavioural error sources. The human subjects study was approved by the Swedish Ethical Review Board in Uppsala (Dnr 2013/089). The reference instrument used throughout the human subject trials was an evidential breath analyser (Evidenzer; Nanopuls AB). Apart from EtOH, the reference instrument was also capable of measuring CO₂ and water concentration in undiluted breath samples. The data collected from human subjects were analysed with respect to compliance with industrial standards.

Results

Results are presented from tests performed on prototype devices using both artificial and human breath.

Resolution, Calibration, and Function Test Cross-sensitivity

Resolution was determined by the EtOH sensor signal noise behaviour. Ideally, "white" noise dominates with a uniform spectral distribution. If the noise magnitude is plotted against the time window
of the measurement, one normally observes a declining characteristic until a minimum is reached. At some point, long-term drift becomes more dominant, which causes the noise to increase. This behavior was observed in our prototypes (Figure 3).

With an integration time of 1 second, the resolution is approximately 0.0009 mg/L. A minimum is observed at approximately 800 seconds, which is more than one order of magnitude lower than the 1 second value.

Figure 4 shows a histogram illustrating the variation of the calibration factor in 115 prototype units. The average value deviated by less than 0.003 mg/L EtOH from the nominal 0.40 mg/L, and the standard deviation was 0.007 mg/L, or 1.75% of the nominal value.

Function tests were performed on four prototypes and 97 breath tests were performed using artificial gas pulses having an EtOH concentration of nominally 0.1 mg/L. The tests were performed at room temperature. The results are summarized in Figure 5, which shows a test result distribution histogram.

Figure 5 illustrates that the measured concentrations have a normal statistical distribution with no major skewness. Compared with the Evidenzer value, the systematic error defined as the deviation between the measured average and the nominal values, was less than 0.01 mg/L, and the standard deviation of the random error was 0.006 mg/L. The units were used in the contactless mode of operation using CO₂ as a tracer gas. The observed variability was considerably smaller than the requirements according to industrial standard EN50436-1,2.

Figure 6 summarizes the cross-sensitivity test results in relation to the requirements of each industrial standard. Unfilled bars illustrate the allowed cross-sensitivity according to different industrial standards and filled bars show the calculated values based on the sensor’s optical filter characteristics. The calculations were confirmed by measurements of the most critical substances, and these results were published recently [17].

The calculated and measured cross-sensitivities of the prototype units fulfilled the requirements of all industrial standards with the exception of methanol, which exceeded the evidential standard requirements by approximately 40%.

Tests with Human Subjects

The variability of end-expiratory CO₂ breath tests is illustrated in Figure 7. As expected, the long- and short-range breath tests exhibited large variations, whereas the variations for the undiluted breath tests were smaller. It is notable that the average of the undiluted values (4.1 ± 0.5 kPa) was significantly lower than the nominal alveolar value (4.8 kPa).

In Figure 8, the results of BrAC determinations for the entire human test population are summarized. In the upper panel, classifier performance is depicted with respect to the Swedish concentration limit of 0.1 mg/L EtOH. The shaded vertical area corresponds to the allowable error according to the function test of EN50436-1,2 centred around the concentration limit. When applying this error band, no FNs or FPs were observed for the short-range tests, whereas one was noted for the long-range tests.

In the lower panel of Figure 8, classifier performance is depicted with respect to the central European concentration limit of 0.25 mg/L. The shaded vertical area corresponds to the allowable error according to the function test of EN50436-1,2. When applying this error band, three FNs and seven FPs were observed for both the short- and long-range tests; therefore, 1.7% of the determinations were falsely classified according to these criteria.

Discussion

Our objective was to evaluate the method and performance of a novel prototype breath alcohol analyzer. The test results using artificial breath samples indicate that the prototype’s performance complies with, or exceeds, the requirements with respect to

![Figure 3: Allan deviation as a function of integration time.](image)
Figure 4: Unit-to-unit variation of calibration factor for a batch of 115 prototype units.

Figure 5: Results of function tests performed on four prototype units.
resolution, calibration factor variation, function test variability, and cross-sensitivity, with methanol being the only deviation with respect to evidential requirements [6].

The human study included 30 individuals and 1465 EtOH breath tests examined with respect to the prototype’s ability to correctly classify the breath tests in relation to the Swedish and central European concentration limits of 0.1 and 0.25 mg/L, respectively. One case of false determination was observed with respect to the Swedish limit, whereas 1.7% of the determinations using CO2 as a tracer gas were falsely classified using the central European limit. When the same prototypes were used in the undiluted mode of operation, no case of false determination was observed regardless of concentration limit.

The significant difference between the observed undiluted CO2 values (Figure 7), and the expected average arterial concentration [22] explains the increased slopes of the regression lines in the diluted modes of operation shown in Figure 8. This observation is in accordance with previous results [24,25] and is related to gas exchange within the respiratory airways.

The prevalence of false determinations was obviously related to the choice of both operational mode and concentration limits. With a low concentration limit, both the short- and long-range modes with CO2 as a tracer gas produced only one false output out of 982 breath tests. At higher concentration limits, the measurement error associated with CO2 variability became more pronounced. In borderline cases, an unobtrusive test may be followed by an undiluted test, which will reduce the risk of false determination to almost zero.

Figure 8 indicates that in both the short- and long-range operational modes, the statistical signal behaviour is heteroscedastic, which was verified by performing the test suggested by Breutsch and Pagan [26]. This behaviour is predictable from Equation (1) and the dominating variability of CO2alv. It is actually a desirable property reducing the dominant error to almost zero at a low EtOH concentration.

Using the technique described in this paper, screening of breath alcohol can be performed at some distance from the mouth of the test subject. It implies a simplified operation for the user or operator compared to present techniques using a mouthpiece. In screening situations where the subjects are normally sober, the vast majority of tests can be performed unobtrusively. When the measurement result is close to the legal limit, and definitely when it is intended as legal evidence, higher accuracy is required. Only then it is necessary to provide an undiluted breath sample, e.g. by using a mouthpiece. The results of the present investigation show that the same type of breath analyser can be used in both the screening and high accuracy modes of operation.
Figure 7: Variation in CO2 for the long- and short-range breath tests, as well as the undiluted tests using a mouthpiece. The error bars represent one standard deviation for tests in each category.

Figure 8: Results of human study of 30 test subjects. Upper: Classifier performance related to the Swedish concentration limit of 0.1 mg/L EtOH. Lower: Classifier performance related to the central European concentration limit of 0.25 mg/L EtOH. See text for more details.
The use of water vapour as tracer gas was patented by Albarda [27] and has been used by Lindberg et al. [28] and Grubb et al. [18]. It has the benefit of enabling high accuracy because of the fact that the inter-individual variations are smaller by a factor of 3-4 compared with CO₂ [14]. However, the background concentration may be significant in most applications and may even exceed the alveolar concentration depending on the local circumstances. Therefore, unobtrusive measurement with high sample dilution is not feasible using H₂O as tracer gas.

It should be noted that physiological error sources are also present in conventional breath tests using a mouthpiece. Depending on the test subject’s lung capacity, a breath test could either represent a deep or a shallow breath, which could give rise to an error exceeding 10% [2,3].

The results reported here demonstrate that the methodological issue raised by Grubb et al. [18] has been effectively resolved. Taking full advantage of the improvements in technology and methodology, there is no longer a contradiction between unobtrusiveness and high accuracy.

References

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