A carbon monoxide ‘single breath’ method to measure total haemoglobin mass: a feasibility study

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Abstract
Anemia is defined by the concentration of haemoglobin (Hb). However, this value is dependent upon both the total circulating haemoglobin mass (tHb-mass) and the plasma volume (PV) – neither of which is routinely measured. Carbon monoxide (CO)-rebreathing methods have been successfully used to determine both PV and tHb-mass in various populations. However, these methods are not yet suitable for ventilated patients. This study aimed to modify the CO-rebreathing procedure such that a single inhalation of a CO bolus would enable its use in ventilated patients. Eleven healthy volunteers performed four CO-rebreathing tests in a randomized order, inhaling an identical CO volume. In two tests, CO was rebreathed for 2 min (optimized CO rebreathing; oCOR), and in the other two tests, a single inhalation of a CO bolus was conducted with a subsequent breath hold of 15 s (Procnew15s) or 30 s (Procnew30s). Subsequently, the CO volume in the exhaled air was continuously determined for 20 min. The amount of CO exhaled after 7 and 20 min was respectively 3.1 ± 0.3 and 5.9 ± 1.1 ml for oCOR, 8.7 ± 3.6 and 12.0 ± 4.4 ml for Procnew15s and 5.1 ± 2.0 and 8.4 ± 2.6 ml for Procnew30s. tHb-mass was 843 ± 293 g determined by oCOR, 821 ± 288 g determined by Procnew15s (difference: P < 0.05) and 849 ± 311 g determined by Procnew30s. Bland–Altman plots demonstrated slightly lower tHb-mass values for Procnew15s compared with oCOR (~21.8 ± 15.3 g) and similar values for Procnew30s. In healthy volunteers, a single inhalation of a CO bolus, preferably followed by a 30 s breath hold, can be used to determine tHb-mass. These results must now be validated for ventilated patients.

KEYWORDS
blood volume, carboxy-haemoglobin, CO rebreathing, ventilated patients
Haemoglobin (Hb) is the oxygen-carrying pigment of the circulation. Its circulating concentration ([Hb]) is routinely measured in clinical practice, and low values are used to define ‘anaemia’ (Beutler & Waalen, 2006). However, [Hb] is determined by the total circulating mass of Hb (tHb-mass) and the volume of plasma (PV) in which it is carried. The measurement of such independent variables has distinct advantages given that PV can change substantially with disease. The importance of tHb-mass measurement in the clinical setting and the advantages over [Hb] have been discussed previously (Otto et al., 2017a,b; Plumb et al., 2016). Indeed, [Hb] correlates poorly with tHb-mass in patients with chronic liver disease or heart failure, in whom PV may be expanded (Otto et al., 2017a). Despite this fact, [Hb] is the major trigger for the transfusion of red blood cells.

Likewise, perioperative changes in tHb-mass and PV are common (Iijima et al., 2013; Makaryus et al., 2018) due to blood loss, administration of red blood cells or haemodilution through administration of intravenous fluids or through salt/water retention due to the ‘surgical stress response’ (Rassam & Counsell, 2005). Fluid distribution between physiological compartments and the impact of hypo/hypervolaemia on the glycocalyx and therefore the functional integrity of the intravascular space also influence PV (Strunden et al., 2011). However, the decision to transfuse blood to a patient in clinical practice in general, and perioperative and critical care settings in particular, hinges on a variety of factors. There is a growing recognition that [Hb] may not be the best clinical indicator to guide such decisions (Plumb et al., 2016; Shander & Ferraris, 2017).

tHb-mass and derived PV can be determined by different dilution methods, in which carbon monoxide (CO) has been found to be the easiest and most precise marker to use (Gore et al., 2005). Using the inhalation of a known volume of CO (thus labelling Hb as carboxyhaemoglobin, COHb) allows the measurement of tHb-mass and, thus, the calculation of PV. In self-ventilating subjects, this is achieved using the so-called ‘optimized CO rebreathing method’ (oCOR). However, this protocol relies upon the participant being alert, able to follow instructions, and able to control their breathing through a closed circuit, which in turn precludes the use of oCOR in participants who are receiving mandatory ventilation from a mechanical ventilator, either under anaesthesia or when sedated in the intensive care unit. We hypothesized that delivery of a single CO bolus into the breathing circuit of a participant without the need for rebreathing could be used to reliably measure tHb-mass, so long as exhaled gas could be analysed. We thus sought to develop such a technique. Here, we describe this development and early data relating to its likely reliability. The primary aim of this study was to develop a new method for measuring tHb-mass in healthy participants simulating a procedure that might be used in participants on a mechanical ventilator (Procvent) and to evaluate the feasibility of this novel method. We aimed to assess reliability compared to the standard oCOR method. We also repeated the oCOR test to quantify the reliability of the standard method within this experiment.

New Findings
- What is the central question of this study?
  Is it possible to modify the CO-rebreathing method to acquire reliable measurements of haemoglobin mass in ventilated patients?
- What is the main finding and its importance?
  A ‘single breath’ of CO with a subsequent 30 s breath hold provides almost as exact a measure of haemoglobin mass as the established optimized CO-rebreathing method when applied to healthy subjects. The modified method has now to be checked in ventilated patients before it can be used to quantify the contributions of blood loss and of dilution to the severity of anaemia.

2 | METHODS

2.1 | Ethical approval

Ethical approval was granted by the South-Central Hampshire B Research Ethics Committee (REC reference: 15/SC/0496) and from the ethics committee of the University of Bayreuth (reference: O1305/1-GB). The study conformed to the standards set by the Declaration of Helsinki, except for registration in a database. Written informed consent was obtained from all participants. The subjects volunteered to participate in the study and were free to withdraw at any time without needing to provide a reason.

2.2 | Subjects

This feasibility study took place at the University Hospital Southampton NHS Foundation Trust, UK, and at the University of Bayreuth, Germany. Eleven healthy non-smoking test subjects (five women, six men) with moderate physical training status took part in the study (for anthropometric data of these subjects see Table 1).

2.3 | Study design

In preliminary tests, we checked whether a single inhalation of a CO bolus could achieve an increase in COHb concentration ([COHb]) that would be sufficient to determine tHb-mass. For this purpose, tHb-mass was measured twice; results from the established CO-rebreathing method (Schmidt & Prommer, 2005), that is, 2 min CO inhalation within a closed spirometry system, were compared to those from a single inhalation with subsequent 10 s breath holding and 15 min collection of expired air. Because the difference in tHb-mass was lower than 50 g
in only six of 13 comparisons, breath holding was prolonged to 15 and 30 s and collection of the expired air to 20 min in the main study.

In the main study, at least four CO-rebreathing tests were carried out by 11 subjects in a randomized order. Two of the tests consisted of the established CO-rebreathing method over 2 min (oCOR). In the third and fourth tests, the subjects inhaled a CO bolus followed by a breath hold for 15 s (Procnew15s) or 30 s (Procnew30s). Afterwards, the volume and CO concentration of the expiratory air were continuously analysed for 20 min, and the whole amount of exhaled air was finally collected in a Douglas bag. tHb-mass was calculated at 2 min intervals using the prevailing [COHb] and the accumulated CO volume in the expired air. Additionally, at the end of the test, tHb-mass was obtained by using the total expiratory volume and the average [COHb] in the Douglas bag. To evaluate possible influences of the test arrangement, that is, collection and analysis of the expired air after inhaling the CO bolus, CO exhalation was determined from six subjects in a fifth test approach using the same methodology as described above after a conventional 2 min CO-rebreathing procedure (oCOR20min).

### 2.4 | Established carbon monoxide rebreathing method

tHb-mass was determined using the optimized CO-rebreathing (oCOR) method as described and modified by Schmidt and Prommer (Prommer & Schmidt, 2007; Schmidt & Prommer, 2005). Briefly, a bolus of 99.97% CO (0.8–1.0 ml CO/kg body mass, depending on the training status) was administered to subjects and rebreathed along with 3 litres of 100% O2 for 2 min. Three arterialized capillary blood samples were taken from a hyperaemic earlobe (Finalgon, Sanofi-Aventis, Frankfurt, Germany) before the rebreathing procedure, and 99.97% CO (0.8–1.0 ml CO/kg body mass, depending on the training & Schmidt, 2007; Schmidt & Prommer, 2005). Briefly, a bolus of CO was administered to subjects and rebreathed along with the three-way valve was turned. Subsequently, the subject deeply inhaled, and CO was administered by the investigator via the access port into the subject’s inhalation path. In contrast to the established method, in which rebreathing occurs in a closed system, this modification presents an open system in which ambient air is inhaled via a three-way valve. The inhaled and expired air passes through a flow sensor (breath-by-breath registration, Metalyzer 3B, Cortex Biophysics GmbH, Leipzig, Germany) and subsequently a small mixing chamber, which is equipped with a CO sensor (Draeger Pak 7000, Liebefeld, Switzerland), and is finally collected in a Douglas bag (Cranial Human Performance Ltd, Birmingham, UK).

After connecting and accustoming the subject to the equipment for at least 10 min in the sitting position, the subject exhaled normally, and the three-way valve was turned. Subsequently, the subject deeply inhaled, and CO was administered by the investigator via the access port into the subject’s inhalation path. The subject held their breath for 15 s (test 3, Procnew15s) or 30 s (test 4, Procnew30s) and breathed normally thereafter for the following 20 min into the Douglas bag. Until the fifth minute after starting the test, the CO concentration and the volume of the exhaled air were monitored at 30 s intervals and thereafter at 1 min intervals until disconnecting the subject from the equipment after 20 min. In the same way as in test 1 and test 2, three capillary blood samples were taken before the test, one sample each was collected after 1 and 2 min, and then further samples were taken every 2 min until min 20.

### 2.5 | Modified method

To adapt the method so that it can be used in everyday clinical practice, several modifications were necessary (see Figure 1). Since a patient frequently cannot put the spirometer into their mouth by themselves, the gas supply was replaced by a mask (Hans Rudolph, Inc, Shawnee, KS, USA) with an access port for the CO supply. This access port is designed in such a way that the manually administered CO from a syringe passes the mask via a small tube directly into the back of the mouth and thus into the test person’s inhalation path. In contrast to the established method, in which rebreathing occurs in a closed system, this modification presents an open system in which ambient air is inhaled via a three-way valve. The inhaled and expired air passes through a flow sensor (breath-by-breath registration, Metalyzer 3B, Cortex Biophysics GmbH, Leipzig, Germany) and subsequently a small mixing chamber, which is equipped with a CO sensor (Draeger Pak 7000, Liebefeld, Switzerland), and is finally collected in a Douglas bag (Cranial Human Performance Ltd, Birmingham, UK).

### 2.6 | tHb-mass calculation

For the established method (tests 1 and 2), tHb-mass was calculated as described previously (Schmidt & Prommer, 2005):

\[
\text{tHb} - \text{mass} = \frac{K \times M_{CO} \times 100}{\Delta \text{COHb} \times 1.39}
\]  \hspace{1cm} (1)

where \(K = \frac{\text{current barometric pressure} \times 760}{\text{current temperature}}\times [1+ (0.003661 \times \text{current temperature})],\)

\[
M_{CO} = CO_{adm} - (CO_{system + lung(After Disconnection)} + CO_{exhaled(After Disconnection)} - M_{Hb} \times \text{CO}_{adm})
\]

\[
= CO \text{ volume administered into the system}
\]

\[
= CO \text{ concentration in the spirometer x (spirometer volume + lung residual volume)}
\]

\[
M_{Hb} = \text{CO diffusing to myoglobin CO}_{exhaled(After Disconnection)}
\]

\[
= \Delta \text{end-tidal CO concentration} \times \text{alveolar ventilation} \times \text{time}
\]
ΔCOHb% is the difference between basal COHb% and COHb% in the blood samples after CO administration. $1.39 \times \text{Hfner number (ml CO x g Hb}^{-1}) \text{ (e.g. Gorelov, 2004).}$

For tests 3 and 4 (Proc_{new}15s and Proc_{new}30s), thb-mass was calculated using formula (1) in two modified ways: (i) for each time point of taking blood after CO inhalation using the corresponding COHb concentrations and accumulated values for CO exhalation ($MCO = CO_{adm} - CO_{exhaled} - MHb$), and (ii) using the COHb concentration at min 20 and the totally exhaled CO volume collected in the Douglas bag. To compare the results of the new methods with those of oCOR, thb-mass was calculated for min 7 (thb-mass_{min7}) as well as using data from the whole test, that is, the mean from the plateau between min 6 and 20 (thb-mass_{plateau}) and for min 20 using the data from the air collected in the Douglas bag (thb-mass_{DouglasBag}).

### 2.7 Statistical analysis

Statistical analysis was performed using IBM SPSS Statistics (version 25 for Windows, IBM Corp., Armonk, NY, USA). Values are presented as the mean ± standard deviation (SD), unless otherwise stated. Categorical variables are presented as frequencies (%).

This was a feasibility study, and a formal power calculation was therefore not required. Test–retest data (repeated measures from the same patient with different analytical methods) are presented using Bland–Altman plots with limits of agreement (Bland & Altman, 1986). Additionally, a specific approach to compute reliability statistics to compare test–retest performance expressed as the typical error of measurement (TE) was used (see Hopkins, 2000).

Student’s paired t-test was used to compare mean values from both tests at identical time points, and a paired t-test was also used to compare the mean values at different time points of the identical test. All tests were two-sided, and statistical significance was set at $P < 0.05$. To minimize the risk for type I errors, a correction for multiple measurements according to Benjamini & Hochberg (1995) was performed.

### 3 RESULTS

All of the tests were conducted without complications or adverse events. All participants inhaled the identical CO volume during the four tests ($63.3 \pm 23.1$ ml; males $82.1 \pm 16.8$ ml, females $44.5 \pm 6.3$ ml). The exhaled CO volume was highest in the first minute of Proc_{new}15s ($6.3 \pm 3.5$ ml; Proc_{new}30s $2.9 \pm 1.6$ ml). This initial phase was followed by an almost linear and parallel increase in both new procedures, showing an accumulation of $8.7 \pm 3.6$ and $5.1 \pm 2.0$ ml in min 7 and $12.0 \pm 4.4$ and $8.4 \pm 2.6$ ml in min 20, respectively. When the expired air was collected after oCOR$_{+20min}$, the values ($3.1 \pm 0.3$ and $5.9 \pm 1.1$ ml) were clearly below those of Proc$_{new}15s$ and Proc$_{new}30s$ (Figure 2) and not different from the volume exhaled 7 min after oCOR.

[COHb] exhibited well-known time-dependent changes, with a fast increase in the first minute followed by a rapid and then decelerating decrease during the rest of the observation period (Figure 3). The values of the new procedures were clearly below those of oCOR$_{+20min}$. In min 7, the CO volume ligated to Hb was $59.5 \pm 22.5$ ml (oCOR), $57.2 \pm 21.5$ ml (Proc$_{new}30s$) and $53.9 \pm 21.6$ ml (Proc$_{new}15s$),
FIGURE 2  Cumulative CO volume exhaled after the three different methods of CO application. n = 6; Procnew15s: single CO bolus inhalation with 15 s breath holding; Procnew30s: single CO bolus inhalation with 30 s breath holding; oCOR +20min: 2 min CO rebreathing followed by an 18 min analysis of exhaled air.

FIGURE 3  Changes in carboxy-haemoglobin ([COHb]) concentration after the inhalation of CO by three different application methods. n = 6; Procnew15s: single CO bolus inhalation with 15 s breath holding; Procnew30s: single CO bolus inhalation with 30 s breath holding; oCOR +20min: 2 min CO rebreathing followed by an 18 min analysis of exhaled air.

corresponding to 94.0 ± 2.1%, 90.4 ± 4.7% and 85.2 ± 6.0% of the inhaled CO volume, respectively.

tHb-mass was calculated for min 1 and 2 and then further measured in 2 min steps until min 20. We found increasing tHb-mass values for Procnew15s until min 6 (Figure 4a) and for Procnew30s until min 8 (Figure 4b) followed by a plateau after both procedures until min 20. Comparing the tHb-mass determined at min 7 yielded similar results for the three methods (Table 2). When tHb-mass from oCOR was compared with tHb-massplateau and tHb-massDouglasBag, we found slightly lower values for Procnew15s and very similar values for Procnew30s (Table 2). Comparison of tHb-mass values obtained with the established method (oCOR) and with oCOR +20min, that is, with collection of the expired air as in Procnews, yielded almost identical results (Table 2).

Bland–Altman plots comparing oCOR and the new procedures demonstrate slightly lower tHb-mass values for Procnew15s (tHb-massplateau: −21.8 ± 15.3 g; tHb-massDouglasBag: −12.2 ± 24.2 g) and very similar values for Procnew30s over a large range of tHb-mass values between 450 and 1300 g (Figure 5).

4 | DISCUSSION

We wished to identify a way to measure tHb-mass in mechanically ventilated patients. We thus explored whether it is possible to reliably measure tHb-mass using a single-breath inhalation (with a 15 or 30 s breath hold) of CO gas and showed for the first time that it is feasible (Procnew). Exhaled gas was collected and measured for the duration of the testing period (20 min). Our data suggest that Procnew with 15 and 30 s breath holds was closely related to the established CO-rebreathing method.

The principle of the CO-rebreathing method is to administer a defined amount of CO by breathing to determine the resulting COHb concentration in the completely mixed blood and to take into account the CO not bound to the Hb, that is CO exhaled and CO diffused to myoglobin. When these conditions are fulfilled, different procedures of the CO method can be applied.

Modifications to the CO rebreathing technique for the measurement of tHb-mass have therefore been made many times since the technique was revived by Fogh-Andersen et al. (1990), notably in 1995 when Burge and Skinner achieved improved precision of the measurement (Burge & Skinner, 1995). The current technique described by Schmidt and Prommer reduced the rebreathing period to only 2 min to improve convenience for participants (Otto et al., 2017a,b; Plumb et al., 2020; Schmidt & Prommer, 2005). The finding that a bolus of CO gas inhaled with a single breath and only rebreathed for 2 min led to valid and reliable results characterized by a typical error between 1% and 2% allowed the method to be used in a variety of different settings. Initially, these were primarily focused on elite sports physiology and performance, but more recently, oCOR has also been used to answer clinical questions (Otto et al., 2017a,b). The high reliability is confirmed in this study with a TE of 1.0% for the standard oCOR method.

When CO is administered for tHb-mass determination in an open spirometry system as we did in this study for the first time, an exact determination of the exhaled CO is mandatory. We determined the exhaled CO volume twice, that is, first by continuously monitoring the volume and CO concentration of the exhaled air, and second by collecting the whole amount of expired air in a Douglas bag and measuring the exhaled CO volume after the test.

To check whether the breathing procedure after the test exerts any unexpected influence on CO exhalation, we compared a 2-min inhalation period with subsequent collection of exhaled air
FIGURE 4  Time course of the calculated tHb-mass after CO breathing and subsequent breath holding for 15 s (a; Procnew15s) and for 30 s (b; Procnew30s). Presented are mean values and individual data of the tHb-mass calculated for different time points of blood sampling. Significant differences from previous values: *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001

TABLE 2  tHb-mass calculated from the three CO application methods

|                  | oCOR(n = 11) | Procnew15s(n = 11) | Procnew30s(n = 11) | oCOR(n = 6) | oCOR+20min(n = 6) |
|------------------|--------------|--------------------|--------------------|-------------|-------------------|
| tHb-massmin7 (g) | 843 ± 293    | 819 ± 285          | 838 ± 301          | 705 ± 209   | 665 ± 194         |
| tHb-massplateau (g) | 821 ± 288*  | 849 ± 311          | 686 ± 200          |             |                   |
| tHb-massDouglasBag (g) | 830 ± 276   | 846 ± 309          | 711 ± 196          |             |                   |

oCOR: established CO-rebreathing method; Procnew15s: single CO bolus inhalation with 15 s breath holding; Procnew30s: single CO bolus inhalation with 30 s breath holding; oCOR+20min: 2 min CO rebreathing followed by an 18 min analysis of exhaled air; tHb-massmin7: tHb-mass calculated with [COHb] determined 7 min after inhalation; tHb-massplateau: Hb-mass calculated with [COHb] determined between min 6 and min 20; tHb-massDouglasBag: tHb-mass calculated with [COHb] determined at min 20 and using the CO volume exhaled into the Douglas bag. Significant difference from oCOR: *P < 0.05.

When breath was held for 15 s, the initial CO exhalation after 1 min was twice as high as that in Procnew30s, indicating that ~10% (Procnew30s ~4%) of the inhaled CO did not diffuse into the blood. After 7 min, that is, when the CO mixing in the blood was completed and therefore used in the established oCOR for blood sampling after the test, the loss of CO was ~13% in Procnew15s and only ~8% in Procnew30s. Although this loss in CO clearly exceeded the CO volume exhaled after oCOR (~4%), these data demonstrate the rapid diffusion of CO from the lungs into the blood, which is also a precondition for the determination of the lung diffusion capacity by means of a deep inhalation of a 0.3% CO-containing gas followed by a 10-s breath hold (Modi & Cascella, 2020). We therefore suggest that tHb-mass might be easily and exactly calculated after a single breath when considering sufficient mixing time of the inhaled CO bolus.

These considerations are supported by the calculated tHb-mass over time. In the new procedures, tHb-mass reached a plateau in min 6 or in min 8 indicating complete mixing (Bruce & Bruce, 2003), that is, that any time point beyond can be used for tHb-mass and blood volume determination (Wachsmuth et al., 2019). In this study, we used the plateau value between min 6 and 20 and compared its mean with the time point usually used in the oCOR (min 7) and with the results obtained in min 20 from the exhaled air collected in the Douglas bag. As shown in Table 2, there is no obvious difference between tHb-mass obtained from the oCOR and that obtained from the new procedures at the time points mentioned above. In the Bland–Altman plot, a small but systematic underestimation by approximately 25 g compared with the reference method becomes obvious in Procnew15s. As such a deviation does not occur between Procnew30s and oCOR, we suggest that the smaller CO volume taken up from the blood during Procnew15s may be the cause. Additionally, the very low limits of agreement in the comparison of both new methods with oCOR indicate that Procnew15s already presents a promising tool for tHb-mass and blood volume determination, and Procnew30s seems to be as exact as the established oCOR.

In future studies, the administration of higher CO volumes than those used for the oCOR may be taken into consideration to compensate for the lower CO uptake during the single-breath application. On the one hand, this procedure increases the [COHb], reducing the measurement error of the CO oximeter (Alexander et al., 2011) but also increases the volume of exhaled CO and thereby
introduces another source of inaccuracy. In the literature, there has also been extensive debate about the merits of having a higher $\Delta$COHb% versus the increased toxicity risk (Alexander et al., 2011; Garvican et al., 2010; Turner et al., 2014). COHb of up to 10% has been described without remarkable side effects in healthy subjects (Schmidt et al., 2020), but to our knowledge, it has never been studied in seriously ill patients. Because CO is endogenously produced and is actually considered for the treatment of various diseases (Motterlini & Otterbein, 2010), we are convinced that the increase in COHb by 4–5%, as achieved in our study, represents a reliable compromise balancing sources of error with minimal patient risk.

4.1 | Practical application

We hypothesize that this method might be used to diagnose and provide more information on the origin of anaemia in intensive care (Magee & Zbrozek, 2013) and the amount of blood loss during surgery (Shoemaker et al., 1996), as well as for distinguishing between dilutional anaemia and genuine anaemia in patients with heart failure (Miller & Mullan, 2015) and liver failure (Plumb et al., 2020). Here, it is of critical importance that the modified method has sufficient accuracy to reveal clinically relevant changes in tHb-mass and their contribution to changes in Hb. Although the reliability of the modified method was not explicitly determined in this feasibility study, the methodological error (typical error, TE; Hopkins, 2000) compared to the established method is 3.2% (Procnew30s) or 3.5% (Procnew15s). This is higher than the TE of the established CO rebreathing methods (TE 2.2%) but close to the TE of the gold standard methods using radio- active markers ($^{51}$Cr, 2.8%; Gore et al., 2005). Since even mild real anaemic states ([Hb] 11.2 g/dl) are associated with a reduction of at least 15% tHb-mass (Wachsmuth et al, 2015), the accuracy of the method should be sufficient to distinguish anaemia due to reduced tHb-mass from that due to dilution. This contention is supported by Otto et al. (2017a) who describe two patients with liver disease who had identical tHb-mass (9.2 g/kg body mass), but with a normal [Hb] (16.1 g/dl) in one and dilutional anaemia ([Hb] 10.7 g/dl) in the other. They also describe two cases of heart failure in which the presence of a severe reduction in tHb-mass (5.2 g/kg) was reflected in a low [Hb] in one (6.9 g/dl), but masked by a contracted PV in another ([Hb] 10.7 g/dl) (Allsop et al., 1998; Otto et al. 2017a). In addition to the frequently occurring dilution anaemia, decompensated heart failure can also be associated with proportional increases in both tHb-mass and PV (Miller, 2016). In all these cases, the determination of the tHb-mass, also when using the modified method, enables a much more precise diagnosis.

The modification described here may permit the CO method to be used in ventilated patients. CO is applied to the inhalation path, and breathing is interrupted in the inhalation position for 15 or 30 s. The CO volume not absorbed by the patient can be determined either by collecting the entire expiratory air for a period of approximately 20 min or by continuous monitoring of the volume of the expired air and its CO concentration. These measurements are carried out until the CO is completely mixed in the blood and a blood sample for the determination of COHb is drawn.

The application of our approach to ventilated patients would offer possible clinical advantages. In most patients treated in an intensive care unit, the [Hb] drops significantly within a few days and transfusions are recommended when [Hb] reaches a threshold of 70 g/l (Watson & Kendrick, 2014) without excluding dilution. Indeed, there is no routine way in which to assess intravascular volume, with central venous pressure being a very poor guide indeed (De Backer & Vincent, 2018). In addition, blood and fluid loss during surgery are imprecisely measured and this, together with altered cardiovascular tone, and the variable administration of packed red blood cells and crystalloid/colloid solutions, makes determination of
intravascular volume (and of true Hb deficit) difficult. Our method may find application in all such situations. Nonetheless, the applicability and validity of this ‘single breath method’ remains to be validated in clinical circumstances.

In such clinical studies CO mixing time must be considered. It is prolonged in patients with polycythaemia (Wachsmuth et al., 2019) and heart failure (Ahlgrim et al., 2018), and perhaps also in other patient groups. This should, however, not be a major problem as the exhaled CO is collected for 20 min and a significant increase in mixing time can be tolerated if the individual COHb plateau is determined for each patient after inhalation of the CO. The use of the new method in patients with pulmonary diffusion disorders could be more problematic if sufficient CO cannot diffuse from the alveoli into the blood within 30 s. Higher CO doses may have to be used in such circumstances, but this can have an adverse effect on the accuracy of the test.

In healthy subjects, there is no risk of interrupting breathing for 30 s, and the risk can be classified as very low also in ventilated patients. Since the oxygen consumption during the 30-s breath interruption is only approximately 150 ml, the arterial O2 saturation does not change during this period (Parkes et al., 2016); but in any case, it must be checked during and after the test. In severely anaemic patients, the test might be used with great caution after extensive validation.

5 CONCLUSION

Using the single-breath method, tHb-mass and blood volume can be determined with approximately the same accuracy as that with established CO-rebreathing methods. We recommend that this method be developed further for use in ventilated patients, that is, patients in intensive care, patients undergoing major surgery, and patients with heart and liver failure.

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COMPETING INTERESTS

W.F.J.S. is a managing partner of the company ‘Blood tec GmbH’, but he is unaware of any direct or indirect conflict of interest with the contents of this paper. The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

AUTHOR CONTRIBUTIONS

L.M.K., J.O.M.P., H.E.M., N.B.W. and W.F.J.S. were involved in the conception and design of the study, the acquisition of data, the analysis and interpretation of the data, and the drafting of the manuscript. S.H., J.S., S.B.K., J.M.O. and M.P.W.G. were involved in the acquisition of data, analysis and interpretation of data for the work, as well as in the critical revision of the manuscript. All authors approved the final version of the manuscript and agreed to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All persons designated as authors qualify for authorship, and all those who qualify for authorship are listed.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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