Effect of variation in the COPD breathing flow pattern on end-tidal CO2 tension: An in vitro study

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Abstract

Aim: The goal of this work was to find out if the variation in breathing flow pattern alone (without change in tidal volume and respiratory rate) changes the end-tidal CO₂ tension (EtCO₂) enough to affect the health of a patient. The influence of I:E ratio on EtCO₂ was also investigated.

Method: Four breathing flow patterns belonging to different individuals diagnosed with COPD were collected from the literature. These were scaled to the same tidal volume, respiratory rate and I:E ratio, leaving shape as the only varying parameter. A programmable piston pump was used to reproduce the breathing flow patterns in an anatomically correct adult upper airway model (3D printed in acrylic, Visijet Ex200). CO₂ was simultaneously bled into the pump chamber at steady rate to mimic metabolic CO₂ production, followed by the measurement of EtCO₂ with and without 30 L/min of nasal high flow therapy.

Results: Breathing flow patterns varying in shape but not in tidal volume, respiratory rate and I:E ratio can produce statistically significant differences in EtCO₂. The variability in EtCO₂ was however found to be small (1 - 2%) and unlikely to be physiologically relevant. A 35% fall in I:E ratio corresponded to a 2% rise in EtCO₂.

Conclusion: Shape alone can cause a statistically significant difference in EtCO₂ however the difference in EtCO₂ is small. A 35% reduction in I:E ratio results in a 2% rise in EtCO₂.

Introduction

Arterial CO₂ tension modulates ventilation via a feedback mechanism. It is thus reasonable to hypothesise that each breathing pattern (though influenced by disease state) is optimized in terms of shape, amplitude and frequency to effect a specific change in CO₂ tension. Previous studies have explored how tidal volume, dead space volume, respiratory frequency and metabolic CO₂ rate influence EtCO₂. In a study by Parot, et al. [1] the respiratory frequency and metabolic CO₂ production in hypercapnic and non-hypercapnic COPD patients were found to be similar however the hypercapnic group showed a lower tidal volume. In hypercapnic individuals, the EtCO₂ correlates well with the ratio of dead space volume to tidal volume [2,3]. EtCO₂ has been reported to correlate well with arterial CO₂ Tension [4].

Critically ill patients suffering from COPD benefit from nasal high flow (NHF) therapy, which is the administration of warmed and humidified air at flow rates up to 8 L/min in neonates [5-7] and 60 L/min in adults [8]. NHF reduces the physiological dead space and respiratory frequency, and improves gas exchange [9-12].

The capnogram of a COPD patient is modulated by the degree of obstruction of the airways [13-15]. This obstruction, which affects the breathing flow pattern and magnitude, causes the capnogram of COPD patients to have a 'shark fin shape', [16] making them distinct from those of healthy individuals, which are more rectangular in shape [17]. Clinical studies on how a variation in the breathing flow pattern alone change EtCO₂ are scarce due to the difficulty in simultaneously maintaining the same tidal volume, peak airflow and I:E ratio over several breaths. In this report, an in-vitro study of the effect of breathing flow pattern, with and without NHF, on EtCO₂ is presented. Also the influence of I:E ratio variation on EtCO₂ is explored.

Methodology

Waveform collection

Four breathing flow patterns obtained from COPD patients (age range = 17 - 77 years, FEV₁% pred = 36 - 63%) were collected from the literature. In three of these flow patterns [18-20], flow rate was plotted against time. In the remainder [21], the flow pattern was presented as a plot of tidal volume versus time which was numerically integrated to yield flow rate versus time plot after acquisition. Together with a healthy adult breathing flow waveform, which was previously used by Van Hove, et al. [22] and Spence, et al. [23] five waveforms were used in all.

A plot of all the waveforms collected is shown in Figure 1a. Note that positive flow represent inspiration and negative flow, expiration. The four COPD waveforms are designated as WF1, WF2, WF3 and WF4 with the healthy waveform labelled as WF5 (unmarked solid line). The COPD flow waveforms have a sharp concavity in the expiratory phase. It is noticeable that they vary in frequency and amplitude.

Normalization of waveforms

All the waveforms were normalized to the same tidal volume and respiratory rate via a linear rescaling of time axis followed by a

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Key words: end-tidal CO2 tension, nasal high flow therapy

Received: October 01, 2017; Accepted: October 17, 2017; Published: October 19, 2017
linear rescaling of the flow rate axis using different scaling factors for inspiratory and expiratory phases. Also, the inspired volume was matched to the expired volume. Two groups of breathing flow waveforms were obtained by scaling each breathing flow to two different I:E ratios i.e. 0.67 (IE1-group) and 0.43 (IE2-group), which corresponded to inspiratory time fractions of 40% and 30%. The fall in I:E ratio from 0.67 to 0.43 equals 35%. This change of 35% in I:E ratio was hypothesized to be sufficient to produce a significant change in EtCO₂. The breathing flow patterns

in the IE1-group and IE2-group are shown in Figures 1b and 1c respectively. Two different I:E ratios were needed to investigate the effect of I:E ratio on EtCO₂. The choice of I:E ratios were not without justification as Tobin, et al. [24] reported an I:E ratio of 0.73 ± 0.03 for healthy adults and 0.53 ± 0.05 for adult COPD patients.

Experimental setup

The experimental set-up is shown in Figure 2b. In the setup, a rigid upper airway model (3D-printed in acrylic, Visijet EX200) - Figure 2a - of an anonymous 44 year male adult (previously used by Spence, et al. [25] and Van Hove, et al. [22]) was connected to a piston pump. A CO₂ source was connected to the barrel of the piston pump via a rotameter. CO₂ was sampled at the trachea using a 20 Hz capnograph (MiniMediCO₂, manufactured by Oridion Medical Ltd., Israel). A mixing test conducted by placing a wire-gauze (to enhance mixing) at the entrance of the piston pump yielded no significant difference in EtCO₂ (< ± 3%). The volume of the piston chamber at the end of expiration (functional residual capacity) was maintained at 2500 ml for all experiments.

Experimental procedure

The pump forces the tracheal flow rate to follow the healthy waveform (WF5) in Figure 1b and Figure 1c. Metabolic CO₂ production was simulated by bleeding CO₂ at a flow rate of 98 scm/min (at a 20 KPa CO₂ source pressure) into the barrel of the piston pump. The end-tidal (EtCO₂) at this setting was 5.4% ± 0.3, which is comparable to the EtCO₂ of a resting healthy adult. CO₂ data was recorded before (zero-therapy, ZT) and during the application of 30 L/min of NHF (NHF30). An identical procedure was performed for all other waveforms with 6 repetitions for each.

Results

The capnogram was recorded for 20 consecutive breathing cycles for each experimental repeat. The 20 cycles of capnogram in repeat one (for healthy waveform (WF5) in IE1-group) have been superimposed and cascaded with those of the other repetitions as shown in Figure 3a. The variations in EtCO₂ from cycle to cycle is due to mixing. Figure 3b shows the average of all 120 capnograms (20 cycles in each of the 6 repeats) for WF5 (in IE1-group) with and without NHF. An identical plot is shown in Figure 3c for WF4 (a COPD waveform). Note the characteristic ‘shark fin’ shape of the COPD waveform (which is due to airway obstruction) [16].

The spontaneous breathing plots (for a no NHF condition) in Figure 3b and Figure 3c show a minimum CO₂ concentration of about 0.2% which is due to rebreathing of dead space CO₂ during inspiration. Note how this falls to 0.04% (atmospheric CO₂ concentration) when NHF of 30 L/min is applied. This indicates a reduction in re-inspired dead space CO₂ by NHF, which is in line with findings by Spence, et al. [23] and Van Hove, et al. [22] that NHF promotes mixing and flushes dead space CO₂. In Figure 3b and Figure 3c the EtCO₂ falls by approximately 20% when NHF of 30 L/min is applied.

The average EtCO₂ (of 120 capnograms) for each waveform (in both IE1-group and IE2-group) have been presented in Figure 4. The error bars represent two standard deviations in EtCO₂. In Figure 4, the EtCO₂ vary slightly between waveforms (e.g. see IE1-group waveforms) however the error bars overlap. Also pairs of plots belonging to the same waveform show some difference in EtCO₂ though error bars overlap (e.g. for label WF4 in Figure 4, compare the IE1-group and IE2-group pair).

The present finding shows that the characteristic shape of the COPD capnogram (‘shark fin’) - due to airway obstruction - can be
reproduced from the corresponding breathing flow pattern. This supports the clinical reports that the breathing flow pattern modifies the shape of the capnogram [16]. Though error bar overlaps in EtCO₂ have been observed between and within the I:E ratio groups (IE-1 and IE-2) it is not conclusive if this means no statistically significant difference in EtCO₂ exists. In what follows a test of statistical difference in EtCO₂ is performed.

Test of statistical significance

A single factor ANOVA test was performed to find if the difference in EtCO₂ was statistically significant between different waveforms of the same I:E ratio. Further, a two sample t-test (assuming unequal variances) on pairs of EtCO₂ belonging to the same waveform but differing in I:E ratio was performed. In both tests, the critical value to confirm the null hypothesis was set to 0.02 instead of 0.05 to indicate a strong evidence against the null hypothesis. The results are presented in Table I, which shows evidence of statistically significant difference in EtCO₂ (p-value < 0.02) within the same group of I:E ratio (single factor ANOVA test, Table 1). Except for WF4, I:E ratio made a statistically significant difference in EtCO₂ (p-value < 0.02) in pairs belonging to the same I:E ratio (two-sample t-test, Table I). WF4 was not distinct in characteristics from the other waveforms.

Another two-sample t-test was performed for each I:E ratio group to find the specific pairings of flow waveforms that showed statistically significant difference in EtCO₂ as the single factor ANOVA test could not determine this. Note that 5 flow waveforms will yield 10 pairs. The results are presented in Table 2 in which 7 pairs (70%) - in bold ink – show statistically significant difference in EtCO₂ (p-value < 0.02) within the IE1-group. Only 4 pairs (40%) in the IE2 group have statistically significant differences in EtCO₂.

It is concluded that the differences in EtCO₂ of waveforms that are similar in all but pattern are statistically significant (p-value < 0.02) (a single factor ANOVA test, Table I). Also, EtCO₂ is sensitive to I:E ratio as 4 out of 5 flow waveforms (WF1, WF2, WF3 and WF5 but not WF4) showed a statistically significant difference in EtCO₂ (p-value < 0.02) (two sample t-test, Table 1). Furthermore, it is deduced from Table 2 that though statistically significant disparities in EtCO₂ was found in each group (Table 1), intragroup differences in EtCO₂ in the higher I:E ratio group (IE1-group, I:E ratio = 0.67) were more frequent (70%) than in the lower I:E ratio group (IE2-group, I:E ration = 0.47) - only 40% of the pairs showed a significant difference in EtCO₂.

Figure 2. (a) Upper airway model (b) UAM is connected to the piston pump and AIRVO™2 device. CO₂ source is connected to piston pump for simulation of metabolic CO₂ production. CO₂ is sampled at the trachea during experiment.

Figure 3. (a) A train of 6 capnograms for the healthy waveform (WF5) (b) Average of 120 capnograms for WF5 (c) Average of 120 capnograms for WF4.

Figure 4. A plot of the average EtCO₂ of 120 capnograms associated with flow waveforms in both groups (IE1-group and IE2-group). The errorbars represent two standard deviations in EtCO₂ over the 120 capnograms.
Shape parameter and EtCO$_2$

Harmonic distortion ($Hd$) describes the degree to which a periodic signal deviates from a sinusoid [26]. It is mathematically expressed as:

$$Hd = \sqrt{H_1^2 + H_2^2 + H_3^2 + ...}$$

(1)

where $H_n$ is the root-mean-square amplitude of the $n$th harmonic (found using Fourier decomposition) of the waveform. $n = 25$ was used as beyond this, changes in $Hd$ was found to be insignificant. $Hd$ was computed for the entire breathing cycle, for the inspiratory phase and then the expiratory phase. EtCO$_2$ correlated better with a product of the inspiratory $Hd$ and the peak inspiratory flow. This product is designated as $IDf$. The EtCO$_2$ and their corresponding $IDf$ are presented in Figure 5. The Spearman’s correlation coefficient ($\rho$) and error (2 standard deviations) in the measurement of EtCO$_2$ are also shown in Figure 5. The coefficient of determination ($R^2$) (0.75 and 0.88) and $\rho$ (0.9 and 1) indicate a strong correlation between IDf and EtCO$_2$ (Figure 5). It is not clear what implication this has for breathing.

In Equation 2, $\text{average}[EtCO2]_{IE2}$ and $\text{average}[EtCO2]_{IE1}$ are respectively, the average EtCO$_2$ in the IE1-group and IE2-group. Using the formula in Equation 2 the average EtCO$_2$ of the IE2-group was found to be 2% greater than that of the IE1-group. It is concluded that a 35% fall in I:E ratio (percentage difference between the I:E ratio of 0.67 (IE1-group) and 0.43 (IE2-group)) produces a 2% rise in EtCO$_2$. The coefficient of variability, defined as the ratio of the standard deviation in EtCO$_2$ within the group to the average EtCO$_2$ was found to be 2% and 1% respectively for the IE1-group and IE2-group.

Discussion

NHF reduces physiological dead space, respiratory frequency, and improves gas exchange [9-11]. The fall in EtCO$_2$ (= 20%) upon the application of 30 L/min of NHF supports the report that NHF improves mixing and reduces the proportion of CO$_2$ in re-inspired dead space air [9,22,23]. The present data suggests that breathing flow waveforms of different patterns, but similar in tidal volume, period and I:E ratio show statistically significant differences in EtCO$_2$. Further, the results indicate that two breathing waveforms that differ only in I:E ratio can produce different EtCO$_2$. If that tidal volume and respiratory rate are fixed, it is more probable to find a statistically significant difference in EtCO$_2$ between a greater I:E ratio group than those of a lower I:E ratio.

To preserve tidal volume, a fall in I:E ratio (longer expiration, shorter inspiration) is matched by a rise in peak inspiratory breathing flow and a fall in peak expiratory breathing flow. In Figure 4, the EtCO$_2$ corresponding to the lower I:E ratio i.e. 0.46 (IE2-group) tended to be greater. Tobin, et al. [24] observed that the inspiratory time (Ti) of 28 COPD subjects (mean age = 67.5 years) was 1 second less than that of healthy adults. Sorli, et al. [27] concluded that CO$_2$ retention (marked by higher EtCO$_2$) in COPD patients is due to shallow breathing, which arises from reduced Ti. This suggests the higher EtCO$_2$ is due to less efficient purging of the dead space during inspiration, i.e. lower ratio of inspired volume to dead space volume resulting in less complete replacement of dead space gas with fresh air (residual volume of expired gas mixes with a smaller volume of fresh gas). This is in unison with the observation by Gorini, et al. [28] who found the highest arterial partial pressure of CO$_2$ amongst COPD patients with the smallest Ti. IE1-group and IE2-group differed in Ti by 0.43 seconds. In the present data, a fall in I:E ratio by 35% (0.67 to 0.43) corresponded to 2% rise in EtCO$_2$. Though the present results agrees qualitatively with findings in the literature [24,27,28] it is concluded that the fall in EtCO$_2$ due to a fall in I:E ratio by 35% is small (2%). The variability in EtCO$_2$ due to flow pattern difference is 1 - 2%. Though statistically significant, it is also small.

The product of inspiratory flow and inspiratory harmonic distortion (IDf) correlate well with EtCO$_2$. The expiratory breathing flow has been reported to be modulated by the inspiratory flow pattern [26]. Inspiratory muscle dysfunction increases EtCO$_2$. Begin, et al. [29] found that airway resistance correlated well with arterial CO$_2$ tension, which also correlates well with EtCO$_2$ [30]. In a study by Loveridge, et al. [31] healthy subjects were observed to show a greater variability in breathing pattern than COPD subjects. Since airway obstruction during COPD influences EtCO$_2$ and the flow pattern, it is speculated that $IDf$ is related to the ventilatory response to CO$_2$ tension.

Limitations of this work

The piston chamber is rigid, unlike the human lung which is compliant (lung compliance is $\approx 200$ mL/cmH$_2$O for healthy young adults). Note that a constant bleed rate differs from the physiologically realistic system in which CO$_2$ flow is distributed over time. Also, this work is limited by the number of breathing flow patterns used.
Conclusion

Breathing patterns similar in tidal volume, respiratory rate, and I:E ratio but differing in shape can produce statistically significant difference in EtCO₂, however the variability is small (1 - 2%). A 35% fall in I:E ratio of breathing waveforms with no change in tidal volume and respiratory rate produces only a 2% rise in EtCO₂. NHF of 30 L/min reduces EtCO₂ by 20%.

Acknowledgment

The authors would like to thank the University of Canterbury for a Doctoral Scholarship, Fisher & Paykel Healthcare for the loan of equipment and advice, and MBIE Smart Ideas grant UOA1403.

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