Comparing the novel microstream and the traditional mainstream method of end-tidal CO$_2$ monitoring with respect to PaCO$_2$ as gold standard in intubated critically ill children

Muhterem Duyu$^{1,2,*}$, Anıl Dogan Bektas$^2$, Zeynep Karakaya$^2$, Meral Bahar$^2$, Aybuke Gunalp$^2$, Yasemin Mocan Caglar$^2$, Meryem Nihal Yersel$^1$ & Özlem Bozkurt$^1$

The objective of this study was to evaluate a novel microstream method by comparison with PaCO$_2$ and the more standard mainstream capnometer in intubated pediatric patients. We hypothesized that the novel microstream method would superior compared to the traditional mainstream method in predicting PaCO$_2$. This was a prospective single-center comparative study. The study was carried out on 174 subjects with a total of 1338 values for each method. Data were collected prospectively from mainstream and microstream capnometer simultaneously and compared with PaCO$_2$ results. Although both mainstream PetCO$_2$ (mainPetCO$_2$) and microstream PetCO$_2$ (microPetCO$_2$) were moderately correlated ($r = 0.63$ and $r = 0.68$, respectively) with PaCO$_2$ values, mainPetCO$_2$ was in better agreement with PaCO$_2$ in all subjects (bias ± precision values of 3.8 ± 8.9 and 7.3 ± 8.2 mmHg, respectively). In those with severe pulmonary disease, the mainPetCO$_2$ and microPetCO$_2$ methods were highly correlated with PaCO$_2$ ($r = 0.80$ and $r = 0.81$, respectively); however, the biases of both methods increased (14.8 ± 9.1 mmHg and 16.2 ± 9.0 mmHg, respectively). In cases with increased physiologic dead space ventilation, the agreement levels of mainPetCO$_2$ and microPetCO$_2$ methods became distorted (bias ± precision values of 20.9 ± 11.2 and 25.0 ± 11.8 mmHg, respectively) even though mainPetCO$_2$ and microPetCO$_2$ were highly correlated ($r = 0.78$ and $r = 0.78$, respectively). It was found that the novel microstream capnometer method for PetCO$_2$ measurements provided no superiority to the traditional mainstream method. Both capnometer methods may be useful in predicting the trend of PaCO$_2$ due to significant correlations with the gold standard measurement in cases with severe pulmonary disease or increased physiological dead space—despite reduced accuracy.

Abbreviations

CO$_2$ Carbon dioxide
ABG Arterial blood gas
PICU Pediatric intensive care unit
MAP Mean airway pressure
OI Oxygenation index
SD Standard deviation
IQR Interquartile range
PASS Power analysis sample size

$^1$Department of Pediatrics, Pediatric Intensive Care Unit, Istanbul Medeniyet University Goztepe Training and Research Hospital, Istanbul, Turkey. $^2$Department of Pediatrics, Istanbul Medeniyet University Goztepe Training and Research Hospital, Istanbul, Turkey. *Email: drmuhteremduyu@gmail.com
The monitoring of carbon dioxide (CO₂) level is essential for diagnosis and therapeutic guidance in mechanically ventilated patients. The current gold standard method for the measurement of partial pressure of carbon dioxide in the blood (PaCO₂) is the arterial blood gas (ABG) method. But ABG is an invasive method and does not provide continuous monitoring.

Capnometers, which continuously monitor PCO₂ levels and display the waveform of PCO₂ in exhaled air non-invasively, provide information on the adequacy of ventilation. Detection of exhaled CO₂ (end-tidal CO₂) has proven to be a valuable mechanism to confirm tracheal intubation and to recognize accidental esophageal intubations, among other critical patient safety benefits. The patient protection enhancements provided by end-tidal PCO₂ (PetCO₂) monitoring also include the detection of invasive airway disconnection, dislodgement or obstruction, prediction of underlying airway or lung pathologies and monitoring the effectiveness of cardiopulmonary resuscitation.

It is possible to measure PetCO₂ by mainstream or sidestream capnometer technologies. The technique is named based on the localization of the PetCO₂ sensor. Mainstream capnometers are devices in which the infrared source and PCO₂ detector are placed between the proximal endotracheal tube (ETT) and the ventilator circuit. On the other hand, sidestream capnometers aspirate samples from the airway through tubing attached to a sampling line and airway adapter between the ETT and the ventilation circuit. Sidestream methods utilize an infrared PCO₂ sensor in a monitor that may be located far from the patients. A new technology for sidestream systems (Microstream, Oridion Medical, Inc., Danville, CA) is now available that uses very low flow rates (50 mL/min) to preserve the accuracy and resolution of the waveform as well as eliminating secretion/moisture-related occlusion problems by the use of special filters.

There are many studies evaluating the accuracy of mainstream, sidestream and microstream capnometer technologies in the literature. Critics of capnometer usage cite multiple studies which demonstrate that PetCO₂ and PaCO₂ do not reliably correlate in some clinical situations. The analyses utilized in these studies are highly variable and fail to consider physiologic dead space/severity of pulmonary disease and/or their effect on the relationship between PaCO₂ and PetCO₂. Additionally, there are few studies that have compared different PetCO₂ monitoring techniques with each other.

This study was undertaken to evaluate the correlations of gold standard PaCO₂ measurements with the microstream and mainstream PetCO₂ capnometers, and to compare the accuracy and results of the latter two methods among ventilated patients in the pediatric intensive care unit (PICU). We hypothesize that, (i) in intubated pediatric patients, the microstream technology will allow better prediction of PaCO₂ compared to the traditional mainstream method, and that (ii) microstream measurements predict PaCO₂ more reliably than mainstream measurements across increased levels of dead space ventilation and in the presence of severe pulmonary disease—after controlling for the expected PetCO₂-PaCO₂ gradient.

Methods

This prospective, single-center comparative study was conducted at the PICU of Medeniyet University, Goztepe Training and Research Hospital (Istanbul, Turkey) between January 2018 and July 2019. All procedures and processes were carried out according to principles mentioned in the Helsinki Declaration and the Good Clinical Practice guideline.

Population. The study evaluated all children aged between 1 month to 17 years that had been intubated with cuffed ETT due to a definite indication for mechanical ventilation. Among these, those who accepted invasive monitoring of arterial blood pressure and provided informed consent (parents or legal guardians) were included in the study. The presence of any one of the following characteristics was defined as grounds for exclusion from the study: patients with tracheostomy, sampling performed with venous blood, non-compliance to study protocols, or obstruction, prediction of underlying airway or lung pathologies and monitoring the effectiveness of cardiopulmonary resuscitation.

Monitoring. The intubations were performed with single-lumen cuffed ETT that was appropriately sized for age and weight. CO₂ in the exhaled air of patients was monitored simultaneously with mainstream (Mainstream EtCO₂; Philips Capnostream M2501A, Germany) and microstream (Microstream EtCO₂; Medtronic Capnostream35, USA) capnometers. The dimensions of the airway adapters to be used were based on the manufacturer’s guidelines. The airway adapters of both methods of measurements were kept in the same location and insertions were performed sequentially between the airway circuit and the proximal ETT. ABG were analyzed at the bedside using an ABL 90 FLEX blood gas analyzer (Radiometer, Medical ApS, Copenhagen, Denmark) within 3 min of collection and without any delay. No additional ABG was performed for the data collection of consecutive samples.

Study protocol and recording. ABG analysis, mainPetCO₂ and microPetCO₂ values and mechanical ventilator parameters were recorded simultaneously. Prior to obtaining each arterial blood gas sample, a researcher checked whether the capnometer adapters were blocked by secretions or moisture. Capnometer adapters were replaced with new ones in the event of any type of blockage. Both capnometer methods were analyzed using continuous steady waveforms of expired CO₂ through the ventilator cycle, in order to ensure the accuracy of readings. A minimum of 4 and a maximum of 8 simultaneous PCO₂ measurements (PaCO₂, mainPetCO₂, microPetCO₂) were planned to be taken from each patient. Patients with a measurement number less than 4 for various reasons (death, extubation, interruption of monitoring etc.) were excluded from the study.
Patients’ demographic characteristics and their clinical and laboratory parameters were identified (gender, age [months] and primary diagnosis). The parameters of mechanical ventilation, including FiO₂ (Fractional inspired oxygen), mean airway pressure (MAP) were recorded in addition to PetCO₂ values (mainPetCO₂, and microPetCO₂), parameters of arterial blood gas analysis (pH, PaCO₂, PaO₂, HCO₃⁻) and oxygenation index (OI) (OI = [FiO₂ × MAP × 100]/PaO₂)²². Lung physiologic dead space volume is defined as wasted tidal volume during respiration (i.e., the volume remaining in the conducting airways [anatomical dead space] and in poorly perfused and non-perfused alveoli [alveolar dead space] that are not participating in gas exchange). A ratio of dead space volume to tidal volume (Vd/Vt) was calculated using the Enghoff modification of the Bohr equation: Vd/Vt = [PaCO₂ - PetCO₂]/PaCO₂²³. Dead space ventilation was calculated separately using PetCO₂ values obtained from microstream and mainstream methods.

For subgroup analyses, patients were grouped with regard to the severity of pulmonary disease and physiological dead space ventilation levels. Severe pulmonary disease was defined as an OI of ≥10 and mild-to-moderate pulmonary disease was defined as an OI of <10²⁴. Determination of Vd/Vt ratio ≥0.4 was defined as increased dead space ventilation²³,²⁵. The consistency of PetCO₂ monitoring (mainPetCO₂ and microPetCO₂) within each patient and subgroup was assessed by examining the relationships between the changes in PaCO₂ and the two PetCO₂ methods in consecutive samples.

Statistical analysis. Analyses were performed by using the IBM Statistical Package for the Social Sciences version 21 (SPSS, Inc., Chicago, IL) or Med Calc version 19.1 (Med Calc Software, Ostend, Belgium). Patient characteristics are described using qualitative variables (using frequencies and percentages) and quantitative variables (using means and standard deviation [SD] or median with interquartile range [IQR], depending on type of distribution). Simple linear regression analysis was performed and Spearman correlation coefficients were calculated for the assessment of relationships between PaCO₂, main-PetCO₂ and micro-PetCO₂. We assessed the agreement between these measurements (bias [mean difference] and precision [SD of the differences]) by the Bland–Altman technique. The results were considered statistically significant in tests resulting in a P value lower than 0.05.

Power analysis. Power analysis was conducted using the Power Analysis Sample Size (PASS) for Windows version 11.0 Package Program. Group sample sizes of 174 were determined to achieve 97% power to detect a difference of 3.6 between the null hypothesis that both group means were 3.8, and the alternative hypothesis that the mean of group 2 was 7.4 (with estimated group standard deviations of 9.0 and 8.3), and with a significance level (alpha) of 0.05 using a two-sided two-sample t-test.

Ethics approval and consent to participate. The study was approved by the institutional review board of our center. The parents of all patients signed an informed consent form before inclusion into the study.

Conference presentation. This study presented at the 15th National Congress of Pediatric Emergency and Critical Care, 18–20 October 2018, Turkey.

Results

The study was performed in 174 patients that provided 1338 measurements for each method. The median age and interquartile range (IQR) of the included subjects was 42 months (IQR: 12–108 mo.). Table 1 shows the characteristics of the study group. Conventional invasive mechanical ventilator modes were used in all patients included in the study (Galileo Mechanical Ventilator, Hamilton Medical AG, Rhäzüns, Switzerland).

The median (range) levels of PaCO₂, mainPetCO₂, and microPetCO₂ were 40.7 (IQR: 35.4–47.3) mm Hg, 38.0 (IQR: 32.0–44.0) mm Hg, and 35.0 (IQR: 29.0–40.0) mm Hg, respectively. The results of the Bland–Altman analysis comparing mainPetCO₂/PaCO₂ and microPetCO₂/PaCO₂ pairs are summarized in Table 2 and illustrated in Fig. 1 in all subject groups—and also according to severity of pulmonary disease. In all subjects (1338 pairs), the mean difference (bias) and SD of the differences (precision) for mainPetCO₂ was 3.8 ± 8.9 mm Hg (95% limits of agreement -13.7 to 21.4 mm Hg) with moderate correlation (r = 0.63, p < 0.001) (Fig. 1A). The mean bias and precision for microPetCO₂ was 7.3 ± 8.2 mm Hg (95% limits of agreement -8.8 to 23.6 mm Hg) with moderate correlation (r = 0.68, p < 0.001) (Fig. 1B). Although both PetCO₂ measurement methods were moderately correlated, mainPetCO₂ was more accurate compared to the microPetCO₂ method overall (in the whole subject group). Additionally, when we evaluated the correlation between mainPetCO₂ and microPetCO₂ throughout all patients, the methods demonstrated a strong level of correlation (r = 0.84, p < 0.001) (Fig. 2).

Study subjects were also compared based on the presence of lung pathology (Table 2). In the mild-to-moderate pulmonary disease group, 1242 measurements from each end-tidal CO₂ method were compared. In this group, the mean bias and precision for mainPetCO₂ was 2.9 ± 8.4 mm Hg (95% limits of agreement – 13.4 to 19.4 mm Hg) with moderate correlation (r = 0.64, p < 0.001) (Fig. 1C). The mean bias and precision for microPetCO₂ was 6.7 ± 7.8 mm Hg (95% limits of agreement – 8.5 to 22.1 mm Hg) with moderate correlation (r = 0.68, p < 0.001) (Fig. 1D). Although both PetCO₂ measurement methods were moderately correlated, mainPetCO₂ was more accurate than microPetCO₂.

In the severe lung disease group, we compared 96 results from each method of measurement. For the mainPetCO₂ and PaCO₂ comparison, the mean bias and precision was 14.8 ± 9.1 (95% limits of agreement – 3.0 to 32.7 mm Hg) (Fig. 1E). Whereas the mean bias and precision between microPetCO₂ and PaCO₂ was 16.2 ± 9.0 mm Hg (95% limits of agreement – 1.4 to 33.9 mm Hg) (Fig. 1F). In the severe lung disease group, almost all PaCO₂ values were higher than PetCO₂ measurements (Fig. 1E,F). Additionally, in this group, both
mainPetCO2 and microPetCO2 were highly correlated with PaCO2 ($r = 0.80, p < 0.001$ and $r = 0.81, p < 0.001$, respectively); however, the biases of both methods increased.

To determine whether the accuracy of the non-invasive PCO2 measurement methods were altered in the presence of high physiologic dead space, we compared the mainPetCO2 and microPetCO2 values with regard to $PaCO2$.

### Table 1. Demographic, clinical and laboratory characteristics of patients (n = 174).

- **PaCO2** = arterial PCO2, **PaO2** = arterial PO2, **mainPetCO2** = mainstream end-tidal PCO2, **microPetCO2** = microstream end-tidal PCO2, **FiO2** = fractional inspired oxygen, oxygenation index = $[(FiO2 \times MAP \times 100)/PaO2]$, IQR: Interquartile range.

| Patients characteristics | Values |
|--------------------------|--------|
| Male sex, no (%)         | 107 (61.5) |
| Age (month), median (IQR)| 42 (12–108) |
| **Primary disease, no (%)** |        |
| Pneumonia                | 39 (22.4%) |
| Multiple trauma          | 29 (16.7%) |
| Status epilepticus       | 22 (12.6%) |
| Shock, multiple organ failure | 19 (10.9%) |
| Postoperative            | 10 (5.8%) |
| Bronchiolitis            | 9 (5.2%) |
| Intracranial mass/hemorrhage | 8 (4.6%) |
| Central nervous system infection | 7 (4.0%) |
| Acute respiratory distress syndrome | 5 (2.8%) |
| Congenital heart disease | 5 (2.8%) |
| Poisoning                | 4 (2.3%) |
| Renal failure            | 4 (2.3%) |
| Others                   | 13 (7.6%) |

| Laboratory values, median (IQR) |
|----------------------------------|
| Arterial blood gas analysis      |
| pH                               | 7.3 (7.3–7.4) |
| PaCO2 (mm Hg)                    | 40.7 (35.4–47.3) |
| PaO2 (mm Hg)                     | 150.0 (115.0–183.0) |
| HCO3⁻ (mmol/L)                   | 22.5 (19.7–25.3) |
| MainPetCO2 (mm Hg)               | 38.0 (32.0–44.0) |
| MicroPetCO2 (mm Hg)              | 35.0 (29.0–40.0) |

| Mechanical ventilator parameters, median (IQR) |
|-----------------------------------------------|
| FiO2 (%)                                      | 40.0 (40.0–50.0) |
| Mean airway pressure (mm Hg)                  | 10.0 (9.0–13.0) |
| Oxygenation Index                             | 2.4 (1.6–3.9) |

### Table 2. Relation between PetCO2 values and severity of pulmonary disease. All CO2 levels in mmHg. LLA = lower limit of agreement, ULA = upper limit of agreement, SD = standard deviation, PCO2 = partial pressure of carbon dioxide, PaCO2 = arterial PCO2, mainPetCO2 = mainstream end-tidal PCO2, microPetCO2 = microstream end-tidal PCO2. aDefinition of mild to moderate pulmonary disease: oxygenation index < 10 (n = 1242 pairs). bDefinition of severe pulmonary disease: oxygenation index ≥ 10 (n = 96 pairs).

| Parameter                                | Mean difference ± SD (mmHg) | 95% LLA (mmHg) | 95% ULA (mmHg) | r      | p        |
|------------------------------------------|-----------------------------|----------------|----------------|--------|----------|
| **In All Subjects (n = 1338 pairs)**     |                             |                |                |        |          |
| PaCO2–MainPetCO2                         | $3.83 \pm 8.99$             | $−13.79$       | 21.46          | 0.63   | < 0.001  |
| PaCO2–MicroPetCO2                        | $7.39 \pm 8.27$             | $−8.83$        | 23.61          | 0.68   | < 0.001  |
| **Severity of pulmonary disease**        |                             |                |                |        |          |
| Mild to moderate pulmonary disease⁴      |                             |                |                |        |          |
| PaCO2–MainPetCO2                         | $2.98 \pm 8.40$             | $−13.49$       | 19.46          | 0.64   | < 0.001  |
| PaCO2–MicroPetCO2                        | $6.70 \pm 7.80$             | $−8.59$        | 22.01          | 0.68   | < 0.001  |
| Severe pulmonary disease⁵                 |                             |                |                |        |          |
| PaCO2–MainPetCO2                         | $14.83 \pm 9.12$            | $−3.06$        | 32.72          | 0.80   | < 0.001  |
| PaCO2–MicroPetCO2                        | $16.24 \pm 9.05$            | $−1.49$        | 33.99          | 0.81   | < 0.001  |
to groups formed according to Vd/Vt ratio (<0.4 vs. ≥0.4). The Vd/Vt ratio was < 0.4 in 1247 of 1338 (93%), and > 0.4 in 91 (7%) measurements.

In the Vd/Vt < 0.4 (the normal physiologic dead space) group, the comparison of mainPetCO2 and PaCO2 values showed a mean bias and precision of 3.0 ± 8.0 mm Hg, with moderate correlation (r = 0.63, p < 0.001). Whereas the mean bias and precision between microPetCO2 and PaCO2 was 6.5 ± 7.0 mm Hg, again with moderate correlation (r = 0.68, p < 0.001). In the Vd/Vt ≥ 0.4 (increased physiologic dead space) group, both mainPetCO2 and microPetCO2 were highly correlated (r = 0.78, p < 0.001 and r = 0.78, p < 0.001, respectively) with increased PetCO2—PaCO2 gradient (bias ± precision values of 20.9 ± 11.2 and 25.02 ± 11.8 mm Hg, respectively). Although both non-invasive PCO2 measurement methods were highly correlated with PaCO2, mainPetCO2 was more accurate than microPetCO2 in both the normal and increased dead space ventilation groups.
To our knowledge, this is the largest cohort study including 174 pediatric patients who received mechanical ventilation in the PICU. The evaluation of 1338 measurements for each method and the comparison of two different PetCO2 monitoring methods with accuracy determined according to simultaneous PaCO2 measurements are among the other strengths of this study. Although different PetCO2 measurement methods have distinct advantages, the accuracy and correlation of these methods in comparison to ABG measurements is without doubt the most vital feature of any method. The microstream capnometer requires in-depth analysis to prove that it contributes to or surpasses available methods by analyzing whether the advantageous properties expressed in the literature are indeed superior in the real-life follow-up of intubated pediatric patients.

Although there are many studies evaluating the accuracy and correlation of various non-invasive PetCO2 measurement methods, the majority of these studies were performed in non-intubated patient groups. In intubated patients, the studies on PetCO2 monitoring are mostly compared with the ABG analysis of a single method and often evaluate the relationship between the severity of lung disease and the accuracy of the method. In our study, two different PetCO2 monitoring methods were evaluated simultaneously, and both mainPetCO2 and microPetCO2 measurements were found to be moderately correlated with PaCO2.

Rozycki et al. reported that mainPetCO2 measurements were highly correlated with PaCO2 in intubated newborns, with a mean bias of -6.9 mm Hg. Similar results have been found in other studies using the mainstream technology in intubated newborns. Microstream is preferred especially in the neonatal age group due to the use of very low flow rates (50 mL/min), causing smaller dead space and allowing measurement from the distal part of the ETT. In the study by Kugelman et al., microPetCO2 was found in adequate agreement with PaCO2, which indicated closer agreement than seen in the current research. Although similar ‘close’ results have been obtained in other studies, Singh and colleagues found similar results to ours in terms of agreement between microPetCO2 and PaCO2. In intubated patients, PetCO2 measurements can be performed from the proximal or distal part of the ETT. To compare the advantages of different PetCO2 measurement technologies in our study, it was thought that the measurements obtained from the same locations would be more guiding. Therefore, in order for one of the methods to gain no advantage due to localization, both PetCO2 measurements were obtained from the same location (proximal part of ETT). In various studies comparing PetCO2 measurements obtained from the distal and proximal parts of the ETT, it has been suggested that distal measurements provide more accurate results; however, several other studies have demonstrated comparable accuracy between proximal and distal measurements.

The first study comparing two different PetCO2 measurement methods in intubated patients was performed by Kugelman and colleagues. This study, which was comprised of 27 infants, showed better correlation between PetCO2 and PaCO2 with distal sampling of expired air using microstream technology against the mainstream method through a proximal port using double lumen ETT. The measurements made in this study were obtained from different locations of the ETT and this situation may have led to an advantage for the microstream method. In our study, although the correlation coefficients of both methods were similar, the agreement level of main PetCO2 measurements was better.

There are various studies investigating the relationships between pulmonary disease and PaCO2-PetCO2 values. These studies have defined pulmonary disease severity according to various parameters, such as OI, arterial-alveolar PO2 gradient and PaO2/FiO2 ratio. In this study we used the OI value to define severe pulmonary disease. Sivan et al. reported in their study that mainPetCO2 and PaCO2 compatibility decreased as lung disease severity increased in neonatal patients. Hagerty et al. evaluated the compatibility of microPetCO2 and PaCO2 in
intubated newborn patients and found that microPetCO₂ and PaCO₂ differences were higher in the pulmonary disease group compared to controls. Different results were reported by other investigators. Tingay et al. found that the PetCO₂ bias was independent of severity of lung disease and similarly Rosycki et al. reported that the degree of lung disease had little influence on the degree of discrepancy between measurement. Kugelman and colleagues reported that although the accuracy of microPetCO₂ decreased with lung disease it still remained good correlation as a useful measure of PaCO₂ in conditions of severe lung disease. The study by McDonald et al. found an overall moderately correlation between PaCO₂ and mainPetCO₂ for all included patients, but the investigators concluded that significant lung disease (defined by PaO₂/FiO₂ < 200) had a negative effect on the correlation. In our study, it was concluded that both PetCO₂ measurement methods highly correlated in patients with severe lung disease, albeit with a significant decrease in measurement accuracy.

The most important parameter contributing to the PetCO₂-PaCO₂ gradient is the increase in physiological dead space due to ventilation-perfusion mismatch. Physiologic dead space ventilation is the sum of anatomic dead space from the conducting airways and alveolar dead space from disease processes and/or therapies employed. The increased gradient between PetCO₂ and PaCO₂ with high PaCO₂ levels are directly proportional to the degree of physiologic dead space. Although typical alveolar CO₂ concentrations are slightly greater than of ABG, PetCO₂ normally 2–5 mmHg lower than PaCO₂ due to mixing of CO₂ containing alveolar gas with exhaled gas devoid of CO₂ from the anatomical dead space. In a patient with lung disease, the addition of alveolar dead space further dilutes PetCO₂ relative to PaCO₂. As a result, PetCO₂ measurements depict greatly reduced results compared to PaCO₂. The normal physiologic dead space to tidal volume ratio (Vd/Vt) is established to be 0.20–0.35. In this study, we provide evidence that physiologic dead space ventilation is a major factor in determining the relationship between capnographic monitoring of PetCO₂ and PaCO₂. Despite multiple earlier publications comparing PetCO₂ and PCO₂ in presence of pulmonary disease and hypercarbia, few studies have examined the effect of change in physiologic dead space on the correlation between PetCO₂ and PaCO₂, across an increased range of Vd/Vt ratios in mechanically ventilated pediatric patients. Our study is the first to investigate the correlations between two different capnometers in patients with increased physiological dead space ventilation.

In patients with a low calculated physiologic dead space to tidal volume ratio (Vd/Vt < 0.4), there is a moderate correlation between both PetCO₂ (measured noninvasively by capnography) measurements and PaCO₂ value. Despite the high correlation between PetCO₂ and PaCO₂ values in patients with high physiologic dead space to tidal volume ratio (Vd/Vt ≥ 0.4), the accuracy of measurements was greatly reduced. Therefore, in the presence of severe pulmonary diseases with increased physiological dead space, it is much more reliable to use PetCO₂ results as a measure of trend rather than absolute value. It is also critical to note that further problems in accuracy may arise with smaller infants or newborns (which were not included in the study population) and reduced volumes or I:E values. In a study including 56 intubated pediatric patients by McSwain et al., it was found that, while the strength of the association diminished slightly as the dead space ratio increased, the correlation still remained strong between the methods. The PaCO₂-PetCO₂ gradient was increased predictably with increasing Vd/Vt. Our findings show that increased physiological dead space as a result of severe pulmonary disease will increase the gradient between PaCO₂ and PetCO₂ in favor of PaCO₂ values, making almost all PaCO₂ results greater than those recorded by PetCO₂. These findings were similar to the outcomes of previous studies performed in newborns and children with pulmonary disease.

To our knowledge, there are no other studies investigating the relationships between PetCO₂ measurements and increased physiological dead space ventilation. There are however, various studies investigating PetCO₂ correlations with hypercarbia as a proxy for increased dead space ventilation. In the study conducted by Kugelman et al., microPetCO₂ was reported as a useful measure of PaCO₂, whereas mainPetCO₂ was distorted on the high range of PaCO₂ level. Rosycki et al. did not find any effect of increased PaCO₂ on mainPetCO₂ measurements.

Our study has several limitations. Non-consecutive ABGs were used for data collection and inadvertent selection bias may have been introduced. In our study, we used proximal measurement method for both PetCO₂ methods. In subsequent studies, the relationship between concurrent microPetCO₂ measurements obtained from the proximal and distal part of the ETT may reveal differences in results which could be crucial for physicians and patients in intensive care units. Although our study reached the highest number of patients and samples in the literature, the number of samples in the subgroups of severe pulmonary disease and increased physiologic dead space ventilation, were relatively low; thus limiting the generalizability of those results. Due to the low number of patients with ARDS, we could not group patients as mild, moderate, severe ARDS with regard to the criteria put forth by the Pediatric Acute Lung Injury Consensus Conference (PALICC); thus, subgroup analyses concerning these groups could not be performed. Also, the number of cases with increased physiologic dead space (Vd/Vt ≥ 0.4) was low, leading to a lack of further subgroup analysis.

**Conclusion**

It was found that the novel microstream capnometer has no superiority to the traditional mainstream method. Although the mainstream and microstream capnometer measurements had similar correlation values with ABG results, the agreement level of the mainstream method was higher. Although the absolute gradient between both PetCO₂ methods and PaCO₂ results demonstrated a consistent increase in the presence of severe pulmonary disease and increased dead space ventilation, both methods showed significant correlations with PaCO₂ values. Therefore, in the presence of severe pulmonary disease and/or increased dead space ventilation, it is possible that both PetCO₂ monitoring methods may be helpful in predicting the trend of PaCO₂ despite limitations in accuracy.

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Author contributions

All authors read and approved the final manuscript. D.M.: study design, statistical analysis, drafted the manuscript, and is responsible for the overall content. D.B.A.: Study design, data extraction and quality assessment. K.Z.: Contributed to the writing of the manuscript and revised the manuscript for important intellectual content. B.M.: Data extraction and quality assessment. G.A.: Data extraction and quality assessment. M.C.Y.: Data extraction and quality assessment. Y.M.N.: Data extraction. B.O.: Data extraction.
