Metabolic acid-base adaptation triggered by acute persistent hypercapnia in mechanically ventilated patients with acute respiratory distress syndrome

INTRODUCTION

Acute respiratory distress syndrome (ARDS) is a common reason for initiating ventilatory support in critically ill patients.\(^{(1)}\) ARDS mortality remains as high as 54% despite advances in critical care.\(^{(2)}\) To minimize ventilator-induced lung injury,\(^{(3)}\) a protective mechanical ventilation based on lower tidal volumes and lower distention pressures is recommended.\(^{(4)}\)
Lung injury increases dead space ventilation.\(^{(5)}\) Furthermore, protective ventilation is associated with reduced effective alveolar ventilation; both factors result in ineffective carbon dioxide clearance from the blood, resulting in hypercapnia.\(^{(6)}\) Currently, there is no consistently demonstrated clinical benefit of hypercapnia in ARDS patients.\(^{(7)}\)

Protective ventilation-induced hypercapnia evokes metabolic responses toward pH normalization within a short period.\(^{(8)}\) In patients with chronic hypercapnic hypoventilation, the metabolic acid-base adaptation is related to plasmatic bicarbonate (\(\text{HCO}_3^-\)) elevation and chloride reduction.\(^{(9)}\) Normal subjects who are acutely exposed to hypercapnia exhibit increased urinary chloride elimination.\(^{(9)}\) In contrast, critically ill patients commonly present with reduced renal chloride removal.\(^{(10,11)}\) Therefore, the mechanism of pH compensation in hypercapnic patients with ARDS has not been established. The aim of this study was to explore the mechanisms involved in pH compensation during acute hypercapnia in patients with ARDS for at least 24 hours.

**METHODS**

The Ethical Committee of the Hospital das Clínicas of Faculdade de Medicina of the Universidade de São Paulo approved this study (approval document number 107.443), and informed written consent was waived. Patient records/information was anonymized and de-identified prior to analysis.

We retrospectively reviewed electronic medical records that were prospectively collected from 1275 patients who were consecutively admitted to our intensive care unit (ICU) from June 2007 to June 2012.

The inclusion criteria were bilateral pulmonary infiltrates on the X-ray, acute onset of hypoxemia, P/F ratio < 300mmHg using a positive end-expiratory pressure (PEEP) ≥ 5cmH\(_2\)O, and no cardiogenic cause of the pulmonary infiltrate.

The exclusion criteria were chronic renal failure on dialysis support, acute kidney injury with any renal replacement therapy mode necessity, bicarbonate infusion and chronic hypercapnia.

The patients were categorized according to the presence or lack of persistent acute hypercapnia. Persistent acute hypercapnia was defined as a partial pressure of carbon dioxide (\(\text{PaCO}_2\)) greater than 50mmHg for more than 24 hours, with at least three arterial blood gas sample analyses during this period, in patients with documented previous normal \(\text{PaCO}_2\) values and without a history of chronic hypercapnia. The hypercapnic group was compared with the control group in order to explore the metabolic compensation to the hypercapnia.

ARDS was defined according to the Berlin conference.\(^{(12)}\) The information that was obtained from the patients’ charts included the following: demographic characteristics (age, gender, weight, height and co-morbidities) and ICU data from the time of admission until the third day after hypercapnia diagnosis [respiratory failure etiology, expected mortality (calculated by the Acute Physiology and Chronic Health disease Classification System - APACHE II)\(^{(13)}\) or the simplified acute physiological score (SAPS 3)\(^{(14)}\) sequential organ failure assessment (SOFA)\(^{(15)}\) score on the first day of ICU stay, need for vasopressors and/or inotropics, laboratory data, fluid balance, diuresis, and daily variations in heart rate, respiratory rate, temperature and mean arterial pressure.

Acid-base arterial blood samples from the day of ICU admission and one day before the hypercapnia installation, with three samples every 8 hours on the day of hypercapnia installation and up to three days after hypercapnia initiation, were analyzed as well as clinical and physiological daily data from these same days. We emphasize that all of the patients were admitted with the diagnosis of ARDS. The laboratorial data that were retrieved during the hypercapnia installation were routinely collected at least every 8 hours until \(\text{PaCO}_2\) stabilization (generally considered when the \(\text{PaCO}_2\) variation is < 3 - 5\%). In the control group, samples were collected at admission and after 24, 48, 72 and 96 hours.

A quantitative physicochemical approach was used to analyze the acid-base variables.\(^{(16)}\) In this approach, after several adaptations\(^{(17)}\) the \([\text{H}^+]\) concentration and, hence, pH, were determined using five independent variables: inorganic apparent strong ion difference (\(\text{SIDai}\)), strong ion gap (\(\text{SIG}\)), lactate, weak acids in plasma (\(\text{Atot}\)), and \(\text{PaCO}_2\) variation.\(^{(18)}\) The standard equations that were used in this study were the following:

\[
\text{SIDai} \ (\text{mEq/L}) = \text{Na}^- \ (\text{mEq/L}) + \text{K}^+ \ (\text{mEq/L}) + \text{Ca}^{2+} \ (\text{mEq/L}) + \text{Mg}^{2+} \ (\text{mEq/L}) - \text{Cl}^- \ (\text{mEq/L})
\]

\[
\text{SIG} \ (\text{mEq/L}) = \text{SIDai} - [2.46 \times 10^{n+6} \times \text{PaCO}_2 \ (\text{mmHg}) + (\text{albumin} \ (\text{g/L}) \times (0.123 \times \text{pH} - 0.631)) + (\text{phosphate} \ (\text{mmol/L}) \times (0.309 \times \text{pH} - 0.469))] - \text{lactate} \ (\text{mEq/L})
\]

\[
\text{Atot} \ (\text{g/dL}) = \text{Albumin} \ (\text{g/dL}) + \text{phosphate} \ (\text{g/dL})
\]
A positive SIG value represents the presence of unmeasured anions, which must be included to determine the measured pH. The standard base excess (SBE) was used in our study to diagnose and quantify the metabolic acid-base variations. The source of SBE variations was analyzed based on SIDai, SIG, lactate, albumin, and phosphate variations.

Statistical analysis

The data distribution was analyzed using the Shapiro-Wilk goodness-of-fit model. The qualitative data, which are shown as occurrences and percentages, were analyzed using Fisher's exact test or chi-square test as appropriate. The quantitative data are presented as the mean and standard deviation values or the medians [25th percentile and 75th percentile], depending on whether the values are parametric or non-parametric, respectively.

The quantitative baseline data were analyzed using non-paired t-test or Mann-Whitney's test as appropriate. The quantitative data of two groups over time were analyzed using interaction analyses and a mixed generalized model with the patient as a random factor for determining the within-subject correlation among repeated observations. The Markov chain Monte Carlo procedure using 1000 simulations to obtain the equilibrium of distributions was used to reach a fixed likelihood of each resulting independent variable. The post hoc analyses for interactions were performed using Mann-Whitney's or Wilcoxon's tests as appropriate. The R free open-source statistical package and Comprehensive-R Archive Network (CRAN)-specific libraries were used to build the graphics and to perform all of the statistical analyses.

RESULTS

There were 49 (4%) patients out of 1275 ICU admissions with ARDS who did not require renal replacement therapy during the study period. A total of 41 patients had the necessary data and were enrolled in the analysis: 26 (64%) patients who developed hypercapnia and 15 (36%) who did not develop hypercapnia as the control group (Figure 1). No patients received diuretics.

The general data for the patients, stratified according to the group, are shown in Table 1. The maximum CO$_2$ levels in the hypercapnic group occurred within the first 48 hours after the patients' admission, with mean values of 60mmHg. Subsequently, a significant difference between groups was observed up to the third day after hypercapnia commencement (Figure 2).

Concomitant with the increasing CO$_2$, the pH levels decreased, with the lowest values observed on the second day of admission. In the ensuing days, the pH gradually increased to values similar to those observed in the control group. The increases in pH were accompanied by elevated SBE and HCO$_3$ elevations, both with a significant difference between the groups (Figure 2). In addition to the SBE elevation, from day 1 to day 3, the PaCO$_2$ slightly decreased, remaining greater than 50mmHg.

The SIDai was greater in the hypercapnic group and was not accompanied by significant sodium and chloride variations, despite a tendency toward elevated sodium (Figure 3) and figure 1S (http://www.rbti.org.br/content/imagebank/pdf/0103-507X-rbti-28-01-0019-suppl01-en.pdf). The analysis of other physiological and laboratory variables during the observation period demonstrated that the serum hemoglobin, calcium and phosphate levels were different between the two groups (Table 1S in http://www.rbti.org.br/content/imagebank/pdf/0103-507X-rbti-28-01-0019-suppl01-en.pdf). Figure 2S (http://www.rbti.org.br/content/imagebank/pdf/0103-507X-rbti-28-01-0019-suppl01-en.pdf) shows the proportional variation of pH according to the respiratory and metabolic determinants. Figure 1S (http://www.rbti.org.br/content/imagebank/pdf/0103-507X-rbti-28-01-0019-suppl01-en.pdf) shows the main acid-base
components of the metabolic adaptation according the patients’ disease severity, using an expected mortality of 20% (median of the expected mortalities) as the cut point.

**DISCUSSION**

The results of our study show a slight elevation of PaCO$_2$ levels in the hypercapnic group before the initiation of hypercapnia, as expected. The SBE and HCO$_3$ levels were higher in the hypercapnic group before hypercapnia initiation, and a subsequent increase in HCO$_3$ was more relevant than the increase in SBE; however, both the SBE and HCO$_3$ levels gradually increased over time (Figure 2) in both groups. The pH initially decreased markedly after hypercapnia initiation and progressively moved toward normalization. The SBE variation in the hypercapnic group could be attributed to an increase in SIDai and a decrease in lactate and in SIG. The SIDai variation was not attributed to chloride variation. In contrast, SIDai improved because of an increase in serum sodium, magnesium and potassium.

Hypercapnia is occasionally required to allow protective ventilation in patients with ARDS. However, the CO$_2$ effect per se (without the concomitant effect of the tidal overdistention reduction) on lung protection and clinical outcomes is controversial.$^{[20-23]}$ Patients with acute and persistent hypercapnia, in the absence of renal failure, evolve a metabolic compensation toward pH normalization in a short time period (approximately 36 hours).$^{[20]}$ This pH compensation makes the tidal volume reduction more acceptable in patients with ARDS.$^{[21]}$

In our patients, the initial reduced pH promptly moved toward normalization after the initiation of hypercapnia, reaching normal values within a period of 24 - 48 hours. The metabolic response was indicated by HCO$_3$ and SBE elevations. The striking initial HCO$_3$ elevation most likely occurred because of a stoichiometric factor, i.e., the elevated CO$_2$ reacted with water, thereby increasing the HCO$_3$ and H$.^{[20]}$ SBE is an independent variable from the acute PCO$_2$ variation;$^{[24]}$ thus, a striking initial elevation did not occur in the SBE in our study. The elevation of SBE and its components occurred in both groups during the observed days but was slightly more accentuated in the hypercapnic group during days 1, 2 and 3, when the mean values of SBE were significantly different between groups. The former group presented a less-increased SBE than that of the control group in the admission, probably secondary to the higher PaCO$_2$ already at this time in the hypercapnic group.

Exploring the SBE and HCO$_3$ elevations after the initiation of hypercapnia, we observed three associated factors in our findings: first, increases in SIDai; second, decreases in lactate; and third, decreases in SIG. The increases in SIDai occurred because of several slight increases in sodium, calcium, magnesium and potassium. We emphasize that serum chloride did not decrease after the initiation of hypercapnia. It is of note that in hypercapnic stable patients with COPD, pH compensation occurred based on a HCO$_3$ elevation and chloride reduction in the blood.$^{[8]}$ Renal chloride excretion improved greatly in acutely hypercapnic sheep,$^{[9]}$ indicating the high relevance.
of chloride modulation in metabolic adaptation triggered by acute hypercapnia in non-critically ill patients. In contrast, in acute critically ill animal models, the acid-base disturbances are marked, especially metabolic acidosis.\textsuperscript{(25,26)}

In critically ill patients, metabolic acidosis is relevant, multifactorial, and related to clinical outcomes.\textsuperscript{(27,28)} These patients present reduced sodium and chloride renal excretion,\textsuperscript{(10,11,29)} together with chloride shift from the extravascular to the intravascular compartments added to exogenous load during fluid resuscitation,\textsuperscript{(30)} frequently resulting in hyperchloremia.\textsuperscript{(27,31)} Therefore, we speculate that these factors most likely differentiate the responses of critically ill patients from stable patients when they are exposed to hypercapnia in terms of chloride and SIDai modulations.

The decrease in lactate after the initiation of hypercapnia was another interesting finding in our study. The lactate behavior in hypercapnic patients is consistent with the study of Carvalho et al.\textsuperscript{(6)} In an experimental model of endotoxemia, lactate production decreased with ongoing hypercapnia between 40mmHg and 60mmHg.\textsuperscript{(32)} A similar reduction in lactate production occurred when hypercapnia was initiated in hypoxemic animals.\textsuperscript{(33)} Several mechanisms are related to this lactate-hypercapnia
Physicochemical variables of the acid-base metabolic component before and after hypercapnia initiation. A) SIDai evolution (mixed model fixed effects $p = 0.646$ for within-group factor analysis, $p = 0.045$ for between-group factor analysis, and $p = 0.224$ for group × time interaction analysis). B) Strong ion gap evolution (mixed model fixed effects $p < 0.001$ for within-group factor analysis, $p < 0.001$ for between-group factor analysis, and $p = 0.007$ for group × time interaction analysis). C) Lactate evolution (mixed model fixed effects $p < 0.978$ for within-group factor analysis, $p < 0.001$ for between-group factor analysis, and $p = 0.975$ for group × time interaction analysis). D) Atot evolution (mixed model fixed effects $p = 0.141$ for within-group factor analysis, $p = 0.010$ for between-group factor analysis, and $p = 0.266$ for group × time interaction analysis). SIDai - inorganic apparent strong ion difference; SIG - strong ion gap; Atot - weak acids in plasma. * Mann-Whitney’s post-hoc analysis $p < 0.05$ versus control group. # Wilcoxon’s post-hoc analysis $p < 0.05$ versus admission day.

In our patients, the presence of SIG acidosis at ICU admission can be observed in figure 3 - Panel B. This metabolic acidosis is common in critically ill infected patients, and its improvement during the first five days of ICU stay is associated with better clinical outcomes. (28) This extra source of metabolic acidosis can be a confounder of the pH evolution interpretation.

Unmeasured anion concentrations also decreased after the initiation of hypercapnia. These unmeasured anions have not been identified in humans. (34) In an animal model of hemorrhagic shock, however, these molecules were highly constituted by Krebs cycle components, such as citrate and acetate. (35) Therefore, the aerobic mitochondrial metabolism modulation of hypercapnia may result in SIG variation by the same mechanism of lactate variation. Furthermore, in figure 2S (http://www.rbti.org.br/content/imagebank/pdf/0103-507X-rbti-28-01-0019-suppl01-en.pdf), subtle variations in each metabolic pH determinant can significantly impact pH variation. In this figure, the more parallel the tested variable line is from the x axis, the more striking is its deviation effect on the pH variation, as is, for example, the SIDai.
Clearly, the PaCO₂ decreased and was associated with the pH normalization during this study in a very important way. This observation probably represents the patient’s ventilatory improvement over time. However, metabolic adaptation also occurred, in a similar manner to the control group, but faster.

Our study has many limitations: the data were retrieved from a prospective collected database; different sources of ARDS and sepsis could influence the metabolic adaptation to hypercapnia in different ways; other sources of metabolic acidosis are additional confounders, mainly in two different groups with different disease severities; individual variations were not considered in our study; and this study was drawn only as an explanatory analysis, as there is a paucity of data in this field in the current literature.

CONCLUSION

In this explanatory study, the results indicate that metabolic acid-base adaptation, which is triggered by acute persistent hypercapnia in patients with acute respiratory distress syndrome, is a complex process. The more rapid standard base excess adaptation than the control group involves decreases in lactate and the strong ion gap and increases in the inorganic apparent strong ion difference, which occur due to slight increases in serum sodium, magnesium, calcium, and potassium but not significant decreases in serum chloride.

Authors contributions

TG Romano analyzed the data and wrote the manuscript. MD Telles collected the data. PV Mendes participated in the data analysis. FG Zampieri designed the study and data analysis. AT Maciel participated in the design of the study and statistical analysis. M Park collected the data, designed the study, analyzed the data and wrote the manuscript. All of the authors approved the final manuscript.

RESUMO

Objetivo: A hipercapnia resultante da ventilação protetora na síndrome do desconforto respiratório agudo desencadeia uma compensação metabólica do pH que ainda não foi completamente caracterizada. Nosso objetivo foi descrever esta compensação metabólica.

Métodos: Os dados foram recuperados a partir de uma base de dados registrada de forma prospectiva. Foram obtidas as variáveis dos pacientes no momento da admissão e quando da instalação da hipercapnia até o terceiro dia após sua instalação. Analisamos 41 pacientes com síndrome do desconforto respiratório agudo, incluindo 26 com hipercapnia persistente (pressão parcial de gás carbônico acima de 50mmHg por mais de 24 horas) e 15 sem hipercapnia (Grupo Controle). Para a realização da análise, utilizamos uma abordagem físico-química quantitativa do metabolismo acidobárico.

Resultados: As médias de idade dos grupos com Hipercapnia e Controle foram, respectivamente, de 48 ± 18 anos e 44 ± 14 anos. Após a indução da hipercapnia, o pH diminuiu acentuadamente e melhorou gradualmente nas 72 horas seguintes, de forma coerente com os aumentos observados no excesso de base padrão. A adaptação metabólica acidobárica ocorreu em razão de diminuições do lactato sérico e do strong ion gap e de aumentos na diferença aparente de strong ions inorgânicos. Além disso, a elevação da diferença aparente de strong ions inorgânicos ocorreu por conta de ligeiros aumentos séricos de sódio, magnésio, potássio e cálcio. O cloreto sérico não diminuiu por até 72 horas após o início da hipercapnia.

Conclusão: A adaptação metabólica acidobárica, que é desencadeada pela hipercapnia aguda persistente em pacientes com síndrome do desconforto respiratório agudo, foi complexa. Mais ainda, aumentos mais rápidos no excesso de base padrão em pacientes com hipercapnia envolveram diminuições séricas de lactato e ions não medidos, e aumentos na diferença aparente de strong ions inorgânicos, por meio de ligeiros aumentos séricos de sódio, magnésio, cálcio e potássio. Não ocorreu redução do cloreto sérico.

Descritores: Acidose respiratória; Síndrome do desconforto respiratório do adulto; Equilíbrio ácido-base; Respiração artificial; Unidades de terapia intensiva
