Introduction

Sulfanilamide, one of the first oral antibiotics, was initially used more than 70 years ago. It was observed that the drug induced both metabolic acidosis and hyperventilation, owing to renal carbonic anhydrase (CA) inhibition. Synthesis of stronger CA inhibitors yielded the more potent sulfonamide, acetazolamide. Pure or mixed metabolic alkalosis is a common finding in the intensive care unit (ICU) and is associated with a detrimental outcome. By inducing metabolic acidosis, acetazolamide might facilitate discontinuation from mechanical ventilation in patients who suffer from chronic obstructive pulmonary disease (COPD) and who have also developed metabolic alkalosis. The evidence supporting the administration of acetazolamide in such a situation, however, remains sparse.

Chronic obstructive pulmonary disease in the intensive care unit

COPD is a progressive and irreversible disease that is defined by an expiratory airflow limitation and caused by a mixture of small-airway disease and lung parenchyma destruction [1]. Cigarette smoking is the most common risk factor for developing the disease. COPD is a major public health issue owing to the associated morbidity and mortality. In the US, COPD is the fourth leading cause of morbidity and mortality [1]. The natural history of the disease is marked by the occurrence of exacerbations affecting the prognosis of patients [2]. COPD is a frequent cause of emergency referrals and ICU hospitalization [3,4]. Initiation of non-invasive mechanical ventilation in the setting of COPD exacerbations reduces mortality [5]. However, initiation of invasive mechanical ventilation may be necessary (for instance, after the failure of an initial trial of non-invasive mechanical ventilation). Acute COPD exacerbations are associated with in-hospital mortality rates of between 24% and 32% [4,6]. Invasively ventilated patients with COPD are at a high risk of prolonged mechanical ventilation [7] and at a high risk of persistent weaning failure. Prolonged invasive mechanical ventilation is associated with an increased hospital mortality [8,9].

A series of factors are associated with persistent weaning failures from mechanical ventilation in patients with COPD: age, severity of initial presentation, associated diseases such as left ventricular dysfunction, metabolic disorders, critical illness polyneuromyopathy, and the existence of ventilator-associated pneumonia [7,9]. Weaning from mechanical ventilation of patients with...
COPD appears to be more successful when undertaken in specialist multidisciplinary wards [10]. Metabolic alkalosis, a common condition in the ICU, is reported to be associated with difficulties in obtaining a successful weaning from mechanical ventilation, especially in patients with COPD [11,12].

**Metabolic alkalosis in the intensive care unit**

The acid-base equilibrium is regulated in the healthy subject by the kidney and the lungs. A disruption of this equilibrium can be due to an alteration of lung function (causing respiratory acidosis or alkalosis by hypo- or hyperventilation) or an alteration of kidney function (inducing an acidic or alkaline charge causing metabolic acidosis or alkalosis) or both [13,14]. Respiratory acidosis and metabolic alkalosis are the two main acid-base disturbances found in patients with COPD during respiratory exacerbations. Metabolic alkalosis results when the H+ concentration in the extracellular compartment is decreased by a loss of non-carbonic acid or an increase in alkali. The treatment of chronic carbonic dioxide retention frequently induces metabolic alkalosis, also known as post-hypercapnic alkalosis. It is a state of persistent metabolic alkalosis after the return of arterial partial pressure of carbon dioxide (PaCO₂) to baseline [11]. Mixed acid-base balance disturbances (the association of chronic respiratory acidosis and metabolic alkalosis) can also be observed. Indeed, in mechanically ventilated patients with COPD, the most frequently observed acid-base disorders are mixed [15,16]. In this context, respiratory acidosis is the consequence of hypercapnia. Respiratory acidosis can be acute or chronic, depending on renal adaptation. In patients with both COPD and respiratory exacerbation, hypercapnia induces an acidic change of the extracellular compartment, a lowering of arterial pH, and an excess of serum bicarbonate.

Metabolic alkalosis is a common disorder in the critically ill and is often associated with other metabolic anomalies (hypokalemia, hypomagnesemia, hypophosphatemia, and hyperalbuminemia) [17,18]. Metabolic alkalosis in patients affected by COPD may diminish the activity of the central nervous respiratory command center and cardiac output [19,20]. In the ICU, metabolic alkalosis has been shown to inhibit the dissociation of oxyhemoglobin and to lead to the development of hypokalemia and hypophosphatemia [19,20]. It has also been suggested that metabolic alkalosis plays a role in prolonging the weaning period and in increasing morbidity and mortality in the critically ill, especially patients with COPD [11,12,21]. Other causes of failure to wean from mechanical ventilation are due to the respiratory control system, mechanics of the lung and chest wall, the respiratory muscles, the cardiovascular system, and gas-exchange properties of the lung [22].

Metabolic alkalosis is frequently iatrogenic, owing to the administration of diuretics or steroids, but may also be the result of permissive hypercapnia induced by lung-protective strategies and by digestive disorders such as vomiting or nasogastric suctioning [11]. Metabolic alkalosis can be treated, after correction of hypokalemia, by the administration of acetazolamide or chloride salts [23]. Correcting metabolic alkalosis may increase minute ventilation and improve oxygenation, which may in turn facilitate weaning from mechanical ventilation [24,25].

**Carbonic anhydrase inhibitors**

CAs are metalloenzymes found in both the vegetable and the animal kingdoms [26]. The enzyme catalyzes an essential physiological function, the interconversion between carbon dioxide (CO₂) and bicarbonate [26]. The zinc ion of CAs plays an important role in catalysis. The active form of the CA enzyme is basic, and a hydroxide radical is bound to the zinc ion. Since the active form of the enzyme is a strong nucleophile, it attacks the CO₂ molecule, forming bicarbonate coordinated to the zinc ion. The bicarbonate ion is then released after being replaced by a water molecule. The CA enzyme family is compromised in vertebrates of 16 isoforms. CA isoforms have wildly different subcellular and tissue distributions throughout the organism [27-29]. The enzyme plays an important role in the alveolocapillary transport of CO₂, in the regulation of the acid-base and hydroelectric equilibrium, and in the control of respiration [30]. CA inhibitors include the classic inhibitor acetazolamide as well as a number of more recent drugs [31]. These drugs are used in varied situations such as in the treatment of hypertension, glaucoma, diabetes, and cancer [26]. All CA inhibitors possess a zinc-binding group by which the drug interacts with the metal ion of the enzyme. Acetazolamide, a sulfonamide derivative, is a non-selective inhibitor of the CA enzyme. Since obstructive pulmonary disease is complicated by an increase in arterial CO₂ levels, the main isozymes of interest in the disease are CA II (which plays a role in regulating gas exchange in the lung and in the erythrocyte) and CA IV (which plays a role in bicarbonate reabsorption in the kidney but also in regulating gas exchanges in the lung) [26]. Acetazolamide might also modulate the activity of slowly adapting pulmonary stretch receptors (SARs) via the inhibition of CA. SARs seem to play a role in the control of respiratory rate and tidal volume [32]. Indeed, SARs play a role in evoking the Hering-Breuer inflation reflexes. This is characterized by an early termination of inspiration when the lungs are inflated and a prolongation of the expiratory pause when a prolonged inflation is applied at the end of inspiration [32]. The inhibition of SAR activity occurs during inhalation of CO₂ [33,34]. In turn, the administration of acetazolamide attenuates or
blocks the inhibitory effects of CO₂ inhalation and could modulate the activity of SARs [32,35]. Acetazolamide is used as a respiratory stimulant in subjects with COPD [30]. The classic explanation of why acetazolamide acts as a respiratory stimulant is through the inhibition of the renal CA enzyme, which in turn induces a decrease of serum bicarbonate and serum pH. The ensuing metabolic acidosis causes a stimulation of peripheral chemoreceptors (carotid artery) and central chemoreceptors (through the acidification of the cerebrospinal liquid), which in turn induces an increase in minute ventilation. However, the effect of CA inhibitors on ventilation and ventilator responses seems more complex [30]. For instance, the effect of the modulation of the activity of the SARs by acetazolamide in the airway remains largely unknown. Hypothetically, this modulation could induce a respiratory-stimulating effect by increasing both respiratory rate and tidal volume. The tissue compartmentalization of CA isoforms and the low selectivity of acetazolamide may explain, in part, the complexity of the effect of the drug in patients with COPD [30,36] and, in turn, might explain why the efficacy of acetazolamide is so moderate in the critically ill patient with COPD. An alternative reason could be that factors other than metabolic alkalosis are more important during discontinuation from mechanical ventilation.

Since acetazolamide is a non-specific CA inhibitor, the administration of the drug is susceptible to induce a host of varied physiological consequences on nervous conduction, ventilation control, oxygen transport and on diuresis. Acetazolamide also seems to have inhibiting effects at the neuromuscular level. Studying hypercapnic conditions in an animal model, Kiwull-Schone and colleagues [28] found that, after administration of low doses of acetazolamide, an increased neuronal drive was necessary to maintain a given tidal volume. Similarly, in healthy subjects, Brechue and colleagues [37] reported that acetazolamide inhibited the Achilles tendon-tap reflex and associated isometric force. Acetazolamide, administered at doses ranging from 250 to 500 mg (3.5 to 7 mg/kg), induces, at sea level, an increase in minute ventilation by 10% to 20% both at rest and during exercise in healthy subjects [38] and an increase in oxygen saturation of hemoglobin by 3% to 6% in subjects with hypoxemia [20,30]. The drug induces an alkaline diuresis by reducing the tubular reabsorption of bicarbonate and by inhibiting the distal secretion of H⁺. After administration of acetazolamide, 30% of filtered bicarbonate is eliminated in the urine, whereas the transepithelial transport of bicarbonate is reduced from 70% to 100% [39,40]. Alkaline diuresis induced by acetazolamide is maximized at 24 hours and is associated with a urinary loss of bicarbonate of about 4 to 6 mmol/L, which in turn induces a decrease of blood pH by 0.05 to 0.1 units. The resulting metabolic acidosis stimulates peripheral and central chemoreceptors associated with an increase in minute ventilation. This respiratory response induces a decrease of PaCO₂ of 5 to 6 mm Hg (provided that there is no mechanical limitation to the increase of tidal volume or respiratory rate) [30]. According to Moviat and colleagues [41], the effect of acetazolamide is mediated by an increased renal excretion of strong ions (bicarbonates), a retention of chloride, and consequently a decrease in the strong ion difference defined by the equation ([Na⁺] + [K⁺] + [Mg²⁺] + [Ca²⁺]) – ([Cl⁻] + [lactates⁻]). This strong ion difference is one of the parameters that govern water dissociation and thus the serum concentration of H⁺.

**Pharmacology**

The plasma half-life of acetazolamide in the healthy subject is between 4 and 8 hours [42]. Absorption of the drug after an oral intake is very fast. Acetazolamide is eliminated mainly by the kidney and the liver. In case of renal or hepatic dysfunction, acetazolamide is therefore susceptible to accumulate [43].

Both furosemide and corticosteroids lessen the effect of acetazolamide on serum bicarbonate concentration. Indeed, both drugs induce metabolic alkalosis by stimulation of distal tubular H⁺ secretion [11]. Additionally, acetazolamide and furosemide are transported by the same carrier-mediated mechanism from the plasma to their site of action localized at the luminal side of renal tubules [44]. In vitro, furosemide interacts with the CA enzyme at the same site as acetazolamide, potentially blocking the interaction between acetazolamide and the CA enzyme [27]. The respiratory effects of acetazolamide at low or moderate doses are well known, mainly in the healthy subject. The effect of acetazolamide is less well known in critically ill patients with COPD. Additionally, these patients often receive loop diuretics during the weaning period [15].

The drug is relatively safe; the main undesirable effects are hypersensibility, blood dyscrasias, dysthyroidism, and gout attacks. These undesirable effects, though potentially severe, are rare [45]. Acetazolamide in patients with cirrhosis is susceptible to induce or aggravate hepatic encephalopathy and therefore should be avoided [43]. Another potential complication of acetazolamide administration is an aggravation of hypercapnia because of red blood cell CA inhibition and impaired CO₂ transport [46]. If the contraindications of the drug are respected, the only real issue faced by physicians in the ICU is hypokalemia. Therefore, blood chemicals should be monitored closely when acetazolamide is administered to the critically ill.

**Clinical data**

Vos and colleagues [47] showed that the administration of acetazolamide to patients with COPD improves
arterial blood gas parameters without significantly changing minute ventilation. On the other hand, Swenson [30] and Teppema and Dahan [36] showed that acetazolamide increases minute ventilation by reducing excess base levels in the healthy subject. It has also been reported that non-responders to acetazolamide (defined by a reduction of PaCO₂ of less than 5 mm Hg) have a more severe airflow limitation than responders [48]. This finding has not been confirmed in later studies [16]. Mazur and colleagues [49] assessed, in 40 invasively ventilated patients with both COPD or asthma and pure metabolic alkalosis, the effect of two dosing regimens of acetazolamide (500 mg per day versus 250 mg four times per day) on the variations of bicarbonate over the course of 72 hours. A significant and lasting decrease of bicarbonate concentration was reported in both treatment groups. These results suggest that the pharmacological effect of acetazolamide lasts longer than expected from the plasma half-life of the drug (5 to 6 hours) in patients with both COPD and pure metabolic alkalosis. However, no clinically relevant endpoint was assessed in this study. It must also be noted that relatively few patients in the ICU were subject to a pure metabolic alkalosis [15]. Another study in a surgical ICU gave similar results when acetazolamide was administered at a dosage of 500 mg per day [50].

Although acetazolamide is used to treat metabolic alkalosis, few data concerning the optimal dosage of the drug in the ICU are available. Usual dosage regimens vary from a single dose of 250 or 500 mg per day [49,50] to multiple doses of 250 or 500 mg every 6 to 8 hours [25,51]. A recent case control observational study showed that acetazolamide, administered at a dosage of 500 mg once a day during the weaning period from mechanical ventilation of patients with both COPD and pure or mixed metabolic alkalosis, significantly decreased the serum concentration of bicarbonate and increased the arterial partial pressure of oxygen/fraction of inspired oxygen (PaO₂/FiO₂) ratio [15]. No effect on respiratory parameters such as minute ventilation or respiratory frequency was observed [15]. This decrease in bicarbonate concentration was statistically significant when compared with the control group but was of little clinical importance (less than 4 mmol/L). Additionally, patients in the control group presented a slight decrease in serum bicarbonate concentration during the weaning period. The moderate effect of acetazolamide in lowering the serum bicarbonate concentration could explain the lack of efficacy of the drug in reducing PaCO₂. Indeed, the pharmacodynamics of the drug in the ICU are not well known. The pharmacodynamics of acetazolamide were recently modeled in 68 mechanically ventilated patients who had COPD and who received single or multiple doses of up to 500 mg of the drug during the weaning period [16]. The main co-variates of interest that were found to influence acetazolamide pharmacodynamics were the Simplified Acute Physiology Score II (SAPS II) at ICU admission, co-prescription of furosemide or corticosteroids, and serum concentrations of chloride. Additionally, co-administration of furosemide was found to significantly decrease bicarbonate elimination. In this pharmacodynamic model, acetazolamide induced a decrease in serum bicarbonate with a dose-response relationship, and the amount of acetazolamide found to induce 50% of the putative maximum effect was 117 ± 21 mg. According to this model, an acetazolamide dosage of greater than 500 mg twice daily is required to reduce serum bicarbonate concentrations of greater than 5 mmol/L in the presence of high serum chloride levels or co-administration of systemic corticosteroids or furosemide.

Conclusions

The efficacy of acetazolamide given to patients with both COPD and metabolic alkalosis during the weaning period remains unknown. In the ICU, the usually employed doses of the drug seem insufficient to have a meaningful effect on the serum concentration of bicarbonate. Surprisingly, up until now, no randomized controlled trial has been undertaken to assess the efficacy of the drug in patients with COPD in the ICU. To our knowledge, at least three randomized controlled studies that aim to determine the efficacy of acetazolamide given to patients with both COPD and metabolic alkalosis during the weaning period are under way (NCT01131377 and NCT01499485 [52] and a French trial that is not yet registered). From a clinical point of view, the compartmentalization of CAs is a promising new subject for the development of drugs modulating these enzymes, especially CA localized in the airway, the erythrocyte, and the diaphragm.

Abbreviations

CA, carbonic anhydrase; CO₂, carbon dioxide; COPD, chronic obstructive pulmonary disease; ICU, intensive care unit; PaCO₂, arterial partial pressure of carbon dioxide; SAR, slowly adapting pulmonary stretch receptor.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

NH and CF were responsible for literature research, manuscript writing, and final approval. SU was responsible for final approval. All authors read and approved the final manuscript.

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Published: 7 August 2012
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doi:10.1186/cc11323
Cite this article as: Heming N et al.: Acetazolamide: a second wind for a respiratory stimulant in the intensive care unit? Critical Care 2012, 16:318.