Severe metabolic acidosis in a patient with an extreme hyperglycaemic hyperosmolar state: how to manage?

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Abstract

Hyperglycaemic hyperosmolar state (HHS) and diabetic ketoacidosis (DKA) are often accompanied by severe metabolic and electrolyte disorders. Analysis and treatment of these disorders can be challenging for clinicians. In this paper, we aimed to discuss the most important steps and pitfalls in analyzing and treating a case with extreme metabolic disarrangements as a consequence of an HHS. Electrolyte disturbances due to fluid shifts and water deficits may result in potentially dangerous hypernatriema and hyperosmolality. In addition, acid-base disorders often co-occur and several approaches have been advocated to assess the acid-base disorder by integration of the principles of mass balance and electroneutrality. Based on the case vignette, four explanatory methods are discussed: the traditional bicarbonate-centered method of Henderson-Hasselbalch, the strong ion model of Stewart, and its modifications 'Stewart at the bedside' by Magder and the simplified Fencl-Stewart approach. The four methods were compared and tested for their bedside usefulness. All approaches gave good insight in the metabolic disarrangements of the presented case. However, we found the traditional method of Henderson-Hasselbalch and 'Stewart at the bedside' by Magder most explanatory and practical to guide treatment of the electrolyte disturbances and in exploring the acid-base disorder of the presented case.

Introduction

Severe acid-base disorders combined with electrolyte disturbances are of ten seen in patients with hyperosmolar hyperglycaemic state (HHS) or diabetic ketoacidosis (DKA). Acid-base disorders and electrolyte disturbances are tightly connected [1]. The aims of this clinical lesson are (i) to give insight in the relation between hyperglycemia, acid-base disturbances and electrolyte homeostasis, (ii) to provide a strategy for the treatment of HHS and DKA during the first hours, and (iii) to compare different methods in analyzing acid-base disorders.

Case

A 38-year-old patient was admitted to the ICU because of a hyperosmolar Hyperglycaemic state with an extreme metabolic (and combined respiratory) acidosis and hypernatriemia (Table 1). Glucose levels were gradually lowered with iv insulin and extensive rehydration. Initially, the metabolic acidosis improved.

Balance between hyperglycemia, acid-base disorders and electrolyte disturbances

The hyperglycaemia of this patient caused severe acid-base and electrolyte disturbances of which the main elements are (i) the effects of ketoacids on other ions and (ii) the effects of the hyperglycemia on extracellular volume and sodium concentrations. By definition electrolytes dissociate in water in anions (negatively charged, predominately chloride) or in cations (positively charged, predominately sodium). However, in pathologic conditions these electrolytes can be extensively lost or gained resulting in a temporary ionic imbalance. To maintain electrochemical neutrality, changes in acid-base equilibrium occur.

The compensatory mechanism to restore electrochemical neutrality includes the dissociation of water in hydrogen (H⁺) and vice versa [1]. This is accompanied by changes in pCO₂ and bicarbonate (HCO₃⁻) levels, depending on the cause of the acid-base disorder.

An absolute shortage of insulin in diabetic ketoacidosis (DKA) results in the accumulation of negatively charged ketoacids from acetyl-Coa serving as a fuel for the brain. These ketoacids anions become apparent in blood when their hepatic production exceeds the renal excretion capacity. To compensate for the accompanying protons from the ketoacids, water and carbon dioxide are formed from bicarbonate to enable respiratory compensation [2]. This equation is the cornerstone of the traditional approach according to Henderson-Hasselbalch for the assessment of acid base disturbances (Textbox). More detailed description and analysis of the acid base disturbance of the presented case will follow below.

Another important element in this pathologic cascade includes sodium and water regulation. Increased glucose levels augment the plasma tonicity because glucose is an effective osmole and both the hyperglycemia and the ketoacids salts cause a decrease in the extracellular volume, factors that contribute to enormous deficits in total body water and changes in plasma sodium concentration [2]. Hyponatriemia may become apparent since sodium, also an effective osmole, remains in the extracellular (the blood) compartment during the glucose-driven water shift. Be aware that normal or increased plasma sodium levels during severe hyperglycemia indicate a substantial loss of total body

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water! As a rule of thumb in clinical medicine, the serum sodium concentration decreases by 3 mmol/L for every 10 mmol/L increase in glucose concentration. Although this correction factor is based on theoretical considerations and has not been well validated, it gives good insight into the severity of water loss [2]. In practice, however, it may vary between 1-4 mmol/L.

The application of this correction factor to our case revealed an extremely dangerous sodium concentration of 182 mmol/L. This is in agreement with an osmolality of 447 mosm/kg and an estimated water shortage of 13 liters of body water.

First-hours treatment plan

Patients in these severe metabolic deranged states need immediate treatment and do not allow much time for assessments. Many guidelines regarding the treatment of HHS and DKA exist and it is beyond the scope of this article to discuss them in detail. However, the extreme hypernatriemia illustrates the importance of careful fluid administration, specifically concerning the prevention of cerebral edema for reasons delineated below.

The initial goal of fluid therapy in a Hyperglycaemic crisis is to stabilize hemodynamics with isotonic solutions. During resuscitation it is important to monitor the corrected plasma sodium levels, but even more important is the monitoring of the effective plasma osmolality (Table 2) [2]. Correction of effective plasma osmolality should be gradual (no more than 4 m Osm/h or 10 m Osm in 2-3 hours) to prevent fast opposing fluid shifts, from extracellular to intracellular, that could lead to cerebral edema when brain cell osmolality is not (yet) adapted to the faster lowering of the extracellular (plasma) osmolality [3,4]. Current expert opinion is based on animal studies and case reports and states that the risk for cerebral edema is determined by changes in effective plasma osmolality [5]. This is considered most important during the first 15 hours of treatment [2,6]. Symptomatic cerebral edema is mostly described in children with diabetic ketoacidosis [7]. It is seldom seen in adults with DKA or HHS [8-10]. However, several guidelines [3,4] advise to consider corrected sodium and effective osmolality and to adjust fluid therapy accordingly, which implies that the reduction in glucose levels is accompanied by increases in (measured) sodium levels in order to safely decrease the plasma osmolality. There is no consensus between different guidelines if the fluid treatment should be based on effective, measured or calculated plasma osmolality, but we suggest to use effective or calculated plasma osmolality since it is quicker in use.

In our patient extensive rehydration with normal saline was started in addition to glucose lowering therapy. Even though plasma sodium remained high, the (calculated) osmolality dropped gradually along with glucose levels, indicating safe and effective rehydration and therapy (Figures 1A and 1B).

Methods in analyzing acid-base and electrolyte disturbances

Several explanatory models aim to help understanding the pathophysiology of metabolic disarrangements. We describe these disarrangements in our case with the traditional model based on the Henderson-Hasselbalch equation, the Stewart approach and with 2 modified methods based on the Stewart approach: 'Stewart at the bedside by Magder' and the simplified Fencel-Stewart approach (Textbox) [11,12].

| Laboratory results | At presentation | After 2 hours | After 16 hours |
|-------------------|----------------|--------------|--------------|
| Sodium (mmol/l)   | 146            | 152          | 168          |
| Potassium (mmol/l)| 4.3            | 2.9          | 5            |
| Chloride (mmol/l) | 90             | 113          | 135          |
| pH                | 6.86           | 6.99         | 7.37         |
| pCO2 (kPa)        | 9.1            | 8.2          | 4.1          |
| Bicarbonate (mmol/l)| 12            | 14.4         | 17.9         |
| Base excess (mmol/l)| -22.1         | -17.3        | -6.5         |
| Lactate (mmol/l)  | 7.6            | 5            | 7.8          |
| Osmol (mOsmol/kg) | 480*           | 453          | 398          |
| Albumin (g/l)     | 28*            | 28           | 19           |
| Glucose (mmol/l)  | 120*           | 92           | 29.5         |
| ionized Calcium (mmol/l)| 1.54       | 1.42         | 1.18         |
| Magnesium (mmol/l)| 1.11*          | 1.11         | 0.9          |
| Phosphate (mmol/l)| 0.35*          | 0.4          | 0.9          |
| Urea (mmol/l)     | 35*            | 32.6         | 28           |

*data are abstracted from first available drawings.

Table 2. Glossary of important formula’s in assessing metabolic disarrangements.

| Formula | Reference value | Case, at presentation |
|---------|----------------|-----------------------|
| Anion gap Correction for albumin | (Na+K]- (Cl+ HCO3 ) per 1 g/dl below/above 4 g/dl albumin add/subtract 2.5 mmol/L to the AG [28] | 5-11 mmol/L | 51.3 |
| Serum osmolality | 2xNa+glucose+urea | 275-300 mOsmol/kg | 470* |
| Osmol gap | calculated- measured osmolality | <10 mOsmol/kg | 8 |
| Corrected sodium | Measured Na + (glucose/3) in mmol/l | 135-145 mmol/L | 180 |
| Delta ratio | (AG-normal value)/(24-HCO3) | 0.8-1.2: ketoacidosis, lactate acidosis >1.2: combined metabolic alkalosis <0.8: combined normal AG metabolic acidosis | >1.2 |

*abstracted from first available drawings.

AG: anion gap
Delta ratio: Delta AG/delta HCO3 (mmol/L)
Na: sodium
K: potassium
Cl: chloride
HCO3: bicarbonate

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Traditional method

The traditional method is a bicarbonate-centered approach. As explained in the textbox, its main principle is that bicarbonate and hydrogen are in equation with carbon dioxide (pCO₂) and that the metabolic acidosis is based on the extent of bicarbonate loss. Either by loss of the ion resulting in a equilibrium shift to the left resulting in hydrogen formation, or by increases in hydrogen pushing the reaction to shift to the right with consequently an increase of pCO₂. Compensatory mechanisms are respiratory to exhale excess pCO₂ or renal to retain bicarbonate. Our patients' low level of bicarbonate combined with an elevated pCO₂ fits with a combined respiratory and metabolic acidosis. The elevated pCO₂ is likely due to respiratory insufficiency as a result of his comatose state.

For more insight into the origin of the metabolic acidosis the anion gap is calculated with correction for the albumin concentration (Table 2). Failure to do so may result in an underestimation of the anion gap due to altered charges of weak anions because of albumin-based changes in pH [13] and finally lead to a false interpretation of the acid-base disorder [14,15].

Our patient had an extremely increased anion gap of 51.3 indicating the presence of other anions than chloride or bicarbonate. Indeed, ketoacids were found, but in relatively low concentrations suggestive for the co-existence of other anions. Other well-known anions that attribute to an increased anion gap are lactate or alcholic ketoacids, or in case of co-existing increased osmole gap ingestion of ethylene glycol, ethanol or methanol [13]. An osmole gap was not detected in the presented case. Assessment of the delta ratio (Table 2) helps identifying the presence of co-existing metabolic acid-base disorders and is used to provide an estimate of the acid load [6,16]. In a 'normal' high anion gap acidosis every 1 mmol/L rise in/of the anion gap is expected to be accompanied by a 1 mmol/L decrease in the bicarbonate concentration, which is indicated by a delta ratio of 0.8-1.2 [16]. In contrast, a delta ratio below 0.8 indicates a combined high anion gap and normal anion gap metabolic acidosis and a delta ratio above 1.2 indicates a combined high anion gap metabolic acidosis and metabolic alkalosis [17,18]. In our case the delta ratio was increased (>1.2) which indicates the presence of a metabolic alkalosis. This metabolic alkalosis could be explained with the hypochloremia that most likely reflects a renal compensation of a probably longstanding existing metabolic acidosis and/or osmotic diuresis.

In conclusion, the extreme increased anion gap could only (partly) be explained by the presence of serum lactate and ketoacids and other, unknown anions should have attributed to the acidosis as well. Moreover, failing renal compensation of the acidosis, in a later stadium of the disease, with increased renal bicarbonate loss, as described by Kamel et al. may have added to the acidosis [6].

Stewart principle

The Stewart principle is based on physicochemical principles of water-based solutions. Its main principle is that all action and anion concentrations must balance, according to the laws of electroneutrality. This is achieved by the dissociation of water as the source of H⁺. It further dictates that pH is determined by three independent variables; the partial pCO₂, the difference between the concentrations of strong cations and anions (SID), and the concentration of weak acids (Atot) [14,15,19]. In the textbox the Stewart approach is further explained. Atot is the sum of non volatile (i.e. non CO₂) weak acids that are
existing in a dissociated form (dissociated albumin and phosphates) and in an associated form accompanied by a proton. They are often referred to as buffers [13,20].

In our patient, analysis of the first obtained sample showed a combined respiratory and metabolic acidosis due to unmeasured anions which generally include lactate, formate, ketoacids, salicylate, and sulfate [21].

Magder and Fencl-Stewart approaches

Next, modified methods based on the Stewart principle and developed for bedside application of physicochemical principles ‘Stewart at the bedside’ by Magder (further denoted as Magder) and the simplified Fencl-Stewart approach are applied to assess the presented case.

Magder included the pCO₂ as a traditional component and defined four other metabolic processes that influence the acid-base balance: the water-effect, the chloride-effect, the albumin-effect and ‘others’ [11]. This approach showed - as also showed in the Textbox and visualized in Figure 1E - a combined respiratory and metabolic acidosis, mostly due to the effect of ‘others’. Similarly to the unmeasured anions of Stewart’s analysis, ‘others’ include the typical negative ions in the differential diagnosis of a wide anion gap acidosis (e.g. lactate, ketoacids, salicylate, sulfate) [11]. In addition, Magder revealed a positive (+15.7) chloride-effect at admission that contributed to a metabolic acidosis and counterbalanced the acidosis. The positive chloride-effect is in agreement with the increased delta ratio as detected with the traditional method.

Lastly, the simplified Fencl-Stewart approach integrated the standard base excess and the Stewart analysis and uses four variables: the standard base excess (SBE), as measured by a bloodgas machine, and calculates the base excess effects of sodium-chloride, albumin and unmeasured ions [12,22]. Sodium-chloride is seen as the principal component of the strong ion difference (SID) and albumin as the principal component of the total of weak anions (Atot). The unmeasured ions may be strong ions (e.g. sulphate, acetate) or weak acids (e.g. phosphate) pCO₂ is considered a separate variable and should be analyzed to assess the respiratory component in acid-base disorders [12].

SBE is a calculated figure derived from pCO₂ and arterial pH [23]. The calculation of SBE assumes a normal plasma protein concentration. However, in the critically ill plasma protein is often lowered, which may limit the accuracy of the SBE [24]. With this in mind this method is useful as a rough estimate of the acid-base disorder.

The assessment of the acid-base disorder of our patient following the simplified Fencl-Stewart approach is demonstrated in Figure 1F. At admission, the true SBE, the SBE corrected for the sodium-chloride and albumin-effect (as also explained in the Textbox), is strongly negative (-43.6), indicating that unmeasured ions are an important factor in causing this extreme acidosis.

Case continued

Several hours into treatment lactate levels were rising again and our patient became hyperchloremic. This, in combination with a diminished diuresis, raised the suspicion of an abdominal compartment syndrome and abdominal CT scan was performed. The scan showed signs of pancreatitis (Figure 2).

At this time point, we analyzed his acid-base disturbance again (see Table 1 for used laboratory findings). We found the anion gap (25.3) (Figure 1C) and the strong ion gap (14.1) (Figure 1D) still increased, but numbers were significantly decreased compared to baseline values. Magder (Figure 1E) showed a metabolic acidosis and a compensatory respiratory alkalosis. The acidosis is explained by a negative chloride-effect (-10.5, recall that this was a positive effect) and the presence of ‘others’, albeit in lower concentrations than initially. The positive albumin- and water-effect (5.01 and 8.40, respectively) indicate a counterbalancing alkalinizing effect. This coincides with a metabolic alkalosis as revealed by the increased delta ratio. The positive water-effect revealed a substantial shortage of body water, still present despite extensive resuscitation.

Analysis using the simplified Fencl-Stewart approach (Figure 1F) showed a negative sodium-chloride effect meaning that the SID has become smaller as a result of the (iatrogenic) hyperchloremia. Again, a positive albumin-effect was observed. The minimal difference between the measured SBE and the true SBE excluded the presence of unmeasured anions, which opposed the findings of the methods above.

To summarize, by assessing our patients’ extreme metabolic disarrangements and acid-base disorder using the four different methods we were able to compare them and to judge their clinical applicability and diagnostic value. Figure 1 clearly demonstrates the changes in laboratory results as assessed by the different methods in the first day after treatment initiation. From this several statements may be derived. First, the traditional approach allowed the detection of the different components of the acid-base disturbances. The traditional method detects the accompanying metabolic alkalosis, as long as all the correct steps are followed. Second, the Stewart approach gives a good physiological insight, but one tends to focus on the strong ion gap (SIG), missing possible accompanying metabolic disorders (e.g. hypochloremia). Another acknowledged drawback is its clinical applicability as it requires complex calculations. Third, Magder seems to give a complete picture of the probable causes of the acid-base disturbances. When critically reviewing the results from Magder, the shift in chloride-effect on the acid-base disorder is striking. Initially the chloride-effect was positive, contributing to an alkalinizing effect, while at our second time point the opposite was demonstrated with the

Figure 2. Abdominal CT-scan of the presented case (see text) showed edema of the pancreas (white arrow) suggestive for an acute pancreatitis.
development of an iatrogenic hyperchloremia. Fourth, the simplified Fenc-Stewart approach is an easy to use method and gives a quick insight in the seriousness of the problem, but does not give any physiological insight in the various components that contribute to the acid-base disturbances.

In conclusion, at this time point the contribution of unmeasured anions to the acidosis had decreased, as assessed with the different methods and in agreement with with the treatment of the hyperglycemia. New, and clearly demonstrated by Magder, is the effect of the iatrogenic induced hyperchloremia to the acid base balance. Moreover, the high anion gap or ‘others’ effect is most likely due to decreased renal acid excretion as a result of renal insufficiency and elevated lactate levels because of an abdominal compartment syndrome.

Conclusion

In conclusion, this case demonstrates the complexity of acid-base and electrolyte disturbances in patients with DKA or HHS.

We showed the importance of understanding the corrected sodium levels and osmolality as indicators of the severity of the metabolic derangement and fluid status of a patient and as parameters to guide choices in fluid therapy.

Four different methods of analyzing acid-base disorders, based on the principle of mass balance and electrochemical neutrality, were compared by step by step analysis of a case with severe HHS and metabolic disarrangements.

Most clinicians are familiar with the traditional bicarbonate-centered approach which, as demonstrated by critical reviewing this extreme case, detected the same acid-base disorders as the Stewart-based approaches as long as the anion gap is corrected for plasma albumin concentrations. In our opinion, the traditional method or Magder are the most easy to use and most insightful methods in analyzing acid-base disorders.

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