Narrowing the Gap between the Anion Gap and the Strong Ion Gap

Berend K* and Lundquist A L

1 Department of Medicine, St. Elisabeth Hospital, Netherlands
2 Division of Nephrology, Massachusetts General Hospital, United States

Submission: February 11, 2017; Published: June 01, 2017

*Corresponding author: Kenrick Berend, Department of Medicine, St. Elisabeth Hospital, p/a: St. Michielsweg 13, Curaçao, Netherlands Antilles, Email: kenber2@me.com

Abstract

Background: Despite its importance in understanding acid-base pathophysiology, many physicians do not comprehend the concept of the strong anion gap (SIG), the core of the Stewart acid-base approach. The quantitative difference between the anion gap and the strong ion gap is not established. This paper will give insight into this difference over a variety of conditions.

Methods: The empiric difference between the SIG and the albumin-corrected anion gap (AGc) was calculated at a wide range of albumin, phosphorus and pH levels.

Results: At an albumin level of 1-3 g/dl and pH from 6.9-7.3, the contribution difference of albumin between the AGc and the SIG will be maximally -0.97 to 0.51mEq/L. In metabolic alkalosis and hypoalbuminaemia, the AGc differs less than 2mEq/L from the SIG. The calculated contribution of phosphorus is higher in the SIG with phosphorus levels >2mmol/L and can be accounted for in the anion gap with the conversion factor 1.76* [phosphorus, in mmol/L].

Conclusion: The SIG and the AGc are nearly identical across a wide range of values, particularly when albumin and phosphorus levels are low. The anion gap will be more precise and incorporate the major components of the SIG using the equation: [Na+] - [Cl-] - [HCO3-] - 2.5* [albumin, in g/dL] - 1.76* [phosphorus, in mmol/L], with an arbitrarily set reference range of 1±5mEq/L.

Keywords: Acid-base; Anion gap; Strong anion gap; Stewart

Introduction

For more than 40 years clinicians have used the anion gap (AG) as a major tool to evaluate acid-base disorders. A positive value of the AG suggests a possible organic acidosis due to endogenous acids or the intake of exogenous acids [1]. Although the concept of the AG was described in 1936 by James Gamble [2], it did not gain widespread recognition by physicians until the 1970s after the introduction of auto analyzers and the rapid availability of multiple analytes. According to Gamble, electrical neutrality in solution demands that the sum of the cations is equal to the sum of the anions, also represented in a Gamblegram (Figure 1). Sodium, chloride, bicarbonate and albumin are quantitatively the major ions in the extracellular fluid compartment and are therefore used to calculate the anion gap. A true “ion gap” does not exist in vivo which makes the anion gap a fundamental tool to evaluate acid-base disorders [1]. In 1978 Peter Stewart challenged the traditional bicarbonate-based approach, by reasoning that his approach offered both a mechanistic explanation and provided the tool to make a more accurate diagnosis [3]. The core of the Stewart approach is the strong anion gap (SIG): Equation 1.
[Na\(^+\)]+[K\(^+\)]+[Ca\(^{2+}\)]+[Mg\(^{2+}\)]-[Cl\(^-\)]-[lactate\(^-\)]-12.2×pCO\(_{2}\)/\((10^{-mmol})-10×[albumin]×0.123×pH-0.631]-[PO\(_4\)\(^3-\)] \text{ in mmol/L}×(0.309×pH-0.469)=+0mEq/L

This complex formula quantitatively accounts for the contribution of weak acids to the electrical charge equilibrium in plasma [3]. On first glance there is a substantial difference from the AG equation: Equation 2.

[Na\(^+\)]-[Cl\(^-\)]-[HCO\(_3\)\(^-\)]=+10mEq/L

Though the SIG gives a wider-ranging picture of the ion balance than the AG, controversial opinions about its real value remain and several differences are observed between the SIG and the AG:

a. The SIG is popular amongst intensivists, though additional parameters can increase variability and error [4,5]. Additionally, small changes in K\(^+\), Ca\(^{2+}\) and Mg\(^{2+}\) are rarely of consequence due to narrow variations in health and disease. Nevertheless, some clinicians include K in the AG. Likewise, one can include Ca and Mg in the AG. The AG may therefore be written like [Na\(^+\)]+[K\(^+\)]+[Ca\(^{2+}\)]+[Mg\(^{2+}\)]-[Cl\(^-\)]-[HCO\(_3\)\(^-\)]; the reference value should increase accordingly. These kations are included in SIG (equation 1). When concentrations of these cations are changed substantially one may include them in the AG as well. Therefore, these cations are not relevant for the discussion.

b. Lactate may be included in the SIG, but clinicians will typically know when a high AG or SIG is caused by lactate, because lactate is often tested in severely ill patients with metabolic acidosis.

c. The formula 12.2×pCO\(_{2}\)/(10\(^{-mmol}\)) is used by blood gas devisors to calculate the bicarbonate concentration and may be therefore replaced by it. This leaves us with the following equation which starts to approximate the more familiar anion gap: Equation 3.

[Na\(^+\)]-[Cl\(^-\)]-[HCO\(_3\)\(^-\)]-10×[albumin]×(0.123×pH-0.631)-[PO\(_4\)\(^3-\)]×(0.309×pH-0.469)=+0mEq/L

d. Thus, the "only" real difference between the SIG and the AG is that albumin and phosphorus are adjusted for changes in pH in the SIG. The purpose of this study is to determine the significance of this difference across a wide range of conditions.

Methods

The quantitative difference between the SIG and the AG, was calculated at a wide range of albumin (from 1 to 5 g/L), phosphorus (from 0.5 to 4 mmol/L) and pH levels (from 6.9 to 7.6) with otherwise standard parameters. For example, with an albumin of 4 g/dl and a pH of 6.9, the albumin effect on the anion gap will be 10mmol/L regardless of the pH. The SIG will change using the formula 10×[albumin]×(0.123×pH-0.631), by 8.71mmol/L. Thus, the Δ AlbSt-10=8.71-10=-1.29mmol/L. Similarly, with a phosphorus of 1mmol/L and a pH of 6.9, the PSt= [PO\(_4\)\(^3-\)]×(0.309×pH-0.469)=1.66emEq/L difference with the AG.

Results

At an albumin level of 1-3 g/dl and a pH from 6.9-7.3, the contribution difference of albumin in the AGc and the SIG will be maximally -0.97 to 0.51mEq/L, suggesting little difference between the two methods (Table 1). In metabolic alkalosis and hypoaalbuminaemia, once again the AGc differs less than 2mEq/L from the SIG. Incorporation of high values of phosphorus in the SIG on the contrary, is significant and there is a linear relationship of the serum phosphorus and the ionic contribution in the SIG. This effect will therefore be more notable at higher phosphorus levels (Table 1).

Table 1: Calculated difference in albumin and phosphorus concentration via the Steward and Anion Gap methods across a physiologic pH range.

<table>
<thead>
<tr>
<th>pH</th>
<th>6.9</th>
<th>7</th>
<th>7.1</th>
<th>7.2</th>
<th>7.3</th>
<th>7.4</th>
<th>7.5</th>
<th>7.6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Albumin(^a)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 g/dL</td>
<td>2.18</td>
<td>2.3</td>
<td>2.42</td>
<td>2.55</td>
<td>2.67</td>
<td>2.79</td>
<td>2.92</td>
<td>3.04</td>
</tr>
<tr>
<td>Δ AlbSt - 2.5</td>
<td>-0.32</td>
<td>-0.2</td>
<td>-0.08</td>
<td>0.05</td>
<td>0.17</td>
<td>0.29</td>
<td>0.42</td>
<td>0.54</td>
</tr>
<tr>
<td>2 g/dL</td>
<td>4.35</td>
<td>4.6</td>
<td>4.85</td>
<td>5.09</td>
<td>5.34</td>
<td>5.58</td>
<td>5.83</td>
<td>6.08</td>
</tr>
<tr>
<td>Δ AlbSt - 5</td>
<td>-0.65</td>
<td>-0.4</td>
<td>-0.15</td>
<td>0.09</td>
<td>0.34</td>
<td>0.58</td>
<td>0.83</td>
<td>1.08</td>
</tr>
<tr>
<td>3 g/dL</td>
<td>6.53</td>
<td>6.9</td>
<td>7.27</td>
<td>7.64</td>
<td>8.01</td>
<td>8.38</td>
<td>8.75</td>
<td>9.11</td>
</tr>
<tr>
<td>Δ AlbSt - 7.5</td>
<td>-0.97</td>
<td>-0.6</td>
<td>-0.23</td>
<td>0.14</td>
<td>0.51</td>
<td>0.88</td>
<td>1.25</td>
<td>1.61</td>
</tr>
<tr>
<td>4 g/dL</td>
<td>8.71</td>
<td>9.2</td>
<td>9.69</td>
<td>10.18</td>
<td>10.68</td>
<td>11.17</td>
<td>11.66</td>
<td>12.15</td>
</tr>
<tr>
<td>Δ AlbSt - 10</td>
<td>-1.29</td>
<td>-0.8</td>
<td>-0.31</td>
<td>0.18</td>
<td>-0.68</td>
<td>1.17</td>
<td>1.66</td>
<td>2.15</td>
</tr>
<tr>
<td>5 g/dL</td>
<td>10.89</td>
<td>11.5</td>
<td>12.12</td>
<td>12.73</td>
<td>13.35</td>
<td>13.96</td>
<td>14.58</td>
<td>15.19</td>
</tr>
<tr>
<td>Δ AlbSt - 12.5</td>
<td>-1.61</td>
<td>-1</td>
<td>-0.38</td>
<td>0.23</td>
<td>0.85</td>
<td>1.46</td>
<td>2.08</td>
<td>2.69</td>
</tr>
<tr>
<td>Phosphorus(^a)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

How to cite this article: Berend K, Lundquist A.L. Narrowing the Gap between the Anion Gap and the Strong Ion Gap. J Anest & Intern Care Med. 2017; 3(1): 555601. DOI: 10.19080/JAICM.2017.03.555601.
Discussion

Body fluid compartments have varying concentrations of non-volatile weak acids, with albumin and inorganic phosphorus the primary components in plasma [3]. The anion gap is used to evaluate metabolic acidosis and is calculated as follows:

\[ [Na^+] - [Cl^-] - [HCO_3^-] \]

Because of the narrow extracellular concentration, \( K^+ \) is often omitted from the calculation. To appreciate the relevance of this correction one can consider a healthy individual with the following serum values:

\[ [Na^+] = 140 \text{ mEq/L}, \quad [Cl^-] = 106 \text{ mEq/L}, \quad [HCO_3^-] = 24 \text{ mEq/L}. \]

The AG = 10 mEq/L, representing primarily albumin. The correction factor for albumin is 2.3-2.5 * [albumin], in g/dL [1]. The albumin corrected AG (AGa) equation can therefore be written as: Equation 4.

\[ [Na^+] - [Cl^-] - [HCO_3^-] - 2.5 \times \text{[albumin, in g/dL]} = 0 + 5 \text{ mEq/L}. \]

This equation gives a better understanding of the principle that there is no actual “gap” between the positive and negative ions and likewise, the “AG” should be zero in-vivo. However, comparing this formula with the SIG, one will observe that a major difference is the inclusion of phosphorus in the SIG and that albumin and phosphorus are both adjusted for changes in pH. The question is therefore if one should include phosphorus as well in the AG.

Albumin

Hypoalbuminemia is common in patients with acid-base disorders. With an albumin of 1-3 g/dL, the contribution difference of albumin between the AGa and the SIG will be maximally -0.97 to 1.6 mEq/L regardless of pH (Table 1). Therefore, the pH correction of albumin used in the SIG formula is irrelevant in the evaluation of metabolic acidosis, as long as the AG is adjusted for albumin.

Phosphorus

When phosphorus levels are normal or reduced, there is little contribution (<2 mmol/L) to the AG (Table 1). If phosphorus levels are elevated, as in renal failure, the ionic contribution, however, becomes more significant. Suppose we have a patient with renal failure and the following plasma electrolytes:

\[ [Na^+] = 140 \text{ mEq/L}, \quad [Cl^-] = 106 \text{ mEq/L}, \quad [HCO_3^-] = 24 \text{ mEq/L}, \quad \text{albumin 4g/dL}, \quad \text{phosphorus of 6mmol/L (reference value 0.97-1.45 mmol/L), and a pH of 7.0}. \]

The anion gap corrected

\[ \Delta \text{Alb}_\text{St} - \text{albumin at a certain level calculated by the Stewart method minus the albumin concentration adjusted for the corrected anion gap method (2.5 g/dL for each g/dL change). Example: with albumin = 4 g/dL at a pH = 6.9 the albumin effect on the anion gap will be 10 mmol/L (regardless of the pH in the anion gap concept). The SIG will change using the formula 10 × [albumin] × (0.123 × pH - 0.631), by 8.71 mmol/L.} \]

\[ \Delta \text{PSt} - \text{phosphorus concentration at a certain level calculated by the Stewart method minus the phosphorus. Example: If the phosphorus is 1 mmol/L and the pH is 6.9, the change in SIG will be 1.66 mmol/L using the formula: } [\text{PO}_4^-] \times (0.309 \times \text{pH} - 0.469). \]

\[ [\text{AlbSt}] = \text{albumin concentration calculated by the Stewart method: } 10 \times [\text{albumin}] \times (0.123 \times \text{pH} - 0.631). \]

\[ [\text{PSt}] = \text{phosphorus concentration calculated by the Stewart method: } [\text{PO}_4^- \text{in mmol/L}] \times (0.309 \times \text{pH} - 0.469). \]

Reference range albumin (3.5-5.5 g/dL)

Reference value phosphorus: 0.97-1.45 mmol/L
for albumin will be 10 mEq/L. When using equation 3 the SIG will be 6.78mEq/L due to the high phosphorus level. The linear relationship of serum phosphorus and the ionic contribution in the SIG can be incorporated in the traditional AG formula (Figure 2). The SIG can thus be replaced by the following adjustment to the AG formula: Equation 5.

\[ AG_{\text{corrected}} = [\text{Na}^+] - [\text{Cl}^{-}] - [\text{HCO}_3^-] - 2.5 \times \text{albumin, g/dL} - 1.76 \times [\text{phosphorus, mmol/L}] \]

and as the median phosphorus level is about 1.2, the reference value will be about -2.5mEq/L. A similar equation has been proposed by Kellum et al. [4].

**Conclusion**

In conclusion, the SIG and AG are almost identical in a variety of physiological acid-base conditions. When phosphorus levels are elevated, one can use the less complex AG equation as an accurate representation of the SIG. To have a better understanding of the pathophysiology and to be more accurate, the anion gap, or perhaps a more logical term, “the ion gap” should be written as equation 5 to become almost identical to the SIG.

**Acknowledgment:**

We are indebted to Jan Willem Boldingh for making Figure 1.

**References**