Arterial Blood Gases: A Simplified Bedside Approach

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Rec date: Feb 09, 2014; Acc date: Jul 07, 2014; Pub date: Jul 09, 2014

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Abstract

Interpretation of arterial blood gases is an essence of critical care. It helps for the assessment of clinical oxygenation, ventilation, and acid-base status in critically ill patients. The four closely inter-related physiological parameters pH, PCO2, HCO3 and PO2 helps to diagnose, monitor, and manage the ICU patients. Its correct interpretation and application necessitates the knowledge of basic applied physiology in relation to these parameters. The present article is an attempt for the simplified approach for the bedside interpretation of ABG.

Keywords: Arterial blood gases; pH; PCO2; HCO3; PaO2

Basic Introduction of Arterial Blood Gases

The term arterial blood gas ("ABG" or simply blood gas, with "arterial" left unstated) refers to a specific set of tests performed on arterial blood sample. It provides four key pieces of information: pH, PO2, HCO3 and PCO2 and thereby clinical oxygenation, alveolar ventilation and acid base status. The name blood gas is really a partial misnomer since H+ and HCO3 are not gases. Interpretation arterial blood gases are an important diagnostic tool for the evaluation of oxygenation, ventilation and acid base status. This article is basically designed to teach the basic principles of arterial blood gas analysis. A background in respiratory physiology is recommended, but not required to understand this approach. The article is aimed to teach a practical and systematic approach to understanding of oxygenation and acid-base disturbances encountered in clinical practice. For trainees, a review of the basic terminology and physiology is included.

The term hypoxia is a reduced O2 delivery to tissues.

The term hypoxemia is a reduced O2 content (CaO2) in arterial blood. A normal PaO2 is dependent on the atmospheric pressure, temperature, inspired O2 concentration (FiO2), and the patient's age. Essentially PaO2 is five times FiO2 when it is less than 50%. A patient can be hypoxic for two basic reasons; oxygen may not be delivered to the alveolar air sacs (hypoventilation) or oxygen in the alveoli may not enter into the blood stream. Hypoxia is not a term which is clinically used, however, we hope that our patients have normoxia, or normal levels of oxygen (generally attached to hemoglobin) with or without oxygen supplementation.

A patient can be hypercarbic (elevated levels of CO2) OR hypocarbic (reduced level of CO2) in the blood which is due to an inability to normally exchange gas in the lungs.

The terms acidemia and alkalemia refer to alterations in blood pH, and are the result of underlying disturbance(s) (metabolic and/or respiratory). The suffix “-emia” is pertaining to the blood. The terms acidosis and alkalosis refer to the processes that alter the acid-base status. There can be one or more than one of these processes simultaneously present in a patient.

Diseases that alter the acid-base status of a patient can be divided:

1. Metabolic
2. Respiratory

Metabolic processes are those that primarily alter the HCO3 concentration in the blood. A decrease in serum HCO3 (an alkali or base) leads to a metabolic acidosis, while an increase in serum HCO3 leads to a metabolic alkalosis [1-3].

**Bicarbonates are of two types but which is clinically important?**

**Actual Bicarbonate:** The plasma HCO3 is a commonly used parameter for interpretation and is easily calculated from PaCO2 and pH using Henderson-Hasselbalch equation.

**Standard Bicarbonate:** This is plasma bicarbonate obtained after the blood has been equilibrated at 37°C with a PaCO2 of 40 mmHg. This is not considered for interpretation ABG. Thus one can afford to forget standard HCO3 in the interpretation context.

Respiratory processes alter the pH by changing the CO2 levels. Remember CO2 is a respiratory acid. CO2 accumulation causes an acid state in the blood (through carbonic acid), and as respirations (respiratory rate and/or tidal volume) increase, the body eliminates more CO2 (acid) and is left with a respiratory alkalosis. In other words, a decrease in ventilation leads to retention and increased levels of CO2, and thus a respiratory acidosis.

In conclusion, pH altering processes can be one of four types: Respiratory acidosis, or alkalemia, metabolic acidosis and/or alkalosis. Again, one or more of these processes may be present in a patient with an abnormal acid-base status. One can have two metabolic processes to gather but NOT two respiratory processes simultaneously.

**Systematic Analysis of ABG**

Arterial blood gases are obtained for three basic purposes:

1. To determine oxygenation and
2. To determine acid-base status.
3. To determine alveolar ventilation
To ensure complete interpretation of these points, it is important to systematically examine each component of the arterial blood gas. This article is aimed to elaborate how to systematically determine oxygenation, and then evaluate the acid-base status.

Assessment of Oxygenation i.e. Alveolar: arterial oxygen gradient: (A-a) \( \text{DO}_2 \)?

(Age and FiO\(_2\) dependent derivative)

An important part of interpreting blood gases is to assess oxygenation. An arterial oxygen concentration of less than 60 mm Hg, associated with an oxygenation of less than 90%, is poorly tolerated in humans; therefore a PaO\(_2\) of less than 60 is termed hypoxicem. However, "normal" oxygenation decreases with age as the lungs become less efficient at diffusing oxygen from the alveolus to the blood. Again, normal oxygenation for age can be estimated

\[
\text{PaO}_2 = 104.2 - (0.27 \times \text{age})
\]

Or more crudely, normal oxygenation for age is roughly 1/3 of the patient's age subtracted from 100. Using this estimation for example a 60-year-old patient should have a PaO\(_2\) of 80. Values less than this would be considered hypoxicem for age.

Calculating the alveolar: arterial oxygen gradient: (A-a) \( \text{DO}_2 \) can determine if hypoxia is a reflection of hypoventilation (in other words, reduced because of a rise in PaCO\(_2\), 1 mm rise of PaCO\(_2\) decreases PaO\(_2\) by 0.9 in alveolar sac) or due to deficiency in oxygenation. Unlike oxygen (for which alveolar concentrations are higher than arterial concentrations), CO\(_2\) is 20 times more freely diffuses across the lung such that the arterial and alveolar concentrations are identical. As patient hypoventilates, CO\(_2\) will accumulate in the body (more CO\(_2\) is produced through metabolism than can be eliminated) and thus the blood (where we measure it as PaCO\(_2\)).

The carbon dioxide displaces the oxygen in the alveolus. This reciprocal relationship between oxygen and carbon dioxide in the alveolus is described by the alveolar gas equation:

\[
\text{PAO}_{2} = 104.2 - (0.27 \times \text{age})
\]

Where \( \text{PAO}_{2} \) = partial pressure of Oxygen in the alveolus.

\( \text{PaO}_{2} \) = partial pressure of Oxygen in the arterial blood.

This equation assumes that the patient is breathing room air (21% O\(_2\)) at atmospheric pressure.

Where do 150 come from?:

(Atmospheric P - water vapor P) x FiO\(_2\). At room temperature, at sea level,

\[\text{Atmospheric pressure} = 760 \text{ mm Hg};\]

In the lung, the air is fully saturated with H\(_2\)O, giving a H\(_2\)O vapor pressure of about 47.

\[\text{Room Air} = 21\% \times \text{at room air, the PAO}_{2} = 0.21(760-47) = 149.7, \text{ or about 150.}\]

And

Where does 1.25 come from?

This is a fudge factor which is derived from the respiratory quotient. The formula actually requires that the \( \text{PACO}_{2}\) be divided by the \( \text{RQ}\), which is expressed as the ratio of \( \text{CO}_2\) produced to \( \text{O}_2\) consumed (and which is related to diet and metabolism). We estimate the \( \text{RQ}\) to be 0.8, and 1.25 is the reciprocal of 0.8.

This value is the partial pressure of \( \text{O}_2\) within the alveolus. Because the \( \text{CO}_2\) freely diffuse from arterial blood to alveolar airspaces, the \( \text{PACO}_2\) is equal to the \( \text{PaCO}_2\), which is measured in the arterial blood gas [2]. The above equation can then be rewritten as

\[
\text{PAO}_2 = 150 - 1.25 \times \text{PaCO}_2
\]

Thus

\[
\text{A-a} \text{ DO}_2 = \text{PAO}_2 - \text{PaO}_2
\]

Or

\[
\text{A-a} \text{ DO}_2 = [150 - 1.25 \times \text{PaCO}_2] - \text{PaO}_2
\]

A normal A-a gradient is 10-20 mm Hg, the gradient increases with advancing age. An increased A-a gradient analyses decreased \( \text{O}_2\) in the arterial blood compared to the \( \text{O}_2\) in the alveolus. This infers a process that interferes with oxygen transfer, or in general terms, suggests physiological ventilation-perfusion mismatch. A normal A-a gradient in the face of hypoxemia suggests the hypoxemia is due to hypventilation and not due to underlying lung disorders. It differentiates ventilatory and oxygenation failure [4,5].

- When the patient is not breathing room air.....

\[
\text{A-a} \text{ gradient} = ((\text{FiO}_2 \times 760-47) - (1.25 \times \text{PaCO}_2)) - \text{PaO}_2
\]

Simplified approach to analyze Acid-Base status

In order to understand the various processes that can co-exist in a patient, one must systematically evaluate the blood gases and serum electrolytes. The article uses 6 simple steps to analyze the acid-base status of the patient.

Normal values and range of ABG parameters

\[
\begin{align*}
\text{pH} & : 7.40 (7.35-7.45) \\
\text{PCO}_2 & : 40 (35-45) \\
\text{HCO}_3 & : 24 (22-26) \\
\text{PaO}_2 & : 100 (90-100)
\end{align*}
\]

Just the numerical values doesn’t tell you normalcy, all values are to be interpreted in the context of each other and with patient’s clinical condition (Table 1).

Remember by heart: (CO\(_2\) is a respiratory acid)

\[
\begin{align*}
\text{pH} \text{ and } \text{HCO}_3 & : \text{Moves in same direction} \\
\text{pH} \text{ and } \text{PCO}_2 & : \text{Moves in opposite direction} \\
\text{HCO}_3 \text{ and } \text{PCO}_2 & : \text{Moves in same direction (simple disorder)} \\
\text{HCO}_3 \text{ and } \text{PCO}_2 & : \text{Moves in opposite directions (Mixed disorder)}
\end{align*}
\]
Step 1. Consider the clinical settings! Anticipate the disorder!

Step 2. Look at the pH?

Step 3. Who is the culprit for changing pH?...Metabolic / Respiratory process

Step 4. If respiratory…… acute and /or chronic And Is metabolic compensation appropriate?

Step 5. If metabolic, Is respiratory compensation appropriate? Anion gap increased and/or normal or both?

Step 6. Is more than one disorder present? Mixed one?

Table 1: Steps in Acid-Base Analysis

Step 1: Consider the clinical settings! Anticipate the disorder!

Clinical assessment based on clinical settings is an essential first step.

From the history, examination and initial investigations make a clinical decision as to what is the most likely acid-base disorder(s).

This is very important a word of caution there are situations where the history may be inadequate, misleading to the range of possible multiple diagnoses.

Difficult to diagnose are mixed disorders: the history and examination alone are usually insufficient in sorting these out.

1. Vomiting: Metabolic alkalosis
2. Diarrhea: Metabolic acidosis
3. Septicemia: Lactic acidosis
4. Hypotension, Hypoxemia, Shock: Lactic acidosis
5. Diabetes mellitus: Ketoadacidosis
6. Pneumonia: Respiratory alkalosis/acidosis
7. Bronchial asthma: Respiratory alkalosis/acidosis
8. Hepatic failure: Respiratory alkalosis, later on metabolic alkalosis
9. CNS disorders: Respiratory alkalosis
10. Renal disorders: Metabolic acidosis

Key point: Metabolic alkalosis and acidosis can exist together with any respiratory either acidosis or alkalosis. Both respiratory disorders can’t occur simultaneously

Step 2: Look at the pH

The pH of the arterial blood gas measurement identifies the disorder as alkalemic or acidic.

pH > 7.4 .......... Alkalosis
pH < 7.4 .......... Acidosis
pH = 7.4 ............. Normal or mixed disorder
pH is abnormal in uncompensated, partially compensated but near normal in compensated disorders (pH will come to normal range)

STEP 3: Who is responsible for this change in pH? Who is the CULPRIT?

HCO₃⁻…… METABOLIC PCO₂ …… Respiratory
> 26 .... Met. Alkalosis > 45 ...... Respiratory Acidosis
< 22 ...... Met. Acidosis < 35 ...... Respiratory Alkalosis

It is essential to determine whether the disturbance affects primarily the arterial PaCO₂ or the serum HCO₃.

- Respiratory disturbances alter the arterial PaCO₂ (normal value 35-45)
- Metabolic disturbances alter the HCO₃ (normal value 22-26) pH returns to normal, not absolutely normal but near normal.
- Initial disturbance and compensation follow “same direction” rule.

If the pH is low (i.e., the primary and controlling disturbance is acidosis causing acidemia) either the PaCO₂ is high or the HCO₃ is low (These are the only ways in which the pH can be low). A high PaCO₂ defines a primary respiratory acidosis and a low HCO₃ defines a primary metabolic acidosis.

Conversely, if the pH is high (i.e., the primary and controlling disturbance is alkalosis causing alkalemia) either the PaCO₂ is low or the HCO₃ is high (These are the only ways in which the pH can be high). A low PaCO₂ defines a primary respiratory alkalosis and a high HCO₃ defines a primary metabolic alkalosis.

Step 4: If respiratory…… acute and /or chronic And Is metabolic compensation appropriate?

If it is a primary respiratory disturbance, Is it acute? And/or Chronic
For 10 mm change in pCO2
pH changes as
Acidosis (↑CO₂) pH ↓ acute by 0.08
Chronic by 0.03
Alkalosis (↓CO₂) pH ↑ acute by 0.08
Chronic by 0.03
HCO₃ Compensates as
Acidosis (↑CO₂) HCO₃↑ acute by 1
Chronic by 3  
Alkalosis (\(\text{\(\mathrm{CO}_2\)}\) \(\text{\(\mathrm{HCO}_3\)}\) \& acute by 2  
Chronic by 5  
For example,  
In an acute respiratory acidosis, if the \(\text{\(\mathrm{PCO}_2\)}\) increases from 40 to 50, you would expect the \(\text{pH}\) to decrease from 7.40 to 7.32.  
In an acute respiratory alkalosis, if the \(\text{\(\mathrm{PCO}_2\)}\) falls from 40 to 30, you would expect the \(\text{pH}\) to increase from 7.40 to 7.48.  
In chronic respiratory disturbances, there are renal mediated shifts of bicarbonate that alter and partially compensate for the \(\text{pH}\) shift for a change in the \(\text{\(\mathrm{PaCO}_2\)}\).  
In a chronic respiratory acidosis, if the \(\text{\(\mathrm{PCO}_2\)}\) increases from 40 to 50, you would expect the \(\text{pH}\) to decrease from 7.40 to 7.37.  
In a chronic respiratory alkalosis, if the \(\text{\(\mathrm{PCO}_2\)}\) decreases from 40 to 30, you would expect the \(\text{pH}\) to increase from 7.40 to 7.43.  

Remember: Keep in mind to suspect if  
• Compensated \(\text{\(\mathrm{HCO}_3\)}\) is > expected: additional metabolic alkalosis is there  
• Compensated \(\text{\(\mathrm{HCO}_3\)}\) is < expected: additional metabolic acidosis is there  

**Compensation limits**  
1. Compensatory \(\text{\(\mathrm{CO}_2\)}\) can go as high as 60 and as low as 10 for metabolic disorders  
2. Compensatory \(\text{\(\mathrm{HCO}_3\)}\) can go as high as 40 and as low as 10 for respiratory disorders  
AND respiratory compensation is faster; takes min to hours to compensate whereas metabolic compensation is slower, may takes days and even week time.  
• Respiratory disorders are better compensated than metabolic ones.  
• Most predictable compensation occurs in Metabolic acidosis.  
• Highly Un-predictable compensation occurs in metabolic alkalosis.  

**Step 5: If metabolic, Is respiratory compensation appropriate? Anion gap increased and/or normal or both?**  
If it is a primary metabolic disorder then is the respiratory compensation adequate or not?  
For metabolic acidosis: Expected \(\text{\(\mathrm{PCO}_2\)} = (1.5 \times [\text{\(\mathrm{HCO}_3\)}]) + 8 + 2\) (Winter’s formula) or  
Expected \(\text{\(\mathrm{CO}_2\)}\) is equal to last two digits of \(\text{pH}\) important and easy to remember.  

For metabolic alkalosis:  
Expected \(\text{\(\mathrm{PCO}_2\)} = 6\) mm for 10 mEq. rise in \(\text{\(\mathrm{HCO}_3\)}\).  

**Uncertain Compensation**  
Remember: Keep in mind to suspect if  
• Compensated \(\text{\(\mathrm{PCO}_2\)}\) is > expected: additional respiratory acidosis is there.  
• Compensated \(\text{\(\mathrm{PCO}_2\)}\) is < expected: additional respiratory alkalosis is there.  

Processes that lead to a metabolic acidosis can be divided into those with an increased anion gap and normal anion gap. The anion gap is the difference between the measured serum cations (positively charged particles) and the measured serum anions (negatively charged particles). (Of course, there is no real gap; in the body the number of positive and negative charges is balanced according to law of electro neutrality. The gap refers to the difference in positive and negative charges among cations and anions which are commonly measured). The commonly measured cation is sodium. (Some people also use potassium to calculate the gap; that results in a different range of normal values, and we will not use potassium to calculate the gap subsequently). The measured anions include chloride and bicarbonate. Thus the anion gap can be summarized as:  
\[
AG = [\text{Na}^+] - ([\text{HCO}_3^-] + [\text{Cl}^-]).
\]

The normal anion gap is 12-18. An increased or even normal anion gap helps for differential diagnosis of a metabolic acidosis. Metabolic acidosis can be normal anion gap (hyperchloremic), and High Anion Gap (normochloremic). Essentially \(\text{HCO}_3^-\) loosing disease like e.g. diarrhea, Renal tubular Acidosis causes Hyperchloremic normal anion gap metabolic acidosis, whereas \(\text{HCO}_3^-\) consuming diseases like e.g. Lactic acidosis, Ketoacidosis, renal failure etc. causes Normochloremic High Anion Gap metabolic acidosis. Various mnemonics exist for the causes of an increased anion gap metabolic acidosis; one of which is MULEPAK. The most common etiologies of a metabolic acidosis with an increased anion gap include:  
- Methanol other alcohols, and ethylene glycol intoxication  
- Uremia (renal failure)  
- Lactic acidosis  
- Ethanol  
- Paraldehyde and other drugs  
- Aspirin  
- Ketones (starvation, alcoholic and diabetic ketoacidosis)  
- Keypoint: The true anion gap is underestimated in hypoalbuminemia (=fall in unmeasured anions); AG must be adjusted. Remember to adjust AG:  
  
  For every 1 g/dl reduction in plasma albumin, anion Gap goes down by 2.5  

**Beware:** A normal anion gap is an increased one in hypoalbuminemia cases though it appears normal numerically.  
Simultaneous analysis of Anion Gap and \(\text{\(\mathrm{HCO}_3\)}\) it’s easy  
It is very helpful in diagnosing mixed disorders. It is determined by Rise and Fall of Anion gap and \(\text{\(\mathrm{HCO}_3\)}\) level from its base level  

**Rationale:**  
For each unit Rise in AG: (above normal level), \(\text{\(\mathrm{HCO}_3\)}\) should Fall by one unit: (below normal level)  
("Normal” values: AG = 12, \(\text{\(\mathrm{HCO}_3\)} = 24\) )
For e.g.:
Is more than one Disorder present?
Proper Clinical history
pH normal, and PCO$_2$ and HCO$_3$ out of range
HCO$_3$ and PCO$_2$ moving in opposite directions
Degree of compensation for primary disorder is inappropriate.
Look at Rise of anion gap and Fall of HCO$_3$.
How do I assess the correctness or laboratory error.
With help of H-H formula it is easy to diagnose,
For example: A report in which pH = 7.30, HCO$_3$ = 30, CO$_2$ = 38.1
though its acidosis neither metabolic nor respiratory since their
changed culprit values do not correlate. But

By Henderson-Hasselbach equation:

\[
H^+ = 24 \times \frac{pCO_2}{HCO_3} = 24 \times \frac{38/30} = 30
80 - \text{Last two digit pH} = H^+
80 - H^+ = \text{last two digit pH (after 7)}
\]

pH should have been 7.50

Ten Commandments of ABG:

1. I shall use only minimal amount of heparin to rinse the syringe. (Heparin changes pH and if excess dilutional effect)
2. I shall always do ALLEN’S test for collateral circulation and, also ensure that the sample sent is arterial and not venous one.
3. I shall ensure there are no air bubbles in the blood.

4. I shall send the sample in ice and analyze it immediately, and keep the total leucocyte count in mind, esp. when there is a delay.
5. I shall always take FiO$_2$ into consideration when interpreting PO$_2$ values. I shall also look at the PCO$_2$ values with care.
6. I shall take the history into consideration before instituting therapy for chronic respiratory failure.
7. I shall always remember the acronym "DOPE" in situations of sudden deterioration of ABG values
   D- Displacement
   O- Obstruction
   P- Pneumothorax
   E- Equipment failure
8. I shall practice gentle mechanical ventilation and not try to bring ABG to perfect normal.
9. I shall treat the patient not the ABG report
10. I shall always correlate ABG report clinically.

References
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