

Let's Talk About Gas: ABG Interpretation Made Easy



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Disclosures

• None.

Objectives

At the end of this presentation, the learner should be able to:

- 1. Analyze an ABG to determine the primary acid-base disorder.
- 2. Recognize compensation and mixed acid-base disorders.
- 3. Formulate differential diagnoses for the various acidbase disorders.
- 4. Apply ABGs to clinical scenarios and decision making.

Henderson-Hasselbalch Equation

$$pH = pK_a + log_{10} [A-] \xrightarrow{PASE} ACID$$

- The ratio of base-acid must stay relatively constant for the pH to stay constant
 - Because this is a logarithmic equation, it takes a fairly large change in acid or base to change the pH

Acid-Base Physiology



Components of the ABG

Component	Measure
рН	Acid-base balance
PaO ₂	Partial pressure of oxygen
PaCO ₂	State of alveolar ventilation
HCO ₃ -	Reflects metabolic component of blood
Alveolar-arterial (A-a) gradient	Gradient between alveolar and arterial oxygen
Base excess*	The amount of acid or base it would take to return the pH back to 7.4

Indications for ABG

- Identify respiratory, metabolic, and mixed acid-base disorders
- Monitoring acid-base status in disorders such as DKA
- Quantification of oxyhemoglobin and oxygen carrying capacity of the patient
- Quantification of levels of dyshemoglobins (methemoglobin, carboxyhemoglobin)

Indications for ABG

- Measuring partial pressures of respiratory gases involved in ventilation and perfusion, e.g....
 - COPD exacerbation
 - Asthma exacerbation
 - Pulmonary embolism
 - Pulmonary fibrosis
 - Pneumothorax
 - In these cases, can measure severity and progression of disease/exacerbation
- Assessment of response to mechanical ventilation

Some Clinical Scenarios Where an ABG is Useful in Acute Care

- Respiratory distress
- Hypoxia
- Airway obstruction
- Sepsis or shock
- DKA
- Renal failure
- Drug overdose or intoxication

- AMS or obtundation
- Monitoring response to invasive and noninvasive ventilation
- Code blue

Contraindications to ABG

- Local infection
- Distorted anatomy
- Abnormal Allen test
- AV fistula
- Severe PVD
- Relative contraindications:
 - Anticoagulation, tPA, severe coagulopathy
- Consider a-line if repeat ABGs will be necessary!



Before You Get an ABG

- Ask yourself...
 - What am I looking for/expecting?
 - Will this help guide or change my management?
 - How?
 - Will comparison be helpful after treatment/intervention?
 - Baseline ABG?

ABG Normal Values

Component	Low Normal	Normal Range	Estimated Normal	High Normal
рН	Acidosis	7.35 – 7.45	7.4	Alkalosis
PaO ₂	Hypoxemia	80 – 100 mmHg	100 – (0.3 x age)	Hyperoxia
PCO ₂	Respiratory alkalosis	35 – 45 mmHg	40 mmHg	Respiratory acidosis
HCO ₃	Metabolic acidosis	22 – 26 mEq/L	24 mEq/L	Metabolic alkalosis
A-a gradient		< 10 mmHg	< 10 mmHg	
Base excess		-3 to +3 mEq/L		

ABG vs. VBG

- Venous blood gases (VBG) are widely used in the emergency setting
 - There is no data to confirm that this level of agreement is maintained in shock states or mixed acid-base disturbances

• Why get a VBG over an ABG?

How do ABG Values Compare to VBG

ABG	VBG
рН	pH + 0.035 units
pCO ₂	pCO ₂ + 5.7 mmHg *correlation dissociates in hypercapnia and shock
HCO ₃	HCO ₃ – 1.41 mmol/L
Base excess (BE)	BE + 0.089 mmol/L
Lactate	Does NOT correlate > 2mM
pO ₂	Does NOT correlate (venous vs. arterial sample)

Clinical Application

Finally...something that makes sense!

Why Do We Care?

- What do acidosis and alkalosis look like?
- What do they imply?
- How dangerous are they?
- What disturbances can they cause in the body?

Respiratory Acidosis

"I can't catch my breath!"

- Rapid, shallow breaths OR bradypnea
- Dyspnea
- Headache
- Disorientation
- Dizziness
- Drowsiness



 $\downarrow pH \uparrow pCO_2$

Retention of CO₂ by the lungs

- Hypoxia
- Hyperkalemia
- Dysrhythmias
- Hyperreflexia
- Muscle Weakness

Metabolic Acidosis

- Confusion
- Drowsiness
- Headache
- Nausea, Vomiting
- Diarrhea



- Decreased BP
- Hyperkalemia
- Vasodilation
- Kussmaul respirations



Decreased ability of kidneys to excrete acid

Clinical Consequences of Acidemia

Cardiovascular	 Impaired cardiac contractility Decreased CO Decreased systemic BP Increased PVR Decreased threshold for arrhythmia
Respiratory	Hyperventilation (compensatory) with possible muscle fatigue
Neurologic	Obtundation/coma
Metabolic	 Insulin resistance Loss of bone and muscle Protein degradation Abnormalities in the release of many hormones Hyperkalemia Inhibition of anaerobic glycolysis Reduction in ATP synthesis

Respiratory Alkalosis

- Rapid, deep breathing
- Lethargy
- Confusion
- Nausea, vomiting
- Numbness and tingling



- Light headedness
- Seizure
- Tachycardia
- Hyperventilation
- Hyperkalemia

↑pH ↓pCO₂

Loss of CO₂ from the lungs

Metabolic Alkalosis

- Restlessness \rightarrow lethargy
- Confusion
- Dizziness
- Irritability
- Nausea, vomiting
- Diarrhea



Tremors

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- Muscle cramping
- Tingling in fingers/toes
- Hypokalemia
- Tachycardia



A decrease in acid or an increase in base

Clinical Consequences of Alkalemia

Cardiovascular	 Reduced coronary blood flow Reduced threshold for angina Decreased threshold for arrhythmia
Respiratory	 Hypoventilation (compensatory) with hypercapnia and hypoxemia
Neurologic	SeizureTetanyLethargy/delirium/stupor
Metabolic	 Hypokalemia Hypomagnesemia Hypophosphatemia Stimulation of anaerobic glycolysis Decreased oxyhemoglobin dissociation

Acid-base Differentials

Causes of Respiratory Acidosis

- ↓Respiratory stimuli
 - Drug overdose
 - Anesthesia
 - Intracranial issues (TBI, Stroke, SAH, etc.)
- Spinal or peripheral nerve issues (C-spine injury, myasthenia gravis, ALS, peripheral neuropathy, etc.)
- Pulmonary issues (ARDS, asthma, pneumonia, COPD, massive PE/Shock, atelectasis)
- Extrathoracic/abdominal distention (obesity, compartment syndrome)

Causes of Metabolic Acidosis

MUDPILES

- Methanol
- Uremia
- DKA
- **P**ropylene glycol (Ativan, Dilantin)
- Isoniazid/Iron
- Lactate
- Ethanol/Ethylene glycol
- Salicylates / Seizures / Starvation

USEDCRAP

- Ureteroenterostomy
- Small bowel fistula
- Excess chloride*
- Diarrhea
- Carbonic Anhydrase Inhibitors
- Renal Tubular Acidosis (RTA)
- Addison's disease / Acetazolamide
- Pancreatoenterostomy

A Word About Chloride...

Fluid	Na	CI	к	Mg	Ca	HC O3	Gluc	A c	Gluc	Osm	рН
Plasma	140	104	4.5	1.25	2.5	24	0.08			290	7.4
0.9% Nacl	154	154								308	5.5
0.45% Nacl	77	77								406	
LR	130	109	4		1.5	28 (as Lac)				273	6.5
P-lyte	140	98	5	1.5				2 7	23	294	7.4
5% dex							5			278	

Causes of Respiratory Alkalosis

- Pain
- Trauma
- Sepsis
- Pulmonary embolism
- Shock

- Drugs
- Pregnancy
- Hyperventilation (think sick first!)
- Mechanical ventilation

Causes of Metabolic Alkalosis



Let's do the math...

ABG Analysis/Calculations

ABG Analysis Big Picture

- 1. Look at the **pH** to determine primary disorder
- Look at PCO₂, use in conjunction with pH to determine primary disorder
 - If pH and PCO₂ move in *opposite* directions, respiratory disorder is primary
 - If pH and PCO₂ move in the same direction, metabolic disorder is primary

ABG Analysis Big Picture

- 3. Look for mixed disorder
 - If both pCO_2 and HCO_3 are $\uparrow \uparrow$ = respiratory acidosis OR metabolic alkalosis
 - If both pCO_2 and HCO_3 are $\downarrow \downarrow$ = respiratory alkalosis OR metabolic acidosis
- 4. Apply Compensation rules
 - Boston rule
 - Winter's formula
- 5. Look at Calculated/Corrected Anion Gap and Delta Gap
- 6. Clinical Application

*paO2 in ABG tells us about the partial pressure of oxygen in blood. It is not used in assessment of acid-base disorders.

ABG Practice: Steps 1-2

- pH low (<7.4) = Acidosis, pH high (>7.4) = Alkalosis
- If pH and PCO₂ move in *opposite* directions, respiratory disorder is primary
- If pH and PCO₂ move in the same direction, metabolic disorder is primary

рН	PCO ₂	HCO ₃	Disorder
7.2	70	28	Respiratory Acidosis
рН	PCO ₂	HCO ₃	Disorder
7.2	30	16	Metabolic Acidosis

ABG Practice: Steps 1-2

рН	PCO ₂	HCO ₃	Disorder
7.5	40	31	Metabolic alkalosis

рН	PCO ₂	HCO ₃	Disorder
7.5	30	16	Respiratory alkalosis

Let's add in the next layer of complexity...

Step 3: Look for mixed disorders

ABG Practice: Step 3

- If both pCO_2 and HCO_3 are $\uparrow \uparrow$ = respiratory acidosis OR metabolic alkalosis
- If both pCO_2 and HCO_3 are $\downarrow \downarrow$ = respiratory alkalosis OR metabolic acidosis
- If pCO_2 and HCO_3 move in opposite direction $\uparrow \downarrow$ = mixed disorder is present

рН	PCO ₂	HCO ₃	Disorder
7.28	55	19	Primary respiratory acidosis, metabolic acidosis

- 1. Look at pH \rightarrow acidosis
- 2. Look at pH and $PCO_2 \rightarrow opposite$, respiratory disorder is primary
- 3. Look at pCO_2 and $HCO_3 \rightarrow$ opposite, mixed disorder present \rightarrow HCO_3 low, implies metabolic acidosis also present

ABG Practice: Step 3

рН	PCO ₂	HCO ₃	Disorder
7.50	24	28	Respiratory alkalosis primary, metabolic alkalosis

- 1. Look at pH \rightarrow alkalosis
- 2. Look at pH and PCO₂ \rightarrow opposite, respiratory disorder is primary
- 3. Look at pCO₂ and HCO₃ \rightarrow opposite, mixed disorder present \rightarrow HCO₃ high, implies metabolic alkalosis also present
Let's add another layer

Step 4: Compensation

Compensation Physiology

- Respiratory system: maintains pH by regulating CO₂
 - Can compensate quickly
- Renal (metabolic) system: regulates pH by excreting H⁺ (acid) or reabsorbing HCO₃⁻ (base)
 Several hours to days to compensate
- With the body's ACUTE compensation, the pH will NOT return to normal, but it may get close

Buffering Systems/Compensation



Buffering Systems/Compensation



Acid-base Nomogram



Compensation

Acid-Base Disorder	Primary Change	Compensatory Change
Respiratory acidosis	$PCO_2\uparrow$	HCO ₃ ↑
Respiratory alkalosis	$PCO_2\downarrow$	$HCO_3\downarrow$
Metabolic acidosis	$HCO_3\downarrow$	$PCO_2\downarrow$
Metabolic alkalosis	HCO ₃ ↑	PCO ₂ ↑

Acid-Base Disorder	Primary Change	Compensa tory Change
Respiratory acidosis	PCO_2 \uparrow	$HCO_3 \uparrow$
Respiratory alkalosis	$PCO_2\downarrow$	$HCO_3 \downarrow$
Metabolic acidosis	$HCO_3\downarrow$	$PCO_2\downarrow$
Metabolic alkalosis	$HCO_3\uparrow$	$PCO_2\uparrow$

ABG Practice: Step 4

рН	PCO ₂		Primary Disorder
7.28	60	35	Respiratory acidosis

- 1. pH: Acidosis
- pCO₂ and pH move in opposite directions → primary respiratory disorder
- 3. pCO_2 and HCO_3 both high \rightarrow respiratory acidosis
- 4. Expected change with respiratory acidosis \rightarrow HCO₃ increases

Acid-Base Disorder	Primary Change	Compensa tory Change
Respiratory acidosis	$PCO_2 \uparrow$	$HCO_3 \uparrow$
Respiratory alkalosis	$PCO_2\downarrow$	$HCO_3 \downarrow$
Metabolic acidosis	$HCO_3\downarrow$	$PCO_2\downarrow$
Metabolic alkalosis	$HCO_3\uparrow$	$PCO_2 \uparrow$

ABG Practice: Step 4

рН	PCO ₂		Primary Disorder
7.5	55	36	Metabolic alkalosis

- 1. pH: Alkalosis
- 2. pCO_2 and pH move in same direction \rightarrow primary metabolic disorder
- 3. pCO_2 and HCO_3 are high = metabolic alkalosis
- 4. Expected change with metabolic alkalosis \rightarrow PCO₂ increases

You're experts! Let's add on...

Calculating for compensation

Calculating for Compensation

- Respiratory Rule #1: pH changes INVERSELY by 0.08 for 10 mm CO₂ in ACUTE cases
 - If CO₂= 50, pH will be 7.32 (0.08 below)
 - If CO₂=30, pH will be 7.48 (0.08 above)

*DO NOT USE IN CHRONIC CASES...pH usually corrects/compensates to normal (7.4).

Calculating for Compensation

Boston Rules (to predict changes in HCO_3^- from $PaCO_2^-$ respiratory disorders):

Change in CO2	Change in HCO3	Condition	Example
10	1	Acute Resp Acidosis	If CO2=50,
			HCO3=25
10	2	Acute Resp Alkalosis	If CO2=30,
			HCO3=22
10	4	Chronic Resp Acidosis	If CO2=50,
			HCO3=28
10	5	Chronic Resp Alkalosis	If CO2=30,
			HCO3=19

Calculating for Compensation

Winter's Formula (metabolic acidosis):
Expected pCO₂ = 1.5 x HCO₃ + 8 +/- 2

Metabolic Alkalosis
Expected pCO₂ = 0.7 x [HCO₃⁻] + 20 (+/- 5)

One last concept(s)...

Step 5: Look at Calculated/Corrected Anion Gap and Delta Gap

Anion Gap (Calculated)

 Anion Gap = the difference between measured cations (Na+, K+) and measured anions (CI- and HCO₃⁻)

= Na - [Cl + HCO₃]

Normal range = 4 – 12 mmol/L

*We call this the calculated anion gap (cAG), because it is calculated from the CMP

- Significance?
 - The law of electrochemical neutrality
 - Provides foresight into managing acidosis

High vs. Normal Anion Gap Metabolic Acidosis



High Anion Gap Metabolic Acidosis (HAGMA)



- Accumulation of or impairment of excretion of acids
- MUDPILES:
 - Methanol
 - Uremia
 - DKA
 - Propylene glycol (Ativan, Dilantin)
 - Isoniazid/Iron
 - Lactate
 - Ethanol/Ethylene glycol
 - Salicylates/Seizures/Starvation

Normal Anion Gap Metabolic Acidosis (NAGMA)

- Mainly from losses of bicarb (HCO₃-)
- USEDCRAP (earlier slide) OR
- ABCDE
 - Addison's
 - Bicarbonate loss (GI or renal think v/d, fistula, ostomy)
 - Chloride excess*
 - Diuretics (acetazolamide)*
 - Extra Renal tubular acidosis (RTA)*

Anion Gap (corrected)

- Corrected Anion Gap (corrAG) = takes into account the unmeasured anions (albumin, sulfate, phos), which may change with acute or chronic illness
- = [2 x Albumin] + [0.5 Phosphate] (+/- 2) OR [3 x Albumin] Normal range = 8 – 12 mmol/L
- Significance?
 - If calcAG > corrAG, there is <u>high gap metabolic</u> acidosis present

Delta Gap

• *Check in the presence of HAGMA to determine if pure HAGMA or coexisting disorder present*

- Delta Gap = $[calcAG corrAG] + HCO_3$
 - Net sum = 24
 - HAGMA only
 - Net sum < 24
 - NAGMA (non-anion gap acidosis) is present
 - Net sum > 24
 - METABOLIC ALKALOSIS is present

Practice: Minding the Gaps

<u>ABG:</u>	
рН	7.18
PCO_2	34 mmHg
HCO_3	12 mEq/L

<u>BMP:</u> Na= 138, K=3.8, CI=115 Albumin=2.3, Phos=1

calcAG = 138 – [115+12] = **11**

corrAG = 2 x 2.3 + [0.5 x1] = **5.1 (+/- 2)**

calcAG > corrAG → HAGMA present

Delta Gap = $[calcAG - corrAG] + HCO_3 = 6 + 12 = 18$

18 < 24...therefore, NAGMA present also

Let's practice...

A 26 YO M with asthma presents to the ED with difficulty breathing x 3 days. It is getting progressively worse. He has tried his regular and rescue inhalers; nothing seems to help. He looks pale and is taking rapid, shallow breaths. On exam, he has diffuse wheezing in all lung fields.

Vitals:

- HR 120
- BP 113/76
- RR 28
- $SpO_2 92\%$
- Temp 37.8C

ABG results:

- pH 7.08
- $pCO_2 80 \text{ mmHg}$
- $HCO_3^- 28 \text{ mEq/L}$

Other results (pertinent):

- Na 138 mEq/L (135-145)
- K 4.0 mmol/L (3.6-5.2)
- CI 106 mEq/L (96-106)
- Albumin 3.8 g/dL (3.5-5.5)
- Phos 3.0 mg/dL (2.8 4.5)

Acute Respiratory Acidosis

ABG results:

- pH 7.08
- pCO₂ 80 mmHg
- HCO₃⁻ 28 mEq/L

Compensation rules...

- Respiratory rule #1
- pCO₂ ↑ by 40 (10x4)
- Expected pH change = $0.08 \times 4 = 0.32$
- Actual pH change = 7.4 7.08 = 0.32

ABG results:

- pH 7.08
- pCO₂ 80 mmHg
- HCO₃⁻ 28 mEq/L

Other results (pertinent):

- Na 138 mEq/L (135-145)
- K 4.0 mmol/L (3.6-5.2)
- CI 106 mEq/L (96-106)
- Albumin 3.8 g/dL (3.5-5.5)
- Phos 3.0 mg/dL (2.8 4.5)

- calcAG = 138- [106+28] = 4
- corrAG = [2x3.8]+[0.5x3] = 9.1
- calcAG < corrAG = no additional HAGMA
- Delta gap = not needed

Acute Respiratory Acidosis

Case 1 DDx + Management

- Asthma exacerbation \rightarrow rapid, shallow breaths \rightarrow retaining CO₂ \rightarrow Acute Respiratory Acidosis
- Management
 - Reverse the respiratory acidosis, augment breathing to help improve gas exchange until the patient improves and can do so on their own
 - How?
 - When would you get a follow-up ABG?

A 35 YO M presents to the ED with gun shot wound (GSW) to the abdomen. He was found down by a civilian about 20 minutes after the shooting who called 911. Upon arrival, he appears pale, diaphoretic, and is experiencing severe abdominal pain. He is slightly altered and cannot tell you where he is. He has no past medical history.

Vitals:

- HR 116
- BP 86/68
- RR 10
- $SpO_2 96\%$
- Temp 37.6C

ABG results:

- pH 7.18
- $pCO_2 34 \text{ mmHg}$
- $HCO_3^- 12 \text{ mEq/L}$

Other results (pertinent):

- Na 132 mEq/L (135-145)
- K 3.6 mmol/L (3.6-5.2)
- CI 92 mEq/L (96-106)
- Albumin 3.2 g/dL (3.5-5.5)
- Phos -2.1 mg/dL (2.8 4.5)

Acute Metabolic Acidosis

ABG results:

- pH 7.18
- $pCO_2 34 \text{ mmHg}$
- HCO₃⁻ 12 mEq/L

Compensation rules...

- Winter's formula (metabolic acidosis)
 - Expected PCO₂ in metabolic acidosis:

= 1.5 x HCO₃ + 8 = **26 (+/- 2)**

 Expected pCO₂ is lower than our actual... what does this mean?

- The expected degree of respiratory compensation is not present...
- There is also a respiratory acidosis

- Respiratory depression/AMS → slower, possibly more shallow breaths → retention of CO₂ → respiratory acidosis
 - Normally, the body would begin breathing more quickly to blow off CO₂ and help compensate for the metabolic acidosis...

ABG results:

- pH 7.18
- pCO₂ 34 mmHg
- HCO₃⁻ 12 mEq/L

Other results (pertinent):

- Na 132 mEq/L (135-145)
- K 3.6 mmol/L (3.6-5.2)
- CI 92 mEq/L (96-106)
- Albumin 3.2 g/dL (3.5-5.5)
- Phos 2.1 mg/dL (2.8 4.5)

- calcAG = 132 [92+12] = 28
- corrAG = [2x3.2]+[0.5x2.1] = 7.45
- calcAG>corrAG = HAGMA present (we already know this)
- Delta gap = [28-7.45]+12 = 32.55
- Delta gap > 24 → metabolic alkalosis also present

HAGMA + Respiratory Acidosis + Metabolic Alkalosis

Case 2 DDx

HAGMA

- MUDPILES:
 - Methanol
 - Uremia
 - DKA
 - Propylene glycol (Ativan, Dilantin)
 - Isoniazid/Iron
 - Lactate
 - Ethanol/Ethylene glycol
 - Salicylates/Seizures/Starvation

Respiratory Acidosis

- ↓respiratory stimulus
- Atelectasis
- Additional injuries?

Metabolic Alkalosis

- GI acid loss
 - Vomiting/NG drainage
 - Diarrhea
 - Ostomy
 - Dehydration
- Renal acid loss
 - Bartter, Gitelman syndrome
 - Diuretics
- Added base
 - Laxatives, milk-alkali syndrome, antacid OD
 - latrogenic bicarb administration
- Endocrine
 - Cushing
 - Steroid excess
 - Hyperaldosteronism

Case 2 Management

- You obtain a lactate level to confirm your suspicion... = 6.2 mmol/L (<2.3).
- His renal function is normal...for now.
- His H/H is 8 g/dL / 26% (13.2-16.6 / 38.3-48.6%).

Case 2 Management

 Acute blood loss → hypovolemic shock → decreased organ perfusion and O2 delivery → interference with aerobic metabolism → increased anaerobic metabolism → production of lactic acid and metabolic acidosis

- How are you going to manage this patient?
- How did the ABG help us in this scenario?
- Will you get a repeat ABG? If so, when?

A 64 YO M with ESRD s/p kidney transplant, Type 2 DM, and chronic HFrEF presents to the ED with 3 days of fatigue, abdominal pain, and shortness of breath. He is unsure, but he may have had a fever. On exam, he appears unwell and has crackles in the L lung base. He missed dialysis today. CXR shows a L basilar infiltrate.

Vitals:

- HR 100
- BP 92/78
- RR 20
- SpO₂ 84%
- Temp 38.0

ABG results:

- pH 7.29
- pCO₂23.3
- HCO₃⁻ 11.1

Acute Metabolic acidosis

• pO₂ 52.9

Other results (pertinent):

- Na 136
- K 5.2
- CI 106
- AG 18
- Glucose 260
- Albumin 2.8
- Phos 3.0
- BUN 89.1
- Cr 4.3
- Lactate 1.3
- Beta-hydroxybutyrate 2.7

ABG results:

- pH 7.29
- pCO₂23.3
- HCO₃⁻ 11.1
- pO₂ 52.9

Compensation rules...

Winters formula

Expected $pCO_2 = 1.5 \times 11 + 8 = 24.5 (+/-2)$

Actual $pCO_2 = 23.3$

ABG results:

- pH 7.29
- pCO₂ 23.3
- HCO₃⁻ 11.1
- pO₂ 52.9

Other results (pertinent):

- Na 136
- K 5.2
- CI 106
- Glucose 260
- Albumin 2.8
- Phos 3.0
- BUN 89.1
- Cr 4.3
- Lactate 1.3
- Beta-hydroxybutyrate 2.7

- calcAG = 136 [106+11] = 19
- corrAG = [2x2.8] + [0.5x3.0] = 7.1
- calcAG>corrAG = HAGMA
 present (we already know this)
- Delta gap = [19-7] + 11 = 23
- Delta gap < 24 = NAGMA also present

Case 3 DDx

<u>MUDPILES = HAGMA</u>

- Methanol
- Uremia
- DKA
- **P**ropylene glycol (Ativan, Dilantin)
- Isoniazid/Iron
- Lactate
- Ethanol/Ethylene glycol
- Salicylates / Seizures / Starvation

<u>ABCDE = NAGMA</u>

Addison's

- Bicarbonate loss (GI or renal think v/d, fistula, ostomy)
- Chloride excess
- Diuretics (acetazolamide)
- Extra Renal tubular acidosis (RTA)
Case 3 Management

- How are we going to correct this patient's HAGMA and NAGMA ?
- 1. Treat DKA
- 2. Dialysis...add bicarb
- 3. Treat underlying cause of DKA...what was going on with this patient?

Are you going to get another ABG? If so, when?

A 23 YO F returns from a volunteer humanitarian trip to Haiti. She presents to her PA with nausea & vomiting x2 days. The vomiting is becoming more frequent, and she can't keep down any liquids. On exam, she appears significantly dehydrated and her respirations are frequent but shallow.

Vitals: Other results (pertinent): **ABG results:** HR 102 pH 7.56 Na – 132 • • $pCO_2 40 \text{ mmHg}$ K - 3.4BP 90/64 • • $HCO_3^- 32 \text{ mEq/L}$ **RR 20** CI - 86• SpO₂ 99% Albumin -2.3• Temp 37.6 C Phos - 2.0Acute Metabolic Alkalosis Glucose – 86 BUN – 32

Cr - 1.0

Lactate - 1.8

ABG results:

- pH 7.56
- pCO₂ 40 mmHg
- HCO₃⁻ 32 mEq/L
- Compensation rules...
- Expected $pCO_2 = 0.7 \times [32] + 20 = 42.4 (+/-5)$
- Actual $pCO_2 = 40$

ABG results:

- pH 7.56
- pCO₂ 40 mmHg
- HCO₃⁻ 32 mEq/L

Other results (pertinent): Na - 132 K - 3.4

CI - 86Albumin - 2.3 Phos - 2.0 Glucose - 86 BUN - 32 Cr - 1.0

Lactate – 1.8

- calcAG = 132 [86+32] = 14
- corrAG = [2x2.3]+[0.5x2.0] = 5.6
- calcAG>corrAG = HAGMA present
- Delta gap = [14-5.6] + 32 = 40.4
- Delta gap > 24 = Metabolic Alkalosis (we already know this)

Acute Metabolic Alkalosis + HAGMA

Case 4 DDx



Case 4 DDx

MUDPILES = HAGMA

- Methanol
- Uremia
- DKA
- Propylene glycol (Ativan, Dilantin)
- Isoniazid/Iron
- Lactate
- Ethanol/Ethylene glycol
- Salicylates / Seizures / Starvation

Beta-hydroxybutyrate = 2.3 (↑)

Case 4 Management

- How are we going to manage this patient?
 - Fluids
 - Antiemetics \rightarrow nutrition
 - Look for infectious causes

Does the ABG change our course of action?

• Will you get a follow up ABG? If so, when?

A 68 YO F presents to the ED with 3 days of progressively worsening cough and shortness of breath. She has been experiencing intermittent fevers. Her appetite is diminished, and she is fatigued. On exam, she has scattered crackles. CXR reveals a multifocal pneumonia.

Vitals:	ABG Results:	Other results (pertinent):
HR 96	• pH 7.55	Na – 134
BP 112/82	• pCO ₂ 22 mmHg	K – 4.2
RR 26	• HCO_{3}^{-1} 16 mEq/L	CI – 100
SpO ₂ 84%		BUN – 32
Temp 37.9 C	Acute Respiratory Alkalosis	Cr – 1.2
· · · · · · · · · · · · · · · · · · ·		Albumin – 3.8
		Phos – 3.0

Lactate -3.2

ABG Results:

- pH 7.55
- pCO₂ 22 mmHg
- HCO₃⁻ 16 mEq/L

Case 5

Compensation rules...Boston Approach for Resp Disorders

Change in CO2	Change in HCO3	Condition	Example
10	1	Acute Resp Acidosis	If CO2=50,
			HCO3=25
10	2	Acute Resp Alkalosis	If CO2=30,
			HCO3=22
10	4	Chronic Resp Acidosis	If CO2=50,
			HCO3=28
10	5	Chronic Resp Alkalosis	If CO2=30,
			HCO3=19

 pCO_2 went from 40 \rightarrow 22 (roughly 20)

We expect HCO₃ to decrease by 4 (using 24 baseline), therefore expecting HCO₃ to be $20 \rightarrow$ there is compensation, and potentially another unaccounted-for process

Is there a mixed disorder present?

ABG Results:

• pH 7.55

Lactate -3.2

- pCO₂ 22 mmHg
- HCO₃⁻ 16 mEq/L

Other results (pertinent): Na - 134 K - 4.2 Cl - 100 BUN - 32 Cr - 1.2 Albumin - 3.8 Phos - 3.0

- calcAG = 134– [100+16] = 18
- corrAG = [2x3.8] + [0.5x3.0] = 9.1
- calcAG>corrAG = HAGMA present

• Delta gap = [18-9.1]+16 = 24.9

 Delta gap = 24.9 ...close enough! Only HAGMA present

Acute Respiratory Alkalosis + HAGMA

Case 5 DDx

 Hypoxic stimulation 2/2 pneumonia leads to hyperventilation to try to correct the hypoxia, at the expense of CO₂ loss → acute respiratory alkalosis

<u>MUDPILES = HAGMA</u>

- Methanol
- Uremia
- DKA
- **P**ropylene glycol (Ativan, Dilantin)
- Isoniazid/Iron
- Lactate
- Ethanol/Ethylene glycol
- Salicylates / Seizures / Starvation

Case 5 Management

- What is an appropriate management strategy to correct the respiratory alkalosis?
 - Supplemental oxygen
- How are we going to correct the metabolic acidosis?
 - Treat the infection!
 - Clear the lactate...ensure perfusion (? fluids)
- How did the ABG help us here?
- Would you get a repeat ABG? If so, when?

In Summary...

- Look at the pH first
 Do pH and PCO₂ change in the same direction?
- Use all components/calculations of the ABG
 - Don't skip the calculations...it may change your management!
- Apply clinically to your patient; don't just treat the numbers
 - Use your differentials...MUDPILES, USEDCRAP, etc.
- Consider what will happen if you start treatment
 - Will treating help or hurt?

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