### Acid-base chemistry and its disturbances

MUDr. Stanislav Matoušek, PhD

#### System of the presentation

- Difficult subject?
- Gradual steps → good understanding, building on what I already know
- Active learning:
  - Slide with questions to solve give it time, try to find solutions by yourself
  - Minimum time is below
  - Answers mostly next slide
- Why active learning?
  - Greater joy and interest (in the end)
  - Deeper knowledge
  - Remembering longer
  - → The extra effort pays off



### Acid-base Chemistry and Physiology Refresher (hopefully)

#### Hydrogen ions

- Is the concentration of hydrogen ions in extracellular fluid (ICF) small, large, huge or minuscule?
- Why is maintaining H<sup>+</sup> concentration within narrow limits much more important than maintaining strict concentrations of let's say iodine or zink<sup>1</sup>?
- Is it more accurate to speak of H<sub>3</sub>O<sup>+</sup> or H<sup>+</sup>? Why?
- What is a hydrogen bond (H bond)?
- Is there more H<sub>3</sub>O<sup>+</sup> or OH<sup>-</sup> in plasma under physiological conditions?
- Minimum time: 2 min

#### Hydrogen ions

- Concentration of H<sup>+</sup> = [H<sup>+</sup>] ~ **1 000 000x** << [Na<sup>+</sup>] minuscule
- Maintaining pH within tight limits is important because of very large reactivity of H<sup>+</sup> and its effect on the conformation of many macromolecules, especially proteins.
- Hydrogen bond special type of weak chemical bond created by H<sup>+</sup>; it binds H<sub>2</sub>0 molecules together →liquefaction of water
- $pH_{plasma,Norm} \approx 7,4 > 7,0 \rightarrow alkaline pH \rightarrow [OH^-] > [H3O^+]$

### Dynamics of H<sub>3</sub>O<sup>+</sup> and OH<sup>-</sup> movement in water



# Amino-acid charge and protein conformation







Glutamic acid:

 $-NH_2 \xrightarrow{}$ 

рК<sub>А</sub> =4,25

0=



– NH,



**Crucial organs:** 









#### pH definition and its consequences

- Would you remember how pH is defined?
- And what are rules of calculating with logarithms? E.g. log(A x B) =
- Try to figure out what these rules imply for the pH behavior: For instance, when H<sup>+</sup> concentration (denoted as [H<sup>+</sup>]) increases two times, how does pH change?<sup>1</sup>
- How does pH change, when the H<sup>+</sup> concentration decreases 10x?
- For straight-A students: How does pH change, when [OH<sup>-</sup>] increases 2x?
- Minimum time: 3 minutes or until completion of all tasks.

• 1) You might find it helpful to know that  $log_{10}(2) = 0.3$ 

# pH definition and its consequences - solution

- pH = log<sub>10</sub>([H<sup>+</sup>])
- log(AB) = log(A) + log(B)
- H<sup>+</sup> concentration increasing twice:  $[H^+]_{New} = 2[H^+]_{Old}$
- From the pH definition and the logarithm calculation rules:  $pH_{New} = -log([H^+]_{New}) = -log(2 \times [H^+]_{Old}) = -log(2) + (-log([H^+]_{Old}) = -0,3 + pH_{Old})$
- Therefore: If H<sup>+</sup> concentration doubles:  $pH_{New} = pH_{Old} 0.3$
- When  $[H^+]_{New} = 1/10 \times [H^+]_{Old} : pH_{New} = -\log(1/10) + (-\log([H^+]_{Old})) =$
- =  $+1 + pH_{Old}$ . If [H<sup>+</sup>] decreases ten times, pH goes up by 1.

#### **Buffers**

- What are buffers and what their effects in a solution are?
- How do buffers influence pH change, when acid or base is added?
- What does pK<sub>a</sub> of a simple buffer denote?
- For straight-A students: Can you write down the mass action equation of a simple buffer?
- Which pH does render a single substance buffer most effective?
- Minimum time: 3 minutes

#### Buffers – Solution 1

- Buffers inhibit the pH change by binding the extra H<sup>+</sup> when their concentration increases (when pH falls) and releasing H<sup>+</sup> when [H<sup>+</sup>] decreases (pH goes up).
- They are crucial for stabilizing pH within certain range!
- Hydrogen ion and buffer react according to the formula:

• HB 
$$\longrightarrow$$
 H<sup>+</sup> + B<sup>-</sup> or:

$$HB^+ \longrightarrow H^+ + B$$

 Reaction equilibrium concentrations can be expressed by the wellknown mass action formula:

$$K_A = \frac{[\mathrm{H}^+] * [\mathrm{B}_i^-]}{[\mathrm{HB}_i]}$$

• This can be expressed in the logarithmic form as well:

$$pH = pK_A + \log_{10} \frac{[B_i^-]}{[HB_i]}$$

 [HB] and [B<sup>-</sup>] are in 1:1 ratio, when pH = pK<sub>A</sub>

(prove this using the previous relationship)

#### Buffers – Solution 2

- Single-substance buffer is most effective when the pH coincides with its pK<sub>A</sub>.
- Efficiency of a buffer at a given pH can be measured by its buffer capacity β.
- When pH and pK<sub>A</sub> fall far, the efficiency of the buffer is constrained by the buffer component that is low in concentration.
- For instance, at acidic pH, there is  $\sqrt[]{B^-]}$ ,
- At alkalic pH, there is  $\downarrow$ [HB]
- For straight-A students: How does the [B<sup>-</sup>]/[HB] ratio change when acid is added at pH far from its pK<sub>A</sub>? Does it change a lot or a bit?



### **Protein Buffers**

- Principal buffers in blood are:
  - Hemoglobin!
  - Albumin, and other proteins of blood plasma

- Key buffer residues are histidine side chains.
- pK<sub>A</sub>'s of individual histidine side chains differ significantly (influence of surrounding aminoacids)



#### **Protein Buffers**



#### **Phosphate Buffer**

- Principal intracellular buffer
- Incl. phosphate residues of DNA
- The 2<sup>nd</sup> dissociation step is important, having pK<sub>A</sub> = 7.2





#### **Bicarbonate Buffer**

 $CO_2+H_2O \Longrightarrow H_2CO_3 \Longrightarrow H^+ + HCO_3^-$ 

Catalyzed by Carboanhydrase – present in stomach, kidneys and erythrocytes.

Equilibrium (mass action) expressed by the Henderson-Hasselbalch equation:

 $pH = pK_A + \log \frac{[HCO_3^-]}{s * pCO_2}$ 

Substituting numerical values (for pCO<sub>2</sub> in kPa) :

$$pH = 6,1 + \log \frac{[HCO_3^-]}{0,22 * pCO_2}$$

Question for straight-A students:  $pK_A$ of the bicarbonate buffer is 6.1. This is quite off the physiological pH = 7.4. Does this lower the buffer capacity of bicarbonate buffer?

#### **Bicarbonate Buffer**

Possible depiction of reaction equilibria (as described by Henderson-Hasselbalch equation):

 $CO_2+H_2O \longrightarrow H_2CO_3 \longrightarrow H^+ + HCO_3^-$ HCO<sub>3</sub>  $CO_2$ CO H<sub>2</sub>CO<sub>3</sub> HCO<sub>3</sub>-H<sup>+</sup> pH= 7,4 Magnifying glass: Concentrations in order of 10 nmol/L

#### Metabolism and the System of Regulating Acid-base Status

- The biggest turnover is in the system of CO<sub>2</sub>
  - Thus pCO<sub>2</sub> can be easily regulated.
     CO<sub>2</sub> behaves as an open system
- Other flows and the relationship with concentrations of other ions (electro-neutrality) are depicted in the figure:



#### **Bicarbonate Buffer**

- It is the principal buffer in terms of longer-term regulation of H<sup>+</sup> balance in the body.
- Lungs regulate pCO<sub>2</sub>
- Kidneys regulate the level of  $HCO_3^-$  in blood plasma and excretion of  $H^+$

	Primary Disturbance	When H+ = 40 nmol/L	Compensation
Respiratory Acidosis	↑pCO <sub>2</sub>	Reacts to the right -个H⁺	Kidneys - 个HCO <sub>3</sub> -, 个BE
Metabolic Acidosis	↓HCO <sub>3</sub> - (or 个个个H+ )	Reacts to the right when primary cause ↓HCO <sub>3</sub> <sup>-</sup> (Reacts to the left when primary cause ↑↑↑H <sup>+</sup> ) –end result - ↑H <sup>+</sup>	Lungs - ↓pCO <sub>2</sub>
Respiratory Alkalosis	↓pCO <sub>2</sub>	Reacts to the left - $\bigvee$ H <sup>+</sup>	Kidneys - $\downarrow$ HCO <sub>3</sub> <sup>-</sup> , $\downarrow$ BE
Metabolic Alkalosis	↑HCO <sub>3</sub> -	Reacts to the left - $\downarrow$ H <sup>+</sup>	Lungs - 个pCO <sub>2</sub>

#### Base Excess - BE

- Base Excess a very precise measure of metabolic disturbances (and metabolic compensations)
- Underlying logic: Lungs regulate pCO<sub>2</sub>. This regulation does not influence the total concentration of base forms of buffers.
- By definition: When pH = 7,4 (norm) and pCO<sub>2</sub> = 5,3 kPa (norm), then BE = 0 mmol/L
- Now, when we add 10mmol/L of acids, part of this amount reacts away with bicarbonate and part with the B- form of nonbicarbonate buffers – BE decreases to -10 mmol/L
- Conversely, taking away 15 mmol/L of acids (H<sup>+</sup>) when BE =0 increases both the level of bicarbonate and the B- of non-bicarbonate buffers – BE increases to +15 mmol/L
- The value of BE is independent of subsequent changes in pCO<sub>2</sub>

#### System of Buffers and Electroneutrality



#### **Electroneutrality:**

When buffer concentration changes – the concentration of strong ions has to change as well

X<sup>-</sup> - (ordinarily) unmeasured ions –e.g. lactate, keto<sup>-</sup>, SO<sub>4</sub><sup>2-</sup>

 $AG = Anion qap = Na^+ + K^+ - Cl^- - HCO_3^-$ 

Parameter used in differential diagnosis of metabolic acidoses

# System of Buffers and Electroneutrality 2 – Example







### Acid-base Balance Disturbances

= Pathophysiology of ABB

Respiratory Disturbances and their Compensation

#### Respiratory Alkalosis and its Compensation





#### **Compensation Diagrams 2**



# Compensation Diagram pCO<sub>2</sub> vs BE – Different



#### "Boston" Rules for Diagnosing ABB Disturbances For straight-A students - optional

Alternative to the compensation diagrams - however, you have to remember them 😕

For straight-A students - optional: This is for  $pCO_2$ expressed in mmHg – convert to a version in kPa ( $pCO_2$  40 mmHg = 5,3 kPa)



#### Respiratory Acidosis and its Compensation



Metabolic Disturbances and their Compensation

#### Metabolic Acidosis-Compensation Diagrams

Respiratory compensation generally develops faster than the metabolic disturbance itself, thus the division into acute and chronic disturbances is missing in diagrams focused more clinically.





# Metabolic Acidosis – Compensation Diagrams

Try to draw the beakers using your memory only and add corresponding arrows illustrating acute and chronic metabolic acidosis into the diagram (full respiratory compensation develops in about 10 h)



#### What is Taken and Assessed?

- Blood Gases Measurement in Arterial Blood (so called "Astrup")
- Serum electrolytes
- Concentrations of buffers (e.g. hemoglobin) and other parameters



#### Blood Gases Measurement – "Astrup"

Assessed by the machine (sensors = selective electrodes):

 $pH = 7,4 \pm 0,04$ 

pCO<sub>2</sub> = 5,3 kPa

pO<sub>2</sub> = 13,3 kPa

- Calculated by the machine:
- [HCO<sub>3</sub><sup>-</sup>] = 24 mmol/l
  - calculated using HH equation
- BE = 0 mEq/I
  - Base Excess, Hb concentration is needed for the calculation.

#### Case study No. 3

- 68 year old male comes to your ambulance.
- History of chronic bronchitis and pulmonary emphysema.
- Mild dyspnea, COVID antigen test negative
- Lab: pH = 7.31
  - pO2 = 8.0 kPa
  - pCO2 = 10.6 kPa
  - HCO3- = 38 mmol/L
  - BE = 12 mmol/L



What kind of acid-base disturbance do we deal with? It this an acute a chronic problem?

#### **Possible Causes of Respiratory Acidosis**

- Decreased alveolar ventilation
  - A) Respiratory center depression
    - Drugs, medicaments (e.g. opioids)
    - Damage or ischemia:
      - Trauma
      - Stroke
      - Tumor
      - Cerebral edema / increased intracranial pressure
  - B) Nerve or muscle disease
    - Myasthenia gravis
    - Polyradiculoneuritis
    - Serious obesity/ Pickwickian syndrome

- C) Lung disease
  - Restrictive diseases
    - ARDS
    - Pulmonary fibroses
  - Obstructive diseases
    - Astma
    - Tumor
    - Foreign body
  - Increase in dead space
    - Pulmonary embolism
    - Pulmonary emphysema
  - Trauma, pneumothorax, serial rib fractures
- Increased pCO<sub>2</sub> in the inspired air



### Case Study No. 2

- You examine a 20 YO student at the hospital admission.
- Cannot concentrate and even could not move her fingers for a brief moment (which scared her). Still feels strange pins and needles in her fingers.
- She has not been seriously ill until now, no medication
- Physical examination normal
- SA: She has split with her boyfriend recently, had been together for 4 years. Difficult to go thru.
- Lab: pH = 7.49
  - pO2 = 13.4 kPa
  - pCO2 = 4.1 kPa
  - HCO3- = 22 mmol/L
  - BE = -1 mmol/l



What acid-base disturbance this is? What kind of acute problem do we see here? What would be your advise her?

### Possible Causes of Respiratory Alkalo

Hyperventilation

- A) At hypoxemia
  - High altitude disease
  - Right-left pulmonary shunting
    - And ventilation-perfusion dysbalance similar to shunt
  - With artificial ventilation
- B) Respiratory center irritation
  - Trauma, inflammation, salicylates.
- C) Panic attack





#### Case Study No. 1

- 38 yo female, DM 1<sup>st</sup> type
- Chills and fever lasting several days
- She has not felt well --> not eaten much
   → not taken much insulin
- During admission day: Abdominal cramps, vomited several times
- Physical exam: BF 30 min<sup>-1</sup>, HF 112 min<sup>-1</sup>, BP 110/70 lying and 100/60 standing, 37 °C,
- Dry mucosae and fruity breath odor

What acid base disturbance do we deal with? Is it a compensated disturbance? What else could be said about her hydration and ion concentrations?



Lab:

pH7.20pO212.8 kPapCO22.8 kPaHCO3-8 mEq/LGlc15 mmol/LNa+148 mEq/LK+5.5 mEq/LCl-110 mEq/LPositive aceton in urine

#### **Possible Causes of Metabolic Acidosis**

- A) Loss of bicarbonates due to increased acid buffering
  - Ketoacidosis
    - Diabetic
    - Alcohol
    - Starving
  - Lactic Acidosis
    - Enormous physical strain
    - Circulatory shock / systemic ischemia
  - Allogenic substances
    - Salicylate poisoning

AG (anion gap) is increased!: Anion of the buffered away acid accumulates in the body.

### B) Loss of bicarbonates into the third space/out of body

- Through intestines
  - Diarrhea
  - Fistulas and stomias
- Through kidneys (loss of regulation)
  - So called Renal tubular acidoses
  - Renal failure (can have 个AG)

The difference in common strong ions reflects  $\downarrow$  HCO<sub>3</sub><sup>-</sup> E.g.  $\uparrow$  Cl<sup>-</sup> (instead of the bicarbonate)so called "hyperchloremic acidoses" (Or there can be e.g.  $\downarrow$  Na<sup>+</sup> or..) AG (anion gap) is normal!

#### Exercise – Metabolic Alkalosis

 Try to derive the beaker chart of metabolic alkalosis and its compensation by yourself. (result can be checked on the next slide)



# Compensation of metabolic alkalosis

- Try do deduce the beakers and the arrows in the compensation diagram yourself.
- Min 2 minutes



#### **Possible Causes of Metabolic Alkalosis**

- Loss of acid by vomiting
   ↑ HCO<sub>3</sub><sup>-</sup> produced by stomach into the blood (when H<sup>+</sup> is secreted into the lumen).
- Increased renal HCO<sub>3</sub><sup>-</sup> production/ increased urine H<sup>+</sup> secretion
  - Hyperaldosteronism
  - So called Bartter syndrome
- Liver failure ( production of urea from NH<sub>4</sub><sup>+</sup> - the reaction would be acidifying)
- Non-adequate infusion of bicarbonates/ Ringer lactate.

# Pathogenesis of Paradoxical Aciduria and Loss of K<sup>+</sup> after Severe Vomiting

- Clinacally important!
- After profuse vomiting, hypochloremic metabolic alkalosis develops
- Under normal circumstances, kidneys should regulate and produce only slightly acidic or alkaline urine.
- Instead, kidneys can worsen the alkalosis
- See next slide:





### Clinical Examples of ABB Disturbances

#### Summary

- Physiology and chemistry
   H+, pH, buffers, buffers
   incorporated into metabolism, HH equation, electroneutrality
- 2. Disturbances divided into respiratory and metabolic
- 3. Clinical examples and causes

	Primary disturbance	Compensation
Respiratory acidosis	↑pCO <sub>2</sub>	Renal - 个HCO <sub>3</sub> -, 个BE
Metabolic acidosis	↓HCO <sub>3</sub> -	Pulmonary - $\downarrow$ pCO <sub>2</sub>
Respiratory alkalosis	↓pCO <sub>2</sub>	Renal - $\downarrow$ HCO <sub>3</sub> <sup>-</sup> , $\downarrow$ BE
Metabolic alkalosis	个HCO <sub>3</sub> -	$Pulmonary - \uparrow pCO_2$

Thank you for your attention