

ACID BASE BALANCE

Normal Mechanism
-Cause of Imbalance
-C
ompensation

Acid base balance

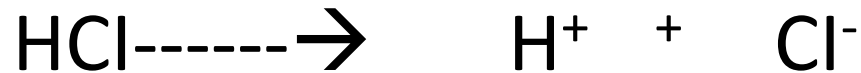
- ▣ For normal functions of body and normal enzyme activity a normal Hydrogen ion concentration(normal PH) is essential.
- ▣ PH is the negative log of Hydrogen (H^+ ion) concentration.
- ▣ A Hydrogen ion is a single free proton released from Hydrogen atom.
- ▣ Normal $[H^+]$ of arterial blood = 0.00004 meq/l which is equal to PH 7.4.

- PH of arterial blood = 7.4
- PH of venous blood = 7.35
- Intra cellular PH is slightly lower than plasma PH.
- PH of urine is 4.5 - 8.0
- **Acidosis** is the PH of body fluid less than normal.
- **Alkalosis** is the PH of the body fluid is more than normal.
- PH limit for human survival is 6.8 – 8.0

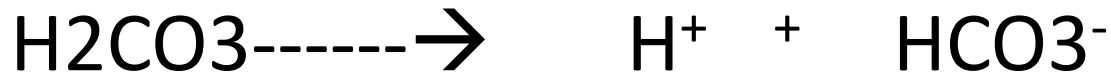
ACIDS

- Acids are molecules that release H^{+} ion (proton) in solutions ("proton donors").

Strong acids dissociate rapidly and release large amount of H^{+} ion .



Weak acids have less tendency to dissociate and release less amount of H^{+} ion



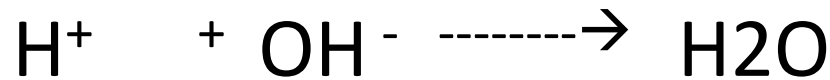
Sources of [H⁺]



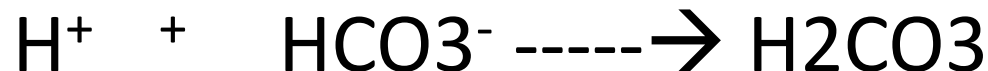
- Carbonic acid formation: the major source of H⁺ is from metabolically produced CO₂
- Inorganic acid produced during nutrient breakdown: dietary **proteins** contain a large quantity of sulfuric acid and phosphoric acid
- Organic acid resulting from intermediary metabolism: lactic acid and fatty acids

BASES

- Bases are molecules that can accept H^+ ion . ("proton acceptors").
- **Strong bases** react strongly with H^+ ion .



- **Weak bases** bind weakly with H^+ ion .



Production of bases in the body:

- Blood alkalinity increases when the level of acid in the body decreases or when the level of base increases e.g:
- A vegetarian diet has a tendency for alkalinity because it produces salts of organic acids such as Na lactate which utilizes H^+ ions making the body alkaline.
- Physiologically imp. base in the body is Bicarbonate ion & biphosphate ion of buffer systems.

Defence against change in PH

- 1. Buffer system of our body.
- 2. Respiratory system of our body.
- 3. Renal control of our body.

MECHANISMS OF REGULATION OF pH

The mechanisms of regulation of blood pH involves the following factors:

(a) *"Front-line" defence*: They are mainly:

- *Buffer systems* in the blood: Which restricts pH change in body fluids.
- *Respiratory mechanisms*: Regulation of excretion of CO_2 and hence, regulation of H_2CO_3 concentration in EC fluid.

(b) *"Second-line" defence*: This is achieved by kidneys (*Renal mechanisms*). Ultimate excretion of excess of acid or base and thus ultimate regulation of concentration of H^+ and HCO_3^- ions in EC fluid.

(c) *Dilution factor*: The acids introduced into and formed in the body are distributed throughout the ECF volume.

Buffer system

- resists sudden changes in pH.
- General Components

buffer contains:

A)~ a weak acid & its salt 20:1 (NaHCO₃/ H₂CO₃)

Addition of strong acid



HCl = strong non volatile acid

H₂CO₃ = weak and volatile acid

Produced H₂CO₃-----→H₂O + CO₂ (exhaled out)

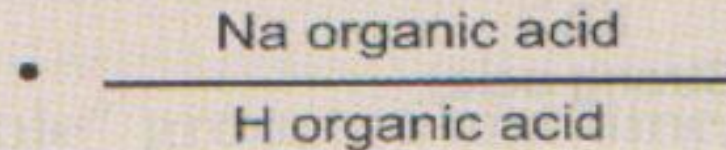
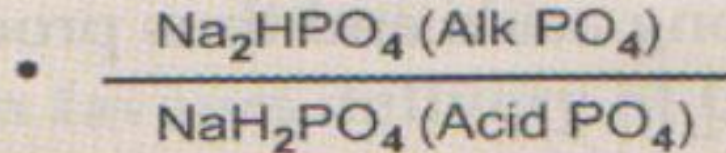
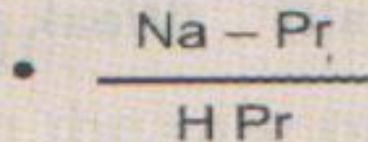
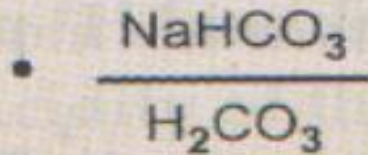
Addition of strong base



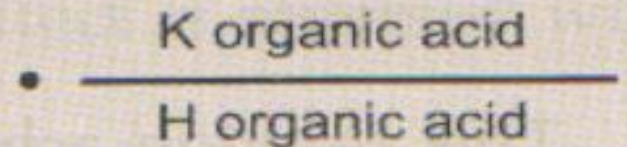
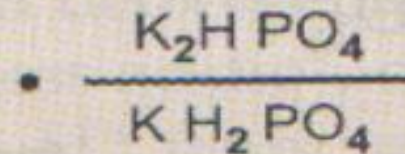
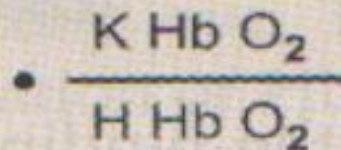
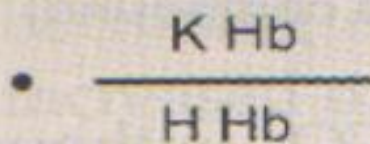
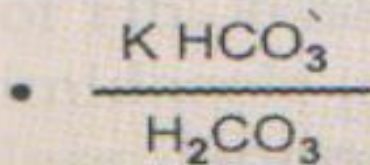
B)~ a weak base & its salt (NH₄OH+NH₄Cl)

- Buffer system reacts within in fraction of a second
- **ONLY KEEP H+ TIED UP.**

- PLASMA BUFFERS



Buffers of RB Cells



BUFFERS OF KIDNEYS

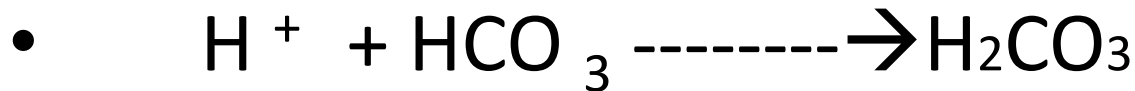
1. Phosphate Buffer system
2. Ammonia and Ammonium buffer systems
3. Urate and citrate also play a minor role

- **VARIOUS BUFFER SYSTEMS:**

- **NaHCO₃ / H₂CO₃ = 20:1**

- Quantitatively most imp buffer system of E.C.F (extra cellular fluid).

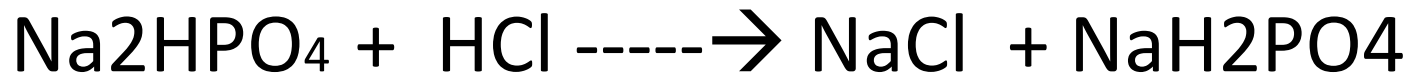
- NaHCO₃ + HL(Lactic acid)



A strong fixed acid -----→ Weak volatile acid
(Lactic Acid)

Phosphate Buffer

Na₂HPO₄ / NaH₂PO₄



Imp Buffer of renal tubular fluid and I C F.

Phosphate are imp because , phosphate are Concentrated due to reabsorption of water but phosphate it self are not reabsorbed.

Protein(zwitter ion)

- **Imp Intra cellular(I C) Buffer system:**
- High concentration with in the cell.

In acidic media:

Act as a base and NH_2 group takes up H^+ ion

•

In basic media:

Act as acid and COOH group give H^+ ion .

- **60%-70% of total buffering of body fluid occurs inside of cell and much of this result from the intra cellular protein.**

Haemoglobin

Haemoglobin (Hb) is not only important in the carriage of oxygen to the tissues but also in the transport of CO₂ and in buffering hydrogen ions

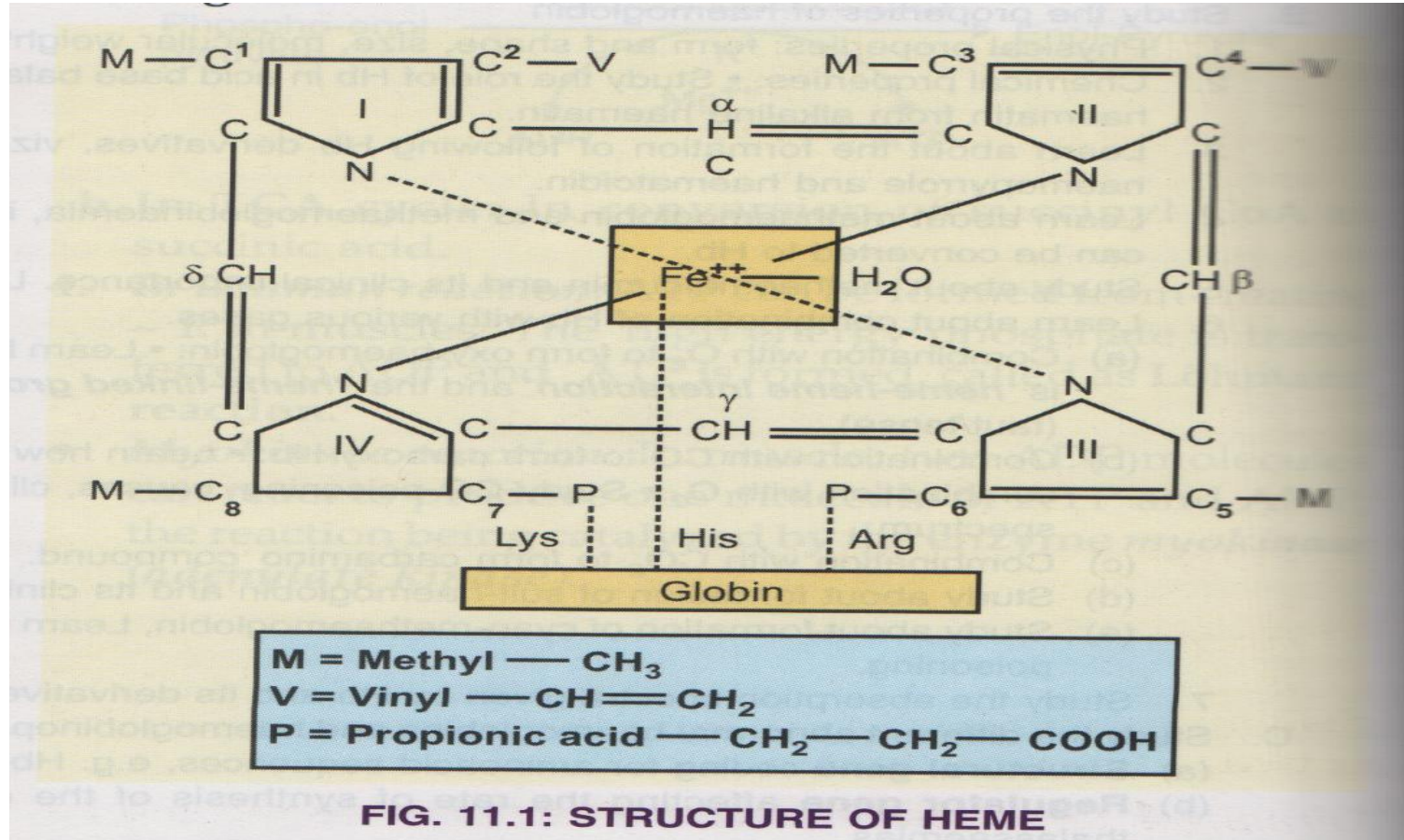
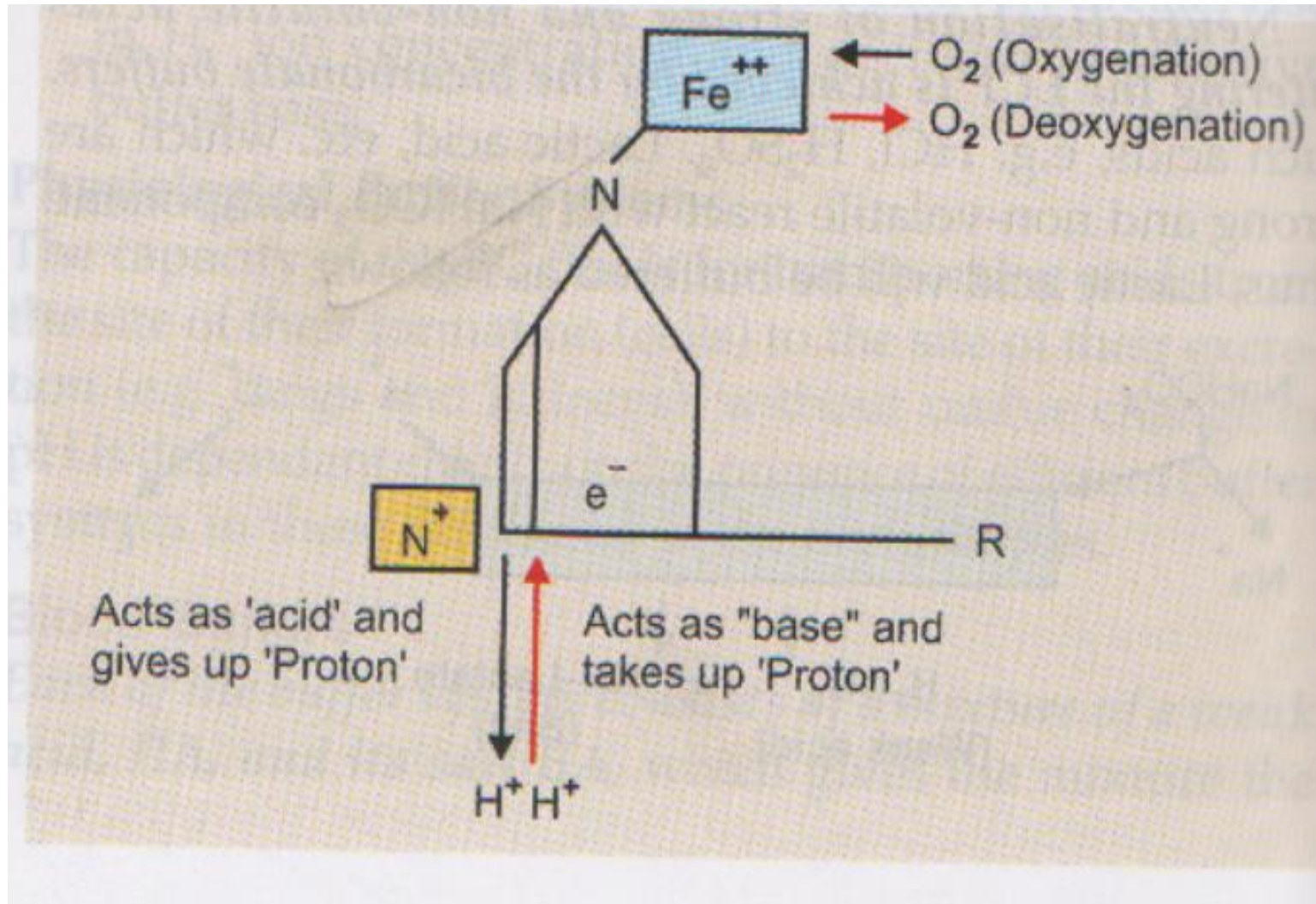


FIG. 11.1: STRUCTURE OF HEME

The buffering capacity of Hb is due to COOH , NH_2 , Guanidine group and imidazole group.



Imidazole contain two group

1. Fe^{++} containing group = Concerned with carriage of O_2 .
2. Imidazole N_2 group: which can give up proton (H^+) and accept H^+ depending on PH of the medium.

So buffering capacity of Hb is due to the presence of imidazole nitrogen group.

Imidazole group of Histidine

Each molecule of Hb contain 38 moles of histidine. In alpha chain histidine at 87 position and in β chain at 92 Position is directly linked with Fe^{++} of haem.

- Guanidine is the functional group on the side chain of Arginine.
- Imadazole group is present in Histadine

Respiratory Regulation

- ▣ Act with in few min (3 – 12).
- ▣ Control PH by altering CO₂ elimination from body by lungs.
- ▣ ↑ blood PCO₂ and (↑ H⁺ ion) stimulates respiratory centre so increases the rate and depth of respiration and so there is ↑CO₂ removal from the body(E C F) through lungs in air.
- ▣ ↓ in blood PCO₂ and (↓ H⁺ ion) depresses R.C.
- ▣ Respiratory centre is sensitive to change in PH and PCO₂.

Respiratory Regulation

- ▣ Respiratory regulation acts rapidly and keeps the Hydrogen Ion concentration from changing too much, until the much more slowly responding kidneys can eliminate the imbalance.
- ▣ **Effect of respiratory control of PH:(50-75%):**
If PH fall from 7.4 to 7, the respiratory system return the PH to the value about 7.2 – 7.3 .

Renal control of acid base balance

- By excreting acidic and basic urine.
- Relatively slow to response.
- Most powerful acid/base regulation system.
- **Mechanism:**
- 80 meq of H^+ non volatile acids excreted/day.
- 4320 meq of HCO_3^- /day.
- Each HCO_3^- require one H^+ to reabsorbed.
- So $4320+80= 4400$ meq of H^+ /day must be secreted by renal tubules.

- 4400 meq of H^+ are secreted by the renal tubules.

Out of this 4400 meq of H^+ ,

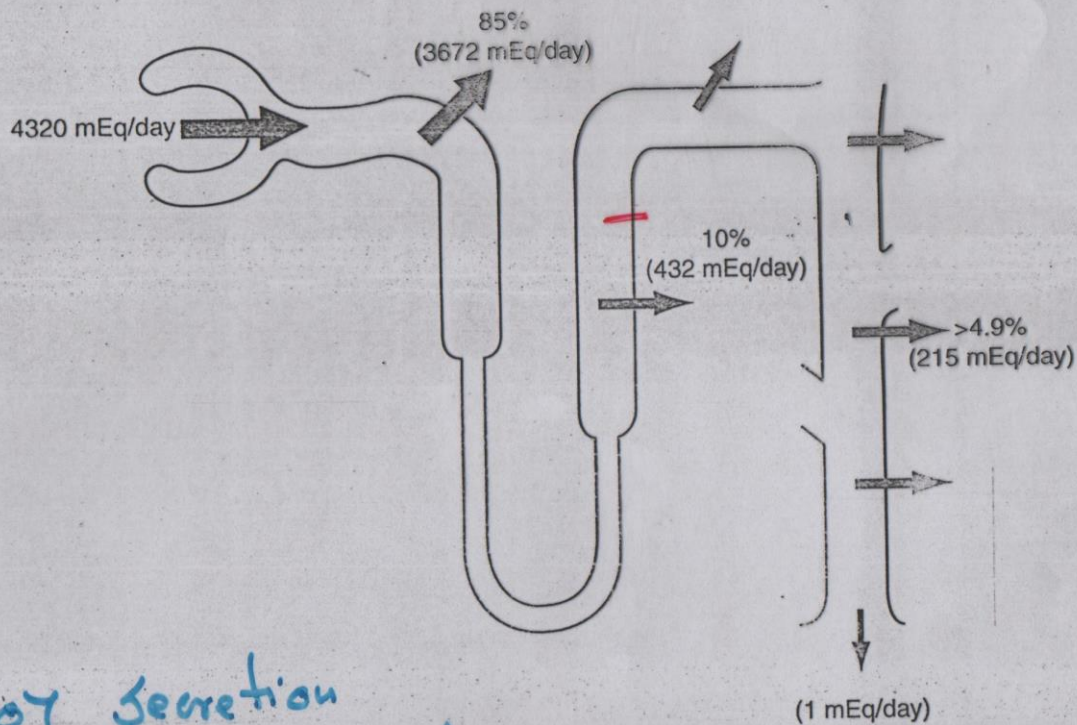
- 80 meq of H^+ are excreted to get rid of Non volatile Acids.
- 4320 meq are required to combine with 4320 meq of HCO_3 filtered in tubule,

Renal control of acid base balance

- **Kidneys regulate the PH by three basic mechanism:**
- Secretion of H^{+lon} .
- Reabsorption of filtered HCO_3^- .
- Production of new HCO_3^- .

- The kidneys control the acid base balance by producing acidic or basic urine.
- **Excreting acidic urine** reduce the amount of acids in extra cellular fluids.
- **Excreting basic urine** reduce the amount of base in the extra cellular fluids.
- Large amount of HCO_3^- are filtered into the tubules i.e. 4320 meq/day
- Large number of H^+ ions are also secreted in the tubular lumen by the tubular epithelial cells to remove 80meq/day of non-volatile acids and to absorb filtered HCO_3^-
- i.e. $4320 + 80 = 4400\text{meq/day}$

Secretion of H^+ and reabsorption of HCO_3^- :



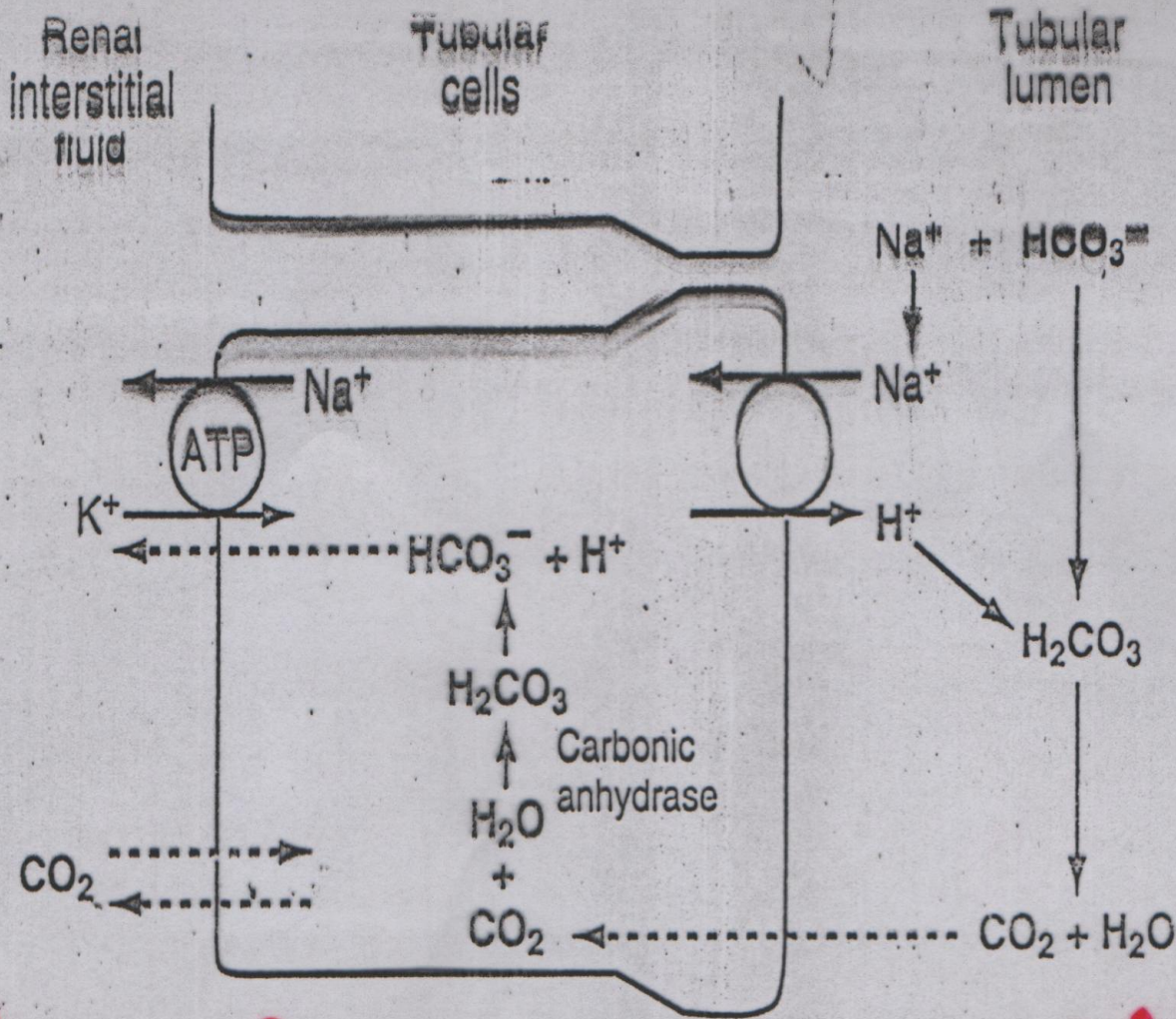
Mechanism of Secretion
of H^+ is different in different
Tubular segments

_____ In Prox tubules H^+ conc can be \uparrow only about 3_4 folds, and Tubular fluid PH can be reduced to only about 6.7 although large amount of H^+ are secreted by this segment.i.e 85% .

_____ In collecting Tubules H^+ conc can be \uparrow up to 900_ fold, decreasing the Tubular fluid PH to about 4.5. although only 5% H^+ secreted in this segment normally.

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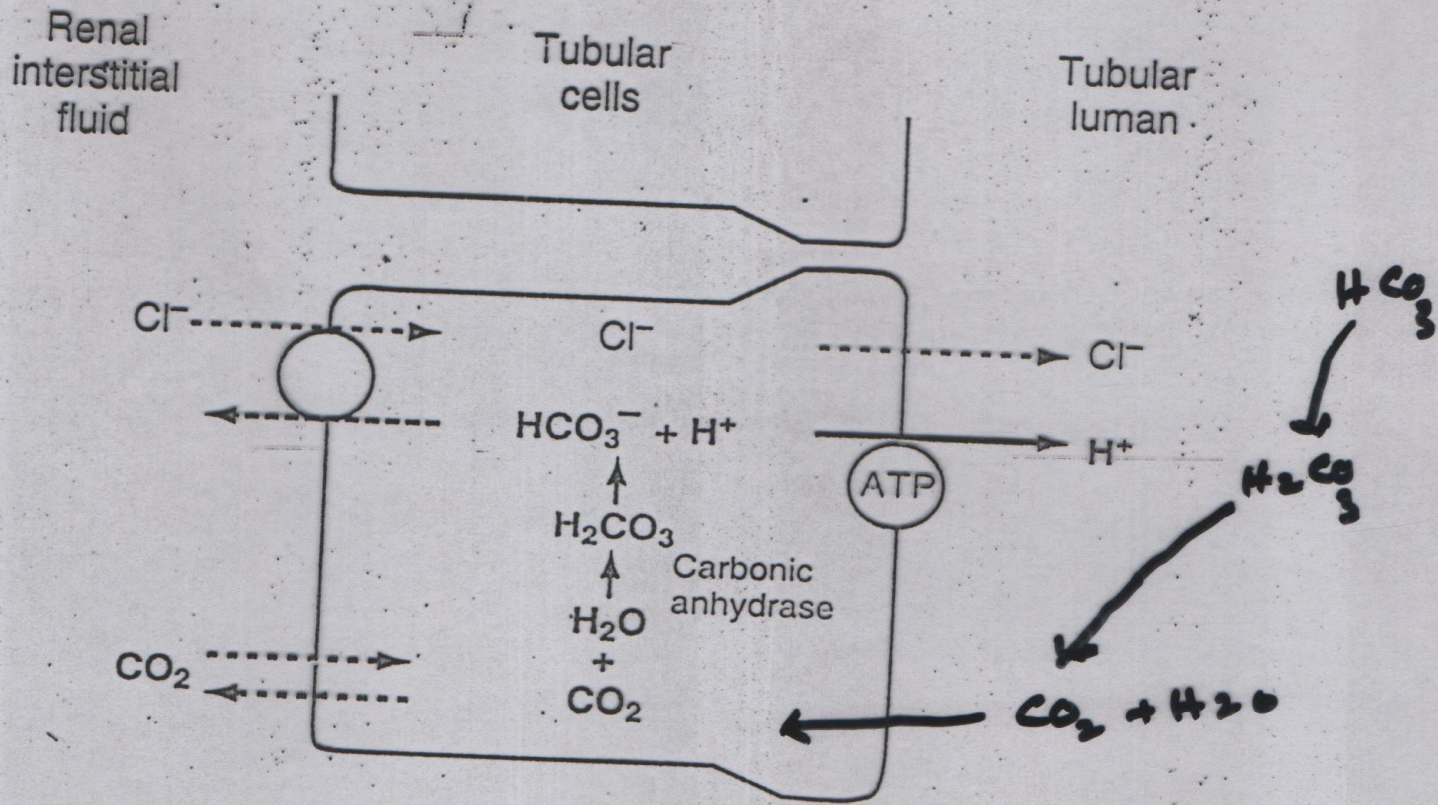
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H⁺ Secretion in Prox tub, Thick Ascending & Early distal tubules.
By Na - H - Counter Transport

The net effect of these reaction is reabsorption of HCO_3^- from Tubules, although the HCO_3^- that has actually entered the extracellular fluid is not the same, as that filtered in to tubules.

II



H^+ secretion in late Tubular segment

- is by Primary active H^+ secretion
- Energy is provided by ATP.

- Once all the HCO_3^- ions has been reabsorbed and is no longer available to combine with H^+ , then any excess of H^+ can combine with HPO_4 and other tubular buffers e.g. Ammonium buffer system
- Whenever a H^+ is secreted into the tubular lumen and combines with a buffer other than bicarbonate, the net effect is addition/regeneration of a new bicarbonate ion into the blood.

Carbonic Anhydrase

Zinc containing metallo enzyme.

Sources: R.B cells. (Never found in plasma)

--- Most of the tissues.

---Parietal cells of the stomach.

---Renal Tubular Epithelial cells.

Also present in small quantity in

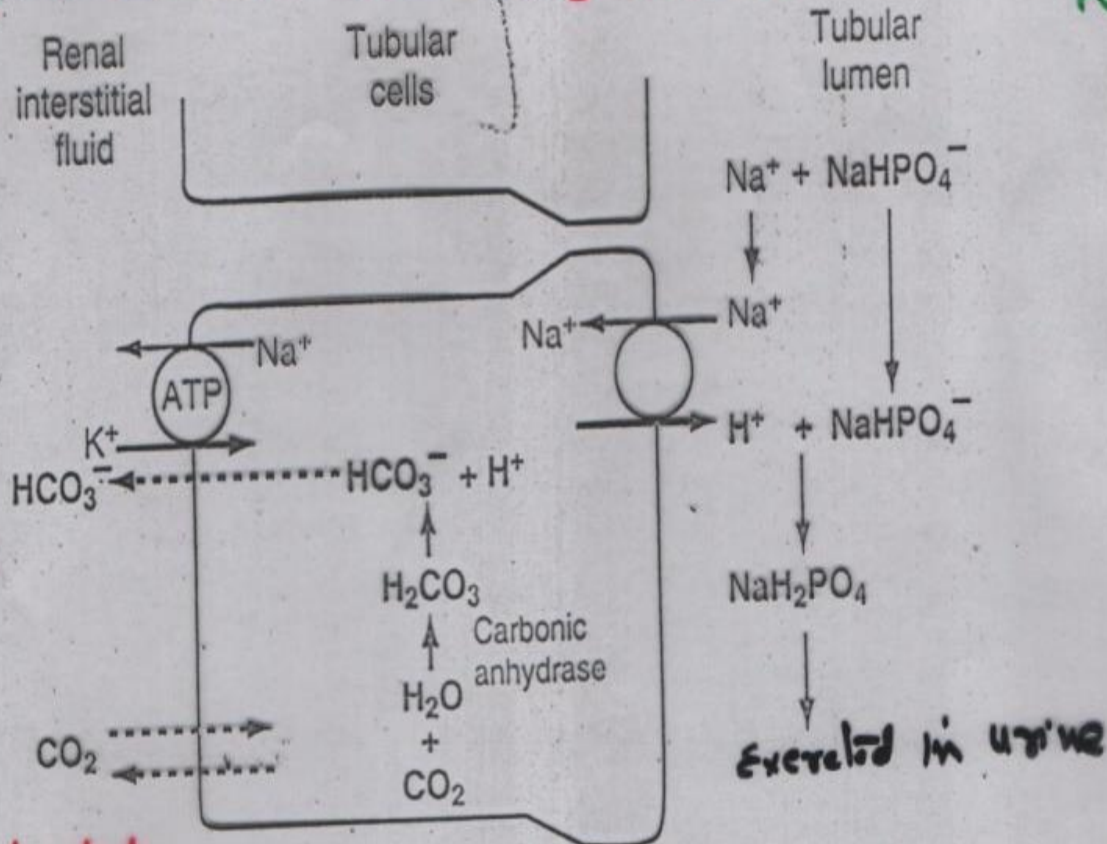
---Muscle tissues.

---Pancreas.

---Spermatozoa.

Combination of Excess of H^+ with urinary buffers and generation of new HCO_3^- ions!.

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Tubular

Renal interstitial fluid

Proximal tubular cells

Tubular lumen

Glutamine

Glutamine

Glutamine

Glutaminase

2HCO_3^-

2NH_4^+

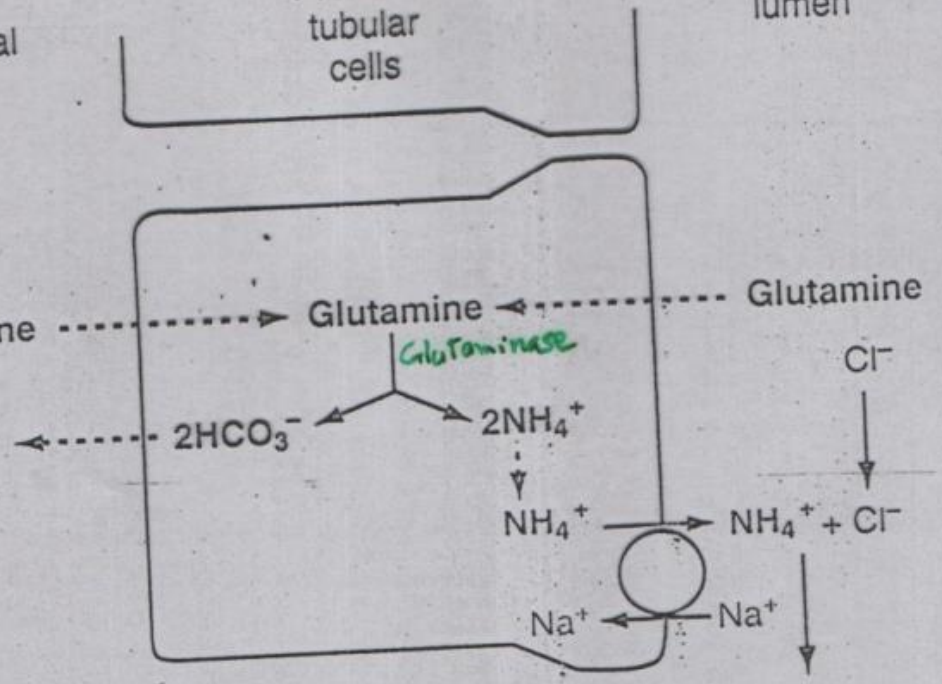
Cl^-

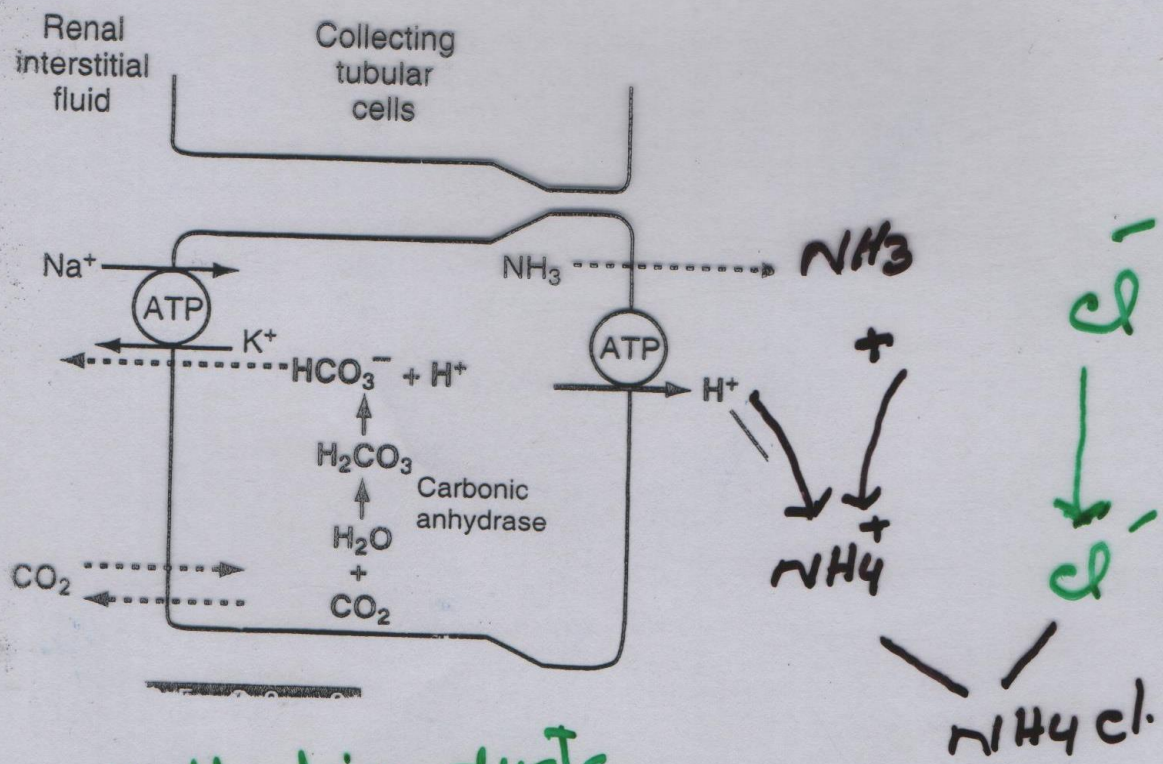
NH_4^+

$\text{NH}_4^+ + \text{Cl}^-$

Na^+

Na^+





Collecting ducts

- Permeable to NH_3
- Not permeable to NH_4^+ .

Thus when there is excess of H^+ , the kidney not only reabsorb all the filtered $H\overline{C}O_3$, but also regenerate new $H\overline{C}O_3$.

- Under normal condition the amount of H⁺ ions eliminated by ammonia buffer system accounts for about 50% of acid excreted and 50% of new bicarbonates generated by the kidneys.
- However in chronic acidosis, the amount of ammonia excreted can increase to as much as 500meq/day.
- Therefore in chronic acidosis, the dominant mechanism by which acid is eliminated is excretion of ammonia.
- This also provides the most important mechanism for generating new bicarbonate during chronic acidosis.

Normal Values

| pH | H ⁺ | Pco ₂ | HCO ₃ ⁻ |
|-----|----------------|------------------|-------------------------------|
| 7.4 | 40 mEq/L | 40 mm Hg | 24 mEq/L |

ACID BASE IMBALANCE

Acid base imbalance can manifest as Acidosis and alkalosis

Acidosis : Which can be

1. Metabolic acidosis
2. Respiratory acidosis

Alkalosis :

Which can be

1. Metabolic alkalosis
2. Respiratory Alkalosis

Metabolic



$$\frac{\text{HCO}_3}{\text{H}_2\text{CO}_3} = \frac{20}{1}$$



Respiratory

Metabolic Acidosis (Primary alkali deficit)

Any type of acidosis except those caused by excessive CO_2

- Fall in pH due to :

A : abnormal accumulation of non-volatile acids in body fluid.

Plasma HCO_3^- is utilized in buffering these acids, so plasma HCO_3^- decrease

B: Increased loss of base :

.

Causes of Metabolic Acidosis

1. Formation of excessive Quantity of Metabolic acids e.g.
 - Diab-Mellitus
 - Starvation
 - High fever
 - violent exercise (L.A)
2. ingestion of acids e.g aspirin and methyl alcohol in large doses.
3. Failure of kidneys to excrete metabolic acids normally formed in the body.
4. Failure of kidneys to reabsorb HCO_3^- resulting in loss of base
5. Severe diarrhea and vomiting of intestinal contents .Large quantity of HCO_3^- are lost .
 - Metabolic Acidosis is the commonest disturbance of Acid base balance observed clinically.

(14) (a) Compensation of Metabolic Acidosis.

(A) Chemical Buffers (Immediately)

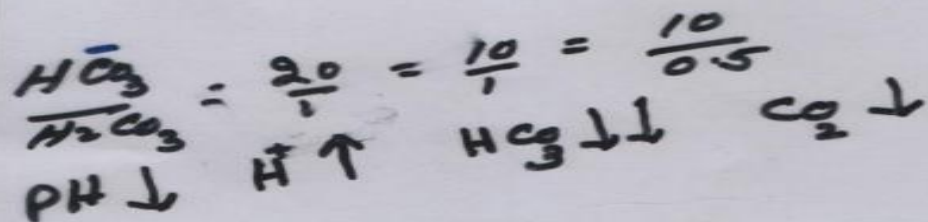
(B) Respiratory Compensation (in minutes)

Acidosis \longrightarrow Stimulation of Resp Centre

\downarrow
 \uparrow Rate and depth of Respiration

\downarrow
 \uparrow Loss of CO_2

\downarrow
 \downarrow Formation of H_2CO_3



As a result, the normal ratio of $\text{HCO}_3^- / \text{H}_2\text{CO}_3$ (20:1) is restored toward normal, as level of both is \downarrow in the blood.

14 (b) Renal (4) Compensation of Metabolic Acidoses

(Hours to days)

(if kidneys are normal)

①

↑ secretion of H^+ (in acidosis) in tubular lumen have two ways.

(i) Complete reabsorption of HCO_3^-

(ii) H^+ in excess of HCO_3^- are lost in combination with urinary buffers
e.g. HPO_4 , NH_3 etc.
 H_2PO_4

So there is ↑ Regeneration of HCO_3^- ions by

(i) HPO_4/H_2PO_4

(ii) NH_3 buffer system

(iii) Glutamine Metab $\begin{cases} 2NH_4^+ \\ 2HCO_3^- \end{cases}$

Acidosis stimulates Glutaminase

Respiration Acidosis

(Primary H_2CO_3 excess)

The underlying abnormality is increase in H_2CO_3 in blood, which follows decreased elimination of CO_2 .

Causes:

Any factor that decrease rate and depth of pulmonary ventilation.

1. Damage to resp centre (Hypoventilation) by trauma , inflammation e.t.c
2. Loss of ventilatory functions due to diseases of plura or lung disease e.g pneumothorax, tumors, emphysema e.t.c
3. Morphine's poisoning
4. Paralysis of diaphragm
5. Certain congenital Heart diseases

6. Condition causing impairment of diffusion of CO_2 across alveolar membrane e.g emphysema , pulmonary oedema , pneumonia

7. Abnormally high conc of CO_2 in atmosphere

- It is an emergency state.
- O_2 should be administered
- Artificial respiration by respirator
- Drugs which stimulate Resp Centre.

(123). Compensation of Respiratory Acidosis: ⁽²⁾

2. Resp Mechanism

① - Chemical buffers

$\uparrow \text{CO}_2 \longrightarrow \uparrow$ stimulation of R.C \longrightarrow
 \uparrow depth and rate of Resp $\longrightarrow \uparrow$ loss of $\text{CO}_2 \longrightarrow \downarrow \text{H}_2\text{CO}_3$

provided the R.C and lungs are normal!

In this Resp acidosis, the Renal Compensation is of prime importance.

③ Renal Mechanism:

(i) $\therefore \uparrow \text{H}^+$ in ECF + Cells $\therefore \uparrow$ Sec of H^+

(ii) $\uparrow \text{H}^+$ in Tubular lumen $\longrightarrow \uparrow \text{HCO}_3^-$ reabsorption

(iii) \uparrow regeneration of HCO_3^-

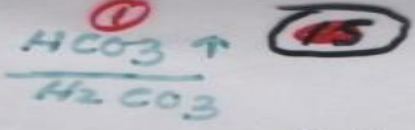
(by glutamine, phosphate buffers & NH_3)

So Ratio of $\text{HCO}_3^- / \text{H}_2\text{CO}_3$ is restored towards 20:1, as level of both is \uparrow in blood. \therefore lowering PH.

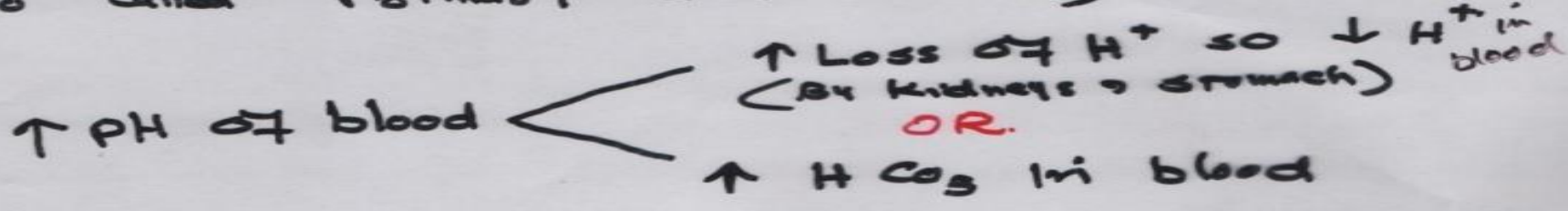
$$\frac{\text{HCO}_3^-}{\text{H}_2\text{CO}_3} = \frac{20}{1} = \frac{20}{2} = \frac{40}{2}$$

PH \downarrow H^+ \uparrow CO_2 $\uparrow\uparrow$ HCO_3^- \uparrow

Metabolic Alkalosis



(Also called Primary alkali Excess)



Causes:

1) vomiting of gastric contents OR suction through N.G. Tube.

2) Administration of Diuretics \rightarrow ChlorThiazide
8+ Causes.

— \uparrow HCO_3 reabsorption associated with
 \uparrow Loss of H^+ so causes alkalosis.

3) Excessive α steroid secretion. \rightarrow Cushing's Syndrome
OR steroid therapy

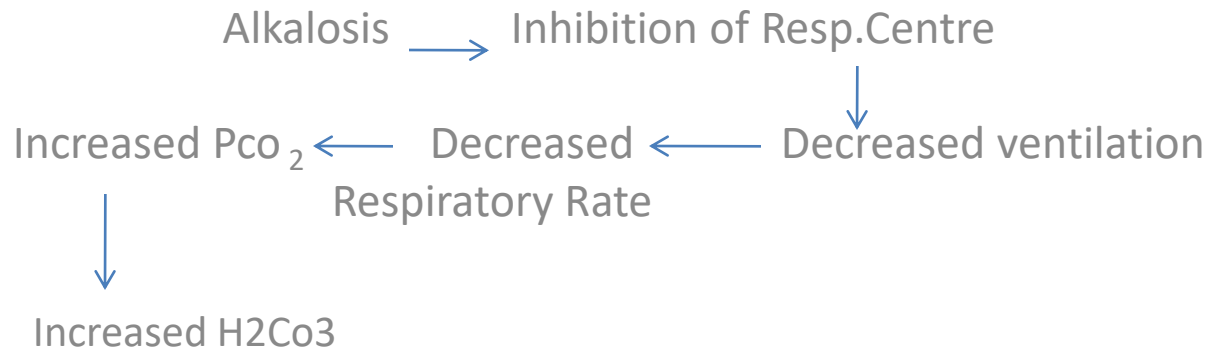
- Steroids causes \uparrow Loss of H^+ (and K^+) in
Exchange for Na^+ , which is reabsorbed,
causing Hypertremia and Hypokalemia.

4) Ingestion of alkaline drug e.g. Sodium Bicarbonate

Compensation of Metabolic Alkalosis

Usually not much helpful and is very difficult to compensate

- A. Chemical Buffers
- B. Respiratory Compensation



$$\text{So } \frac{\text{HCO}_3}{\text{H}_2\text{CO}_3} = \frac{20}{1} = \frac{40}{1} = \frac{40}{2}$$

So HCO₃/H₂CO₃ Will be restore to 20:1 as level of both is increased

PH ↑ H ↓ HCO₃ ↑ ↑ Co₂ ↑

Renal Compensation

- Increased HCO_3 in plasma

so

- Increased Renal filtration

So

- Increased loss of HCO_3

Why metabolic alkalosis is difficult to compensable

Respiratory Compensation: As soon as the depression of respiratory centre by alkalosis causes retention of PCO_2 and increased H_2CO_3 , the increase PCO_2 stimulates the respiratory centre, so the compensation is not of great significance.

Renal compensation: Due to slight respiratory compensation, increased Hydrogen ions will lead again to increased reabsorption of filtered HCO_3 . So again compensation is not very effective.

Respiratory Alkalosis $\frac{HCO_3^-}{H_2CO_3} \downarrow$
(Primary H_2CO_3 deficient)

- It is a rise in pH due to primary decrease in blood PCO_2 and H_2CO_3 .

Causes:

Hyper ventilation (Any cause) e.g

- Psychological causes – e.g Hysteria
- CNS Diseases: Meningitis, Encephalitis
- Patient on respirator
- Hyper pyrexia

Pulmonary causes:

- Pneumonia
- Asthma
- C.C.F
- Pulmonary embolism
- Salicylate poisoning By large doses e.g in rheumatic fever.

↑ Loss of CO_2 due to hyper ventilation result in decreased H_2CO_3 .

So ratio of $\frac{\text{HCO}_3^-}{\text{H}_2\text{CO}_3}$ ↑ i.e pH is ↑

③ $\frac{HCO_3}{H_2CO_3} \downarrow$

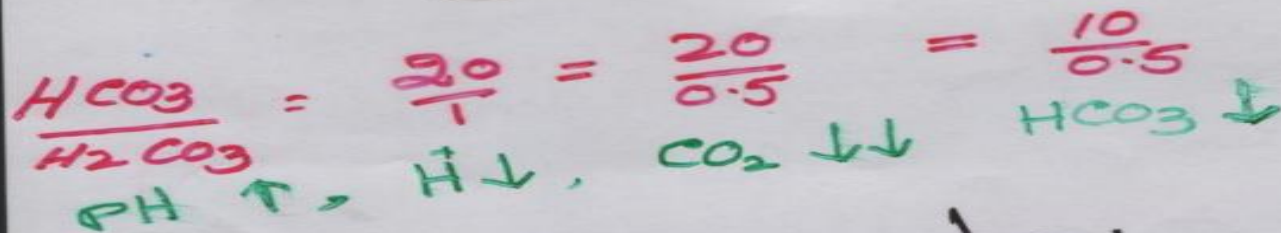
Compensation of Respiratory Alkalosis: - (19)

① Chemical Buffers

② Renal Compensation: - (Main Compens)

(i) \uparrow Filtration of HCO_3^- , but as there is less H^+ secretion due to alkalosis, and more loss of HCO_3^- reabsorption.

(ii) \downarrow HCO_3^- regeneration.



Respiratory Compensation:

Alkalosis and low PCO_2 depresses Resp Center.
So \downarrow rate and depth of resp.
So \downarrow expiration of CO_2 , so \uparrow Blood PCO_2 .

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Metabolic Acidosis :: (Primary Alkali Deficit)

$$\frac{HCO_3^-}{H_2CO_3} = \frac{20}{1} = \frac{10}{1} = \frac{10}{0.5}$$

pH ↓ H ↑ HCO₃ ↓↓ CO₂ ↓

Metabolic Alkalosis :: (Primary Alkali Excess)

$$\frac{HCO_3^-}{H_2CO_3} = \frac{20}{1} = \frac{40}{1} = \frac{40}{2}$$

pH ↑ H ↓ HCO₃ ↑↑ CO₂ ↑

Respiratory Acidosis :: (Primary H₂CO₃ Excess)

$$\frac{HCO_3^-}{H_2CO_3} = \frac{20}{1} = \frac{20}{2} = \frac{40}{2}$$

pH ↓ H ↑ CO₂ ↑↑ HCO₃ ↑

Resp Alkalosis :: (Primary H₂CO₃ Deficit)

$$\frac{HCO_3^-}{H_2CO_3} = \frac{20}{1} = \frac{20}{0.5} = \frac{10}{0.5}$$

pH ↑ H ↓ CO₂ ↓↓ HCO₃ ↓

Table 30-3**Characteristics of Primary Acid-Base Disturbances**

| | pH | H ⁺ | Pco ₂ | HCO ₃ ⁻ |
|-----------------------|-----|----------------|------------------|-------------------------------|
| Normal | 7.4 | 40 mEq/L | 40 mm Hg | 24 mEq/L |
| Respiratory acidosis | ↓ | ↑ | ↑↑ | ↑ |
| Respiratory alkalosis | ↑ | ↓ | ↓↓ | ↓ |
| Metabolic acidosis | ↓ | ↑ | ↓ | ↓↓ |
| Metabolic alkalosis | ↑ | ↓ | ↑ | ↑↑ |

The primary event is indicated by the double arrows (↑↑ or ↓↓). Note that respiratory acid-base disorders are initiated by an increase or decrease in PCO₂, whereas metabolic disorders are initiated by an increase or decrease in HCO₃⁻.

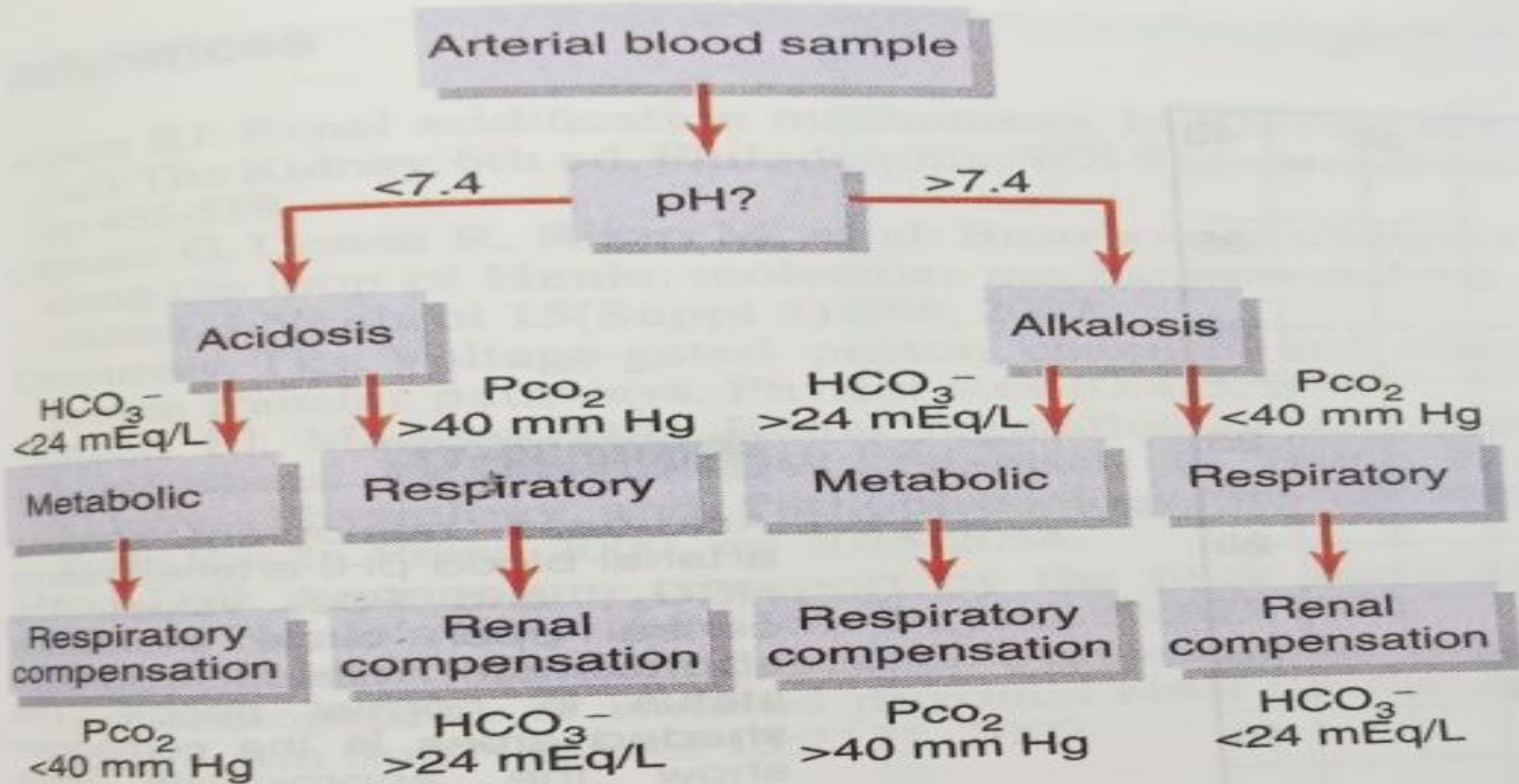


Figure 30-10

Analysis of simple acid-base disorders. If the compensatory responses are markedly different from those shown at the bottom of the figure, one should suspect a mixed acid-base disorder.

- **1. The normal ratio between the alkaline phosphate and acid phosphate in plasma is 4 : 1**
- **2. At pH 7.4, the ratio of bicarbonate : dissolved CO₂ is 20 : 1**
- **3. Quantitatively, the most significant buffer system in plasma is Carbonic acid-bicarbonate buffer system**

4. Buffering action of haemoglobin is mainly due to its Histidine

- **5. Respiratory acidosis results from Retention of carbon dioxide**

- Anion Gap
- Alkali Reserve