

Serum electrolyte Anions

major extracellular anion
← Chloride

The major extracellular anion ☐

Function in body: ☐

Maintaining osmolality ☐

Blood volume and ☐

Electric neutrality ☐

Regulation: Na^+ و HCO_3^- مع Cl^-
~~bicarbonate~~

Cl is usually shifted according to Na and bicarbonate ☐

Excess chloride in the body is excreted in urine and sweat, excessive sweating will induce the release of aldosterone which will conserve Na and Cl ☐

لذا في سوائل الجسم 40 و Cl^- 20
فيما في رشح بس

Chloride

Chloride maintains electrical neutrality in two ways: ☐

Na is reabsorbed along with Cl in the proximal tubules. Na ☐
reabsorption is limited by the amount of Cl- available

Electroneutrality is also maintained by chloride through the chloride ☐
shift.

Carbon dioxide generated by cellular metabolism within the tissue diffuses ☐
out into both the plasma and the red cells

In the red cell, CO₂ forms carbonic acid (H₂CO₃), which splits into H⁺ and ☐
HCO₃⁻ (bicarbonate).

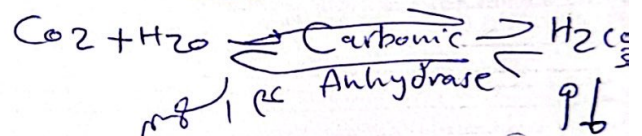
Deoxyhemoglobin buffers H⁺, whereas the HCO₃⁻ diffuses out into the ☐
plasma and Cl diffuses into the red cell to maintain the electric balance of
the cell.

9)

reference range Na 135 - 145

Cl⁻ → 98 - 107

HCO₃⁻ → 23 - 29



Chloride applications $\text{H}^+ + \text{HCO}_3^- \rightarrow \text{H}_2\text{O} + \text{CO}_2$

Chloride disorders are often the result of the same causes that ☐
disturb Na levels because chloride passively follows Na

renal tubule
acidosis

There are a few exceptions. ☐

Hyperchloremia may also occur when there is an excess loss of bicarbonate as a ☐
result of GI losses, RTA or metabolic acidosis

Hypochloremia may occur with excessive loss of chloride from prolonged vomiting, ☐
diabetic ketoacidosis, aldosterone deficiency or salt-losing renal diseases.

A low serum level of chloride may be encountered in conditions associated with high ☐
serum bicarbonate concentrations such as compensated respiratory acidosis or
metabolic alkalosis.

↑ ions
↑ bodies
↑ Na⁺
↑ H₂O
↑ HCO₃⁻
↑ Cl⁻
↑ hyperchloremia

Determination of the chloride

Specimen: serum or plasma, whole blood samples, urine (24-hr) or sweat may be used ☐

* Lithium heparin is the anticoagulant of choice. ☐

Hemolysis does not cause significant change in serum or plasma values as a result of decreased levels of intracellular chloride (marked hemolysis, decrease due to dilutional effect). ☐

Methods: there are several methodologies includes: ☐

ISE (most commonly used where an ion-exchange membrane is used to selectively bind Cl^- ions) ☐

Ion selective electron-

Amperometric coulometric titration ☐

Mercurimetric titration ☐

Colorimetry ☐

Amperometric coulometric titration method using coulometric generation of silver ions (Ag^+ which combine Cl^- to quantitate the Cl^- ion concentration) ☐

$\text{Ag}^+ + \text{Cl}^-$

AgCl

Reference range

Reference Range

TABLE 15-10 REFERENCE RANGES FOR CHLORIDE

Plasma, serum	98-107 mmol/L	Ref Range Cl^- in Blood
Urine (24-h)	110-250 mmol/day, varies with diet	

(بانيمايوتو) (diabetes Rel)

Bicarbonate

في قنصل الكلى

Is the second most abundant anion in the ECF

The total CO₂ comprises the bicarbonate (90%), carbonic acid and dissolved CO₂ so total CO₂ measurement is indicative of HCO₃⁻ measurement

Bicarbonate is the major buffering system in the blood where carbonic anhydrase in RBCs converts CO₂ and H₂O to carbonic acid



Bicarbonate diffuses out of the cells in exchange for chloride to maintain ionic charge neutrality within the cell

Bicarbonate regulation

Most of the filtered bicarbonate ion is reabsorbed in the kidneys. (85% in proximal tubules and 15% in the distal) in the form of CO₂ (due to low permeability of tubules to bicarbonate)

Normally nearly all the bicarbonate ions are reabsorbed from the tubules, with little lost in the urine

When bicarbonate ions are filtered in excess of hydrogen ions available, almost all excess HCO₃⁻ flows into the urine.

In alkalosis, with relative increase in bicarbonate ion compared to CO₂, the kidneys increase excretion of HCO₃⁻ into the urine, carrying along a cation such as sodium. This loss of HCO₃⁻ from the body helps correct pH

In acidosis, the excretion of H into the urine is increased and HCO₃⁻ reabsorption is virtually complete

Alkalosis

Clinical applications

Acid-base imbalances cause changes in bicarbonate and CO₂ levels. A decreased bicarbonate/ CO₂ occurs in metabolic acidosis leads to exhalation of CO₂ by the lungs (hyperventilation), which lowers pCO₂.

Elevated total CO₂ concentrations occur in metabolic alkalosis as bicarbonate is retained, often with increased pCO₂, as a result of compensation by hypoventilation.

Typical causes of metabolic alkalosis include:

- Severe vomiting ☐
 - Hypokalemia ☐
 - Excessive alkali intake ☐
- Handwritten notes:
 - HCl secretion
 - H^+ secretion
 - K^+ depletion
 - H_2O depletion

Method

reference range
 الی فستوب در ال لاسون
 آرما آرکست ال لاسون
 Specimen: venous serum or plasma. ☐

Serum or lithium heparin plasma is suitable for analysis. ☐

The sample is capped until the serum or plasma is separated and the sample is analyzed immediately ☐

If the sample is left uncapped before analysis, CO₂ escapes. Levels can decrease by 6 mmol/L per hour ☐

Two common methods are ISE and an enzymatic method. ☐

ISE for measuring total CO₂, uses an acidic reagent to convert all the forms CO₂ to CO₂ gas and measured by a pCO₂ electrode ☐

The enzyme method alkalizes the sample to convert all forms of CO₂ to HCO₃⁻. ☐

Method

HCO₃⁻ is used to carboxylate phosphoenolpyruvate (PEP) of phosphoenolpyruvate (PEP) □
carboxylase, which catalyzes the formation of oxaloacetate.

Phosphoenolpyruvate + HCO₃⁻ $\xrightarrow{\text{PEP carboxylase}}$ Oxaloacetate + H₂PO₄⁻ (Eq. 15-4) s a result of the action of □
 This is coupled to the following reaction in which NADH is consumed as are suff
 of the action

Oxaloacetate + NADH + H⁺ $\xrightarrow{\text{MDH}}$ Malate + NAD⁺ (Eq. 15-5) □
 The rate of change in the

absorbance
 of NADH is proportional to the
 Concentration of HCO₃⁻ □
 Reference ranges
 Carbon dioxide, venous 23-29 mmol/L (plasma, serum). □

Lactate

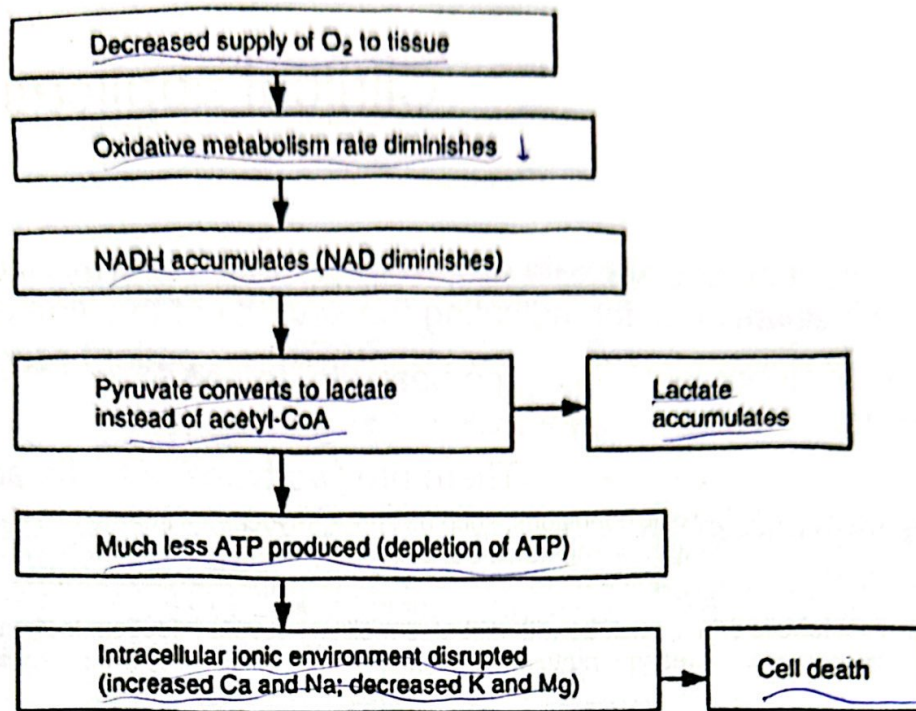
Lactate is a by-product of an emergency mechanism that □
 produces a small amount of ATP (2 moles)

Under hypoxic conditions, acetyl CoA formation does not □
 occur and NADH accumulates, favoring the conversion of
 pyruvate to lactate through anaerobic metabolism.

The accumulation of excess lactate in blood is an early □
 sensitive and quantitative indicator of the severity of oxygen
 deprivation (more than pH)

1 NADH = 2.5 ATP

□



QX Antibiotic

Regulation

It is not regulated as with potassium and calcium □

As oxygen delivery decreases below a critical level, blood □
lactate concentration rise rapidly and indicate tissue hypoxia
earlier than pH

The liver is the major organ for removing lactate by converting □
lactate back to glucose by a process called (gluconeogenesis)

97

Clinical application

Measurement of blood lactate are useful for metabolic monitoring ☐
in critically ill patients, for indicating the severity of the illness, and
patient prognosis

There are two types of lactic acidosis: ما في نقل O_2

Type A is associated with hypoxic conditions, such as shock, myocardial infarction, severe ☐
congestive heart failure, pulmonary edema, or severe blood loss

Type B is of metabolic origin, such as with diabetes mellitus, severe infection, leukemia, ☐
liver or renal disease, and toxins (ethanol, methanol, or salicylate poisoning).

Determination of lactate

Special care should be practiced when collecting and handling ☐
specimens for lactate levels

الربطة لا اسحب ابداً

A tourniquet should not be used because venous stasis will increase ☐
lactate levels

If a tourniquet is used, blood should be collected immediately and the ☐
patient should not exercise the hand before and during collection

After sample collection, glucose is converted to lactate by a way of ☐
anaerobic glycolysis and should be prevented:

Heparinized blood may be used but must be delivered on ice and the plasma must ☐
be quickly separated

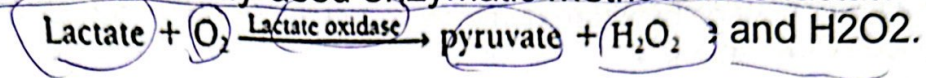
Iodoacetate or fluoride will inhibit glycolysis without affecting coagulation ☐

لا بأس
lactate
معدل

Method

Current enzymatic methods make lactate determination ☐ readily available.

The most commonly used enzymatic method uses lactate ☐



One of two couple reactions may then be used. Peroxidase ☐

may be used $\text{H}_2\text{O}_2 + \text{H donor} + \text{chromogen} \xrightarrow{\text{Peroxidase}}$ from H_2O_2

colored dye + $2\text{H}_2\text{O}$ (E)

Reference range

ENZYMATIC METHOD, PLASMA

Venous	0.5–2.2 mmol/L (4.5–19.8 mg/dL)
Arterial	0.5–1.6 mmol/L (4.5–14.4 mg/dL)
CSF	1.1–2.4 mmol/L (10–22 mg/dL)

Anion gap (AG)

Routine measurement of electrolytes usually involves only Na^+ , K^+ , Cl^- and HCO_3^- □

These values may be used to approximate the anion gap (AG), which is the difference between unmeasured anions and unmeasured cations. □

$$\text{AG} = (\text{Na}^+ + \text{K}^+) - (\text{Cl}^- + \text{HCO}_3^-)$$

There is never a "Gap" between total cationic charges and anionic charges □

AG is useful in indicating an increase in one or more of the unmeasured anions in the serum and also in the form of quality control for the analyzer used to measure these electrolytes. □

Anion gap (AG)

There are two commonly used methods for calculating the anion gap □

$$\text{AG} = \text{Na}^+ - (\text{Cl}^- + \text{HCO}_3^-)$$

With a reference range of 7-16 mmol/L □

The second method: □

$$\text{AG} = (\text{Na}^+ + \text{K}^+) - (\text{Cl}^- + \text{HCO}_3^-)$$

It has a reference range of 10-20 mmol/L □

Anion gap (AG)

- An elevated anion gap may be caused by:
- Uremia/renal failure, which leads to PO_4^{3-} and SO_4^{2-} retention ☒ 1
 - Ketoacidosis, as seen in cases of starvation or diabetes ☒ 2
 - Methanol, ethanol, ethylene glycol poisoning, or salicylate ☒ 3
 - Lactic acidosis ☒ 4
 - Hypernatremia ☒ 5
 - Instrument error ☒ 6

CASE STUDY 15-2

A 60-year-old man entered the emergency department after 2 days of "not feeling so well." History revealed a myocardial infarction 5 years ago, when he was prescribed digoxin. Two years ago, he was prescribed a diuretic after periodic bouts of edema. An electrocardiogram at time of admission indicated a cardiac arrhythmia. Admitting lab results are shown in Case Study Table 15-2.1.

Questions

1. Because the digoxin level is within the therapeutic range, what may be the cause for the arrhythmia?
2. What is the most likely cause for the hypomagnesemia?
3. What is the most likely cause for the decreased potassium and ionized calcium levels?
4. What type of treatment would be helpful?

CASE STUDY TABLE 15-2.1 LABORATORY RESULTS

VENOUS BLOOD

Digoxin: 1.4 ng/mL, therapeutic 0.5-2.2 (1.8 nmol/L, therapeutic 0.6-2.8)

Na^+ : 137 mmol/L

K^+ : 2.5 mmol/L

Cl^- : 100 mmol/L

HCO_3^- : 25 mmol/L

Mg^{2+} : 0.4 mmol/L

Ion/free Ca^{2+} : 1.0 mmol/L

Na	135-145
K	3.4-5.0
Cl	98-107
HCO_3^-	23-29
Mg	0.63-1.0
Ca/ionized	1.16-1.32