

Regulation of Parathyroid gland Calcium, Magnesium and Phosphate

بِسْمِ اللَّهِ نَبْدَأُ 

يعطيكم العافية جميعا ان شاء الله
هاد راح يكون تفريغ اول شايتر
من مادة القاينل

Calcium level ← monitoring
* كثير منهم عندي اعلى

خصوصاً عند الأشخاص :
1. critically ill patient.

2. have a surgery

لعمريان حايصير عندهم تسنج
بالعضلات

Calcium → There are 3 forms of calcium:

1. free or ionized.

active form ← حيواري
Calcium. هذا

2. bind with albumin (40%).

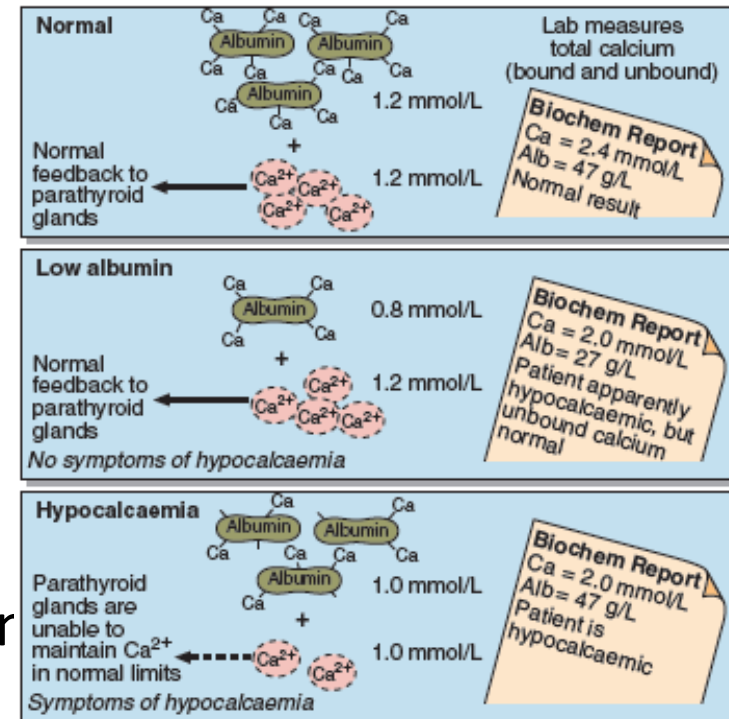
3. bind with phosphate, bicarbonate, citrate, lactate → insoluble form. (5%).

له صلا ليقبض منه.

❑ The ionized/free calcium is essential for myocardial contraction whereas protein bound and citrate-bound calcium had no effect

❑ It is important to maintain ionized calcium at a near normal concentration during surgery and in critically ill patients.

❑ Decreased ionized Ca conc. in blood can cause neuromuscular irritability which may become clinically apparent as irregular muscle spasms, called tetany.



① ionized calcium.

or

② total calcium + calcium bind with albumin

← شرط أساسي فابقدر أعمل استخدم ال
total calcium alone.

① total calcium → normal
+ calcium albumin → normal } → Then the level of calcium normal.

② total calcium → normal → have a problem.
calcium albumin → low يكون ياف عنده ارتفاع calcium. لا

③ ionized calcium → normal → the problem because of albumin.
total calcium → low

④ total calcium → high → hypercalcemia
calcium albumin → normal

⑤ total calcium → low → hypocalcemia.
calcium albumin → normal

99% of stored
calcium in the
body → in the bone.

Regulation of Calcium

- ① → parathyroid hormone. ②
- ❑ Three hormones, PTH, vitamin D, and calcitonin, are known to regulate serum calcium by altering their secretion rate in response to changes in ionized calcium

→ any decrease of ionized calcium → signal to parathyroid gland to secrete PTH.

- ❑ PTH secretion in blood is stimulated by a decrease in ionized calcium and, conversely PTH secretion is stopped by an increase in ionized calcium.

secretion له بروتين ← Bone ← PTH موجوده على الـ receptors ← يكون في

- ❑ PTH exerts three major effects on both:

- ❑ Bone: activates bone resorption, break down osteoclast and release Ca to ECF

- ❑ Kidney: conserves calcium by increasing tubular reabsorption of calcium ions and stimulates renal production of active vitamin D.

receptors in PTH in kidney → 1. increase tubular reabsorption.

→ 2. activate α hydroxylase enzyme → activation of vitamin D

Activation of Vitamin D: Vitamin D is activated from an inactive form to an active form. The process involves two steps: (1) 25-hydroxylation in the liver and (2) 1- α -hydroxylation in the kidney.

Regulation of Calcium

→ in hypocalcemia.

❑ Vitamin D₃, a cholecalciferol, is obtained from the diet or exposure of skin to sunlight, hydroxylated in liver to 25-OH- (inactive form), activated in the kidney by 1- α -hydroxylase to form (1,25-OH)₂-D₃, the biologically active form which will:

← Vitamin D active form.

- ① ❑ Increase calcium absorption in the intestine and
- ② ❑ Enhance the effect of PTH on bone resorption.

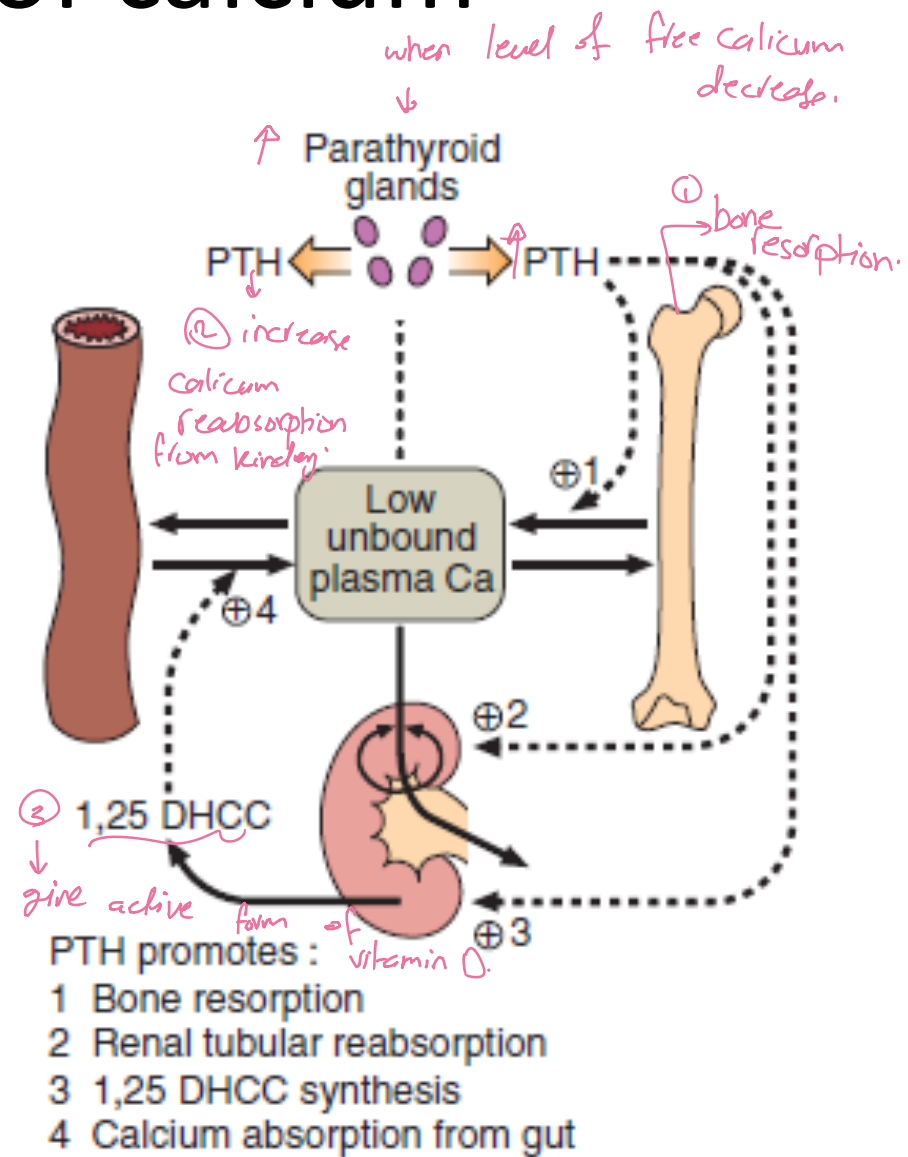
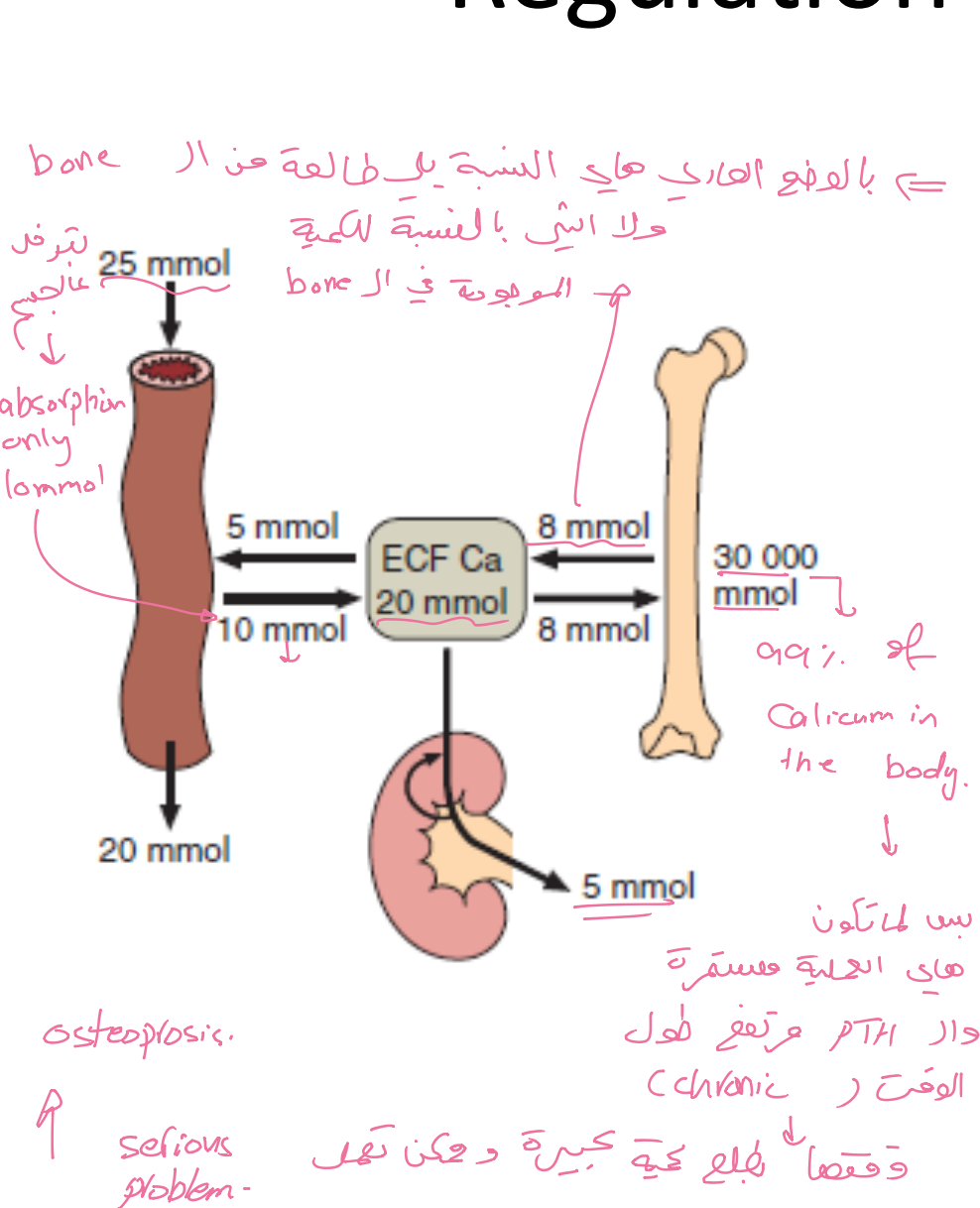
→ secreted in hypercalcemia:

❑ Calcitonin, which originates in the medullary cells of the thyroid gland, is secreted when the concentration of Ca in blood increases (not at normal levels)

❑ Calcitonin reduces Ca by inhibiting the actions of both PTH and vitamin D.

① increase excretion of calcium from kidney. ← low level of calcium

Regulation of calcium



Distribution

- About 99% of calcium in the body is part of bone.
- The remaining 1% is mostly in the blood and other ECF. Little is in the cytosol of most cells.
mainly in blood
- The concentration of ionized calcium in blood is 5,000-10,000 times higher than in the cytosol of cardiac or smooth muscle cells.
release of insulin ← وصار يسهل عليه ان

- Maintenance of this large gradient is vital to maintain the essential rapid inward flux of calcium ions
له اخل الخلية. ← eflux

- Calcium in blood: 45% circulates as free calcium ions (ionized calcium), 40% is bound to albumin, and 15% is bound to anions (bicarbonate, citrate, phosphate, and lactate)
حكيانهم فوق

- Distribution can change in disease as conc. Of citrate, bicarbonate, lactate, phosphate and albumin can change dramatically during surgery or critical care.
problem in kidney or liver → change in normal distribution of calcium.

This calcium cannot be reliably calculated from total calcium measurement (acutely ill patients)

Calcium bound with albumin ← حكيانها (لازم حصة اقصي)
lactate بالدم ← hypoxia ← بالتالي ارجح ترفع نسبة ال ← حكيانها (لازم حصة اقصي)
bind with ionized calcium → hypocalcemia

Causes of hypocalcemia

parathyroid gland
ج. الغدة ← No parathyroid Hormon.

❑ ① Primary hypoparathyroidism-glandular aplasia, destruction, or removal

❑ ② Hypomagnesemia (in hospitalized patients)

→ Cause impaired parathyroid Hormon Secretion.

❑ Hypermagnesemia causes hypocalcemia by three mechanisms:

① ❑ it inhibits the glandular secretion of PTH across the membrane parathyroid gland

② ❑ it impairs PTH action at its receptor site on bone

③ ❑ It causes vitamin D resistance "elevated Mg levels may inhibit PTH release and target tissue response. Perhaps leading to hypocalcemia and hypercalciuria" → high level of calcium in urine.

PTH ← إنتاج الـ PTH
activation of enzyme
that give active form of vitamin D → That help in absorption of Ca^{+} from the intestine.
albumin. ← فاني تصنع الـ albumin.

❑ Hypoalbuminemia (total calcium only, ionized not affected): chronic liver disease, nephrotic syndrome, malnutrition: for each 1g/dL decrease in serum albumin, there is a 0.2 mmol/L (0.8 mg/dL) decrease in total calcium levels

total calcium
إجمالي الكالسيوم

⇒ nephrotic syndrome → loss of albumin in large quantity from kidney.

Causes of hypocalcemia

❑ **Acute pancreatitis:** the cause appears to be a result of increased intestinal binding of calcium as increased intestinal lipase activity occurs

بسبب ارتفاع الـ lipase activity
في الأمعاء
intestine in calcium.

❑ **Vitamin D deficiency and malabsorption** can cause decreased absorption, which leads to increased PTH production or secondary hyperparathyroidism.

PTH ← بسبب نقص فيتامين D
activation of active form from vitamin D.
نقص في امتصاص الكالسيوم
activation قبل في ارتفاع مستوى PTH

❑ **Renal disease** (Altered concentrations of calcium, phosphate, albumin, magnesium and hydrogen ion (pH):

❑ In chronic renal disease, secondary hyperparathyroidism frequently develops as the body tries to compensate for hypocalcemia caused either by hyperphosphatemia (phosphate binds and lowers ionized calcium) or altered vitamin D metabolism

+ hypoalbuminemia + hypermagnesemia + acidosis.

↓
decreases total calcium but ionized will be normal

Causes of hypocalcemia

- ❑ **Pseudohypoparathyroidism**: a rare hereditary disorder in which PTH target tissue response is decreased (end organ resistance)

PTH مستقبلات نقصان
parathyroid hormone defect

- ❑ PTH production responds normally to loss of calcium, however, without normal response (decreased cAMP (Adenosine 3',5'-cyclic phosphate) production), calcium is lost in the urine or remains in the bone storage pool

لأنه قابلية
تأثير لا Calcium وإعادة بناء في bone
①
②
④

- ❑ Patients often have common physical features, including short stature, obesity. Shortened metacarpals and metatarsals, and abnormal calcification.

- ❑ **Rhabdomyolysis**: as with major crush injury and muscle damage, may cause hypocalcemia as a result of increased phosphate release from the cells, which binds to calcium ions.

لأنه داخل ال tissue في phosphate

بعض يطلق ويرتفع ال
Calcium (فشل بسبب أنه ال Calcium - لي جوا الخلايا فلع لأنه مكونا نسبة قليلة جدا
هو الخلايا ففشل يمكن تهر أي تغيير).

۷۴ جن اہم ار patient
یہ لازم آئے
Control
Calicium
عندہم

Surgical and intensive care

- ❑ Controlling calcium concentrations may be critical in open heart surgery when the heart is restarted and during liver transplantation because large volumes of citrated blood are given.

لے بیو خندا عیات کثیرہ جن

وحدات الدم . سے عشان احفظ الدم تكون موجود فيه citrate سے یہ بھری وحدات الدم

بہتر ال citrate پر ہوا سے ar citrate سے بعض hypo سے بہتر المرفق عندہ سنجار و size

- ❑ Ionized Ca measurements are the measurement of greatest clinical value.

- ❑ Hypocalcemia occurs commonly in critically ill patients, that is, those with sepsis, thermal burns, renal failure, or cardiopulmonary insufficiency (abnormalities of acid-base regulation and losses of protein and albumin)

حديث الولادة .

Neonatal monitoring

- Typically blood-ionized calcium concentrations in neonates are high at birth and then rapidly decline by 10-20% after 1-3 days. After about 1 week, ionized calcium concentrations in the neonate stabilize at levels slightly higher than in adults.

أول عانيول
يكون ال

Calcium
عالي
جدا

stable → ارتفاع بسيط عن ال
adult

يمكن الانخفاض يكون سريع جدا

و يكون ال neonate مش قادر يعوضه ← hypo

بعد كم يوم
يبدأ ليخف
من 2-3 ما

- The concentration of ionized calcium may decrease rapidly in the early neonatal period because the infant may lose calcium rapidly and not readily reabsorb it.

- Several possible etiologies have been suggested: abnormal PTH and vitamin D metabolism, hypercholesterolemia, hyperphosphatemia, and hypomagnesemia.

4

5

1

2

3

Symptoms of hypocalcemia

- ❑ Neuromuscular irritability and cardiac irregularities are the primary groups of symptoms that occur with hypocalcemia.
- ❑ Neuromuscular symptoms include parasthesia, muscle cramps, tetany, and seizures.
- ❑ Cardiac symptoms may include arrhythmia or heart block.
- ❑ Symptoms usually occur with severe hypocalcemia, in which total calcium levels are below 1.88 mmol/L (7.5 mg/dL).

Normal level (8.6 - 10.8)

sever
hypocalcemia

دليل على انه في

Treatment of hypocalcemia

له لازم أدرج السبب لهاد الإخفاض + ١

Severity of decrease.

❑ Oral or parenteral calcium therapy may occur, depending on the severity of the decreased level and the cause.

or diet
↓

انخفاض

تناول الأطعمة

داخلية (البنفسج) إلى خضراوات

❑ Vitamin D may sometimes be administered in addition to oral calcium to increase absorption.

يمكن
يكون

هو السبب

بعد الإخفاض

يمكن يكون الـ هو السبب

❑ If hypomagnesemia is a concurrent disorder, magnesium therapy should also be provided

له يعطى بين الـ و تدخل المشكلة

Hypercalcemia

- ❑ Although ^①either total or ^②ionized calcium measurements serious cases, ionized calcium is more frequently elevated in asymptomatic hyperparathyroidism.
لے آحد اسباب ارتفاع ال Ca^{+}
- ❑ In general, ionized calcium measurement elevated in 90-95% of cases of hyperparathyroidism whereas total calcium is elevated in 80-85% of the cases

Causes of hypercalcemia

① ☐ Primary hyperparathyroidism (in old women)-adenoma or glandular hyperplasia

↓
cancer in parathyroid gland

② ☐ hyperthyroidism, because of the proximity of the parathyroid gland to the thyroid gland, hyperthyroidism can sometimes cause hyperparathyroidism.

③ ☐ Benign familial hypocalciuria

low level of calcium in urine (mainly reabsorption of calcium).

④ ☐ Malignancy:

☐ Ca is sole biochemical marker for many tumors.

① ☐ Many tumors produce PTH-related peptide (PTH-rP), which binds to normal PTH receptors and causes increased calcium levels

☐ Assays to measure PTH-rP are available because this abnormal protein is not detected by most PTH assays

↓
level 2 یا 1
1. normal or 2. low.

↓
specific antigen
antibodies reaction

detection

Causes of hypercalcemia

- 5 ☐ **Thiazide diuretics** increase calcium reabsorption
↳ hypokalemia + Hyponatremia → Hypercalcemia.
- 6 ☐ **Prolonged immobilization** may cause increased bone resorption. This cause is further compounded by renal insufficiency
- 7 ☐ **Multiple myeloma**
- 8 ☐ **Increased vitamin D**

Symptoms of hypercalcemia

- ❑ A mild hypercalcemia (2.62-3.00 mmol/L) is often asymptomatic.
- ❑ Moderate or severe calcium elevations include:
 - ❑ Neurologic symptoms: mild drowsiness or weakness, lethargy, and coma depression,
① ② ③ ④
 - ❑ GI: constipation, nausea, vomiting, anorexia, and peptic ulcer disease.
① ② ③ ④
 - ❑ Hypercalcemia may cause renal symptoms of nephrolithiasis and nephrocalcinosis.
① ② ③ ④ ⑤ ⑥
- ❑ Hypercalciuria can result in nephrogenic diabetes insipidus, which causes polyuria and results in hypovolemia, which further aggravates the hypercalcemia.
① ② ③
- ❑ Hypercalcemia can also cause symptoms of digitalis toxicity.

Treatment of hypercalcemia

لأنه لازم أعرف سبب الارتفاع

- ❑ Depends on the level of hypercalcemia and the cause.
- ❑ Often people with primary hyperparathyroidism are asymptomatic. Postmenopausal women with primary hyperparathyroidism may have estrogen deficiency → السبب بالـ hyperparathyroidism → give estrogen therapy حل المشكلة
- ❑ **Parathyroidectomy** may be necessary in some hyperparathyroidic patients ^{علاج جراحية}
- ❑ Often, estrogen replacement therapy reduces calcium levels.
- ❑ Patients with moderate to severe hypercalcemia are treated by:
 - ❑ Encouraging salt and water intake to increase calcium excretion and avoid dehydration
 - ❑ Thiazide diuretics should be discontinued.
 - ❑ Biphosphanate (a derivative of pyrophosphate) are the main drug class used to lower calcium levels by preventing bone resorption, achieved by its binding action to bone.
لأنه يمنع الـ calcium عن طريق ارتباطه مع الـ bone.
of calcium from bone

Determination of Calcium

- ❑ For total calcium determinations is either ^①serum or ^②lithium heparin plasma collected without venous stasis.

← بدون و استغنى كثير

- ❑ Anticoagulant such as **EDTA** or **oxalate** bind calcium tightly and interfere with the measurement, so, their use should **be avoided**

← لانها ترتبط مع ال calcium الموجود بالعينة .

- ❑ The proper collection of samples for ionized calcium measurements requires greater care. Because loss of CO₂ will increase pH, samples must be collected anaerobically.

← ترتبط ال calcium في هائي الحالة . (turbidity)

بفطيرة منع

- ❑ Although heparinized whole blood is the preferred sample, serum from sealed evacuated blood collection tubes may be used if clotting and centrifugation are done quickly (<30 minutes) and at room temperature.

← بعد استئني رهير عندي clotting رهدين بعد Centrifugation و بجلد العينة .
أكثر
← لازم

Determination of Calcium

- ☐ No liquid heparin products should be used. Most heparin anticoagulants (sodium, lithium) partially bind to calcium and lower ionized calcium concentrations
- ☐ Dry heparin products are available titrated with small amounts of Ca or Zn ions that essentially eliminates the interference by heparin.
- ☐ For analysis of calcium in urine, an accurately timed urine collection is preferred, acidified with approximately 1 ml of HCl (6M) for each 100 ml of urine

Method

- ❑ The two commonly used methods for total calcium analysis are:
- ❑ Use of ortho-cresolphthalein complexone (CPC) or arsenzo III dye to form a complex with calcium.
 - measured total calcium
 - ❑ Prior to the dye binding reaction, calcium is released from its protein carrier and complexes by acidification of the sample.
 - acidification ← زيادة الحموضة
 - ❑ The CPC method uses 8-hydroxyquinoline to prevent magnesium interference.
 - ← لأنه يثبط تفاعل مع زئبق ال reagent
- ❑ Use ISEs:
 - ion selective electrode.
→ ionized calcium.
 - ❑ Use membranes impregnated with special molecules that selectively, but reversibly, bind calcium ion.
 - ← بعض قراءة potential ← بعض قراءة calcium ← كانت القراءة أعلى.
 - ❑ As calcium ions bind to these membranes, an electric potential develops across the membrane that is proportional to the ionized calcium concentration.
 - ← معمم
- ❑ AAS remains the reference method for total calcium
 - ↓
atomic absorption spectroscopy

Reference Ranges

TOTAL CALCIUM (SERUM, PLASMA)

Child	2.20–2.70 mmol/L (8.8–10.8 mg/dL)
Adult	2.15–2.50 mmol/L (8.6–10.0 mg/dL)

أعلى مستوى
يكون الأنتم
في حالة غو

IONIZED CALCIUM (SERUM)

Neonate	1.20–1.48 mmol/L (4.8–5.9 mg/dL)
Child	1.20–1.38 mmol/L (4.8–5.5 mg/dL)
Adult	1.16–1.32 mmol/L (4.6–5.3 mg/dL)
Urine (24-hour)	2.50–7.50 mmol/day (100–300 mg/day), varies with diet

Magnesium

- ❑ The average human body (70 kg) contains 1 mole (distributed as 24 g) of magnesium
 - ❑ 53% in bone
 - ❑ 46% in muscle and other organs and soft tissue
 - ❑ less than 1% in serum and RBC's (1/3 bound to albumin)
- ❑ Similar to calcium, it is the free ion that is physiologically active in the body
- ❑ The role magnesium in the body is:
 - ① ❑ It is an essential cofactor of more than 300 enzymes
 - ❑ The most significant findings are the relationship between abnormal serum magnesium levels and cardiovascular, metabolic, and neuromuscular disorders. Although serum levels may not reflect total body stores of Mg, serum level is useful in determining acute changes in the ion

Regulation

- ❑ Rich sources of Mg in the diet: raw nuts, dry cereal, and “hard” drinking water.
(1) (2) (3)
- ❑ Other sources include vegetables, meats, fish, and fruit.
(4) (5) (6) (7)
- ❑ Processed foods have low levels of magnesium that may cause an inadequate intake.
→ low
- ❑ The small intestine may absorb 20-65% of the dietary Mg, depending on the need and intake.
→ depend on need and intake.
- ❑ The overall regulation of body magnesium is controlled largely by the kidney which can reabsorb magnesium in deficiency states or readily excrete excess magnesium in overload states.
(1) (2)
- ❑ Henle loop is the major renal regulatory site, where 50-60% of filtered Mg is reabsorbed in the ascending limb.
→ reabsorption.

Regulation

زيادة عن الحد ← بعمل excretion
نقصان عنه ← بعمل reabsorption

- ❑ The renal threshold for magnesium is about 0.60-0.85 mmol/L (close to normal serum conc.), so slight excesses of magnesium in serum are rapidly excreted by the kidneys

- ❑ Normally only about 6% of filtered Mg is excreted in the urine per day

- ❑ Magnesium regulation appears to be related to that of calcium and sodium

↓ Calcium → ↑ PTH → ↑ reabsorption of mg from kidney + absorption of mg from intestine.

- ❑ Parathyroid hormone (PTH) increases the renal reabsorption of magnesium and enhances the absorption of magnesium in the intestine.

- ❑ Changes in ionized calcium have a far greater effect on PTH secretion.

- ❑ Aldosterone and thyroxine apparently have the opposite effect of PTH in the kidney, increasing the renal excretion of magnesium.

لازم اعرف من ديف المشككة
 ↓ ال cat قليل ← بزيادة PTH ← بعمل hypermagnesi (↑ mg)
 ← من ال mg ← hypo or hyper ← بعمل hypocalcemia (↓ cat)

① كبحار العيون
الأعشرفية ←
② hospitalized patient

Hypomagnesemia

- ❑ Most frequently observed in hospitalized individuals in intensive care units of those receiving diuretic therapy or digitalis therapy
- ❑ Hypomagnesemia is rare in nonhospitalized individuals
- ❑ There are many causes of hypomagnesemia
 - ① ❑ Reduced intake Poor diet/starvation, Prolonged magnesium-deficient IV therapy, chronic alcoholism.
 - ② ❑ Decreased absorption: due to GI disorders as malabsorption syndrome, surgical resection of the small intestine, nasogastric suction, pancreatitis, prolonged vomiting, diarrhea, laxative abuse, neonatal (due to surgical procedure), primary (due to selective malabsorption of the ion), congenital (autosomal recessive disorder)
 - ❑ A chronic congenital hypomagnesemia with secondary hypocalcemia occurs due to specific transport protein defect in the intestine

يرتفع الـ
lipas
enzyme
وبنية السبيل الـ
magnesium
بالـ
intestin
↓
hypo

قوى للأعشرف

نتيجة الـ
hypomagnesemia

Causes of hypomagnesemia

③ ☐ Increased Excretion

☐ **Renal:** Tubular disorder, Glomerulonephritis, Pyelonephritis

↳ No reabsorption of magnesium. → excreted in urine.

☐ **Endocrine:** Hyperparathyroidism (increased calcium)

② Hyperaldosteronism (increase of Mg excretion and water retention (pseudohypomagnesemia), hyperthyroidism (increase excretion cause intracellular shift of ions), hypercalcemia, diabetic ketoacidosis (increase renal loss due to glycosuria)

☐ **Drug induced** (increase renal loss of Mg): diuretics, antibiotics (gentamicin), cyclosporine, and cisplatin (nephrotoxic), digitalis (interfere with Mg reabsorption)

☐ **Miscellaneous:** Excess lactation (loss in milk)), Pregnancy (may cause a hyperexcitable uterus, anxiety and insomnia)

↓
(تسم الحمل) preeclampsia
mg ↓ (نقص)

+ hypertension

③

diuretic effect
↓
increase excretion of magnesium in urine
↳ hypo

cause polyuria
↓
I see ↓ Mg
magnesium
↓
hypomagnesemia

Symptoms of hypomagnesemia

راجع لفيال asymptomatic كدوا قیل عن 0.5 ے تبلیش زفهر الاعراض

- ❑ A patient who is hypomagnesemic may be asymptomatic until serum levels fall below 0.5 mmol/L. normal level = 0.63
- ❑ A variety of symptoms can occur. The most frequent involve cardiovascular, neuromuscular, psychiatric, and metabolic abnormalities

Cardiovascular	Psychiatric
1. Arrhythmia	1. Depression
2. Hypertension	2. Agitation
3. Digitalis toxicity	3. Psychosis
Neuromuscular	Metabolic
1. Weakness	1. Hypokalemia
2. Cramps	2. Hypocalcemia
3. Ataxia	3. Hypophosphatemia
4. Tremor	4. Hyponatremia
5. Seizure	
6. Tetany	
7. Paralysis	
8. Coma	

كله :
hypo

Symptoms of hypomagnesemia

❑ Cardiac:

- ❑ Symptoms result primarily from the ATPase enzyme's Mg. requirement for Mg
- ❑ Mg loss leads to decreased intracellular K levels because of a faulty NaK pump (ATPase)
- ❑ This change in cellular RMP causes increased excitability that may lead to cardiac arrhythmia and digitalis toxicity

→ sodium - potassium ATPase

→ resting membrane potential

❑ Normal nerve and muscle cell stimulation: (↑ contractility + excitation).

- ❑ Requires magnesium and ATPase for normal calcium uptake following contraction.
- ❑ Requires magnesium to assist with the regulation of acetylcholine, a potent neurotransmitter

① co factor

② after contraction

need mg

release of me c lines

❑ Metabolic disorders:

- ❑ Mg deficiency can impair PTH release and target tissue response, resulting in hypocalcemia. Mg therapy alone may restore both ions levels to normal
- ❑ Serum levels of the ions must be monitored during treatment.

→ to avoid hypermagnesemia

Treatment of hypomagnesemia

- ❑ The preferred treatment of hypomagnesemia by oral intake is Mg-lactate, Mg oxide, MgCl or an antacid that contains Mg.
- ❑ In severely ill patients, a MgSO₄ solution is given parenterally
2x sulfate
- ❑ Before initiation of therapy; renal function must be evaluated to avoid inducing hypermagnesemia during treatment

Hypermagnesemia and its causes

- ❑ less frequently than hypomagnesemia
- ❑ The most common cause is **renal failure** (GFR <30 severe elevations are usually a result of the combine effects of decreased renal function and ^①increased intake of commonly prescribed magnesium-containing medication, such as antacid, ^②enemas, or cathartics.
- ❑ ^{دور العجزة} Nursing home patients are at greatest risk for this occurrence.
- ❑ **Decreased excretion**: ^①acute or chronic renal failure, ^②hypothyroidism, hypoadosteronism, hypopituitarism (IGH)
↳ impaired growth hormon.
- ❑ **Increased intake**: Antacids, enemas, cathartics, therapeutic-eclampsia, cardiac arrhythmia
- ❑ **Miscellaneous**: ^①dehydration (pseudohypermagnesemia, corrected by dehydration), ^②bone carcinoma, bone metastases (high Mg due to bone loss)
↳ 53% in bones of mg.

Causes of hypomagnesemia

- ❑ **Endocrine disorders:** Thyroxine and growth hormone cause a decrease in tubular reabsorption of Mg and of either hormone may cause a moderate elevation in serum Mg.
- ❑ **Adrenal insufficiency** may cause a mild elevation as a result of decreased renal excretion of Mg
- ❑ $MgSO_4$ may be used therapeutically with preeclampsia, cardiac arrhythmia, or myocardial infarction
تسبب الحمل
1 *2*
3
- ❑ Mg is a vasodilator, and can decrease uterine hyperactivity in eclamptic states and increase uterine blood flow (maternal hypermagnesemia)
1 *2*
له اذا حائل مراقبة خلال العلاج
- ❑ Neonatal hypermagnesemia due to the immature kidney of the newborn (Premature infants are at great risk)

Symptoms of hypermagnesemia

- ❑ Hypermagnesemia typically do not occur until the serum level exceeds 1.5 mmol/L.
- ❑ Most frequent symptoms involve cardiovascular^①, dermatological^②, GI^③, neurologic^④, neuromuscular^⑤, metabolic^⑥, and hemostatic abnormalities^⑦.
- ❑ Mild to moderate symptoms may occur when serum levels are 1.5-2.5mmol/L: hypotension, bradycardia, skin flushing, increased skin temperature, nausea, vomiting, and lethargy.
vasodilation.
- ❑ Life-threatening symptoms, such as ECG changes^①, heart block^②, asystole^③, sedation^④, coma^⑤, respiratory depression or arrest^⑥ and paralysis^⑦, can occur when serum levels reach 5.0 mmol/L.

Symptoms of hypermagnesemia

❑ Elevated Mg levels may inhibit PTH release and target tissue response. This may lead to hypocalcemia and hypercalciuria

❑ Normal hemostasis is a calcium-dependent process that may be inhibited as a result of competition between increased levels of magnesium and calcium ions. Thrombin generation and platelet adhesion are two processes in which interference may occur.

→ very important for coagulation. → ↓ Ca²⁺ → Coagulation
فالمهم
لبنك
هو

Treatment of hypermagnesemia

- ❑ If Mg excess associated with increased intake one should discontinue the source of Mg. ①
- ❑ Severe symptomatic hypermagnesemia requires immediate supportive therapy for cardiac, neuromuscular, respiratory or neurologic abnormalities. ① ② ③ ④
- ❑ Patients with renal failure require hemodialysis. غسيل كلوي
- ❑ Patients with normal renal function may be treated with a diuretic and IV fluids ① ②

لـ لازم انتبه لا hymolized لأنه في كحلات
 جيرة جوا الخلايا من الكالسيوم لومار hymolysis ← الحة - نفي القراءة.

- 1 2

Method

- ❑ The three most common methods for measuring total serum Mg are colorimetric:
 - ❑ Calmagite: Mg binds with calmagite to form a reddish-violet complex that is read at 532 nm
 - ❑ Formazen dye: Mg binds with the dye to form a colored complex that is read at 660 nm
 - ❑ Methylthymol blue: Mg binds with the chromogen to form a colored complex
- ❑ Most methods use a calcium shelter to prohibit interference from Ca.
- ❑ The reference method for measuring magnesium is AAS.

atomic absorption
spectroscopy.

Limitations of Mg determination

- ❑ Although the measurement of total Mg conc in serum remains the usual diagnostic test for the detection of magnesium abnormalities, it has limitations:

Because approximately 25

- ❑ Because approximately 25% of magnesium is protein bound,
total magnesium may not reflect the physiologically active
free ionized magnesium.

حذف يكون عند depletion خارج الخلية حاي زيادة الجسم قاعد بفرجه برا.

- ❑ Because magnesium is primarily an intracellular ion, serum concentration will not necessarily reflect the status of intracellular magnesium (depletion of 20% of cellular Mg, serum magnesium concentrations may remain normal)

TABLE 13-15. REFERENCE RANGE FOR MAGNESIUM

Serum, plasma	0.63–1.0 mmol/L (1.2–2.1 mEq/L)
---------------	---------------------------------

Phosphate → main intracellular anion.

❑ Found everywhere in living cells: DNA, RNA, in most coenzymes - NADPH

❑ The most important reservoirs of biochemical energy are ATP, creatine, phosphate, and phosphoenolpyruvate.

❑ Phosphate deficiency can lead to ATP depletion, which is ultimately responsible for many of the clinical symptoms observed

❑ Alteration in the concentration of 2,3-bisphosphoglycerate (2,3-BPG) in red blood cells affect the affinity of hemoglobin for oxygen, the concentration of inorganic phosphate indirectly affects the release of oxygen from hemoglobin

Handwritten notes for the 2,3-BPG block:
- decrease affinity of HB to O_2 → تستخذه الخلايا غشائاً تحولاً NADH to ATP (في الحالة الطبيعية)
- لا يفس عندي مشكلة باد phosphate - راجع تا شرح على تنفس الخلية.
- metabolizing tissue يكون قريب عند ال

❑ Transcellular shifts of phosphate are a major cause of hypophosphatemia in blood. Once phosphate is taken up by the cell, it will be used in the synthesis of phosphorylated compounds. As these phosphate compounds are metabolized, P_i slowly leaks into the blood, where it is regulated principally by the kidney

Handwritten notes for the hypophosphatemia block:
- regulation by kidney.
- لا يتأثر بعض inorganic phosphate
- يستخذه / الفضلات غشائاً تنتج phosphate
- كرياتين phosphate
- زي ال

Regulation

- ❑ Phosphate in blood may be absorbed in the intestine from dietary sources, released from cells into blood and lost from bones. In healthy individuals, all these processes are relatively constant and easily regulated by renal excretion or reabsorption of phosphate.

↳ if there high level ↳ if there low level

- ❑ Many factors can alter phosphate concentrations in the blood:
 - ❑ The loss of regulation by the kidneys will have the most profound effect
 - ❑ The most important factor is PTH, which overall lowers blood conc. By increasing renal excretion.
 - ❑ Vitamin D acts to increase phosphate in the blood by increasing both phosphate absorption in the intestine and phosphate reabsorption in the kidney.
 - ❑ Growth hormone, which helps regulate skeletal growth, can affect circulating concentration of phosphate
 - ❑ Excessive secretion or administration of growth hormone, phosphate concentrations in the blood may increase because of decreased renal excretion of phosphate.
 - ❑ Calcitonin, acid-base status, can also affect renal regulation of Phosphate

PTH ↓ ↓ ↓ ↑ phosphate.

Distribution

- ❑ Although the concentration of all phosphate compounds in blood is about 12 mg/dl (3.9 mmol/L), only about 3-4 mg/dL is inorganic phosphate.
- ❑ Phosphate is the predominant intracellular anion, with variable concentrations depending on the type of cell.
- ❑ About 80% of the total body pool of phosphate is contained in bone, 20% in soft tissues, and less than 1% is active in the serum/plasma.

Hypophosphatemia

- ❑ Hypophosphatemia occurs in about 1-5% of hospitalized patients.
- ❑ The incidence of hypophosphatemia increases to 20-40% in patients with:
 - ❑ diabetic ketoacidosis, chronic obstructive pulmonary disease (COPD), asthma, malignancy, long-term treatment with total parenteral nutrition (TPN), inflammatory bowel disease, anorexia nervosa, and alcoholism.
- ❑ The incidence increases to 60-80% in:
 - ❑ ICU patients with sepsis.
 - ❑ increased renal excretion (*Hyperparathyroidism*)
 - ❑ Hyperparathyroidism (*increase excretion of phosphate*).
 - ❑ Decreased intestinal absorption (*Vitamin D deficiency*).
 - ❑ vitamin D deficiency
 - ❑ Antacid use (*bind with phosphate and decrease its absorption*).
- ❑ Although most cases are moderate and seldom cause problems severe hypophosphatemia ($<1\text{mg/dl}$ or 0.3 mmol/L) requires monitoring and possible replacement therapy

Hyperphosphatemia

- ❑ Patients at greatest risk for hyperphosphatemia are those with acute or chronic renal failure
- ❑ An ^①increased intake of phosphate or ^②increased release of cellular phosphate may also cause hyperphosphatemia
- ❑ Neonates: not developed mature PTH and vitamin D metabolism, hyperphosphatemia is caused by increased intake such as from cow's milk or laxatives.
- ❑ Increased breakdown of the cells as with severe infections, intensive exercise, neoplastic disorders, or intravascular hemolysis
↳ cancer → precursor of lymphocyte cells.
- ❑ Because immature lymphoblasts have about 4 times the phosphate content of mature lymphocytes, patients with lymphoplastic leukemia are especially susceptible to hyperphosphatemia.

Determination of inorganic phosphorus

- ❑ Specimen. ^① Serum or ^② lithium heparin plasma is acceptable for analysis.
- ❑ Oxalate, citrate, or EDTA anticoagulants should not be used because they interfere with the analytic method.
لے بٹفاعلوں سے اجتناب رکھنا۔ reagent کی بے ساختہ استعمال نہ کرنا۔
- ❑ Hemolysis should be avoided because of the higher concentrations inside the red cells.
- ❑ Circulating phosphate levels are subject to circadian rhythm, with highest levels in late morning and lowest in the evening.
Urine analysis for phosphate requires a 24-hour sample collection because of significant diurnal variations.

Methods and reference ranges

Reagent used \rightarrow molybdenum.

- ❑ Phosphorus determination methods involve the formation of an ammonium phosphomolybdate complex. This colorless ^① complex can be measured by ultraviolet absorption at 340 nm or can be ^② reduced to form molybdenum blue, a stable blue chromophore, which is read between 600 and 700 nm.

- ❑ Normal ranges:

SERUM, PLASMA

Neonate	1.45–2.91 mmol/L (4.5–9.0 mg/dL)
Child	1.45–1.78 mmol/L (4.5–5.5 mg/dL)
Adult	0.87–1.45 mmol/L (2.7–4.5 mg/dL)
Urine (24-hour)	13–42 mmol/day (0.4–1.3 g/day)

يعطيكم العافية جميعا
ما تنسوننا من دعواتكم
زميلتكم مرام الزبيدي

CASE STUDY 13-2

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 13-2.1.

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 13-2.1. 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

HCO₃⁻: 25 mmol/L

Mg²⁺: 0.4 mmol/L

Ion/free Ca²⁺: 1.0 mmol/L

CASE STUDY 13-3

An 84-year-old nursing home resident was seen in the emergency department with the following symptoms: nausea, vomiting, decreased respiration, hypotension, and low pulse rate (46). Physical exam showed the skin was warm to the touch and flushed. Admission lab data are found in Case Study Table 13-3.1.

Questions

1. What is the most likely cause for the patient's symptoms?
2. What is the most likely cause for the hypermagnesemia?
3. What could be the cause for the hypocalcemia?

CASE STUDY TABLE 13-3.1. LABORATORY RESULTS

		RESULT	REFERENCE RANGE
Serum	Total protein	5.6 g/dL	6.0–8.0 g/dL
	Albumin	3.0 g/dL	3.5–5.0 g/dL
	Total calcium	8.2 g/dL	8.6–10.0 g/dL
	BUN	45 mg/dL	5–20 mg/dL
	Creatinine	2.3 mg/dL	0.7–1.5 mg/dL
Plasma	Magnesium	4.0 mmol/L	0.63–1.0 mmol/L
	Na ⁺	129 mmol/L	136–145 mmol/L
	K ⁺	5.3 mmol/L	3.4–5.0 mmol/L
	Cl ⁻	96 mmol/L	
	HCO ₃ ⁻	16 mmol/L	

CASE STUDY 13-4

Consider the following laboratory results from three adult patients:

Questions

- Which set of laboratory results (Case A, B, or C) is most likely associated with each of the following diagnoses:
 - Primary hyperparathyroidism
 - Malignancy
 - Hypomagnesemic hypocalcemia

CASE STUDY TABLE 13-4.1. LABORATORY RESULTS

CASE	REFERENCE RANGES				
	ION Ca^{++} 1.16–1.32 mmol/L	TOTAL Mg^{++} 0.63–1.0 mmol/L	PO_4^{--} 0.87–1.45 mmol/L	HEMATOCRIT 35–45%	INTACT PARATHYROID HORMONE 13–64 ng/L
A	1.44	0.90	0.85	42	100
B	1.08	0.50	0.90	40	25
C	1.70	0.98	1.43	30	12