



Amino acid metabolism and plasma proteins

يعطيكم العافية جميعا ان شاء الله تكون دراستكم
ماشية تمام ركزوا كثير على هاد الشابتة لأنه
الدكتورة حكت كثير مهم

Amino acids

➤ Amino acids in blood are used in:

① ➤ Synthesis of plasma, intracellular and structural proteins

② ➤ Synthesis of nonprotein nitrogen containing compounds: purines, pyrimidines, porphyrins, creatine, histamine, thyroxine, epinephrine and coenzyme NAD → all of them synthesis from amino acid.

③ ➤ Body energy: 12-20% of energy is due to proteins

④ ➤ The ammonium produced during deamination of amino acids is converted into urea in liver

metabolism of
amino acid

↓
produce ammonium

↓
urea cycle

↓
excreted from
the body

209
F. J.

non-polar amino acid

proline

Contain amide

↳ Non-polar.

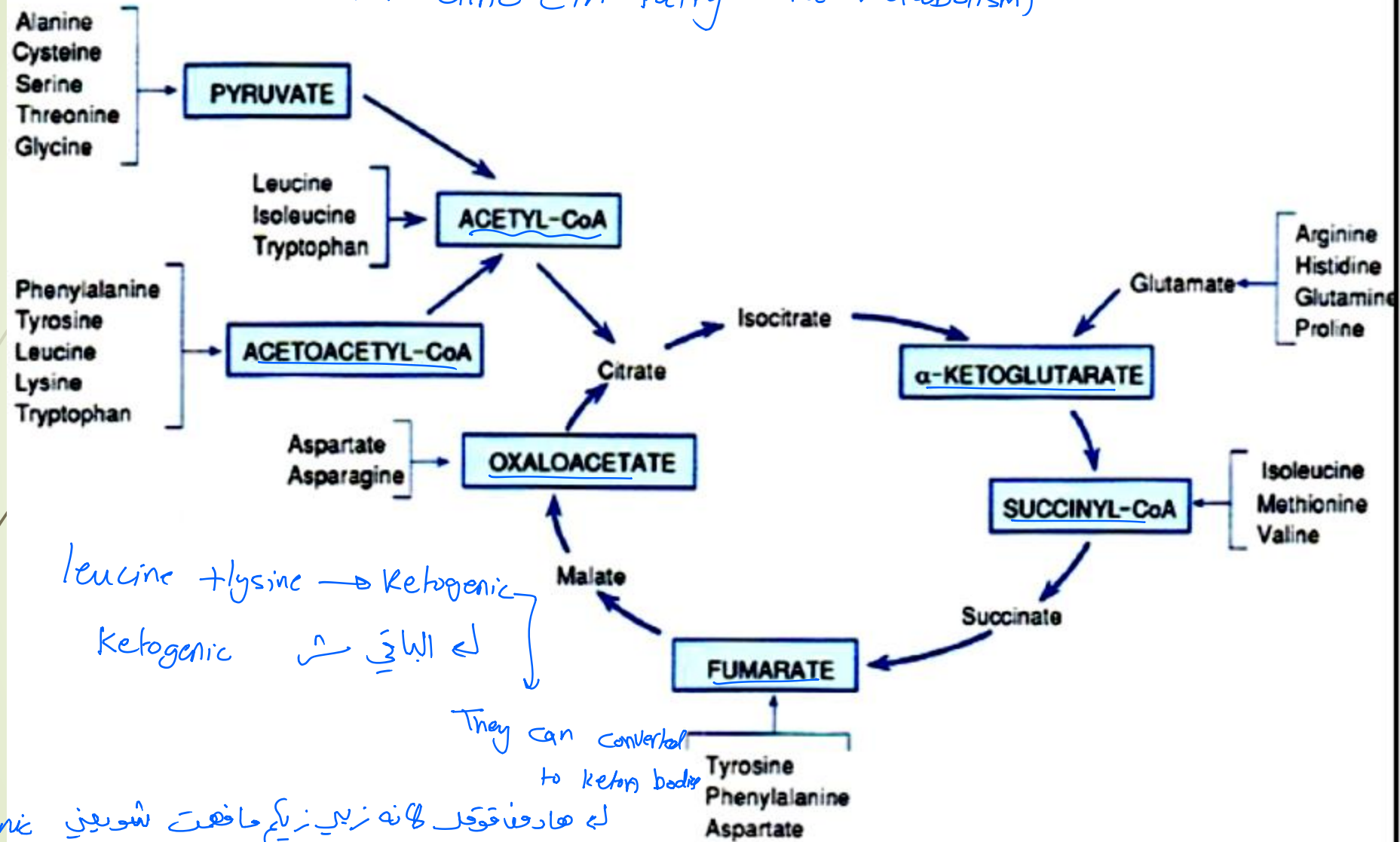
$\text{N} \rightarrow \text{N within cycle}$

alpha helices

lysine
+
Arginine
+
histidine
↓
basic
amino
acid

carboxylic acid

amino acid don't used in protein synthesis → 1. Ornithine (in urea cycle)
 2. Carnitine (in fatty acid metabolism)

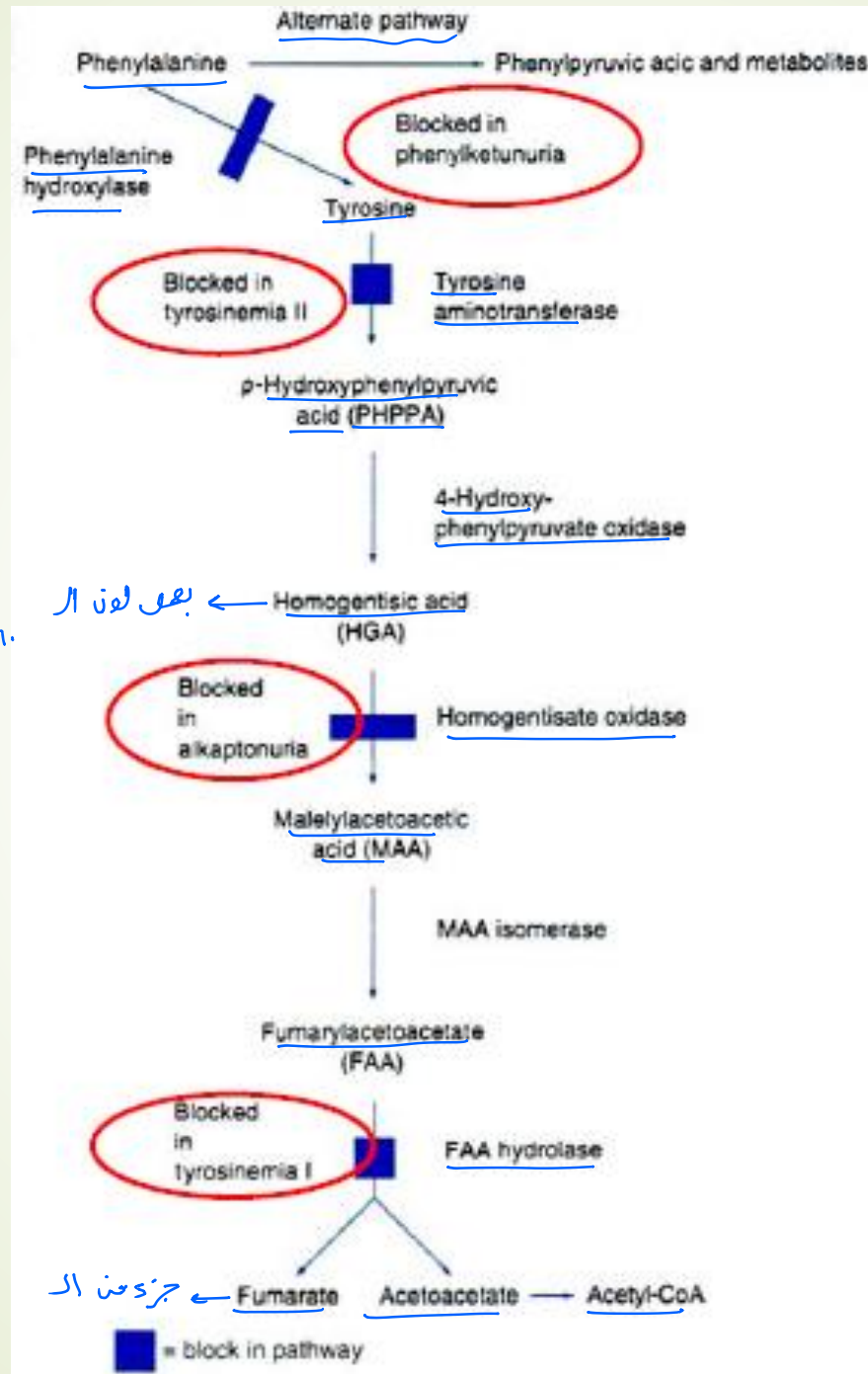


Aminoacidopathies

المشاكل التي يمكن تصنيفها إلى 1) نقص enzyme المولود به و 2) membrane transport system of amino acids. (metabolic pathway)

- Can be in the activity of specific enzyme in the metabolic pathway ①
- Membrane transport system for amino acids ②
- Diseases to talk about:
 - Phenylketonurea
 - Maple syrup urine disease (MSUD)
 - Homocystinuria
 - Argeninosuccinic aciduria and citrullinemia
 - Cystinurea

خطوة
شرحها
الخطوة



urine
brown.

crebs
cycle.

phenylalanine
metabolism

Alternative ←

↓

block
pathway
أو defect في ال
الإنزيم

Phenylketonuria (PKU)

- عرفه ورأى لاجتاجه ← 2 gens عشان يظهر عندي
An autosomal recessive genetic disorder characterized by a deficiency in the hepatic enzyme phenylalanine hydroxylase (PAH)
- The PAH gene is located on chromosome 12 → mutation مش حلوبوة مكان حدد.
↓
phenylalanine تحول ال
to
tyrosine
(hydroxylation process.)
- More than four hundred disease-causing mutations have been found in the PAH gene
400
- PAH is necessary to metabolize the amino acid phenylalanine to tyrosine
↓
essential amino acid → لانه لايستطيع الجسم
البنفسه تحويله للجسم
- When deficient, phenylalanine accumulates to a level $> 1200 \mu\text{mol/L}$ and metabolized by alternative pathways.

Phenylketonuria (PKU)

- The metabolites which are detected in blood and urine include:
- Phenylpyruvic acid (which known as phenylketone): which is the product of deamination of phenylalanine ↓ double bond oxygen
- phenyllactic acid: which is the reduction product of phenylpyruvic acid ↳ OH
- Phenylacetic acid which is produced by decarboxylation and oxidation of phenylpyruvic acid
- And phenylacetylglutamine: which is the glutamine conjugate of phenylacetic acid
- These metabolites give urine musty odor لم رائحة سيئة لـ urine .

Phenylketonuria (PKU)

deficiency of the enzyme
phenylalanine
عند ۱۲۰۰

- Variants of the disease result from partial deficiencies of PAH activity and are typically classified as:
 - Mild PKU if phenylalanine levels are between 600 and 1200 $\mu\text{mol/L}$
 - Non-PKU mild hyperphenylalaninemia which present with phenylalanine levels in the range of 180-600 $\mu\text{mol/L}$ and no accompanying accumulation of phenylketones.
- The normal limits for serum phenylalanine levels for full term, normal weight newborns range from 1.2 to 3.4 mg/dL (70-200 $\mu\text{mol/L}$)

حافظی
symptoms.

defect of the enzyme → severe or mild mutation of the enzyme
activity of the enzyme

Phenylketonuria (PKU)

× عن المشكلة ما تكون بال enzyme
↓
defect in cofactor

- A rarer of the disease occurs when PAH is normal but there is a defect in the biosynthesis of the **cofactor tetrahydrobiopterin (BH4)** by the patient which is necessary for proper activity of the enzyme (for PA, tyrosine and tryptophan hydroxylation)
- It results in hyperphenylalaninemia, that are not responsive to dietary treatment
- Examination of urinary proteins is helpful in diagnosis
- Although cofactor defects are rare, they must be identified so that appropriate treatment can be initiated
- Patients must be given the active cofactor a long with the neurotransmitter precursor L-dopa and 5-OH tryptophan

treatment → cofactor + L-dopa + tryptophan.

Phenylketonuria (PKU)

الضرر يبدأ من الأسبوع الثاني لثالث
من الولادة.

- Left untreated, this condition can cause problems with brain development, leading to progressive mental retardation^① and seizures^②
- In infants and children, the deterioration of brain function begins in the second or third week of life
- Brain damage can be avoided if the disease is detected at birth and the infant is maintained on a diet containing very low level of phenylalanine and high levels of tyrosine
- There is no cure. Damage done is irreversible so early detection is crucial.

علاج
فائي

بعض الطفل يجب فائيه
phenylalanine
+
high level of tyrosine.

Screening for PKU

بوختن ال spores of bacillus دجھائی agar
وچھائی ا.ا. بائینا یا phenylalanine

inhibitor of growth ← Bacillus
Metabolism of Bacillus

positive result ← growth ← agar
Bacillus growth ← phenylalanine
↓
filter paper ← blood of the infant

➤ The Guthrie bacterial inhibition assay:

➤ Spores of the organism Bacillus subtilis are incorporated into an agar plate that contains β 2-thienylalanine, a metabolic antagonist to B. subtilis growth.

➤ A filter paper disk impregnated with blood from the infant is placed on the agar

➤ If the blood level exceeds the range of 2-4 mg/dL, the phenylalanine counteracts the antagonists and bacterial growth occurs.

➤ To avoid false-negative results, the infant must be at least 24 hours old to ensure adequate time for enzyme and amino acid levels to develop

➤ The sample should be taken before administration of antibiotics or transfusion of blood or blood products

➤ Premature infant can show false positive results due to the immaturity of the liver enzyme systems

↓
liver enzyme
نقص و کثافت

positive result ← growth
Bacillus growth ← phenylalanine
↓
filter paper ← blood of the infant

فکین ریلو عنی
false negative result.

① عمر الطفل أقل من
24 ساعة.

↓
amino acid
عنی
نقص و کثافت

② لازم ال sample
اسحبها قبل
أعطيه أي antibiotic
أو عينة نقل دم

Other screening methods

- Microfluorometric assay: The direct measurement of phenylalanine in dried blood filter disks:

■ This method is quantitative, more adaptable to automation, and is not affected by presence of antibiotics.

لائي ما استفيدت
microorganism.

- The procedure is based on the fluorescence of complex formed of phenylalanine-ninhydrin-copper in the presence of dipeptide (i.e. L-leucyl-L-alanine).
- The test requires pretreatment of the filter paper specimen with trichloacetic acid (TCA)
- The extract is then reacted with microtiter with a mixture of ninhydrin, succinate, and leucylalanine in the presence of copper tartarate.
- The fluorescence of the complex is measured using excitation/ emission wavelengths of 360 nm and 530 nm, respectively
- For quantitative methods, HPLC or tandem mass spectrometry (MS/MS) are used

Other screening methods

- Urine testing for phenylpyruvate can be used for diagnosis in questionable cases and for monitoring of dietary therapy
- The test which may be performed by tube or reagent strip test involves the reaction of ferric chloride with phenylpyruvic acid in urine to produce a green color ① ②
- Prenatal diagnosis and detection of carrier status in families with PKU is now available using DNA analysis
- Analysis using cloned human PAH cDNA, has revealed the presence or numerous restriction fragment length polymorphism in the PAH gene

استوف اذا الشخص
حاضر للمرض أو حامل
↓
يؤخذوا عينة
من السائل بـ ١٠ حولين
الطفل ويحلوا
DNA analysis

Tyrosinemia and related disorders

↳ high level of tyrosine → نتيجة defect في الenzymes
metabolic pathway.

- A range of familial metabolic disorders of tyrosine catabolism is characterized by excretion of tyrosine and tyrosine catabolites in urine
- The defect in inherited tyrosine abnormalities is either a deficiency in tyrosine aminotransferase, resulting in tyrosinemia II, a deficiency of 4-hydroxyl-phenylpyruvic acid oxidase, leading to tyrosinoma type III or, more commonly a deficiency of fumarylacetoacetate (FAC) hydrolase, resulting in tyrosinemia I
- The absence of these enzymes results in abnormally high levels of tyrosine and, in some cases, increases in PHPPA and methionine
- The elevated tyrosine leads to liver damage, which may be fatal in infancy or to cirrhosis and liver cancer later in life
- The incidence of tyrosinemia I is approximately 1 of 100,000 births.

Diagnosis

- The disease is diagnosed by elevated tyrosine level using MS/MS coupled with a confirmatory test for an elevated level of the abnormal metabolite succinylacetone

→ mass spectroscopy

← بنتج عن ال tyrosin

Alkaptonuria

→ homogenisate oxidase enzyme defect.

↓
homogenetic acid will accumulation.

- This disorder is one of the original inborn errors of metabolism that showed a pattern of familial inheritance
- The biochemical defect in alkaptonuria is a lack of homogenisate oxidase in the tyrosine catabolic pathway
- This disorder occurs in about 1 of 250,000 births.
- of HGA in the urine, which oxidizes to produce a dark polymer.
- Alkaptonuria patients have no immediate problems but late in the disease, the high level of HGA, gradually accumulates in the connective tissue, causing generalized pigmentation of these tissues (ochronosis) and an arthritis-like degeneration

↓
clear color in urine

①

①
عالمه لون بار urine ← پس اذا تركته فترة

2- يميل لون ال urine (أغث) (darker)

↓

Homogenetic acid. → polymerization ال نتيجة ← brown Black color

Maple syrup urine disease (MSUD)

↓
branched chain amino acid → metabolism
↓
ketacid
لأفيسيلها
تتأخر في التمثيل
الناجيه عنه

- also called branched-chain ketoaciduria
- An autosomal recessive metabolic disorder affecting branched-chain amino acids. It is one type of organic acidemia
- MSUD is caused by a deficiency of the branched chain a-keto acid dehydrogenase complex (BCKDH), leading to a buildup of the branched-chain amino acids (leucine, isoleucine and valine) and their toxic by-products (ketoacids) in the blood, urine and cerebrospinal fluid (CSF).
- The disease is characterized in an infant by the characteristic maple syrup or burnt sugar odor of the urine, breath, and skin.

لأفيسيلها

Cause acidosis.

Maple syrup urine disease (MSUD)

اول اسبوع الطفل تكون لُبَيْرٍ بعد هيك بيلت يظهر عليه اعراض

- Typically infants with this inherited abnormality appear normal at birth but, by age 4-7 days, develop lethargy^①, vomiting^②, and signs of failure to thrive^③
- Central nervous system (CNS) symptoms follow including muscle rigidity^①, stupor^②, and respiratory irregularities^③
- If left untreated, the disease causes severe mental retardation^①, convulsions^②, acidosis^③, and hypoglycemia^④
- In the classic form of the disease, death usually occurs during the 1st year.
- In less severe variants, the activity of the decarboxylase is approximately 25% of normal. Although this still results in a persistent elevation of the branched amino acids, the levels frequently can be controlled by limiting dietary protein intake

Screening method

① أول فحص

- A modified Guthrie test is commonly used for this neonatal screening.
- The metabolic inhibitor to *B. subtilis*, included in the growth media, is 4-azaleucine
- In a positive test for MSUD, an elevated level of leucine from a filter paper disk impregnated with the infant's blood will overcome the inhibitor and bacterial growth occurs.

لـ ٤ ازل
thiongalanine.

إذا صار في growth agar صناعي
leucine
فرفع عنه.

Screening method

- Microfluorometric assay for branched-chain amino acids, using leucine dehydrogenase, can be used for mass screening
 - The filter paper specimen is treated with a solvent mixture of methanol and acetone to denature the hemoglobin.
①②
 - Leucine dehydrogenase is added to an aliquot of this sample extract. The fluorescence of the NADH produced in the subsequent reaction is measured at 450 nm, using an excitation wavelength of 360 nm.
 - A confirmed diagnosis is based on finding increased plasma and urinary levels of the three branched-chain amino acids and their ketoacids, with leucine being in highest concentration
له الأعلى
 - A leucine level above 4 mg/dL is indicative of MSUD. The presence of alloisoleucine, an unusual metabolite of isoleucine is characteristic.
- Measurement of leucine and its metabolites is also possible using tandem mass spectrometry. MSUD can be diagnosed prenatally by measuring the decarboxylase enzyme concentration in cells cultured from amniotic fluid.
ف يكون حولين الغفل

Isovaleric acidemia

↓
problem in metabolism of leucine.

- Isovaleric acidemia results from a deficiency of the enzyme isovaleryl-CoA dehydrogenase in the degradative pathway of leucine

↓
Convert isovaleryl CoA to β -Methylcrotonyl CoA

isovaleryl
CoA
راحت
بیش
تترام

- The resultant elevation of the glycine conjugate of isovaleric acid, isovalerylglycine, produces a characteristic "sweaty feet" odor.

- The abnormal organic acid levels can be identified by chromatography or MS/MS.

①

↓
mass spectroscopy.

Homocystinuria

methionine may be converted in the body to homocysteine then to cysteine.

- Homocysteine is an intermediate amino acid in the synthesis of cysteine from methionine.
- The usual cause of the hereditary disease, homocystinuria, is an impaired activity of the enzyme cystathionine β -synthase, which results in elevated plasma and urine levels of the precursors homocysteine and methionine.
- Newborns show no abnormalities, but physical defects develop gradually with age.
- Associated clinical findings in the late childhood include thrombosis resulting from toxicity of homocysteine to the vascular endothelium, osteoporosis, dislocated lenses in the eye resulting from the lack of cysteine synthesis essential for collagen formation and mental retardation.

حرف و رائی

① جلفان

②

③

Screening test

هناك 2 enzyme! cofactor فحين يكون سبب المشكلة
هناك 2 cofactor وحين يكون الانزيم نفسه

cofactor: هنا المشكلة عند بـ response ← هنا انما اعطيه اياه و هذا المرض

- The enzyme cystathionine β -synthase requires vitamin B6 (pyridoxine) as its cofactor.
- There are two forms of the disease:
 - A vitamin B6-responsive form, in which treatment consists of therapeutic doses of vitamin B6 → defect in cofactor
 - A vitamin B6-unresponsive form, in which the treatment is a diet low in methionine and high in cysteine → defect in enzyme.
- The incidence of homocystinuria is approximately 1 of 200,000 births
- Screened in infants by Guthrie test using L-methionine sulfoximine as the metabolic inhibitor.
 - Increased plasma methionine levels from affected infants will result in bacterial growth
 - A level of methionine greater than 2 mg/dL using an HPLC procedure confirms positive results on the screening test

Neonatal screening

- Alternately screening programs can use MS/MS to test for methionine

→ mass spectrometry.

تستات من هان لما نموت
شكرو الوضع 🤔

- Elevations in urinary homocystine can be detected by the cyanide-nitroprusside spot test

→ reducing agent

هو ال
مقطع تفاعل
مع ال cystine

- Cystine and homocystine reduced by sodium cyanide to their free thiol form, cysteine and homocysteine, which can then react with sodium nitroprusside to produce a red-purple color

→ homo ل cystine يا عندي

2 cysteine
مربوطين ب
disulfide bond

- Because cysteine also produces a positive result, the presence of homocysteine must be confirmed with a silver nitroprusside test. Silver nitrate reduces homocysteine but not cysteine, allowing only homocysteine to react with nitroprusside and produce a reddish color.

2 homocystine
مربوطين ب
disulfide bond

- Cystine remains in the oxidized form, which does not react with sodium nitroprusside.

→ when we add sodium cyanide they will be reduce to cysteine and homocysteine

Neonatal screening

- Elevations of homocysteine are of interest in the cardiovascular risk. It was found that approximately 50% of individual with untreated homocystinuria with significantly elevated levels of plasma homocysteine (200-300 $\mu\text{mol/L}$) had experienced a thromboembolic event before the age of 30.
- Mild homocysteine elevation ($>15 \mu\text{mol/L}$) occurs in 20-30% of patients with atherosclerotic disease
- In addition to cystathionine- synthase deficiency hyperhomocystinemia can be caused low folate concentration, vitamin B1 deficiency, decline in renal function, and genetic alteration of the enzyme, methylenetetrahydrofolate reductase (MTHFR), which converts homocysteine back to methionine.
- Although there is evidence of endothelial dysfunction in patients with elevated homocysteine levels, there is a disagreement with whether mild hyperhomocystinemia is a causative factor in the development of atherosclerotic disease or a consequence of the disease process

المسألة هي
عن ال
enzyme.

defect on arginine succinic acid
 → lyase → cause arginine succinic aciduria
 or
 → synthetase → cause citrullinemia

Argininosuccinic aciduria and Citrullinemia

- Results from inherited enzyme deficiencies in the urea cycle, arginine succinic aciduria results from the deficiency in argininosuccinic acid (ASA) lyase and a decrease in activity of ASA synthetase causes citrullinemia

- Symptoms include vomiting and high ammonia levels, and mental retardation is associated with some of the conditions

↓
 لا نه
 urea cycle
 فيما مشكلة ← تبصير ammonia
 تتراكم جوا الجسم داخل جزء تبصير هو الدماغ

- MS/MS technology has allowed measurement of the affected metabolites.

- Citrulline is the diagnostic marker for both citrullinemia and argininosuccinic aciduria

① ← في هاي الحالة ال citrulline فابكون عرتفع كثير.
 ② high level of ornithine + arginine.
 بس في هاي الحالة ال citrulline فابكون كثير مرتفع

- Citrulline is dramatically elevated in citrullinemia, while in argininosuccinic aciduria, the increase in citrulline is milder and increases in ornithine and arginine are seen in older infants

Cystinuria → transporter defect.

* all nutrients in our body have transporter in the kidney.

↓ Cystin / lysine / arginine / ornithine → all of them have transporter → any defect of these transporter will increase secretion of them

relatively insoluble

↓ increase concentration
 ➤ Caused by a defect in the amino acid transport system rather than a metabolic enzyme deficiency

نتيجة اذ
 في transporter
 ↓

➤ Normally amino acids are freely filtered by the glomerulus and then actively reabsorbed in the proximal renal tubules

↓
 precipitate in kidney tubules

➤ In cystinuria, there is 20-30-fold increase in the urinary excretion of cysteine as a result of genetic defect in the renal resorptive mechanism

↓
 Cause urinary calculi
 ← حصى بالكلى

↓
 Treatment:

1. high fluid intake
2. alkalinization of urine
3. penicillamine drug use.

↓
 (regular doses of penicillamine)

➤ The transport mechanism is not specific for cysteine. Excretion of the other amino acids, lysine, arginine, and ornithine, is also significantly elevated as a result of deficient resorption.

بالنسبة لـ arginine + ornithine and lysine ← كلهم عبارة عن basic amino acid
 ← بالأي حاي أي مشكلة بار solubility بيقوم ← بطلعوا من الجسم بدون أي مشاكل .

Cystinuria

- Of the four, cysteine is relatively insoluble, when it reaches high levels in the urine, it tends to precipitate in the kidney tubules and form urinary calculi.
- The formation of cysteine calculi can be minimized by a ^① high fluid intake and alkalinizing the urine, which makes cysteine relatively more soluble^②
- If this does not succeed, ^③ treatment with regular doses of penicillamine can be initiated
- cystinuria can be diagnosed by testing the urine for cysteine using cyanide-nitroprusside, which produces a red-purple color on reaction with sulfhydryl groups.
- False-positive results as a result of homocystine must be ruled out.

Proteins

- total protein test. (7-8 g/dL)
- albumin test. (4 g/dL).

■ General characteristics

- Proteins are an essential class of compounds comprising 50-70% of the cell's dry weight.
- Proteins are found in all cells of the body as well as in all fluids, secretions and excretions

■ Molecular Size

- Biological active proteins are macromolecules that range in molecular weight from approximately 6000 for insulin to several million for some structural proteins.

■ Structure

- All proteins comprise covalently linked polymers of amino acids.
- The carboxy group of one amino acid combines with the amino group of another amino acid by peptide bond
- In human serum, proteins average about 100-150 amino acids in the polypeptide chain
- The conformation of a protein is determined by interaction between a polypeptide and its aqueous environment

↓
3 dimension structure

Protein structure and characteristics

- The primary structure is crucial for the function and molecular characteristics of the protein.
- Denaturation can be caused by ^①heat, ^②hydrolysis by strong acid or alkali, ^③enzymatic action, exposure to urea or other substances, or exposure to ultraviolet light ^④
^⑤Non polar solvent.
- **The nitrogen content** of serum protein varies somewhat, the average is approximately 16%. This characteristic is used in one method of total protein measurement
عرفتها نسبة نيتروجين في بروتين
- Each protein has its own **isoelectric point (pI)** which is the pH at which a protein has no net charge which help in isolation of proteins by electrophoresis
[يمكن يحدد في عيني HLB + defect = غير يحدد من خلال ال isoelectric point.]
- Proteins are antigenic
may cause allergic.
- Protein form colloidal solution or micelles because they are charged produces an envelope of water around it which make possible to precipitate using different concentrations of salt and nonpolar solvents.
[solubility في ال water أعلى = سبب ال HLB من حولين ال protein و جعل ترسيب ال.]

Proteins

■ Classification of proteins :

→ only amino acid.

- Simple proteins: contain peptide chains that on hydrolysis yield only amino acids (can be globular or fibrous)
- Conjugated proteins: comprise a protein (apoprotein) and a nonprotein moiety (prosthetic group)
 - The prosthetic group may be lipid, carbohydrate, porphyrins or metal

■ Function of protein

- ① ■ Plasma proteins and tissue proteins share the same amino acid pool and so important in tissue nutrition. → مصدر الطاقة.
- ② ■ Distribution of water among the compartments of the body by osmotic force of plasma proteins
- ③ ■ They act as buffers within the plasma and interstitial tissue.
↳ Hemoglobine.

General function of proteins

- ④ ➤ Many plasma proteins functions as specific transporters of metabolic substances as thyroxine-binding globulin and albumin
- ⑤ ➤ لتمييز الخلايا Several proteins are glycoproteins which function to distinguish which cells are native and which are foreign to the body
- ⑥ ➤ Many cellular proteins act as receptors for hormones.
- ⑦ ➤ Certain hormones (e.g. growth hormone and adrenocorticotrophic hormone (ACTH)) are themselves proteins
- ⑧ ➤ Proteins also serve a structural role as collagen. elastine.
- ⑨ ➤ Some proteins (enzymes)

Plasma proteins

Prealbumin (transthyretin)

لا تقدر ابيض من انواع مختلفة من البروتينات عن طريق ال electrophoresis (الجل) ← علة فصل تبعها على الاختلاف بار charge. + size

تبيين عندي different bands

← كمية قليلة جداً بالدم.

لـ أحل band هو Prealbumin ← علة بال electrophoresis العادي ما آشوفها

- It is rarely observed as a distinct band on routine cellulose acetate electrophoresis patterns of serum, although it can be exhibited by high-resolution electrophoresis (HRE) or immunoelectrophoresis

Function:

- Prealbumin is used in the body to transport thyroxine and triiodothyronine, in addition to the transportation of retinol (vitamin A)

Negative acute phase protein inflammation
← يعني تنخفض لما يكون عندي inflammation.

- Prealbumin is **decreased** in hepatic damage, acute phase inflammatory response, and tissue necrosis

لـ لأنه ال half-life تبعه كيرة قصيرة.

- A low prealbumin level is a sensitive marker of poor protein nutritional status which results in a decrease in the level of the proteins originating in the liver, including prealbumin (short half life, 2 days)

- Prealbumin is **increased** in patients receiving steroids, in alcoholism, and in chronic renal failure

Albumin → synthesis in Liver.

- Albumin is the protein present in highest concentration in the serum that is synthesized in the liver. 60% of plasma protein albumin.
- Because of its high concentration in blood, albumin is responsible for nearly 80% of osmotic pressure
- Albumin binds bilirubin, salicylic acid, fatty acids, calcium, magnesium ions, cortisol and some drugs. This characteristic is also exhibited with certain dyes, providing a method for the quantitation of albumin

unconjugated

← يمكن استخدامه
للاختبار
1. albumin
2. quantitation

↓
1. warfarin
2. sulfonyleureas.
3. salicylic acid.
4. digoxin.

Albumin

Decreased conc. of serum albumin may be caused by the following:

① ➤ **An inadequate source** of amino acids (malnutrition and muscle-wasting disease)

② ➤ **Liver disease**, resulting in the ability of hepatocytes to synthesize albumin.
 The increase in globulins that occurs in the early cirrhosis will balance the loss in albumin to give a total protein concentration within acceptable limits. The decline in serum albumin is insignificant in viral hepatitis.

③ ➤ **Gastrointestinal loss** as interstitial fluid leaks out in inflammation and disease of the intestinal mucosa

④ ➤ Loss in the urine in **renal disease**. → damaged in glomerulus.

↩ درائي ➤ **Analbuminemia**: the absence of albumin because of genetic origin resulting from an autosomal recessive trait

↩ 2 gens لاني عشان يظهر المرض عدي ➤ **Bisalbuminemia**: the presence of albumin that has unusual molecular characteristics demonstrated by the presence of two albumin bands instead of the single band usually seen by electrophoresis

acute cases
 حاد 2
 يهربي انخفاض
 chronic cases
 مبراني اد

انخفاض بسيط

defect
 albumin.

normal
 albumin

و حدة لا و حدة لا

Albumin

فان في عذبة زيادة
production of
albumin.

- Increased serum levels are seen in **Dehydration** (vomiting, diarrhea).
- Administering fluids to treat the dehydration will decrease serum albumin levels back to normal.

- The **earliest method** for its determination involved the salting out of the **globulins with sodium sulfate** leaving the albumin in solution

حجما كبير دينا ← lipophilicity
أكثر ← يبين على العينة
sodium sulfate
يحل عذبة
salting out of
globulin ↓
يحل عذبة ال
albumin
بار solution
يضيف عليها
biuret reagent
↓
يوز كمية ال
albumin
الموجودة.

- The albumin was then determined by the **biuret color development**. The method commonly used today involves a **dye binding and shift in color** when a dye is bound by albumin
- When more information about proteins is needed, an electrophoretic pattern is obtained, and albumin is calculated as percentage of the total protein (usually, approximately 60%)
← تكون ال band تبقته كثير كبيرة.
- At birth, the reference value for serum albumin averages 39g/L. the concentration falls to 28.4 g/L at about 9 months and then begins to increase slowly until adult values of 35-55 g/L are reached.

← 100 ml → 100 g / 100 ml
← نسبة البروتين 35-55 g/L وحسب 39 g/L

في عينة ال electrophoresis حكيما كل band تباع عن ال prealbumine و يكون خفيفة
وال band بل بعدا يكون ال albumin و يكون - large
ال band ال α_1 -Antitrypsin و

α 1-Antitrypsin \rightarrow 90% of α 1 proteins.

elastic enzyme. Protein lung

- Its main function to neutralize trypsin-like enzymes (as elastase) α_1 -Antitrypsin is a major component (90%) of the fraction of serum protein that migrates electrophoretically immediately following albumin.
- A deficiency of α_1 -Antitrypsin is associated with severe, degenerative emphysematous pulmonary disease due to proteolytic activity of proteases from leukocytes in the lung during periods of inflammation

← defect في α_1 -antitrypsin ← تبني في الـ liver بس ما بيسره releases الـ lung
 ← ليس يتراكم جوا الـ liver
- **Juvenile hepatic cirrhosis** is also correlative disease in α_1 -Antitrypsin deficiency. The protein is synthesized but not released from the hepatocyte.

من الكبد
- Increased levels of α_1 -Antitrypsin are seen in inflammatory reactions, pregnancy and contraceptive use



3

إذا كانت على activation enzyme ← بعد emphazema له وقت الأشخاص.

α_1 -Antitrypsin

لدراسة ال globulin في ال electrophoresis ← إذا كانت موجودة فها وقت الأشخاص
فاعة α_1 -antitrypsin band

- The discovery of abnormal α_1 -Antitrypsin levels is most often made by the lack of an α_1 -globulin band on protein electrophoresis.
- The discovery is followed with one of the quantitative methods. A widely used method is radial immunodiffusion ←
- Immunonephelometric assays by automated instrumentation are also available. Phenotyping can be accomplished by immunofixation → antigen-antibodies reaction.

ل رجة على العين ← antibodies specific ل α_1 -antitrypsin ← تترفع ال antigen

• precipitation. الوجة على و بـ ال

α 1-Fetoprotein

يتم تصنيعه في المراحل
الأولى من بداية الحمل
عند الجنين .

يُنتج بكميات عالية

يسير بكميات متدنية

يسير بكميات متدنية

- α 1-fetoprotein (AFP) is synthesized initially by the fetal yolk sac and then by the parenchymal cells of the liver
- It peaks in the fetus at about 13 weeks of gestation (3 mg/ml) and recedes at 34 weeks gestation. At birth, it recedes rapidly to adult concentration, which are normally very low
- The methods commonly used for AFP determinations are radioimmunoassay and enzyme labeled immunoassay
- function:
 - It has been proposed that the protein protects the fetus from immunolytic attack by its mother, modulates cell growth transport compounds such as steroids and is required for the functional development of the female reproductive system
- AFP is detectable in the maternal blood up to month 7 or 8 of pregnancy (transmitted across the placenta). AFP in maternal serum is a screening test

①

②

③

مساعدة على نمو الخلايا .

بعد الفحص مفاد الأم ← يسير بكميات منخفضة ← الحالة مرضية وراثية عند الطفل

α 1-Fetoprotein

■ Elevated AFP level include:

- ① القحة بالي ليفر الفقر ■ neural tube defects, atresia of the gastrointestinal tract and fetal distress in general. Its use in determining neural tube defects before term is an important reason for its assay.
- It is also increased in ataxia-telangiectasia, tyrosinosis, and hemolytic disease of the newborn
- maternal serum AFP is also increased in the presence of twins. حادي twins

■ Low levels of maternal AFP indicate an increased risk for Down's syndrome and trisomy 18. ②

- are also affected by maternal weight, which reflects blood volume (inverse relationship), race (10% higher in African Americans), and diabetes (lowered value) تباثر بوزن الحامل. $\rightarrow \uparrow \text{weight} \rightarrow \uparrow \text{blood volume} \rightarrow \downarrow \alpha_1 \text{feto-protein}$
- Serum levels of AFP can also be used as a tumor marker (high in hepatocellular carcinoma (80%) and certain gonadal tumors in adults) face. ①

①
حسب بكمي ارق شخاض
لكون مرتفع

Ceruloplasmin

have enzymetic activity. ← نجزن الـ copper

90% of copper in the body. ← يوجد على هيئة

الحديد (حسينا المخلوطة)
في الشاير
في قية.

↓
defecancy in
copper

↓
from defecancy
anemia

↓
Treatment by
give copper.

- Ceruloplasmin is copper-containing α 2-glycoprotein that has enzymatic activities (ie. Copper oxidase, histaminase and ferrous oxidase)
- It is synthesized in the liver. 90% or more of total serum copper is found in ceruloplasmin
- The early analytic method of ceruloplasmin determination was based on its copper oxidative capacity ^① → convert apoferritin to ferritin.
- Most assays today use immunochemical methods, including radial immunodiffusion and nephelometry ^② (antigen-antibodies reaction).
- Low concentrations of ceruloplasmin at birth gradually increase to adult levels and slowly continue to rise with age. Adult females have higher concentrations than males and pregnancy, inflammatory processes, malignancies, oral estrogen and contraceptives cause an increased serum concentration.

Ceruloplasmin

ازا كان في low level فها ال Ceruloplasmin
مصنوعا copper عشا لاتي مكان
تخزن فيه ← بصر يفلح بال urine
← تترسب في أماكن بالجسم
(skin, brain, liver, cornea).

low concentration:

- Certain diseases or disorders are associated with low serum concentrations. In Wilson's disease, an autosomal recessive inherited disease, the levels are typically low (0.1 g/L) ←
- Total serum copper is decreased, but the direct reacting fraction is elevated and the urinary excretion of copper is increased
- The copper is deposited in the skin, liver and brain, resulting in hepatic cirrhosis and neurological damage. Copper also deposits in the cornea, producing the characteristic Kayser-Fleischer rings ←
فان اختلفت من ال GI
فان توفد ال pfeifer عشان اتمعه
- Low ceruloplasmin is also seen in malnutrition, malabsorption, severe liver disease, nephrotic syndrome, and Menke's syndrome (kinky hair disease), in which a decreased absorption of copper results in a decrease in ceruloplasmin.
← ما عند synthesis ال
بصر له
فقد الدم

Transferrin (Siderophillin)

→ β -globulin. ← علاقة عكسية

مع iron deficiency

↓
* low level of iron
→ ↑ Transferrin

* high level of iron.
→ ↓ Transferrin.

بجفاف - تركيزه
باعتلاف الجينات

- Transferrin, a glycoprotein, is synthesized primarily by the liver.
 - Transferrin is the major component of the β -globulin fraction and appears as distinct band on high-resolution serum protein electrophoresis
 - Genetic variation of transferrin has been demonstrated by electrophoresis on polyacrylamide gel
 - Precise and accurate analytic methods used for the quantitation of transferrin include immunodiffusion and immunonephelometry
- Function:
- The major fractions of transferrin are the transport of iron and the prevention of loss of iron through the kidney and deposition in the tissue during temporary increases in absorbed iron or free iron. Transferrin transports iron to its storage sites (ferritin) and to bone marrow that synthesize hemoglobin
- ① and hemosedrin (in Liver). ②

Transferrin (Siderophillin)

→ Negative acute protein in inflammatory conditions
يعني ينخفض في حالات الـ inflammation

- The most common form of anemia is iron deficiency anemia where transferrin in serum is normal or increased.
- A decreased transferrin level reflects an overall decrease in the synthesis of protein (as seen in liver disease or malnutrition and protein-losing disorders such as nephrotic syndrome).
(1) (2) (3)
- Transferrin, a negative acute phase protein, is also decreased in inflammation. A deficiency of plasma transferrin may result in the accumulation of iron in apoferritin.
(4)
- Patients with hereditary transferrin deficiencies have been shown to have significant hypochromic anemia. An increased of iron bound to transferrin is found in hereditary disorder of iron metabolism, hemochromatosis, in which excess iron is deposited in the tissue, especially the liver and the pancreas. This disorder is associated with bronze skin, cirrhosis, diabetes mellitus, and low plasma transferrin levels.
(1) (2) (3) (4)

→ hemochromatosis.

transferrin ← بعد الحديد
يعني saturation
iron
في
تتراكم في
tissue

Lipoprotein

— proteins bind with triglyceride + cholesterol

- They are complexes of proteins and lipids whose function is to transport cholesterol, triglycerides, and phospholipids in the blood
- Lipoproteins are subclassified according to the apoprotein and specific lipid content
 - On high-resolution serum protein electrophoresis, high-density lipoproteins (HDL) migrate between the albumin and the α_1 -globulin zone
 - Very-low-density lipoprotein (VLDL) migrate at the beginning of the β -globulin fraction (pre- β)
 - The low density lipoproteins (LDL) appear as a separate band in the β -globulin region

Fibrinogen

از ابدی اعلیٰ assessment

بسته ال thrombin

نمیفته ال plasma

بسیار بکول ال fibrinogen

Clotting (تعیین الوقت) در سیر علی

بال اخذ عشان

یحل

clotting



هو ال بصری

quality

and quantity

of fibrinogen.

- One of the largest proteins in blood plasma. It is synthesized in the liver and it is classified as a glycoprotein because it has considerable carbohydrate content.

→ converted to fibrin → close the injury

- The function of fibrinogen is to form fibrin clot when activated by thrombin, therefore, fibrinogen is virtually all removed in the clotting process and is not seen in serum.

- Fibrinogen customarily has been determined as clot-table protein.

در حالت بحیه الجبر ← معناتو قدر اسر الجرح بشل أخض

- Fibrinogen concentration is proportional to the time required to form a clot after the addition of thrombin to citrated plasma

Fibrinogen

→ positive acute phase protein inflammation.

- Degradation products of fibrinogen and fibrin are determined by immunoassay methods such as immunodiffusion, nephelometry and radioimmunoassay
③ ① ②
- Fibrinogen is one of the acute phase reactants (significantly increased in plasma during acute phase of inflammatory process)
①
- Fibrinogen levels also rise with pregnancy and the use of birth control pills
② ②
- Decreased values generally reflect extensive coagulation, during which the fibrinogen is consumed

Troponin (found in skeletal muscles, cardiac muscles)

↳ good indication for patient have myocardial infarction.

- Troponin is complex of three proteins that bind to the thin filaments of striated muscle (cardiac and skeletal) but are not present in smooth muscles
(1) (2) → specific for myocardial tissue
- The complex consists of troponin T, troponin I, and troponin C
- Cardiac troponin T in serum begins to rise within 3-4 hours following the onset of myocardial damage, peak in 10-24 hours, and remain elevated for 10-14 days following AMI
- Cardiac troponin I is also highly specific for myocardial tissue
- Because cTnI, like cTnT does not normally circulate in the blood and it is 13 times more abundant in the myocardium than CK-MB on a weight basis, cTnI is a very sensitive indicator of even a minor amount of cardiac necrosis

Troponin I حساس من ال CK-MB في حالات ال myocardial Infarction.

بالنسبة لقلب قديم
بالنسبة لآدمي
very low concentration
وصف ال myocardial
أكثر 13 مرة

Total protein abnormalities

Hypoproteinemia

decrease intake or
increase excretion.

- Occurs in any condition where a negative nitrogen balance exists
- Plasma proteins is excessive loss as in renal disease (ie. Nephrotic syndrome) leakage into the GIT in inflammation of the digestive system and in loss of blood in open wounds, internal bleeding, or extensive burns
- Decreased intake either because of deficiency of protein in diet (malnutrition) or through intestinal malabsorption due to structural damage
- A decrease in serum proteins as a result of decreased synthesis is also seen in liver disease
- Hypoproteinemia may result from accelerated catabolism of proteins, such as in burns, trauma, or other injuries

Hyperproteinemia

- An elevation of all protein fractions is observed is dehydration which may result from a variety of conditions, including vomiting, diarrhea, excessive sweating, diabetic acidosis, and hypoaldosteronism

- Hyperproteinemia may be result of excessive production, primarily of the γ-globulins

↓
Paraproteins

↓
albumin و normal globulin.

- The appearance of monoclonal protein or paraprotein in the serum and often in the urine as well

← جيع القرح ← Serum + urine. لبرتنجوا بال
infection bacterial. و viral سوا

Hyperproteinemia

نوع من أنواع Cancer

- The most common disorder is multiple myeloma, in which the neoplastic plasma cells proliferate in the bone marrow
- The paraproteins in this case is usually IgG and IgA. IgD and IgE paraproteins rarely occur. Paraproteins in multiple myeloma may reach a serum concentration of several grams per deciliter
immunoglobulin G + A
- Not all paraproteins are associated with multiple myeloma. IgM paraprotein is often found in patients with Waldenstrom's macroglobulinemia, a more benign condition.
- Many disorders including chronic inflammatory states, collagen vascular disorders, and other neoplasms, may be associated with paraproteins
① ②
- Polyclonal increases in immunoglobulins are seen in the serum and urine in many chronic diseases

TABLE 8-5. PROTEIN LEVELS IN SELECTED DISEASE STATES

TOTAL PROTEIN	ALBUMIN	GLOBULIN	DISEASE
<u>N, ↓</u>	<u>↓</u>	↑	<u>Hepatic Damage</u> <ul style="list-style-type: none"> • <u>Cirrhosis</u> β-γ bridging • <u>Hepatitis</u> ↑ γ-globulins • <u>Obstructive jaundice</u> ↑ α_2-, β-globulins <u>Burns, Trauma</u> <u>Infections</u> <ul style="list-style-type: none"> • <u>Acute</u> ↑ α_1-, α_2-globulins → γ-globulin ↓ • <u>Chronic</u> ↑ α_1-, α_2-, γ-globulins ← γ-globulin
↓	<u>↓</u>	N	<u>Malabsorption</u> <u>Inadequate Diet</u> <u>Nephrotic Syndrome</u> ↑ α_2 -, β -globulins; ↓ γ -globulins
<u>↓</u>	N	<u>↓</u>	<u>Immunodeficiency Syndromes</u>
↓	↓	↓	<u>Salt Retention Syndrome</u> → ↑ water retention
<u>↑</u>	<u>↑</u>	<u>↑</u>	<u>Dehydration</u>
<u>↑</u>	N	<u>↑</u>	<u>Multiple Myeloma</u> <u>Monoclonal and Polyclonal Gammopathies</u>

↑ = increased; ↓ = decreased; N = normal levels.

CASE STUDY 8.2

Immediately following the birth of a baby girl, the attending physician requested a protein electrophoretic examination of the mother's serum. This was done on a sample that was obtained on the mother's admission to the hospital the previous day. An electrophoretic examination was also performed on the cord-blood specimen. Laboratory reports are shown in Case Study Table 8-2.1.

The appearance of the mother's electrophoretic pattern was within that expected for a healthy person. The electrophoretic pattern of the cord-blood serum resembled the one shown in Figure 8-13C.

Questions

1. What protein fraction(s) is/are abnormal in the mother's serum and the cord-blood serum?
2. An abnormality in this/these fraction(s) is/are most often associated with what disease?
3. What other test(s) may be done to confirm this abnormality?

CASE STUDY TABLE 8-2.1 ELECTROPHORESIS (VALUES g/dL)

	ADULT REFERENCE VALUES	MOTHER'S SERUM	CORD BLOOD
Albumin	3.5–5.0	4.2	3.3
α_1 -Globulins	0.1–0.4	0.3	0.0
α_2 -Globulins	0.3–0.8	1.2	0.4
β -Globulins	0.6–1.1	1.3	0.7
γ -Globulins	0.5–1.7	1.3	1.0

CASE STUDY 8-3

A 76-year-old woman was admitted to the hospital with gangrene of her right toe. She was disoriented and had difficulty finding the right words to express herself. On evaluation, it was revealed she lived alone and was responsible for her own cooking. A daughter who lived in the area said her mother was a poor eater, even with much encouragement. An ECG, performed on admission, showed possible ectopic rhythm with occasional premature supraventricular contractions. The cardiologist suspected a possible inferior myocardial infarction of undetermined age. Lab results are shown in Case Study Table 8-3.1.

Questions

1. In this patient, what is the clinical value of the troponin I measurements?
2. What is a possible explanation for the elevated myoglobin?
3. What condition is indicated by the low prealbumin value?

CASE STUDY TABLE 8-3.1 LABORATORY RESULTS

Day 1

CK-total	187 U/L	(40–325)
CK-MB Mass	6 µg/L	(<8)
Troponin I	16.3 µg/L	(0–2)
Prealbumin	15 mg/dL	(17–42)
Albumin	2.7 g/dL	(3.7–4.9)

Repeat (5 Hours Later)

CK-total	180 U/L
CK-MB mass	5.4 µg/L
Troponin I	17.5 µg/L

Day 2

CK-total	177 U/L	
CK-MB mass	4.5 µg/L	
Troponin I	13.7 µg/L	
Myoglobin	<500 µg/L	(<76)

CASE STUDY 8-5

A 45-year-old man was undergoing continuing evaluation of possible recurrence of a plasmacytoma that had originally presented with a compression fracture of a vertebra. He had been treated with local radiation and chemotherapy. His serum protein electrophoresis showed normal amounts of albumin, α_1 , α_2 , and β fractions. The γ fraction demonstrated a slight monoclonal band in the fast γ region (close to β). Protein electrophoresis of concentrated urine showed a single monoclonal band that migrated slightly less than the serum band. (Case 8-5 courtesy of Dr. R. McPherson, Chairman, Clinical Pathology,

Medical College of Virginia Hospitals, Virginia Commonwealth University Health System)

Questions

1. Does the presence of the monoclonal band in the serum indicate the recurrence of the patient's tumor?
2. What further information is obtained from a urine protein electrophoresis?
3. What other test is needed to confirm the type of urinary protein?

CASE STUDY 8-8

A 36-year-old woman complained of intermittent blurred vision and numbness and weakness in her left leg that had persisted for more than three weeks. On examination, vertical nystagmus (involuntary back-and-forth or circular movements of eyes) was noted on upward gaze. CSF was drawn and the specimen was clear and colorless with normal cell count. The CSF total protein level was 49 mg/dL with an IgG of 8.1 mg/dL. Electrophoresis of the patient's serum and CSF revealed the following pattern:



Questions

1. What is the significance of the protein bands indicated by the arrows?
2. What conditions would produce this type of protein electrophoresis pattern?
3. What other tests would be helpful in the investigation of this patient's diagnosis?
4. What laboratory test can be useful for monitoring the course of this patient's condition?