Plasma proteins

Defivition

They are used as a biomarker What is the meaning of biomarker ? لتشخيص الامراض مثل الانزيمات For example MI (myocardial

infraction) ? CHEST PAIN

- ECG , May be normal
- ceriaten kinase MP
- Troponin
- Both elevated > have MI

- Plasma contains >300 different proteins, their levels are affected by many pathological conditions.
- Mostly synthesized in the liver The only one not synthesised in the liver is : Gamma globulin's AB synthesised by cytotoxic killer lymphocytes.
- Some are produced in other sites
- A normal adult has 6-8g/dl of plasma proteins اكثر تواجدًا ال albumin
- The proteins of the plasma are a complex mixture that includes not only simple proteins but also conjugated proteins such as glycoproteins and various types of lipoproteins. Deficiency of plasma proteins: 1. Liver disease Lipids + proteins
- Functions
- Transport (Albumin, prealbumin, globulins)
- Maintain plasma oncotic pressure (Albumin)
- Defense (Immunoglobulins and complement)
- Clotting and fibrinolysis (Thrombin and plasmin)
- Buffering pH
- Catalytic functions (enzymes as LPL)
- Signal proteins

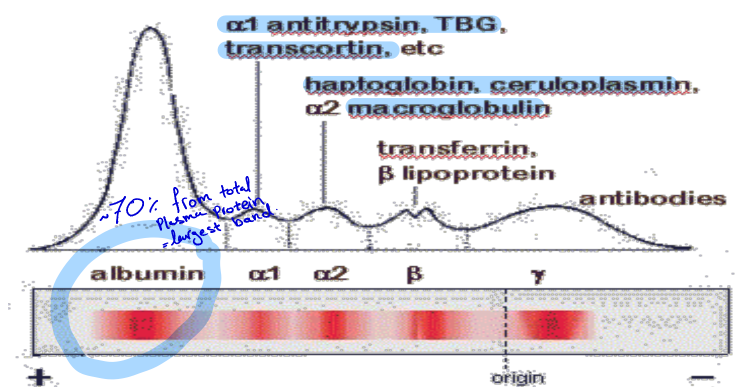
- 2. Malabsorption
- 3. Mal nutrition

Measurement of Plasma Proteins

A-Quantitative measurement of a specific protein by chemical or immunological reactions



B- Semiquantitative measurement by electrophoresis: proteins are separated by their electrical charge in electrophoresis (five separate bands of proteins are observed, these bands change in disease).



Types of Plasma Proteins

- Prealbumin Doesn't have a band in electrophoresis due to the low concentration
- Albumin

Tumor marker But is best preferred in the case of following up

- α 1-Globulins: as α 1-Antitrypsin, α -fetoprotein
- α 2-Globulins: as Ceruloplasmin, haptoglobin

Emphysema

- β Globulins: as CRP, transferrin, β 2-microglobulin
- γ Globulins

Prealbumin (Transthyretin)

- A transport protein for: thyroid hormones and retinol
- Migrates faster than albumin in electrophoresis
- Separated by immunoelectrophoresis
- Lower levels found in: liver disease, nephrotic syndrome, acute phase inflammatory response, malnutrition
 Short half-life (2 days)

Albumin

- Most abundant plasma protein (3.5-5.5 g/l) in normal adult
- Synthesized in the liver as preproalbumin and secreted as albumin
- Half-life in plasma: 20 days ~ 3 weeks
- Decreases rapidly in injury, infection and surgery
- 40% of albumin found in plasma, 60% in the extracellular space

Functions

- May cause edema in cases - Maintains oncotic pressure:
 - The osmotic pressure exerted by plasma proteins that pulls water into the circulatory system

Hypoalbunemia

• swelling of RBCs > edema

- Maintains fluid distribution in and outside cells and plasma volume (80% of plasma oncotic pressure is maintained by albumin)
- A non-specific carrier of hormones, calcium, free fatty acids, drugs, etc.
- It is by pinocytosis in the cells where it is hydrolyzed to amino acids
- Nutritive function
- Buffering function
- Useful in treatment of liver diseases, hemorrhage, shock and burns

Synthesis of albumin

- The liver produces albumin, it represents about 25% of total hepatic protein synthesis.
- Albumin is initially synthesized as a preproprotein Inactive form
- Its signal peptide is removed as it passes into rough endoplasmic reticulum, and a hexapeptide at the resulting amino terminal is subsequently cleaved off farther along the secretory pathway.
- Mature human albumin consists of one polypeptide chain of 585 amino acids and contains 17 disulfide bonds
- It has an ellipsoidal shape, which means that it does not increase the viscosity of the plasma as much as an elongated molecule such as fibrinogen does.
 In cases of liver diseases we give the patient albumin infusion to the blood that may cause viscosity but due to its ellipsoidal shape it prevent it to high viscosity.
- Has a relatively low molecular mass about 69 kDa

<u>Clinical significance of albumin</u> Blood brain barrier

- Ratio between the unconjugated bilirubin and albumin 2:1 so there will be high binding site.
- Albumin- free fatty acid complex can not cross the blood brain barrier, hence fatty acids can not be utilized by the brain.
- Loosely bound bilirubin to albumin can be easily replaced by drugs like aspirin ^{unconjugated bilirubin} مخاف بحالات الاطفال يزيح الدواء المعطى ال
- In new born if such drugs are given, the released bilirubin gets deposited in brain causing Kernicterus.

Advance jaundice with mixed signs

Protein bound calcium

- Calcium level is lowered in conditions of hypoalbuminemia
- Serum total calcium may be decreased
- Ionic calcium remains the same
- Tetany does not occur
- Calcium is lowered by 0.8 mg/dl for a fall of 1g/dl of albumin

50% ionised & 50% binded to the albumin (preserved)



Drug interactions

-Two drugs having same affinity for albumin when administered together, can compete for available binding sites with consequent displacement of other drug, resulting in clinically significant drug interactions. As phenytoin, dicoumarol interactions if both medication

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<u>Oedema</u> Hypoalbuminemia

Causes

- Decreased albumin synthesis (liver cirrhosis, malnutrition)
- Increased losses of albumin
 - Increased catabolism in infections
 - Excessive excretion by the kidneys (nephrotic syndrome).
 - Severe burns (plasma loss in the absence of skin barrier)
 - Excessive loss in bowel

Effects

- Edema due to low oncotic pressure
 - Albumin level drops in liver disease causing low oncotic pressure
 - Fluid moves into the interstitial spaces causing edema
- Reduced transport of drugs and other substances in plasma
- Reduced protein-bound calcium
 - Total plasma calcium level drops
 - -Ionized calcium level may remain normal

Hyperalbuminemia

- No clinical conditions are known that cause the liver to produce large amounts of albumin
- The only cause of hyperalbuminemia is <u>dehydration</u> and high protein diet Decrease the volume of the blood

<u>α1-antitrypsin</u>

- Called α 1- antiprotease, single polypeptide [394 amino acids (52 kDa)].
- Synthesized by the liver and macrophages
- An acute-phase protein that inhibits proteases (trypsin, elastase, and other proteases) by forming complexes with them.
- Infection leads to protease release from bacteria and leukocytes. (Marrophage)
- Normally α1-antitrypsin protects the lung tissues from the released active elastase from macrophages.
- In its deficiency, the active elastase destroys the lung tissue by proteolysis.

Types of α₁-Antitrypsin Even if little decrease of its activity

- Over 30 types are known (the most common is M type).
- Genetic deficiency of α 1-antitrypsin (synthesis of the defective α 1antitrypsin occurs in the liver but it cannot secrete the protein) \rightarrow its accumulation in hepatocytes and its deficiency in plasma

Clinical consequences of a1-antitrypsin deficienc

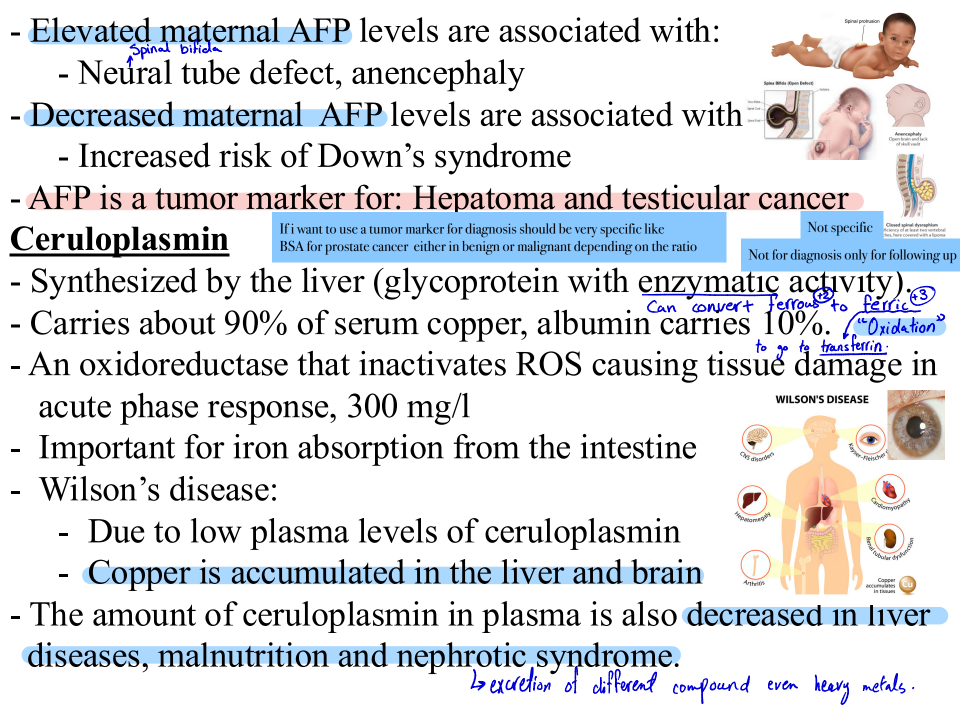
- Neonatal jaundice
- Childhood liver cirrhosis
- Pulmonary emphysema in young adults

Laboratory Diagnosis

- Lack of α 1-globulin band in protein electrophoresis
- Quantitative measurement of α 1-antitrypsin by: radial immunodiffusion and isoelectric focusing.

<u>α-Fetoprotein (AFP)</u>

- Synthesized in the developing embryo and fetus by the parenchymal cells of the liver.
- AFP levels decrease gradually during intra-uterine life and reach adult levels at birth (normal level is 1 μ g/100 ml).
- Function is unknown but it may protect fetus from immunologic attack by the mother.
- No known physiological function in adults



<u>Haptoglobin</u>

- Synthesized by the liver (glycoprotein).
- Binds to free hemoglobin to form complexes that are metabolized in the RES, when bound to hemoglobin, it is cleared from the plasma about 80 times faster than normally.
- Limits iron losses by preventing Hb loss from kidneys
 Plasma level decreases during hemolysis and increases in inflammation.

Binding capacity to the ironBinding capacity of ferritin inisnlow only carry 3 iron atomsthe iron is 4300 atoms of iron

It originally low molecular weight if it is not binded to binding to the Hb it will pass through the glomurilic pathway so it will cause renal failure

Transferrin May cause blood toxicity thats why its binding capacity is low when the iron is high

- A major iron-transport protein in plasma, 76 kDa
 - 30% saturated with iron
- Plasma level drops in:
 - Malnutrition, liver disease, inflammation, malignancy
- Iron deficiency results in increased hepatic synthesis

- A negative acute phase protein

Positive acute phase protein in cases of surgery, malignancy and inflammation is due to increase in the negative acute phase proteins مش فاهمة قصد الدكتور بالريكورد بس هيك قال حرفي

<u>β2– Microglobulin</u>

- A component of human leukocyte antigen (HLA)
- Present on the surface of lymphocytes and most nucleated cells
 Filtered by the renal glomeruli due to its small size but most
- (>99%) is reabsorbed
- Elevated serum levels are found in
 - Impaired kidney function
- May be a tumor marker for:

For kidney function we test:

- urea
- creatine
- Uric acid
- بعتمد اكثر اشي على الكيرياتين لانه اليوريا بتقدر ترتفع بحالات
 اكل كميات كبيرة من اللحمة بحالات المنسف
- Leukemia, lymphomas, multiple myeloma

<u>C-reactive protein (CRP)</u>

- An acute-phase protein synthesized by the liver (so named because it reacts with the polysaccharide of the capsule of pneumococci, important for phagocytosis
- High plasma levels are found in many inflammatory conditions such as rheumatoid arthritis
- -A marker for ischemic heart disease

<u>a2– Macroglobulin</u>

- Major component of $\alpha 2$ proteins
- Comprises 8–10% of the total plasma protein in humans.
- Tetrameric protein with molecular weight of 725 kDa. Supplesized by hepatocytes and macrophages
- Synthesized by hepatocytes and macrophages
- Inactivates all proteases and thus is an important in vivo anticoagulant.
- Carrier of many growth factors
- Normal serum level-130-300 mg/dl
- Concentration is markedly increased in nephrotic syndrome, since other proteins are lost through urine in this condition.

Hypergammaglobulinemia

- May result from stimulation of
 - B cells (Polyclonal hypergammaglobulinemia)
 - Monoclonal proliferation (Paraproteinemia)
- Polyclonal hypergammaglobulinemia: Not spesic
 - Stimulation of many clones of B cells produce a wide range of antibodies
 - γ -globulin band appears large in electrophoresis
 - Clinical conditions: acute and chronic infections, autoimmune diseases, chronic liver diseases

Spesific.

Monoclonal Hypergammaglobulinemia:

- Proliferation of a single B-cell clone produces a single type of Ig
- Appears as a separate dense band (paraprotein or M band) in electrophoresis
- Paraproteins are characteristic of malignant B-cell proliferation
- Clinical condition: multiple myeloma

Positive acute phase proteins (acute phase reactants)

- Plasma protein levels increase in:

Infection, inflammation, malignancy, trauma, surgery

- Synthesized due to body's response to injury as α_1 -antitypsin, haptoglobin, ceruloplasmin, fibrinogen and C-reactive protein
- Mediators cause these proteins to increase after injury as cytokines (IL-1, IL-6), tumor necrosis factors α and β, interferons, platelet activating factor
 Functions: 1. Bind to polysaccharides in bacterial walls
 2. Activate complement system 3. Stimulate phagocytosis.
 <u>Negative acute phase proteins</u>
- These proteins decrease in inflammation

Albumin, prealbumin and transferrin ψ

- Mediated by inflammatory response via cytokines and hormones
- Synthesis of these proteins decrease to save amino acids for positive acute phase proteins

Abnormal proteins

Why unusual ?

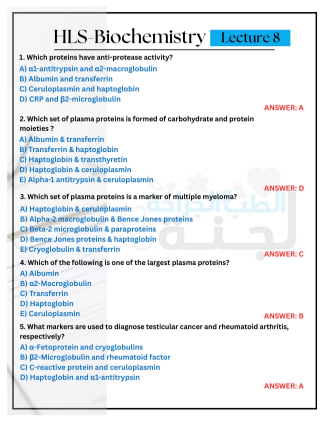
1- Bence Jone's proteins

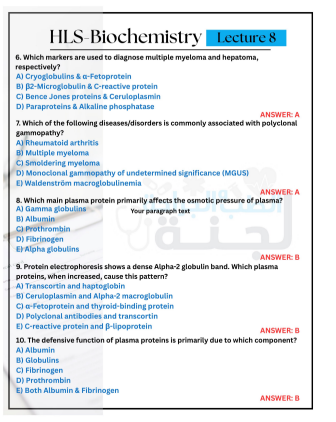
With heating will cause dissolution Other proteins with boiling & heating will cause COAGULATION

- Abnormal proteins (monoclonal light chains).
- Present in the urine of a patient suffering from multiple myeloma (50% of patients)
- Molecular weight 45 kDa
- Identified by heat coagulation test
- Best detected by zone electrophoresis and immunoelectrophoresis

2- <u>Cryoglobulins</u>

- These proteins coagulate when serum is cooled to very low temperature
- Commonly monoclonal IgG or IgM or both
- Increased in rheumatoid arthritis, multiple myeloma,
 lymphocytic leukemia, lymphosarcoma and systemic lupus
 erythematosus





HLS-Biochemistry Lecture 8

11. Which plasma proteins can be used as biomarkers for the diagnosis of multiple myeloma?

- A) 82-Microglobulin paraprotein and g-fetoprotein
- B) a2-Macroglobulin, Bence Jones proteins, and B2-microglobulin
- C) Bence Jones proteins cryoglobulins and paraprotein
- D) B2-Microglobulin, α -fetoprotein, and α 2-macroglobulin
- E) q1-Antitrypsin, paraprotein, and C-reactive protein

ANSWERC

12. Protein electrophoresis shows a dense Beta (B) globulin band. Which plasma proteins, when increased, cause this pattern?

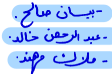
- A) Transcortin, haptoglobin, and ceruloplasmin
- B) Transcortin, paraprotein, and alpha transcortin
- C) g-Fetoprotein, transcortin, and thyroid-binding protein
- D) Thyroid-binding protein, polyclonal antibodies, and g-fetoprotein
- E) C-reactive protein, transferrin, and B-lipoprotein

ANSWER E

13. What is the major component of the Alpha-2 (g2) protein fraction in plasma? A) a1-Antitrypsin B) Ceruloplasmin C) Haptoglobin D) g2-Macroglobulin E) Transferrin







ANSWER: D