Liver disease in children

Dr. Ahmed Al-Sarkhy MD, MHSc, FAAP, FRCPC
Pediatric gastroenterology & hepatology consultant
College of medicine & KKUH
King Saud university

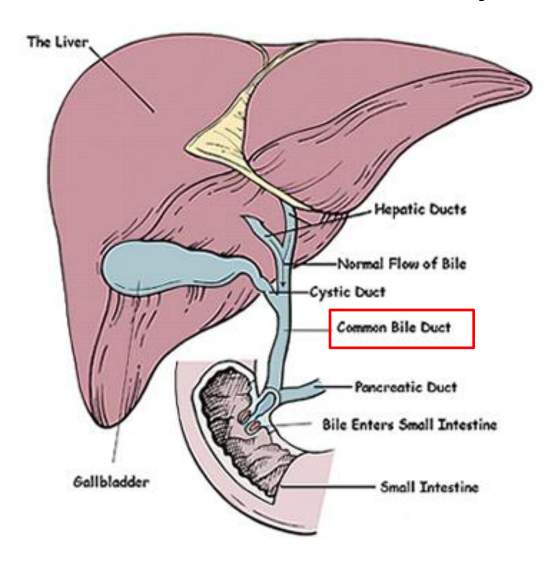
Objectives

- To understand the anatomy & physiology of liver & biliary tree
- To be able to read & interpret the basics of liver function tests
- To be able to recognize the variable presentations of acute & chronic liver disease
- To know the most common conditions causing neonatal liver diseases & chronic liver diseases in children
- To know how to diagnose these conditions appropriately

PART - 1

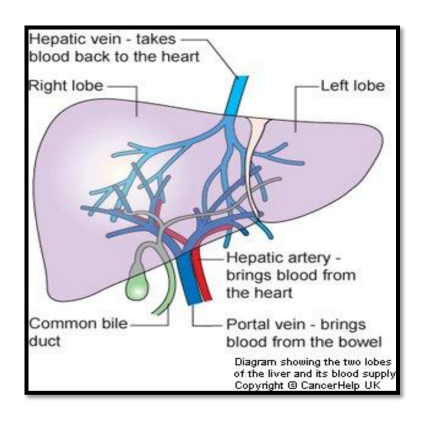
NORMAL ANATOMY & PHYSIOLOGY OF THE LIVER

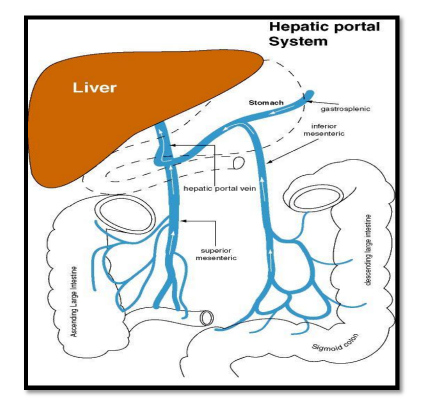
Liver anatomy



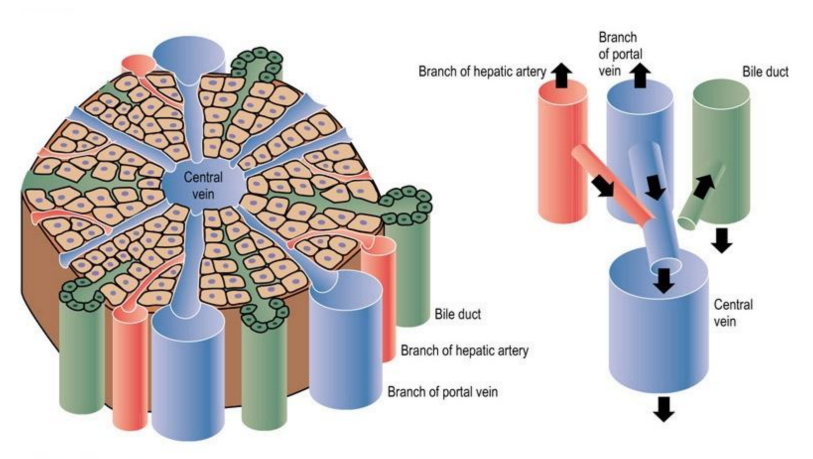
Liver blood supply

- Liver has <u>dual (bouble) Blood supply</u> resources;
 - 70% from portal vein (nutrients)
 - 30% from Hepatic artery (oxygenated blood)



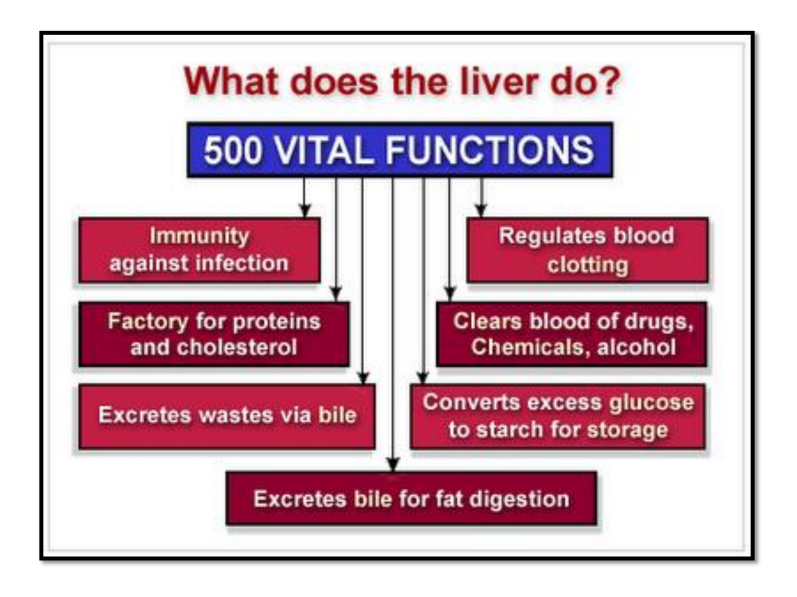


Liver Histology



© Fleshandbones.com Davies et al: Human Physiology

Liver functions



What are the liver function markers?

Liver enzymes # LFTs

- Enzymatic markers:
 - ALT
 - AST
 - ALP
 - GGT

- Synthetic function markers:
 - Glucose
 - Bilirubin (indirect/direct)
 - Bile acids
 - Albumin
 - Clotting factors (PT &PTT)
 - Urea (formed from NH3& AAs)

- The laboratory findings of <u>liver injury</u> can be divided broadly into two patterns:
 - 1) Cholestatic or obstructive bile duct injury:

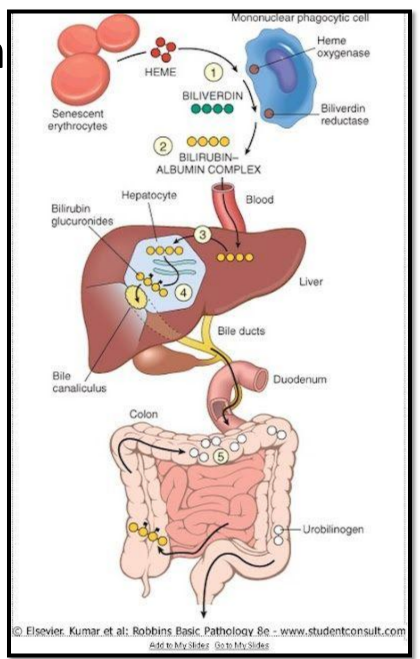
GGT /ALP > AST/ALT

2) Hepatocellular or liver cell injury:

ALT/AST > GGT/ALP

• There is often <u>considerable overlap</u> between injury types in a patient who has liver disease.

Bilirubin metabolism



Causes of un-conjugated hyperbilli

Haemolysis;

- COOMB +ve:
 - ABO incompt., Rh incompt
 - Autoimmune
 - Drug-induced HA
- COOMB ve;
 - RBC membrane defects Hbpathy- Enzyme defects
 - HUS
 - Wilson disease

Non-haemolysis;

- 1- Breast feeding/Milk Jaundice
- 2- Criglar Najjar syndrome
- 3- hypothyroidism
- 4- Gilbert syndrome,
- 5-Pyloric stenosis

Un-conjugated hyperbilirubinemia with normal LFTs

Criggler Najjar syndrome:

- reduction in glucouronyl transferase enzyme (type 2) or OR totally absence (type 1)...
- Differences??

Gilbert syndrome:

- older children & adults, observed when sick or dehydrated
- not need treatment

Conjugated hyperbilirubinemia with normal LFTs

- 1- Dubin Johnson syndrome
- 2- Rotor disease

- Present with jaundice at any age, mild
- Not require treatment

QUESTIONS FROM PART 1

PART-2

Liver disease in children

Liver disease in children

- Variable : age dependant
 - Infants: <u>Biliary atresia (BA)</u>, Neonatal hepatitis, metabolic liver disease, genetic disorders (progressive familial intraheptaic cholestasis (PFIC)
 - Older children = adults liver diseases: Viral Hepatitis, Wilson disease, Auto-immune hepatitis, ect...
- The main presenting symptoms of liver disease is jaundice
- Any jaundice after 2 weeks of age should be investigated (MCQ)

Types of liver diseases

- Liver disease can be:
- 1- Primary cholestatic/obstructive or
- 2- Hepato-cellular dominant picture
- 3- MIXED PICTURE-Usually the case

Cholestatic liver disease

- Cholestasis → chole= bile Stasis=stagnation
- The obstruction of bile flow either:
 - Mechanical block(biliary atresia, stones...) or
 - Functional block (receptor & transporter levels):eg:
 progressive familial intrahepatic cholestasis (PFIC)
- Cholestasis is <u>characterized by</u> an accumulation of compounds that cannot be excreted through the bile
 - Conjugated/direct bilirubin → jaundice (Cholestasis # jaundice)
 - Enzymes (ALT/AST>GGT/ALP)→ high liver enzymes in serum
 - Bile salts → itchiness
 - Cholestrol → xanthomas

Presentation of cholestasis

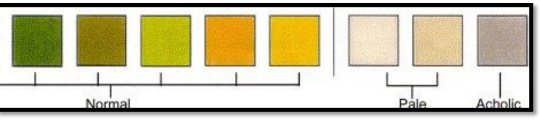
- Jaundice (accumulation of conjugated bilirubin)
- Pale stool (Acholic stool)... Why?? (MCQ)
- Dark and foamy urine (bile salts in the urine)
- Pruritis (accumulation of bile salts under the skin)
- Xanthomas depositions (accumulation of cholestrol in the skin)
- Hepatomegaly +/- Splenomegaly (Portal HTN, Storage disease, infiltrative process)
- Failure to thrive (FTT)/ poor weight gain
- Incidental lab finding

Signs of cholestatic liver disease









Evaluation of infants with cholestatic liver disease

- <u>STEP1</u>: Confirm the <u>presence of cholestasis</u> (Clinically: (jaundice, acholic stool, pruritis, & lab: direct hyperbilli)
- STEP 2: Rule out <u>surgical obstruction</u> such as Biliary atresia, Choledocal cyst and GB stones (Abdm US, HIDA scan)
- STEP 3: Investigate the treatable medical conditions:
 - Infections: UTI, TORCH infections
 - Endocrine: hypothyrodism, panhypopituitarism
 - Metabolic disorders (Galactosemia, Tyrosenemia)
- STEP 4: Further studies for other causes (genetic/metabolic)

Hepato-cellular liver disease

- Necrosis of hepatocytes following a <u>viral</u>, ischemic or toxic <u>insult</u> to the liver will cause primarily an elevation of enzymes found within the hepatocyte (<u>ALT and AST</u>)
- In hepatocellular disease, the serum levels of GGT and AP do not rise to the same degree as the aminotransferases (in general)

<u>Causes</u> of liver disease in neonates & infants (both types)

- · Cholestatic disorders
 - Biliary atresia
 - —Choledochal cyst
 - —Paucity of intrahepatic bile ducts (eg, Alagille syndrome)
 - —Progressive familial intrahepatic cholestasis syndromes (Byler disease and syndrome)
 - —Benign recurrent intrahepatic cholestasis
 - -Caroli disease and syndrome
 - —Inspissated bile (S/P hemolytic disease)
 - —Cholelithiasis

- Idiopathic neonatal hepatitis and mimickers
 - Cystic fibrosis
 - —Alpha 1-antitrypsin deficiency
 - Hypopituitarism/hypothyroidism
 - —Neonatal iron storage disease

- Viral hepatitis or other infectious diseases in the neonate
 - -Cytomegalovirus
 - —Herpes simplex virus/herpes zoster virus/human herpesvirus 6
 - —Epstein-Barr virus
 - -Parvovirus B19
 - —Rubella
 - —Reovirus—type 3
 - —Adenovirus
 - —Enterovirus
 - —Bacterial sepsis/urinary tract infection
 - —Syphilis
 - —Tuberculosis
 - —Toxoplasmosis

Causes of liver disease in neonates & infants

- Metabolic disease

 Disorders of peroxisomal function (Zellweger syndrome)
 Disorders of bile acid metabolism
 Disorders of urea cycle (arginase deficiency)
 Disorders of amino acid metabolism (tyrosinemia)
 Disorders of lipid metabolism (Niemann-Pick type C/Gaucher/Wolman)
 Disorders of carbohydrate metabolism (galactosemia, fructosemia, type IV glycogen storage disease)
 - Toxic/pharmacologic injury (eg, acetaminophen, total parenteral nutrition, hypervitaminosis A)
 - Tumors (intra- and extrahepatic)

QUESTIONS from part 2

PART-3

SPECIFIC LIVER DISEASES IN INFANTS

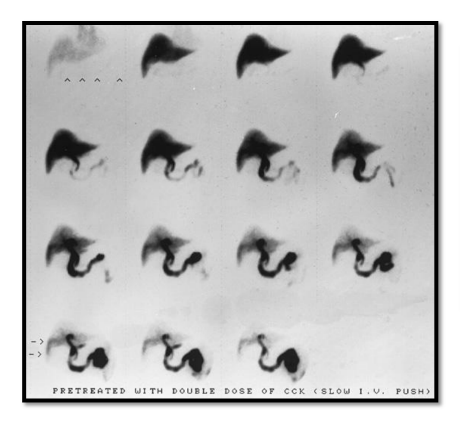
Biliary Atresia (BA)

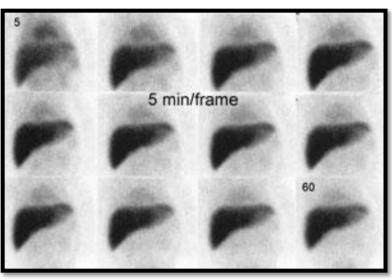
- Biliary atresia is an obstructive disease of the biliary tree (mainly extra-hepatic) secondary to idiopathic inflammatory/autoimmune process??
- It leads to gradual <u>fibrosis</u> and ultimate <u>obliteration of the</u>
 <u>biliary tract</u> → <u>biliary cirrhosis</u> → liver failure → <u>infant death</u>
 within 2 years If not treated (surgery or liver transplantation)
- The most frequent indication worldwide for liver transplantation among infants and children

BA - Diagnosis

- Clinical presentation:
 - It presents with signs of cholestasis (jaundice, acholic stool, pruritis, FTT) in the <u>first 2-6 weeks of life (MCQ)</u>
- Abdominal US: <u>rule out other causes</u> of biliary <u>obstruction</u> (choledochal cyst, GB stones...)
- Hepato-biliary scintography = nuclear scan (HIDA scan):
 - shows good uptake of tracer and then <u>NO excretion</u> into the intestine, even 24 hours later (next slide)

Hepato-biliary scintigraphy (HIDA scan)





HIDA scan in BA patient

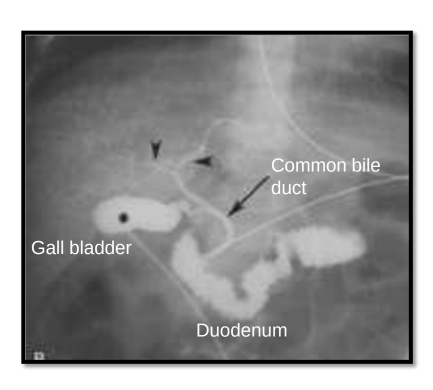
NORMAL HIDA SCAN

BA - Diagnosis

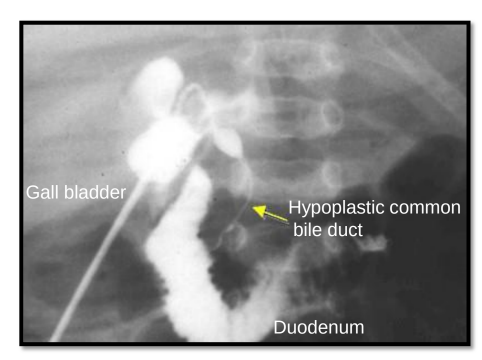
A liver biopsy:

- confirms the diagnosis by revealing characteristic findings (proliferation of the interlobular bile ducts, periportal fibrosis, and bile plugs in canaliculi and ductules)
- <u>Definitive diagnosis</u> is confirmed by Intra-operative cholangiogram

<u>Definitive diagnosis</u> is confirmed by Intra-operative cholangiogram

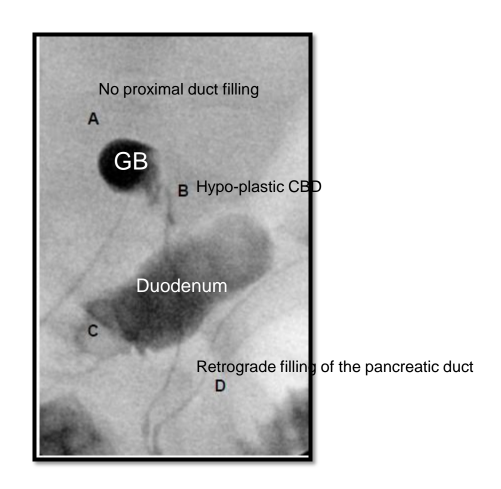


Normal study



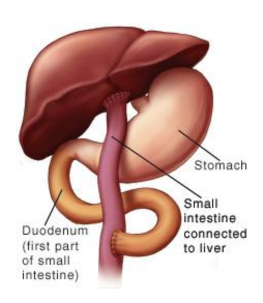
Abnormal study (hypoplastic common bile duct)

Intra-operative cholangiogram



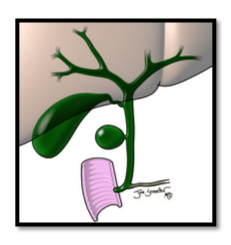
BA Management

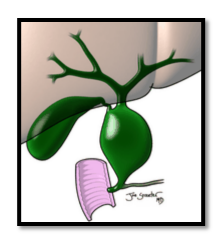
- Surgical correction (Kasai procedure or porto-entero-stomy):
- Should be done <u>before 2 months of age (MCQ)</u>
 - after this age, there is increased risk of fibrosis & subsequent cirrhosis → decrease the chance for surgery success)
- Liver transplantation if:
 - Kasai failed, or
 - if late presentation (> 3 months), or
 - picture of decompensated liver disease

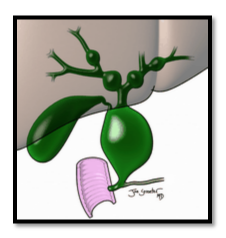


Choledocal cyst

- Cystic dilatation of the biliary tree at different levels ->
 obstructive picture
- Present with cholestasis picture, abdominal mass or asymptomatic, biliary stones or biliary carcinoma in adults



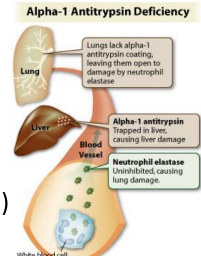




Treatment: surgical excision

Alpha-1 Antitrypsin deficiency

- A-1 AT is a <u>protease</u> (such as elastase, trypsin) <u>inhibitor</u> that protect lung from neutrophil elastase destruction → its deficiency cause <u>neonatal liver</u> disease & adult emphysema lung disease (lung dis. is rare in children)
- AR disease (rare in our community)
- Abnormal mutation (Pi MM→ Pi ZZ → form abnormal A-1 AT protein → failed excretion from liver (trapped) → cholestatic liver disease
- Dx: A-1 AT level, phenotyping (pi ZZ) and confirm with Liver biopsy (seen in special stain)
- Treatment: supportive
- Prognosis: varies (improve over time> chronic liver disease)



Neonatal Hepatitis

- "Idiopathic" neonatal hepatitis = an <u>aetiology has not been</u> <u>identified</u>
- The list gets smaller overtime (new advancement in diagnostic modalities = more genetic & metabolic causes are discovered daily)
- Management of these infants involves supportive measures till specific cause found

?? Questions PART 3

Part 4

SPECIFIC LIVER DISEASES IN OLDER CHILDREN

Liver disease in older children = adults !!

- Infectious (Viral, Bacterial, Protozoal)
- Toxic/medications (drugs, TPN)
- Ischemia (CR arrest, hypotention)
- Metabolic disorders (CHO, FAT, Amino Acids)
- Autoimmune: AIH
- Genetics; Wilson disease
- Vascular (thrombosis)
- Infiltrative/Malignancy (leukemia, primary liver tumours)

Acute hepatitis

- Five primarily viruses: hepatitis A, B, C, D, and E
- The clinical presentation of viral hepatitis varies with the pathogen (hepato-cellular injury >> mixed)
- HEPATITIS A: (MCQs)
- Presentation:
 - flu-like illness, Anorexia, fever, vomiting, abdominal pain, darkening of the urine, following ingestion of contaminated food or contact with infected patient (oral-fecal route)
- Hepatitis A is often <u>an-icteric</u> (no jaundice) in young children (<5 y) and frequently is unrecognized

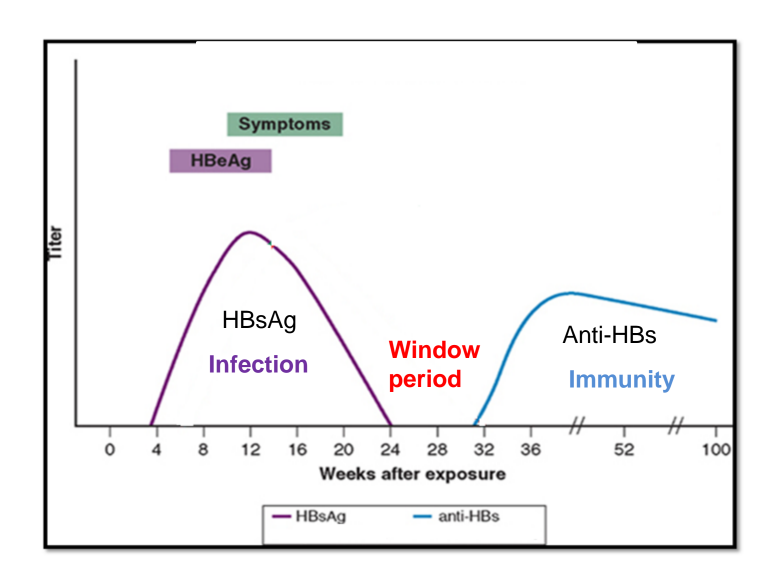
HEPATITIS A

- Diagnosis of acute infection is based on the presence of <u>anti-HAV IgM</u> antibody in serum (MCQ)
- The disease typically is <u>self-limited</u> in children and often is clinically not clear
- <u>No chronic carrier</u> state is identified (full recovery or rarely death from fulminant liver failure)
- Treatment is supportive (IVF, Antipyretics)
- Prevention: Hep. A vaccine: 2 doses (18 ms & 24 months)

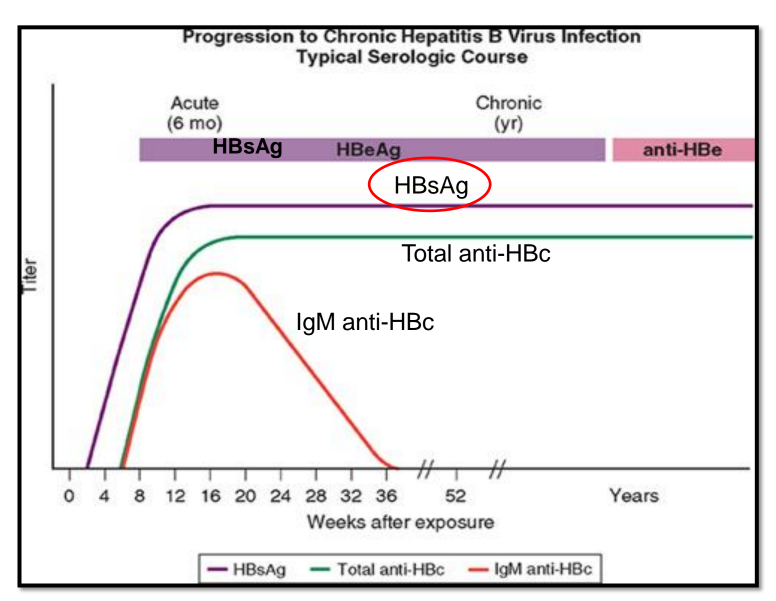
Hepatitis B

- Hepatitis B virus (HBV) infection can cause <u>both acute and chronic</u> <u>hepatitis</u>
- It can progress to cause cirrhosis and hepatocellular carcinoma if not treated (take long time to happen)
- **Risk of transmission:** primarily <u>vertical</u> (mother to baby) in children or via contaminated blood + other risk factors..
- Diagnosis: Hepatitis B surface antigen (HBsAg)
- Chronic HBV infection is associated with the <u>persistence of HBsAg</u> and HBV DNA for > 6 moths

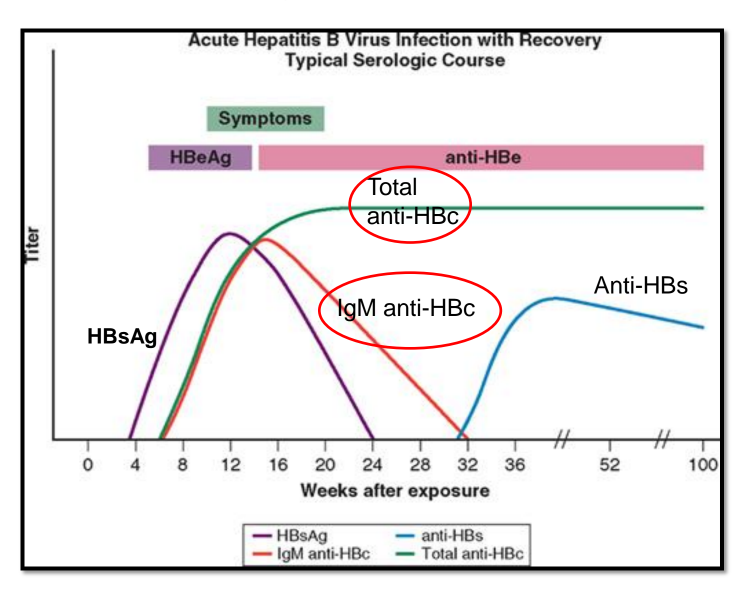
HBV serology markers



Chronic hepatitis



HBV serology markers.. recovery



Hepatitis B serological markers

(for fun, not exam!!)

HBsAg anti-HBc anti-HBs	negative negative negative	Susceptible
HBsAg anti-HBc anti-HBs	negative positive positive	Immune due to natural infection
HBsAg anti-HBc anti-HBs	negative negative positive	Immune due to hepatitis B vaccination
HBsAg anti-HBc IgM anti-HBc anti-HBs	positive positive positive negative	Acutely infected
HBsAg anti-HBc IgM anti-HBc anti-HBs	positive positive negative negative	Chronically infected
HBsAg anti-HBc anti-HBs	negative positive negative	Interpretation unclear; four possibilities: 1. Resolved infection (most common) 2. False-positive anti-HBc, thus susceptible 3. "Low level" chronic infection 4. Resolving acute infection

Treatment

- Newborn of Hep BsAg-positive mothers (MCQ):
 - Hep. B Immunoglobulins (within 12 hrs of birth)
 - Hep. B Vaccine after birth (within 7 days after birth, then at 1 month & 6 months)

- Older children: antiviral meds
 - ??? Wait & observe (spontaneous recovery, new better antiviral meds)

Hepatitis C

- Hepatitis C virus (HCV) causes acute hepatitis, which progresses to chronic disease
- End-stage liver disease can occur in up to 10 %
- Fulminant hepatitis rarely has been described
- Risk of transmission similar to hepatitis B
- Diagnosis is based on the detection of <u>anti-HCV antibodies</u> and confirmed by PCR for <u>HCV RNA</u>
- Prophylaxis: no vaccine yet
- Treatment: antiviral Rx (new generation, > 95% effective)

Hepatitis D

- Hepatitis D virus (HDV) infection occurs only in patients who have HBV infection
- HDV usually <u>aggravates liver disease</u> in a patient who has <u>hepatitis B</u> and always should be considered in those who have particularly aggressive HBV disease
- Associated primarily with <u>intravenous drug abuse</u>

Hepatitis E

 Hepatitis E virus (HEV) occurs in epidemics in parts of the world that have <u>poor sanitary conditions</u>

It can be a particularly <u>devastating disease in pregnant women</u>

?? Questions PART 4

Chronic hepatitis

Definition:

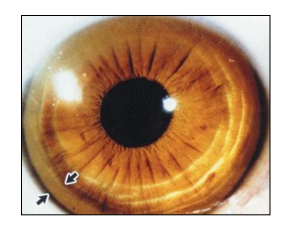
- an inflammatory condition of the liver in which the biochemical and histologic abnormalities <u>persist for more than 6 months</u> <u>from any disease</u>.
- Chronic hepatitis in children can be caused by:
 viral infection (Hep B & C); autoimmune process; exposure to
 hepatotoxic drugs; or metabolic, or systemic disorders
- Can progress to CLD if the primary disease not treated well

Chest 3 Loss of body hair 4 Abdomen: inspection Scars Distension Movement Veins Testicular atrophy Gynaecomastia A Abdominal swelling in asc Spider naevi A Face 2 Jaundice Spider naevi Parotid swelling Rhinophyma Dilated abdominal wall veins 5 Abdomen: palpation Hepatomegaly Splenomegay Xanthelasma and A (see opposite) jaundiced sclera in a patient with chronic cholestasis Hands 6 Abdomen: percussion Clubbing Dupuytren's contracture * (see opposite) Leuconychia Smooth nails (from scratching) Bruising Flapping tremor (when arms outstretched www.1aim.net Abdomen: auscultation and hands dorsiflexed Bowel sounds -see opposite) Hepatic bruit Legs Observation Bruising Unkempt Oedema · Smell of alcohol or fetor hepaticus Palmar erythema · Encephalopathy

Signs of CLD

Wilson disease (a must to know)

- AR disorder
- caused by a <u>defect in biliary copper excretion</u>
- Excessive copper accumulation in the:
 - liver → leads to cirrhosis
 - Other organs: cornea, kidneys, and brain, resulting in <u>extra-hepatic</u> manifestations of the disease
- Wilson disease SHOULD be included in the differential diagnosis of any child who presents with liver disease, neurologic abnormalities, behavioural changes (treatable condition)



Wilson disease

- Wilson disease may present as fulminant hepatic failure, usually in association with a hemolytic crisis due to the toxic effect of copper on red blood cells.
- Definitive diagnosis requires evaluation of:
 - 24-hour urinary copper excretion and
 - copper quantification in liver tissue obtained by biopsy
- **Therapy** is **chelating** therapy of the copper with <u>penicillamine</u>, which allows for its excretion into the urine (early diagnosis = better prognosis)

AIH

- AIH is a hepatic inflammation associated with the presence of circulating <u>autoantibodies against liver cells</u> in the absence of other recognized causes of liver disease
- Other autoimmune diseases may coexist, including: thyroiditis, DM

• Dx:

- High transaminases +
- High <u>autoimmune markers (anti SMA, KLM)</u>
- High serum gamma globulin concentrations
- Liver biopsy
- Rx: Immunosuppressive medications e.g.: steroids....

Ischemic hepatitis

- Ischemic hepatitis results from congestive heart failure, shock (eg, dehydration), asphyxia, cardio-respiratory arrest, or seizures.
- The disorder is due to hypotension/hypoperfusion to the liver
- Typically, aminotransaminases are elevated in the absence of other markers of severe liver disease.
- Ischemic hepatitis may resemble infectious hepatitis, but it is distinguished easily by rapidly decreasing aminotransaminases levels in the days following the initial insult without increasing coagulopathy or hyperbilirubinemia.

Infiltrative disorders

- Infiltrative disorders of the liver are observed with leukemia, lymphoma, and neuroblastoma (more common than primary liver tumers)
- Primary liver tumors: Hepatoblastoma, hepatocarcinoma, and hemangioendothelioma
- Presentation: hepatomegaly or abdominal distension or mass
- Serum <u>alpha-fetoprotein</u> levels usually are elevated.
- Dx by CT scan or MRI
- Surgical excision of a solitary tumor or radiation/chemotherapy is the treatment of choice.

THE END

QUESTIONS

TABLE 6. Miscellaneous Physical Findings Associated With Liver Disease

Infants

- Microcephaly: congenital cytomegalovirus, rubella, toxoplasmosis
- Characteristic facies: arteriohepatic dysplasia (Alagille syndrome)
- Cataracts: galactosemia
- Retinal pigmentation and posterior embryotoxon: Alagille syndrome
- Abnormal auscultation of lungs: cystic fibrosis
- Neuromuscular abnormalities (tremors, flaccidity): lipid storage disease, Wilson disease, disorders of oxidative phosphorylation

Children

- Pruritus: chronic cholestasis
- Hemangiomas: hemangiomatosis of the liver
- Kayser-Fleischer rings: Wilson disease
- Glossitis: cirrhosis
- Enlarged kidneys: congenital hepatic fibrosis or polycystic disease
- Arthritis and erythema nodosum: liver disease with chronic inflammatory bowel disease
- Arthritis, acne, fatigue: autoimmune hepatitis