Autoimmune Hepatitis

- Chronic hepatitis with immunologic abnormalities
- -Histologic features are similar to chronic viral hepatitis
- -Indolent or severe course
- -Dramatic response to immunosuppressive therapy

Features:

- 1-Female predominance (70%)
- 2-Negative serelogy for viral Ags.
- 3-↑serum lg (>2.5 g/dl)

- 4-High titers of autoantibodies (80% of cases)
- 5-The presence of other autoimmune diseases as RA, thyroiditis, sjogern syndrome, UC in 60% of the cases

The type of autoantibodies

1-Antismooth myuscle abs

anti actin

anti troponin

anti tropomyosin

2-liver/kidney microsomal Abs

anti cytochrome P-450 components

anti UDP-glucuronosyl

transferases

3-Anti – soluble liver / pancreas antigen

<u>Outcome</u>

Mild to severe chronic hepatitis
Full remission is unusual
Risk of cirrhosis is 5% which is the
main cause of death

Nonalcoholic Fatty Liver Disease

Types:

- 1.Steatosis (Fatty liver)
- 2.Steatohepatitis
 hepatocyte destruction
 parenchymal inflammation
 progressive pericellular fibrosis

Predisposing factors:

1-Type 2 DM
2-Obesity: body mass index
> 30 kg /m2 in caucasians
> 25 kg /m2 in Asians
3-Dyslipidemia (↑ TG, ↑LDL, ↓HDL)

Pathogenesis

.Metabolic syndrome

- . Insulin resistance
- . Obesity
- . Dyslipidemia

Mechanism of fatty accumulation

- 1.Impaired oxidation of fatty acids
- 2.Increased synthesis & uptake of FFA
- 3. Decreased hepatic sec. of VLDL
- . ↑TNF , IL6 , chemokine →liver inflammation & damage

Clinically

- -NAFLD is the most common cause of incidental ↑ in transaminases
- -Most pts. are asymptomatic
- -Non-specific symptoms
 Fatigue, malaise, RUQ discomfort
- -Severe symptoms
- -Liver Bx is required for dx.
- -NAFLD m.b a significant contributer to cryptogenic cirrhosis

Hemochromatosis

- Excessive accumalation of body iron (liver & pancreas)
- -1ry or 2ry (genetic or acquired)

Causes of acquired hemosidrosis:

- 1-multiple transfusions
- 2-ineffective erythropoiesis (thalassemia)
- 3-increased iron intake (Bantu sidrosis)
- 4-chronic liver disease

-Features:

- 1-Micronodular cirrhosis (all patients)
- 2-D.M (75 80%)
- 3-skin prigmentation 75-80%)
- 4-cardiomegaly, joints disease, testicular atrophy

Symptoms appear 5th – 6th decades not before age 40

- -M:F ratio 5 7: 1
- -Genetic hemochromatosis (4 variants)
- -The most common form is aut. recessive disease of adult onset caused by mutation in the HFE gene on chr.6

Pathogenesis

- -1ry defect in intestinal absorption of dietary iron.
- -Total body iron 2-6gm in adults 0.5gm in liver mostly in hepatocytes
- -In disease >50gm Fe accumulated → 1/3 in liver

-In herediatary hemochromatosis there is a defect in regulation of intestinal absorption of dietary iron leading to net iron accumulation of 0.5 – 1 gm/yr

- The gene responsible is HFE gene located on chr.6 close to HLA gene complex
- -HFE gene regulates the level of hepcidin hormone synthesized in liver
- -Hepcidin → (-) Fe. absorption from intestine
- -HFE gene deletion causes iron overload

-Two mutation can occur in HFE gene:

- 1-Mutation at 845 nucleotide → tyrosine substitution for cystine at AA 282
 (C282 Y)
- 2-aspartate substitution for histidine at AA 63 (H63D)
- 10% of pts. have other gene mutations

- -Carrier rate for C282Y is 1/70
- -Homozygosity is 1/200
- -80% of pts. are homozygous for (C282Y) mutation & have the highest incidence of iron accumulation
- -10% of pts. are either homozygous for H63D mutation or compound heterozygous for C282Y/H63D mutation

Excessive Fe deposition → toxicity of the tissues:

- 1. Lipid peroxidation
- 2. Stimulation of collagen formation
- 3. DNA damage

Morphological changes:

1-Deposition of hemosiderin in diffferent orgens

Liver

Pancreas

Myocardium

Pituitary

Adrenal

Thyroid & parathyroid

Joints

Skin

2-Cirrhosis

3-Pancreatic fibrosis

- -No inflammation
- -Fibrosis
- -Cirrhosis
- -Synovitis
- -Polyarthritis(pseudogout)
- -Pigmentation of liver
- -fibrosis of pancreas & myocardium
- -Atrophy of testes

Clinical presentation

M:F 5-7:1 5-6 the decades Hepatomegaly Abdominal pain Skin pigmentation D_.M Cardiac dysfunction Atypical arthritis Hypogonadism ↑serum Fe ferritin HCC 200x ↑in the risk

Wilson Disease

- -aut. Recessive disorder of Cu metabolism
- -mutation in ATP7B gene on chr. 13 which encodes an ATPase metal ion transporter in Golgi region
- -> 80 mutations
- -Gene freq. 1:200
- -Incidence is 1:30000

<u>Pathogenesis</u>

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Main source of Cu is from diet
Absorption of ingested Cu (2-5 mg/d)
Complex with albumin
Hepatocellular uptake
Incorporation with \alpha-2-globulin to form
Ceruloplasmin
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Sec. into plasma
(90 – 95% of plasma Cu)
Hepatic uptake of ceruloplasmin
Lysosomal degradation
Secretion of free Cu into bile
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 In Wilson disease absorbed Cu. Fails to enter the circulation in the form of ceruloplamin & the biliary excertion of Cu. is \u22c4 Defective function of ATP-7B → failure of Cu. excretion into bile & inhibits sec. of ceruloplasmin into the plasma → Cu. accumulation in liver

- Cu. Accumulation in the liver reults in:-
- 1-Production of free radicals
- 2-Binding to sulfhydryl groups of cellular proteins
- 3-Displacement of other metals in hepatic metalloenzymes

- -By the age of 5yrs. Cu. Spills over to circulation causing hemolysis & involvement of other organs as brain & cornea also kidneys, bones joints & parathyroid glands
- -Urinary exc. Of cu. ↑

Morphology

Liver

- 1-Fatty change
- 2-Acute hepatitis
- 3-chronic hepatitis
- 4-cirrhosis
- 5-massive hepatic necrosis

(rhodanine stain or orcein stain)

Brain:

Toxic injury to basal ganglia esp. the putamen causing atrophy & cavitation

Eye: kayser- fleischer rings

green – brown depositis of Cu. in descemet membrane in the limbus of the cornea

(hepatolenticular degenceration)

- Clinically
- -Presentation > 6 yrs of age
- Most common presentation is acute on chronic hepatitis
- -Neuropsychiatric presentation can occur behavioral changes Frank psychosis
 - Parkinson disease- like syndrome

- <u>DX</u>
- 1- ↓ in serum ceruloplasmin level
- 2- ↑ in urinary exc. Of Cu.
- 3- ↑ hepatic content of copper
 - > 250 mg/gm dry wt.

<u>α-1-Antitrypsin Deficiency</u>

- Aut. Recessive disorder
- freq. 1:7000 in N. American white population
- α -1-antiryrpsin is a protease inhibitor as elastase, cathepsinG , proteinase 3 which are released from neutrophils at the site of inflammation
- -The gene pi. Is located on chr.14

- At least 75 forms of gene mutation are present
- -The most common genotype is pi.MM present in 90% of individuals
- -PiZZ genotype →↓ level of α-1antitrypsin in blood (only 10% of normal) are at high risk of developing clinical disease

Pathogenesis

- The mutant polypeptide (PiZ) is abnormally folded & polymerizes causing its retention in the ER of hepatocytes
- -Although all individual with Pizz genotype accumulate α -1-AT-Z protein only 10% of them develop clinical liver disease . This is due to lages in ER protein degradation pathway

- -The accumulated α-1-AT-Z is not toxic but the autophagocytic response stimulated within the hepatocytes appear to be the cause of liver injury by autophagocytosis of the mitochondria
- -8-10% of patients develop significant liver damage

Morphology

- -Intracytoplasmic globular inclusions in hepatocytes which are acidophilic in H&E. sections
- -The inclusions are PAS-+ve & diastase resistant
- -Neonatal hepatitis cholestasis & fibrosis
- -Chronic hepatitis
- -Cirrhosis
- -Fatty change
- -Mallory bodies

Clinical features

- -Neonatal hepatitis with cholestatic jaundice appears in 10–20% of newborns with the disease
- -Attacks of hepatitis in adolescance
- -Chronic hepatitis & cirrhosis
- -HCC in 2-3 % of Pizz adults + cirrhosis

Reye Syndrome

- -Fatty change in liver & encephalopathy
- -< 4 yr.
- -3 5 d after viral illness
- -↑liver & abn. LFT Vomiting lethargy. 25% may go into coma

Pathogenesis

- -Derangement of mitochondrial function along or in combination with viral infection & salicylate
- -Microvesicular steatosis
- -Brain edema
- -Absent inflammation
- -Sk. Muscles, heart, kidneys fatty change

<u>Budd – Chiari Syndrome</u>

- -Thrombotic occlusion of the hepatic vein
- -Hepatomegaly
- -Wt.gain
- -Ascitis
- -Abd. Pain

Causes:

- 1-PCV
- 2-Pregnancy
- **3-Postpartum**
- 4-Oral contraceptive
- 5-PNH
- 7-Mechanical obstruction
- 8-Tumors as HCC
- 9-Idiopathic in 30% of the cases

Morphology

- -Swollen liver, red with tense capsule
- -centrilobular congestion & necrosis
- -Fibrosis
- -Thrombi

- Clinically
- Mortality rate is high if not treated

Primary sclerosing cholangitis

-Inflammation, obliterative firosis, & segmental dilation of the obstructed intra hepatic & extra hepatic bile ducts -In PSC, UC coexists in 70% of patients -in patients of UC, 4% develop PSC -3-5 the decades M: F 2:1

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- -asymptomatic pts. -
- -persistent ↑ serum alkaline phosphatase -
- fatigue, pruritis, jaundice, wt loss, ascitis, bleeding, encephalopathy
- -antimitochondrial Abs < 10% of cases -
- -Antinuclear cytoplasmic Abs in 80% of cases

<u>Morphology</u>

- -Concentric periductal onion-skin fibrosis & lymphocytic infilrate
- -Atrophy & obliteration of bile ducts
- -Dilation of bile ducts inbetween areas of stricture
- -Cholestasis & fibrosis
- -Cirrhosis, cholangiocarcinoma (10 15%)

Pathogenesis

- -Exposure to gut derived toxins
- -Immune attack
- -Ischemia of biliary tree

Secondary biliary cirrhosis

-Prolonged obst. To extrahepatic biliary tree

-Causes:

- 1-cholelithiasis
- 2-biliary atresia
- 3-malignancies
- 4-stricutres

Primary biliary Cirrhosis

- chronic, progressive & often fatal cholestatic liver disease
- -Non-suppurative granulomatous destruction of medium-sized intrahepatic bile ducts, portal inflammation & scarring

- -Age 20-80yrs (peak 40-50yrs)
- -F>M
- -Insidious onset
- -Pruritis, jaundice
- -Cirrhosis over 2 or more decdes

- -↑Alkaline phosphatase & cholesterol
- -Hyperbilirubinemia = hepatic decompansation
- -Antimitochondrial Abs > 90%
- Antimitochondrial pyruvate dehydrogenase
- -Associated conditions: sjogern synd. Scleroderma thyroiditis, RA, Raynauds phenomenon. MGN, celiac disease.

Morphology

- -interlobular bile ducts are absent or severely destructed (florid duct lesion)
- -intra epithelial inflammation
- -Granulomatous inflammation
- Bile ductular proliferation
- Cholestesis
- Necrosis of parenchyma
- Cirrhosis

Sinusoidal Obstruction Syndrome (Veno-occlusive disease)

- Originally described in Jamaican drinkers of bush-tea containing pyrrolizidine alkaloids
- -This occurs in the first 20-30 days after bone marrow transplantation
- . Which is caused by:
- 1-Drugs as cyclophosphamide
- 2-Total body radiation

.Incidence

-20% in recepients of allogeneic marrow transplant

-Clinical presentation

Mild – severe

Death if does not resolve in 3 months

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Mechanism

- Toxic injury to sinusoidal endothelium →emboli
- →blockage of bl. Flow
- Passage of blood into space of Disse
 - → ↑ stellate cells → fibrosis

Peliosis Hepatis

- -sinusoidal dilatation
- -Causes:
- 1-anabolic steroids
- 2-oral contraceptive
- 3-danazol
- -Pathogenesis Unknown

- -Asymptomatic
- -Intra abdominal hemorrhage
- -Liver failure

-reversible

Liver tumors

- Benign
- Most common is cavernous hemagioma
- Usually <2cm
- Subcapsular
- Liver cell adenoma
- Young female
- Hx of oral contraceptive intake
- May rupture esp. during pregnancy causing severe intraperitoneal hemorrhage
- Rarely may contain HCC
- Misdx. Of HCC

Liver Nodules

Focal noudular hyperplasia

- Well demarcated hyperplastic hepatocytes with central scar.
- Non-cirrhotic liver
- Not neoplasm but nodular regeneration
- Local vascular injury
- Females of reproductive age
- No risk of malignancy
- 20% of cases have cavernous hemagnioma

Macroregenerative Nodules

- Cirrhotic liver
- Larger than cirrhotic nodules
- No atypical features,
- Reticulin is intact
- No malignant potential

Dysplastic nodules

- Larger than 1 mm
- Cirrhotic liver
- Atypical features, pleomorphism and crowding
- High proliferative activity
- High or low dysplasia
- Precancerous (monoclonal, +ve gene mutations)
- Types:
- 1. Small cell dysplastic nodules
- 2. Large cell dysplastic nodules

Hepatocellular carcinoma

- 5.4% of all cancers
- Incidence:
 - <5/100000 population in N&S America</p>
 N& central Europe
 Australia
 - 15/100000 population in Mediterranean 36/100000 population in Korea, Taiwan mozambique, china

- Blacks > white
- M:F ratio
 - 3:1 in low incidence areas. >60yr
 - 8:1 in high incidence areas. 20-40yr

Predisposing Factors

- Hepatitis carrier state
 vertical transmission increases the risk
 200X
 cirrhosis mau be absent
 young age group (20-40yr)
- 2. >85% of cases of HCC occur in countries with high rates of chronic HBV infection

- 3-Cirrhosis
 In western countries cirrhosis is present in 85-90% of cases >60yr
 HCV & alcoholism
- 4. Aflatoxins
- 5. Hereditary tyrosinemia (in 40% of cases)
- 6. Hereditary hemochromatosis

Pathogenesis

- Repeated cycles of cell death & regeneration HBC, HCV, gene mutations, Genomic instability
- Viral integration
 HBV DNA intergration which leads to clonal expansion
- 3. HBV DNA intergration which leads to genomic instability not limited to integration site.

4. HBV

X-protein which leads to transactivation of viral & cellular promoters,

Activation of oncogenes,

Inhibition of apoptosis

- 5. Aflatoxins (fungus Aspirgillus flavus) mutation of p53
- 6. Cirrhosis

HCV

Alcohol

Hemochromatosis

Tyrosinemia (40% of pts. Develop HCC despite adequate dietary control

Morphology

- 1. HCC
- 2. CC
- 3. Mixed

- Unifocal
- Multfiocal
- Diffusely infiltrative

- Vascular invasion is common in all types.
- Well ---- Anaplastic
- Fibrolamellar carcinoma

20-40 yr. M=F
No relation to HBV or cirrhosis
better prognosis
single hard scirrhous tumor

Cholangiocarcinoma are desmoplastic

<u>metastasis</u>

Vascular – lungs, bones, adrenals, brain, in 50% of cholagiocarcinoma

C/P
 abd. Pain, malaise, wt. loss
 increase α-feto protein in 60 – 75% of pts.

- α-feto protein increases also with:
 1-yolk sac tumor
- 2- cirrhosis,
- 3-massive liver necrosis,
- 4-chronic hepatitis,
- 5-normal pregnancy,
- 6-fetal distress or death
- 7- fetal neural tube defect.

Prognosis

- Death within 7 -10 months
- Causes:
- 1-Cachexia
- 2-GI bleeding
- 3-Liver failure
- 4-Tumor rupture and hemorrhage

THE END