Board Review 2018

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Disclosures

none

Liver Diseases

disclosure

none

Alcoholic Liver disease

- Risk Factors for Alcoholic Liver Disease
 - Amount of alcohol consumed
 - Duration of alcohol consumption
 - Gender
 - Viral hepatitis
 - Nutrition
 - Iron overload
 - Genetics

Alcoholic Hepatitis

- Typically seen in malnourished patients
- Frequently precipitated by a period of binge drinking
- Prodrome: (2–3 weeks)
 - Anorexia
 - Nausea
 - Fatigue
 - Weight loss

Alcoholic Hepatitis

- Persistence of Alc. Hep. is associated with relentless progression to cirrhosis over months to years.
- Complications can be identical to those of cirrhosis.
- Poor prognostic signs:
 - Advanced age, jaundice, azotemia, and coagulopathy.

Alcoholic Hepatitis

- Clinical manifestations
 - Hepatomegaly, mild fever, jaundice
 - More severe cases: ascites, encephalopathy
- Lab
 - Increased AST&ALT \rightarrow not more than 10x normal
 - Increased AST/ALT ratio (2-3:1)
 - Decreased albumin
 - Prolonged PT

Alcoholic hepatitis-treatment

- Abstinence
- Bed rest
- Nutrition
- +/- steroids

Liver question

What is most commonly used to assess the prognosis of patients with alcoholic hepatitis?

Answer: Maddrey discriminant function analysis

- Discriminant function = 4.6(prothrombin time-control) + serum bilirubin (mg/dL)
- Discriminant function >32 effectively identifies patients whose risk of death is higher than 50%
 - Consider steroids

- Clinical
 - Nonalcoholic (<20g alcohol/day)
 - Exclusion of viral, autoimmune, genetic, and druginduced liver disease.
- Nonalcoholic Steatohepatitis (NASH)
 - Chronic inflammatory condition in people who don't have significant alcohol history.
 - Characteristics: steatosis, hepatocellular necrosis, and inflammation.

Fat liver—pale yellow coloring



- clinical manifestations
 - Central obesity (apple shaped not pear-shaped)
 - Abd. Obesity (waist >40" in men and 34.5" for women)
 - NIDDM
 - +/- hyperlipidemia
 - Most patients are asymptomatic
 - Occasional RUQ discomfort, malaise, fatigue
 - Hepatomegaly \rightarrow 75% of patients

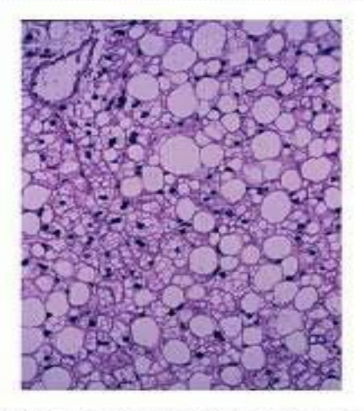
- Lab
 - Elevated aminotransferase (<300UI/L)
 - AST/ALT ratio <1
 - Mild elevation alkaline phosphatase and GGTP

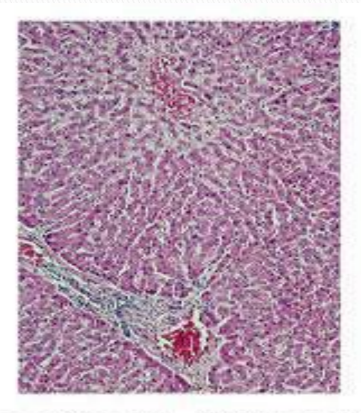
Diagnosis

- Findings of fatty infiltrate on imaging studies.
- Exclusion of other liver diseases by history, physical, and serology.
- Alcohol consumption should be <40g/week.
- Liver biopsy is the definitive method of diagnosis.
 Not indicated in asymptomatic patients with normal AST, ALT.

- Histologic finding
 - Steatosis-macrovasicular mild to severe
 - Inflammation
 - Hepatocyte injury- focal necrosis and ballooning
 - Hepatocyte degeneration- mallory hyaline
 - Fibrosis- varying degree

Liver





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Fatty liver

Normal liver

Management

- Directed at associated risk factors.
- Gradual weight loss.
- Control of hyperglycemia and hyperlipidemia.
- Discontinue suspected meds.
- Alcohol use <20g/day. Alcohol abstinence if significant fibrosis
- HAV and HBV vaccination
- Avoid drugs that may promote steatohepatitis (amiodarone,tamoxifen)

Viral Hepatitis

Hepatitis **B**

A DNA virus.

- Risks in US: sexual promiscuity and IVDA
 - Many immigrants likely contracted at birth or young childhood
- Prevention:
 - Hep B immune globulin should be given to household and sexual contacts of patients with acute hepatitis B.
 - Infants and previously unvaccinated should receive hep B vaccine.

Hep B Serologic Markers

1.HBsAG

- 2. Anti–HBs
- 3. IgM anti-HBc

1.Current infection

2.Immunity (immunization or resolved infection)3. Recent infection, occasionally reactivation

4. IgG anti-HBc5.HBeAg and/or HBVDNA>105 viral copies/mL

4. Remote infection5. Active viral replication

Interpretation of Hep B serologic panel- examples

- HBsAg +
- Anti-HBc +
- ▶ IgM anti-HBc +
- Anti-HBs –

- HBsAg +
- Anti-HBc +
- IgM anti-HBc -
- Anti-HBs

Acutely infected

Chronically infected

Нер В

Treatment –when?

- If pt at increased risk of progression:
 - LFTs >2x normal,
 - active viral replication (HBV DNA increased),
 - and active disease identified in liver biopsy specimens

Hep B treatment

- Interferon
 - Pegylated—once weekly and better efficacy
- Oral agents
 - Lamivudine, Adefovir, Entecavir
 - Become popular for treatment of chronic hepB
 - Few side effects
 - Adefovir \rightarrow nephrotoxicity
 - Useful in pts with decompensated cirrhosis

Hepatitis question

- > 21 yo presents to the ER with abdominal pain, fatigue, and loss of appetite. He admits to iv heroin use and drinks 2-3 beers/d.
- PE: mild icterus, hepatomegaly-tender, +needle tracks antecubital
- Lab: Tbili: 5.6mg/dL; AST & /ALT 950 & 1280 alkPhos 115; albumin 3.4
 - HBsAg HAV IgM –
 - HBsAb + HCV Ab -
 - HBclgM –
- What lab is most likely to make a diagnosis?

Hepatitis question

- What lab is most likely to make a diagnosis?
- a. Antimitochondrial and anti smooth muscle Ab
- b. HCV RNA
- c. HCV RIBA
- d. HBc IgG
- e. HAV total

Answer: b

- > Pt with signs and symptoms of acute hepatitis
- Initial serology shows immunity to HepB otherwise negative
- With active iv drug use acute Hep C must be considered
 - HCV ab may take up to 6 weeks to develop
- RIBA confirms + HCV ab
 - No longer recommended by CDC
- HAV total and HBV signal prior exposure and not for acute disease concerns

Hepatitis C

Leading indication for liver transplantation

Diagnostic tests

- anti-HCV: indicates current infection or previous exposure with clearance.
- "gold standard" presence of HCV RNA by PCR—now the preferred test, bypassing RIBA.
- Level of RNA does not correlate with severity of disease.
- Genotyping: genotype 1most common in US

HCV testing recommendations(CDC)

Adults born from 1945 through 1965 should be tested once (without prior ascertainment of HCV risk factors)

- Currently injecting drugs; Ever injected drugs
- Have certain medical conditions, including persons:
 - who received clotting factor concentrates produced before 1987
 - who were ever on long-term hemodialysis
 - with persistently abnormal alanine aminotransferase levels (ALT)
 - who have HIV infection

HCV

- Were prior recipients of transfusions or organ transplants, including persons who:
 - were notified that they received blood from a donor who later tested positive for HCV infection
 - received a transfusion of blood, blood components, or an organ transplant before July 1992
- HCV testing based on a recognized exposure is recommended for:
 - Healthcare, emergency medical, and public safety workers after needle sticks, sharps, or mucosal exposures to HCVpositive blood

Children born to HCV-positive women

Hepatitis question

- 56 yo male presents to his pcp for an annual physical. He is in good health, but is found to have ALT of 86 and a skin rash diagnosed by his dermatologist as porphyria cutanea tarda. The pcp should order which blood test to help explain the findings?
- a. Hep A IgM
- b. Hep B surface Ab
- c. Hep C Ab
- d. Hep E Ab

Answer: c

- Hep C associated with a variety of dermatologic findings
- HCV therapy may result in resolution of the skin findings.

Hepatitis C

Subgroup of pts likely to develop progressive liver disease

- 1. duration of infection
- 2. alcohol intake >50g/d
- 3. coinfection with HIV or HBV
- 4. male sex

Pts with cirrhosis due to HCV generally have disease >20 years.

Hepatitis question

- 19 yo college student presents with 8 days of N/V/D and fatigue. She recently returned from a 2 week mission trip to Haiti.
- PE: low grade fever, tender hepatomegaly, mild scleral icterus
- Lab: Tbili 4.9; AST 1280 ALT 1980; Alk 99 INR 0.9
- Which of the following lab tests is most likely to reveal the diagnosis?
- a. HAV IgM
- b. HAV total
- c. CMV stool PCR
- d. HBsAb
- e. HCV Ab

Answer: a

- Pt presents with acute hepatitis
- Recent travel to endemic area
- HAV total: only reveals prior infection and immunity
- *CMV* unlikely with no history of immunosuppression
- HBsAb describes immunity rather than acute
 - Given her age, likely vaccinated as baby
- No clear risk factors for HCV exposure

Hepatitis D

- A defective virus
 - Requires the presence of HBsAg to replicate

Hepatitis E

- Single stranded RNA
- The highest incidence of HEV infection is in Asia, Africa, Middle East, and Central America.
- HEV is the second most common cause of sporadic hepatitis in North Africa and the Middle East.

Hepatitis E

- HEV is spread by fecally contaminated water in endemic areas
- Person-to-person transmission is uncommon
- HEV can be transmitted by blood transfusion, particularly in endemic areas

Cirrhosis--Complications

Portal hypertension

- an increase in hepatic venous pressure gradient.
- In cirrhosis it occurs through an increase in resistance to portal venous outflow
 - Due to distortion of liver
 - ~30% of the increase is through potentially reversible vascular factors---where pharmacotherapy targets

Esophageal varices

- risk factors for hemorrhage from esophageal varices:
 - radius of varix,
 - thickness of varix wall
 - pressure gradient between the varix and the esophageal lumen.

Esophageal varices

Recommendations for treatment of esophageal varices

Primary prophylaxis: all patients with cirrhosis should have EGD for screening.

If no varices repeat endoscopy in 2-3 years.

-1st line therapy : nonselective beta blockers (propranolol or nadolol)

-2nd line therapy: endoscopic band ligation

 Control of bleeding: best managed by endoscopic means preferable band ligation.

- begin octreotide, continue for up to 5 days.

-2nd line therapy: TIPS

Esophageal varices

 Secondary prophylaxis: prevent rebleeding. Essential—80% of patients who bleed will have a rebleed within 2 years.

1st line therapy: endoscopy and beta blockers.

other: liver transplantation

EGD esophageal varices



Liver question

- A 47 yo female presents with new onset ascites that has developed over ~ 4 months. She denies ETOH, +tobacco. She is obese but no other medical problems.
- +fatigue, decreased appetite, dyspnea
- Diagnostic paracentesis: ascites albumin 1.5g/dL, ascites protein 2.6g/dL, and ascites cell count 101 neutrophils/mm3. Her serum albumin is 2.9.
- US is limited due to body habitus, reveals patent portal and hepatic veins.

Liver question

- What is the next most appropriate diagnostic step?
- a. Exploratory laparotomy
- b. Echocardiogram
- c. Cytologic analysis of the fluid
- d. Triple phase CT scan of the liver

Answer: b

- Pts SAAG (serum-ascites albumin gradient) is elevated at 1.4g/dL
 - Differential: cirrhosis, CHF, pericardial disease, Budd Chiari, and veno-occlusive disease
 - Mildly elevated protein in her ascitic fluid necessitates cardiac evaluation

Ascites

- Pathogenesis: renal retention of sodium and movement of this extra fluid into the peritoneal space.
- diagnostic paracentesis is essential for patients who present with ascites.
 - the difference between serum albumin and ascitic albumin help determine portal hypertension (1.1g/dL or greater). Could be liver or heart disease. (SAAG)
 - A protein of 2.5g/dL or more favors heart disease.
- cell count of more than 250 neutrophils/mm3 is spontaneous bacterial peritonitis (SBP).

Management of Ascites

- Iow sodium diet
- fluid restriction: only necessary if serum sodium is <125mEq/L</p>
- diuretic therapy:
 - urinary sodium excretion is used to determine the efficacy of therapy.
 - If urinary sodium excretion is more than 30mEq/d, spironolactone alone may be used.

If urinary sodium excretion is between 10-30mEq/L then a combination of spironolactone and furosemide is used.

If urinary sodium excretion is < 10mEq/L then large volume paracentesis is usually required.

Spontaneous Bacterial peritonitis

- End-stage liver disease
- No secondary source
- Clinical manifestations
 - Fever
 - Abdominal pain/tenderness
 - Altered mental status
- Index of suspicion should be high

SBP- Diagnosis

- +bacterial culture
- And/or pmn >250 cells/mm3

Hepatic encephalopathy

Pathogenesis:

 Ammonia and manganese considered etiologic factors for encephalopathy.

Clinical features:

 range from 0—no overt encephalophy to IV patient in a coma.

Precipitating factors:

 GI bleed, infection, large protein meal, use of sedatives, electrolyte abnormalities or hypoxia, constipation, and hypoglycemia.

Hepatic encephalopathy-Management

- dietary: limit protein based on level of encephalopathy.
 - Long-term restriction of dietary protein of < 1g/kg daily should be avoided.
- Nonabsorbable disaccharides:
 - Lactulose, may help remove dietary and endogenous ammonia.
 - Pt should have 2-3 semiformed stools/day.
- Antibiotics:
 - neomycin, metronidazole, and rifaximin have been used for treatment.

Liver diseases

Liver question

- 46 yo asymptomatic male has a brother with hemochromatosis
- Exam is normal. He drinks 2 beers/day
- Lab: fe 180ug/dL, Transferrin sat 88%, ferritin 1200ug/L. CBC nl, AST 52 US normal.
- HFE gene test + C282Y/C282Y mutation

Most appropriate next step would be:

- a. Liver biopsy
- b. Therapeutic phlebotomy
- c. Stop etoh and repeat iron studies in 1 year
- d. MRI of the liver

Answer: a

- Ferritin <1000ug/L and normal AST→phlebotomy
- Ferritin >1000ug/L and/or elevated AST→ liver biopsy and then phlebotomy
- Normal ferritin \rightarrow repeat ferritin q 2–3 years

Hemochromatosis

- autosomal recessive disorder with increased intestinal absorption of iron.
- Excess iron is deposited in the liver, pancreas, and other organs.
- About 1 in every 250 white persons in the US is homozygous for the mutation.

Hemochromatosis

- Suspect in pts with elevated iron sat, ferritin, or family hx.
- Most pt asymptomatic
- Cirrhosis, heart failure, hypogonadism, and arthritis
- HFE gene mutation
 - Autosomal recessive dz
 - 85% homozygous for C282Y mutation

Hemochromatosis-Treatment

- reserved for patients with evidence of iron overload, indicated by an increase in the serum concentration of ferritin.
 - therapeutic phlebotomy: simple, relatively inexpensive and effective.
 - avoid supplements with iron
 - avoid raw fish due to risk of Vibrio vulnificus infection
 - avoid alcohol
- If diagnosed and treated before diabetes and cirrhosis develops survival rate is normal

Liver question

- 18 yo male is seen for 6 month hx of abnormal liver tests. He is asymptomatic.
- Recent poor school performance and ADD
- PE: mild obesity, no stigmata of chronic liver dz.
- Lab: AST 65, ALT 87 bili 1.2 ALP 120. Hepatitis panel, ANA negative. Ceruloplasmin 19.2 (nl 22.9-43.1) Eye exam neg. 24h urine Copper >40. bx: mild steatosis, minimal inflammation copper >250mcg/g
 What is the most likely diagnosis?

What is the most likely diagnosis?

- a. Primary biliary cirrhosis
- b. Wilsons disease
- c. Drug induced liver disease
- d. Autoimmune liver disease
- e. Fatty liver

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Wilson's disease

- Inherited
- Excess copper
- Hepatic, neurologic, and psychiatric manifestations
- Gene mutation
 - ATP7B genes
- All ethnic groups
- ▶ ~1 in 30,000

Wilson's disease

Diagnosis

- Reduced ceruloplasmin
- Increased urinary excretion of copper
- Presence of K–F rings
- Elevated hepatic copper level
- Treatment
 - Copper-chelating medications

Wilson's

- Kayser-Fleischer rings (KF)
- Seen with slit-lamp



Liver question

- 16 yo presents with AST and ALT elevation for 4 months. Originally felt to be mono, due to fatigue and low grade fever. However, monospot was negative.
- > PE: no stigmata of chronic liver disease
- Lab: AST 356 ALT 435. Tbili 1.1 PT 13.2, hepatitis panel neg, ANA 1:640, Anti-smooth muscle Ab 1:320, AMA normal Liver bx: cirrhosis with increased lymphoplasmocytes
 What is the likely diagnosis?

What is the likely diagnosis?

- a. Autoimmune hepatitis
- b. Primary biliary cholangitis
- c. Wilsons disease
- d. Acute viral hepatitis
- e. Primary sclerosing cholangitis

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Autoimmune Hepatitis

- Occurs in children and adults
 - 3.6 to 1 female to male
- All ethnic groups

Clinical Manifestations

- Asymptomatic \rightarrow liver failure
 - Subclinical
- Present with cirrhosis

Diagnosis

- Aminotransferase elevation
- ANA
 - Anti-smooth muscle antibody
- hypergammaglobulinemia
- Histology: nonspecific
 - Portal mononuclear cell infiltration
 - Lymphoplasmacytic
 - fibrosis

Treatment -Autoimmune Hepatitis

- Liver transplant
- Prednisone
- azathioprine

Autoimmune Hepatitis: Typical lab

- Increased AST and Alt 100% Increased gamma globulin
- and IgG 90% 83%
- Mild hyperbilirubinemia
 - \cdot <3 mg/dL
- Alkaline phosphatase increase 67%
 - <2x normal
- ANA, SMA, or anti–LKM1 87%

Primary Biliary Cirrhosis

- Cholestatic liver disease
- 90% women
- 95% will be AMA + (anti-mitochondrial Ab)
- Fatigue common
- Pruritis 30–50%
- Frequently being picked up in pts with asymptomatic lab abnormalities
- IgM high

Alpha1-antitrypsin (AAT) deficiency

- Autosomal co-dominant disorder with lung and liver injury
- Can cause premature emphysema and liver disease
- Pt with cirrhosis due to AAT have a significant increased risk of HCC up to 30%
- Diagnosed by phenotyping. Liver damage does NOT correlate with serum AAT levels (unlike lung). Diagnosis confirmed with biopsy
- No effective medical treatment for the liver manifestations of AAT deficiency.