Outline/Objectives

• Cirrhosis
  – Etiology
  – Progression
• Complications of Cirrhosis
  – Portal Hypertension
    » Varices
    » Ascites
    » Hepatohydrothorax
    » Spontaneous Bacterial Peritonitis
    » Hepatorenal Syndrome
  – Hepatic Encephalopathy
• Therapeutic Options
  – Endoscopic Therapy
  – Surgical Shunts
  – Liver Transplantation
What is Cirrhosis?

• End stage of chronic liver disease
• Regenerative Nodules surrounded by fibrous septa
• Disruption of the architecture
• Common Etiologies
  – Alcohol
  – Viral hepatitis
  – Non-alcoholic Fatty Liver Disease
  – Genetic/Metabolic
Pathway to Cirrhosis

Injury

Inflammation  Fibrosis

Resolution

Cirrhosis

Viral Hepatitis
Alcohol
Steato
PBC
???????
Cirrhosis-pathogenesis

- Stellate cell or Ito cell: normally, sit in Space of Disse and store Vit. A; during cirrhosis, transform into myofibroblasts that make and deposit collagen
- Probably stimulated by inflammation, cytokines, and toxins
- Collagen types I and III are deposited in all portions of lobule
- Ultimately, architecture and vasculature are disrupted and diffusion of solutes is impaired
CIRRHOTIC LIVER
CIRRHOTIC LIVER
Decreased clearance of Estrogen
Increased Free Estrogen due to decreased sex steroid-binding globulin

Bones
osteoporosis
& osteomalacia
Esp: PBC, PSC
Complications

• **Portal Hypertension**
  – Varices
  – Ascites
  – Hepato-renal Syndrome

• **Liver Failure**
  – Hepatic Encephalopathy
  – Protein Loss
    » Coagulopathy
    » Decrease muscle mass
    » Infection
**LIVER: Blood Flow**

- **High Flow**
  - Mesenteric vessels

- **Low pressure**
  - Vast network of sinusoids
Portal Hypertension

- **Pre hepatic**
  - Portal Vein or Splenic Vein Thrombosis

- **Intra hepatic**
  - Pre-sinusoidal
    - Schistosomiasis
  - Sinusoidal
    - Cirrhosis
  - Post Sinusoidal
    - Veno-occlusive disease

- **Post hepatic**
  - Obstruction of vena cava of hepatic vein (Budd-Chiari)
Cirrhosis: Portal Hypertension

Effects on blood flow

1. Fibrosis restricts blood flow; increased portal vein pressure
2. Collaterals acquire increased pressure, affecting spleen, esophagus, stomach (varices), gastropathy
3. Ascites due to fluid shift into peritoneum
4. Shunting of blood from liver decreases of metabolism of "toxins"
5. Portal bacteremia not cleared induces peritonitis
What Pressure Defines Portal HTN?
Portal Vein-Hepatic Vein Pressure Gradient

- Catheter (with deflated balloon) in Hepatic Vein measures “free HV pressure”
- Inflate Balloon in Hepatic Vein measures Portal Vein Pressure
- PV-HV pressure > 10 mmHG = “significant” portal HTN Normal gradient is less than 7
Portal Hypertension
Pathophysiology

• Increased Resistance to inflow/outflow
  – Fixed scarring of the liver
  – ? reversible elements: sinusoidal blood vessels

• Increased Flow to portal system
  – Increased splanchnic flow (vasodilatation/NO)
  – Increased cardiac output
    » Low SVR
  – Increased blood volume

• Decreased Albumin: hepatic synthetic dysfunction
  » Decreased oncotic pressure, fluid leaks out of vascular space
PHYSICAL EXAM FINDINGS

Left image: A close-up of a red area on the skin.
Right image: A pattern of veins on the skin with a centimeter ruler for measurement.
Esophageal Varices

- Risk of bleeding proportional to size and degree of portal hypertension
- Even with optimal therapy, death from initial bleed may be greater than 20%
- Non-selective Beta blockers (Propranalol, Nadolol)
- Use of Isosorbide Mononitrate controversial
Gastric Varices
Portal Gastropathy

Normal

Gastropathy
Esophageal Varices

- **Bleeding Control: Acute**
  - Stabilize hemodynamics
  - Decrease portal Pressure
    » Octreotide, Somatostatin
  - Endoscopy
    » Sclerotherapy: past
    » Banding: present
- **TIPS**
- **Surgical Shunt**
- **Chronic therapy**
  » Beta Blockers/Nitrates
  » Banding ablation
- **Transplantation**
Varices/ Banding
TIPS

Transjugular Intrahepatic Porto-Systemic Shunt
TIPS

- Functional side to side shunt
- Produces connection from Portal Vein to Hepatic Vein
  - Increase Hepatic vein pressure, RV volume
  - Decrease Portal pressure, hepatic perfusion
- TIPS improves sodium and water handling
  - Hepatic hydrothorax
  - Refractory ascites
- Caution:
  - CHF, Bili > 4, Inc Creat, PSE, Older pt
Surgical shunts

• Limited indications: for endoscopic, medical failure, not OLT candidates
  – Cirrhosis: Child’s A
  – Budd Chiari syndrome
  – Non-cirrhotic portal hypertension

• Selective vs non-selective
  – Goal is to preserve portal perfusion
Surgical Shunts

- Porto-caval
  - End to side
    - Nonselective total shunt

- Meso-caval
  - Side to side
    - Nonselective partial shunt

- Spleno-renal
  - Selective shunt
Blakemore Tube

- Historically, an important way to stabilize a patient with variceal bleeding prior to:
  - Surgery
  - Transplant
- Now, only used in emergencies
  - Prior to TIPS
  - To Transport a patient from hospital to hospital
Ascites
Clinical Diagnosis

• History: increasing abdominal girth

• Physical Examination:
  – shifting dullness, fluid wave
  – very poor in detecting modest amounts of ascites

• Radiology: ultrasound, CT scan more sensitive
Ascites

- Differential dx:
  - Portal hypertension
  - Hepatic (or portal vein) occlusion
  - Heart failure
  - Peritoneal inflammation
    - TB peritonitis
    - Carcinomatosis (sometimes chylous ascites)
  - Ovarian Cancer
  - Nephrogenic ascites (nephrotic syndrome)
  - Pancreatic ascites
  - “Other” (Schistosomiasis, non-cirrhotic portal HTN, polycystic liver disease,
Portal HTN and Ascites

Cirrhosis
  ↓
Splanchnic arterial vasodilation
  ↓
Arterial underfilling
  ↓
Stimulation of systemic vasoconstrictors*
  ↓
Renal vasoconstriction

Late Stages of Cirrhosis
  ↓
Decrease in local vasodilators with increase of local vasoconstrictors
  ↓
Hepatorenal Syndrome

Early Stages of Cirrhosis
  ↓
Increased systemic and local vasodilators
  ↓
Preserved renal perfusion

Portal hypertension

Decreased systemic vascular resistance
Probability of Survival After Ascites Diagnosed

- At risk 263 149 100 56 39 26
- Months
- No complications
- Dilutional hyponatremia
- Refractory ascites
- Hepatorenal syndrome
- Spontaneous bacterial peritonitis
- Months
Characteristics of Ascites due to Portal Hypertension

- Transudate; i.e., ascites protein < 3 g/dl; most < 1 g/dl
- WBC < 50 cc/mm$^3$; mostly mononuclear
- Normal ascitic fluid amylase
- Serum - ascites Albumin gradient (SAG) > 1.1 g/dl due to portal HTN
Ascites: Treatment

- **Bedrest**
  - Na\(^+\) restriction; 1.5-2 gms/day
  - fluid restriction: 1.5 liters if Na\(^+\) < 120
- **Diuretics (maximum doses):**
  - Spironolactone 400mg, Furosemide (200 mg)
  - Amiloride, HCTZ, Metolazone, Zaroxyln
- **Large-volume paracentesis**
- **TIPS**
- **Surgery:** Liver transplantation
  - Leveen or Denver Shunt (historical value, ? If valuable now, radiologists now place these
Large-Volume Paracentesis

- **Advantages:** Fast, ↓ hospital time, less expensive
  - Patients should have normal creatinine
  - Better if volume overloaded (peripheral edema)

- **Disadvantages:**
  - Precipitate renal insufficiency
  - Removes proteins (e.g., opsonins)

- **Use of volume expansion**
  - Albumin: 6 gms/liter of ascites removed
  - May not be required for < 2-3 liter paracentesis
LeVeen/Denver peritoneo-venous shunt

- **Coagulopathy:**
  - DIC almost universal
  - severity can be limited by replacing ascites with saline

- **Infection**
  - generally requires removal of shunt

- **Occlusion**
  - Venous side of shunt

- **Heart failure**
  - Volume overload
Hepatic Hydrothorax

- Ascites leaks through rents in the diaphragm
- Diagnosis: Fluid should have characteristics similar to ascites
- Treatment: AVOID CHEST TUBES. Surgical repair not usually effective
- TIPS is treatment of choice for diuretic-refractory cases
- Liver transplantation
SBP
Spontaneous Bacterial Peritonitis

- Infection independent of another intra-abdominal source
- Monomicrobial
- Enteric flora enters portal circulation, not cleared
- Ascites WBC > 500 or 250 with greater than 50% polys
- Culture negative neutrocytic ascites
- Culture positive neutrocytic ascites
- Culture positive non-neutrocytic ascites
  - If gm negative: treat
  - If gm positive: likely contaminant
Spontaneous Bacterial Peritonitis: Prevention

- **Risks:**
  - GI bleeding/hypotension
  - Advanced liver disease
  - Previous history!
- Early treatment of other infections
- Prophylactic antibiotics to GI bleeders
- Volume expand with Albumin
  - Effective to reduce hepato-renal syndrome
- Oral Quinolones, Bactrim can prevent recurrence when given chronically
- Liver transplantation
Spontaneous Bacterial Peritonitis: Treatment

- Most common organisms are E. coli, Klebsiella, Pneumocococcus, Enterococcus
- Broad Spectrum antibiotics and then narrow antibiotic spectrum if culture results are known
- Re-tap after 48 hours to confirm response to therapy
Portal HTN and HRS

- Early-Decrease in SVR is compensated by increased HR, CO
  Stimulation of RA and SNS, ADH
- Late-Splanchnic circulation is resistant to AngII, Vasopression,
  pressure is maintained by local vasoconstriction
**Hepato-Renal Syndrome**

- **Etiology:** Unclear, but likely an exaggeration of mechanisms involved in ascites formation
- **Precipitants:**
  - Gi Bleed
  - Nephtotoxins (NSAID’s, Aminoglycosides, sepsis)
  - Iatrogenic (diuresis, paracentesis)
- **Diagnosis:**
  - Euvolemic patient
  - Urine output < 800 cc/day, $U_Na < 10$ mEq/l,
  - “clean urine sediment”
- **Treatment:** TIPS, MARS, Glypressin, Terlipressin, Transplantation
  - Midodrine 5-15mg po tid
  - Octreotide sq 100-200 mcg sq tid
Portal Systemic Encephalopathy: Hepatic Encephalopathy

- Inability to clear “encephalopathogenic agents” (Ammonia, Gaba, Mercaptans, endogenous Benzos)
  - Cirrhosis
  - Portal Hypertension
  - Shunting (TIPS, surgical shunt)
  - Protein load
    - Usually GI bleed, Gastropathy, less common PO proteins
- Acute Liver Failure:
  - PSE defines fulminant Hepatic Failure
  - Cerebral Edema (not in chronic!!)
  - Emergency Liver Transplantation is therapy
Treatment

• Decrease encephalopathic agent:
  – Lactulose 30cc po q 2 until effect (traps NH3 in colon or NH3 incorporated into bacterial proteins)
  – Rectal Tube
    » Tap water Enema
    » Lactulose 200 cc in 300 cc tap water

• Decrease production/block EA:
  – Rifaximin
  – Neomycin 500 mg po q 6 (watch Creatinine and hearing)
  – Flagyl 500 mg po q 8 (neuropathy, antabuse effect)

• “Brain Stabilizers”
  – Zinc and/or L-Carnitene

• Cathartic:
  – Mg Citrate, Miralax, Go-lytely
HepatAssist® Bioartificial Liver Support System
Indications for Liver Transplantation

- Manifestations of Portal HTN not controlled by alternative measures
  - Esophageal and/or gastric variceal bleeding
  - Bleeding from portal hypertensive gastropathy
  - Hepatic encephalopathy
  - Spontaneous Bacterial Peritonitis
  - Significant Ascites, hydrothorax
  - Hepatocellular Carcinoma
Summary

- Cirrhosis is the end stage of many diseases that cause liver inflammation.
- Complications of cirrhosis are related to loss of synthetic function and portal hypertension.
- Portal Hypertension is the cause of variceal bleeding, ascites and hepatorenal syndrome in patients with liver disease.
- All offered therapies other than liver transplant are supportive, not curative of end-stage liver disease.