25% of cirrhotics have varices
   25-33% with varices will bleed at some point
   25% mortality with variceal bleed
   66% incidence of rebleeding without intervention
   50% of pts with varices who bleed, bleed from other causes

normal portal pressure 8mm, higher 2/3 develop varices, 1/3 bleed > 12mm
varices bleed by rupture, not erosion

Location of blockage

prehepatic: portal V, splenic V, tumor at porta
intrahepatic: cirrhosis (alcoholic, hepatitis, toxic injury)(polycystic liver and Caroli’s may be complicated by portal hypertension)
posthepatic: Budd-Chiari

Encephalopathy

cause unclear; nitrogenous compounds contribute
sepsis, constipation, dehydration, blood in gut
no diagnostic test (serum NH₄, EEG non-diagnostic)
treat dehydration, decrease protein intake, cleanse gut (lactulose, induces diarrhea, does not decrease colonic flora), treat other causes

Ascites

when sinusoidal pressure exceeds colloid oncotic pressure
induced by physiologic stress, IV fluids (decreased colloid oncotic press)
complications
   spontaneous bacterial peritonitis
   hepatorenal syndrome (extreme prerenal azotemia)
control:
   medical therapy is treatment of choice
   Na/H₂O restriction, spironolactone (promotes Na diuresis), then give loop diuretic (lasix) if necessary
   large volume paracentesis (several liters) is safe
   replace albumin?
peritoneal-venous shunt
   clog early with protein
   sepsis within 6 weeks
   increases chance of variceal bleeding (increases circulating volume)
   contraindicated: bacterial peritonitis, uncontrolled coagulopathy, CHF
transjugular intrahepatic porta-systemic shunt (TIPS):
   trade encephalopathy for ascites
   doesn’t always control, high occlusion rate and encephalopathy like shunt
last resort
umbilical hernia with uncontrolled ascites 30% morbidity, 5% mortality, v 15 and 0 when ascites is controlled

Bleeding

each bleeding admission carries a 25-50% mortality
limit crystalloid, transfuse RBC, FFP, platelets (rarely, only if drops < 50K, mechanism: splenic engorgement and sequestration)
maintain tissue perfusion, monitor urine output, don’t overload (variceal pressure parallels CVP, increases bleeding)
prior use of vasopressin: intense constriction all arterial beds, stopped 80% of bleeds
risk peripheral, myocardial ischemia (NTG ameliorates latter)
somatostatin/octreotide drug of choice, 85% success, less side effects, safer as effective as vasopressin
50mcg bolus, then 50mcg/h X 24h after endoscopy
EGD: 50% of UGI bleeds in cirrhotics are non-variceal
varices source if active bleeding or stigmata and no other source
start somatostatin as soon as possible after accurate Dx, not before scope with encephalopathy protect airway with intubation before scope
Rx: banding treatment of choice v sclerosing, safer occasionally requires TIPS emergency shunt last resort
support after bleeding stops: coag. correction, FFP, vit K, (platelets) hepatitis profile, angio if evaluation for transplant appropriate

Child-Pugh (Gardner numeric score modification)

original classification to assess for portacaval shunt

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th></th>
<th>C</th>
</tr>
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<tbody>
<tr>
<td>Bilirubin</td>
<td>&lt;2</td>
<td>&gt;3</td>
<td></td>
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<tr>
<td>albumin</td>
<td>&gt;3.5</td>
<td>2.8-3.5</td>
<td>&lt;2.8</td>
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<td>ascites</td>
<td>none</td>
<td>controlled</td>
<td>uncontrolled</td>
</tr>
<tr>
<td>encephalopathy</td>
<td>none</td>
<td>minimal</td>
<td>advanced</td>
</tr>
<tr>
<td>nutrition</td>
<td>excellent</td>
<td>good</td>
<td>poor</td>
</tr>
</tbody>
</table>

point score: A=1, B=2, C=3 (15 max)
Class (total score): A 5-6, B 7-9, C 10-15

Hepatic coma

decrease protein to 50g/d, control bleeding, lactulose cathartic (acidifies colon, decreases ammonia absorption)
non-absorbable antibiotics: kanamycin, neomycin
**MELD** (model for end-stage liver disease score)

developed for TIPS but general applicability for liver transplant, more precise than Child Cr, tot bili, INR, etiology of cirrhosis factors in score

**Definitive treatment**

2/3 of bleeders will rebleed, most early, <6w
25% of bleeders will die on that admission
definitive treatment indicated with first bleed
beta block decreases bleeding by decreasing cardiac output
prophylaxis for varices that haven’t bled
endoscopic Rx
  - banding treatment of choice, as effective as sclerosing with fewer complications,
  - lowest incidence of encephalopathy
  - sclerosing: multiple sessions, >60% rebleed, 1/3 fail, 30% complications (ulceration, stricture, perforation, fever, mediastinitis, CNS (embolization of sclerosant)
  - rarely done anymore

surgery
  - surgical treatment is rarely done anymore since the advent of endoscopic treatment, octreotide and TIPS
  - total shunt: 90% prevention of rebleed, 40% encephalopathy
  - selective: eg distal splenorenal, same mortality, effect, decreased encephalopathy
    - Childs class A, B candidates, not C, high mortality
    - low thrombosis (8%, technical error), promotes, does not decrease ascites
    - less rebleeding, encephalopathy than portacaval
  - distal splenorenal does not require splenectomy v central splenorenal
  - partial shunt: short, 8-10mm (larger, 16mm clot) straight supported PTFE
  - non-shunt: esophageal transection, variceal ligation, devascularization (done mostly outside US)

transplant: treatment for liver failure only, not bleeding

**TIPS** (transjugular intrahepatic portacaval shunt)

not indicated for prehepatic causes; polycystic liver or Caroli’s (risk hemorrhage)
surgery superior to TIPS Child class A, B
99% successful technically, 9-50% complications, 3-13% mortality
30-40% encephalopathy (v 10-15% for distal splenorenal)
less rebleeding than Endoscopic banding
33-73% occlusion at 1y, 18% rebleed
  - US monitor every few months, dilate or restent

role: refractory bleeding (after emergency shunt)
  - bridge to transplant
  - Child class C
  - refractory ascites (trade for encephalopathy)

23 December 2008
Splenic Vein thrombosis

due to pancreatitis, cancer
most do not result in varices
if coronary vein joins splenic proximal to the thrombosis, may result in esophageal varices
isolated gastric varices, < 10% bleed
anticoagulation increases risk of bleed
sclerosis and banding not effective for gastric varices
routine EGD not indicated
Rx with splenectomy if bleeding

Portal Vein thrombosis

congenital cavernous transformation, neonatal omphalitis
normal liver function with esophageal varices
mostly extrahepatic portal vein thrombosis in children, liver spared
excellent results with shunt
Endoscopic Rx, distal splenorenal shunt (with normal liver function prevents encephalopathy)

Budd-Chiari (hepatic vein thrombosis)

etiology:
  coagulopathy: polycythemia, myeloproliferative, estrogen, disease, paroxysmal nocturnal hemoglobinuria, factor V Leiden, essential thrombocytosis, antiphospholipid syndr, protein C, S deficiency, antithrombin III deficiency
  IVC occlusion: right atrial myxoma, pericarditis, IVC membrane (most common cause in Asia: thrombolysis, angioplasty)
  liver mass
  hi dose chemotherapy
  (no association with right heart failure or cirrhosis)
presentation
  abdominal pain, ascites, new onset hepatomegaly (liver congested, capsule stretches)
Dx: duplex US, CT, angio.
Rx: depends of status of liver, Bx
  no central necrosis treat symptomatically
  lifelong anticoagulation for hypercoagulable state
  necrosis: do shunt (portacaval, mesoatrial) or transplant (in fulminant hepatic failure)
    75% long term transplant survival
  shunt effective in preserving liver function, controlling ascites, 85% symptomatic relief

50% of patients with portal hypertension and GI stoma develop parastomal varices: local measures, resect stoma