Rethinking Common Labs: Pearls for the Hospitalist (plus some zebras)

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No disclosures

ANION GAP





GOLDMARK - The new MUDPILES...

- Glycols Ethylene/propylene glycol
- <u>O</u>xoproline
- L- lactate
- D- lactate
- Methanol
- Aspirin. Salicylates
- Renal failure
- <u>K</u>etoacidosis- AKA, DKA, starvation ketosis.

Mehta. Lancet, 2008

ANION GAP

60 yo malnourished pt admitted to ICU with sepsis from SBE and spinal abscess. Placed on acetaminophen 1000 mg QID for back pain.

- On Hosp day #4 , AG increases from 8 to 16.
- LFTS/lactate/ serum β-hydroxybutyrate all nl. No urine ketones
- What test to order?

Check 5-oxoproline level (also known as pyroglutamic acid)

• Dx: 5-oxoproline AG metabolic acidosis

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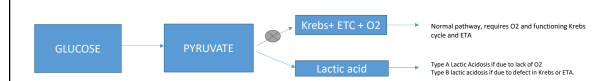
ANION GAP

55 F with extensive bowel resection for SBO 10 y ago. Over past 6 mo family notes episodic confusion after consuming large pasta meals. Pt admitted with confusion.

- AG= 20; lactate= 1 mmol/L (nl)
- What test is next?
- D- Lactate 7 mmol/L (elevated)- send out
- Dx: D- lactic acidosis due to short gut.
- Standard lactate (lactic acidosis) lab is L-Lactate
- SB malabsorption → excess carbs in colon
 → bacteria ferment to D- lactate.
- Rx oral antibiotic

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•32 yo with HIV on AZT. Lactate 12 mmol/L.

ARV's may poison the ETC.

•45 yo DM with AKI with SCr 8 mg/dl. Lactate 13.

On metformin 1000 bid

•23 yo hyperemesis gravidarum. Confusion. 30lb wt loss. Lactate 14. *Thiamine undetectable*

•67 yo metastatic RCC. Lactate 16.

Neoplasm related lactic acidosis.

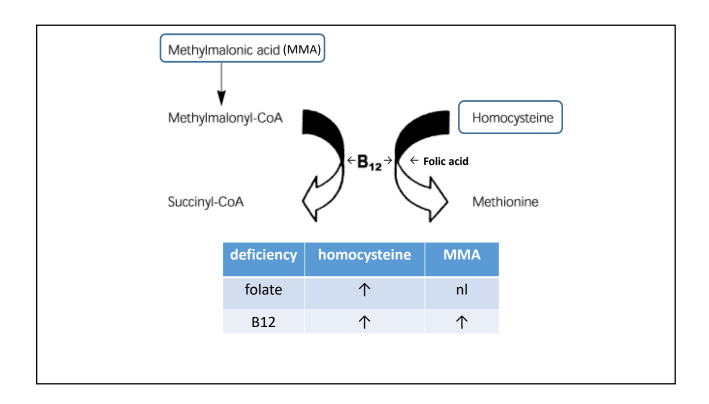
•19 yo runner uses topical analgesic. Confused. Lactate 9.

Salicylate toxicity.

•45 yo rescued from house fire. Lactate 17.

Cyanide toxicity.

B12



B12 deficiency

 72 yo with pancytopenia, AST 200 U/L, ALT 100U/L, LDH 800 U/L, T bili 3.5 mg/dL, indirect 3.0. hyperseg PMNs. MCV 120 fL. Ineffective hematopoiesis(bone marrow hemolysis) Dx? Textbook B12 deficiency

 81 yo with atrophic gastritis. + macrocytic anemia. MCV 118. Intrinsic factor ab + and parietal cell ab +. Dx? Pernicious anemia- 3 tests: intrinsic factor ab, parietal cell ab, gastrin

 22 yo vegan with leg weakness and paresthesias. Abuses "whippits"- nitrous oxide. Dx? Subacute combined degeneration of the cord.

-posterior cord (position and vibration sense)
-lateral corticospinal tract (motor, spasticity)

B12 deficiency

67 yo on metformin. B12 is 167 (ref >190 ng/L)

Metformin causes b12 deficiency

• 23 yo with abd pain, SBO. B12 177 ng/L.

Dx: Crohn's. B12 absorbed in TI

 34 yo with blistering rash, fatigue. B12 150 ng/L, ferritin 8 mcg/L, vit D 25-OH is 6 ng/mL.

Celiac. Rash is DH, check celiac ab panel, tissue transglutaminase IgA. Duodenal biospy.

Anemia

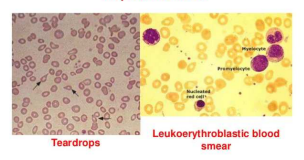
- Anemia DDX: marrow defect, hemolysis or blood loss
- DDX: marrow defect
- Nutritional (B1, B6, folate, iron, B12, malnutrition in general)
- · Decreased erythropoietin (ESRD, inflammation, autoantibody)
- · Endocrine- hypothyroidism, hypogonadism, adrenal insufficiency
- · Sideroblastic anemia- defect hemoglobin synthesis. Ringed sideroblasts on BMBX. + pappenheimer bodies- peripheral smear. MDS, Cu def , ETOH, meds (linezolid)
- · Primary bone marrow problem:
- 1. myelophthisic anemia- due to marrow infiltration. Peripheral smear= "leukoerythroblastic"= nucleated RBC, myelocytes, tear drop RBCs. Causes includes
 primary myelofibrosis or marrow infiltration from tumor (e.g., breast or prostate cancer) or infection (e.g., TB). Autoimmune connective tissue disease can cause
 marrow fibrosis and a myelophthisic anemia.
- 2. MDS, aplastic anemia (pancytopenia from bone marrow failure), pure red cell aplasia
- DDX: Hemolysis
- A. Congenital 1, hemoglobinopathy- sickle cell, thalassemia 2, membrane- hereditary spherocytosis 3, enzyme- G6PD deficiency
- B. Acquired
- . 1. autoimmune- warm AIHA, cold agglutinins, paroxysmal cold hemoglobinuria(PCH). Alloimmune (transfusion reaction, delayed or acute)
- . 2. MAHA- micro/macroangiopathic HA- prosthetic valve malfunction, march, AVM, TMA (MAHA + thrombocytopenia)
- 3. hypersplenism, liver disease (spur cell anemia)
- · 4. infection- malaria, babesia, clostridium perfringens, bartonella
- . 5. copper excess, Wilson's disease, rapid osmotic IVF
- 6. PNH- paroxysmal nocturnal hemoglobinuria the only intrinsic cause of hemolysis that is acquired and not congenital

Anemia

- 53 yo with newly diagnosed widely metastatic prostate cancer presents with fatigue. Found to have Hg 7. Peripheral blood smear reveals nucleated RBCs, myelocytes and tear drop cells. Bone Marrow biopsy reveals diffuse metastatic prostate cancer.
- This is a myelophthisic anemia from diffuse prostate cancer involvement of the marrow.
- The smear is called a leukoerythroblastic smear- immature WBC, nucleated RBC and tear drop RBC.
- Primary myelofibrosis would produce the same leukoerythroblastic smear.

Anemia- Leukoerythroblastic smear

Myelofibrosis



Anemia- Hemolysis

- Hemolysis- suggested by increased reticulocyte, LDH, indirect bilirubin and decreased haptoglobin
- Extravascular v. intravascular- intravascular hemolysis suggested by dark urine (heme pigment, + u/a" blood"), + plasma free hemoglobin and urine hemosiderin
- Peripheral smear- spherocytes (HS, warm AIHA), schistocytes (TMA), bite/blister cells (GGPD), RBC agglutination (cold agglutinin disease), inclusions (infection)
- DDX: Hemolysis
- A. Congenital 1. hemoglobinopathy- sickle cell, thalassemia 2. membrane- hereditary spherocytosis 3. enzyme- G6PD deficiency
- B. Acquired
- 1. autoimmune- warm AIHA, cold agglutinins, paroxysmal cold hemoglobinuria(PCH). Alloimmune (transfusion reaction, delayed or acute)
- 2. MAHA- micro/macroangiopathic HA- prosthetic valve malfunction, march, AVM, TMA (MAHA + thrombocytopenia)
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- 5. copper toxicity, Wilson's disease, rapid hypoosmotic IVF (DW5)
- 6. PNH- paroxysmal nocturnal hemoglobinuria the only intrinsic cause of hemolysis that is acquired and not congenital
- 1. Warm AIHA- + spherocytes, extravascular hemolysis. ddx meds (PCN), autoimmune. Coombs (DAT)+, IgG type. Ab coat RBC and are removed in spleen
- 2. Cold agglutinin disease- + agglutination of RBC. Intravascular hemolysis. False increase MCV. Coombs(DAT)+, C3 type. DDX: lymphoma, infection.
- 3. G6PD deficiency- bite cells on peripheral smear, Heinz bodies on peripheral smear (special stain, not seen on regular peripheral smear), G6PD level decrease
- 4. PNH- episodic intravascular hemolysis. Check peripheral blood flow cytometry for CD55, CD59 (RBC membrane proteins that are absent in PNH)
- 5. Thalassemia- target cells on peripheral smear

Hemolysis

45 yo with neurosyphilis on continuous IV PCN. Presents with fatigue on antibiotic day # 11.

- Hg 6 mg/dl, tbili 3.4 mg/dL (indirect 3.0), LDH 788, retics 30%, haptoglobin low. DAT+, lgG type. Smear shows spherocytes. Dx?
- warm AIHA from PCN

67 yo man with malfunctioning mechanical AoV. Hg 9, Tbili 4 mg/dL (indirect 3.2), LDH 1000 U/L. Smear shows schistos. Dx?

MAHA from AoV

32 yo M presents with fatigue, back pain and dark urine a day after starting dapsone. Hg 6 mg/dL, LDH 1200 U/L, + bite cells on smear. Dipstick urinalysis shows "blood."Dx?

G6PD deficiency- Heinz body+ on special stain as well

Hemolysis

76 yo presents with abd pain and found to have Budd Chiari. Hg 8 mg/dL, LDH 700 U/L, haptoglobin undetectable. Urine hemosiderin is positive confirming intravascular hemolysis. Flow cytometry shows absence of CD55 and CD59. Dx?

PNH: Rx: anti- C5 (eculizumab)

23 yo with sickle presents with sickle pain episode. Retic 40%.

ABG: CO-Hg= 6% (carbon monoxide)

Heme → biliverdin +CO → bilirubin.
 purple → green → yellow.

Increase heme turnover \rightarrow increase CO production.

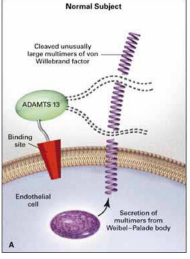
Heme breakdown produces CO

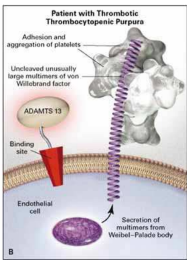
Hemolysis-TMA

- · MAHA- microangiopathic hemolytic anemia. Suggested by hemolysis with schistocytes on peripheral smear
- TMA- thrombotic microangiopathy- suggested by MAHA + thrombocytopenia. TMA pathology is microvascular thrombosis
- TMA leads to microangiopathic hemolysis by shearing of RBC passing through small blood vessels with microthrombosis
- TMA leads to thrombocytopenia by platelet activation and consumption
- TTP- thrombotic thrombocytopenic purpura is prototypical TMA.
- Acquired TTP is an autoimmune disease with antibodies against the ADAMTS13 protein. This causes depletion of ADAMTS13
- ADAMTS13 protein cleaves large VWF proteins attached to endothelial cells. Large VWF proteins not cleaved (because of lack of ADAMTS13) cause microthombosis by aggregation of plts with resultant MAHA and thrombocytopenia.
- Evaluation for TTP: check ADAMTS13 level and ADAMTS13 antibody. With TTP the ADAMTS13 level is usually <5% normal.
- · Treatment:
- · Start PLEX, to remove ADAMTS13 ab and to replenish ADAMTS13 molecule via FFP as replacement fluid. FFP contains ADAMTS13
- Start prednisone and consider rituximab to decrease ADAMTS13 antibody production
- · Consider caplacizumab- a monoclonal antibody that blocks VWF-platelet interaction.
- If there is no immediate access to PLEX, then give FFP prior to transfer to facility that has access to PLEX.
- FFP will provide the ADAMTS13 molecule but will not remove the anti-ADAMTS13 antibody (PLEX required)
- FFP is never a substitute for PLEX, only a temporizing measure.

TTP- thrombotic thrombocytopenic purpura

- MAHA is hemolysis + schistocytes
- TMA =MAHA + thrombocytopenia. TMA is microthrombosis in small vessels.





TTP

32 yo presents with confusion. Exam - petechiae. Hg 10 mg/dL, plts 45K, LDH 600 U/L, indirect bili 3 mg/dL, hapto - zero, retics 14%. periph smear: 12 schistos / HPF. Cr 0.9 mg/dL

TTP suspected...

TTP

Case cont'd

- ADAMTS13 level and α -ADAMTS13 Ab drawn.
- PLEX (plasma exchange) started with FFP used as replacement fluid.
 - Rationale for PLEX (with FFP):
 - FFP replacement fluid provides the ADAMTS13
 - PLEX removes the ADAMTS13 Ab
- ADAMTS13 level returns low at 3% and the α -ADAMTS 13 Ab returns elevated. **TTP dx confirmed.**
- Tx: PLEX and prednisone, +/- rituxan to \downarrow ADAMTS13 ab.
- TTP is the one TMA that usually does not present with AKI.

Primary TMA Syndromes- other than TTP

- 21 yo presents with abd pain and bloody diarrhea.
- Hg 11 mg/dL, plt 62, SCr 4 mg/dL; indirect bili 3 mg/dL, LDH 1100 U/L, haptoglobin- undetectable. + schistos on smear
- ADAMTS13, α-ADAMTS13 drawn and return WNL. Pex started after lab draw Dx?
- Shiga toxin from stool returns positive. PLEX stopped. Supportive care
- ST- HUS (hemolytic uremic syndrome)
- 55 yo drinks tonic H2O daily. Presents with anuric renal failure. Hg 9 mg/dL, plt 32. + schistos. Labs drawn and PLEX started
- ADAMTS13 level is nl and α -ADAMTS13 ab is negative. Dx?
- Drug induced (immune) TMA also known as DITMA from quinine.
- PLEX stopped. Supportive care

Primary TMA Syndromes

- TTP- hereditary and acquired
- HUS (Shiga Toxin TMA)
- DITMA- Drug induced TMA. Immune vs dose dependent
- Complement Mediated TMAinherited or acquired mutation in alternative complement pathway
- Metabolism TMA- inherited mutation in MMACHC gene. Can appear like B12 def. Low plasma MMA and homocyteine
- Coag TMA- inherited mutation in TM, plasminogen, DKGE

VS

TMA mimics

- Also presents with MAHA and thrombocytopenia. Treat underlying cause.
- Pregnancy- HEELP
- Malignant HTN
- DIC
- Neoplasm
- CTD- lupus, APLS, Scleroderma renal crisis
- Stem cell transplant

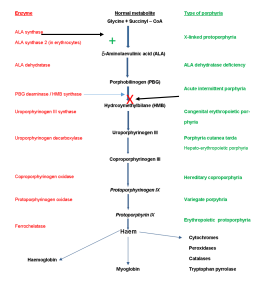
TMA

- 28 yo 39 wk pregnant F presents with oliguric AKI with Scr 5.5 mg/dL, plts 68K, LDH 600, and schistos on peripheral smear. HELLP is considered and fetus is delivered emergently. 1 day after delivery no improvement: Scr 6.7 mg/dL and plts 45K. ADAMTS13 labs drawn. PLEX started. No improvement. ADAMTS13 returns normal.
- Complement medicated TMA. Rx: Eculizumab (anti-complement antibody). High index of suspicion. Early addition of Eculizumab can preserve renal function.

28 F with multiple ER visits for acute N/V/abd pain. Pain is severe, epigastric, colicky, lasting several days and requires opiates. Intermittent lower ext weakness. Labs reveal hyponatremia. Workup unrevealing. Most recent episode followed new diet.

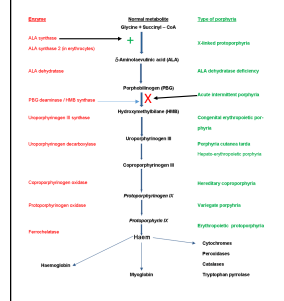
WHAT DISEASE SHOULD BE IN YOUR DDX?

Acute Porphyria- focus on AIP



- Acute porphyria- ALAD, AIP, HCP (+/- blistering rash), VP (+/- blistering rash)
- AIP is acute intermittent porphyria
- Cutaneous porphyria (blistering)- PCT and HEP, CEP, HCP, VP
- Cutaneous porphyria (acute nonblistering)- EPP, XLP
- All 4 acute porphyrias(AIP is prototype) present similarly. <u>Neuropathic pain including abd pain and</u> <u>often weakness and hypona</u>.
- AIP is autosomal dominant with variable penetrance.
- Meds that stimulate the enzyme ALA synthase (ALAS) precipitate attacks of AIP in patients deficient in HMB synthase.

Acute Porphyria



- For dx send spot U porphobilinogen. This test is nearly 100% sn and 100% sp for AIP, HCP and VP. ALAD is extremely rare. 10 case reports only.
- Other tests: urine total porphyrin, urine ALA and plasma and fecal porphyrins.
- Tx: IV D5/D10 until IV hematin available then 4 mg/kg qd for at least 4 d. Inhibits ALA synthase. Stop offending meds that stimulate ALA synthase.
- Pb poisoning mimics acute porphyria. Pb blocks ALA dehydratase. ↑ALA is thought to be the toxic metabolite that causes pain, neuropathy.
- Purple urine is from oxidized porphobilinogen, porphyrins

Acute Porphyria

- 28 F with multiple ER visits for acute N/V/abd pain. Pain is severe, epigastric, colicky, lasting several days and requires opiates. Intermittent lower ext weakness. Labs reveal hyponatremia. Workup unrevealing. Most recent episode followed EtOH ingestion.
- If you could send only 1 test which would it be?
 - a. urine total porphyrins
 - b. 24 hour urine porphobilinogen (PBG)
 - c. spot (random) urine porphobilinogen (PBG) with urine creatinine
 - d. plasma porphyrins

Acute Porphyria

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 - c. spot (random) urine porphobilinogen (PBG) with urine creatinine
 - d. plasma porphyrins

Acute Porphyria- Tests

- Urine porphobilinogen (PBG)- spot or 24 h urine. Spot is best.
- Urine total porphyrins
- Urine ALA
- Plasma Porphyrins
- Fecal Porphyrins
- For AIP: blood erythrocyte PBG deaminase- low in 90% pts with AIP. Helpful to diagnosis AIP if asymptomatic.
- The additional tests help differentiate between AIP, VP and HCP.

Porphyria

- · Acute Porphyria- presents with abdominal pain, neuropathy
- · Acute Porphyria- AIP, HCP, VP, ALAD
- · Cutaneous Porphyria- chronic blistering or acute non- blistering
- · To diagnose Acute Porphyria (e.g., AIP)- check spot urine porphobilinogen (PBG) with creatinine
- Urine PBG is highly sensitive and specific for AIP if obtained when patient is symptomatic
- If patient is symptomatic (abdominal pain, neuropathy) and urine PBG is normal then diagnosis of acute porphyria unlikely.
- If patient is symptomatic and urine PBG is elevated then patient has acute porphyria. AIP is most likely. Urine porphyrins can differentiate HCP from VP.
- AIP is autosomal dominant. Only has ~10% penetrance.

Acute Porphyria –Treatment.

- · AIP is prototype
- Review patient medications for those that can precipitate porphyria attacks.
- Infuse D10W at ~100 ml/hr as temporizing measure while awaiting Hematin
- Start Hematin 3-4 mg/kg IV qd. Usually will require at least 4 doses
- Stop D10 after Hematin started.
- Consider givosiran as outpatient to reduce frequency of attacks

Eosinophilia

- Eosinophilia- defined as absolute eosinophil count (AEC) ~>300 cells/ mcl
- · Hypereosinophilia (HE)- defined as:
 - 1. AEC > 1500 on 2 occasions separated by > 1 month

and/or

- 2. evidence of tissue hypereosinophilia (e.g., bone marrow with >20% eos)
- Hypereosinophilic Syndrome (HES)- is HE + eosinophil mediated tissue damage
- · Primary HES- underlying clonal neoplasm
- Secondary HES- any cause of HE with tissue damage. For example, Loeffler's myocarditis from Strongyloides infection.

Eosinophilia- etiology

- NAACP
- Neoplasm- Primary hypereosinophlic syndrome, lymphoma, leukemia
- Addison, AI, asthma, allergies
- CVD- Churg-Strauss (CSS) is eGPA(eosinophilic granulomatosis with polyangiitis)
- Parasites- strongyloides, toxocara, schistosomiasis, filariasis, hookworm, trichinella
- Other- APBA, cholesterol emboli, eosinophilia myalgia syndrome, DRESS
- Other- GI dzs (e.g, eosinophilic esophagitis) Skin (E.fasciitis)
- 35 yo with asthma with foot drop, dark urine. Peripheral eos= 35%. ANCA+. Dx?

eGPA

• 23 yo with CF with migrating infiltrates. Eos- 18%. IgE 1050 kU/L. Dx? Tx?

APBA. Pred and voriconazole

• 56 yo on pre-op testing for renal transplant has unexplained eosinophilia of 12%. Must check strongyloides ab. If positive then treat with ivermectin.

Eosinophilia

- NAACP
- Neoplasm- hypereosinophlic syndrome, lymphoma, leukemia
- Addison, AI, asthma, allergies
- CVD- Churg-Strauss (CSS) is EGPA(eosinophilic granulomatosis with polyangiitis)
- Parasites- strongyloides, toxocara, schistosomiasis, filariasis, hookworm, trichinella
- Other- APBA, cholesterol emboli, eosinophilia myalgia syndrome, DRESS
- Other- GI dzs (e.g, eosinophilic esophagitis) Skin (E.fasciitis)
- Immune suppression (e.g. prednisone) in pt with occult strongyloides can lead to strongyloides hyperInfxn syndrome and death.
- 24 yo with malaise, nausea, weight loss and hyperpigmentation. CBC shows eos 16%. Am cortisol = 1 mcg/dL. Dx?

Adrenal Insufficiency

67 yo s/p LHC (cardiac cath) last wk. Has ARF, livedo reticularis, and peripheral cholesterol emboli causing AKI

LFT's- Hepatocellular vs Cholestatic pattern

Which set is most elevated -- AST/ALT or alk phos?

- If alk phos, then DDX is cholestatic DDX:
 - Anatomic obstruction -CBD stone, sludge, stricture. Cholangiocarcinoma, ampullary mass (pancreatic head mass)
 - Autoimmune PBC, PSC
 - Meds: some meds more likely to present with this pattern (e.g., amoxicillin)
 - Infiltrationamyloid, granulomatous(sarcoid, Infxn- TB, (fungal-Crypto, histo)), tumor, mass lesion
 - Right HF

LFT's- Hepatocellular vs Cholestatic pattern

43 yo with UC with abd pain. Tbili 2.1 mg/dL, AP 350 U/L, AST 88 U/L, ALT 90 U/L. ERCP and MRCP show ductal dilation and beaded appearance. Dx?

Primary Sclerosing Cholangits (PSC)

55 F with Hashimoto's thyroiditis presents with fatigue, pruritis. Tbili 1.7 mg/dL, AP 500 U/L, AST/ALT- 55 U/L. MRCP nl. AMA+. Dx?

Primary Biliary Cirrhosis (PBC)

Which set is most elevated -- AST/ALT or alk phos?

- If alk phos, then DDX is cholestatic DDX:
 - Anatomic obstruction -CBD stone, sludge, stricture. Cholangiocarcinoma, ampullary mass (pancreatic head mass)
 - Autoimmune PBC, PSC
 - Meds: some meds more likely to present with this pattern (e.g., amoxicillin)
 - Infiltrationamyloid, granulomatous(sarcoid, Infxn- TB, (fungal-Crypto, histo)), tumor, mass lesion
 - Right HF

LFT's- Hepatocellular vs Cholestatic pattern

62 yo presents with HIV, CD4 45 presents with pancytopenia, AP 600 U/L, Tbili 2 mg/dL, ALT/AST 77 U/L. Dx?

liver bx shows granulomas and histoplasma

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LFT's- Hepatocellular vs Cholestatic pattern

45 yo on amoxicillin for 1 wk presents with n.v. abd pain. Tbili 2.1 mg/dL, alt/ast 100 U/L, alk phos 550 U/L.

Medication rxn

Which set is most elevated -- AST/ALT or alk phos?

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 - Right HF

LFT's- Hepatocellular vs Cholestatic pattern

55 yo presents with subacute RUQ pain. LFTs- nl except for elevated alk phos 175 U/L.

CT shows liver abscess.

Which set is most elevated -- AST/ALT or alk phos?

- If alk phos, then DDX is cholestatic DDX:
 - Anatomic obstruction -CBD stone, sludge, stricture. Cholangiocarcinoma, ampullary mass (pancreatic head mass)
 - Autoimmune PBC, PSC
 - Meds: some meds more likely to present with this pattern (e.g., amoxicillin)
 - Infiltrationamyloid, granulomatous(sarcoid, Infxn- TB, (fungal-Crypto, histo)), tumor, mass lesion
 - Right HF

Cirrhosis

VS.

liver failure

- ETOH
- · Hepatitis B, C
- NASH
- Hemochromatosis, Alpha-1 AT, Wilson's, CF
- Autoimmune hepatitis, PBC, PSC, celiac, sarcoid
- Right CHF, Budd Chiari
- Meds: MTX, amiodarone
- · Secondary biliary cirrhosis
- Rare:
- idiopathic ductopenia
 - veno-occlusive dz(VOD) " small vessel Budd Chiari". s/p BMT
- HHT

- Acute hepatitis A,B,C,D,E
- EBV, CMV, adeno, HSV, VZV
- Leptospirosis
- Vascular- shock liver, right CHF, Budd Chiari
- · Wilson's
- · AIH, celiac
- Toxins- ETOH, Tylenol, amanita, CCl4
- Meds: idiosyncratic usually
- Illicit Drugs- cocaine, methamphetamine
- Tumor infiltration- SCC, breast, lymphoma, melanoma
- Pregnancy: HELLP, acute fatty liver of pregnancy
- Other: HLH, VOD, heat stroke

Liver Cases:

• 45 yo N/V/abd pain. WBC 22K.

Tbili 17 mg/dL, AP 335, AST 290 U/L, ALT 120 U/L. Dx?

ETOH hepatitis

• 32 yo started PTU 4 wks ago, now confused,

liver failure

med

INR 4, Tbili 10 mg/dL, AST/AST 500 U/L. Dx?

• 19 yo h/o tremor presents in liver failure.

Tbili 17 mg/dL, INR 3, AP low at 28. Haptoglobin undetectable c/w hemolysis.

Wilson's

Serum ceruloplasmin low at 7. 24 h urine copper elevated.

Liver bx shows ↑copper. Eye exam: pigmented outer cornea. Dx?

• 65 F with acute liver failure. Hb 17 mg/dL.

Erythematous painful distal extremities (erythromelalgia) x 6 mo. Epo level suppressed. Doppler - acute thrombus in hepatic veins and IVC. JAK-2 positive. Dx? related

P. Vera

Budd Chiari

Liver Cases (cont)

 34 yo F with Tbili 17 mg/dL, alk phos 222 U/L, ALT 778 U/L, AST 700 U/L. ANA+, total IgG elevated, ASMA+, f-actin+. Liver bx confirms dx. Dx?

AIH

• 23 yo with liver failure. + conjunctival suffusion, stiff neck, ARF. Recent honeymoon in Hawaii. Walked barefoot in stream. Dx?

Leptospirosis

• 34 yo with liver failure. Exam shows numerous vesicles and erosion on labia. What med should be started ASAP?

IV acyclovir

• 45 yo with h/o IVDA presents with abd pain, n,v, hepatitis. Hep C ab negative, hep C PCR 3,000,000. Dx?

Acute Hep C

Liver Cases (cont)

- 45 yo lifelong nonsmoker dx with emphysema 5 yrs ago presents with abnl LFTs.
- Has ZZ genotype, α -1 antitrypsin level is 12 (nl >80)

Exam - displaced PMI, dark skin. Testosterone low.

α1 AT deficiency - Has h/o tender red lesions on shins (e.nodosum). Dx?

- 42 yo M scientist presents with low libido. AST 60 U/L, ALT 90 U/L.
 - What labs would you send? You make a dx but he defers tx. 3 yrs later he dies of the plague from working with a modified Yersinia. pestis in his lab. Dx?

Ferritin, Fe/TIBC, genetic tests hemochromatosis

Coagulopathy

Coagulopathy

DDX: elevated INR

- cirrhosis/liver failure
- warfarin
- vitamin K deficiency- malabsorption, decrease po; decreased colonic bacteria (s/p antibiotic)
- DIC

Dx?

Dx?

Dx?

- Liver produces vit K dependent factors 2,7,9,10
- Liver produces vit K independent factor 5
- Endothelial cells make factor 8

Coagulopathy - liver failure, Vit k deficiency, DIC?

INR 3	Factor 2 (liver:vit K dep)	Factor 7 (liver: vit K dep)	Factor 5 (liver:vit K indep)	Factor 8 (endothelial)
Cirrhosis/liver failure	\	\	\	nl/↑
Vit K deficiency	\downarrow	\downarrow	nl	nl/↑
DIC	\downarrow	\downarrow	\downarrow	\downarrow

CASE	Factor 2 (> 80%)	Factor 7 (> 80%)	Factor 5 (>80%)	Factor 8 (>80%)	
32 yo acetamin OD, INR 4.	12% (↓)	16% (↓)	19% (↓)	120%	Liver failure
45 yo severe malnut. INR 2.5	20% (↓)	22% (↓)	93%	102%	Vitamin K def
54 yo admitted with sepsis. INR 3	14% (↓)	18% (↓)	14% (↓)	19% (↓)	DIC

Hepatitis B labs

Labs:

- hepatitis B surface antigen (hep B s ag)
- hepatitis B core antibody- (hep B c ab) IgM or IgG
- hepatitis B surface antibody (hep B s ab)

Cases:

- s Ab pos, c ab neg, s Ag neg:
- s Ab pos, c ab lgG pos, s Ag neg
- s Ab neg, c ab IgG pos, s Ag pos
- s Ab neg, c ab IgM pos, sAg pos
- Isolated c ab **pos** 4 possibilities
- - remote hep B Infxn- cleared. Now immune. Hep B s ab has waned below level of detection

· Dx: vaccine

• Dx: immune • Dx: chronic

• Dx: acute

- chronic hep B Infxn- infected remotely. Hep B s ag has waned below level of detection.
- acute hep B- in window. Hep B c ab would be IgM type.

