Evaluation of patients with elevated liver function tests (LFT)



ESIM 16, Saas-Fee, 2012 Stefan Lindgren, professor, M.D., Ph.D., FACP, FRCP, FEFIM (hon) Lund university, University Hospital Skane, Malmö, Sweden

How extensive should you be?

Elevated LFT in a 62 year old woman with arthrosis/osteoporosis, type 2 diabetes, hypertension and hyperlipidemia.

Extensive medication list.

Overweight. No liver stigmata.

Focused investigation?

28-year old woman with slight, pressingdiscomfort in her right upper abdomen.Previously healthy. On oral contraception since3 years.

Bil 22, ASAT 0.8, ALAT 1.0, GT 1.2, ALP 2.6, INR 1.0

When should we use imaging?

1. Is the entire liver affected?

2. Are focal changes present?

3. Are the bile-ducts affected?

Liver imaging methods

Ultrasound MR incl. MRCP ERC PTC CT

Which method should you choose?





Focal changes

Ultra sound



Echogenicity Penetrance

Contrast enhanced ultra sound in a woman with cirrhosis



CT and MR



MRC, primary sclerosing cholangitis



Which diseases are most likely in patients with elevated LFT?

Parenchyma Only transaminases and GGT Fatty liver alcohol obesity Chronic viral hepatitis Haemochromatosis AAT-deficiency Autoimmune hepatitis Drug induced

Transaminases and GGT + ALP PBC

Drug induced

Focal

Transaminases and ALP/GGT Focal nodular hyperplasia Adenoma HCC

Metastases (Hemangioma)

Bile ducts

<u>Transaminases and</u> <u>ALP/GGT</u> Bile duct stone Cholangiocarcinoma

Pancreatic cancer

PSC

Diagnosis of chronic liver disease

- Elevated LFT (80%)
- Spiders, palmar erythema
- Icterus
- Decompensation (oedema, ascites, encefalopathi, variceal bleeding, infection)
- General symptoms (pain, cholangitis, itching, arthralgia, fatigue, anorexia)







Clinical evaluation of a patient with suspected/known chronic liver disease

- Etiology?
- Prognosis (cirrhosis? portal ht?)
- Acute and long-term management

Accidentally detected elevated LFT

Normal range: Mean ± 2 SD i.e. 2.5% of healthy individuals have some elevated LFT.

20 tests – 65% have at least one elevated test!

Intra-individual variation

Retesting of accidentally detected elevated LFT within 3 weeks will give normal results in 30% of patients.

(Lazo M, Ann Intern Med 2008)

How common?

Elevated ALT in 99/19877 recruits to US Air Force. Specific explanation in 12/99

(Kundrotas, LW Dig Dis Sci 1993)

249 blood donors with elevated ALT - alcohol 11-48%, steathosis 22-56%, HCV 17-20%, diverse 4-8%, no specific diagnosis 2-4%.

(Hultcrantz R, Scand J Gastroenterol 1986) (Katkow WN, Ann Intern Med 1991)

Contribution of liver biopsy to diagnosis

354 patients with elevated LFT (ALT, GGT and/or ALP) >6 months, after exclusion of patients with clinical or serological evidence of liver disease

• Steathosis 66%

• Lever biopsy contributed to clinical decisions in 18%

(Shelly MM, J Hepatol 2001)

Conclusion

- Etiological diagnosis is usually possible without biopsy.
- Most patients with unclear diagnosis after careful history, physical examination and analysis of biochemical and serological tests have alcoholic liver disease or steathosis.

Clinical management

- Hepatocellular pattern pre-dominant (ALT, AST)
- Cholestatic pre-dominance (GGT, ALP)

Principles

- Consider retesting once within 3 weeks.
- Avoid further extended check-ups. Clinical decision!
- Keep extra-hepatic explanations in mind.

Medical history

- Drugs!!
- Contact with blood
- Other known diseases
- Alcohol
- Specific symptoms

Other diseases

- Cardiovascular
- Pulmonary
- Inflammatory systemic disease
- Thyroid disease
- Myositis (AST, CK)
- Malabsorption, coeliac disease
- Metabolic syndrome
- Addison

Physical examination

Low sensitivity!

- Spiders, palmar erythema?
- Signs of extrahepatic disorders? (jugolar veins, BP, atrial fibrillation, joints, skin, thyroid dysfunction)
- Oedema, ascites?
- Hepatomegali, splenomegali?

Isolated elevation of bilirubin

- Hemolysis
- Defect conjugation

 unconjugated
 Gilbert (3-7%)
 (Crigler Najjar typ 2)

 UDP glucoronyl transferase

 (conjugated: Dubin-Johnson, Rotor)



Other isolated biochemical findings

- GGT usually not liver/bile ducts (drugs, alcohol, obesity)
- ALP bone, growing teen-agers, metastases, osteomalacia, Paget.
- Always interpret GGT and ALP together!
- ALP isoenzymes

Phosphatidylethanol in blood (HPLC)

- Estimates mean alcohol consumption during 2 weeks
- An abnormal phospholipid generated in cellmembranes *only* by ethanol
- Specificity as a marker of alcohol consumption 100%
- No false positive results
- Correlates with amounts of alcohol consumed over >7 days

Cholestatic predominance

- Must always be investigated!
- Intrahepatic/extrahepatic
- Ultrasound, (CT), MRCP, ERCP
- PBC AMA, IgM
- PSC IBD? MRCP
- Drugs

Hepatocellular predominance

Chronic viral hepatitis HCV – antibodies, RNA HBV – HBsAg, anti-HBsAg, Hb_cAg e-antigen, DNA

Hepatocellular predominance 2

Autoimmune hepatitis – IgG, ANA, SMA (LKM, ANCA, SLA)

Note! 20% will not be ANA or SMA positive

Hepatocellular predominance 3

Steathosis – ALT (GGT)

NASH – increasing AST with fibrosis

Hepatocellular predominance – metabolic liver disease

- AAT-deficiency Upper middle age Plasma protein analysis Isoelectric focusing
- Wilson

Unusual! 5-25 years, up to 40 years Low ceruloplasmin in 85% tU-copper Mutation analysis





Haemochromatosis

- Manifest liver disease in middle age elderly Heterozygotes in 10% of the population – no disease
- HFE homozygotes (C282Y) 0.5%
- Compound heterozygotes
- Low penetrance
- Transferrin saturation >50% (>45% in females)
- Ferritin (acute phase reactant! alcohol!)
- Ferritin <1000 no fibrosis

Haemchromatosis with fibrosis



Summary

Bilirubin	ANA, SMA, AMA
AST	HCV-antibodies
ALT	HBsAg
GGT	TSAT
ALP	Plasma protein analysis (AAT, ceruloplasmin, Ig)
PK/INR	

Always!!

Cardiovascular causes of liver disease

Acute, less often chronic liver disease

- Ischemic hepatitis (forward failure)
- Right heart failure
- Budd-Chiari
- Portal thrombosis
- Occlusion of a. hepatica

When should we perform a liver biopsy?

Etiology? Fibrosis! Prognosis? Indication for treatment?



Liver stiffness, transient Elastography

FIBROSCAN

To acquisition system





FIG. 3. Liver stiffness values for each fibrosis stage. The

How important is it to separate NASH from steathosis?

History/Physical exam/LFT Biopsy

Fatty liver/NASH



