# Alcohol and the brain

Prof. Hanan Hagar

Pharmacology Unit

College of Medicine

KSU

# Ethyl alcohol (ethanol)

Ethyl alcohol (ethanol) is the most commonly abused drug in the world.

### **Pharmacokinetics**

- is a small <u>lipophilic</u> molecule
- readily crosses all biological membranes
- Rapidly & completely absorbed from GIT
- Has large Vd (distributed to all body tissues)
- Volume of distribution = Total body water (0.5-0.7 L/kg).
- Crosses placenta and excreted in milk

### Metabolism of ethanol

Metabolism in gastric mucosa & liver.

- Oxidation of ethanol to acetaldehyde via <u>alcohol dehydrogenase</u> or CYP-450 (CYP-2E1).
- Oxidation of acetaldehyde to acetate via <u>aldehyde dehydrogenase</u> which also reduces NAD+ to NADH.
- Acetate ultimately is converted to CO<sub>2</sub> + water.

### Metabolism of ethanol

 At low ethanol conc. minor metabolism by microsomal ethanol-oxidizing system (MEOS) mainly CYP-450 (CYP2E1).

 Upon continuous alcohol use, this enzyme is stimulated and contribute significantly to alcohol metabolism & tolerance.

# Alcohol Metabolism (the major pathway) 90-98% in the liver

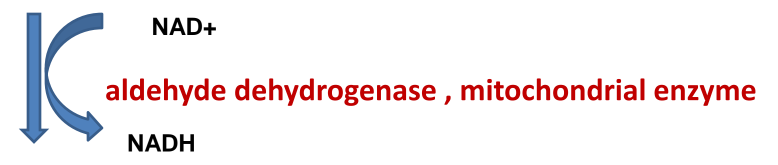
CH<sub>3</sub>CH<sub>2</sub>OH (Ethanol)

NAD+

Alcohol dehydrogenase , cytosolic enzyme

NADH

CH<sub>3</sub>CHO (Acetaldehyde) more toxic than alcohol

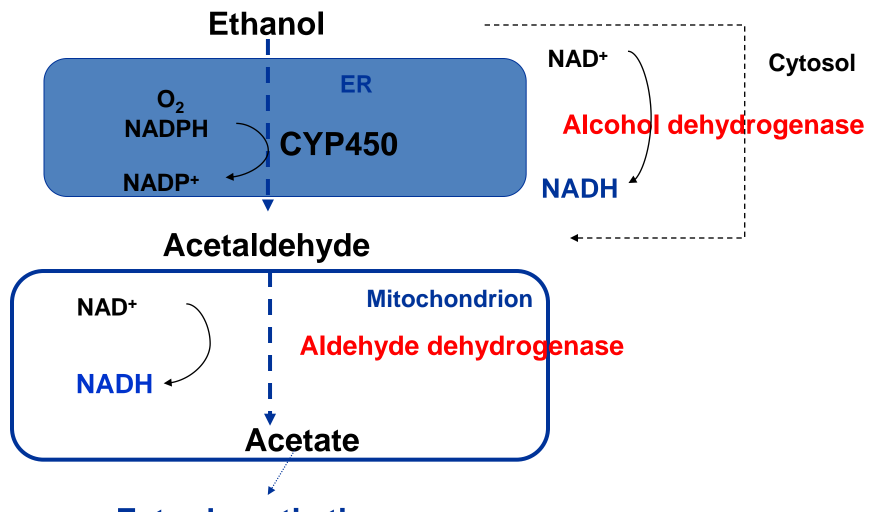


**CH3COOH** (Acetic acid)



CO2 +water

## Hepatic Cellular Processing of alcohol



**Extra-hepatic tissue** 

NAD+/NADH: nicotinamide adenine dinucleotide

### Metabolism of ethanol

• Acute alcohol consumption inhibits CYP450 2E1 so decrease metabolism of other drugs taken concurrently as (warfarin, phenytoin).

• Chronic alcohol consumption induces liver microsomal enzymes CYP450 2E1, which leads to significant increases in ethanol metabolism (Tolerance) & metabolism of other drugs as warfarin taken with it (Drug interactions).

### Genetic variation of alcohol metabolism

### Aldehyde Dehydrogenase Polymorphism

- Asian populations (including Chinese, Japanese, Taiwanese, Korean) have genetic variation in aldehyde dehydrogenase resulting in a variant allele.
- The rate of metabolism of alcohol is slower than other populations.
- Can develop "Acute acetaldehyde toxicity" after alcohol intake characterized by nausea, vomiting, dizziness, vasodilatation, headache, and facial flushing and prevent them from becoming alcoholic.

### **Alcohol excretion**

- Excreted unchanged in urine (2-8%).
- Excretion unchanged via lung (basis for breath alcohol test).
- Rate of elimination is
  - Zero-order kinetic.
  - Is not concentration-dependent.
  - i.e. rate of elimination is the same at low and high concentration.

### Mechanism of action of alcohol

- It is a CNS depressants
- Acute alcohol causes:
  - Enhancement the effect of GABA (inhibitory neurotransmitter) on its GABA receptors in brain leading to CNS depression
  - Inhibition of glutamate action (excitatory neurotransmitter) on NMDA receptors leading to disruption in memory, consciousness, alertness.

#### Chronic alcohol leads to

up-regulation of NMDA receptors & voltage sensitive Ca channels (Ca influx to nerve cells) leading to alcohol tolerance & withdrawal symptoms (tremors, exaggerated response & seizures).

#### **Acute actions of alcohol**

#### In mild-moderate amounts

### **CNS** depression

- relieves anxiety, euphoria (feeling of well-being).
- Nystagmus, slurred speech, impaired judgment, ataxia
- Sedation, hypnosis, loss of consciousness.

### **CVS** depression

- Myocardial contractility depression
- Vasodilatation due to:
  - ✓ Vasomotor center depression
  - ✓ Direct smooth muscle relaxation caused by acetaldehyde

### **Acute actions of ethanol:**

#### In severe amounts

- Severe CNS depression
- Respiratory depression.
- Respiratory acidosis
- Nausea, vomiting, aspiration of vomitus.
- CVS depression
- Volume depletion
- Hypotension
- Hypothermia
- Coma, death.

# Chronic ethanol abuse (alcoholism) is associated with many complications

- Tolerance, dependence, addiction, behavioral changes
- Liver: hepatic cirrhosis & liver failure.
- **CVS:** hypertension, myocardial infarction
- **CNS:** cerebral atrophy, cerebellar degeneration, and peripheral neuropathy. Wernicke encephalopathy or Korsakoff psychosis may occur.
- GIT system: irritation, inflammation, bleeding, nutritional deficiencies
- Endocrine system: gynecomastia & testicular atrophy
- Hematological disorders, neoplasia.

# Chronic alcohol use (Alcoholism)

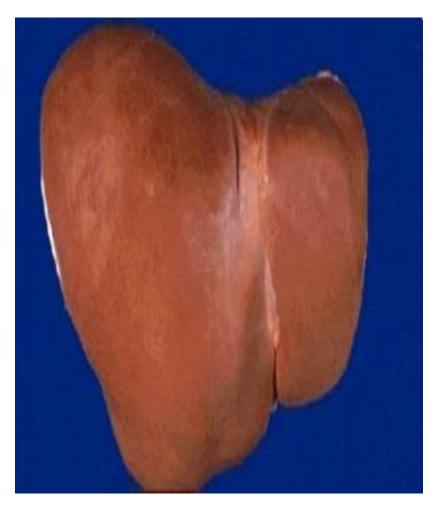
### Liver

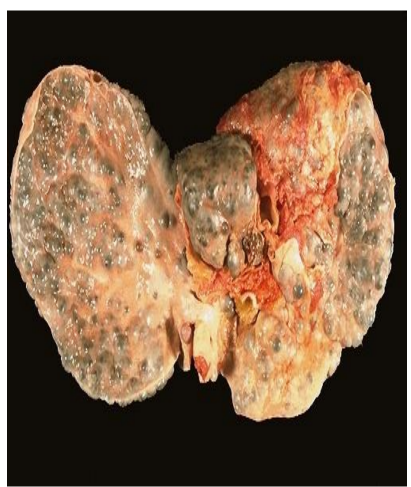
# The most common medical complication of alcoholism occurs with liver

- Reduction of gluconeogenesis
- Fatty liver/ alcoholic steatosis
- Hepatitis
- **Hepatic cirrhosis**: jaundice, ascites, bleeding, encephalopathy.
- Irreversible liver failure.

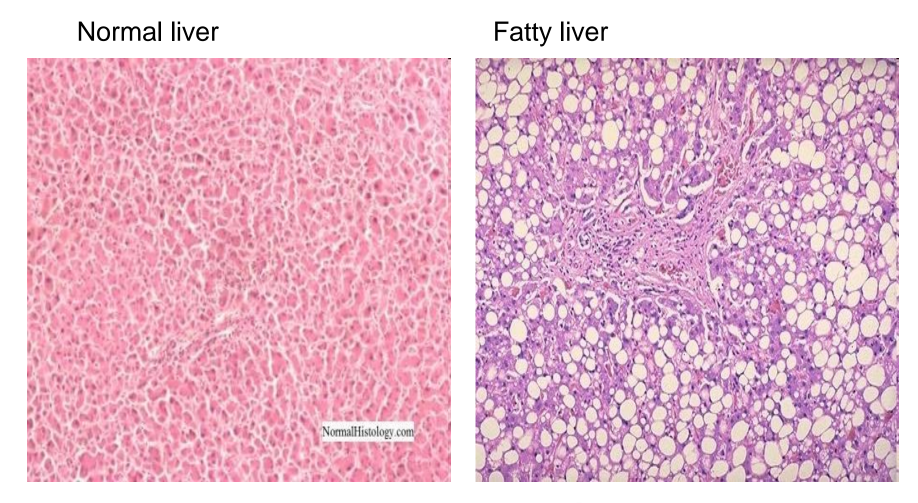
# **Healthy Liver**

# Liver in chronic alcoholics



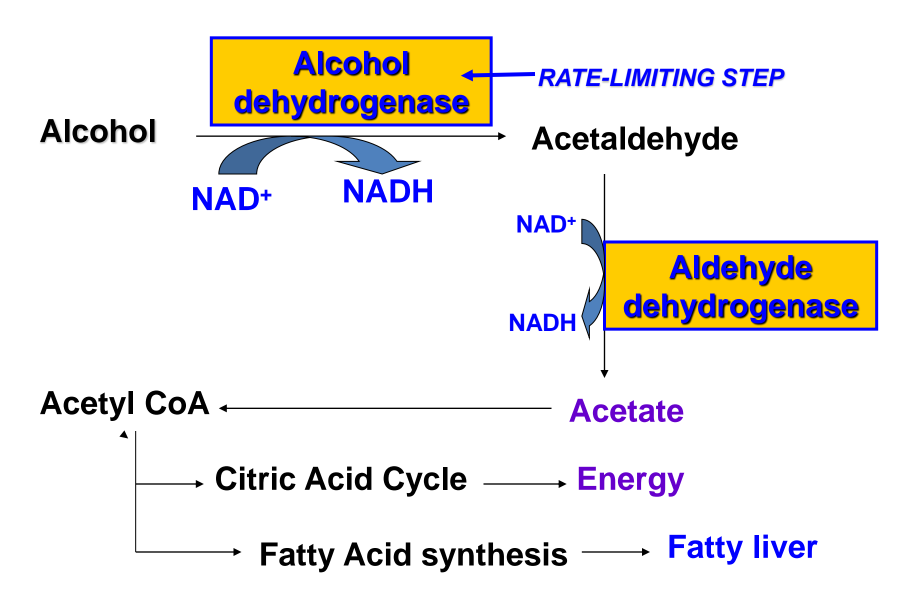


# Healthy Liver vs Fatty Liver



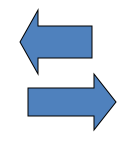
Acetaldehyde is more toxic than alcohol → causing inflammation and fat cell proliferation

# **Hepatic Ethanol Metabolism**

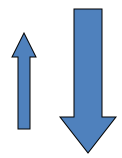


# Alcoholic Liver Disease

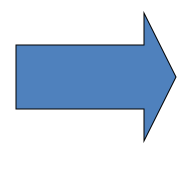
Steatosis



Normal



Steatohepatitis



Cirrhosis



# Gastrointestinal system

- Gastritis, hemorrhagic esopahgitis, ulcer diseases, pancreatitis (due to direct toxic action on epithelium).
- Diarrhea
- Deficiency of vitamins.
- Exacerbates nutritional deficiencies
- weight loss, and malnutrition
- In heavy drinkers: increased risk of oral and esophageal cancer.

# Cardiovascular System

Chronic alcohol abuse can lead to cardiomyopathy

- Cardiac hypertrophy
- Congestive heart failure.
- **Arrhythmia** (due to potassium and magnesium depletion)
- **Hypertension**: due to increased calcium & sympathetic activity.

## Hematological complications:

- Iron deficiency anemia (due to inadequate dietary intake & GIT blood loss).
- **Megaloblastic anemia**: (due to folate deficiency, malnutrition, impaired folate absorption).
- Hemolytic anemia.
- Bone marrow suppression
- Thrombocytopenia (suppressing platelet formation, prolong bleeding times).
- Impaired production of vitamin-K dependent clotting factors leading to prolonged prothrombin time.

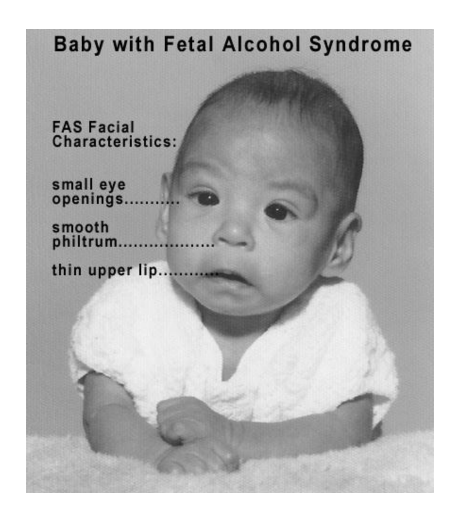
22

## Fetal Alcohol Syndrome: Irreversible

- Ethanol rapidly crosses placenta
- Pre-natal exposure to alcohol causes:
- Intrauterine growth retardation (due to hypoxia)
- Congenital malformation (teratogenesis):
  - Microcephaly
  - Impaired facial development
  - Congenital heart defects
  - Physical and mental retardation.



# Fetal Alcohol Syndrome (FAS)





The right brain is an infant's brain that had pre-natal exposure to alcohol.

## **Endocrine system:**

## > Hypogonadism:

In women: ovarian dysfunction, amenorrhea, anovulation, hyperprolactinemia, infertility.

In men: gynecomastia, decreased muscle & bone mass, testicular atrophy, sexual impotence due to inhibition of luteinizing hormone (LH), decrease in testosterone, estradiol, progesterone.

➤ Hypoglycemia & ketoacidosis due to impaired hepatic gluconeogenesis & excessive lipolytic factors, especially increased cortisol and growth hormone.

25

## **Central Nervous System**

- Tolerance
- Physiological and psychological dependence
- Addiction: dopamine, serotonin and opioids are involved.
- Neurologic disturbances
- Wernicke-Korsakoff syndrome

### Wernicke-Korsakoff syndrome

It is a combined manifestation of **two** disorders:

Wernicke's encephalopathy: characterized by

- ocular disturbances unsteady gait
- changes in mental state as confusion, delirium, ataxia

Korsakoff's psychosis: impaired memory & cognitive and behavioral dysfunction.

### Cause: thiamine (vitamin B1) deficiency due to:

- ☐ inadequate nutritional intake
- decreased uptake of thiamine from GIT
- ☐ decreased liver thiamine stores

Treated by: thiamine + dextrose-containing IV fluids. 27

### **Alcoholism Tolerance**

Chronic consumption of alcohol leads to tolerance That develops due to:

Metabolic tolerance (pharmacokinetic): due to induction of liver microsomal enzymes.

Functional tolerance (Pharmacodynamic): due to change in CNS sensitivity.

## Alcoholism withdrawal symptoms

- Autonomic hyperactivity & craving for alcohol
- Vomiting, thirst
- Profuse sweating, severe tachycardia
- Vasodilatation, fever
- Delirium, anxiety, insomnia, agitation, tremors.
- transient visual/ auditory illusions, violent behavior, hallucinations.
- Grand mal **seizures** (after 7-48 hr alcohol cessation)
- Due to super-sensitivity of glutamate receptors & hypoactivity of GABA receptors are possibly involved.

## Management of alcoholism withdrawal

- Substituting alcohol with a long-acting sedative/hypnotic drug then tapering the dose.
- **Benzodiazepines** as (chlordiazepoxide, diazepam) or lorazepam that is preferable (shorter duration of action).
- Efficacy: IV/ po
- Manage withdrawal symptoms & prevent irritability, insomnia, agitation & seizures.
- Dose of BDZs should be carefully adjusted to provide efficacy & avoid excessive dose that causes respiratory depression & hypotension.

### Management of alcoholism withdrawal

- Fluoxetine
- Clonidine & Propranolol: inhibits the action of exaggerated sympathetic activity.
- **Acamprosate:** a weak NMDA receptor antagonist & GABA activator, reduce psychic craving.

## To prevent alcohol relapse:

### **Disulfiram therapy**: 250 mg daily

- Disulfiram-induced symptoms that render alcoholics afraid from alcohol drinking.
- Inhibits hepatic aldehyde dehydrogenase, this will increase blood level of acetaldehyde.
- Acetaldehyde produces extreme discomfort, vomiting, diarrhea, flushing, hotness, cyanosis, tachycardia, dyspnea, palpitations & headache.

## Alcohol and drug interactions

- Acute alcohol use causes inhibition of liver microsomal enzymes thus decreases metabolism of some drugs and increases their toxicities e.g. bleeding with warfarin
- Chronic alcohol use induces liver microsomal enzymes and increases metabolism of drugs such as warfarin, propranolol and etc.

• Acetaminophen + alcohol (chronic use): risk of hepatotoxicity due to increased production of free radical metabolite of acetaminophen.

- **NSAIDs** + **alcohol:** Increase in the risk of developing a major GIT bleeding or ulcers.
- Narcotic drugs (codeine and methadone) + alcohol: risk of respiratory and CNS depression.
- Alcohol suppresses gluconeogenesis, which may increase risk for hypoglycemia in diabetic patients.