



Pharmacology

Subject :

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Hala AL Beshtawe

وَقُلْ رَبِّ زِدْنِي عِلْمًا

General Pharmacology

Medication use during pregnancy Clinical Toxicology

Faculty of Medicine
The Hashemite University
Arwa Al Anber(MD, PhD)
Office :1018



The **Thalidomide Disaster** كان يستعمل لتقليل اعراض الاستغراب النفس للمرأة الحامل

On October 1, 1957, Chemie Grünenthal GmbH marketed Thalidomide in West Germany. It was originally used as a sleeping tablet and **was considered safe for pregnant women**. It **later relieved morning sickness** in pregnant women. Thousands of women used it, and it appeared to work until the following year, when thousands of kids were born with **birth defects**. They varied from **brain damage** to **limb deformities**.



Medication use during pregnancy

- Drug therapy in pregnancy may cause anxiety to both providers and patients due to uncertainty regarding dosing and safety.
- The needs for medications are high as the incidence of maternal chronic illnesses such as hypertension or diabetes are rising.
- More than 40% of pregnancies are not planned. To prevent the use of unsafe medications in (early) pregnancy, Women with chronic conditions or recurrent episodes of illness should be treated with pregnancy-compatible drugs, even if no pregnancy is planned.

* حالات اميتر لاستعمال ادوية عند امراض الحامل ومن الامراض المزمنة
* معظم الكوارث الدوائية تحدث عندما تكون المرأة لاتعلم بحملها او لم يتحلا له مما يجعل المولود اكثر خطرا من تناول الحبل و استعمال
Contraceptive pills
* مواقع الحمل عند استعمال ادوية تكون خطره على الحبيب

Medication use during pregnancy

- Most therapies for chronic diseases use pharmacologic medications that have not been thoroughly investigated in pregnant and lactating women, so the exact hazards and benefits of such interventions are unknown.

* لا توجد دراسات كافية لمدى تأثير الأدوية خاصة التي تستعمل في الأمراض المزمنة على المرأة الحامل والمحس
ما يحتمل فوائدها وتأثيرها الدقيقة غير معروفة

- **Medication use during pregnancy is of concern because:**
- The physiology of pregnancy affects the pharmacokinetics of drugs.
- Some medications can reach the fetus and cause harm.

قلقت
على سبل الجنين بمرور مستويات albomix عند المرأة الحامل وذلك يؤثر على كثير من الأدوية

Pharmacokinetics of Drugs in Pregnancy

- Stomach pH, transit time, metabolism, uptake, and efflux transport processes are altered during pregnancy, thereby potentially changing the drug's bioavailability.
وقت العبور (الامتصاص)
- Drug **distribution, metabolism and elimination** maybe altered during pregnancy.
- One of the biggest **physiologic changes** in pregnancy is the **expansion of plasma volume by approximately 50%**.

Fetal Development and Vulnerability

- Human embryogenesis, and the early stages of cell division and differentiation in the human embryo, is dependent on the ability of ^① stem cells to divide, ^② migrate, and ^③ specialize.
- This complex mechanism results in **teratogenicity and vulnerability**. The **teratogenic period** could be **particularly severe from day 31 to 71** after the last menstrual period in a 28-day cycle.

FDA Pregnancy Drug Risk Categories

Pregnancy Category	Description
A	Appropriate human studies - no risk
B	Insufficient human studies, but animal research suggests safety <u>or</u> : Animal studies show issues but human studies show safety
C	Insufficient human studies, but animal studies show problems <u>or</u> : No animal studies, and insufficient human studies
D	Human studies, with/without animal research show fetal risks, but the drug is important to some women to treat their conditions
X	Fetal risks are evident; there are no situations where the risk/benefit justifies use

These categories were not clear and often led to wrong conclusions. As of June 2015, all new FDA-approved drugs have a new pregnant label that included a risk summary, clinical considerations, and supporting data.

FDA Pregnancy Drug Risk Categories

After

2015

The U.S. Food and Drug Administration (FDA) implemented the Pregnancy Lactation Labeling Rule (PLLR).

Pregnancy Lactation Labeling Rule (PLLR)

- The PLLR provides a ^{طاق} set framework for drug manufacturers to provide information about the **risks and benefits of using prescription drugs and biologic products during pregnancy and lactation.**
- Prescribing information for clinicians has been updated in the labeling subsections:
 1. **Pregnancy:** ^{طوع} narrative risk summary of the maternal and fetal risks based on available human, animal, and pharmacologic data.
 2. **Lactation:** the amount of drug in breast milk and ^{الخطر المحتمل} potential effects on the breastfed infant
 3. **Females and Males of Reproductive Potential:** the need for pregnancy testing, contraception recommendations, and information about infertility as it relates to the drug.

Table 1. Drugs Identified With Known Risk of Teratogenicity

Human teratogen	Identifiable or Related Outcome
1 Alcohol	Fetal alcohol syndrome: IUGR and FTT; decreased muscle tone and poor coordination; developmental delay; and craniofacial abnormalities
2 Angiotensin converting enzyme inhibitors	Oligohydramnios; hypocalvaria; IUGR; renal effects (renal tubular dysplasia, anuria/oliguria, and hyperkalemia, end-stage renal failure); neonatal hypotension; cardiovascular abnormalities (e.g. patent ductus arteriosus, aortic arch obstructive); fetal death
3 Carbamazepine	10 × increased risk of neural tube defects; fetal anticonvulsant syndrome (IUGR, developmental delay, craniofacial defects, fingernail hypoplasia)
4 Cocaine ليس حواء	Placental abruption, fetal loss, low birth weight, microcephaly, limb and urinary tract malformations, poor neurodevelopmental performance
5 Coumarin anticoagulants	Fetal warfarin syndrome (nasal hypoplasia, eye abnormalities [i.e. optic atrophy, microphthalmia, and blindness]); epiphyseal stippling, hypoplasia of the extremities and fingernails; low birth weight; developmental retardation; fetal hemorrhage
6 Diethylstilbestrol (DES)	Clear cell adenocarcinoma and benign adenosis in exposed offspring
7 Methotrexate (Folic acid antagonists)	Central nervous system (i.e. anencephaly, neural tube defects); cardiovascular (tetralogy of Fallot); craniofacial (i.e. absence of lambdoid, coronal sutures, and frontal bone, low set ears, depressed/wide nasal bridge); long webbed fingers and absence of digits; growth and mental retardation
8 Phenytoin	Fetal anticonvulsant syndrome: IUGR; dysmorphic craniofacial features (i.e. microcephaly, low nasal bridge, cleft lip and cleft palate, maxillary hypoplasia); limb defects (i.e. hypoplastic nails and distal phalanges); cardiac defects
9 Isotretinoin	Spontaneous abortion; craniofacial abnormalities (i.e. microcephalus, hydrocephalus, deformity of ears, face, limbs); thymic hypoplasia; cardiac defects
10 Lithium	Tricuspid valve malformation (Ebstein's anomaly)
11 Misoprostol	Association with limb and neural tube defects
12 Tetracyclines	Weakened fetal bones, tooth enamel dysplasia, permanent tooth discoloration
13 Thalidomide	Limb, ear, cardiovascular and gastrointestinal anomalies
14 Valproate	Neural tube defects; fetal valproate syndrome: dysmorphic facial anomalies including microcephaly, hypertelorism, prominent forehead, low flat nasal bridge, low-set or odd-shaped ears

FTT, failure to thrive; IUGR, intrauterine growth retardation

Teratogenic Drugs

Only drug name is required

Clinical Toxicology

Introduction to Toxicology

Toxicology studies the harmful effects of chemical substances on all biological systems.

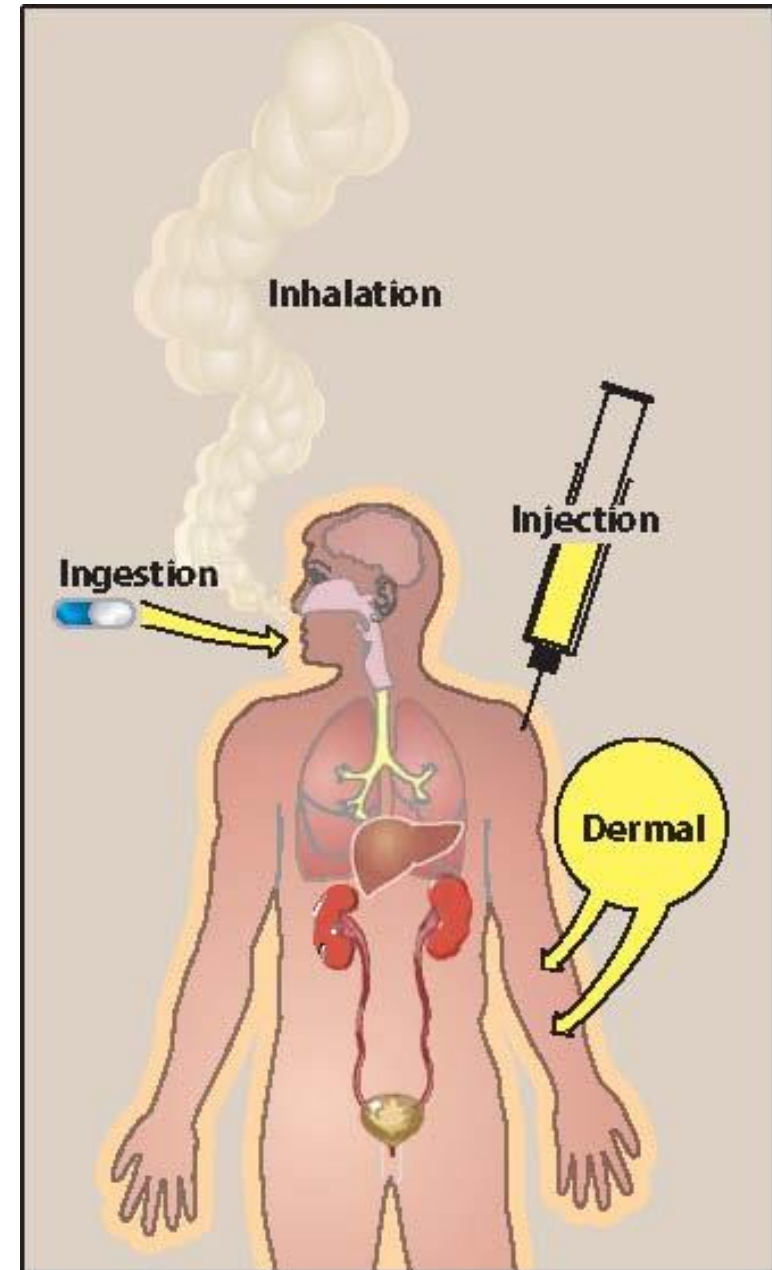
Occupational toxicology ^{تعرضه خلال العمل} is the study of substances that are found at work.

Environmental toxicology : is the study of potentially harmful effects of substances found in the environment as pollutants on living organisms .

Clinical Toxicology

• Toxins can be:

1. Inhaled
2. Insufflated (snorted) — الشَّم (تستعمل من قبل الدمييين) —
3. Orally ingested
4. Injected — (٧) —
5. Absorbed dermally — (٨) —



Emergency treatment of poisoned patient

* عند دراستنا لعالم السموم ما يهمنا هو علاج المريض الذي تضرر منه السموم وليس السموم

- **Treat the patient not the poison:**
 - I. **Airway breathing circulation (ABC):** give oxygen, IV access, heart monitoring
 - II. **Life threatening toxic effects** such as profound changes in blood pressure, heart rhythm, respiration and body temperature.
 - III. **Correction acid/ base and electrolytes disturbance.**
- If the patient have altered mental status, consider giving the “coma cocktail” (**dextrose**^① for hypoglycemia), **naloxone**^② (possible opioid or clonidine toxicity, and **thiamine**^③ for ethanol-induced Wernicke encephalopathy).

* لعرضك انك تعلم في طوري احدى الاستشحيات دمج مريض مع عليه في انت لا تعلم من الحالة الذي هو معها ما بنا بظنية مركب تسمى كوكتيل (cocktail) بهذا المركب يتكون من (3) مواد وهي

- ① dextrose يتعمل لمن احد حرمة كبريه من (insuline) وصارمه (hypoglycemia)
- ② naloxone يتعمل اذا كان المريض قد احد جرعه كبريه من المواد المسكرة (مريض) opioid
- ③ thiamine يتعمل لمن احد كبريه من الكحول clonidine

المرضى من حول المريض أو من جسمه حاله
النسب

Decontamination

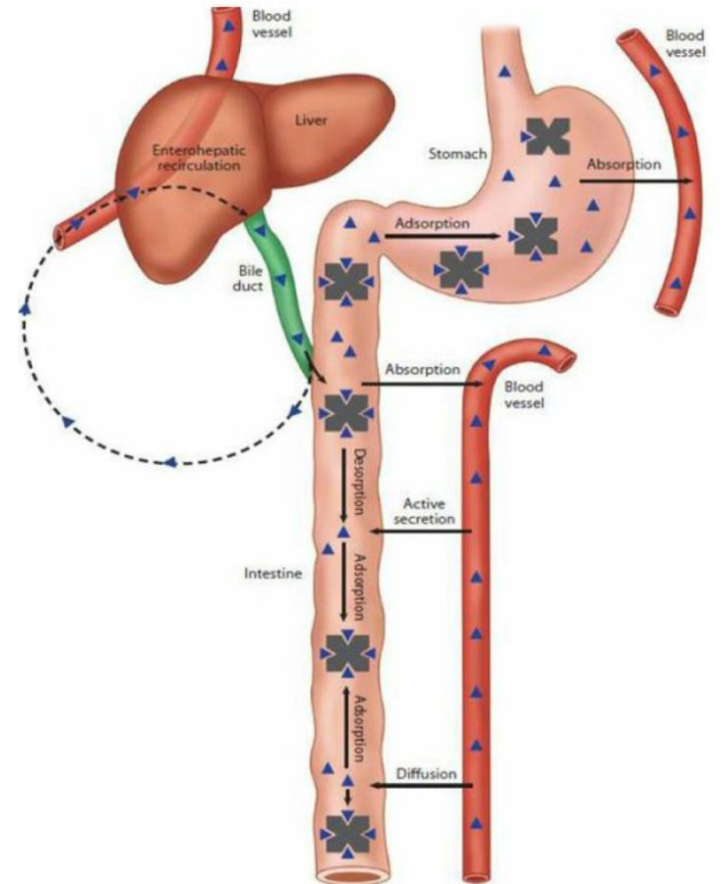
- Once patient is stabilized decontamination can occur.
- Eyes and skin exposure: flushing and washing → وقعت عليه مادة كيميائية
- Gastrointestinal decontamination:
 1. Gastric lavage → غسل المعدة
 2. Activated charcoal → يرتبط بالمركبات الضارة بجلودها خارج الجسم
 3. Whole bowel irrigation → غسل الأمعاء إذا حرمتمت المادة من المعدة ووجهت الأمعاء

Elimination enhancement

1. **Hemodialysis** e.g., methanol, salicylate and lithium.
 2. **Urinary alkalization**(by administration of sodium bicarbonate): enhances urinary excretions of salicylate and phenobarbital.
- ❖ **Serum pH shouldn't increase more than 7.55**
3. **Multiple dose activated charcoal:** e.g., phenobarbital, digoxin, and carbamazepine)

Activated Charcoal

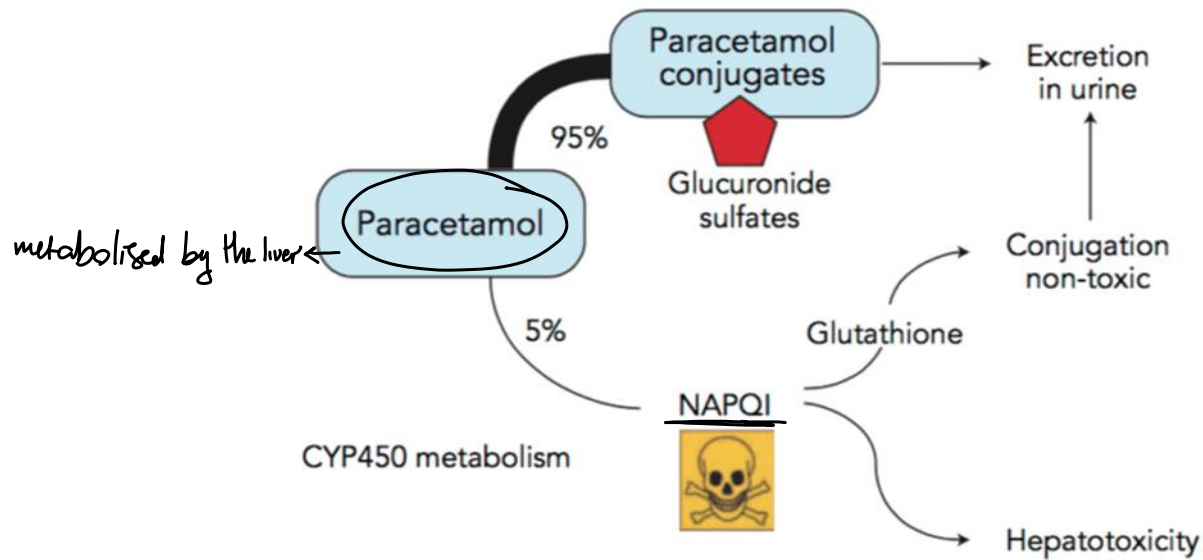
- Charcoal is extremely porous and has high surface area which create gradient a cross the lumen of the gut.
- Medications traverse from area of high concentration to area to area of low concentration, promoting absorbed medication to cross back into the gut to be adsorbed by activated charcoal, Also, activated charcoal blocks the reabsorption of medication that undergo enterohepatic recirculation by adsorbing the substance to activated charcoal.
- It doesn't effectively adsorb metals(e.g., lead, iron, lithium), corrosives, and alcohols.
- The most important contraindication is a not fully conscious patient with no swallowing reflex.



Select Pharmacological and occupational toxicities

- **Acetaminophen** → paracetamol
- **Alcohols**
- **Carbon monoxide**
- **lead**

Acetaminophen (Paracetamol)



At high doses normal metabolic pathways become saturated leading to the production of hepatotoxic metabolites (NAPQI).

As hepatic stores of glutathione are depleted and NAPQI binding to hepatocytes induces cell death and hepatic necrosis.

NAPQI: *N*-acetyl-*p*-benzoquinone imine

Acetaminophen (or Paracetamol)

يستخدم لعلاج تسبب ال

Phase 1 (0 to 24 hours): loss of appetite, nausea, vomiting, general malaise

Phase 2 (24 to 72 hours): abdominal pain, increased liver enzymes

Phase 3 (72 to 96 hours): liver necrosis, jaundice, encephalopathy, renal failure, death

Phase 4 (>4 days to 2 weeks): complete resolution of symptoms and organ failure

- Antidote is N-acetylcysteine (NAC).
- NAC replenishes hepatic glutathione and may also act as a glutathione substitute, combining directly with the toxic metabolite.
- Intravenous NAC is most effective when it is initiated within 8 to 10 hours of ingestion

Alcohols

1. Methanol (wood alcohol) and ethylene glycol:

• **Methanol** is found in windshield washer fluid and airplane fuel. It is metabolized to **formic acids** (toxic)

* الكينول يوجد في المصابيح و ربيط السيارات يتحلل مادة (toxic) Formic Acid تسمى

• **Ethylene glycol** is found in radiator antifreeze. It is metabolized to toxic metabolites such as **Glycolate**.

يتحلل مادة سامة تسمى

• **Primary alcohols are not toxic but lead to CNS depression. However, their metabolites are toxic.**

* الكحول نفسه غير سام لكن ال metabolites هي السامة.

If untreated:

• Methanol may cause blindness, metabolic acidosis, seizures and coma.

• Ethylene glycol may cause renal failure, hypocalcemia, metabolic acidosis

and heart failure.

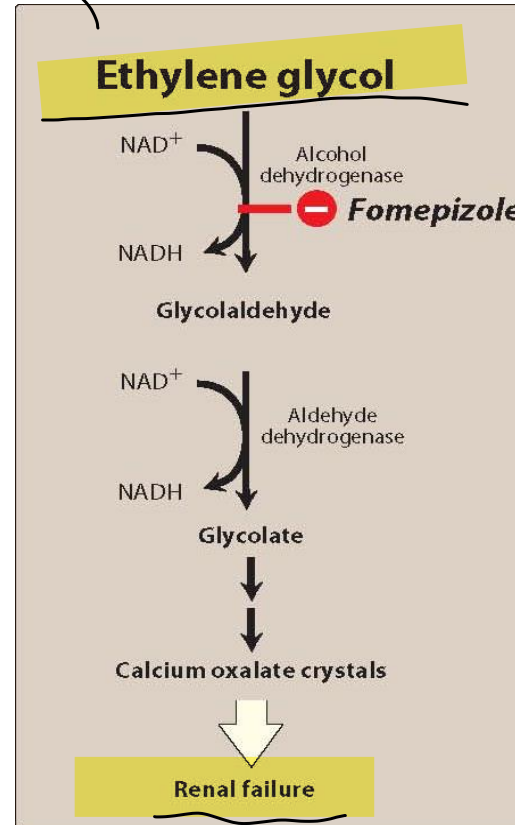
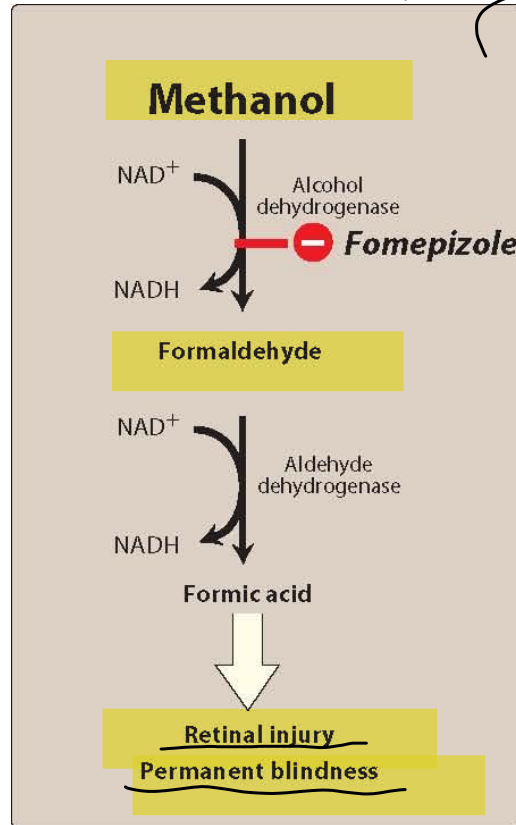
(1) العشى (2)

(3) (4)

(1) (2) (3)

Methanol and ethylene glycol

وقبل الحذر (غير ذلك لا)



Treatment:

← يمنع ال (metabolism)

- Antidote is Fomepizole.
- Hemodialysis to remove toxic acid that are already produced
- Administration of cofactors as folate for methanol, and thiamine and Pyridoxine for Ethylene glycol.

Alcohols

يستخدم في المنزل لتطهير الأسطح

2. Isopropanol (rubbing alcohol) : 2ndary alcohol, It is metabolized to acetone.

- It leads to CNS depression and GI irritation
- No antidote — لا يوجد مضاد
- Supportive care

Carbon Monoxide

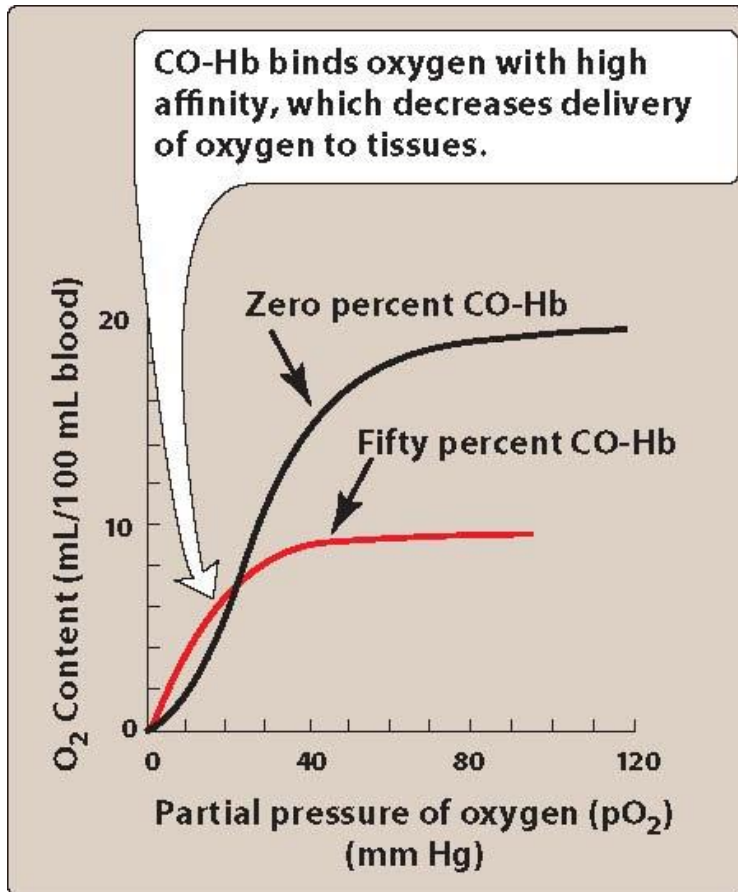
دائماً تحدث حالات الاحتراق بسبب الصدمات في التربة

لن نحوسه في تعاضل الموسوي لانا سأمه
لحقا

- **Carbon monoxide is colorless odorless and tasteless gas.**
- Sources: automobile, poorly vented furnaces, fireplaces and wood burning stoves ad kerosene space heaters, charcoal grills and generators.
- Binds to hemoglobin rabidly and forms carboxyhemoglobin.
- **The binding affinity of Carbon monoxide to hemoglobin is 230 to 270 greater than that of oxygen.**

* ال (affinity) ل CO << ال (O₂) ال الهيموجلوبين

Effect of CO on the oxygen affinity of hemoglobin



Co-Hb: carbon monoxyhemoglobin.

Treatment:

- To remove the source of carbon monoxide.
- Give 100% oxygen
- Oxygenation in hyperbaric Chamber

العلا }
له اطلعه من المكان الذي كان فيه
اعطيه اكسجين

Lead ← يرداد حالته عند الاطفال لاهم صكديا كوا من دهان الاسطح

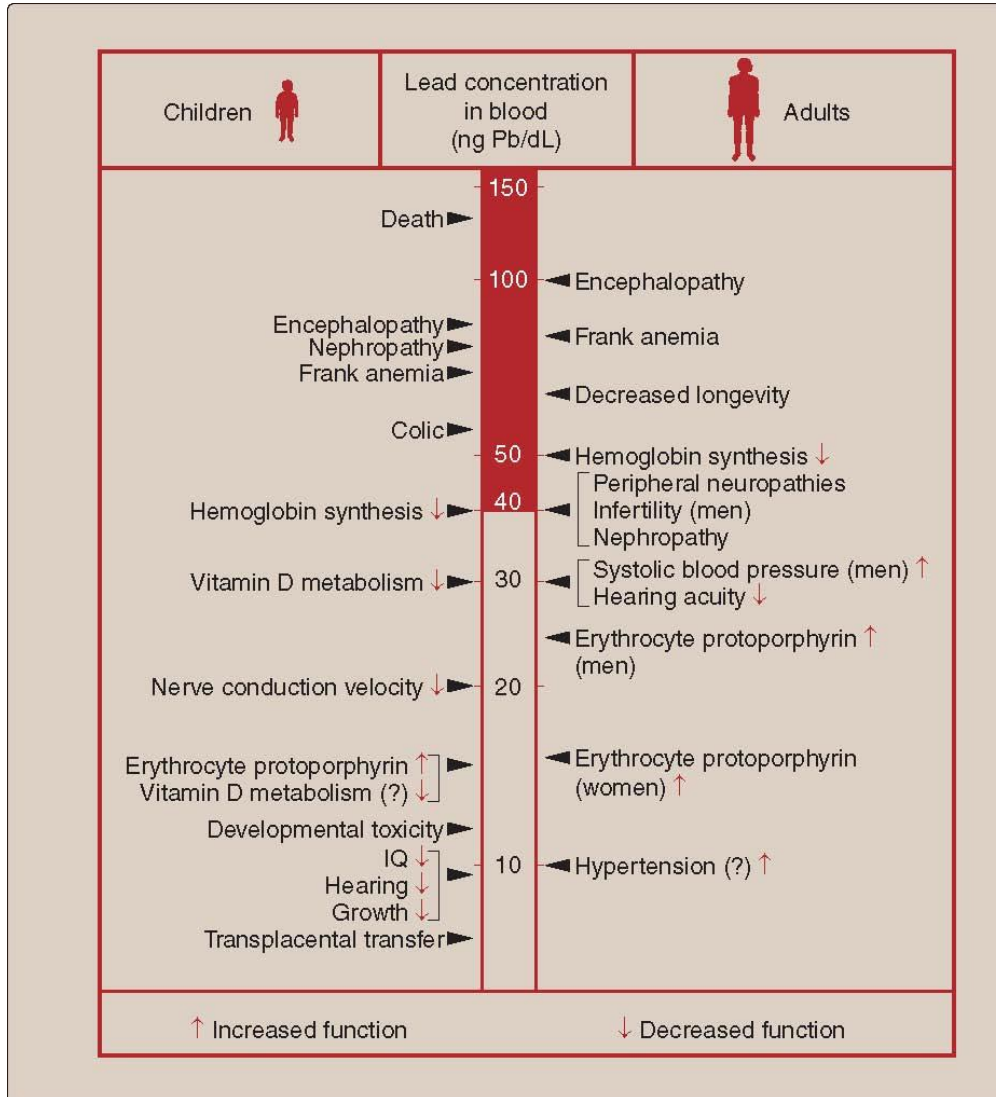
+ → الطفل يتصّب ال lead اكثر (١٧٤٠٠٠) من حجم البالغ

- **Source of exposure is old paint (houses constructed prior 1978) , industrial pollution, food, and contaminated water.**

- **Adults absorb 10%** of ingested lead, whereas **children absorb 40%.**

له يرد ترس الاطفال (Brain) اكثر لاهم في صرطه

Lead effects in children and adult



Treatment:

The first step in treating lead poisoning is to remove the source of the contamination.

② Chelation therapy: recommended for children with a blood level of 45 mcg/dL or greater and adults with high blood levels of lead or symptoms of lead poisoning.


Chelation agents contain **sulphydryl groups** that bind or chelate lead, and the resulting complex is excreted either renally or hepatically.

The chelation agents:

- ① Succimer (DMSA) is given orally
- ② Dimercaprol and edetate (EDTA) are administered parenterally. (10)

Antidotes:

For a variety of chemicals and poisons, specific chemical antidotes have been developed.



Poison	Antidote
Acetaminophen	N-acetylcysteine
Benzodiazepine	Flumazenil
Carbon monoxide	Oxygen(hyperbaric chamber)
Heparin	<u>Protamine sulfate</u>
Iron	Deferoxamine
Lead	Succimer (DMSA), Dimercaprol, edetate (EDTA)
Methanol and ethylene glycol	Fomepizole
Opiates	Naloxone
Warfarin	Vitamin K
Organophosphates	Atropine, pralidoxime