

 Cyanide (CN) is one of the oldest and most rapidly fatal toxins known to human.

 $H - C \equiv N$

- Its use as an agent of **homicide** and of **suicide** is well documented from the days of the **ancient Egyptians** to modern times.
- Fortunately, it also is one of the few poisons for which specific and effective antidotes are available.
- Cyanide occurs both **naturally** and as the result **of human activities**. **Many plants**, including **fruits** and **vegetables**, can release cyanide when ingested. Well-known sources include the pits of cherry, apricots, almond etc.



CYANOGENIC GLYCOSIDES

Amygdalin

Amygdalin (Almond seed) HCN in Hydrangea, Linum (Linseed) Prunus (Wild cherry) Sorghum vulgare (Jowar) Sorghum sudanese (Sudan grass) Gossypol (cotton seed)

CYANIDE PLANTS

250 mg CN/100g plant tissue

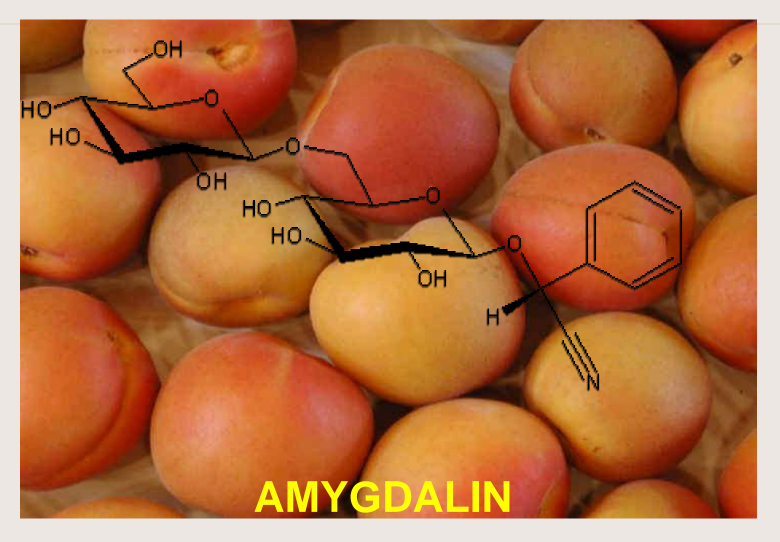
Almond

Cassava Cassava 104 mg CN/ 100 g plant tissue



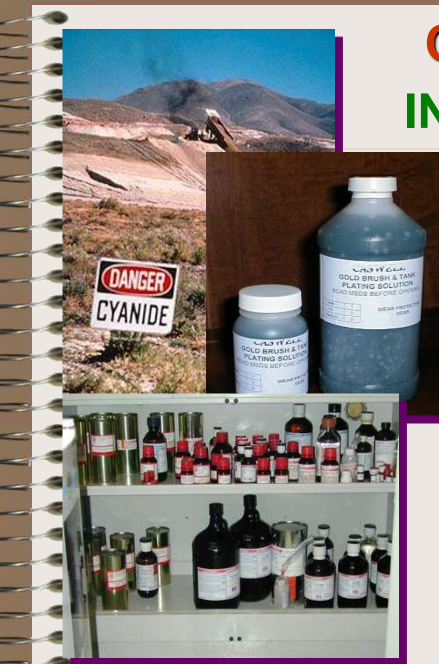
N N N N N

CYANIDE PLANTS



Sources and uses

- The many industrial sources include fumigants, insecticides, pesticides, and the production of plastic and rubber, electroplating, metallurgy and the processing of photographic film.
- The presence of HCN in various industrial gases results from an incomplete combustion of nitrogen-containing organic compounds, and its presence often is not suspected until an accident occurs.



CYANIDE INDUSTRY

ELECTROPLATING

HARDENING METALS

GOLD EXTRACTION

LABORATORIES

CYANIDE FIRE

 Victims of smoke inhalation have been shown to have toxic levels of both carbon monoxide and hydrogen cyanide.



CYANIDE/CARBON MONOXIDE Toxic twins

CYANIDE RODENTICIDE/FUMIGANT



FERATOX/CYANIDE PASTE



Cyanide can be released from a **large number** of different **compounds** (all having the **CN** group). $[:c \equiv N:]^{-}$

- Hydrogen cyanide (HCN)
- Sodium cyanide (NaCN)
- Potassium cyanide (KCN)
 - Calcium cyanamid (CaCN₂) etc
- Hydrogen cyanide the base compound of the group - is a colorless gas or liquid with a faint, bitter almond odor.



HYDROGEN CYANIDE CYANOGEN CHLORIDE

POTASSIUM CYANIDE SODIUM CYANIDE

Liquid hydrogen cyanide

TOXICITY

INGESTION

LETHAL DOSES

60-90 mg Hydrogen Cyanide (HCN)

200 mg Potassium Cyanide (KCN)

TOXICITY

INHALATION

Concentration (mg.m³)

Effect

300 200 150 120-150 50-60

20-40

immediately lethal
lethal after 10 minutes
lethal after 30 minutes
lethal after 30-60 minutes
20 minutes to 1 hour without
effect
light symptoms after several
hours

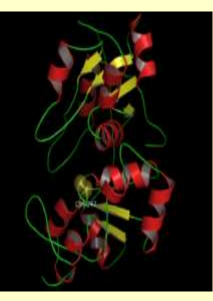
Mechanism of action

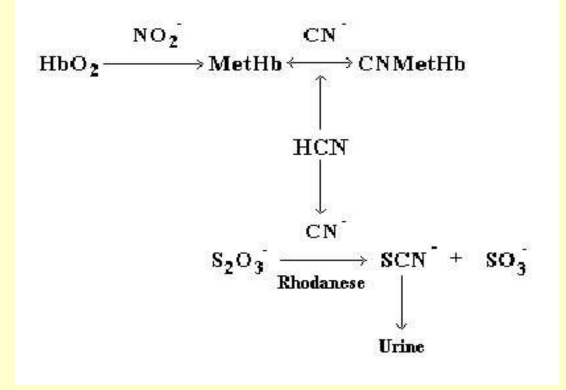
 Cyanide is rapidly absorbed through the lungs, skin and gastrointestinal tract because of its unionized state and low molecular weight. Symptoms occur within seconds of inhalation and within minutes of ingestion of cyanide salts.

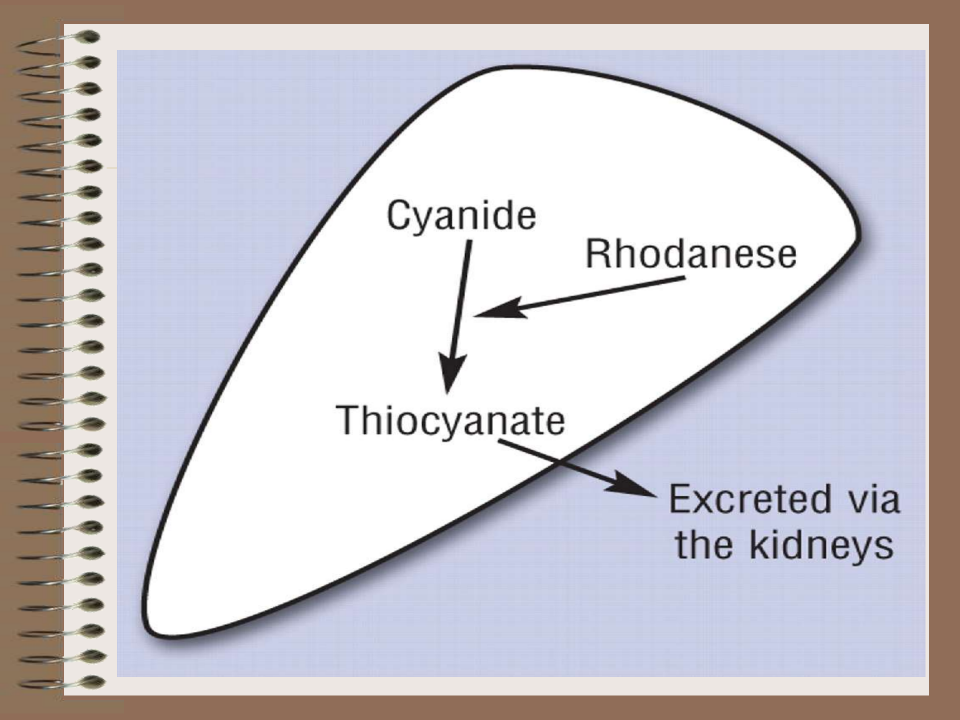
Cyanide can be **metabolized** by **five mechanisms**.

Only **two** of these five **ways** are **clinically important**:

 The major process of detoxification of the cyanide (80%) occurs mostly in the liver. The mitochondrial enzyme rhodanase catalyses the transfer of sulfur from thiosulfate to cyanide, forming the less toxic thiocyanate, which is excreted in the urine.



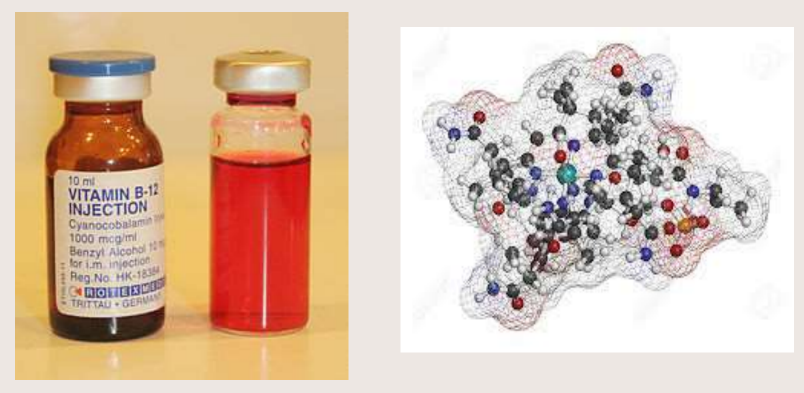








2. Cyanide **forms cyanocobalamin** through a reaction with **hydroxocobalamin** (**Vit.B**₁₂).



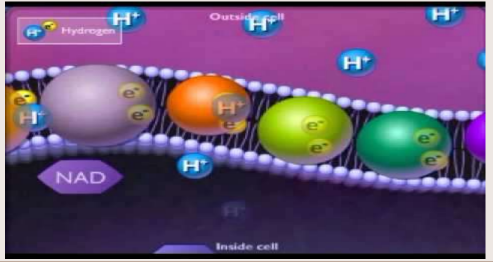


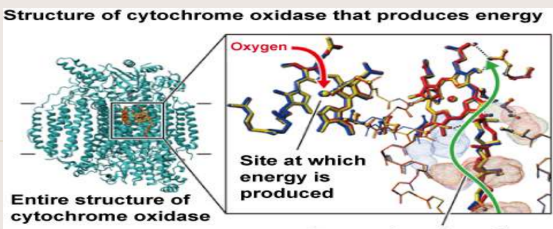
3.Cyanide is oxidized and incorporated into choline and methionine.

 4.Cyanide reacts with cystine, which product is excreted.

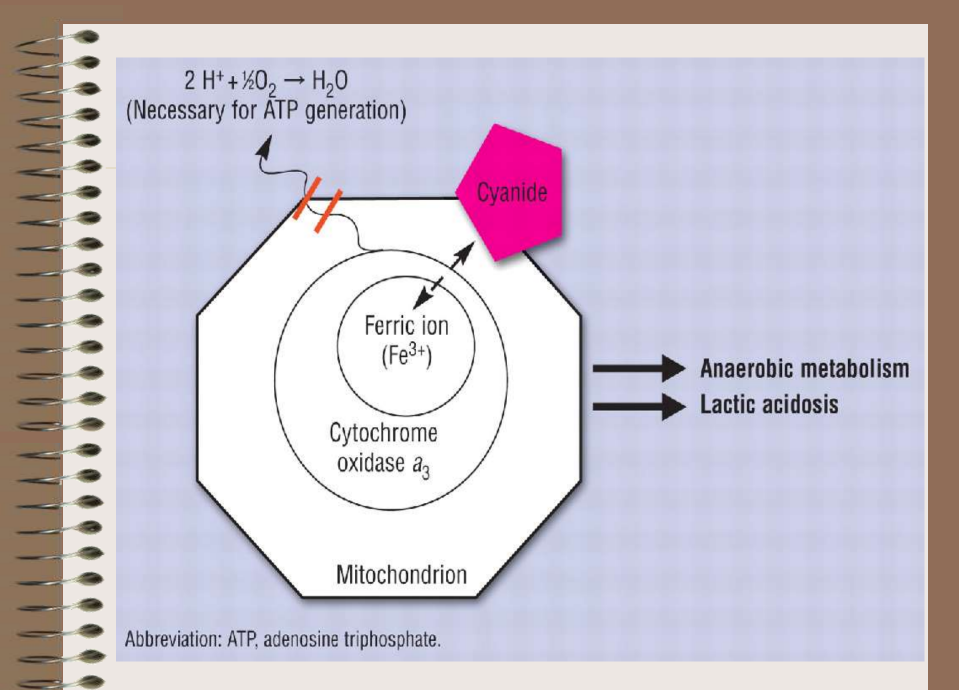
 5.Small amount of the cyanide is excreted unchanged in the lung and urine;

The toxicity of cyanide arises from its ability to inhibit the enzymes of the cellular respiration. Cyanide can form stable complex with the ferric ion in cytochrome oxidase. So, this poison inhibits the final step of oxidative phosphorylation, preventing the production of adenosine triphosphate (ATP) and results in anaerobic metabolism. Cellular anoxia results from the inability to use oxygen and a severe lactic acidosis ensues.

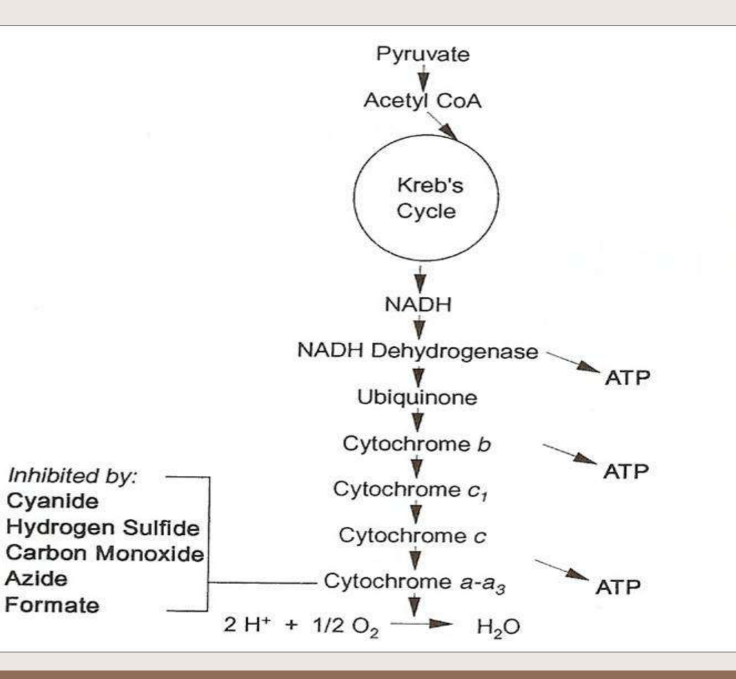




- Energy transfer pathway
- CN has high affinity for metals
 - Complexes with metallic cations at catalytic sites of several enzymes
- Binds ferric (3+) iron of mitochondrial cytochrome oxidase (cytochrome a-a3)
- cytochrome a-a3 mediates transfer of electrons to molecular oxygen (final step in oxidative phosphorylation)

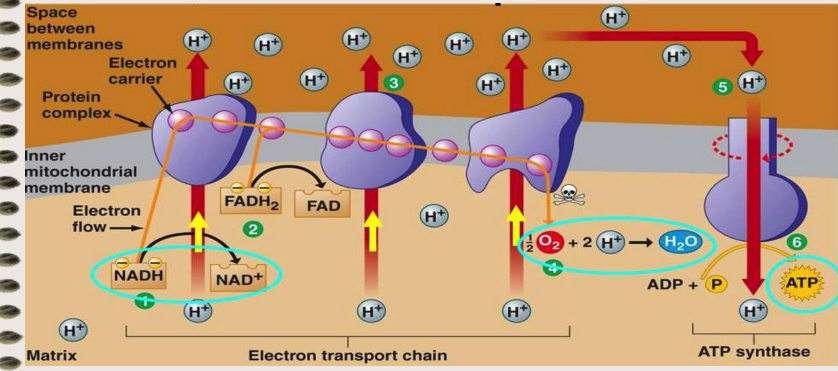


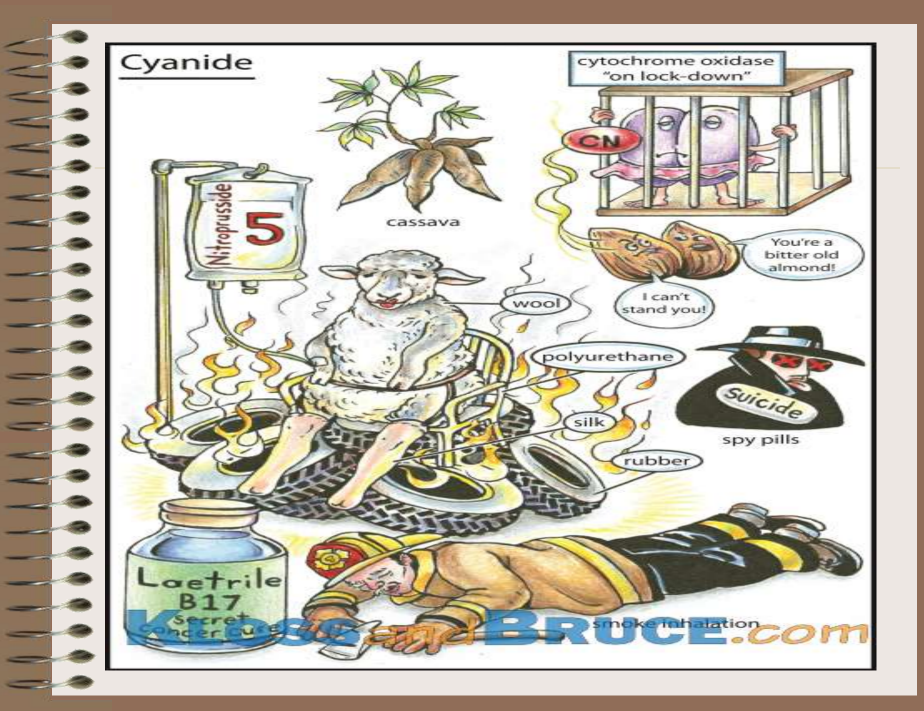




Blockade of oxidative phosphorylation

- Tissue anoxia
- Anaerobic metabolism
- Lactic acidosis •Arterialization of venous blood





Acute effects

Acute exposure to lower concentration (5 to 50 mg/m³) of hydrogen cyanide cause a variety of effects in humans - weakness, headache, nausea, increased rate of respiration and eye and skin irritation.

Acute effects

• Acute cyanide poisoning has primarily central nervous system effects. After inhalation - dry mouth, irritated eyes, itchy nose and throat and metallic taste is rapidly followed by air hunger and hyperpnea. Headache, flushed skin, stertorous breathing, agitation, muscle rigidity, ataxia, aphasia and confusion progress to loss of consciousness, and coma. Finally, bradycardia and hypotension lead to a terminal cardiovascular collapse.

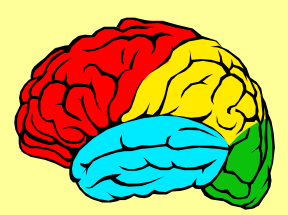
Acute effects

- Pulmonary edema and lactic acidosis have been observed after cyanide ingestion.
- Ingested cyanide follows a similar course, with vomiting caused by the strong irritant effect on the gastric mucosa.





- Headache
- Dizziness
- Seizures
- Coma



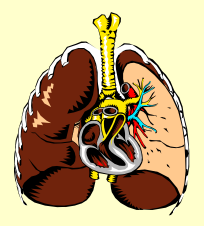
Cardiovascular

- Hypertension, tachycardia
- Hypotension, later in course
- Bradycardia
- Asystole
- Cardiovascular collapse





- Pulmonary
 - Dyspnea
 - Tachypnea
 - Pulmonary edema
 - Apnea
 - Gastrointestinal –Nausea, vomiting
 - -Caustic effects





Chronic exposure

- Chronic exposure involves the CNS (headache, tremor and loss of visual acuity) and endocrine systems (enlarged thyroid gland) and irritation to the eyes and skin.
- dermatitis in workers who are chronically exposed to cyanide solutions because the solution is strongly irritating and causes severe itching;
- cyanide rash papules and vesicles;
- blotchy eruption on the face;
- cyanide induced Parkinson's disease;
- visual disturbances;
- depletion of vit. B12;
- endocrine toxicity enlarged thyroid gland and altered thyroid function;

Diagnosis

An odor of bitter almonds, but up to 40% of the population is unable to detect cyanide by odor.

- A cherry red colour of venous blood, arising from poor oxygen utilization.
- > Metabolic acidosis.
- \rightarrow Absence of cyanosis.



Differential Diagnosis (DD)

- **Ingestion** with altered LOC (level of consciousness) and acidosis
 - Salicylates
 - Iron
 - Beta-adrenergic antagonists (beta-blockers)
 - Cocaine
 - Isoniazid (prophylaxis and treatment of tuberculosis)
 - toxic alcohols

Differential Diagnosis (DD)

- Inhalational Exposures
 - -hydrogen sulfide
 - -carbon monoxide
 - -simple asphyxiants

TREATMENT

- When cyanide has been inhaled the patient must be removed from the toxic atmosphere, contaminated clothing removed and exposed skin washed. When cyanide has been ingested, gastric aspiration and lavage should be carried out.
- The treatment begins with artificial respiration with 100 percent oxygen in patient with respiratory difficulty or apnea.
- The role of the **antidotes** in the therapy of cyanide intoxications is exceptional important, but they are potentially dangerous in the absence of cyanide ions. So, the diagnosis must be absolutely sure.

Treatment Antidotes

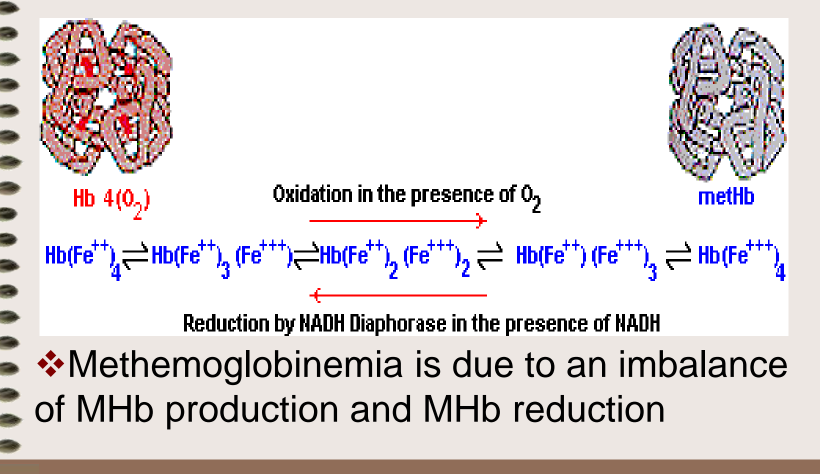
1.Methemoglobin-producers

Formation of **methemoglobin competes** for CN bound to **cytochrome oxydase**. Methemoglobin **removes the CN** from the enzyme and **restores its function** because the **iron (FeO3+) is oxidized** in methemoglobin. This leads to the **formation** of **cyanmethemoglobin**, which has a **low toxicity**.

The methemoglobinemia higher than 25% induced by methhemoglobin-producers can intensify tissue hypoxia.

What is methemoglobinemia?

Oxidation of iron within heme from Fe²⁺ to Fe³⁺



Toxins causing MHb

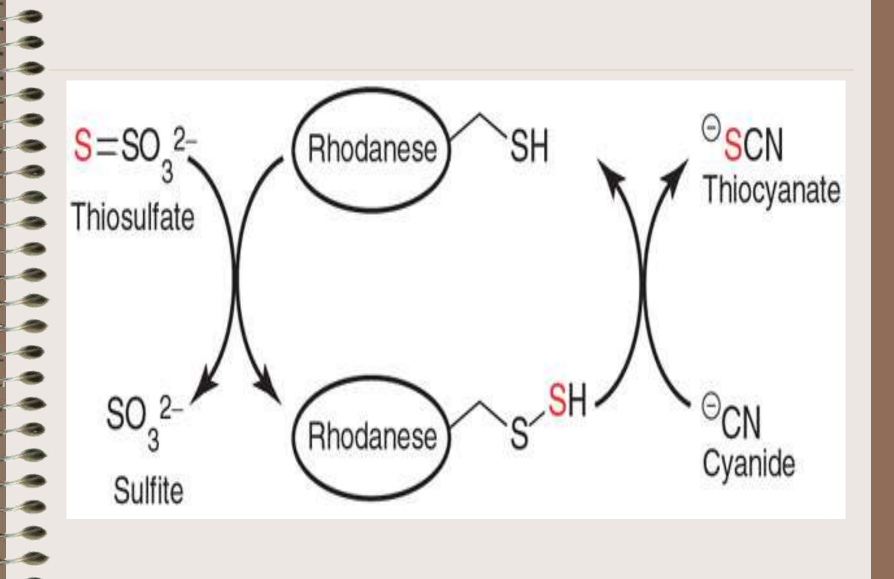
- Chloroquine (malaria, amebiasis)
- dapsone (treating leprosy)
- local anaesthetics
- methylene blue
- Metoclopramide (emesis)
- nitrates
- nitrites
- NTG (nitroglycerin)

- Nitroprusside (congestive heart failure, hypertension)
- Phenacetin (analgesic, pain, fever)
- Pyridium (pain releiver)
- Rifampin (tuberculosis)
- sulfonamides
- vitamin K₃
- Chlorhexidine (antiseptic agent)

T r e a t m e n t Antidotes 1. Methemoglobin-producers

- Amyl nitrite 0.2-0.4 ml via Ambu bag (first-aid measure).
- Sodium nitrate, 300 mg (10 ml of 3% solution), i.v. should be administered to adults to attain a desired methemoglobin level of approximately 25%. A dose of 10 mg/kg for children is recommended.
- DMAP (4-Dimethylaminophenol) 5 ml of 5% solution (250 mg or 3-4 mg/kg) i.v. for 1 minute

2. Cyanide is detoxified by conversion to thiocyanate
Sodium thiosulfate (50 ml of 25% solution) should be immediately given i.v.







Nitrites

N N N N N N N N N

- Therapeutic induction of methemoglobinemia
- NO2 + Hb = MHb
- Methemoglobin binds strongly to CN- and removes it from tissues
- **CN- + MHB = cyanomethemoglobin**
- cyanomethemoglobin is relatively non-toxic



Sodium Thiosulfate

donates sulfur molecule to rhodanese (enzyme which catalyzes formation of thiocyanate)

Na2S2O3 + HCN + O = HSCN

Synergistic effect

Oxygen

Synergy of 100% O₂ with nitrites/thiosulfate



3.Hydroxycobalamin (vitamin B12) 10ml of 40% hydroxocobalamin solution (vit.B12) (4 g) i.v. for 20 minutes.

Reduces cyanide to cyanocobalamin

$$B12_a + CN^- = B12$$

When combined with sodium thiosulfate end product is thiocyanate

≻Na2S2O3 + B12 = HSCN + B12a

- Recycling of hydroxycobalamin
- ➢ Renally cleared
- Synergistic effect of thiosulfate and B12a

TREATMENT Antidotes

4.CoNa2EDTA (dicobalt edetate) 20 ml of 1.5% solution (300 mg) i.v. for 1 minute. This chelating agent should be reserved for cases of severe poisoning because of its side effects, which include nausea vomiting, retrosternal pain, cardiac arrhythmia and facial edema. It may lead to loss of calcium and magnesium ions.

5. Glucosa - 25 g i.v. may be given a few minutes

