

# Occupational Hematologic Disease

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# Hematologic disorders

- Haematological problem is the most important health effect, such as benzene-induced aplastic anaemia
- Effects on the blood are direct but of less significance than the effects on other organ systems, such as lead-induced anaemia
- Secondary effect of a workplace hazard (polycytemia)

# Screening

- History
- Physical Examination
- Examination of Blood
- Bone Marrow Examination
- Other Examinations

# **History**

- Attention to nonspecific symptom of anemia
- A history of jaundice or pigmenturia
- Symptom of infection
- Symptom of thromcytopenia
- Drug history
- → Work history
- History of exposure to any potential toxic substance

# **Physical Examination**

- Examination of skin and mucous membranes
- Cyanosis
- Abnormal bleeding or petechiae
- Examination of the fundi and mucous membrane
- Examination of the lymph nodes and spleen
- Examination of the bone (bone tenderness)

# **Examination of Blood**

- Measurement of Hb, Hct, MCV,MCH
- Examination of peripheral blood smear :
  - Number of cells
  - Morphology
  - Differential count
- Retic count

#### **Bone Marrow Examination**

- Observation of erythroid of myeloid cells at all stages of maturation
- Presence of cells foreign to the marrow
- Presence of abnormal hematopoetic cells
- Chromosomal study

# **Other Examinations**

- Platelet count
- Bleeding time
- PT
- PTT
- Serum iron
- TIBC
- **LDH**
- Billirubin (Indirect)
- Uric acid

#### **Hemato Toxins Effect**

- Blood cell survival (Denaturation HB & Hemolysis)
- Metabolism(Porphyria)
- Formation(Aplasia)
- Morphology and Function (Preleukemia & leukemia)
- Coagulation (through development of thrombocytopenia)

# Methemoglobinemia and Hemolysis Produced By Oxidant Chemical

- Methemoglobin is formed by oxidation of ferrous Hb to ferric Hb.
- Oxidation of Hb was toxic to RBC and could be followed by an acute hemolysis
- Individuals with G6PD-D to be much more susceptible to oxidant stress than others.

### **Aniline**

- Historically, most work-related episode of methemoglobinemia and hemolytic anemia were a result of exposure to aromatic nitro and amino compound.
- Intermediate in the synthesis of aniline dyes, accelerator and antioxidant in rubber industry, Production of pesticides ,plastics, paints and varnishes.

# **Aniline**

Fat-soluble and readily penetrate the skin, even the through clothing

Vapor form may entry to the body through the lung

# **Clinical Presentation**

%Methemoglobinemia	Symptoms
10-30	Cyanosis, mild fatigue, tachycardia
30-50	Weakness, breathlessness Headache, exercise intolerance
50-70	Altered consciousness
>70-80	Coma, death

#### **Clinical Presentation**

- Patient appear deeply cyanotic.
- Freshly drawn blood appear dark maroonbrown and does not become red after exposure to air.
- Normoxia or mild hypoxia at pulse oximetry.
- Reticulocytosis in PBS(Polychromasia,possibly nucleated red cells).
- Polycythemia in chronic methemoglobinemia.

#### **Treatment**

- Removal of the offending agents
- Decontamination
- Mild intoxication(<%20) observation</p>
- Moderate to severe (>%30) oxygen%100,methylene blue solution %1
- Exchange transfusion

#### **Prevention**

- Minimize atmospheric and cutaneous exposure to oxidizing chemicals such as coal tar.
- Biologic monitoring in the workplace by measuring methemoglobin levels and reticulocyte counts.
- Screening for G6PD deficiency 1-2 months after the hemolysis has resolved.

# HEAVY METALS

- Arsine
- Lead
- Copper
- Cadmium

# **Arsine**

- Arsine poisoning:
  - Volatile, colorless, Nonirritating gas
  - Exposure sources:
    - Refining of metals
    - Processes of galvanizing , soldering-plating
    - Treatment of metals with acid
    - Semi conductor industry

# Signs & Symptoms

- Acute arsine poisoning are caused by acute and massive IV hemolysis +/\_delayed 2-24 h after exposure
- Nausea
- vomiting
- abdominal cramping
- headache
- malaise
- dyspnea
- Tea-colored urine

# **Physical Exam**

- Peculaire garlickly odor of arsine,fever,tachycardia,tachypnea and hypotension
- Jaundice, generalized nonspecific abdominal tenderness
- Acute tubular necrosis(oliguria or even anuria is usually evident within a day)
- CNS dysfunction, including alterations of mental status.
- Acute myopathy is seen with muscle tenderness and elevations of creatine phosphokinase

# **Laboratory Findings**

- Hemoglobinuria
- Decreased plasma haptoglobin & increased free HB levels
- Dramatic elevation of lactate dehydrogenase (LDH).
- Reticulocytosis/leukocytosis/ platelet count may be low
- Altered renal function AND increased serum Cr
- Arsenic levels in blood and urine are useful as indicators of exposure rather than as guidelines for therapy.

### **Treatment**

- Vigorous hydration
- Exchange transfusion if plasma Hb levels>400-500mg/dl
- If oliguric renal failure ensues, exchange transfusion is considered the treatment of choice because the hemolytic reaction itself may persist for up to 4 days after a single acute exposure.

#### **Treatment**

- There is no role in acute management for chelating agents such as British Anti-Lewisite because it cannot bind arsine or stibine to any appreciable degree.
- Hemodialysis if ARF developed
- All patients must be monitored closely until all evidence of hemolysis has resolved and renal function has stabilized.
- All survivors of acute arsine poisoning must be evaluated for at least 1 years
- Reduction of exposure or removal from exposure in chronic arsine poisoning

### Lead

- Suppression erythropoiesis and heme synthesis
- Hemolytic anemia
- Pathogenesis of lead induced hemolysis is related to inhibition of pyrimidin 5"nucleotidase
- Rarely is hemolysis the major presenting sign of acute lead poisoning

# Signs & Symptoms

- Abdominal pain
- Constipation
- Vomiting
- Anorexia
- Peripheral neuropathy
- Lead encephalopathy (convalsion, headache, confusion)
- Anemia
- Lead line
- Stippling of retina

# **Laboratory Finding**

- Anemia
- Mild increase of retic count
- Elevated of urinary levels of ALA & coporoporpherine III
- Elevated of urinary or blood levels of lead

# Copper

- Manufacture of wire, sheet metal, pipe, and other metal products,
- In agriculture as a fungicide, and as a preservative for wood, leather, and fabrics
- Mildly elevated serum copper levels
- IV hemolysis ,methemoglobinemia and RF
- Treatment is supportive

# Cadmium

- Battery and solar cell manufacture,
- Electroplating,
- Silver soldering,
- work around cadmium-containing pigments and coatings.

- Most notably on its nephrotoxic effects
- Acute anemia related, in part, to a hemolytic process
- Occupational cadmium exposure has not yet been definitively implicated as a cause of hemolytic anemia in workers.

# Chemical Causes Acquired Porphyria

- Hexachlorobenzene.....fungicide
- Vinyle chloride.....plastic
- Lead.....paint compound
- Aluminum.....phosphorus binder
- Chlorophenol..... Cleanser and disinfectant
- Polyhalogenated Aromatic Hydrocarbons
- Dioxin/PCBs /DDT

# **Acquired Porphyria**

- Neurotoxicity
  - Abdominal colic
  - Constipation
  - Autonomic dysfunction
  - Sensory motor neuropathy
  - Psychiatric problem
- Cutaneous photosensivity
  - Repetitive vesiculation
  - Scarring
  - Deformity
  - Hypertricosis of sun exposed area

#### Hexacholorobenzene

- In adults
  - Hyperpigmentation
  - Hypertricosis
  - Bulla
  - Weakness
  - Hepatomegaly
- In infants
  - Weakness
  - Convulsion
  - Cutaneous annular erythema

### Lead

- Blood lead level greater than 60 µg/dL
- The classic acute intermittent porphyria triad:
- Abdominal pain, constipation, and vomiting all representing the neurotoxic effects of excess 8-aminolevulinic acid and porphobilinogen.
- This triad is seen with equal frequency in lead intoxication.
- Other shared characteristics include: neuromuscular pains, paresis or paralysis, paresthesias, diarrhea, and seizures.

- The major differences between the two diseases are :
- 1- An increase in neuropsychiatric signs in acute intermittent porphyria compared with lead intoxication
- 2- Anemia, which is present in lead intoxication but virtually absent in porphyria

# **Aplastic Anemia**

- Idiopathic
- Secondary

(Drugs, Chemicals, Radiation, Infection, Immunologic)

Drugs:

chloramphenicol, acetazolamide, phenylbutazone, phenytoin, sulfonamide, cytotoxic drugs such as antimetabolites and alkylating agents ....

Benzene, ionizing radiation, and cytotoxic drugs such as antimetabolites and alkylating agents are dose-dependent basis

# Chemical Causes Aplastic Anemia

- **Benzene....** pesticides, rubber, fabric, solvents for glues, ink, paint
- lonizing Radiation
- **▼ TNT.....**production of explosive
- Lindane.....pesticide
- Arsenic......manufacture of glass, paint, weed killer, pesticide
- Ethylene glycol monornethyl ......Production of paints, lacquers, dyes, Inks, cleaning agents

#### Inhibition of cell production

- Benzene poisoning
  - Exposure sources
    - Manufacture of explosive
    - Dye industry
    - Dry cleaning
    - Shoe making industry
    - Production of cosmetic soap, perfume, drugs
    - Rubber manufacturing, petroleum and chemical production
    - printing
    - Steel working

#### Sign and Symptom

- Acute poisoning
  - Headache, dizziness, vertigo
- Chronic poisoning
  - shortness of breath, paller, petecheia, purpura, infection, painful mouth
  - angina
- Physiacal examination
  - Pale
  - Petechiae, bruises
  - Fever

#### Benzene

- Bone marrow:
  - Hypo cellularity with fatty replacement cytogenetic abnormalities are associated with benzene exposure, But specific chromosome changes are not.
- Prognosis:
  - Up to 40% patient may recover completely after removal of exposure
  - If hypo cellularity persists for more than several months, recovery is not likely to occur.
- Exposure is also associated with the development of acute non-lymphocytic leukemia and chronic myelogenous leukemia
- Treatment :supportive(transfusion,GFE,GCSF,GMCSF), Androgens, Allogenic BM trasplantation

#### **Ionizing Radiation**

- Risk of aplastic anemia is increased until 3-5 years after exposure, after which there is a marked decline in incidence
- Chromosomal aberrations may signify excessive exposure but is not predictive of aplastic anemia or leukemia
- Recovery may be seen after a prolonged period of aplastic anemia lasting 3-6 weeks.
- If recovery does not occur, permanent injury to the stem cell population will result in dysplasia or in leukemia.

#### Induction of leukemia

- Radiation
- Benzene
- Formaldehyde (myeloid leukemias)
- Butadiene (leukemia)
- Ethylene oxide(lymphoid tumors)
- Styrene
- Petroleum refining work

#### Multiple myeloma

- 15% of all hematologic cancers
- MM involves predominantly bone (skull), bone marrow and kidney
- Malignant proliferation of B-lymphocyte-derived cells that synthesize and secrete immunoglobulins
- Incidence of MM increases sharply with increasing age
- The peak incidence of multiple myeloma is between ages 55 and 65, and fewer than 2% of cases occur before the age of 40.
- Multiple myeloma is equally common in men and women

- Exposure to petroleum products, organic solvents, heavy metals, pesticides, and asbestos, high-dose radiation exposures implicated
- Workers thought to be at risk include :
- Agricultural workers
- Chemicals workers
- Miners
- Smelters
- Stokers
- Furniture workers

#### Non-Hodgkin's Lymphoma

- Pesticide (lindane or pentachlorophenol)
- Pentachlorophenol as a preservative
- Butadiene
- Benzene

Genotoxic and oxidative stress mechanisms, systemic immunosuppressive effects

## Agents Caused Toxic Thrombocytopenia

Agents Use Mechanism

Toluene diisocyanate Polymerizing agent Immune

DDT / Insecticide Megacarycyte hypoplasia

Turpentine Organic solvent Immune

Vinyl chloride Plastics Liver insufficiency with hypersplenism

## Platelet Aggregation & Hypercoagulation

- Some environmental substance caused platelet aggregation and hypercoagulation
  - Methyl mercury
  - Cadmium
  - Triethyl lead
- Some pesticide inhibit platelet aggregation
  - Chlorophenyl
  - Arochlor

#### Polycythemia

- Metal cobalt
- Chronic exposure to asphyxiants, especially carbon monoxide.
- Secondary polycythemia also may be seen with lung diseases due to environmental factors

#### **Neutrophils**

- Transient neutrophilia is common in a host of acute occupational and environmental illnesses, such as:
- Metal fume fever
- Alveolitis caused by irritant gases

- Neutropenia may be seen in the setting of injury to the bone marrow due to:
- lonizing radiation
- Arsenic
- Benzene
- Other agents affecting stem cells or the marrow microenvironment

### Occupational Exposure to Anticancer Drugs

- Oncology nurses and pharmacist are at risk for exposure to potentially mutagenic agents.
- The agent most commonly implicated in mutagenic potential are Alkylating drugs.

#### Prevention

- Worker education
- Gloves and protective Clothing
- Laminar hood for preparing drugs
- Proper waste disposal
- Periodic atmospheric checks & Biologic monitoring

