

Anemia (4)

Congenital Hemolytic Anemias/26.Oct/2015

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Congenital Hemolytic Anemias: Subtypes

- 1- Membrane defects: HS
- 2- Enzymopathies: G6PD Deficiency, PK Def
- 3- Hemoglobinopathies: B-Thal, SS

Anemia (4): Congenital Hemolytic Anemias

Case 4

18 yr old male presented to ER with headaches, dizziness, red urine and severe loin pain few hours after he ate fresh “fool” beans.

He looked jaundiced and sweaty.

His BP 90/60, Pulse rate 120.

He had no splenomegaly.

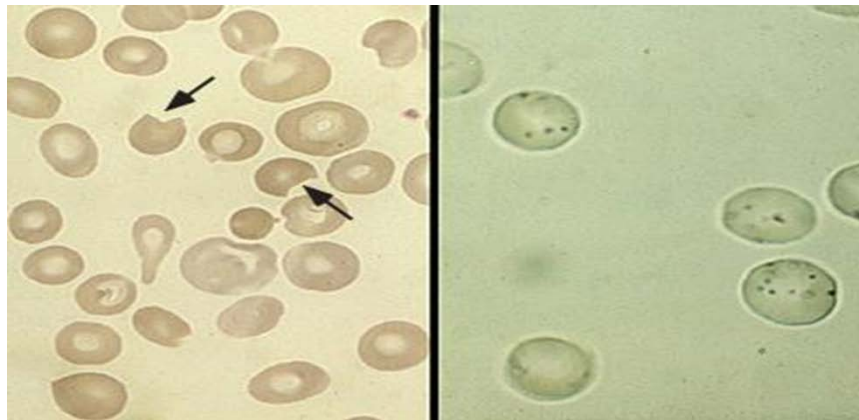
Hb 9 g/dl, WBC 16K, Plt 280K. Retics 9%.LDH 3000, Bilirubin 5; mostly indirect.

Case 4 Investigations & findings & Diagnosis

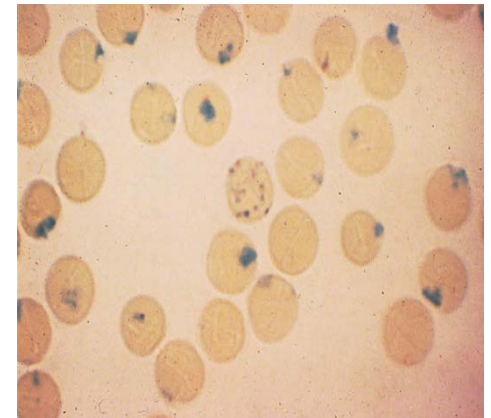
Urine



Bld film

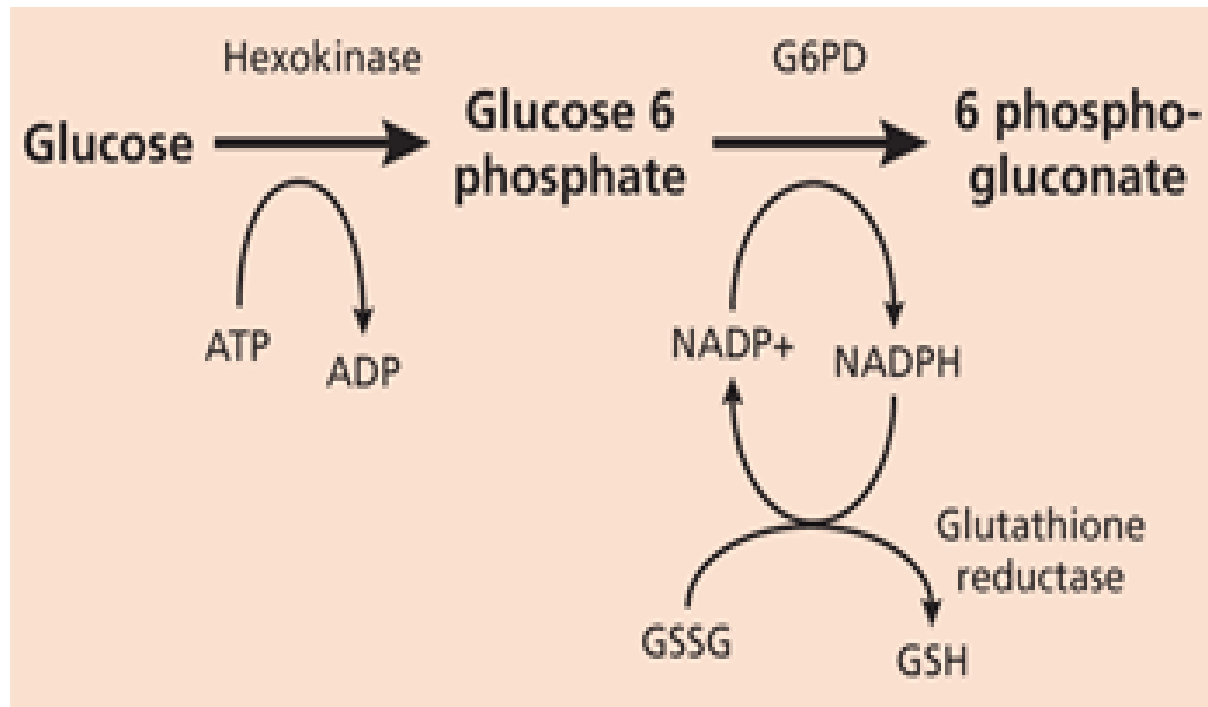


Heinz
Bodies

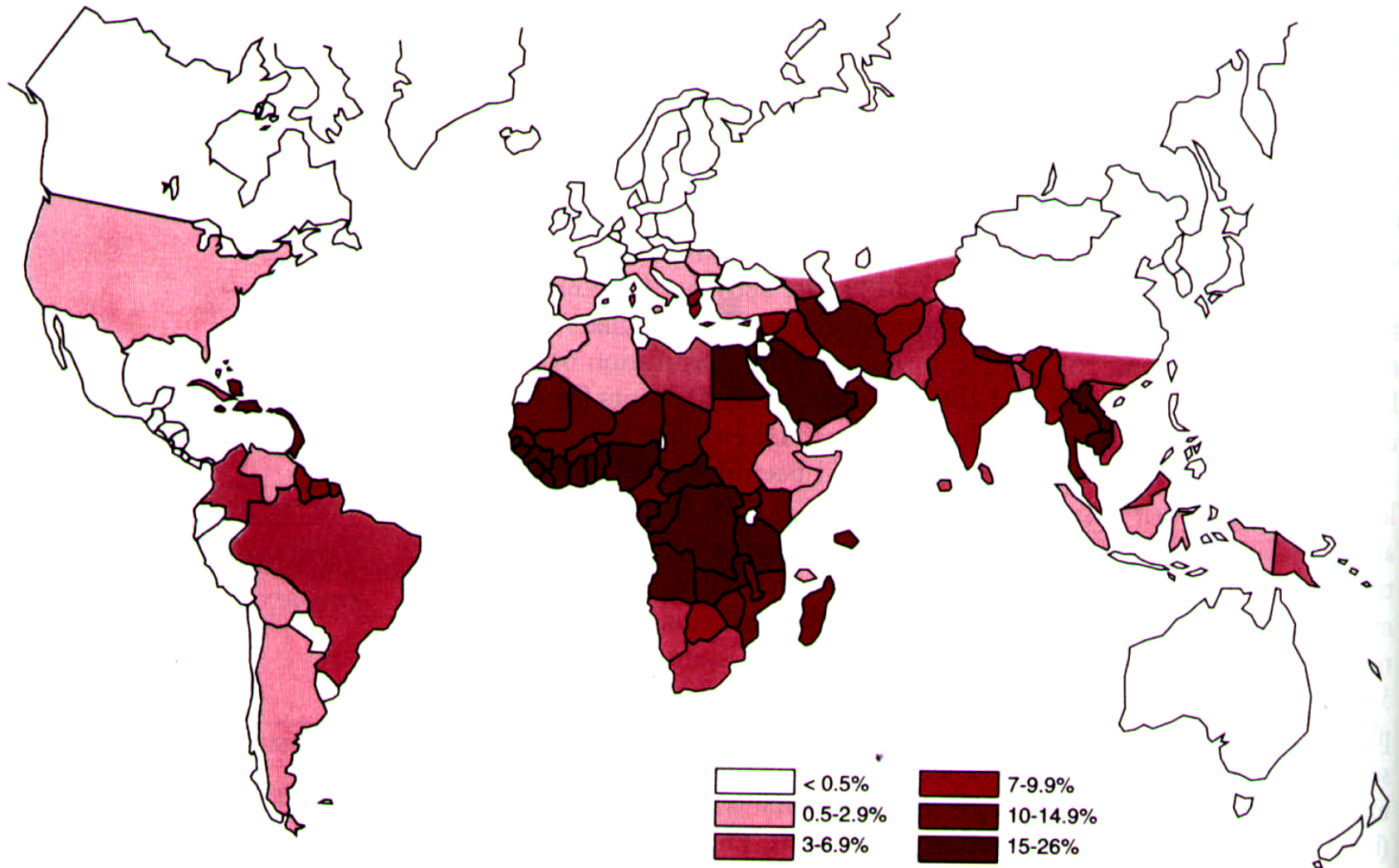


Diagnosis of case 4: G6PD deficiency: hemolytic anemia induced by fava(broad) beans (Favism)

Pentose Phosphate Pathway



Prevalence of G6PD: > 400 mill people/ Malaria belt



Clinical Features:

- **Disease from completely asymptomatic to severe intravascular hemolysis upon exposure to oxidant stress.**
- **Common precipitating factors:**
 - **Drugs: Primaquine– Nalidixic acid – sulpha drugs –**
 - **Infections**
 - **Diabetic ketoacidosis**
 - **Favism: hemolysis after exposure to Fava beans, occurs in Gd^{Med} variant**

Clinical Syndromes: G6PD Deficiency

- 1- **Neonatal Jaundice**: severe/ Kernicterus /, 1-3 day after birth.
- 2- **Favism**: acute intravascular Hemolysis due exposure broad bean (Vicia fava), the offending agent is **divicine**, it produces free Oxygen radicals and autooxidation.
- 3- **Infection** which promote the formation of H_2O_2 following oxygen burst in neutrophils and macrophage may result in hemolysis
- 4- **Drug induced hemolysis**

Genetics

- 1989 - more than 400 variants of G6PD
- Variants divided 5 classes according to the residual enzyme activity based (WHO).
- Mediterranean and African (A-) variants - by far the most clinically significant.
- Enzyme activity scarcely detectable in the Mediterranean type

Genetics

- Majority of the variants - from a single point-mutation resulting in amino acid substitution in gene encoding for G6PD located at the Xq28 region on the tip of the long arm of the X-chromosome
- G6PD Mediterranean is frequently caused by mutation (**563 C-->T**)

Therapy

- Avoid precipitating factors.
- Blood transfusion in severe hemolysis.
- Maintenance of good urine output during hemolytic episodes
- Folic acid.
- Exchange transfusion in newborn

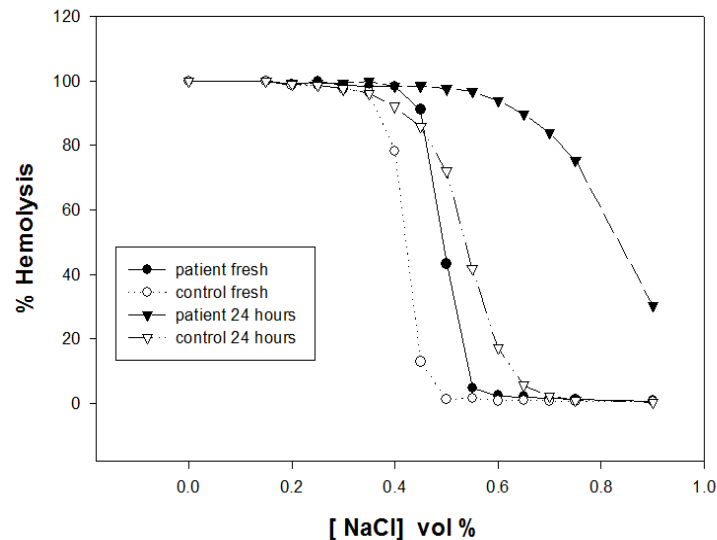
Case 4 B

36 yr old lady presented with “anemia syndrome” and splenomegaly. She was mildly jaundiced. Hb 8g/dl, retics 10%, WBC, Plt were normal. LDH 1160, Bilirubin 3mg/dl d 1. DAT –ve.

Bld film:
spherocytosis/splnc
conditioning



Osmotic fragility test



Abd. US



Hereditary Spherocytosis

- Prevalence and inheritance
 - Common In Northern Europeans
 - Clinical severity is **highly variable**, but uniform within a given family
 - Typically the autosomal dominant *homozygous is very severe or lethal*
 - some recessively inherited
 - Supportive therapy
 - No consensus for splenectomy indications

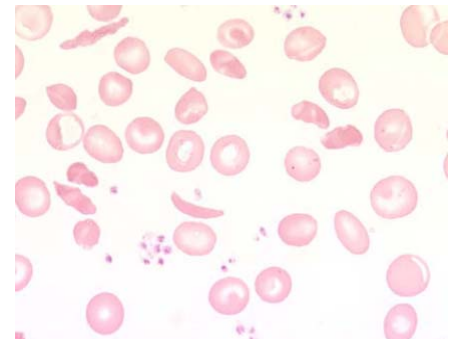
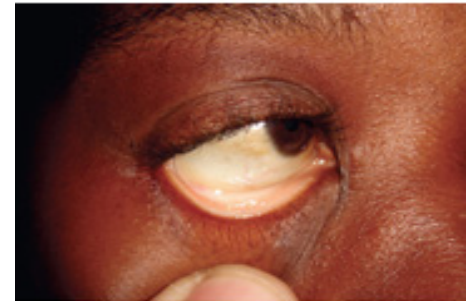
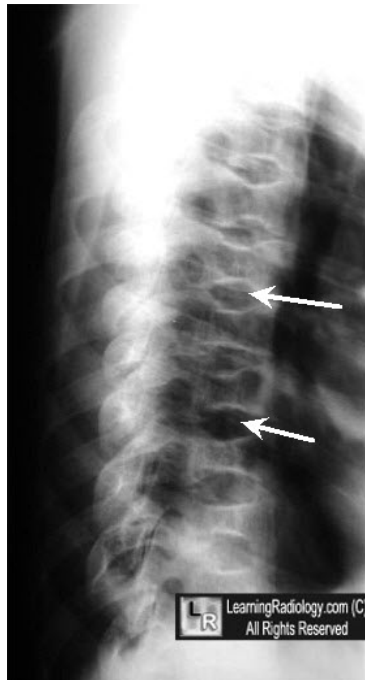
Hereditary Spherocytosis

- Molecular pathology
 - Partial deficiency of spectrin
 - Combined deficiency of spectrin and ankyrin
 - Molecular Defects:
 - mutations of **ankyrin: most common**
 - mutations of **band 3** protein
 - mutations of **protein 4.2** (common in Japanese)
 - Others: β & α spectrin, protein 4.9 are rare

Case 4 C

18 yr old male complains of acute pain in his back, Dizziness, Fatigue, Shortness of breath and Headaches for the last 6 hours. He has had similar attacks in the past. P/E

Xray spine



Sickle Cell Disease

- Inherited as autosomal recessive
- Point mutation in beta globin gene ($\beta 6$ Glu \rightarrow Val)
- Gene occurs in 8% of African-Americans

Sickle Cell Anemia Clinical Effects

- Chronic hemolytic anemia
 - Gallstones (bilirubin)
 - Risk of red cell aplasia (Parvovirus)
 - Decreased vascular tone
- Susceptible to infection
 - Functional asplenia
 - Infarcted tissue
 - Numerous manipulations
- Vaso-occlusion
- Autosplenectomy

Vascular beds susceptible to injury

- Brain
- Lung
- Ankle
- Erectile vasculature of the penis

Infectious complications of Sickle cell anemia

1- Related to absent spleen

- Pneumococcus infections
- Hemophilus infections
- Dramatically improved with the use of prophylactic penicillin in childhood

2- Others: staphylococcal, Osteomyelitis

Sickle Cell Anemia Vaso-occlusion: Unique pathophysiologic feature

- Causes acute and chronic organ damage
- Acute complications
 - Sickle cell vaso-occlusive pain crisis
 - Hepatic crisis
 - Splenic crisis
 - Priapism
- Chronic organ damage
 - Stroke
 - Chronic lung disease with pulmonary hypertension
 - Renal failure
 - Avascular necrosis of bone

Sickle cell: avascular necrosis of the hip



Sickle Cell Anemia Vaso-occlusive Events (Pain Crisis)

- Precipitating factors
 - Hypoxia
 - Acidosis
 - Fever
 - Infection
 - Dehydration
 - Exposure to cold
- Perceived factors
 - Exposure to cold 34%
 - Emotional stress 10%
 - Physical exertion 7%
 - Pregnancy 5%
 - Alcohol consumption 4%
 - Not identified 40%

Sickle cell vaso-occlusive crisis

- Serious complication of sickle cell anemia
- Risk of acute event (<48 hours)
 - Acute chest syndrome
 - Splenic sequestration
 - Massive hemolysis
 - Risk of sudden death

Sickle Cell Anemia Painful Events: Management Principles

- Correct fluid/electrolyte abnormalities; use hypotonic fluid and limit volume to avoid overhydration
- Treat any underlying illness
- Opioid analgesics (meperidine is not recommended)
- Blood transfusion is not indicated for an uncomplicated pain episode
- Incentive spirometry should be used during waking hours

Prevention of Painful Episodes

- Hydroxyurea increases Hgb F
 - Reduces the frequency of painful episodes, acute chest syndrome, RBC transfusions and hospitalizations
- Non-pharmacologic approaches have not been fully evaluated
- Prophylactic transfusions showed a decreased incidence of painful crisis in pregnancy

Addiction and pseudo-addiction

- Addiction (abuse)
 - Overwhelming involvement with obtaining and using mind-altering drug
- Pseudo-addiction
 - Relief seeking behavior misidentified as addictive behavior

The “difficult” patient with sickle cell disease

- Approximately 5% of patients account for 25-50% of hospitalizations
- High use group has >10 hospitalizations/year
- Hospital stay is longer (10 days vs. 6 days)
- Young, poor, unemployed, male
- Difficult interpersonal interactions
- May be associated substance abuse
- Higher death rate

Acute Chest Syndrome: Clinical Findings

- Etiology - multifactorial
 - Rib infarct causing splinting/atelectasis
 - Pulmonary fat embolism
 - Infection (mycoplasma, chlamydia, viral)
- Indistinguishable from pneumonia
 - Pleuritic chest pain, fever, cough, tachypnea, hypoxia
- Laboratory diagnosis
 - Worsening anemia
 - Infiltrate on chest radiograph

Acute Chest Syndrome: Outcome

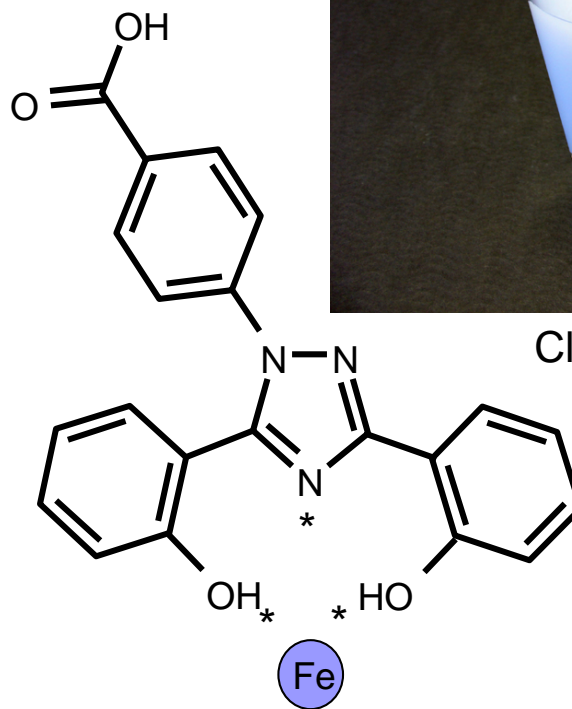
- Complete recovery 91%
 - Weaned of supplemental O₂ 3.1±1.9 days
 - Hospital discharge 5.4±2.3 days
 - Chronic respiratory disease 3%
-
- Death 6%

Acute Chest Syndrome: Prevention and Treatment

- Incentive spirometry
- Treat possible underlying infection
- Bronchodilators and supplemental oxygen
- RBC transfusion therapy
 - Simple transfusion
 - Exchange transfusion for:
 - Multiple lobes involved
 - Rapidly progressing
 - Worsening hypoxia

Deferasirox : Oral Iron Chelator in chronic blood transfusion

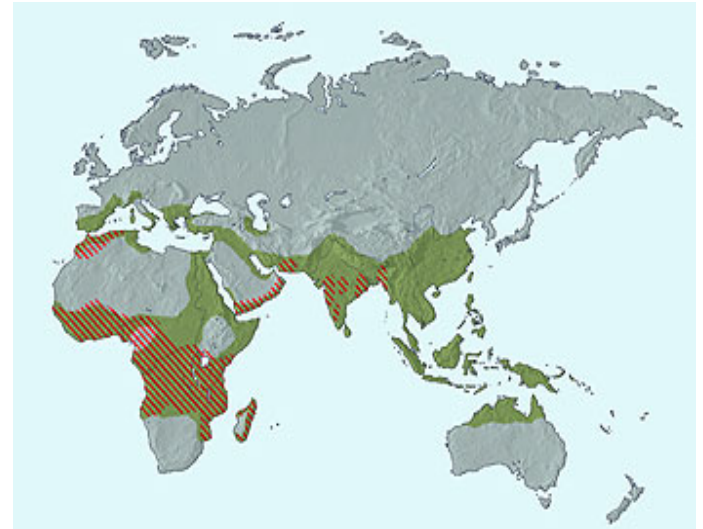
- Tridentate* iron chelator
 - An oral, dispersible tablet
 - Administered once daily
 - Highly specific for iron
- Chelated iron excreted mainly in feces (< 10% in urine)



Clinical trial formulation
or preparation

Sickle Cell Trait

- Protection against malaria
- Genitourinary complications
 - Hyposthenuria/
papillary sloughing
 - Painless hematuria
 - UTI during pregnancy
- Vaso-occlusive complications
 - Splenic infarction with hypoxia
 - Sudden death
 - Rhabdomyolysis



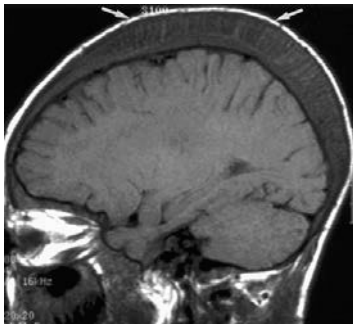
Sickle cell trait areas shown in orange stripes

Case 4 D

13 yr old male complains of skin pigmentation, abdominal swelling and pallor. He has been receiving blood transfusions since the age of 9 months. Stunted growth. Hb 6, MCV 55, retcs16%,s.Ferritin 5000.

P/E

Xray



Beta Thalassemia

Clinical Syndrome	Genotype	Hgb (g/dl)	Hgb Analysis
Minor (Trait)	β/β^+ or β/β°	10-13	\uparrow Hgb A2, \uparrow Hgb F
Intermedia	β^+/β^+	7-10	\uparrow Hgb A2, $\uparrow\uparrow$ Hgb F
Major (Cooleys)	β^+/β° or β°/β°	< 7	\uparrow Hgb A2, $\uparrow\uparrow\uparrow$ Hgb F

Most commonly reported mutations in the B-globin gens in Jordanians

- **Eight mutations constituted about 86% of the Jordanian thalassemic mutations**
- **These mutations were IVS1-110 (G>A) (25%), IVS2-1 (G>A) (15%),**
- **IVS2-745 (C>G) (14.2%), IVS1-1 (G>A) (10%), IVS1-6 (T>C) (8.3%), codon 37 (G>A) (6.3%),**
- **codon 39 (C>T) (4.6%), and codon 5 (-C) (3.8%)**

Beta Thalassemia: Clinical Manifestations/ complication

Osteoporosis

Extramedullary erythropoiesis/ tumor effect

Iron overload: skin, heart, liver, endocrine organs

Dilated cardiomyopathy secondary to severe anemia

Growth and development delayed

Large splenomegaly

Treatment/ Prevention of B thal major

- Blood Transfusion
- Iron chelation: deferroxamine (parenteral)
- ?splenectomy
- Allo-BMT
- Supportive
- **Prevention**

Oral deferasirox

