

# Hemoglobinopathies

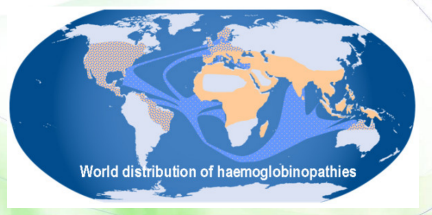
Prof. Mamoun Ahram Hematopoietic-lymphatic system

### Resources

- This lecture
- Mark's Basic Medical Biochemistry, Ch. 44

### What are hemoglobinopathies?

- Hemoglobinopathies: Disorders of human hemoglobin.
- The most common genetic disease group in the world (5% of people are carriers) with substantial morbidity (about 300,000 born each year).
- Hemoglobin disorders account for 3.4% of deaths in children < 5 years.



### Hereditary hemoglobins disorders

- Quantitative abnormalities are abnormalities in the relative amounts of  $\alpha$  and  $\beta$  subunits (thalassemias).
- Qualitative abnormalities: mutations resulting in structural variants.
  - Over 800 variants have been identified.
- Hereditary persistence of fetal hemoglobin (HPFH): impairment of the perinatal switch from to globin.

# **Quantitative abnormalities** (thalassemias)

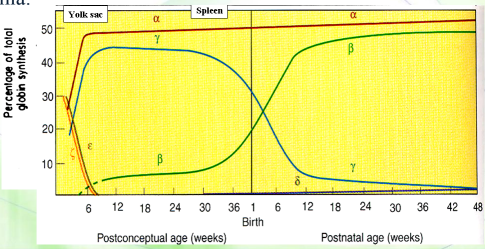
### Thalassemias

- Thalassemias: the most common human single-gene disorder.
- They are caused by a reduced amount of either the or protein, which alters the ratio of the : ratio.



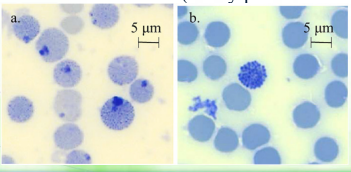
### The Alpha-Thalassemias

- Alpha-thalassemia: underproduction of the -globin chains.
- HbA (22), HbF (22), and HbA2 (22) are all affected in α-thalassemia.



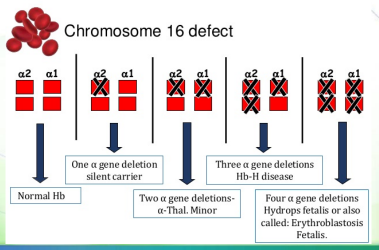
### HbH

- With the reduction of chain production, and -chain production is established, homotetramers of (4 or HbH) are formed.
- The HbH tetramers have a <u>high</u> affinity towards oxygen and are <u>highly</u> unstable (meaning that they denature, aggregate and precipitate resulting in the formation of Heinz bodies).
- The main type of mutation is deletion (rarely point mutations)



### Variable severity

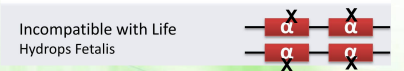
- With -thalassemias, the level of -globin production can range from none to very nearly normal levels.
- This is due in part to the fact that each individual has 4 genes.



### -thalassemia major

### Hydrops fetalis

- 4 of 4 genes are deleted.
- The predominant fetal hemoglobin is a tetramer of -chains.
- 4 or Hb Bart: a homotetramer of .
- Hb Bart has a high affinity towards oxygen.
- This situation is called hydrops fetalis.
- Stillbirth or death shortly after birth occurs.







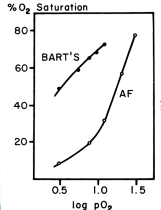


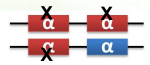
Fig. 4.—Oxygen dissociation curves of the hemoglobin components of cord blood: Hb-Bart's and the Hb-A and F mixture

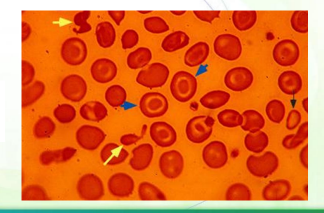
### -thalassemia intermedia and

### Hemoglobin H disease

- 3 of 4 genes deleted.
- Mild to moderate hemolytic anemia in adults.
- A high level of 4 tetramer is present.
- Clinically, it is known as hemoglobin H disease.
- The disease is not fatal.

Hb H Disease: Symptomatic Hemolytic and Microcytic anemia Splenomegaly





### -thalassemia minor and silent carrier

- Thalassemia trait: If 2 of the 4 genes are inactivated.
  - The individuals are generally asymptomatic.
- Silent carrier: 1 of 4 genes deleted.
  - Individuals are completely asymptomatic.

### Summary of -thalassemias

Genotype	α-globin gene number <sup>a</sup>	Name	Phenotype
αα / αα	4	Normal state	None
αα / α–	3	Silent carrier	None (values for Hb and MCV may be near the lower limits of normal)
/αα or α-/α-	2	Thalassemia trait	Thalassemia minor: asymptomatic, mild microcytic anemia
/α-	1	Hb H disease	Thalassemia intermedia: mild to moderate microcytic anemia
/	0	Alpha thalassemia major	Thalassemia major: hydrops fetalis

<sup>\*</sup>Number of normal alpha globin genes

### The beta-thalassemias

- -globins are deficient and the -globins are in excess and will form -globin homotetramers.
- Main type of mutation is point mutations, mutations within the promoter or LCR, translation initiation codon, splicing positions, or polyadenylation termination signal.
- The -globin homotetramers are extremely insoluble, which leads to premature red cell destruction in the bone marrow and spleen.

### -thalassemia major and minor

#### -thalassemia major

- A complete lack of HbA is denoted as 0-thalassemia or -thalassemia major.
- Affected individuals suffer from severe anemia beginning in the first year of life and need blood transfusions.
  - Long-term transfusions lead to the accumulation of iron in the organs, particularly the heart, liver and pancreas and, finally, death in the teens to early twenties.

#### -thalassemia minor

- Individuals heterozygous for thalassemia with one normal -globin gene and a mutated gene are termed thalassemia minor.
- Individuals with beta-thalassemia minor are generally asymptomatic.

### Classification and types of -thalassemia

Common genotypes	Name	Phenotype
β/β	Normal	None
β/β <sup>0</sup> β/β <sup>+</sup>	Beta thalassemia trait	Thalassemia minor: asymptomatic, mild microcytic hypochromic anemia
β+/β+ β+/β <sup>0</sup> βE/β+ βE/β <sup>0</sup>	Beta thalassemia intermedia	Variable severity Mild to moderate anemia Possible extramedullary hematopoiesis Iron overload
β°/β°	Beta thalassemia major (Cooley's Anemia)	Severe anemia Transfusion dependence Extramedullary hematopoiesis Iron overload

0: complete lack of chain

+: some expression of chain

: normal expression of chain

E: HbE

### Qualitative abnormalities

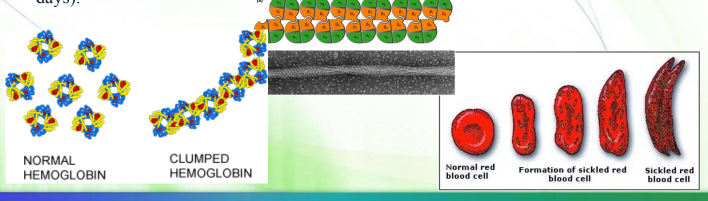
## Classification of molecular mutations

- Mutations in surface residues
  - Usually asymptomatic (e.g. HbE); an exception is HbS
- Mutations in internal residues
  - Often producing unstable hemoglobin and Heinz bodies and causing hemolytic anemia (e.g. Hb Hammersmith, Hb Constant Spring (Hb CS))
- Mutations stabilizing methemoglobin
  - Stabilizing heme-Fe+3; resulting in cyanosis
- Mutations at α1-β2 contacts
  - Altered oxygen affinity (mainly higher; a condition known as polycythemia)

### Sickle cell hemoglobin (HbS)

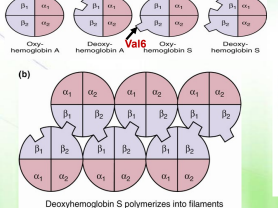
- It is caused by a change of amino acids in the 6th position of globin (Glu to Val).
- The hemoglobin is designated 2 s2 or HbS.
- The hemoglobin tetramers aggregate into arrays upon deoxygenation in the tissues.
- This aggregation leads to deformation of the red blood cell.

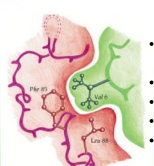
• It can also cause hemolytic anemia (life span of RBCs is reduced from 120 days to <20 days).



### How does the fiber form?

- Fiber formation only occurs in the deoxy or T-state.
- The mutated valine of 2 chain is protruded and inserts itself into a hydrophobic pocket on the surface of 1 chain.





#### Variables that increase sickling

Decreased oxygen pressure (high altitudes)

Increased pCO2

Decreased pH

Increased 2.3-BPG

Dehydration (why?)

### Sickle cell trait

- It occurs in heterozygotes (individuals with both HbA and HbS), who are clinically normal, but their cells sickle when subjected to low oxygen.
- Advantage: selective advantage from plasmodium falciparum that causes malaria. Why?

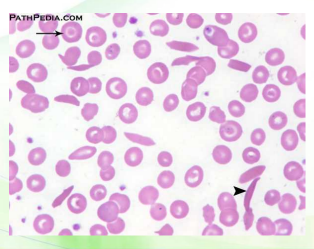


### Hemoglobin C (HbC

- (HbC) is also due to a change at the 6th position of globin replacing the glutamate with lysine (designated as c).
- This hemoglobin is less soluble than HbA so it crystallizes in RBCs reducing their deformability in capillaries.
- HbC also leads to water loss from cells leading to higher hemoglobin concentration.
- This problem causes only a minor hemolytic disorder.

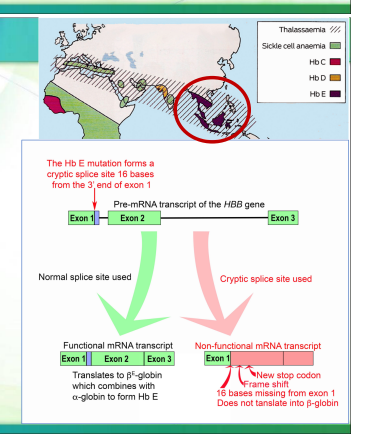
### HbSC disease

• Individuals with both c and s mutations have HbSC disease, a mild hemolytic disorder that may have no clinical consequences, but is clinically variable.



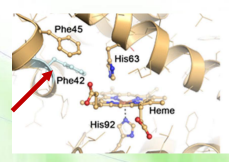
### Hemoglobin E

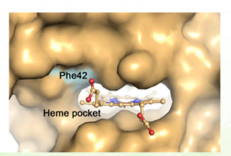
- It is common in Southeast Asia
- It has both quantitative and qualitative characteristics.
- It is caused by a point mutation in <u>codon</u> <u>26</u> that changes glutamic acid (GAG) to lysine (AAG) creating an <u>alternative</u> <u>RNA splice</u> site and a defective protein.
  - Individuals with this mutation make only around 60% of the normal amount of globin protein.
  - Mild disease but can be severe if coinherited with beta-thalassemia.



### Hb Hammersmith

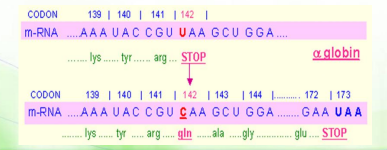
- Hb Hammersmith results from a point mutation that leads to formation of unstable hemoglobin and denaturation of the globin protein.
- The most common point mutation of Hb Hammersmith substitutes an internal <u>phenylalanine with a serine</u> within the globin, reducing the hydrophobicity of the heme-binding pocket, heme positioning, and oxygen binding affinity causing cyanosis.





### Hb Constant Spring (Hb CS)

- Hemoglobin Constant Spring (Hb CS) is an abnormal Hb caused by a mutation at the termination codon of the  $\alpha$ 2-globin gene leading to the production of unstable mRNA and protein products.
  - The anemia is usually moderate.
- Heterozygotes have the genotype ( $\alpha\alpha/\alpha\alpha CS$ ) and have  $\alpha$  -thalassemia trait phenotype.
- It is commonly found among Southeast Asian and Chinese people.
- If co-inherited with  $\alpha$ -thalassemia, it leads to an  $\alpha$  -thalassemia intermedia syndrome.



### Mutations at $\alpha 1$ - $\beta 2$ contacts

• Hb Cowtown: Substitution of <u>His146</u> (responsible for the Bohr Effect) to Leucine produces more hemoglobin in the R state (increased affinity).

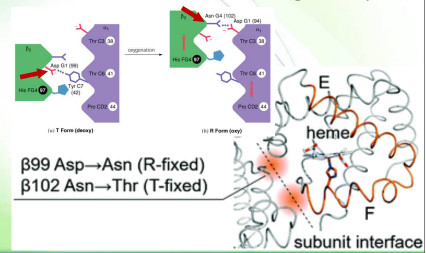
Elimination of hydrogen bonds between the chains can also alter the quaternary

structure:

#### Decreased cooperativity:

• Hb Yakima: stabilization of the R state (Asp G1 (99) to His).

• Hb Kansas: stabilization of the T state (Asn G4 (102) to Thr).



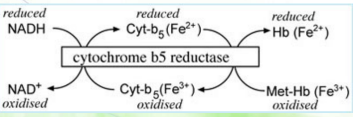
### **Altered Oxygen Transport**

### Methemoglobin (HbM)

• Oxyhemoglobin can undergo reversible oxygenation because its heme iron is in the reduced (ferrous, Fe+2) state.

• During oxygen release from heme, Fe+2 is oxidized to Fe+3, forming methemoglobin (HbM), except that the enzyme methemoglobin reductase reduces iron back.

• If not, a condition known as methemoglobinemia develops.

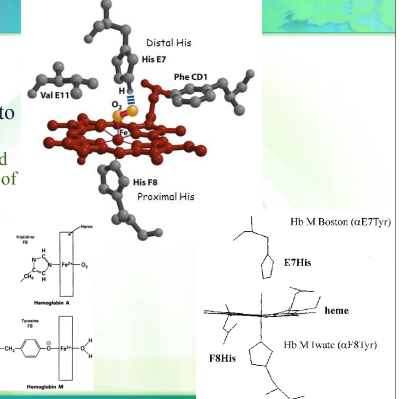


Methemoglobin reductase AKA NADH-Cytochrome b5 reductase

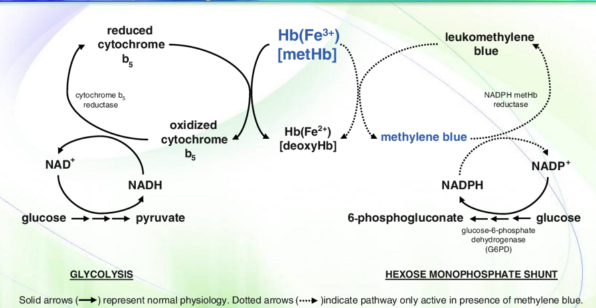


### Why HbM?

- Some mutant globins ( and ) bond with heme in such a way as to resist the reductase.
  - Hb Boston: distal histidine is mutated into a tyrosine resulting in oxidation of ferrous iron by tyrosine's oxygen.
  - HbM Iwate: proximal histidine is replaced by a tyrosine.
- A deficiency of the reductase enzyme.
- Certain drugs or drinking water containing nitrates.



### Treatment (methylene blue)



# Hereditary persistence of fetal hemoglobin

### (HPFH)

- Persons with HPFH continue to make HbF as adults.
- Because the syndrome is benign most individuals do not even know they carry a hemoglobin abnormality.
- Many HPFH individuals harbor large deletions of the and -coding region of the cluster.
- There is no deletion of the fetal globin genes.
- Think: treatment for -thalassemia!!!!

#### Switching from fetal to adult hemoglobin

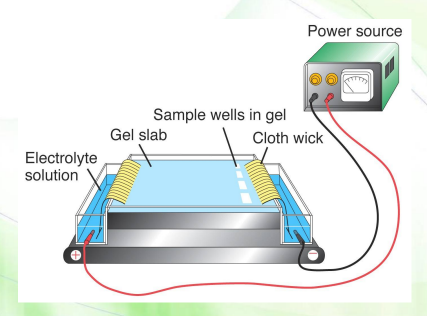
Xunde Wang & Swee Lay Thein ⊠

Nature Genetics 50, 478–480(2018) | Cite this article

**1102** Accesses  $\mid$  **5** Citations  $\mid$  **9** Altmetric  $\mid$  Metrics

The switch from fetal to adult hemoglobin relies on repression or silencing of the upstream  $\gamma$ -globin gene, but identification of the transcriptional repressors that bind to the sites at which a cluster of naturally occurring variants associated with HPFH (hereditary persistence of fetal hemoglobin) are found has been elusive. A new study provides mechanistic evidence for the direct binding of BCL11A and ZBTB7A, two previously identified  $\gamma$ -globin gene repressors.

### Hemoglobin Electrophoresis



### Mutation and migration

- Amino acid substitution in abnormal Hbs results in an overall change in the charge of the molecule.
- Therefore, Hb migration in a voltage gradient is altered.
- Electrophoresis of hemoglobin proteins from individuals is an effective diagnostic tool in determining if an individual has a defective hemoglobin and the relative ratios of the patient's hemoglobin pattern.

### Examples

- In Sickle Cell hemoglobin, replacement of a negatively-charged glu in the standard HbA by a neutral val in HbS results in a protein with a slightly reduced negative charge.
- In homozygous individuals, the HbA tetramer electrophoreses as a single band, and the HbS tetramer as another single band.
- Hemoglobin from a heterozygous individual (with both alleles) appears as two bands.
- Since HbC contains a lysine instead of the normal glutamate, HbC will travel even faster to the cathode.

### Results

- Lanes 1 and 5: Hb standards
- Lane 2: normal adult
- Lane 3: normal neonate
- Lane 4: homozygous HbS
- Lanes 6 and 8: Sickle cell trait
- Lane 7: HbSC disease

