Module II: BLOOD AND IMMUNITY:

Learning Objectives:

By the end of the CBL, students will be able to

- Correlate the changes in structure and function of the different types of Hemoglobin in health and in disease states.
- Correlate different Hb electrophoresis pattern
- Relate normal and abnormal haemostatic mechanisms with the coagulation cascade
- Relate blood and blood components transfusion and transfusional reactions

CLINICAL CASE

Moula Bux and his wife are sitting in the waiting room of their family physician, Dr. ABC. They are a young couple, who are first cousins and have been married for just two years and have a son, Sumar, whom they have brought in for a follow-up visit to the doctor. They are concerned because Sumar has recently been suffering from repeated infections. Although the boy seemed to be a happy and healthy newborn, he has grown increasingly listless over the past few months. He has lost much of his appetite and his complexion has become pale. The parents believe that their son has become anemic due to his poor diet; both Moula Bux and his wife themselves suffered from jaundice as babies.

The family is called in to see the physician, who, after exchanging pleasantries, reviews the boy’s symptoms and examines him. On examination he is short stunted and pale. He has prominent facial and frontal bones. His spleen is enlarged and palpable 4cm below his left costal margin. The blood reports reveal:

<table>
<thead>
<tr>
<th>Table 1. Blood Sample Results</th>
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</thead>
<tbody>
<tr>
<td>Hemoglobin (Hb)</td>
</tr>
<tr>
<td>MCV</td>
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<tr>
<td>MCHC</td>
</tr>
</tbody>
</table>
Poikilocytic red cells
(elliptocytes¹, schistocytes²,
target cells³, tear drop⁴,
spherocytes⁵ & hypochromic⁶)
usually present
in Thalassemia Major.