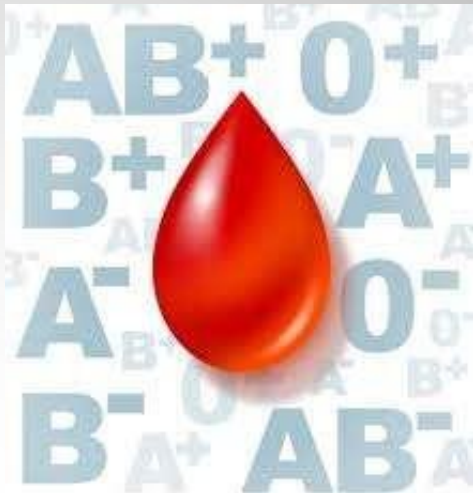


BLOOD GROUPING AND CROSS MATCHING



BLOOD GROUP

- Blood grouping is based on type of antigen present on the red blood cells.
- There are more than 300 blood group systems but ABO and Rh(Rhesus) are of importance from clinical point of view.
- Other blood group systems are MNS , Lutheran , Kell , Lewis , Duffy , Kidd etc.

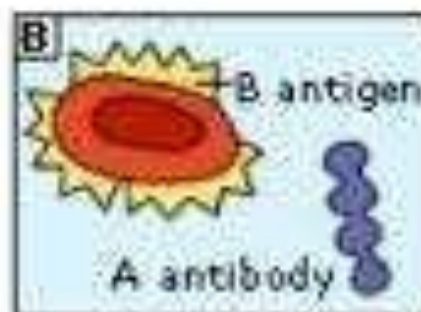
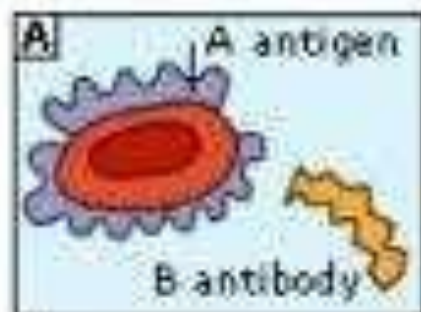
ABO SYSTEM

- Discovered by Karl Landsteiner in 1900.
- The red cells contain different types of Antigen(Agglutinogen) while plasma contains antibody(Agglutinins)
- Genes that control the system are present on chromosome 9

LANDSTEINER'S LAW

If an antigen(Ag) is present on a patient's RBC, the corresponding antibody(Ab) should not be present in patient's plasma under normal condition

ABO blood grouping system

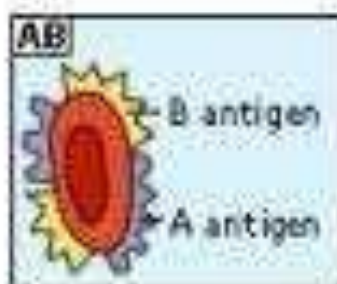


Blood group A

If you belong to the blood group A, you have A antigens on the surface of your RBCs and B antibodies in your blood plasma.

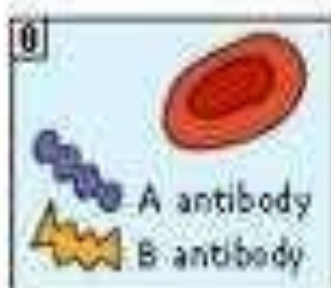
Blood group B

If you belong to the blood group B, you have B antigens on the surface of your RBCs and A antibodies in your blood plasma.



Blood group AB

If you belong to the blood group AB, you have both A and B antigens on the surface of your RBCs and no A or B antibodies at all in your blood plasma.



Blood group O

If you belong to the blood group O, you have neither A or B antigens on the surface of your RBCs but you have both A and B antibodies in your blood plasma.

Major ABO Blood Group

ABO Group	Antigen Present	Antigen Missing	Antibody Present
A	A	B	Anti-B
B	B	A	Anti-A
O	None	A and B	Anti-A&B
AB	A and B	None	None

Methods of blood grouping:

1) Slide method

2) Tube method

Tube method - better method
but takes longer

Sample in tube with antiserum ---
incubate it --- centrifuge it ---examine it
macroscopically and microscopically for
agglutination.

SLIDE METHOD

REQUIREMENTS:

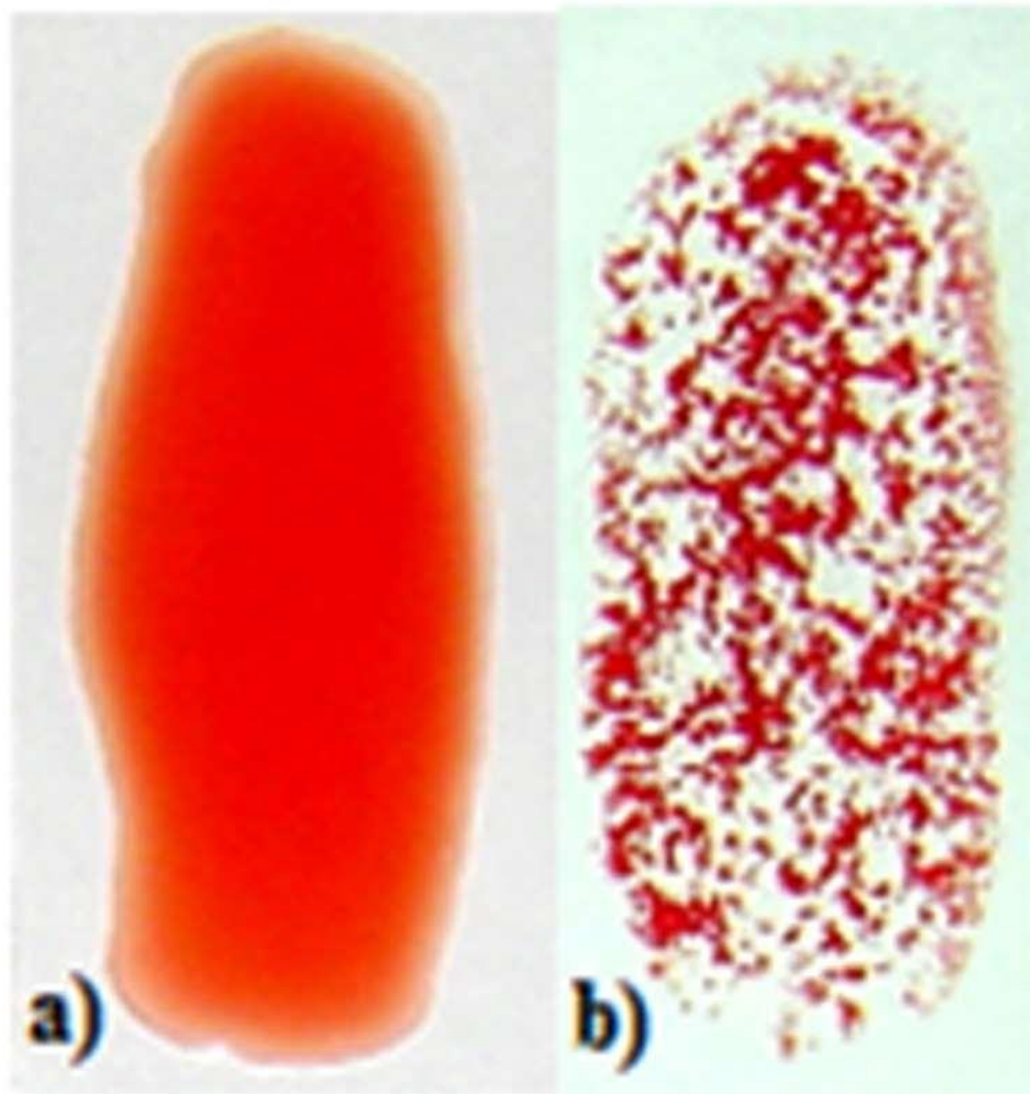
- 1) 3 slides
- 2) Antisera A , B
- 3) Blood samples

PROCEDURE:

- 1) Take 2 clean slides and mark them 1, 2 .
- 2) Put one drop of antisera A on slide 1 , one drop of antisera B on slide 2.
- 3) Add one drop of blood to each and mix well with stick
- 4) Wait for 5 min and observe.

OBSERVATION:

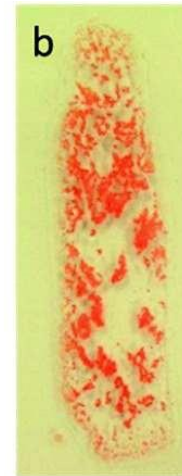
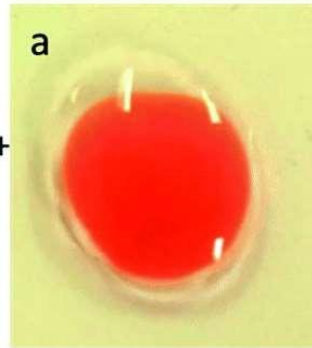
- If any agglutination occurs it is visible with naked eyes as dark reddish clumps of different sizes.
- If agglutination is minimal it can be confirmed by examining it under microscope.



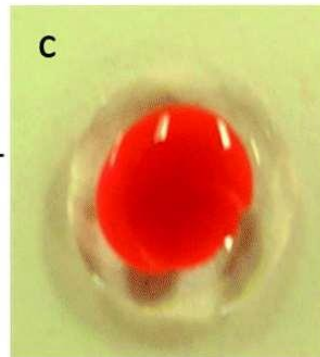
INTERPRETATION:

- 1) Agglutination with antisera A not with antisera B - group A
- 2) Agglutination with antisera B not with antisera A - group B.
- 3) Agglutination with both antisera A and B - group AB
- 4) No agglutination in any slide - group O

Blood sample A+
interacted with
antibody-A



Blood sample A+
interacted with
antibody B



Universal donor - blood group O as no Ag so no agglutination.

Universal recipient - blood group AB as both A and B Ags present so agglutination occurs in both as no Abs present in serum.

Rh TYPING

HISTORY:

- 1939 - Levine and Stetson defined D antigen (Rh factor)

- 1949 - Landsteiner and Weiner discovered anti Rh (named after Rhesus monkey)

Rh TYPING

- Rh blood group system is second in significance after ABO system.
- Genes that control the system are present on chromosome 1
- Consists of over 50 related Ags.
- Important genes are D,C,E,c,e.

- All Rh antigens are controlled by 2 genes -
- RHD gene - determines expression of D
- RHCE - encodes for C, c and E, e.

- RhD is a strong antigen (immunogenic) and other antigens are less antigenic than D and are of less clinical significance.
- Therefore, in practice Rh negative and Rh positive depends on presence of D antigen on the surface of red cells which is detected by strong anti-D serum.
- Occasionally, Anti - D, C, E, c, e may develop in case of pregnancy or transfusion.

□ Rh positive

There is presence of D antigen.

These individuals constitute 80% of population.

Rh negative:

There is absence of D antigen. These individuals constitute 17% of population.

Cc and Ee antigen:

These are weak antigens and therefore risk of sensitisation is less than that of D antigen.

Rh antibody:

- Unlike ABO system there is no naturally occurring antibodies against Rh antigens in Rh negative individuals.

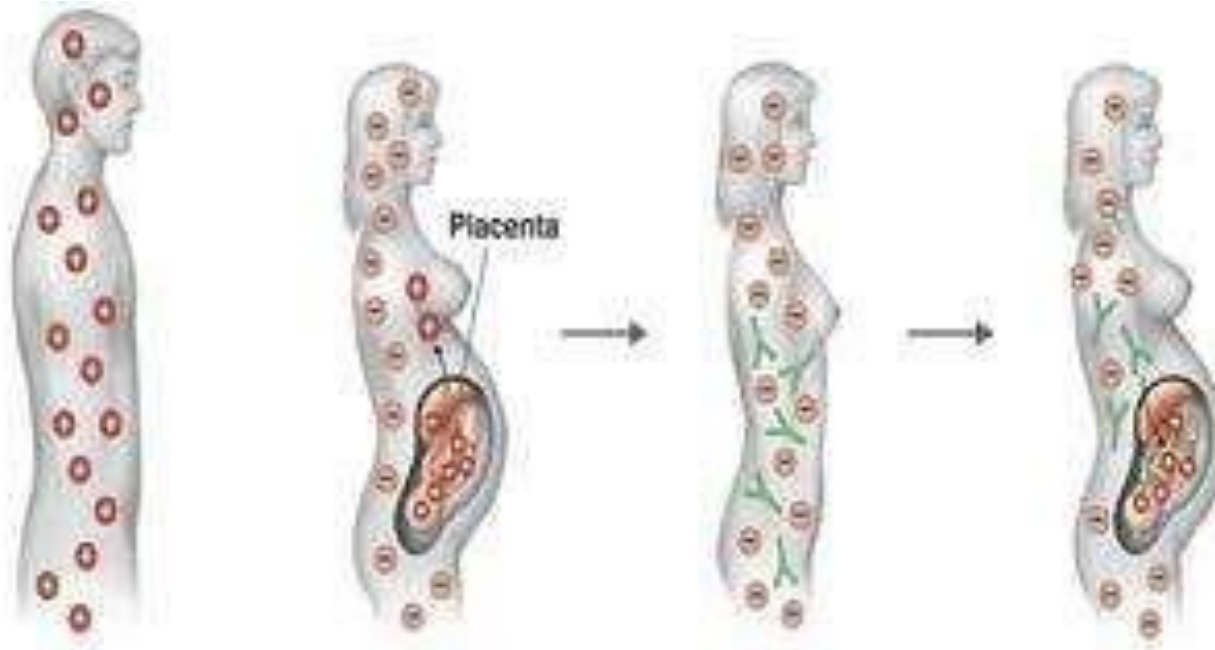
Immune Abs:

- Rh Abs develop against Rh Ag after exposure to Rh Ags following transfusion or pregnancy.
- But can be detected by enzyme treatment or coomb test(antiglobulin test)

SIGNIFICANCE:

- Rh incompatibility results in haemolytic tranfusion reaction.
- Haemolytic disease of newborn.

ERYTHROBLASTOSIS FETALIS



1 Rh⁺ father.

2 Rh⁻ mother carrying her first Rh⁺ fetus. Rh antigens from the developing fetus can enter the mother's blood during delivery.

3 In response to the fetal Rh antigens, the mother will produce anti-Rh antibodies.

4 If the woman becomes pregnant with another Rh⁺ fetus, her anti-Rh antibodies will cross the placenta and damage fetal red blood cells.

TECHNIQUES:

1) slide method

2) Tube method

SLIDE METHOD:

- Place one drop of anti D on slide.
- Add one drop of blood and mix well with stick
- Wait for 5 min and observe.

RESULT:

- Agglutination indicates Rh positive blood samples.

IMPORTANCE OF BLOOD GROUPING AND Rh TYPING:

- In blood transfusion
- Haemolytic disease of newborn.
- Paternity dispute
- Medicolegal issues
- Immunology, genetics, anthropology
- Susceptibility to various disease (blood group O - peptic ulcer
Blood group A - gastric ulcer)

CROSS MATCHING

- Also known as compatibility testing.
- It is the most important test before a blood transfusion is given.
- The primary purpose of cross matching is to detect ABO incompatibilities between donor and recipient.
- This is carried out to prevent transfusion reactions by detecting Abs in recipient's serum.

□ Two main functions of cross matching test:

1) It is a confirm ABO compatibility between donor and recipient.

2) It may detect presence of irregular Ab in patient's serum that will react with donor RBCs.

- Cross matching test can be
 - 1) major
 - 2) minor

MAJOR CROSS MATCH TEST:

Mixing the patient's plasma with donor RBCs.

MINOR CROSS MATCH TEST:

- mixing the donor's plasma with patient's RBCs.

SCREENING TESTS BEFORE BT:

- Malaria
- Syphilis
- HBV
- HCV
- HIV



LOCOMO

THANK YOU