

# ABO discrepancies Recognition and Resolution

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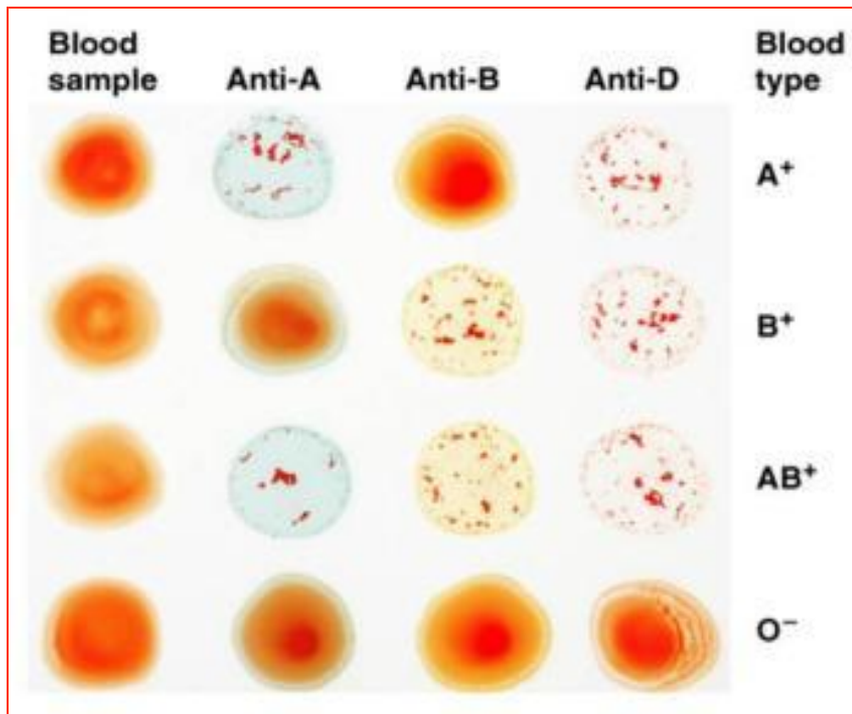
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# Recognition and resolution of ABO discrepancies

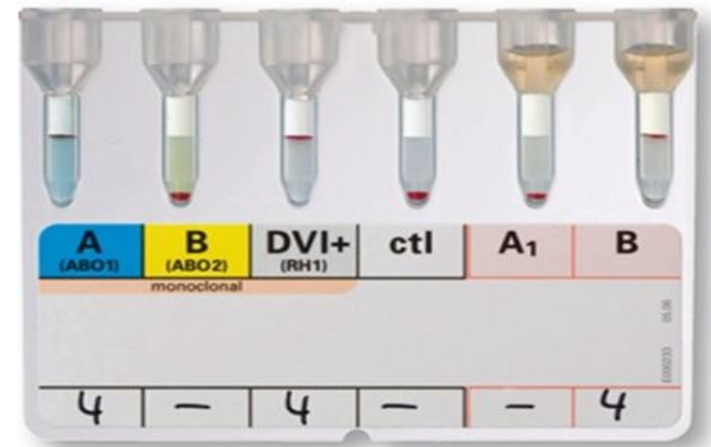
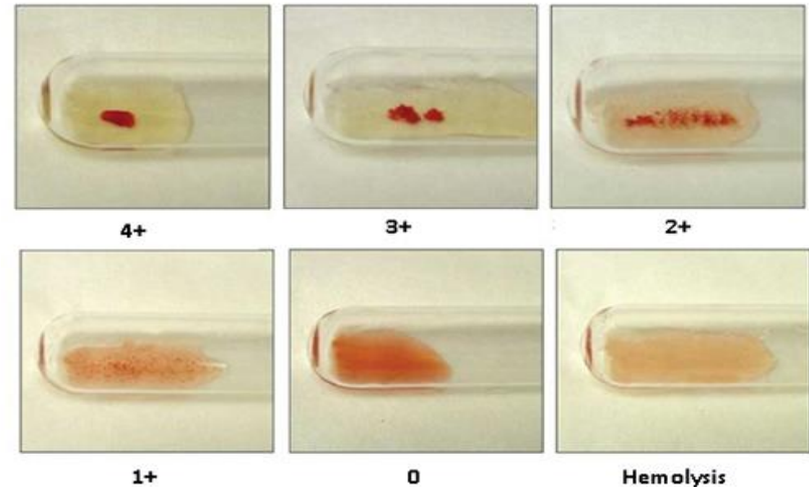
- Discrepancy: results of Forward do not agree with Reverse grouping



	Patient RBCs	Patient RBCs	Patient Plasma	Patient Plasma
	Anti-A Antibody	Anti-B Antibody	Type A RBCs	Type B RBCs
O				
A				
B				
AB				

# Recognition and resolution of ABO discrepancies

- Discrepancies may be indicated when following observations are noted:
  1. Agglutination strengths of reactions are weaker than expected
    - ✓ agglutination reactions in Forward grouping → are 3+ to 4+
    - ✓ agglutination reactions in Reverse grouping → are 2+ to 4+
  2. Expected reactions in Forward & Reverse grouping are missing
    - ✓ e.g., group O individual → missed one or both reactions in serum testing with reagent A<sub>1</sub> and B cells
  3. Extra reactions are noted in either Forward or Reverse grouping



# Recognition and resolution of ABO discrepancies

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- The first step in the resolution → identify the source of problem: .
  1. technical problems
  2. sample-related problems

# Technical errors in ABO typing

□ **Technical errors** can be classified into several categories:

- ① **Identification and documentation errors** ② **Reagent and equipment errors** ③ **SOP errors**

## Practical Application: Guidelines for Investigating ABO Technical Errors

### Identification or Documentation Errors

Correct sample identification on all tubes  
Results are properly recorded  
Interpretations are accurate and properly recorded

### Reagent or Equipment Errors

Daily quality control on ABO typing reagents is satisfactory  
Inspect reagents for contamination and hemolysis  
Centrifugation time and calibration are confirmed

### Standard Operating Procedure Errors

Procedure follows manufacturer's directions  
Correct reagents were used and added to testing  
Red blood cell suspensions are at the correct concentration  
Cell buttons are completely suspended before grading the reaction

# Technical errors in ABO typing

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- When a **technical error** is discovered and corrected → ABO discrepancy can be quickly resolved with repeated testing
  - ✓ if discrepancy still exists ⇨ possibility of a **problem related to sample** itself (related to patient or donor) should be considered.

# Sample-related ABO discrepancies

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- can be divided into 2 groups:
  1. Problems that affect **RBC testing** (Forward grouping)
  2. Problems that affect **serum (plasma) testing** (Reverse grouping)
  
- A logical approach is ⇨ determine the **side** of ABO test (RBC testing or serum testing)
  - ✓ **strengths of agglutination** reaction (in both Forward & Reverse grouping) → is a key point
  
- ❖ the most commonly encountered ABO discrepancies in Lab ⇨ relating to weak or missing ABO Abs in **serum testing**

# Sample-related ABO discrepancies

## Overviews of ABO Discrepancies

### PROBLEMS WITH RED CELL TESTING

Extra antigens  
Group A with acquired B antigen  
B(A) phenotype  
Polyagglutination  
Rouleaux  
Hematopoietic progenitor cell transplants

Missing or weak antigens  
ABO subgroup  
Pathologic etiology  
Transplantation

Mixed-field reactions  
Transfusion of group O to group A, B, or AB  
Hematopoietic progenitor stem cell transplants  
A<sub>3</sub> phenotype

### PROBLEMS WITH SERUM/PLASMA TESTING

Extra antibodies  
A subgroups with anti-A<sub>1</sub>  
Cold alloantibodies  
Cold autoantibodies  
Rouleaux  
IVIG

Missing or weak antibodies  
Newborn  
Elderly  
Pathologic etiology  
Immunosuppressive therapy for transplantation

# Sample-related ABO discrepancies

## □ Discrepancies Associated with **Red Cell Testing** (forward grouping):

1. extra antigens present
2. missing or weak antigens
3. mixed-field reactions

### Extra antigens

Group A with acquired B antigen

B(A) phenotype

Polyagglutination

Rouleaux

Hematopoietic progenitor cell transplants

### Missing or weak antigens

ABO subgroup

Pathologic etiology

Transplantation

### Mixed-field reactions

Transfusion of group O to group A, B, or AB

Hematopoietic progenitor stem cell transplants

A<sub>3</sub> phenotype

# Extra antigens present

## – Acquired B Antigen

<i>Group A with Acquired B Antigen</i>			
ABO Testing Results			
Patient Red Cells with		Patient Serum with Reagent Red Cells	
Anti-A	Anti-B	A <sub>1</sub>	B
4+	1+	0	4+

### □ EVALUATION OF ABO TESTING RESULTS

1. agglutination of patient's RBCs with anti-A is strong (4+).
2. agglutination of patient's RBCs with anti-B is 1+ ⇨ weaker than usually expected (3+ to 4+) → in RBC testing result → group AB
3. The results of serum testing reactions → group A.

### □ CONCLUSION:

- ✓ group A with acquired B

# Extra antigens present

## – Acquired B Antigen

### □ Background information

- ✓ only in group A<sub>1</sub> individuals with **diseases of lower GI tract** → cancers of colon and rectum, intestinal obstruction, or gram-negative septicemia
- ✓ the most common mechanism:
  - ❖ a bacterial deacetylating enzyme → alters **A ID-sugar** (N-acetylgalactosamine) by removing acetyl group → resulting sugar (galactosamine) resembles B ID-sugar (D-galactose) ⇨ cross-reacts with many anti-B reagents
  - ❖ the observation was linked to → use of ES-4 mAb anti-B clone at pH 6.5–7.0
    - if formulation of clone acidified to pH 6.0 → acquired B antigen not observed

# Extra antigens present

## – Acquired B Antigen

### □ RESOLUTION OF ABO DISCREPANCY

1. Determine the patient's **diagnosis** and **transfusion history**.

- ✓ First step: obtain more information about patient ⇨ may provide additional clues about cause of ABO discrepancy

2. Test **patient's serum** against **autologous RBCs**.

- ✓ In acquired B Ag ⇨ Anti-B in patient's serum, does not agglutinate autologous RBCs

3. Test RBCs with:

- ✓ additional monoclonal anti-B reagents ⇨ from other manufacturers (that not to react with acquired B Ag)
- ✓ a source of human polyclonal anti-B

➤ for transfusion purposes ⇨ patients should receive RBCs of group A

# Extra antigens present

## – B(A) Phenotype

<i>B(A) Phenotype</i>			
ABO Testing Results			
Patient Red Cells with		Patient Serum with Reagent Red Cells	
Anti-A	Anti-B	A <sub>1</sub>	B
1+	4+	4+	0

### □ EVALUATION OF ABO TESTING RESULTS

1. agglutination of patient's RBCs with anti-A is weak (1+).
2. agglutination of patient's RBCs with anti-B is strong (4+) ⇒ results of serum testing → are typical of a group B individual.

### □ CONCLUSION:

- ✓ a group B with an extra reaction with anti-A in RBC testing ⇨ a possible **B(A) phenotype**

# Extra antigens present

## – B(A) Phenotype

### □ BACKGROUND INFORMATION

- B(A) phenotype → observed as a result of ↑sensitivity of mAb reagents
  - ✓ These reagents ⇨ can detect trace amounts of A or B antigens that are nonspecifically transferred by glycosyltransferase enzymes
- B gene enzyme (Galactosyltransferase) → **transfers trace amounts** of N-acetylgalactosamine (ID-sugar for A-Ag) + D-galactose (ID-sugar for B-Ag) to H-Ag
  - ✓ trace amounts of A-Ags ⇨ are detected with certain mAb reagents
  - ❖ a similar mechanism → can cause A(B) phenotype

### □ RESOLUTION OF ABO DISCREPANCY

1. Determine patient's **diagnosis** and **transfusion history**.
2. Test RBCs with:
  - ✓ additional mAb anti-A reagents from other manufacturers or
  - ✓ a source of human polyclonal anti-A

# Missing or Weakly Expressed Antigens

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- In this category of ABO discrepancies → RBCs demonstrate **weaker** or **no reactions** with anti-A and anti-B reagents
  
- Phenomena associated with this category include:
  1. **ABO subgroups**
  2. **Weakened A and B** antigen expression → in leukemia or Hodgkin's disease

# Missing or Weakly Expressed Antigens

## – Subgroup of A

<i>Subgroup of A</i>			
ABO Testing Results			
Patient Red Cells with		Patient Serum with Reagent Red Cells	
Anti-A	Anti-B	A <sub>1</sub>	B
0	0	0	3+

### □ EVALUATION OF ABO TESTING RESULTS

1. No agglutination of patient's RBCs with anti-A and anti-B reagents ⇒ patient appears to be a group O phenotype.
2. The results of serum testing → typical of a group A individual

### □ CONCLUSION: reactions are characteristic of **a missing Ag** (in RBC testing):

- ✓ Serum testing results → are those expected in a group A individual.
- ✓ Anti-A (found in group O individuals) → is absent in serum testing.

# Missing or Weakly Expressed Antigens

## – Subgroup of A

### □ BACKGROUND INFORMATION

- ✓ weak or missing reactions with antiserum reagents → correlate with subgroups of A and B:
  - ✓ Subgroups of A: are <1% of group A population
  - ✓ subgroups of B: are even rarer

### □ RESOLUTION OF ABO DISCREPANCY

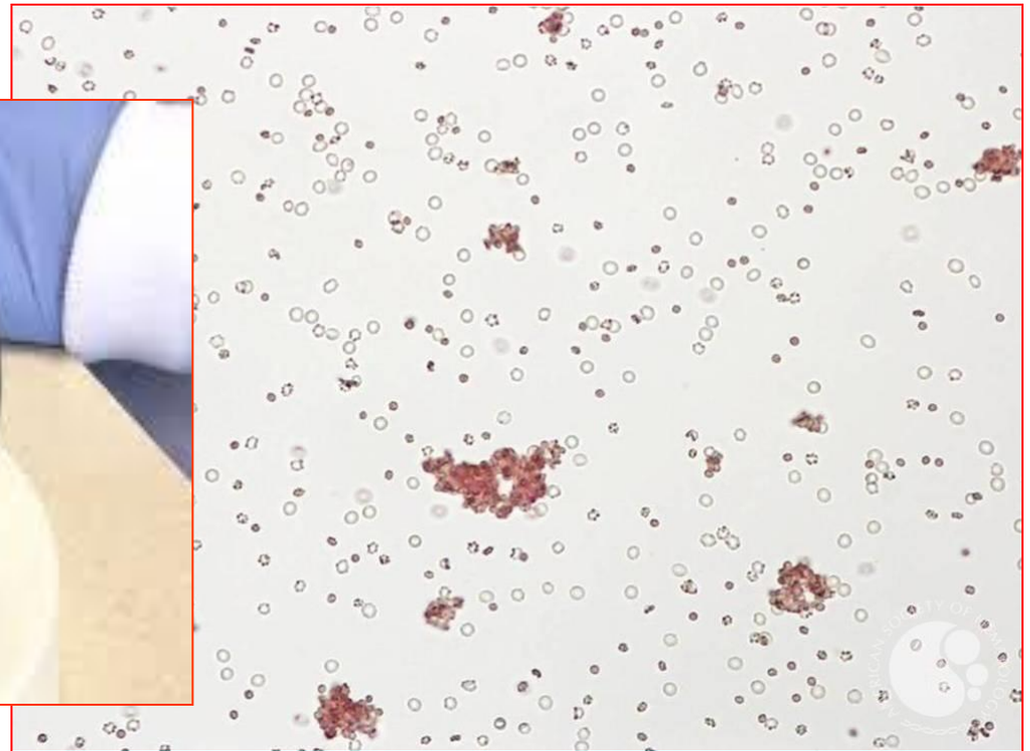
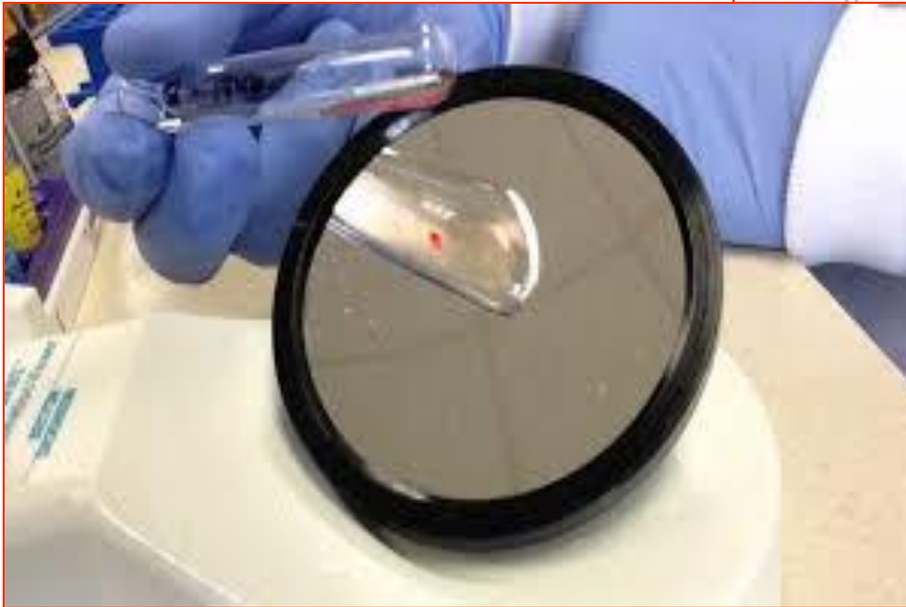
1. Determine the patient's **diagnosis** and **transfusion history**.
2. **Repeat RBC testing** with:
  - ✓ extended incubation times ⇨ may enhance Ag-Ab reaction
  - ✓ human polyclonal anti-A,B or monoclonal blend anti-A,B

Additional Testing Results	
	Anti-A,B
Patient red cells	1+
Conclusion: Probable subgroup of A	

Additional Testing Results	
	Anti-A,B
Patient red cells	0
Next Step: Perform adsorption and elution studies with anti-A; these studies assist in determining the presence of A antigens on the patient's red cells	

# Mixed-Field (MF) Reactions

- MF reactions → can occur with either anti-A or anti-B reagents:
  - ✓ a MF-reaction contains: agglutinates + a mass of un-agglutinated RBCs
  - ✓ MF-reaction → is due to presence of 2 distinct cell populations



# Mixed-Field (MF) Reactions

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- MF-reactions can occur in:
  1. transfusion of group O RBCs to group A, B, or AB individuals,
  2. recipients of HPC transplants,
  3. individuals with A<sub>3</sub> phenotype,
  4. patients with Tn-polyagglutinable RBCs

# Mixed-Field Reactions

## Group B Patient Transfused with Group O RBCs

### ABO Testing Results

Patient Red Cells with		Patient Serum with Reagent Red Cells	
Anti-A	Anti-B	A <sub>1</sub>	B
0	2+mf	4+	0

#### ❑ EVALUATION OF ABO TESTING RESULTS

1. The strength of agglutination reaction with anti-B → is weaker than expected for group B individuals ⇨ Mixed-field reaction (a 2+ agglutination with a sufficient number of un-agglutinated cells)
2. The results of serum testing → are typical of a group B individual.

#### ❑ CONCLUSION: a group B individual possibly transfused with group O RBCs

# Mixed-Field Reactions

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## □ BACKGROUND INFORMATION

- ✓ In certain situations, ABO-identical RBC products might not be available for transfusion → group O RBC products are transfused
  - ❖ If many group O RBC units are transfused ⇨ MF- reactions may appear in ABO red cell testing.

## □ RESOLUTION OF ABO DISCREPANCY

- 1) Determine the patient's **diagnosis** and recent **transfusion history**.
- 2) Determine whether the patient is a recent **HPC** recipient.
- 3) Investigate **pre-transfusion ABO phenotype history**, if possible.

# ABO Discrepancies Associated with Reverse grouping

- ABO discrepancies that affect **serum testing** include:
  1. presence of additional Abs (other than anti-A and anti-B)
  2. absence of expected ABO Ab reactions
- ✓ The most commonly encountered ABO discrepancies → absence of expected ABO Ab reactions.

## PROBLEMS WITH SERUM/PLASMA TESTING

### Extra antibodies

- A subgroups with anti-A<sub>1</sub>
- Cold alloantibodies
- Cold autoantibodies
- Rouleaux
- IVIG

### Missing or weak antibodies

- Newborn
- Elderly
- Pathologic etiology
- Immunosuppressive therapy for transplantation

# Additional Antibodies in Reverse grouping

- detection of anti-A<sub>1</sub>

## Group A<sub>2</sub> with Anti-A<sub>1</sub>

ABO Testing Results			
Patient Red Cells with		Patient Serum with Reagent Red Cells	
Anti-A	Anti-B	A <sub>1</sub>	B
4+	0	2+	4+

### ❑ EVALUATION OF ABO TESTING RESULTS

1. agglutination pattern with anti-A and anti-B reagents → typical of a group A individual.
2. results of serum testing with A<sub>1</sub> and B red cells → indicate a group O individual

### ❑ CONCLUSION

- ✓ an extra reaction in serum testing with reagent A<sub>1</sub> RBCs (2+) ⇨ possible explanations include: anti-A<sub>1</sub>, cold allo-Ab, cold auto-Ab, or rouleaux.

# Additional Antibodies in Reverse grouping

## - detection of anti-A<sub>1</sub>

### ❑ RESOLUTION OF ABO DISCREPANCY

1. Determine the patient's **diagnosis** and **transfusion history**.
2. Test the patient's RBCs with **anti- A<sub>1</sub> lectin**

Additional Testing Results	
Patient Red Cells Tested with Anti-A <sub>1</sub> Lectin	Conclusion
0	Subgroup of A; suspect anti-A <sub>1</sub> antibody

3. Test the patient's serum with 3 examples of **A<sub>1</sub> and A<sub>2</sub> reagent RBCs** ⇨ to confirm presence of **anti- A<sub>1</sub> antibody**

Additional Testing Results					
Patient Serum Tested with					
A <sub>1</sub> Cells	A <sub>1</sub> Cells	A <sub>1</sub> Cells	A <sub>2</sub> Cells	A <sub>2</sub> Cells	A <sub>2</sub> Cells
2+	2+	2+	0	0	0

### ❑ CONCLUSION: ABO discrepancy resulting from **group A<sub>2</sub> with anti-A<sub>1</sub>**.

- ✓ Agglutination is observed with A<sub>1</sub> RBCs → providing evidence for anti-A<sub>1</sub>.  
Anti- A<sub>1</sub> may be present in 1-8% of group A<sub>2</sub> phenotype.

# Additional Antibodies in Reverse grouping

## - Rouleaux

<i>Rouleaux</i>			
ABO Testing Results			
Patient Red Cells with		Patient Serum with Reagent Red Cells	
Anti-A	Anti-B	A <sub>1</sub>	B
4+	4+	2+	2+

### ❑ EVALUATION OF ABO TESTING RESULTS

1. Strong agglutination reactions are observed in RBC testing → consistent with expected results of group AB.
2. Serum testing results → consistent with those of a group O.

### ❑ CONCLUSION

- ❖ possibility of **extra reactions in serum** testing → because of allo-Ab, auto-Ab, or rouleaux.

# Additional Antibodies in Reverse grouping

## - Rouleaux

### □ BACKGROUND INFORMATION

- ❖ Rouleaux can produce false-positive agglutination:
  - ✓ RBCs resemble stacked coins under microscopic examination.
  - ✓ ↑ concentrations of serum proteins → can spontaneous rouleaux of RBCs.
  - ✓ Diseases associated with rouleaux → MM and WM
- ❖ problems with rouleaux:
  - 1) extra reactions in serum testing in ABO phenotyping
  - 2) extra reactions in ABO RBC typing (if unwashed RBC suspensions are used)

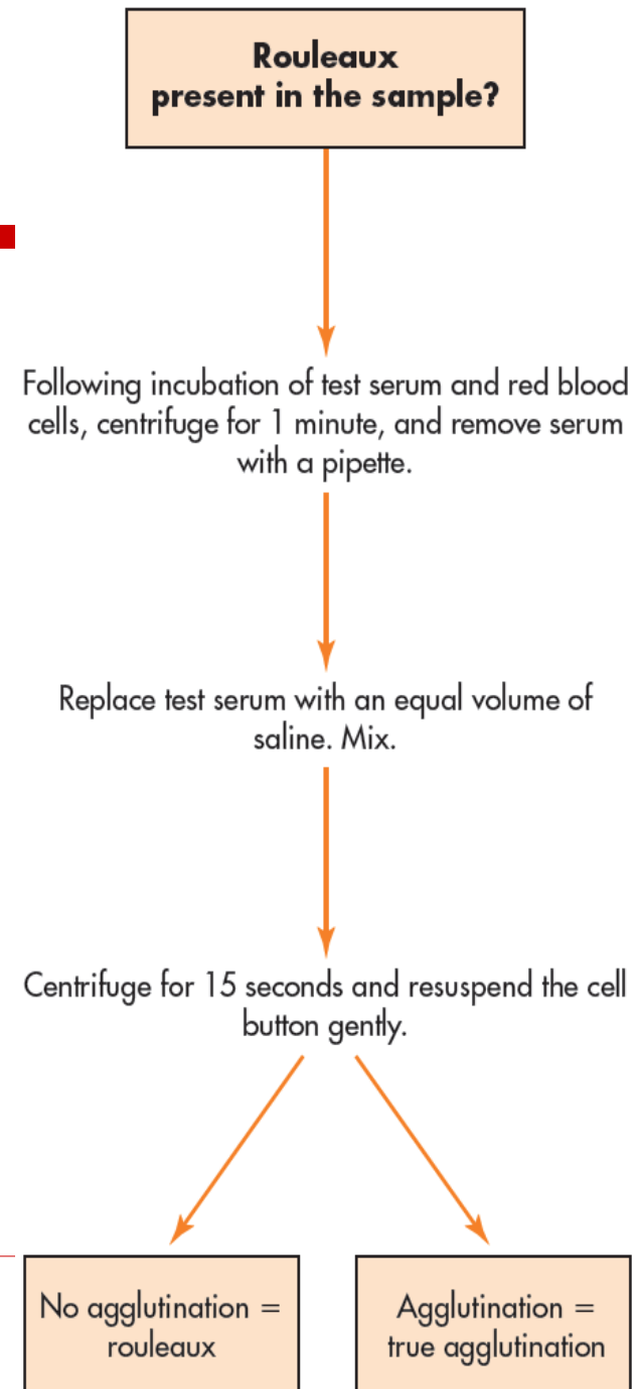


# Additional Antibodies in Reverse grouping

## - Rouleaux

### RESOLUTION OF ABO DISCREPANCY

1. Determine the patient's **diagnosis** and **transfusion history**.
2. **Wash** RBC suspension and repeat the phenotyping.
3. Perform **saline replacement technique** ☞ for distinguish true agglutination from rouleaux



## Missing or Weak ABO Antibodies in Serum/Plasma Testing

- ABO Abs may be missing or weakened in certain patient-related situations.

### *Missing or Weak ABO Antibodies in Serum or Plasma Testing*

ABO Testing Results			
Patient Red Cells with		Patient Serum with Reagent Red Cells	
Anti-A	Anti-B	A <sub>1</sub>	B
0	0	0	0

- EVALUATION OF ABO TESTING RESULTS

1. agglutination pattern with anti-A and anti-B reagents → group O.
2. results of serum testing with reagent A<sub>1</sub> and B red cells → group AB.

- CONCLUSION

- ✓ missing serum reactions with reagent A<sub>1</sub> or B cells.

# Missing or Weak ABO Antibodies in Serum or Plasma Testing

## □ BACKGROUND INFORMATION

- ❖ investigation of patient's **history** (including **age**, **diagnosis**, **Ig levels**) ⇨ provides clues to missing reactions in serum testing.
  - ✓ patient's **age** is an important factor ⇨ ↓ concentrations of ABO Abs in newborns and elderly adults.
  - ✓ patient's **diagnosis** is essential ⇨ ↓ Ig levels are associated with several pathologic states.
- ❖ patient's diagnosis, Ig levels and serum protein electrophoretic patterns → are helpful data in identification & resolution of cause for this ABO discrepancy.
- ❖ for cord blood and infants <4 months → only Forward grouping

# Missing or Weak ABO Antibodies in Serum or Plasma Testing

## ❑ RESOLUTION OF ABO DISCREPANCY

1. Determine patient's **diagnosis**, **age**, and **Ig levels**, if available.
2. **Incubate serum testing** for 15 minutes at RT → then centrifuge and examine for agglutination ⇨ **incubation step** often solves the problem.
3. If the results are still negative → place serum testing at **4° C** for 5 minutes with an autologous control.
  - ✓ The autologous control validates the test by ensuring that positive reactions are not attributable to a cold autoantibody.

### Interpretation of Additional Testing Results

4° C	A <sub>1</sub> Red Cells	B Red Cells	Autologous Red Cells	Conclusion
Patient serum	Pos	Pos	Neg	Group O
Patient serum	Pos	Pos	Pos	Cold autoantibody

Thank You

