MONOCLONAL ANTI–D BLEND (IgG/IgM)
BLOOD GROUPING SERA

For use by slide, tube and microplate methods

INTRODUCTION
First described in 1939, the RhD antigen is surpassed in importance only by the antigens of the ABO blood group system. Transfusion of RhD positive blood to a RhD negative recipient or failure to administer prophylactic anti-D to an RhD negative woman following birth of a RhD positive baby, can result in the production of anti-D. Consequently, establishing the correct RhD group is fundamental to safe transfusion practice. Certain individuals exhibit a quantitative reduction in the expression of their RhD antigen and are categorised as weak D (D\text{u}). Others display a qualitative variation in D antigen expression and are referred to as D category or partial D. Weak D individuals may also be partial D.

The recent availability of potent high quality IgM monoclonal anti D reagents and a greater awareness of the clinical importance of partial D phenotypes, especially D\text{VI}, has resulted in a change to RhD testing policies in the UK.

This reagent is designed to be used in conjunction with Anti D IgM reagent to optimise the detection of weak D and D variant red cells.

PRINCIPLE OF THE SERA AND TEST PROCEDURE
The test procedures recommended for the use of this sera are based upon the agglutination (clumping) of red blood cells carrying the D antigen in the presence of an IgM Anti–D antibody. The sera has been optimised for use as supplied by the recommended techniques without further dilution or additions.

COMPOSITION
The main component of this reagent is derived from the In Vitro culture of the human/mouse heterohybridomas LDM3 which secretes IgM anti-D and ESD1 which secretes IgG anti-D. The formula also contains less than 0.1% azide and EDTA

PRECAUTIONS
1. All blood products should be treated as potentially infectious. The source used to produce this sera has been tested and found to be negative for HIV and HCV antibodies and HBsAg in microbiological tests required by the UK. No known regime of testing can completely guarantee that any product derived from human blood is incapable of transmitting infectious agents. Care should be exercised in the use and the disposal of the container and its contents.
2. The sera contains less than 0.1% w/v sodium azide. Sodium azide may be toxic if ingested and may react with lead and copper plumbing to form highly toxic explosive salts. On disposal flush with large quantities of water.
3. The reagent should be stored at 2-8°C at all times. The product should be clear. Turbidity may indicate bacterial contamination. The product should not be used beyond the stated expiry date.
4. For in vitro diagnostic use only.
5. UNDER NO CIRCUMSTANCES SHOULD THE PRODUCT BE DILUTED PRIOR TO USE.

ADDITIONAL REAGENTS REQUIRED
PBS pH 7.0 +/- 0.2
LISS, Low Ionic Strength Saline
Reagent red cells for use in RhD grouping
RhD reagent control
Coombs control (Anti D)
Polyspecific Anti Human Globulin Reagent

ADVICE TO USERS
It is recommended that a positive control (ideally R\text{r} cells) and a negative control (rr cells) should be tested in parallel with each batch of tests. It is not required to use a diluent control in parallel with all tests using this sera. Only in typing the red cells of patients known to have auto-antibodies or protein abnormalities or apparently low grade D\text{u} is use of a diluent control recommended. This should be tested in parallel with the sera.

TEST PROCEDURES
General Information
This reagent has been standardised for use by the techniques described below and therefore suitability for use in other techniques cannot be guaranteed. Users are advised to confirm reagent suitability before using alternative techniques.
RECOMMENDED TECHNIQUES

1. Slide Techniques
   1.1 Prepare a 30–45% suspension of red blood cells in homologous plasma/serum or PBS pH 7.0 may be used.
   1.2 To one volume of sera on a clean, labelled glass slide add an equal volume of the test red cell suspension.
   1.3 Mix well by rocking slide for 30 seconds and incubate the test at room temperature for 5 minutes with occasional mixing.
   1.4 Read macroscopically for agglutination. This may be facilitated by the use of a diffuse light source.
   Note: Do not confuse drying effects with weak positive results.

2. Tube Technique – Immediate spin
   2.1 To one volume of sera in a labelled test tube, add an equal volume of a 2–3% suspension of test red blood cells in PBS pH 7.0 or 1.5 – 2% in LISS.
   2.2 Mix the test well.
   2.3 Centrifuge immediately at 1000g for 10 seconds or for a suitable alternative force and time.
   2.4 Gently agitate the tube to dislodge the cells and examine macroscopically for agglutination.

3. Tube Technique – LISS
   3.1 To one volume of sera in a labelled test tube add an equal volume of a 1.5-2% suspension of test red blood cells in LISS.
   3.2 Mix the test well and incubate at 37°C for 15-20 minutes.
   3.3 Centrifuge at 1000g for 10 seconds or for a suitable alternative force and time.
   3.4 Gently agitate the tube to dislodge the cells and examine macroscopically for agglutination.

4 LIS/NIS Indirect Anti human Globulin Test
   After reading the recommended immediate spin direct agglutination test, re-incubate the test for a further 20 minutes at 37°C before completing the indirect anti globulin test by the procedure described below
   OR After reading the recommended LISS tube test complete the indirect antiglobulin test, without further incubation by the procedure described below.
   4.1 Wash the test 4 times with a large excess of PBS pH 7.0 +/- 0.2. Allow adequate spin time to sediment the red cells and ensure residual saline is removed at the end of each wash to leave a “dry” cell button.
   4.2 Add two drops of polyspecific anti human globulin reagent to each tube and mix thoroughly
   4.3 Centrifuge at 1000g for 10 seconds or suitable alternative.
   4.4 Gently agitate the tube to dislodge the cells and examine macroscopically for agglutination.

QUALITY CONTROL
Quality control of reagents is essential and should be performed with each series of groups and with single groups. It is recommended that the following cell samples are used to control the reactions of these reagents. Other cell types may be suitable but should be selected with care.
OR r red cells should be used as positive control.
Orr red cells should be used as negative control.
A reagent control is required for this anti D

TECHNICAL NOTES
Patient Testing
For RhD grouping of patients, the UK Blood transfusion service recommended that two different Anti D reagents should be used. Neither of these should agglutinate DVI red cells by any method recommended for use. In addition “follow on” indirect anti human globulin tests for samples giving negative direct agglutination results are not required.

Donor Testing
For RhD grouping of donors the UK blood transfusion service recommend that the selection of blood grouping reagents should be made to maximise the detection of weak D and partial D red cells. The sensitivity of the grouping procedure should be such to detect all D antigens of clinical significance. When in doubt of a donors RhD status it is safer to consider them RhD positive.

LIMITATIONS
Some examples of Dweak or partial D may not be directly agglutinated when using the slide method. Indeterminate results in the slide method should be confirmed using a tube method.
False positive or false negative results may occur through contamination of test materials or any deviation from the recommended technique.
Certain tests performed on unwashed samples (e.g. cord) or samples stored and tested at 20°C or below may exhibit false positive reactions due to the potentiators used in the formulation of this reagent.

REFERENCES:

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