Anti-Skin Antibody Test System

QUALITY CONTROL
1. Positive and negative serum controls must be included in each day's testing to confirm reproducibility, sensitivity and specificity of the test procedure.
2. The negative serum control should result in little (+) or no fluorescence. If this control shows bright fluorescence, either the control, antigen, conjugate or technique may be at fault.
3. The positive serum controls should result in bright 3+ to 4+ fluorescence. If these controls show little or no fluorescence, either the control, antigen, conjugate or technique may be at fault.
4. In addition to positive and negative serum controls, a PBS control should be run to establish that the conjugate is free from non-specific staining of the antigen substrate. If the antigen shows bright fluorescence in the PBS control repeat using fresh conjugate. If the antigen still fluoresces, either the conjugate or antigen may be at fault.

RESULTS
1. Diffuse staining throughout the tissue is considered non-specific and should be considered a negative result.
2. Staining of the basement membrane (BM) of the epidermis is considered positive and is associated with 70% of bullous pemphigoid cases.
3. Staining of the intercellular substance (ICS) of the prickle cell layer of the epidermis is considered positive 90% of pemphigus cases.
4. The titre is the highest dilution of the patient's serum, showing a weak 1+ fluorescence of the ICS or BM. Titres of 1/20 or greater are clinically relevant for both patterns.

TEST LIMITATIONS
1. No diagnosis should be based on a single serologic test since various host factors must be taken into consideration.
2. Patients with Lydel's toxic neurolysis, extensive burns and myasthenia gravis may demonstrate specific staining of the antigen substrate. If the antigen shows bright fluorescence that can be visually identified.

LITERATURE REFERENCES

INSTRUCTIONS FOR USE
Anti-Skin Antibody (ASA) Test System

INTENDED USE
The Bio-Diagnostics Anti-Skin Antibody Test Kit is an immunofluorescent antibody (IFA) test to detect the presence of antibodies to skin, in human serum.

SUMMARY AND EXPLANATION
The in-vitro detection of skin antibodies by the indirect immunofluorescent technique has been established as an aid in the diagnosis of skin and systemic diseases (4). Monkey oesophagus is the recommended substrate for IFA. Monkey oesophagus is used for the detection of both basement membrane antibodies and intercellular substance antibodies (1,5). The intercellular substance antibody has been associated with the presence of a variety of disorders of the skin (3). The detection of the basement membrane antibody has been associated with the presence of a variety of bullous pemphigoid autoimmune disorders of the skin (3).

PRINCIPLE OF THE TEST
Skin antibodies are not organ or species specific. The primary test reaction involves circulating anti-epidermal antibodies present in the patient's serum, which attach to their homologous epidermal antigens. This occurs during the incubation period whilst the serum covers the antigen surface. A secondary reaction then follows a rinsing period that removes all unbound human antibody. The reagent used in the secondary reaction is a fluorescein labelled anti-human globulin conjugate containing Evans Blue Counterstain. The antigen surface is then thoroughly rinsed free of unbound conjugate and viewed under the appropriate fluorescent microscope for various morphological patterns of epidermal fluorescence that can be visually identified.
**Anti-Skin Antibody Test System**

**WARNINGS AND PRECAUTIONS**

1. All human components have been tested by radioimmunoassay for (HBsAg) and HTLVIII/LAV by an FDA approved method and found to be negative (not repeatedly reactive). However, this does not assure the absence of (HBsAg) or HTLVIII/LAV. All human components should be handled with appropriate care.

2. The controls included in the kit contain 0.1% sodium azide as a preservative. Although this is at a low concentration, these reagents should be considered toxic. They should not be ingested or allowed to come into contact with either the skin or the mucous membranes. Sodium azide may also cause the formation of potentially explosive lead or copper azides in sinks.

3. Do not use components beyond their expiration date.

4. Follow the procedural instructions exactly as they appear in this insert to ensure valid results.

5. Do not use components beyond their expiration date.

6. Handle slides by the edges since direct pressure on the antigen wells may damage the antigen.

7. Once the procedure has started do not allow the antigen in the wells to dry out. This may result in false negative test results, or unnecessary artefacts.

**KIT CONTENTS**

**SLIDE**
- Monkey oesophagus substrate antigen slides (S4205, S8205, S5205 or S0205)

**CONJ**
- FITC Conjugate (for use with Primate substrates) with Evans Blue Counterstain: JS02
- This reagent contains antibodies that will react with the human IgG (H+L) Immunoglobulin class.

**CONTROL**
- ASA basement membrane Positive Control no: C009
- ASA intercellular Positive Control no: C010

**CONTROL**
- Universal Negative Control no: C000

**IFA/DFA PBS**
- Buffer Pack no: R002

**MM**
- Mounting Medium no: R005

**ADDITIONAL MATERIALS REQUIRED BUT NOT PROVIDED**
- Disposable pipettes
- Moisture Chamber
- Distilled Water
- Paper Towels – lint free

**REAGENT PREPARATION**


2. Reconstituted Buffer does not require refrigeration.

**STORAGE AND STABILITY**

The IFA Test System components (except PBS) must be stored at a temperature of +2°C to +8°C. Do not freeze the test kit. The stability of the kit is as indicated by the expiry date on the packaging under the above storage conditions. This applies to unopened and opened reagents. Phosphate Buffered Saline is stable at room temperature storage. The reconstituted Buffer does not contain preservatives and should be stored at 2-8°C. Care should be taken to avoid contamination.

**SPECIMEN COLLECTION**

Serological specimens should be collected under aseptic conditions. Haemolysis is avoided through prompt separation of the serum from the clot. Serum should be stored at 2-8°C if it is to be analysed within a few days. Serum may be held for 3 to 6 months by storage at -20°C or lower. Lipoaemic and strongly haemolytic serum should be avoided. When specimens are shipped at ambient temperatures, addition of a preservative such as 0.01% thiomersal or 0.1% sodium azide is strongly recommended.

**TEST INSTRUCTIONS**

**Screening:** dilute test sera 1/20 (1 part patient sample to 19 parts diluent) in PBS. N.B. Although this dilution factor is suggested, each laboratory should determine their individual screening dilution.

**Titration:** set up doubling dilutions of serum starting at 1/20, (i.e. 1/20, 1/40, 1/80, 1/160, 1/320, etc.).

1. Once slides reach room temperature tear slide envelope at notch. Carefully remove the slide and place on a blotter. Wipe the back of the slide with dry lint free paper towel. Apply sufficient pressure to moisture side faces absorbent side of blotter. Wipe the back of the slide with dry lint free paper towel. Apply sufficient pressure to slide while wiping to absorb buffer. Do not allow tissue to dry.

2. Place a drop of diluted serum (20 to 30µl) and controls over the antigen wells.

3. Place slide with patient’s serum and controls in a moist chamber for 30 minutes at room temperature (approximately 18-24°C).

4. Remove slide from moisture chamber and tap the slide on its side to allow the serum to run off onto a piece of paper towel. Using a wash bottle, gently rinse remaining sera from slide being careful not to aim the rinse stream directly onto the well.

5. Wash in PBS for 5 minutes. Repeat using fresh PBS.

6. Place a blotted on the lab table with absorbent side up. Remove slide from PBS and invert so that tissue side faces absorbent side of blotted. Line up the wells to blotter holes. Place slide on top of the blotter. Wipe the back of the slide with dry lint free paper towel. Apply sufficient pressure to slide while wiping to absorb buffer. Do not allow tissue to dry.

7. Deliver 1 drop (20-30µl) of conjugate per antigen well. Repeat steps 3-6.

8. Place 4-5 drops of mounting medium on slide.

9. Apply a 22 x 70 mm coverslip. Examine the slide under a fluorescent microscope. Note: To maintain fluorescence, store mounted slide in a moisture chamber placed in a dark refrigerator.