For detection of antibodies to HIV-1 in serum or plasma samples.

NAME

The GENELABS DIAGNOSTICS HIV-1 BLOT 1.3 is a qualitative enzyme immunoassay for antibodies to HIV-1 in human serum or plasma.

This kit is supplied for research purposes only. It is not intended for use in the diagnosis or prognosis of disease. In particular, this test cannot be used to evaluate blood specimens for the purpose of donor screening, or as a confirmatory diagnostic.

INTRODUCTION

The Genelabs Diagnostics HIV-1 BLOT 1.3 Western Blot is an informational research test on serum or plasma specimens. The separated specific viral antigens incorporated onto the strips via electrophoretic and electrotransblot procedures, will also allow for further delineation of the antibody responses to specific viral proteins. Each strip also includes an internal sample addition control to minimize the risk of false negatives due to operational errors and to ensure the addition of samples.
The nitrocellulose strips are incorporated with separated, bound antigenic proteins from partially purified inactivated HIV-1 using electrophoretic blotting. Individual nitrocellulose strips are incubated with diluted serum or plasma and controls. Specific antibodies to HIV-1 if present in the specimens, will bind to the HIV-1 proteins on the strips. The strips are washed to remove unbound materials. Antibodies that bind specifically to HIV-1 proteins can be visualized using a series of reactions with goat anti-Human IgG conjugated with alkaline phosphatase and the substrate, BCIP/NBT.

### KIT COMPONENTS

1. **NITROCELLULOSE STRIPS**
   - Incorporated with HIV-1 viral lysate.
   - Keep dry and away from light.
   - Available in 18 or 36 strips

2. **NON-REACTIVE CONTROL**
   - Inactivated normal human serum non-reactive for Hepatitis B surface antigen (HBsAg), antibodies to HIV-1 and HCV. Contains sodium azide and thimerosal as preservatives.
   - 1 vial (80 ul)

3. **STRONG REACTIVE CONTROL**
   - Inactivated human serum with high titered antibodies to HIV-1 and non-reactive for HBsAg and anti-HCV. Contains sodium azide and thimerosal as preservatives.
   - 1 vial (80 ul)

4. **WEAK REACTIVE CONTROL**
   - Inactivated human serum with low titered antibodies to HIV-1 and non-reactive for HBsAg & HCV. Contains sodium azide and thimerosal as preservatives.
   - 1 vial (80 ul)

5. **STOCK BUFFER CONCENTRATE (10X)**
   - Tris buffer with heat inactivated normal goat serum. Contains thimerosal as preservative.
   - 1 bottle (20 ml)

6. **WASH BUFFER CONCENTRATE (20X)**
   - Tris with Tween-20. Contains thimerosal as preservative.
   - 1 bottle (70 ml)

7. **CONJUGATE**
   - Goat anti-human IgG conjugated with alkaline phosphatase.
   - 1 vial (120 ul)

8. **SUBSTRATE**
   - Solution of 5-bromo-4-chloro-3-indolyl-phosphate (BCIP) and nitroblue tetrazolium (NBT).
   - 1 bottle (100 ml)
9. BLOTTING POWDER
   Non-fat dry milk (1g each)

10. Incubation Tray, 9 wells each.

11. Instruction Manual

12. Forceps

Volume of reagents provided are sufficient for 4 runs.

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**PRECAUTIONS TO USERS**

**Caution:** Handle all assay specimens, positive and negative controls as potentially infectious agents.

1. Substituting reagents even between lots, may affect results.
2. **FOR RESEARCH USE ONLY, NOT FOR USE IN DIAGNOSTIC PROCEDURES.**
3. Do not use kit components beyond the expiry date.
4. Avoid microbial contamination of reagents when opening and removing aliquots from the original vials or bottles.
5. Gloves and lab coats must be worn.
6. Do not pipette by mouth.
7. Wipe spills quickly and thoroughly with sodium hypochlorite solution.
8. Autoclave all used and contaminated materials at 121°C at 15 p.s.i. for 30 minutes before disposal.
9. It is highly recommended that this assay be performed in a biohazard cabinet.
10. Decontaminate all used chemicals and reagents in sodium hypochlorite solution.
11. We do not recommend re-use of incubation trays.

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**STORAGE INSTRUCTIONS**

**A. Antigen strips**
   Avoid unnecessary exposure of antigen strips to light.

**B. Reagents**
   Store all reagents at 2-8°C.
   For best results, dispense all reagents while cold and return to 2-8°C storage as soon as possible.

**CAUTION:** Avoid unnecessary exposure of substrate to light.
MATERIALS REQUIRED BUT NOT PROVIDED

- Rocker platform *
- Pipettors and tips
- Aspirator with sodium hypochlorite trap *
- 56°C Water bath (optional)

* Not required if using Autoblot System 36.

SPECIMEN HANDLING AND STORAGE (OPTIONAL)

Sera can be inactivated but this is not a requirement for optimal test performance.

Inactivate as follows:
1. Loosen caps of serum containers.
2. Heat serum to 56°C for 30 mins in a water bath.
3. Allow serum to cool before retightening caps.
4. Serum can be stored frozen until analysis.

We recommend that the sera should not undergo repeated freeze-thaw cycles prior to testing.

PREPARATION OF REAGENTS

1. DILUTED WASH BUFFER
   (a) Dilute 1 volume of WASH BUFFER CONCENTRATE (20X) with 19 volumes of reagent grade water. Mix well.

2. BLOTTING BUFFER
   (a) BLOTTING BUFFER should be prepared fresh prior to use.
   (b) Dilute 1 volume of STOCK BUFFER CONCENTRATE (10X) with 9 volumes of reagent grade water. Mix well.
   (c) Add 1 g of BLOTTING POWDER to every 20ml of the DILUTED STOCK BUFFER prepared in step 2(b) above. Mix well.

3. WORKING CONJUGATE SOLUTION
(a) Prepare WORKING CONJUGATE SOLUTION by diluting CONJUGATE 1:1000 into BLOTTING BUFFER, for example, 5ul CONJUGATE to 5ml BLOTTING BUFFER.
(b) WORKING CONJUGATE SOLUTION should be prepared fresh prior to use.

4. SUBSTRATE SOLUTION (ready to use)
   (a) Dispense directly the required volume from the bottle. Use a clean pipette. Cap tightly after use.

**RECOMMENDED ASSAY PROCEDURE - RAPID ASSAY**

*Note:* Aspirate all used chemicals and reagents into trap containing sodium hypochlorite.

1. Using forceps, carefully remove required number of STRIPS from the tube and place numbered side up into each well. Include strips for Strong Reactive, Weak Reactive and Non-Reactive controls.

2. Add 2ml of DILUTED WASH BUFFER to each well.

3. Incubate the strips for at least 5 minutes at room temperature (25 ± 3°C) on a rocking platform. Remove buffer by aspiration.

4. Add 2ml of BLOTTING BUFFER to each well followed by 20ul each of sera or controls to appropriate wells.

5. Cover the tray with the cover provided and incubate for 1 hour at room temperature (25 ± 3°C) on the rocking platform.

6. Carefully uncover the tray to avoid splashing or mixing of samples. Aspirate the mixture from the wells. Change aspirator tips between samples to avoid cross-contamination.

7. Wash each strip 3 times with 2ml of DILUTED WASH BUFFER allowing 5 minutes soak on the rocking platform between each wash.

8. Add 2 ml of WORKING CONJUGATE SOLUTION to each well. Cover tray and incubate for 1 hour at room temperature (25 ± 3°C) on the rocking platform.


10. Add 2 ml of SUBSTRATE SOLUTION to each well. Cover tray and incubate for 15 minutes on the rocking platform.
11. Aspirate the SUBSTRATE and rinse the strips several times with reagent grade water to stop the reaction.


13. Mount strips on worksheet (non-absorbent white paper). Do not apply adhesive tape over the developed bands. Observe the bands and grade the results. For storage, keep the strips in the dark.

**ALTERNATIVE RECOMMENDED PROCEDURE - OVERNIGHT ASSAY**

**Note:** Aspirate all used chemicals and reagents into trap containing sodium hypochlorite.

1. Using forceps, carefully remove required number of STRIPS from the tube and place numbered side up into each well. Include strips for Strong Reactive, Weak Reactive and Non- Reactive controls.

2. Add 2ml of DILUTED WASH BUFFER to each well.

3. Incubate the strips for at least 5 minutes at room temperature (25 ± 3°C) on a rocking platform. Remove buffer by aspiration.

4. Add 2ml of BLOTTING BUFFER to each well followed by 20ul of each of sera or controls to appropriate wells.

5. Cover the tray with the cover provided and incubate overnight (16-20 hours) at room temperature (25 ± 3°C) on the rocking platform.

6. Carefully uncover the tray to avoid splashing or mixing of samples. Aspirate the mixture from the wells. Change aspirator tips between samples to avoid cross-contamination.

7. Wash each strip 3 times with 2ml of DILUTED WASH BUFFER allowing 5 minutes soak on the rocking platform between each wash.

8. Add 2 ml of WORKING CONJUGATE SOLUTION to each well. Cover tray and incubate for 30 minutes at room temperature (25 ± 3°C) on the rocking platform.


10. Add 2 ml of SUBSTRATE SOLUTION to each well. Cover tray and incubate for 15 minutes on the rocking platform.
11. Aspirate the substrate and rinse the strips several times with reagent grade water to stop the reaction.


13. Mount strips on worksheet (non-absorbent white paper). Do not apply adhesive tape over the developed bands. Observe the bands and grade the results. For storage, keep the strips in the dark.

### SUMMARY RECOMMENDED ASSAY PROTOCOLS

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Qty</th>
<th>Rm Temp Rapid Assay</th>
<th>Rm Temp Overnight Assay</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrocellulose strip</td>
<td>1</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Wash Buffer</td>
<td>2ml</td>
<td>5 mins</td>
<td>5 mins</td>
</tr>
<tr>
<td>Blotting Buffer</td>
<td>2ml</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Specimen</td>
<td>20ul</td>
<td>60 mins</td>
<td>Overnight (16 - 20 hours)</td>
</tr>
<tr>
<td>Wash Buffer</td>
<td>3 x 2ml</td>
<td>3 x 5 mins</td>
<td>3 x 5 mins</td>
</tr>
<tr>
<td>Conjugate</td>
<td>2ml</td>
<td>60 mins</td>
<td>30 mins</td>
</tr>
<tr>
<td>Wash Buffer</td>
<td>3 x 2ml</td>
<td>3 x 5 mins</td>
<td>3 x 5 mins</td>
</tr>
<tr>
<td>Substrate (Ready to use)</td>
<td>2ml</td>
<td>15 mins</td>
<td>15 mins</td>
</tr>
<tr>
<td>Distilled Water</td>
<td>2ml</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

**Note:** All incubations are to be carried out on a rocking platform. Alternatively GENELABS offers an Autoblot System 36 which is designed to perform all Genelabs Diagnostics Western Blot assays automatically. Please contact your nearest distributor for more information.

### AMOUNT OF REAGENTS REQUIRED FOR VARIOUS NUMBER OF STRIPS

<table>
<thead>
<tr>
<th>Reagents</th>
<th>NUMBER OF STRIPS TO BE USED</th>
</tr>
</thead>
<tbody>
<tr>
<td>1X Wash Buffer (ml)</td>
<td>3  6  9  15  20  27  36</td>
</tr>
<tr>
<td>1X Blotting Buffer (ml)</td>
<td>60 100 140 240 300 400 520</td>
</tr>
<tr>
<td>Conjugate</td>
<td>20 40 60 80 100 120 160</td>
</tr>
<tr>
<td>Substrate (ml)</td>
<td>11 17 23 35 45 59 77</td>
</tr>
<tr>
<td>Blotting Powder (g)</td>
<td>11 17 23 35 45 59 77</td>
</tr>
</tbody>
</table>

### REFERENCE STANDARDS

We recommend that the Non-Reactive, Strong Reactive and Weak Reactive controls be run with every assay regardless of the number of samples tested.
1. **NON-REACTIVE CONTROL**
No HIV-1 specific bands should be observed on the Non-Reactive control strips. The band for the serum control should be visible (Fig 1c).

2. **STRONG REACTIVE CONTROL**
All relevant molecular weight bands must be evident. Figure 1a provides a guide to the relative positioning of bands visualised with the Genelabs Diagnostics HIV-1 Blot 1.3 and permits identification of bands observed for the STRONG REACTIVE CONTROL. The bands are p17, p24, p31, gp41, p51, p55, p66 and gp120/gp160. Other bands associated with core antigens (ie. p39, p42) may also be visible. Be careful not to misinterpret these as gp41. The envelope antigens, gp41, gp120/gp160 appear as diffuse bands as they are typical of glycoproteins. The serum control band will be visible.

3. **WEAK REACTIVE CONTROL**
Weak bands at p24 and gp120/160 should appear. Some additional weak bands may or may not be present. The serum control band will be visible (Fig 1b).

### INTERPRETATION OF BANDS

The presence or absence of antibodies to HIV-1 in a sample is determined by comparing each nitrocellulose strip to the assay control strips tested with the NON-REACTIVE, STRONG REACTIVE and WEAK REACTIVE controls.

The Figure 1a on Page 11 is suggested as an aid to identify the various bands which develop on the strip reacted with the STRONG REACTIVE Control.

<table>
<thead>
<tr>
<th>Molecular Weight</th>
<th>Gene</th>
<th>Antigen</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>gp 160</td>
<td>ENV</td>
<td>Polymeric form of gp41</td>
<td>Broad diffuse glycoprotein</td>
</tr>
<tr>
<td>gp 120</td>
<td>ENV</td>
<td>Outermembrane</td>
<td>Diffuse glycoprotein</td>
</tr>
<tr>
<td>p66</td>
<td>POL</td>
<td>Reverse Transcriptase</td>
<td>Discreet band</td>
</tr>
<tr>
<td>p55</td>
<td>GAG</td>
<td>Precursor protein</td>
<td>Discreet band</td>
</tr>
<tr>
<td>p51</td>
<td>POL</td>
<td>Reverse Transcriptase</td>
<td>Discreet band just below p55</td>
</tr>
<tr>
<td>gp41</td>
<td>ENV</td>
<td>Transmembrane</td>
<td>Diffuse glycoprotein</td>
</tr>
<tr>
<td>p39</td>
<td>GAG</td>
<td>Fragment of p55</td>
<td>Discreet band</td>
</tr>
<tr>
<td>p31</td>
<td>POL</td>
<td>Endonuclease</td>
<td>Doublet</td>
</tr>
<tr>
<td>p24</td>
<td>GAG</td>
<td>Core protein</td>
<td>Broad band</td>
</tr>
<tr>
<td>p17</td>
<td>GAG</td>
<td>Core protein</td>
<td>Broad band</td>
</tr>
</tbody>
</table>

Some of the different antigens mentioned in the Table above are derived from the same precursor protein and may have overlapping epitopes. This should be considered when interpreting the pattern, for example:-
1. It is unlikely to detect gp41 in the absence of gp160 because the gp160 is the polymeric form of gp41 and the concentration of gp160 is higher than gp41 on the HIV-1 Blot 1.3.

2. The p55 band is generally detected when there is strong reactivity to p24 and/or p17. The bands seen as p42 and p39 are both GAG fragments and should not be interpreted as gp41 (ENV).

3. The POL bands p66, p51 and p31 are generally detected simultaneously. However the sensitivity of p66 and p31 are greater than p51.

4. HIV-2 cross reactivity is variable but typically shows reactivity with GAG and/or POL antigens. However, there can be cross reactivity with the gp160 band in some cases, but rarely with gp41.

5. There is also a high molecular weight band around 160KD that is presumed to be a GAG-POL precursor protein. This is seen with some high titered HIV-2 or Indeterminate (GAG Reactive Only) sera but the band pattern is a sharp discreet band which is different from the diffuse band of ENV gp160.

**LIMITATIONS OF THE PROCEDURE**

Deviation from the recommended procedure may lead to aberrant results.

**LIMITED EXPRESSED WARRANTY DISCLAIMER**

The manufacturer makes no warranty other than that the test kit will function as a Research Use Only assay within the specifications and limitations described in the Product Instruction Manual when used in accordance with the instructions contained therein. The manufacturer disclaims any warranty, express or implied, including such express or implied warranty with respect to merchantability, fitness for use or implied utility for any purpose. The manufacturer is limited to either replacement of the product or refund of the purchase price of the product. The manufacturer shall not be liable to the purchaser or third parties for any damage, injury or economic loss howsoever caused by the product in the use or in the application thereof.

**REFERENCE**


FIGURE 1

Viral specific bands as visualized with:

a) Strong Reactive Control
b) Weak Reactive Control
c) Non-Reactive Control