



## HTLV BLOT 2.4 WESTERN BLOT ASSAY INSTRUCTIONS FOR USE

### IVD

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### REF

(18 tests kit): 0711088018  
 (36 tests kit): 0711088036

### NAME AND INTENDED USE

The **MP Diagnostics HTLV Blot 2.4** is a qualitative enzyme immunoassay intended for confirming the presence of and differentiating antibodies to HTLV-I and HTLV-II in human serum and plasma. It is intended for use as a supplemental (additional, more specific) test for human serum and plasma samples with repeatedly reactive results by an FDA licensed HTLV-I/II donor screening test. The MP Diagnostics HTLV Blot 2.4 is intended for use in a manual mode or a semi-automated mode using the MP Diagnostics AutoBlot System 20. This test is not intended for use in medical diagnosis.

### INTRODUCTION AND EXPLANATION OF THE TEST

#### Background

Human T-cell Lymphotropic Viruses (HTLVs) are pathogenic retroviruses that may cause severe hematological and neurological diseases in infected individuals. The HTLV family has two well-studied members: HTLV-I and HTLV-II. HTLV-I is known as the etiological agent of adult T-cell leukemia/lymphoma (ATL), HTLV-associated myelopathy/tropical spastic paraparesis (HAM/TSP), and HTLV-associated uveitis. Although less pathogenic than HTLV-I, HTLV-II infection has been associated with leukemia and neurological disease but the causal relationship remains uncertain.

Studies of the geographic distribution of HTLV-I infection reveal that the HTLV-I virus is highly prevalent in Japan, Africa, the Caribbean Islands, and South America. Recent epidemiological studies in the United States and Europe confirm the presence of a mixed prevalence of both HTLV-I and HTLV-II among different high-risk populations, such as intravenous drug users and transfusion recipients. The viruses can be transmitted through sexual contact, through contaminated blood products, and from mother to child via breastfeeding.

Screening tests for HTLV-I/II are available although limited. Repeatedly reactive specimens from screening tests require additional more specific tests to confirm HTLV seropositivity including discrimination of HTLV-I and HTLV-II seropositives. These supplemental assays (i.e. type-specific peptide EIAs, ELISAs, or Western blots) must be capable of identifying antibodies to core (gag) and envelope (env) proteins of HTLV-I and HTLV-II. Western blot strips incorporating HTLV-I native viral antigens are one such commonly used supplemental test.

Screening of whole blood donations for the presence of antibodies to HTLV-I/II has been required in the United States since 1988. Simple yet specific and sensitive supplemental serological tests are therefore needed to enable rapid confirmation and differentiation of HTLV-I and HTLV-II seropositive samples. A supplemental test is essential to provide additional key information necessary for donor counseling, follow-up testing, and/or treatment.

#### Virology

HTLV-I and HTLV-II are type C human oncoviruses with single-stranded RNA genomes that are approximately 8,900 base pairs in length. The HTLV-I/II genomes include gag and env genes which encode structural core proteins p19 and p24, as well as envelope proteins gp46 and p24<sup>ie</sup>, respectively. Like other human retroviruses, the HTLV-I/II *pol*

genes encode a reverse transcriptase to allow transcription of the RNA genome into a complementary DNA strand, which is then integrated into the host genome by a *pol* encoded integrase.

#### Diagnosis

HTLV-I/II infections are generally diagnosed by antibody tests (e.g., Enzyme-linked immunosorbent assay (ELISA), Chemiluminescence assay (CLIA), Western blot, Immunofluorescence assay (IFA)). Due to the inclusion of cross-reactive antigens, most assays detect both HTLV-I and HTLV-II antibodies, although sensitivity for HTLV-II may be lower<sup>2</sup>. Natural or recombinant, type-specific, envelope proteins, in IFA or Western blot format permit the differentiation of HTLV-I from HTLV-II antibodies<sup>5</sup>. The two virus types may also be distinguished by polymerase chain reaction (PCR) or *in-situ* hybridization directed against HTLV-I proviral DNA or RNA sequences.<sup>6</sup> Quantitative PCR studies have also determined that the proviral DNA load in both HTLV-I and HTLV-II ranges from approximately 10<sup>4</sup> to 10<sup>7</sup> per peripheral blood mononuclear cell.<sup>5,6</sup>

#### Epidemiology

HTLV-II is endemic at levels up to five percent (5%) of the general population in central Africa, several Caribbean basin and South American countries, and in southern Japan.<sup>7</sup> Transmission is from mother to child, predominantly by breastfeeding; through sexual intercourse, predominantly in the male-to-female direction; and via parenteral exposure by blood transfusion or needle sharing. In the United States, first time blood donor HTLV-II seroprevalence is about one per ten thousand, and risk factors include maternal or sexual links to HTLV-II endemic areas. In contrast, HTLV-II seroprevalence is about two per ten thousand, and predominant risk factors are injection drug use and sexual contact with an injection drug user (IDU)<sup>8,9</sup>. In a 2012 study, the rate of overall HTLV infection (undifferentiated) in all US donors was determined as 1:35,313<sup>10</sup> Researchers have estimated that there may be as many as ten to twenty million persons with HTLV-I infection in the world; a more conservative estimate might be between one to five million<sup>11</sup>.

HTLV-II is endemic in certain North,<sup>11</sup> Central<sup>12</sup> and South<sup>13,14</sup> American Indian tribes, with some of the highest seroprevalence values (up to fifty percent) documented in tribes with the least contact with contemporary civilization, such as the Brazilian Kayapo. This led to the hypothesis that HTLV-II was already endemic in these tribes before they migrated across the Bering Land Bridge over ten thousand years ago. A single report of HTLV-II among Mongolians has not been supported by other studies of the same population<sup>15</sup>. However, clusters of HTLV-II infection have been conclusively demonstrated among isolated Pygmy tribes in central Africa<sup>16,17</sup>. Genetic similarities between Pygmy and Native American HTLV-I isolates have not been explained<sup>18,20</sup>.

An early study that differentiated HTLV-I from HTLV-II using a competitive HTLV-I/II ELISA technique reported a high seroprevalence of both HTLV-I and HTLV-II among IDU in the New Jersey area<sup>21</sup>. In New Orleans, approximately twenty-five percent (25%) of IDU tested were HTLV-II positive by PCR and another two percent (2%) were infected with HTLV-I<sup>22</sup>. Sixteen percent (16%) of San Francisco IDU are HTLV seropositive, and most of these appear to be infected with HTLV-II<sup>23</sup>. A study of primarily white IDU from the Staten Island, New York area, reported a seroprevalence of eleven percent (11%) for HTLV-I and an additional nine percent (9%) for HTLV-I<sup>24</sup>. Finally, measurement of HTLV-I/II antibodies in sera from the CDC-sponsored HIV Sentinel Counties Survey yielded undifferentiated HTLV-I/II prevalence among IDU in methadone treatment centers ranging from 0.4% (Atlanta) to 17.6% (Los Angeles)<sup>25</sup>. Interestingly, there was little concordance in the ranking of cities by HIV prevalence versus HTLV-I/II prevalence.

Based upon the 2000 U.S. Census data, it is estimated that the total number of HTLV-II infected persons in the United States is approximately 197,000. This includes 56,000 in the general population (U.S. population 281,422,000 X 0.02% blood donor prevalence<sup>9</sup>), 100,000 among IDU (1 million IDU X 10% prevalence<sup>26</sup>) and 41,000 among American Indians (4,119,000 Native American/Alaska natives X 1% prevalence<sup>27</sup>).

#### Disease Associations

HTLV-I causes ATL, a malignancy of mature CD4<sup>+</sup> T-lymphocytes that presents most commonly as lymphoma with skin involvement and hypercalcemia.<sup>28</sup> HTLV-I is the causative agent of HAM, a slowly progressive spastic paraparesis that is characterized by weakness in the legs, diffuse hyperreflexia, clonus, loss of vibration sense, and detrusor insufficiency leading to bladder dysfunction. HTLV-I may also be associated with a wider spectrum of neurological manifestations that do not meet diagnostic criteria for HAM. Sensory neuropathy,<sup>27,29</sup> gait abnormalities,<sup>30,31</sup> bladder dysfunction,<sup>27,30,31</sup> erectile dysfunction,<sup>32,33</sup> amyotrophic lateral sclerosis (ALS),<sup>34</sup> mild cognitive deficits<sup>35</sup>, and rarely, motor neuronopathy<sup>36,37,38</sup> have all been reported among HTLV-I-infected individuals without HAM. HTLV-I infection has also been implicated in a spectrum of autoimmune conditions such as uveitis, arthritis, and pneumonitis, although there is good epidemiologic evidence of association only with uveitis and arthritis<sup>37,41,42</sup>.

#### Manual Method – Equipment

- Titration pump/aspirator with sodium hypochlorite waste trap
- Belco Rocker or equivalent rocking platform (capable of a speed of 12 to 16 oscillations per minute and a tilt angle of 5 to 10 degrees)

#### Semi-Automated Method – Equipment

- MP Diagnostics AutoBlot System 20.

The integrated protocol to run the MP Diagnostics HTLV Blot 2.4 assay using the MP Diagnostics AutoBlot System 20 is available directly from MP Biomedicals, LLC. Please contact MP Biomedicals' Customer Service.

#### PREPARATION OF REAGENTS – Manual Use Only

##### 1. DILUTED WASH BUFFER

- (a) DILUTED WASH BUFFER should be prepared fresh prior to use.
- (b) Dilute 1 volume of WASH BUFFER CONCENTRATE (20x) with 19 volumes of reagent water. Mix well by inverting or stirring.

Wash Buffer Preparation Chart		
Number of Strips	Volume of Wash Buffer Concentrate	Volume of Reagent Water
9 – 12	10 mL	190 mL
13 – 18	15 mL	285 mL
19 – 24	18 mL	342 mL
25 – 30	23 mL	437 mL
31 – 36	27 mL	513 mL

##### 2. BLOTTING BUFFER

- (a) Reconstitute each bottle of LYOPHILIZED STOCK BUFFER with 100 mL reagent or deionized water. Mix well to dissolve. This RECONSTITUTED STOCK BUFFER is stable for 6 weeks if stored at 2°C to 8°C.
- (b) BLOTTING BUFFER should be prepared fresh prior to use. Add 1 mL (one packet) of BLOTTING POWDER to every 20 mL of the RECONSTITUTED STOCK BUFFER prepared in step 2(a) above. Mix well by inversion or stirring to ensure powder dissolves completely.
- (c) Stir again before dispensing.

Blotting Buffer Preparation Chart		
Number of Strips	Packets of Blotting Powder	Volume of Reconstituted Stock Buffer
9	2	40 mL
10 – 12	3	60 mL
13 – 18	4	80 mL
19 – 24	5	100 mL
25 – 30	7	140 mL
31 – 36	8	160 mL

##### 3. WORKING CONJUGATE SOLUTION

- Note: Prepare solution in polypropylene container / beaker.
- (a) WORKING CONJUGATE SOLUTION should be prepared fresh prior to use.
  - (b) Prepare WORKING CONJUGATE SOLUTION by diluting CONJUGATE with BLOTTING BUFFER in the ratio of 1:1000 (e.g., 10 µl CONJUGATE to 10 mL BLOTTING BUFFER, etc.).

Working Conjugate Preparation Chart		
Number of Strips	Volume of Conjugate	Volume of Blotting Buffer
9	20 µL	20 mL
10 – 12	26 µL	26 mL
13 – 18	38 µL	38 mL
19 – 24	50 µL	50 mL

HTLV-II was initially isolated from two patients with unusual hairy T-cell leukemia. It was subsequently determined that at least one of these patients had a dual disorder: HTLV-II negative T-cell hairy cell leukemia and HTLV-II positive CD8<sup>+</sup> lymphoproliferative syndrome<sup>43</sup>. Although the old literature reports suggests HTLV-II is associated with a myelopathic syndrome similar to HTLV-I related HAM is derived from four cases from the HTLV Outcomes Study (HOST) cohort and about a dozen cases of HTLV-II classical HAM, some with virologic evidence of HTLV-II in cerebrospinal fluid<sup>44-46</sup>. The role of HTLV-II in neurological diseases is less clear<sup>47</sup>.

#### Explanation of the Test

The **MP Diagnostics HTLV Blot 2.4** is intended as a supplemental (additional more specific), test to confirm the presence of anti-HTLV-I/II antibodies in blood donor specimens repeatedly reactive on an FDA licensed screening test and to differentiate between HTLV type-I and HTLV type-II infections for donor notification and counseling. The possible serological profiles defined by the HTLV Blot 2.4 include the following: HTLV-I Seropositive, HTLV-II Seropositive, HTLV-I/II Seropositive, Seronegative and Indeterminate.

The **MP Diagnostics HTLV Blot 2.4** uses a combination of HTLV-I/II genetically engineered proteins (i.e., recombinant antigens) and HTLV-I viral proteins derived from native, inactivated viral particles (i.e., viral lysate). The differentiation between HTLV-I and HTLV-II is accomplished through the use of rgp46-I, a unique HTLV-I envelope recombinant protein, and rgp46-II, a unique HTLV-II envelope recombinant protein. Both proteins are derived from the central region of the external glycoprotein, gp46, of HTLV-I and HTLV-II respectively. GD21, a common yet specific HTLV-I and HTLV-II epitope envelope recombinant protein derived from a truncated region of p21e (rgp21), is also used to enhance the specificity of envelope antibody detection: GD21 has demonstrated better specificity over p21e<sup>48</sup>, an earlier version of the recombinant antigen. The antigenicity exhibited by these recombinant proteins is either common to HTLV-I and HTLV-II antibodies or type specific to one of the two viral types to allow confirmation and differentiation in a single assay. Additional differentiation between HTLV viral types is effected using gag proteins p19 and p24. If p19 is greater than or equal to p24, HTLV-I infection is suggested, and if p24 is greater than p19, HTLV-II infection is suggested<sup>49-71</sup>.

#### DESCRIPTION OF SYMBOLS USED

The following are graphical symbols used in, or found on, **MP Diagnostics** products and packaging.

	Catalogue Number <i>Synonym for this:</i> Reference Number Re-order Number
	Do not reuse
	Caution
	Consult instructions for use

#### CHEMICAL & BIOLOGICAL ASSAY PRINCIPLES

HTLV-I viral proteins, derived from native, inactivated viral particles (viral lysate) and HTLV-II genetically engineered proteins, are incorporated into the nitrocellulose strips.

Individual nitrocellulose strips are incubated with diluted serum or plasma specimens; specific antibodies to HTLV-I/II, if present in the specimen, will bind to the HTLV-I/II proteins on the strip.

The strips are washed to remove unbound materials, and the remaining antibodies, bound to the HTLV proteins on the strips, are visualized using a series of reactions with goat anti-human IgG conjugated with alkaline phosphatase and the substrate, BCIP/NBT.

25 – 30	62 µL	62 mL
31 – 36	74 µL	74 mL

#### 4. SUBSTRATE SOLUTION (ready-to-use)

- (a) Transfer the required volume from the bottle using a clean pipette. Cap tightly after use.

#### ASSAY PROCEDURE

**NOTE:** This section describes the **Manual Assay Procedure**. The automated assay procedure using the MP Diagnostics AutoBlot System 20 is provided as an addendum. To receive a copy, please contact MP Biomedicals, LLC.

- Note:** a) Aspirate all used chemicals and reagents into waste trap containing sodium hypochlorite.
- b) Add samples and controls carefully to avoid mixing up the order of the addition of samples and controls.**
- c) All incubation steps are to be carried out on a rocking platform.
- d) New disposable trays should be used for each assay; do not reuse trays.

**Caution: Adding sample or control directly to the strip may result in the formation of dark patches on the strip in the location where the sample/control was added. To ensure the proper addition of sample or control:**

- i. Nitrocellulose strips should be added with the numbered end of the strip facing up and located at the top of the tray well (the side furthest away from the operator). It is strongly recommended that the strip numbers be placed in ascending order and the tray wells be numbered.
- ii. Sample should be added only after BLOTTING BUFFER is added.
- iii. Tilt the tray slightly by elevating the top of the tray. Add the sample at the bottom of the tray well where the Blotting Buffer has collected. When all the samples have been added, return the tray back to its original flat position. Always ensure that the strips are kept wet during the process.

#### Procedure:

1. Add 2 mL of DILUTED WASH BUFFER to each well. **2 mL**
2. Using forceps, carefully remove a nitrocellulose strip from the tube and place numbered side up into the first well in the tray. The number should be placed at the top of the tray well. Repeat this process until the correct number of strips has been added to the tray. Include strips for Strong Reactive Control I, Strong Reactive Control II and Non-Reactive Control. **20 µL**
3. Incubate the strips for 5 minutes at room temperature (25°C ± 3°C) on a rocking platform with a speed of 12 to 16 oscillations per minute and a 5° - 10° tilt. Remove buffer using a vacuum pump aspirator with sodium hypochlorite waste trap. **5 minutes 25°C ± 3°C**
4. Add 2 mL of BLOTTING BUFFER to each well. **2 mL**
5. Add 20 µL each of test sample or control to appropriate wells. **20 µL**
6. Cover the tray with the cover provided and incubate for 1 hour at room temperature (25°C ± 3°C) on a rocking platform with a speed of 12 to 16 oscillations per minute and a 5° - 10° tilt. **60 minutes 25°C ± 3°C**
7. Carefully uncover the tray to avoid splashing or mixing of samples. Tilt the tray to aspirate the mixture from the wells. Change the manual aspirator tips (if possible) between samples to avoid cross-contamination. **15 minutes 25°C ± 3°C**
8. Wash each strip 3 times with 2mL of DILUTED WASH BUFFER allowing 5 minutes to soak on the rocking platform between each wash. **3 x 2 mL**
9. Add 2 mL of WORKING CONJUGATE SOLUTION to each well. **2 mL**
10. Cover tray and incubate for 1 hour at room temperature (25°C ± 3°C) on the rocking platform. **60 minutes 25°C ± 3°C**
11. Aspirate WORKING CONJUGATE SOLUTION from the wells. Wash. (See Step 8). **3 x 2 mL**
12. Add 2 mL of SUBSTRATE SOLUTION to each well. **2 mL**
13. Cover tray and incubate for 15 minutes on the rocking platform. **15 minutes 25°C ± 3°C**
14. Aspirate the SUBSTRATE and rinse strips 3 times with reagent or deionized water to stop the reaction. Rinse the strips by adding 2 mL of reagent water, manually rocking the tray gently to ensure that the strips are fully covered with water, and immediately aspirating. Repeat this procedure an additional 2 times. **3 x 2 mL**

Of the proteins applied to the nitrocellulose strips, five are used to confirm the presence of antibodies against HTLV-II. These are the following: rgp46-I, rgp46-II, GD21, p19 and p24.

Type-specific recombinant envelope protein rgp46-I is specific for HTLV-I, while rgp46-II is specific for HTLV-II; these antigens are used to differentiate between HTLV-I and HTLV-II infections<sup>72,73</sup>.

GD21, a third recombinant envelope protein, is broadly immunoreactive with sera or plasma from HTLV-I and HTLV-II infected individuals<sup>68-72, 74</sup>.

Two gag proteins, p19 and p24, which are reactive to HTLV-I and cross-reactive to HTLV-II, are used to confirm the presence of antibodies. It has been found that reactivity against p19 is greater than, or equal to, reactivity against p24 in subjects who had HTLV-I infection confirmed by PCR. Conversely, p24 bands were stronger than p19 bands in persons who had PCR confirmed HTLV-II infection<sup>67-71</sup>.

KIT COMPONENTS		
Component	Description	Quantity Provided
	<b>NITROCELLULOSE STRIPS</b> Incorporated with HTLV-I viral lysate, HTLV-I and II recombinant envelope antigens, and a sample addition control (anti-human IgG) band. Keep dry and away from light.	Available in 18 or 36 strips
	<b>NON-REACTIVE CONTROL</b> Inactivated normal human serum, non-reactive for anti-HCV, anti-HIV-1/2, anti-HTLV-I/II and HBSAg. Contains sodium azide and thimerosal as preservatives.	1 vial (80 µL)
	<b>STRONG REACTIVE CONTROL I</b> Inactivated human serum with high titer antibodies to HTLV-I and non-reactive for anti-HCV, anti-HIV-1/2 and HBSAg. Contains sodium azide and thimerosal as preservatives.	1 vial (80 µL)
	<b>STRONG REACTIVE CONTROL II</b> Inactivated human serum with high titer antibodies to HTLV-I and non-reactive for anti-HCV, anti-HIV-1/2 and HBSAg. Contains sodium azide and thimerosal as preservatives.	1 vial (80 µL)
	<b>LYOPHILIZED STOCK BUFFER</b> To be reconstituted in reagent grade water. Tris buffer with heat inactivated animal and non-animal proteins. Contains thimerosal as preservative.	1 or 2 bottles (each to be reconstituted to 100 mL)
	<b>WASH BUFFER CONCENTRATE (20x)</b> Tris with Tween-20. Contains thimerosal as preservative.	1 bottle (70 mL)
	<b>CONJUGATE</b> Goat anti-human IgG conjugated with alkaline phosphatase. Contains sodium azide as preservative.	1 vial (120 µL)
	<b>SUBSTRATE</b> Solution of 5-bromo-4-chloro-3-indolylphosphate (BCIP) and nitroblue tetrazolium (NBT).	1 bottle (100 mL)
	<b>BLOTTING POWDER</b> Non-fat dry milk.	10 packets (1g each)
	Instructions for Use (IFU)	1 copy
	Protein Finder	1 piece
	Intensity Finder	1 piece
	Forceps	1 pair
	Disposable 9-well incubation tray (manual use only and packed separately from the kit)	2 or 4 trays

15. Using forceps, gently remove strips and place on laboratory paper towel or Office Printer paper. Allow to dry for 30 minutes. Alternatively, allow the strips to dry in the wells of the tray for 3 hours.
16. Mount strips on the Report Sheet or equivalent worksheet (non-absorbent white paper). If using adhesive tape for mounting, do not apply over the developed bands. Observe the bands (See Interpretation of Results) and grade the results within 24 hours of drying. For storage, keep the strips in the dark.

SUMMARY OF ASSAY PROTOCOLS		
Reagents	Qty	Duration
Nitrocellulose strip	1 strip	-
Wash Buffer	2 mL	5 mins
Blotting Buffer	2 mL	-
Specimen	20 µL	60 mins
Wash Buffer	3 x 2 mL	3 x 5 mins
Conjugate	2 mL	60 mins
Wash Buffer	3 x 2 mL	3 x 5 mins
Substrate (Ready to use)	2 mL	15 mins
Reagent or Deionized Water	3 x 2 mL	-

#### QUALITY CONTROL

The Non-Reactive Control, Strong Reactive Control I, and Strong Reactive Control II must be run with the assay regardless of the number of samples tested. **Figure 1** shows the appearance of these control strips.

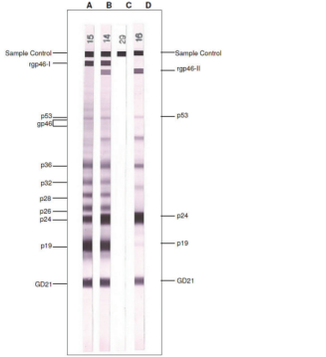


Figure 1: Examples of control strips

- A. Strong Reactive Control I. (Reactive for HTLV-I only)
- B. HTLV-II serum
- C. Non-Reactive Control
- D. Strong Reactive Control II (Reactive for HTLV-II only)

In order for the results obtained from any assay to be considered valid, the following conditions must be met:

1. **NON-REACTIVE CONTROL**  
No HTLV-I viral specific bands, rgp46-I, rgp46-II or GD21 should be observed on the Non-Reactive control strip. The sample control (anti-human IgG) band should be visible.

HTLV Blot 2.4 Report Sheet		1 piece
<b>WARNINGS AND PRECAUTIONS</b>		
	<b>CAUTION: Test kit should be handled only by qualified personnel trained in laboratory procedures and familiar with their potential hazards.</b>	
This kit contains materials of human origin. No test method can offer complete assurance that human blood products will not transmit infection. Follow established laboratory policy and applicable CDC/NIH biosafety and/or OSHA/NIH hazardous material spill guidelines for appropriate hazardous chemical and/or biological spill response and clean-up.		

**HANDLE ASSAY SPECIMENS, STRONG REACTIVE CONTROL I, STRONG REACTIVE CONTROL II, AND NON-REACTIVE CONTROL AS POTENTIALLY INFECTIOUS AGENTS.** It is recommended that the kit components and test specimens be handled with universal precautions as if capable of transmitting infectious disease. Refer to guidelines from the current CDC/NIH *Biosafety in Microbiological and Biomedical Laboratories* or equivalent, for safe practices in handling specimens. Specimens should be disposed of in accordance with established safety procedures.

The Strong Reactive Control I, Strong Reactive Control II and Non-Reactive Control contain both thimerosal and sodium azide as preservatives; the Lyophilized Stock Buffer and the Wash Buffer Concentrate contain thimerosal and the Conjugate contains sodium azide. Sodium Azide may react with lead or copper plumbing to form highly explosive metal azides. Build up in piping has led to laboratory explosions. Therefore, dilute and/or flush with copious amounts of water when disposing down the drain. Check with your local, regional, or national ordinances accordingly.

The ingredients present in the kit components are, in their pure form, a dangerous substance. However, their low concentrations, as prepared in these kit components, are not considered a dangerous substance. Sodium azide < 0.1% w/v is below the regulatory threshold limits according to OSHA standard 29 CFR 1910.1200.

The substrate, BCIP/NBT, can potentially be irritating to the skin and eyes.

Pursuant to EC regulation 1272/2008 (CLP), hazardous components are classified and labelled as follows:

Component:	Nitrocellulose strips
<b>Signal Word:</b>	<b>Danger</b>
<b>Pictogram:</b>	
<b>Hazard Statements:</b>	H228 Flammable solid
<b>Precautionary Statements:</b>	P210 Keep away from heat/sparks/open flames/hot surfaces. – No smoking. P280 Wear protective gloves/protective clothing/eye protection/face protection.
<b>Supplemental Statements:</b>	EUH 210 Safety Data Sheet is available on request
<b>Contains:</b>	100% Nitrocellulose

Component:	WASH BUFFER CONCENTRATE (20x)
<b>Signal Word:</b>	Warning
<b>Pictogram:</b>	
<b>Hazard Statements:</b>	H373 May cause damage to organs through prolonged or repeated exposure.
<b>Precautionary Statements:</b>	P260 Do not breathe dust/ fume/ gas/ mist/ vapours/ spray. P501 Dispose of contents/ container in accordance with local/regional/national/international regulations.
<b>Supplemental Statements:</b>	EUH 210 Safety Data Sheet is available on request
<b>Contains:</b>	0.1% Thimerosal

2. **STRONG REACTIVE CONTROL I**  
The relevant HTLV bands must be present are p19, p24, rgp46-I and GD21. The sample control (anti-human IgG) band should be visible.

**Note:** Although uncommon, a gp46 viral band may be present. If present, it appears as a diffuse band. Because of the rarity of gp46 and misreading of viral bands in this molecular weight range, viral gp46 is not used as part of the assay's interpretative criteria.

