





The major HTLV-I/II gene products that have been identified are listed in Table 1.

Table 1: Listing of major HTLV-I/II Gene Products

Band	Gene product	HTLV-I/II
rgp46-I	Recombinant env glycoprotein	I
rgp46-II	Recombinant env glycoprotein	II
p53	Precursor of gag protein	I
gp46	Outer env glycoprotein	I
p36	gag protein intermediate	I
p32	gag protein intermediate	I
p28	gag protein intermediate	I
p26	gag protein intermediate	I
p24	Major gag capsid protein	I
p19	Major gag matrix protein	I
GD21	Recombinant transmembrane ENV protein	I/II

The above bands are the only bands that should be read and considered in the interpretation of test samples.

Use the following guidelines to determine the interpretation of test samples:

#### SERONEGATIVE INTERPRETATION:

- No reactivity to HTLV specific proteins; or
- Any combination of gag proteins excluding p24 (p19, p26, p28, p32, p36, p53); or
- Any single gag protein other than p19 or p24 (p26, p28, p32, p36, p53).

	GD21 Recombinant env Protein	p19 Major gag Matrix	p24 Major gag Capsid	rgp46-I Recombinant env Protein	rgp46-I Recombinant env Protein	Non-major gag Proteins* (p26, p28, p32, p36, p53)
Seronegative						
		X				X
						X
						X

#### HTLV-I SEROPOSITIVE:

(Note: The non-major gag proteins (p26, p28, p32, p36, p53) may or may not be present and are not utilized in determining HTLV-I seropositivity)

- Reactivity to p19, GD21 and rgp46-I; or
- Reactivity to p19, p24 and GD21, with reactivity to p19 greater than or equal to p24;

	GD21 Recombinant env Protein	p19 Major gag Matrix	p24 Major gag Capsid	rgp46-I Recombinant env Protein	rgp46-I Recombinant env Protein
HTLV-I Seropositive					
	X	X		X*	X
	X	X			X
	X	X*	X		

\*reactivity to p19 ≥ p24  
\*\* low level reactivity

= 20), multiparous women (n = 10), high rheumatoid factor (n = 20), Hashimoto's disease (n = 10), and Sjogren's disease (n = 10). Specimens were tested for HTLV-I and HTLV-II positive specimens. Specimen results are shown in Table 7. Of the 200 spiked samples, all but one of the samples remained positive; one sample from the EBV population was resulted as indeterminate. Of the 200 unspiked samples tested, 4 were resulted as positive; one each from the HIV, dialysis, hemophilic, and Sjogren's populations. The specimen from the dialysis population was determined to be a true positive based on subsequent testing, and was further excluded from calculations.

Table 7: Effect of Unrelated Medical Conditions

Potentially Interfering Medical Condition	Number of Specimens Tested	Number of Specimens Positive in Unspiked Population	Number of Specimens Negative in Spiked Population
HIV	20	1	0
HCV	20	0	0
HBV	20	0	0
EBV	20	0	0
CMV	20	0	0
Influenza Vaccine	10	0	0
Hemophilic	20	1	0
Dialysis	19	0	0
Multiparous Women	10	0	0
Elevated Rheumatoid Factor	20	0	0
Hashimoto's disease	10	0	0
Sjogren's disease	10	1	0

The high strip background obscured the reading of bands in 7 out of 200 cross-reactive samples (3.5%). This affected both the unspiked samples and spiked HTLV-positive samples. The principal observation was the appearance of randomly occurring critical bands with many of the unspiked samples. This occurred across all twelve populations, and it caused many unspiked samples to be interpreted as indeterminate, rather than the expected negative interpretation. From this data, it is reasonable to conclude that the presence of the potential cross reactants increases the frequency of indeterminate results and may produce erroneous results.

#### LIMITED EXPRESSED WARRANTY DISCLAIMER

The manufacturer makes no expressed warranty other than that the test kit will function as an in-vitro diagnostic assay within the specifications and limitations described in the Instructions for Use (IFU) when used in accordance with the instructions contained therein. The manufacturer disclaims any warranty expressed or implied, including such expressed or implied warranty with respect to merchantability, fitness for use or implied utility for any other purposes. 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 however caused by the product in the use or in the application thereof.

#### TECHNICAL PROBLEMS / COMPLAINTS

Should there be a technical problem / complaint, please do the following:

- Note the kit lot, strip lot number and the expiration date.
- Retain the kits and the results that were obtained.
- Contact MP Biomedicals' Customer Service.

#### HTLV-II SEROPOSITIVE:

(Note: The non-major gag proteins (p26, p28, p32, p36, p53) may or may not be present and are not utilized in determining HTLV-II seropositivity)

- Reactivity to p24, GD21 and rgp46-II; or
- Reactivity to p19, p24 and GD21, with reactivity to p24 greater than p19\*\*

	GD21 Recombinant env Protein	p19 Major gag Matrix	p24 Major gag Capsid	rgp46-II Recombinant env Protein	rgp46-I Recombinant env Protein
HTLV-II Seropositive					
	X		X	X	
	X		X	X	X*
	X	X	X	X	
	X	X	X*		

\*reactivity to p24 > p19  
\*\* low level reactivity

#### HTLV-I/II SEROPOSITIVE:

(Note: The non-major gag proteins (p26, p28, p32, p36, p53) may or may not be present and are not utilized in determining HTLV-I/II seropositivity)

- Reactivity to GD21, p19, p24, rgp46-I and rgp46-II

HTLV-I/II Seropositive	GD21 Recombinant env Protein	p19 Major gag Matrix	p24 Major gag Capsid	rgp46-II Recombinant env Protein	rgp46-I Recombinant env Protein
	X	X	X	X	X

#### INDETERMINATE:

- Reactivity to HTLV specific bands that do not meet the criteria for HTLV-I seropositive, HTLV-II seropositive, HTLV-I/II seropositive or seronegative. The list below includes some, but not all, of the indeterminate band pattern patterns:

Common Indeterminate Band Patterns					
p19 only p19, p24 rgp46-I, GD21 GD21, p24, rgp46-I	p24 only GD21, p19 rgp46-I, rgp46-II GD21, p19, rgp46-II	GD21 only GD21, p19, rgp46-II p19, p24, p26, p28, p32, p53			

- The p36 band is not associated with HTLV-II infection. A band with similar molecular weight may appear with HTLV-II samples and should be disregarded.
- HTLV-I gag indeterminate Western Blot patterns (HGIP) refer to the presence of p19, p26, p28, p32, p36, p53 (in various combinations) but absence of p24 and any ENV proteins. While HGIP would be interpreted as HTLV seroindeterminate based on 1990 guideline<sup>62</sup>, various studies suggested that HGIP should be interpreted as seronegative especially with healthy blood donors.<sup>63</sup>
- Comparison of reactivity is based on intensity and broadness of band. If intensity of bands is similar, the reactivity is determined by comparing broadness of band.
- The p24 gag protein from the HTLV-I viral lysates cross-reacts with HTLV-II antibodies and is therefore used as an HTLV-II diagnostic marker.<sup>67-71</sup>
- If an indeterminate result occurs, a fresh sample should be obtained for repeat HTLV antibody testing.

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low level bands. Donor specimens seropositive by the criteria of the HTLV Blot 2.4 using s bands only should be retested using a fresh sample to confirm infection.

#### SPECIFIC PERFORMANCE CHARACTERISTICS

- MP Diagnostics HTLV Blot 2.4 Performance Characteristics in Known Positive Population and Normal Blood Donors**  
The performance of the MP Diagnostics HTLV Blot 2.4 was evaluated in clinical studies on blood donor populations by comparison of HTLV Blot 2.4 results with those obtained from matched plasma specimens tested using the California Department of Public Health (CDPHL) HTLV Supplemental Algorithm.<sup>1</sup> Sensitivity was evaluated using a known positive population, characterized as archival specimens from deferred blood donors who had tested repeatedly reactive by at least one licensed HTLV screening assay and were confirmed positive through additional, research use supplemental assays, including, IFA, Western blot and RIPA. Specificity was evaluated using archival specimens from normal volunteer blood donors that had tested HTLV non-reactive by a licensed screening assay. The MP Diagnostics HTLV Blot 2.4 testing was performed at three, geographically distinct clinical testing sites.

#### 1.1 Sensitivity in Known Positive Population

A total of 200 repository specimens from a well-characterized, known positive population were evaluated at three geographically distinct clinical testing sites. These specimens were from deferred donors that had previously tested repeatedly reactive using a licensed screening assay in conjunction with research use HTLV supplemental testing, including ELISA, IFA, Western blot and RIPA. The summary results from testing the known positive population are shown in Table 1.

Table 1: MP Diagnostics HTLV Blot 2.4 and CDPHL HTLV Supplemental Algorithm Results for 200 Known Positive Specimens

MP Diagnostics HTLV Blot 2.4	CDPHL Algorithm				
	HTLV-I POS	HTLV-II POS	IND	NEG	Total
	HTLV-I POS	HTLV-II POS	IND	NEG	Total
HTLV-I POS	79	1	1	0	81
HTLV-II POS	0	100	0	4	104
HTLV-I/II POS	9	1	0	0	10
IND	1	2	0	1	4
NEG	0	0	0	1*	1
Total	89	104	1	6	200

\* One sample was negative by both the MP Diagnostics HTLV Blot 2.4 and the CDPHL HTLV Algorithm.

A greater number of known positive specimens were identified as positive by the MP Diagnostics HTLV Blot 2.4 than by the CDPHL HTLV Algorithm (195 versus 193, respectively). Additionally, the MP Diagnostics HTLV Blot 2.4 identified more samples as reactive (i.e., positive or indeterminate) than the CDPHL HTLV Algorithm (199 versus 194, respectively). Of the 195 specimens identified as Positive by the HTLV Blot, 185 (81 + 104) were interpreted as HTLV-I Positive or HTLV-II Positive, and 10 (9 + 1) were HTLV-I/II Positive. Additionally, the HTLV Blot 2.4 identified more samples as reactive (i.e., Positive or Indeterminate) than the CDPHL Algorithm (199 versus 194, respectively). Of the six samples identified as Negative by the CDPHL Algorithm, four were identified as HTLV-II Positive by the MP Diagnostics HTLV Blot 2.4.

Although these 200 specimens were previously identified as Positive for HTLV antibodies using the CDPHL algorithm, six specimens were Negative and one was Indeterminate on retesting by the CDPHL Algorithm. This Indeterminate specimen was determined to be HTLV-I Positive by the MP Diagnostics HTLV Blot 2.4. One sample was negative by both the HTLV Blot 2.4 and the CDPHL Algorithm.

In this study, the sensitivity of the MP Diagnostics HTLV Blot 2.4 was 97.5%<sup>62</sup> (195/200) with a 95% CI of 94.26 - 99.18%. The indeterminate rate for this study was 2% (4/200).

<sup>1</sup> A licensed, HTLV supplemental assay was not available at the time of testing. The CDPHL HTLV Supplemental Algorithm consists of a series of in-house developed, HTLV supplemental assays. The HTLV Algorithm includes the following assays in sequence: ELISA; IFA; Western Blot; & RIPA. The number of assays that a sample will be tested with is dependent upon the sample results within the HTLV Algorithm.

<sup>2</sup> Sensitivity was calculated as follows: TP/(TP+FN) x 100% where TP = true positives, that is, the number of specimens positive by MP Diagnostics HTLV Blot 2.4; and FN = false negatives, that is, the number of specimens indeterminate or negative by MP Diagnostics HTLV Blot 2.4.

#### 1.2 Specificity in Normal Blood Donors Testing HTLV Non-reactive by a Licensed Screening Assay

A total of 200 repository specimens from a normal blood donor population were evaluated at three geographically distinct clinical testing sites. These specimens were from blood donors that had previously tested HTLV non-reactive using a licensed HTLV screening assay. The summary results from testing the HTLV screening assay negative population are shown in Table 2.

Table 2: MP Diagnostics HTLV Blot 2.4 and CDPHL HTLV Supplemental Algorithm Results on HTLV Screening Assay Negative Population

MP Diagnostics HTLV Blot 2.4	CDPHL Algorithm			
	POS	IND	NEG	Total
	POS	IND	NEG	Total
POS	0	0	0	0
IND	0	0	43	43
NEG	0	0	157	157
Total	0	0	200	200

The MP Diagnostics HTLV Blot 2.4 identified 157 as negative and 43 as indeterminate; there were no positive samples identified in this population. Of these 200 specimens tested by the CDPHL HTLV Algorithm, 15 were repeatedly reactive by ELISA. The majority of these repeatedly reactive samples were resolved at the Western blot stage of the CDPHL HTLV Algorithm, based on non-reactivity from both the IFA and Western blot. Two of these 15 samples, however, showed reactivity with the p21e protein on the Western blot and were subjected to additional testing using RIPA. A non-reactive result on the RIPA for these 2 specimens resulted in an overall call of negative by the CDPHL Algorithm. All but one of these 15 specimens was negative by a single MP Diagnostics HTLV Blot 2.4 assay.

In this study the indeterminate rate of MP Diagnostics HTLV Blot 2.4 for licensed HTLV-I/II ELISA negative specimens was 21.5% (43/200)

#### 2. Comparative Testing of Repeatedly Reactive Specimens Identified by Specific Licensed HTLV-II Screening Tests

A total of 200 repeatedly reactive samples were evaluated at three geographically distinct clinical testing sites. These specimens were from blood donors that had previously tested repeatedly reactive using the Abbott PRISM HTLV-I/II CLIA. Of these 200 samples, the MP Diagnostics HTLV Blot 2.4 identified 3 as positive, 88 as negative and 109 as indeterminate or equivocal (Table 3). Comparatively, the CDPHL HTLV Algorithm identified 3 as inconclusive and 197 as negative. Follow-up testing that was available on one donor confirmed that the MP Diagnostics HTLV Blot 2.4 had correctly identified that specimen as positive. Additionally, three inconclusive CDPHL HTLV Algorithm samples that were identified as negative by the MP Diagnostics HTLV Blot 2.4 were confirmed as negative during donor follow-up; the CDPHL HTLV Algorithm result of inconclusive was due to a false positive western blot that used the less specific p21e recombinant.

In this study the indeterminate rate of MP Diagnostics HTLV Blot 2.4 for Abbott PRISM HTLV-I/II false positive specimens was 55% (109/197)

Table 3: Performance of MP Diagnostics HTLV Blot 2.4 against the CDPHL algorithm with samples that are RR on Abbott PRISM HTLV-I/II screening test

MP Diagnostics HTLV Blot 2.4	CDPHL Algorithm			
	POS	IND	NEG	Total
	POS	IND	NEG	Total
POS	0	0	3	3
IND	0	0	109	109
NEG	0	3	85	88
Total	0	3	197	200

A total of 105 preselected repository samples that were repeatedly reactive using the Abbott HTLV-I/II HTLV-II Microelisa System were evaluated at one clinical testing site as well as in-house at MP Biomedicals, LLC. Of these 105 samples, the MP Diagnostics HTLV Blot 2.4 identified 50 as positive, 18 as indeterminate and 37 as negative (Table 4). Comparatively, the CDPHL HTLV Algorithm identified 50 as positive, 51 as negative and 4 as inconclusive or equivocal. The percent positive agreement of the MP Diagnostics HTLV Blot 2.4 with the CDPHL HTLV Algorithm was 100% and the overall percent agreement was 82.18% (95% CI of 73.30 to 89.08%). The four CDPHL HTLV Algorithm

inconclusive results were due to the presence of p21e; all sample results were resolved as negative by the MP Diagnostics HTLV Blot 2.4 due to the inclusion of GD21, a more specific envelope reagent.

In this study the indeterminate rate of MP Diagnostics HTLV Blot 2.4 for Avioq HTLV-I/II Microelisa System false positive was 35% (18/51)

Table 4: Performance of MP Diagnostics HTLV Blot 2.4 against the CDPHL algorithm with samples that are RR on Avioq HTLV-I/II Microelisa System

MP Diagnostics HTLV Blot 2.4	CDPHL Algorithm			
	POS	IND	NEG	Total
	POS	IND	NEG	Total
POS	50	0	0	50
IND	0	0	18	18
NEG	0	4	33	37
Total	50	4	51	105

Among the 50 positive specimens, the HTLV Blot 2.4 identified 18 as HTLV-I and 29 as HTLV-II, and three specimens as HTLV-I/II Undifferentiated (see Table 5). In comparison, the CDPHL Algorithm identified 15 as HTLV-I, 29 as HTLV-II, and 8 as HTLV-I/II Undifferentiated. These data indicated overall agreement between the HTLV Blot 2.4 and the CDPHL algorithm to differentiate HTLV-I and HTLV-II infections with concordant differentiation by the HTLV Blot 2.4 of 13/15 specimens categorized as HTLV-I by the CDPHL algorithm and 27/27 specimens categorized as HTLV-II by the CDPHL algorithm.

Table 5: Differentiation of positive specimens against the CDPHL algorithm for those RR using the Avioq HTLV-I/II Microelisa System

MP Diagnostics HTLV Blot 2.4	CDPHL HTLV Algorithm			
	HTLV-I POS	HTLV-II POS	HTLV-II POS Undifferentiated	Total
	HTLV-I POS	HTLV-II POS	HTLV-II POS Undifferentiated	Total
HTLV-I POS	13	0	5	18
HTLV-II POS	0	27	2	29
HTLV-II POS Undifferentiated	2	0	1	3
Total	15	27	8	50

#### 3. Reproducibility

The reproducibility of the MP Diagnostics HTLV Blot 2.4 assay was established in a study that assessed assay reproducibility within operators, within site, and between lots. This study tested two replicates of a three-member panel at three clinical sites with each of three product lots over multiple days by three operators. The three-member panel consisted of one HTLV-I antibody specimen, one HTLV-II antibody specimen, and one specimen non-reactive to antibodies for HTLV-I/II. For each of the three kit lots, there were a total of 54 HTLV Blot 2.4 strips tested with each panel member. Reproducibility was calculated as percent agreement of positive results / negative results.

In this study, no strips were incorrectly interpreted. These data demonstrate that the MP Diagnostics HTLV Blot 2.4 assay is reproducible across multiple sites, operators and lots.

#### 3.1 Reproducibility of Test Strip Interpretation at 24 Hours post assay

Between June 16, 2013 to December 10, 2015, 950 unique HTLV RFR plasma specimens were evaluated with the HTLV Blot 2.4. Of the 945/950 (99.5%) per-protocol specimens evaluated, 491/945 (52.0%) were HTLV seropositive, 271/945 (28.7%) seronegative, 180/945 (19.0%) indeterminate and 3/945 (0.31%) invalid. HTLV Blot 2.4 test result interpretation agreement within 3 & 24 hours post assay is summarized below:

HTLV Blot 2.4 Test Result	Result Interpretation within 3 Hrs. n (%)	Result Interpretation Agreement at
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