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National Laboratory for HIV Reference Services Sexually Transmitted and Bloodborne Infections National Microbiology Laboratory Public Health Agency of Canada

HTLV Serology Quality Assessment Program <u>Summary for Panel HTLVSER 2022Apr19</u>

2022Apr19 HTLV Serology Panel							
Panel Sample	True Status	Labs Reporting Incorrect Status					
А	Negative						
В	HTLV-I Ab Positive						
С	HTLV-II Ab Positive						
D	HTLV-I Ab Positive						
E	Negative						

Summary of finding observed for the 2022Apr19 panel:

1) Participant HV22 and HV80 did not return results in this test event.



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HTLV Serology Quality Assessment Program Final Report for Panel HTLVSER 2022Ap19

Issued 2022-July-14

Introduction

The NLHRS distributed the 2021Oct29 and 2022Apr19 panels on October 20, 2021. This final report is specific to the 2022Apr19 panel only and is publicly available; however, the identity of participants has not been disclosed. The deadline for results submission was April 19, 2022. The preliminary report was issued on May 20, 2022.

Panel Samples, HTLV Test Kits, and Data Entry

- Panel Composition
 - The 2022Apr19 panel consisted of five samples: two HTLV negative (A, E), two HTLV-I positive (B,D), and one HTLV-II positive sample (C). Samples B, C and D were diluted 1 in 2 with defibrinated human plasma (Basematrix 53, Seracare Life Sciences). Testing and characterization by the NLHRS are presented in Appendix 1. Panels were sent to 18 participants including the NLHRS on October 20, 2021.
 - The metrological traceability and uncertainty is not applicable for this panel.
- HTLV Test Kits
 - Five different assays were used by the 16 participants (excluding the NLHRS) who returned results (Appendix 2).
- Data entry
 - Results entry for this panel utilized an in-house developed website.

Homogeneity and Stability

- The homogeneity and stability of the 2022Apr19 HTLV serology panel was assessed by comparing the participants' results (including the NLHRS) with the results of the panel's characterization performed by the NLHRS prior to the test event.
- There was no indication of heterogeneity or instability of the panel samples as the results submitted by the participants are consistent with the expected results from the NLHRS characterization of each panel member (Figure 1 and Appendix 1).

Results

- Evaluation Criteria:
 - Negative samples: HTLV non-reactive/negative in the final HTLV serology interpretation with assay results supporting the interpretation.
 - Positive samples: HTLV reactive/positive in the final HTLV serology interpretation with assay results supporting the interpretation. Participants must provide a recommendation for further action for samples that they could not determine the true serology status for based on the assay used in their testing.
- Qualitative Group Analysis (Figure 1):
 - Sample A (Negative) 16/16 participants provided either a correct serology status and/or recommendation.
 - Sample B (HTLV-I Ab Positive) 16/16 participants provided either a correct serology status and/or recommendation.
 - Sample C (HTLV-II Ab Positive) 16/16 participants provided either a correct serology status and/or recommendation.
 - Sample D (HTLV-I Ab Positive) 16/16 participants provided either a correct serology status and/or recommendation.
 - Sample E (Negative) 16/16 participants provided either a correct serology status and/or recommendation.

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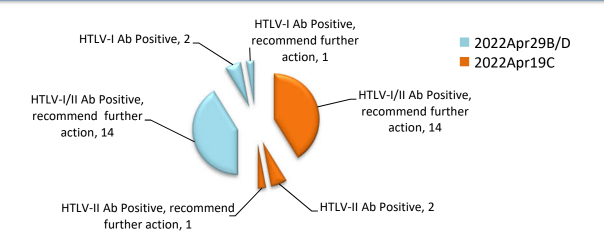


Figure 1: The final HTLV serology status of the positive samples in the 2022Apr19 HTLV serology panel submitted by participants using HTLV screening and confirmatory assays (including NLHRS).

Findings

Two participants did not return results for this test event.

Since the 2021Apr19 test event, we have noticed several of the Abbott Architect users adopting the newer Abbott Alinity platform. We anticipate this trend will continue in the future as more laboratories adopt the newer Abbott platform.

We value each laboratory's participation in these QA test events and your suggestions for improvement. The NLHRS is committed to improve all aspects of the HTLV serology proficiency testing program in order to provide quality proficiency testing to our participants.

If you have any comments, suggestions or concerns, please contact us at:

nlhrs.qap-peq.lnsrv@phac-aspc.gc.ca

Thank you for your participation in the NLHRS HTLV Serology Quality Assurance Program

John Ad

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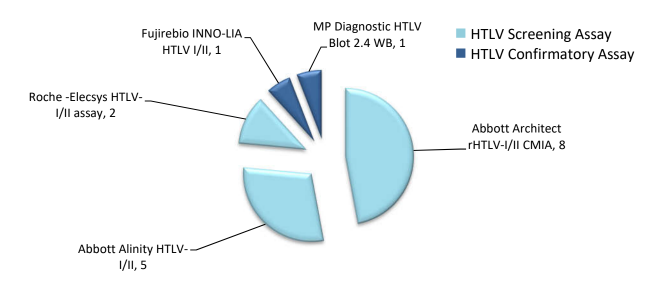
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Appendix 1: NLHRS characterization of the 2022Apr19 HTLV serology panel samples

The NLHRS 2022Apr19 HTLV Panel Sample Testing Results											
		NLHRS Testing									
Sample Final Status Fujirebio INNO-LIA HTLV I/II Sco											
		Interpretation	p19 I/II	p24 I/II	gp46 I/II	gp21 I/II	p19 I	gp46 I	gp46 II		
Α	Negative	Negative	-	-	-	-	-	-	-		
В	HTLV-I Ab Positive	HTLV-I Positive	++	++	++	+++	++	++	-		
C	HTLV-II Ab Positive	HTLV-II Positive	++	+++	+++	+++	+	-	++		
D	HTLV-I Ab Positive	HTLV-I Positive	++	++	++	+++	++	++	-		
E	Negative	Negative	-	-	-	-	-	-	-		

N/T: Not tested



Appendix 2: Summary of assays used by the participants in the 2022Apr19 HTLV test event

Appendix 3: Summary of bands detected in samples B, C, and D by the Fujirebio INNO-LIA HTLV-I/II and MP Diagnostic HTLV Blot 2.4 WB assays in the 2022Apr19 HTLV test event

Fujirebio INNO-LIA HTLV-I/II	Frequency of Bands Detected									
Sample	p19 I/II	p24 I/II	gp46 I/II	gp21 l/ll	p19-l	gp46-l	gp46-II			
2022Apr19B	2	2	2	2	2	2	-			
2022Apr19C	2	2	2	2	1	-	2			
2022Apr19D	2	2	2	2	2	2	-			

MP Diagnostic HTLV Blot 2.4 WB	Frequency of Bands Detected										
Sample	rgp46-I	rgp46-II	p53	gp46	p36	p32	p28	P26	P24	P19	GD21
2022Apr19B	1	-	1	1	1	1	1	1	1	1	1
2022Apr19C	-	1	1	-	1	1	-	-	1	-	1
2022Apr19D	1	-	1	1	1	1	1	1	1	1	1

Appendix 4: Troubleshooting

Troubleshooting; common causes of outlying and/or aberrant results in serology and molecular Laboratories.

Type of Error	Possible Cause(s)	Pre-Analytical	Analytical	Post- Analytical					
Sample	Can occur during specimen reception or testing. May result in	✓	✓						
mix-up	outlying/aberrant results for one or all samples mixed-up.	•	•						
	 Incorrect test ordering by physician 	✓							
	 Incorrect shipment address 	✓							
	 Selecting the wrong assay for data entry 	\checkmark							
	 Interchanging results for two or more specimens 			\checkmark					
	• Entering incorrect results			✓					
	• Entering values in the incorrect field (e.g., OD as S/Co)			✓					
Transcription	• Entering values in the incorrect unit (e.g., IU/mL instead of			✓					
	log10 copies/mL)			v					
	• Using a comma instead of a dot to denote a decimal point			✓					
	Selecting the incorrect assay interpretation or analyte			✓					
	• Failure to recommend follow-up testing where necessary			✓					
	It is recommended all results that are manually transcribed or entered electronically be checked by a second								
	individual to avoid transcription errors.								
	Sporadic test results identified as outlying and/or aberrant can be classified as random events. Possible causes of								
	random error include:								
	 Incorrect sample storage/shipping conditions 	✓	✓						
Outlying	Incorrect test method	✓	✓						
and/or	Insufficient mixing of sample, especially following freezing		✓						
Aberrant	Poor pipetting		✓						
Results (<u>random error</u>)	Ineffective or inconsistent washing		✓						
(<u>random enor</u>)	Transcription errors	✓		✓					
	Cross-contamination or carryover	✓	✓						
	Presence of inhibitors to PCR		✓						
	A series of test results identified as outlying and/or aberrant may be due to a systematic problem. Systematic								
	problems may be due to:								
	• Reagents contaminated, expired, or subject to batch variation		✓						
	Instrument error or malfunction		✓						
	Insufficient washing		✓						
Outlying	Incorrect wavelength used to read the assay result		✓						
and/or Aberrant Results (<u>systematic</u> <u>error</u>)	Cycling times too long/short or temperature too high/low		✓						
	Incubation time too long/short or temperature too high/low		✓						
	Insufficient mixing/centrifuging before testing		✓						
	 Incorrect storage of test kits and/or reagents 	✓							
	Contamination of master-mix, extraction areas or equipment		✓						
	Ineffective extraction process		✓						
	Degradation of master-mix components		✓						
	Suboptimal primer design (in-house assays)		✓						
				l					

This table was modified from a report produced by the National Reference Laboratory (NRL), Melbourne, Australia.