Individuals using assistive technology may not be able to fully access the information contained in this file. For assistance, please send an e-mail to: ocod@fda.hhs.gov and include 508 Accommodation and the title of the document in the subject line of your e-mail.

HTLV I/II Reagent Kit

Human T-Lymphotropic Virus Types I and II (E coli, Recombinant) Antigen and Synthetic Peptides

Revised June 2019

Instructions must be carefully followed. Reliability of assay results cannot be guaranteed if there are any deviations from these instructions.

■NAME

Alinity s HTLV I/II Reagent Kit

Human T-Lymphotropic Virus Types I and II (E coli, Recombinant) Antigen and Synthetic Peptides

■INTENDED USE

The Alinity s HTLV I/II assay is a chemiluminescent microparticle immunoassay (CMIA) used for the qualitative detection of antibodies to human T-lymphotropic virus Type I and/or human T-lymphotropic virus Type II (anti-HTLV I/HTLV II) in human serum and plasma specimens on the Alinity s System.

The Alinity s HTLV I/II assay is intended to screen individual human donors, including volunteer donors of whole blood and blood components, and other living donors for the presence of anti-HTLV I/HTLV II. The assay is also intended for use in testing serum and plasma specimens to screen organ donors when specimens are obtained while the donor's heart is still beating, and in testing serum specimens to screen cadaveric (non-heart-beating) donors. It is not intended for use on cord blood specimens.

SUMMARY AND EXPLANATION OF THE TEST

Human T-lymphotropic virus Type I (HTLV I) and Type II (HTLV II) are closely related but distinct retroviruses that can infect humans. HTLV I causes adult T-cell leukaemia (ATL) and HTLV I-associated myelopathy/tropical spastic paraparesis (HAM/TSP).¹ Although HTLV II is less pathogenic than HTLV I, it has been associated with a neurological disease similar to HAM/TSP^{2,3} and with chronic inflammatory arthropathy.¹

HTLV I infection is endemic in south Japan⁴, the Caribbean⁵, in some regions of Africa⁶. Central and South America⁷ and also found in Melanesia,⁸ the Middle East,⁹ and central and northern Australia.^{10,11} HTLV II infection is endemic to a number of indigenous American Indian populations.^{7,12} Both HTLV I and HTLV II are distributed worldwide.

HTLV I and HTLV II were the first discovered human retroviruses, 13,14 both viruses belonging to the oncovirus subfamily of retroviruses.¹⁵ Unlike HIV retroviruses, HTLV I and HTLV II show minimal genetic variation, mainly in the env, which defines the HTLV subtypes.¹⁶ HTLV I has six reported subtypes (subtypes A to F).¹⁷ HTLV II has four reported subtypes (subtypes A to D).^{18,19} However, there is no reported association of a particular HTLV I or HTLV II subtype with a specific disease.19,20

Transmission of HTLV I and HTLV II infection occurs via transfusion of infected cellular blood products, ²¹⁻²⁶ via breast feeding, ²⁷⁻³⁰ sexual contact,³¹ and sharing of contaminated needles and syringes by intravenous drug users.^{32,33} Mother-to-child transmission of HTLV II has recently been reported.34

HTLV I and HTLV II antibodies develop within 4 to 8 weeks after infection. Most individuals infected with HTLV I and HTLV II are asymptomatic, and the infection is lifelong.35

HTLV I/HTLV II antibody assays are used to identify individuals infected with HTLV I or HTLV II and to prevent transmission of the virus to recipients of blood, blood components, and organs.

■BIOLOGICAL PRINCIPLES OF THE PROCEDURE

This assay is a two-step immunoassay for the qualitative detection of antibodies to HTLV I and HTLV II in human serum and plasma using chemiluminescent microparticle immunoassay (CMIA) technology

Sample, HTLV I/HTLV II coated paramagnetic microparticles, and assay diluent are combined and incubated. The antibodies to HTLV I/HTLV II present in the sample bind to the HTLV I/HTLV II synthetic peptides and HTLV II recombinant antigen coated microparticles. The mixture is washed. HTLV I/HTLV II synthetic peptides and HTLV I recombinant antigen acridinium-labeled conjugate is added to create a reaction mixture and incubated. Following a wash cycle, Pre-Trigger and Trigger Solutions are added.

The resulting chemiluminescent reaction is measured as relative light units (RLU). There is a direct relationship between the amount of antibodies to HTLV I/HTLV II in the sample and the RLU detected by the system optics.

The presence or absence of antibodies to HTLV I/HTLV II in the sample is determined by comparing the chemiluminescent RLU in the reaction to the cutoff RLU determined from an active calibration.

For additional information on system and assay technology, refer to the Alinity s System Operations Manual, Section 3.

REAGENTS

Kit Contents

Volumes (mL) listed in the table below indicate the volume per cartridge.

| REF | 06P0760 |
|------------------------------|---------|
| Tests per cartridge | 500 |
| Number of cartridges per kit | 10 |
| Tests per kit | 5000 |
| MICROPARTICLES | 27.0 mL |
| CONJUGATE | 26.5 mL |
| ASSAY DILUENT | 26.7 mL |

MICROPARTICLES HTLV I/HTLV II synthetic peptides and HTLV II recombinant antigen coated microparticles in TRIS buffered saline. Minimum concentration: 0.1% solids. Preservatives: ProClin 950 and sodium azide.

CONJUGATE HTLV I/HTLV II synthetic peptides and HTLV I recombinant antigen acridinium-labeled conjugate in HEPES buffer with protein (bovine) stabilizer and surfactant. Maximum concentration: Peptides 100.0 ng/mL each, antigen 10.0 ng/mL. Preservative: ProClin 950.

ASSAY DILUENT TRIS buffer with surfactant. Preservatives: ProClin 950 and sodium azide.







HTLV I/II 06P07 FDA R03 **B6P07E**

06P0760

REF

1 of 10

Alinity s HTLV I/II Reagent Kit 06P07

Warnings and Precautions

- IVD
- For In Vitro Diagnostic Use
- Performance characteristics of this product have not been established for laboratory diagnosis of HTLV I/II infection.

Safety Precautions

CAUTION: This product requires the handling of human specimens. It is recommended that all human-sourced materials be considered potentially infectious and handled in accordance with the OSHA Standard on Bloodborne Pathogens. Biosafety Level 2 or other appropriate biosafety practices should be used for materials that contain or are suspected of containing infectious agents.³⁶⁻³⁹

| The following warnings and precautions apply to: MICROPARTICLES | | | | |
|---|--|--|--|--|
| $\langle \mathbf{\hat{b}} \rangle$ | | | | |
| WARNING | Contains methylisothiazolone and sodium | | | |
| | azide. | | | |
| H317 | May cause an allergic skin reaction. | | | |
| EUH032 | Contact with acids liberates very toxic gas. | | | |
| Prevention | | | | |
| P261 | Avoid breathing mist / vapors / spray. | | | |
| P272 | Contaminated work clothing should not be | | | |
| | allowed out of the workplace. | | | |
| P280 | Wear protective gloves / protective | | | |
| | clothing / eye protection. | | | |
| Response | | | | |
| P302+P352 | IF ON SKIN: Wash with plenty of water. | | | |
| P333+P313 | If skin irritation or rash occurs: Get medical | | | |
| | advice / attention. | | | |
| P362+P364 | Take off contaminated clothing and wash it | | | |
| | before reuse. | | | |
| Disposal | | | | |
| P501 | Dispose of contents / container in | | | |
| accordance with local regulations. | | | | |

| The following warning | s and precautions apply to: CONJUGATE | | | |
|-----------------------|--|--|--|--|
| \Diamond | | | | |
| WARNING | Contains polyethylene glycol octylphenyl ether (Triton X-405) and methylisothiazolone. | | | |
| H317 | May cause an allergic skin reaction. | | | |
| H319 | Causes serious eye irritation. | | | |
| Prevention | | | | |
| P261 | Avoid breathing mist / vapors / spray. | | | |
| P264 | Wash hands thoroughly after handling. | | | |
| P272 | Contaminated work clothing should not be allowed out of the workplace. | | | |
| P280 | Wear protective gloves / protective clothing / eye protection. | | | |
| Response | | | | |
| P305+P351+P338 | IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing. | | | |
| P337+P313 | If eye irritation persists: Get medical advice / attention. | | | |

| - | |
|--|--|
| IF ON SKIN: Wash with plenty of water. | |
| | |
| If skin irritation or rash occurs: Get medical | |
| advice / attention. | |
| Take off contaminated clothing and wash it | |
| before reuse. | |
| | |
| Dispose of contents / container in | |
| accordance with local regulations. | |
| | |

| \Diamond | | |
|----------------|--|--|
| WARNING | Contains polyethylene glycol octylphenyl ether (Triton X-100), methylisothiazolone and sodium azide. | |
| H317 | May cause an allergic skin reaction. | |
| H319 | Causes serious eye irritation. | |
| EUH032 | Contact with acids liberates very toxic gas | |
| Prevention | | |
| P261 | Avoid breathing mist / vapors / spray. | |
| P264 | Wash hands thoroughly after handling. | |
| P272 | Contaminated work clothing should not be | |
| | allowed out of the workplace. | |
| P280 | Wear protective gloves / protective clothing / eye protection. | |
| Response | | |
| P305+P351+P338 | IF IN EYES: Rinse cautiously with water fc several minutes. Remove contact lenses, i present and easy to do. Continue rinsing. | |
| P337+P313 | If eye irritation persists: Get medical advice //attention. | |
| P302+P352 | IF ON SKIN: Wash with plenty of water. | |
| P333+P313 | If skin irritation or rash occurs: Get medical advice / attention. | |
| P362+P364 | Take off contaminated clothing and wash it before reuse. | |
| Disposal | · · | |
| P501 | Dispose of contents / container in accordance with local regulations. | |

Safety Data Sheets are available at www.transfusion.abbott or contact your local representative.

For a detailed discussion of safety precautions during system operation, refer to the Alinity s System Operations Manual, Section 8.

Reagent Handling

- Do not invert reagent cartridges.
- Upon receipt, reagent cartridges can be used immediately or stored in an upright position.
- If a reagent cartridge is dropped, place in an upright position for 1 hour before use to allow bubbles that may have formed to dissipate.
- Reagents are susceptible to the formation of foam and bubbles. Bubbles may interfere with the detection of the reagent level in the cartridge and cause insufficient reagent aspiration that may adversely affect results.

For a detailed discussion of reagent handling precautions during system operation, refer to the Alinity s System Operations Manual, Section 7.



Reagent Storage

• Do not freeze.

| | | Maximum | |
|----------|-------------|------------|---------------------|
| | Storage | Storage | Additional Storage |
| | Temperature | Time | Instructions |
| Unopened | 2 to 8°C | Until | Store in upright |
| | | expiration | position. |
| | | date | |
| Opened | 2 to 15°C | 15 days | Store in upright |
| | | after | position. |
| | | opening* | Discard after 15 |
| | | | days. |
| | | | If cartridge does |
| | | | not remain upright |
| | | | during storage off |
| | | | the system, discard |
| | | | the cartridge. |
| | | | Do not reuse |
| | | | original reagent |
| | | | caps or |
| | | | replacement caps |
| | | | due to the risk of |
| | | | contamination and |
| | | | the potential to |
| | | | compromise |
| | | | reagent |
| | | | performance. |

* Includes time on board the system.

Reagents may be stored on or off the system. If removed from the system, store reagents with new replacement caps in an upright position at 2 to 15°C. For reagents stored off the system, it is recommended that they be stored in their original trays or boxes to ensure they remain upright.

For information on unloading reagents, refer to the Alinity s System Operations Manual, Section 5.

Indications of Reagent Deterioration

Deterioration of the reagents may be indicated when a calibration error occurs or a control value is out of the specified range. Associated test results are invalid, and samples must be retested. Assay recalibration may be necessary.

For troubleshooting information, refer to the Alinity s System Operations Manual, Section 10.

■ INSTRUMENT PROCEDURE

The Alinity s HTLV I/II Assay File must be installed on the Alinity s System prior to performing the assay.

For detailed information on assay file installation and viewing and editing assay parameters, refer to the Alinity s System Operations Manual, Section 2.

For information on printing assay parameters, refer to the Alinity s System Operations Manual, Section 5.

For a detailed description of system procedures, refer to the Alinity s System Operations Manual.

SPECIMEN COLLECTION AND PREPARATION

FOR ANALYSIS

Specimen Types

The specimen types listed below were verified for use with this assay. Other specimen types and anticoagulants have not been verified with this assay.

| Specimen Types | Anticoagulants |
|----------------------------|-----------------------------|
| Serum | Not Applicable |
| (including serum separator | |
| tubes) | |
| Plasma | Dipotassium EDTA (including |
| | plasma preparation tubes) |
| | Tripotassium EDTA |
| | Lithium heparin (including |
| | plasma separator tubes) |
| | Sodium citrate |
| | Sodium heparin |
| | ACD-A |
| | ACD-B |
| | CP2D |
| | CPD |
| | CPDA-1 |

• Liquid anticoagulants may have a dilution effect resulting in lower S/CO values for individual specimens.

- Performance has not been established for the use of plasmapheresis specimens.
- Performance has not been established for the use of umbilical cord blood or bodily fluids such as urine, saliva, semen, amniotic fluid, cerebrospinal fluid, or pleural fluid.
- Performance has been established for the use of cadaveric serum specimens (including specimens collected post-mortem, non-heart-beating) that have been collected up to 24 hours after death.⁴⁰ Follow general standards and/or regulations for collection, storage and handling.
- Performance has not been established for the use of cadaveric plasma specimens.
- Testing of cadaveric serum specimens from patients with plasma dilution due to transfusions of > 2000 mL of blood or colloids within 48 hours, or > 2000 mL of crystalloids within 1 hour (or any combination thereof) prior to collection of the specimens has not been verified.
- The system does not provide the capability to verify specimen types. It is the responsibility of the operator to verify that the correct specimen types are used with the assay.

Specimen Conditions

- Do not use:
 - heat-inactivated specimens
 - pooled specimens
 - grossly hemolyzed specimens
 - specimens with obvious microbial contamination
 - specimens with fungal growth
- For accurate results, serum and plasma specimens should be free of fibrin, red blood cells, and other particulate matter.
- To prevent cross contamination, use of disposable pipettes or pipette tips is recommended.



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Preparation for Analysis

Failure to follow the specified centrifugation procedure may give erroneous or inconsistent test results.

- Clear, nonhemolyzed specimens should be used when possible. Specimens containing visible particulate matter may give erroneous or inconsistent test results.
- Specimens should be free of bubbles. Remove bubbles with an applicator stick before analysis. Use a new applicator stick for each specimen to prevent cross contamination.
- Prior to centrifugation, previously frozen specimens must be mixed gently and thoroughly after thawing.
- All specimens must be centrifuged between 30 000 75 000 g-minutes.
- All specimens must be tested or retested within 48 hours of initial centrifugation. After 48 hours, these specimens need to be recentrifuged between 30 000 - 75 000 g-minutes.

The acceptable time and force ranges that meet this criterion are listed in the table below.

| | Centrifugation Time | | |
|---|---------------------|-------------|-----------------|
| | (Minutes) | RCF (x g) | g-Minutes |
| | 10 | 3000 | 30 000 |
| | 15 | 2000 - 3000 | 30 000 - 45 000 |
| | 20 | 1500 - 3000 | 30 000 - 60 000 |
| _ | 25 | 1300 - 3000 | 32 500 - 75 000 |

Convert rpm to RCF as follows: RCF = $1.12 \times r_{max} (rpm/1000)^2$ Convert RCF to rpm as follows:

| $rpm = 1000 \text{ x } \sqrt{\frac{RCF}{1.12 \text{ x } r_{max}}}$ | | | | |
|--|---|--|--|--|
| RCF- | The relative centrifugal force generated during centrifugation. | | | |
| rpm - | The revolutions per minute of the rotor on which the specimens are being spun (usually the digital readout on the centrifuge will indicate the rpm). | | | |
| Centrifugation | The time should be measured from the time the | | | |
| Time - | rotor reaches the required RCF or rpm to the time it begins decelerating. | | | |
| r _{max} - | Radius of the rotor in millimeters. The radius measured is dependent on whether the rotor is a fixed angle rotor or a swinging bucket rotor. This value is typically provided with the rotor by the manufacturer. For the fixed angle rotor, r_{max} is the measure of the distance from the rotor axis (center) to the bottom of the specimen tube in the rotor or rotor adapter. For the swinging bucket rotor, r_{max} is the measure of the distance from the rotor dapter. For the swinging bucket rotor, r_{max} is the measure of the distance from the rotor axis (center) to the bottom of the specimen tube in the rotor axis (center) to the bottom of the specimen tube in the rotor adapter or bucket at full extension. | | | |
| g-minutes - | NOTE: If custom tube adapters (i.e., adapters not defined by the centrifuge manufacturer) are used, then the radius (r_{max}) should be manually measured in millimeters and the RCF calculated. The unit of measure for the product of RCF (× g) and centrifugation time (minutes). | | | |

Specimen Storage

| | | Maximum | |
|--------------|--------------|----------|----------------------------|
| Specimen | | Storage | |
| Туре | Temperature | Time | Special Instructions |
| Living Donor | Room | 7 days | Specimens may be stored on |
| Serum/ | temperature | | or off the clot, red blood |
| Plasma | (15 to 30°C) | | cells, or separator gel. |
| | 2 to 8°C | 14 days | Specimens may be stored on |
| | | | or off the clot, red blood |
| | | | cells, or separator gel. |
| | -20°C or | 3 months | Remove serum or plasma |
| | colder | | from the clot, red blood |
| | | | cells, or separator gel. |

- Living donor specimens stored at -20°C or colder for greater than 3 months may be used for informational purposes. (e.g., lookback testing, discordant sample testing, clinical and validation testing).
- Storage at a combination of 15 to 30°C and 2 to 8°C may not exceed 14 days (inclusive of shipping time) and cannot exceed the maximum durations listed in the table above.
- Performance has not been established for living donor specimens • that have undergone more than 6 freeze/thaw cycles.

| | | Maximum | |
|-----------------------------------|--|----------------|---|
| Specimen | . | Storage | Constal Instantion |
| Type Cadaveric Serum | Temperature Room temperature (15 to 30°C) | Time 3 days | Special Instructions If specimens are not processed directly after initial centrifugation, it is recommended to remove the supernatant from the clot, red blood cells, or separator gel until further processing. |
| | 2 to 8°C | 14 days | If specimens are not processed directly after initial centrifugation, it is recommended to remove the supernatant from the clot, red blood cells, or separator gel until further processing. |
| | -20°C or Colder | 3 months | If specimens are not processed directly after initial centrifugation, it is recommended to remove the supernatant from the clot, red blood cells, or separator gel until further processing. |

- Performance has not been established using cadaveric specimens stored at -20°C or colder for greater than 3 months.
- Storage at a combination of 15 to 30°C and 2 to 8°C may not exceed 14 days (inclusive of shipping time) and cannot exceed the maximum durations listed in the table above.
- Performance has not been established for cadaveric specimens that have undergone more than 6 freeze/thaw cycles.

Specimen Shipping

Package and label specimens in compliance with applicable state, federal, and international regulations covering the transport of clinical specimens and infectious substances.



■ PROCEDURE

Materials Provided

06P07 Alinity s HTLV I/II Reagent Kit

Materials Required but not Provided

- Alinity s HTLV I/II Assay File
- 06P0703 Alinity s HTLV I/II Calibrator Kit
- 06P0720 Alinity s HTLV I/II Assay Control Kit
- 06P0724 Alinity s HTLV I/II Release Control Kit
- Alinity Trigger Solution
- Alinity Pre-Trigger Solution
- Alinity s Concentrated Wash Buffer

For information on materials required for operation of the system, refer to the Alinity s System Operations Manual, Section 1.

For information on materials required for maintenance procedures, refer to the Alinity s System Operations Manual, Section 9.

Assay Procedure

For a detailed description of how to run an assay, refer to the Alinity s System Operations Manual, Section 5.

- Primary tubes may be on board the system for up to 10 hours.
- If using primary or aliquot tubes, refer to the Alinity s System Operations Manual, Section 4 to ensure sufficient specimen is present.
- To minimize the effects of evaporation, verify adequate sample cup volume is present prior to running the test.
- Maximum number of replicates sampled from the same sample cup: 10
 - \leq 3 hours on the reagent and sample manager:
 - Sample volume for first test: 300 µL
 - Sample volume for each additional test from same sample cup: 100 μL
 - > 3 hours on the reagent and sample manager:
 - Replace with a fresh aliquot of sample.
- Refer to the Alinity s HTLV I/II Calibrator Kit, Assay Control Kit, and/or Release Control Kit package inserts for preparation and usage.
- For general operating procedures, refer to the Alinity s System Operations Manual, Section 5.
- For optimal performance, it is important to perform routine maintenance as described in the Alinity s System Operations Manual, Section 9. Perform maintenance more frequently when required by laboratory procedures.

Calibration

For instructions on performing a calibration, refer to the Alinity s System Operations Manual, Section 5.

Three replicates of Alinity s HTLV I/II Calibrator 1 are automatically tested by the system. The calibrator must be priority loaded.

Each assay control must be tested to evaluate the assay calibration. Once a calibration is accepted and stored, it may be used for 14 days. During this time, all subsequent samples may be tested without further calibration unless:

- A reagent kit with a new lot number is used.
- Daily quality control results are outside of quality control limits used to monitor and control system performance.

This assay may require recalibration after maintenance to critical parts or subsystems or after service procedures have been performed.

Quality Control Procedures

Assay Controls

The Alinity s HTLV I/II Assay Controls must be tested once every 24 hours when the system is being used.

Assay control values must be within the ranges specified in the Alinity s HTLV I/II Assay Control Kit package insert. When the assay control values are within range, sample results are generated, and a valid release control result is required to release test results. If an assay control value is not within range, sample results are not generated for in-process or scheduled samples. For troubleshooting information, refer to the Alinity s System Operations Manual, Section 10.

Release Controls

The Alinity s HTLV I/II Release Control must be tested in order to release test results.

The release control is tested at user-defined intervals. For configuring the release control, refer to the Alinity s System Operations Manual, Section 2. For manually ordering the release control, refer to the Alinity s System Operations Manual, Section 5.

The release control must meet specifications defined in the Alinity s HTLV I/II Release Control Kit package insert in order to validate the system functionality and release test results. If the release control does not meet specifications, refer to the Alinity s System Operations Manual, Section 10, for additional information.

Other Controls

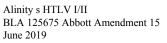
Additional controls may be tested at operator's discretion in accordance with local, state, and/or federal regulations or accreditation requirements and your laboratory's quality control policy. For additional information on configuring customer controls, refer to the Alinity s System Operations Manual, Section 2.

Invalidate controls: Additional controls may be tested anywhere within a run as an invalidate control. Specifications may be assigned to invalidating controls. If an invalidate control fails to meet assigned specifications, no sample results are calculated or provided by the system. When an invalidate control meets assigned specifications, sample processing continues, and a valid release control result is required to release test results.

Non-validating controls: Additional controls may be tested anywhere within a run as a non-validating control. Specifications may be assigned to non-validating controls. A valid release control result is required to release test results. If the user-assigned specifications for the non-validating control(s) are not met and the release control specifications are met, there will be no effect on sample processing. In this case, reactive sample results must not be considered invalid.

Quality Control Guidance

Refer to "Basic QC Practices" by James O Westgard, Ph.D. for guidance on laboratory quality control practices. $^{\rm 41}$





RESULTS

Calculation

The Alinity s System calculates results for the Alinity s HTLV I/II assay using the ratio of the sample RLU to the cutoff RLU (S/CO) for each specimen and control.

Cutoff RLU = Calibrator 1 Mean RLU x 0.40

The cutoff RLU is stored for each reagent lot calibration.

S/CO = Sample RLU/Cutoff RLU

Interpretation of Results

The cutoff is 1.00 S/CO.

Initial Results

| Initial Result (S/CO) | Interpretation | Retest Procedure |
|-----------------------|----------------|-------------------------|
| < 1.00 | Nonreactive | No retest required. |
| | | Specimen considered |
| | | negative for antibodies |
| | | to HTLV I and HTLV II. |
| <u>≥</u> 1.00 | Reactive | Retest in duplicate. |

Final Interpretation

| Retest Result (S/CO) | Final Results | Final Interpretation |
|----------------------|---------------|--|
| Both results < 1.00 | Nonreactive | Specimen considered negative for antibodies |
| | | to HTLV I and HTLV II. |
| One or both results | Repeatedly | Specimen should be |
| <u>></u> 1.00 | Reactive | further tested by |
| | | supplemental methods |

Supplemental methods may include other HTLV I/HTLV II specific immunoassays and/or immunoblot assays per FDA regulations.

Customers outside the US must follow their country's government recommendations and regulations for specimens found to be repeatedly reactive.

SPECIFIC PERFORMANCE CHARACTERISTICS

Representative performance data are provided in this section. Results obtained in individual laboratories may vary.

Reproducibility

A study was performed based on guidance from CLSI EP15-A2.⁴² Testing was conducted using 3 lots of the Alinity s HTLV I/II Reagent Kit, Calibrator Kit, Assay Control Kit, and Release Control Kit. Panel members and controls were tested twice a day for 5 days in replicates of 4 at 3 sites.

Flags

Some results may contain information in the Flags field. For a description of the flags that may appear in this field, refer to the

· Potential interference has not been evaluated for substances other

CHARACTERISTICS - Interference section of this package insert.False reactive results can be expected with any test kit. Falsely

elevated results may be observed due to non-specific interactions

(refer to the SPECIFIC PERFORMANCE CHARACTERISTICS section of

antibodies to HTLV I/HTLV II is strong, it is recognized that presently available methods for HTLV I/HTLV II antibody detection are not sensitive enough to detect all potentially infectious units of blood or possible cases of HTLV I/HTLV II infection. A nonreactive test result

Alinity s System Operations Manual, Section 5.

this package insert).

does not exclude infection.

■ LIMITATIONS OF THE PROCEDURE

than those described in the SPECIFIC PERFORMANCE

• Although the association of infectivity and the presence of

Refer to the **SPECIMEN COLLECTION AND PREPARATION FOR ANALYSIS** section of this package insert for specimen limitations.

| | | | | | | | | | Wit | hin- | | | | | | |
|------------------------------|------|------|-------|--------|-------|--------|-------|--------|-------|--------------------|-------|---------|-------|--------|--------|------------------------|
| | | | With | in-Run | Betwe | en-Run | Betwe | en-Day | Labor | atory ^a | Betwe | en-Site | Betwe | en-Lot | Reprod | ucibility ^b |
| | | Mean | | | | | | | | | | | | | | |
| Sample | N | s/co | SD | %CV | SD | %CV | SD | %CV | SD | %CV | SD | %CV | SD | %CV | SD | %CV |
| Low HTLV I Antibody | 360 | 1.71 | 0.057 | 3.3 | 0.011 | 0.7 | 0.000 | 0.0 | 0.058 | 3.4 | 0.036 | 2.1 | 0.066 | 3.9 | 0.097 | 5.7 |
| High HTLV I Antibody | 359° | 8.68 | 0.288 | 3.3 | 0.000 | 0.0 | 0.058 | 0.7 | 0.294 | 3.4 | 0.037 | 0.4 | 0.496 | 5.7 | 0.582 | 6.7 |
| Low HTLV II Antibody | 360 | 1.67 | 0.058 | 3.4 | 0.015 | 0.9 | 0.000 | 0.0 | 0.059 | 3.6 | 0.006 | 0.4 | 0.134 | 8.0 | 0.147 | 8.8 |
| High HTLV II Antibody | 360 | 8.39 | 0.302 | 3.6 | 0.070 | 0.8 | 0.036 | 0.4 | 0.312 | 3.7 | 0.068 | 0.8 | 0.827 | 9.9 | 0.893 | 10.6 |
| Positive Control 1 (HTLV I) | 360 | 2.45 | 0.081 | 3.3 | 0.000 | 0.0 | 0.018 | 0.7 | 0.083 | 3.4 | 0.000 | 0.0 | 0.181 | 7.4 | 0.200 | 8.2 |
| Positive Control 2 (HTLV II) | 360 | 2.85 | 0.100 | 3.5 | 0.000 | 0.0 | 0.006 | 0.2 | 0.100 | 3.5 | 0.000 | 0.0 | 0.183 | 6.4 | 0.212 | 7.4 |
| Negative Control | 360 | 0.18 | 0.021 | NA | 0.000 | NA | 0.007 | NA | 0.022 | NA | 0.007 | NA | 0.024 | NA | 0.034 | NA |

%CV = Coefficient of Variation expressed as a percentage; N = Number of Replicates; NA = Not Applicable: %CVs are not meaningful when S/CO approaches zero; SD = Standard Deviation

^a Includes within-run, between-run, and between-day variability

^b Includes within-run, between-run, between-day, between-site, between-lot and the site-lot interaction variability.

^c One replicate was missing due to pressure monitoring error.

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Specificity

A total of 9365 fresh serum specimens and 6512 fresh plasma specimens from volunteer whole blood donors were collected at 3 distinct blood centers. The initial and repeat reactive rates for the serum specimens were 0.03% (3/9365) and 0.03% (3/9365), respectively. The initial and repeat reactive rates for the plasma specimens were 0.03% (2/6512) and 0.03% (2/6512), respectively. Repeatedly reactive specimens were further tested using a supplemental HTLV immunoblot. Based on supplemental test results, 1 specimen was positive, 2 specimens were negative, and 2 specimens were indeterminate.

Specificity based on assumed zero prevalence of antibody to HTLV I/HTLV II in whole blood donors was estimated in this study to be 99.99% (15 864/15 866) with a 95% confidence interval of 99.95% to 100.00%.

| Specimen Category | Number Tested | IR (% of Total) (95% Cl) | RR (% of Total) (95% Cl) | Number Positive by Supplemental Testing (% of RR) | Specificity (%) ^a (95% Cl) |
|------------------------------------|------------------|--------------------------------|--------------------------------|---|--|
| Volunteer Blood Donors – Serum | 9365 | 3 (0.03) (0.01 – 0.09) | 3 (0.03) (0.01 – 0.09) | 1 (33.33) | 99.99 (9360/9361) (99.94 – 100.00) |
| Volunteer Blood Donors – Plasma | 6512 | 2 (0.03) (0.00 - 0.11) | 2 (0.03) (0.00 – 0.11) | 0 (0.00) | 99.98 (6504/6505) (99.91 – 100.00) |
| Total Donors | 15 877 | 5 (0.03) (0.01 – 0.07) | 5 (0.03) (0.01 – 0.07) | 1 (20.00) | 99.99 (15 864/15 866) (99.95 – 100.00) |

IR = Initially Reactive, RR = Repeatedly Reactive, CI = Confidence Interval ^a Specimens determined to be positive (n = 1) or indeterminate (n = 2) by supplemental testing were excluded from the specificity calculations. Additionally, there were 8 specimens that were Alinity s HTLV I/II nonreactive and indeterminate by supplemental testing that were excluded from the specificity calculations.

For total donors, IR rate not reactive on retest was estimated to be 0.00% (0/15 872) with a 95% confidence interval of 0.00% to 0.02%. IR Rate Not Reactive on Retest = $100\% \times (Number \text{ of } IR - Number \text{ of } RR) / (Number Tested - Number of RR)$

Sensitivity

A total of 1717 specimens from the categories shown in the table below were tested using the Alinity s HTLV I/II assay at 3 clinical sites. Repeatedly reactive specimens from individuals with HTLV I/II associated disease, individuals at increased risk of HTLV I/II infection, and individuals from HTLV I/II endemic areas were tested using a supplemental HTLV immunoblot. Sensitivity was estimated to be 100.00% (706/706) with a 95% confidence interval of 99.48% to 100.00% for preselected positive specimens and HTLV I/II associated disease.

| Specimen Category | Number Tested | Number Positive | Number RR (% of Total) | Number RR Positive by Supplemental Testing (% of RR) | Sensitivity (%) (95% Cl) |
|---|------------------|--------------------|---------------------------|--|---|
| Preselected Anti-HTLV I Positive ^a | 461 | 461 | 461 (100.00) | 461 (100.00) | 100.00 (461/461) (99.20 – 100.00 |
| Preselected Anti-HTLV II Positive ^a | 141 | 141 | 141 (100.00) | 141 (100.00) | 100.00 (141/141) (97.42 - 100.00) |
| Preselected Anti-HTLV I/II Positive Undifferentiated ^a | 4 | 4 | 4 (100.00) | 4 (100.00) | 100.00 (4/4) (NA) ^g |
| Individuals with HTLV I/II Associated Disease ^b | 100 | 100 | 100 (100.00) | 100 (100.00) | 100.00 (100/100) (96.38 - 100.00) |
| Subtotal | 706 | 706 | 706 (100.00) | 706 (100.00) | 100.00 (706/706) (99.48 – 100.00 |
| Individuals at Increased Risk of HTLV I/II Infection ^c | 502 | 11 ^d | 11 (2.19) | 11 (100.00) | NA ^g |
| Individuals from HTLV I/II Endemic Areas ^e | 509 | 10 ^f | 10 (1.96) | 10 (100.00) | NA ^g |
| Total | 1717 | 727 | 727 (42.34) | 727 (100.00) | 100.00 (727/727) (99.49 – 100.00 |

NA = Not Applicable; RR = Repeatedly Reactive; CI = Confidence Interval

- ^a Preselected anti-HTLV I/II positive specimens were determined by detecting the presence of antibodies to both gag (p24 and/or p19) and env (native gp46 and/or gp67/68) antigens using research use only Western blot and/or RIPA. HTLV I and HTLV II type differentiation was determined using research use only HTLV I and HTLV II IFA.
- ^b Individuals with HTLV I/II associated diseases category included ATL patients (50) and HAM/TSP patients (50).
- ^c The following risk factors were included: intravenous drug users, multiple sex partners, and STD clinic patients.
- ^d All 11 specimens that were positive by supplemental testing were anti-HTLV II positive specimens.
- ^e Individuals from HTLV I/II endemic areas included specimens from the following areas: Congo (100), Haiti (204), and Peru (205).
- ^f The 10 specimens that were positive by supplemental testing included 9 anti-HTLV I positive specimens, and 1 undifferentiated anti-HTLV I/II positive specimen.
- ^g The sensitivity calculation and/or confidence interval are not meaningful due to the small number of specimens.

Other Specimen Conditions or Disease States

A total of 225 specimens from individuals with other specimen conditions or disease states unrelated to HTLV I/II infection were evaluated. Of the 225 specimens, 1 was repeatedly reactive using the Alinity s HTLV I/II assay and a commercially available HTLV I/II assay and was anti-HTLV I/II positive by supplemental testing.

| Specimen Category | Number Tested | IR (% of Total) | RR (% of Total) | Number Positive by Supplemental Testing (% of Repeatedly Reactive) |
|--|------------------|-----------------------|-----------------------|--|
| Other Specimen Conditions or Disease States ^a | 225 | 1 (0.44) | 1 (0.44) | 1 (100.00) ^b |

IR = Initially Reactive; RR = Repeatedly Reactive

^a The specimens included the following: Anti-HIV-1/HIV-2 Positive (10), Anti-HCV Positive (10), HBV Positive (10), Co-infected CMV/EBV/HSV (10), Anti-*T* pallidum Positive (10), Anti-VZV Positive (10), Rheumatoid Factor Positive (10), Anti-ds DNA Positive (10), Pregnant Females (14), Multiparous Females (10), Hyper IgG/IgM (10), Influenza Vaccine Recipients (10), Hemodialysis Patients (10), HAMA Positive (10), *E coli* Infection (10), Heterophilic Antibody Positive (12), Anti-*gonaccoccus* Positive (9), Anti-*C trachomatis* Positive (10), Anti-*T gondii* Positive (10), Anti-nuclear Antibody Positive (10), Fungal (Yeast) Infection (10), and Anti-Rubella Positive (10).

^b One anti-HCV positive specimen was positive by supplemental testing.

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Interference

Potentially Interfering Endogenous Substances

A study was performed based on guidance from CLSI EP07-A2.43

No interference was observed using the Alinity s HTLV I/II assay from potentially interfering substances at the levels shown below.

| Potentially Interfering Substance | Interferent Level |
|-----------------------------------|-------------------|
| Conjugated Bilirubin | ≤ 20 mg/dL |
| Unconjugated Bilirubin | ≤ 20 mg/dL |
| Hemoglobin | ≤ 500 mg/dL |
| Triglycerides | ≤ 3000 mg/dL |
| Total Protein | ≤ 12 g/dL |

In addition, a negative and two positive controls were spiked with biotin to a concentration of 4250 ng/mL. No interference was observed using the Alinity s HTLV I/II assay.

The effect of potentially interfering substances has only been evaluated for those listed in this package insert.

PERFORMANCE CHARACTERISTICS OF CADAVERIC SPECIMEN TESTING

Reproducibility

Twenty-two nonreactive cadaveric donor serum specimens and 23 nonreactive living donor serum specimens were spiked with human plasma reactive for anti-HTLV I or anti-HTLV II to create low-level reactive specimens.

Each specimen was tested once per day for 6 days using each of 3 lots of the Alinity s HTLV I/II Reagent Kit. Total %CV values were determined.

| | | | | Total ^a | |
|--------------|------------------------|-------------------------|--------------|--------------------|-----|
| Analyte | Specimen Category | Number of Replicates | Mean S/CO | SD | %CV |
| Anti-HTLV I | Cadaveric ^b | 396 | 4.03 | 0.214 | 5.3 |
| | Living Donor | 414 | 4.72 | 0.242 | 5.1 |
| Anti-HTLV II | Cadaveric ^b | 396 | 4.10 | 0.297 | 7.3 |
| | Living Donor | 414 | 4.57 | 0.322 | 7.0 |

^a Total variability contains within-specimen, between-lot and lot-specimen interaction variance components.

^b Cadaveric serum specimens were collected up to 23.1 hours after death.

Specificity

Specificity was determined by testing 55 cadaveric serum specimens and 55 living donor serum specimens. Each specimen was tested once using each of 3 lots of the Alinity s HTLV I/II Reagent Kit.

| Specimen Category | Lot | Nonreactive | Repeatedly Reactive | Specificity (%) (95% CI) |
|------------------------|-------|-------------|------------------------|-----------------------------|
| Cadaveric ^a | Lot 1 | 55 | 0 | 100.00 (93.51 – 100.00) |
| | Lot 2 | 55 | 0 | 100.00 (93.51 – 100.00) |
| | Lot 3 | 55 | 0 | 100.00 (93.51 – 100.00) |
| Living Donor | Lot 1 | 55 | 0 | 100.00 (93.51 – 100.00) |
| | Lot 2 | 55 | 0 | 100.00 (93.51 – 100.00) |
| | Lot 3 | 55 | 0 | 100.00 (93.51 – 100.00) |

CI = Confidence Interval

^a Cadaveric serum specimens were collected up to 23.7 hours after death.

Analytical Sensitivity

Cadaveric serum specimens and living donor serum specimens were spiked with human plasma reactive for anti-HTLV I or anti-HTLV II to create low-level reactive specimens. Each specimen was tested once, within 24 hours of spiking, using each of 3 lots of the Alinity s HTLV I/II Reagent Kit. All specimens were reactive on all 3 reagent lots.

| | Specimen | | Number of | Mean | Sensitivity (%) |
|--------------|------------|-------|-----------|------|------------------|
| Analyte | Category | Lot | Specimens | s/co | (95% CI) |
| Anti-HTLV I | Cadaverica | Lot 1 | 52 | 4.50 | 100.00 |
| Anti-IIIEVI | Cauaveric | LULI | 52 | 4.50 | (93.15 – 100.00) |
| | | Lot 2 | 52 | 4.70 | 100.00 |
| | | LOUZ | 52 | 4.70 | (93.15 – 100.00) |
| | | Lot 3 | 52 | 4.69 | 100.00 |
| | | LOUS | 52 | 4.09 | (93.15 - 100.00) |
| | Living | Lot 1 | 53 | 5.12 | 100.0 |
| | Donor | LOUI | 22 | 5.12 | (93.28 - 100.00) |
| | | Lot 2 | 53 | 5.37 | 100.0 |
| | | LOUZ | 55 | 5.57 | (93.28 – 100.00) |
| | | Lot 3 | 53 | 5.12 | 100.0 |
| | | LOUS | 55 | 5.12 | (93.28 – 100.00) |
| Anti-HTLV II | Cadaverica | Lot 1 | 53 | 4.76 | 100.0 |
| AIIU-HILV II | Cauavence | LOUI | 55 | 4.70 | (93.28 – 100.00) |
| | | Lot 2 | 53 | 4.68 | 100.0 |
| | | LOUZ | 55 | 4.00 | (93.28 – 100.00) |
| | | Lot 3 | 53 | 4.75 | 100.0 |
| | | LULS | 55 | 4.75 | (93.28 – 100.00) |
| | Living | Lot 1 | 53 | 5.05 | 100.0 |
| | Donor | LOUI | 53 | 5.05 | (93.28 - 100.00) |
| | | Lot 2 | 53 | 4.93 | 100.0 |
| | | LOI Z | 53 | 4.93 | (93.28 - 100.00) |
| | | Lot 2 | E2 | 4 74 | 100.0 |
| | | Lot 3 | 53 | 4.74 | (93.28 - 100.00) |

CI = Confidence Interval

^a Cadaveric serum specimens were collected up to 22.6 hours after death.

Cadaveric Specimen Storage

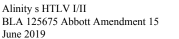
Cadaveric specimen storage was determined by testing a minimum of 12 low-level reactive specimens, prepared by spiking nonreactive cadaveric serum specimens to a target S/CO value near the cutoff with human plasma reactive for anti-HTLV I and anti-HTLV II, and a minimum of 12 nonreactive cadaveric serum specimens. Each specimen was tested at Day 0, and then subjected to either 2 to 8°C storage for 14 days, room temperature (15 to 30°C) storage for 3 days, -20°C or colder storage for 3 months, or 6 freeze/thaw cycles. Nonreactive specimens were evaluated by calculating the differences between the mean S/CO of Day 0 and the mean S/CO of each storage condition and related timepoint. Reactive specimens were evaluated by calculating the mean S/CO of each storage condition and related timepoint. There were no changes to the interpretation; the data demonstrate that cadaveric serum specimens can be stored at the following conditions when tested using the Alinity s HTLV I/II assay.

| | Storage Condition | Timepoint | Nonreactive Specimens Upper Limit of 2-sided 95% Cl of Differences | Anti-HTLV I Reactive Specimens Lower Limit of 2-sided 95% CI of % Differences | Anti-HTLV II Reactive Specimens Lower Limit of 2-sided 95% CI of % Differences | |
|---|--------------------------------------|-----------|--|--|---|--|
| | Room Temperature (15 to 30°C)ª | 3 days | 0.02 S/CO | -10.0% | -2.4% | |
| - | 2 to 8°C ^a | 14 days | 0.01 S/CO | -2.7% | -0.7% | |
| | –20°C or colder $^{\rm b}$ | 3 months | 0.00 S/CO | -0.9% | 4.6% | |
| | Freeze/Thaw ^a | 6 cycles | 0.02 S/CO | -10.8% | -10.0% | |

CI = confidence interva

^a Cadaveric serum specimens were collected up to 41.8 hours after death.

^b Cadaveric serum specimens were collected up to 21.4 hours after death.







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Alinity s HTLV I/II BLA 125675 Abbott Amendment 15 June 2019 Note for number formatting:

- A space is used as thousands separator (example: 10 000 specimens).
- A period is used to separate the integer part from the fractional part of a number written in decimal form (example: 3.12%).

| Rey to Symbols | |
|---------------------------|---|
| <u>í</u> i | Consult instructions for use |
| | Manufacturer |
| T | Sufficient for |
| X | Temperature limitation |
| | Use by/Expiration date |
| ASSAY DILUENT | Assay Diluent |
| CONJUGATE | Conjugate |
| CONTAINS: AZIDE | Contains Sodium Azide. Contact with acids liberates very toxic gas. |
| DISTRIBUTED IN THE USA BY | Distributed in the USA by |
| INFORMATION FOR USA ONLY | Information needed for United States of America Only |
| שעו | <i>In Vitro</i> Diagnostic Medical Device |
| LOT | Lot Number |
| MICROPARTICLES | Microparticles |
| PRODUCT OF GERMANY | Product of Germany |
| REF | List Number |
| SN | Serial Number |

Key to Symbols

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Revised June 2019

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HTLV I/II Calibrator Kit

Revised April 2019.

HTLV I/II REF 06P0703 G93465R02 S6P0S0

Instructions must be carefully followed. Reliability of assay results cannot be guaranteed if there are any deviations from these instructions.

NAME

Alinity s HTLV I/II Calibrator Kit

INTENDED USE

The Alinity s HTLV I/II Calibrator is used to calibrate the Alinity s System when it is used for the qualitative detection of antibodies to human T-lymphotropic virus Type I and/or human T-lymphotropic virus Type II (anti-HTLV I/HTLV II) in human serum and plasma.

REAGENTS

Kit Contents

CAL 1 2 bottles of HTLV I/II Calibrator 1 contain recalcified, inactivated, human plasma reactive for anti-HTLV I. Preservatives: ProClin 950 and sodium azide.

| | | | Target Value |
|------------|------------|------------------|--------------|
| Calibrator | Quantity | Color | (S/CO) |
| CAL 1 | 2 x 1.6 mL | Red ^a | 2.50 |

S/CO = Sample to Cutoff

a Dye: Red D&C No. 33

Standardization

The HTLV I/II Calibrator 1 is standardized to an Abbott internal reference standard.

Warnings and Precautions

For In Vitro Diagnostic Use

CAUTION: This product contains human-sourced and/or potentially infectious components. Refer to the **REAGENTS** section of this package insert. No known test method can offer complete assurance that products derived from human sources or inactivated microorganisms will not transmit infection. Therefore, all humansourced materials should be considered potentially infectious. It is recommended that these reagents and human specimens be handled in accordance with the OSHA Standard on Bloodborne Pathogens. Biosafety Level 2 or other appropriate biosafety practices should be used for materials that contain or are suspected of containing infectious agents.¹⁻⁴

The human plasma used in the calibrator is reactive for anti-HTLV I and nonreactive for HBsAg, anti-HIV-1/HIV-2, anti-HCV, and HIV-1 RNA or HIV-1 Ag.

| The following warn | ings and precautions apply to: CAL 1 | |
|------------------------------------|---|--|
| $\langle \mathbf{\hat{v}} \rangle$ | | |
| WARNING: | Contains methylisothiazolone and sodium azide | |
| H317 | May cause an allergic skin reaction. | |
| EUH032 | , | |
| Prevention | | |
| P261 | Avoid breathing mist / vapors / spray. | |
| P272 | Contaminated work clothing should not be | |
| | allowed out of the workplace. | |
| P280 | Wear protective gloves / protective | |
| | clothing / eye protection. | |
| Response | | |
| P302+P352 | IF ON SKIN: Wash with plenty of water. | |
| P333+P313 | If skin irritation or rash occurs: Get | |
| | medical advice / attention. | |
| P362+P364 | Take off contaminated clothing and wash | |
| | it before reuse. | |
| Disposal | | |
| P501 | Dispose of contents / container in | |
| | accordance with local regulations. | |

Safety Data Sheets are available at www.transfusion.abbott or contact your local representative.

For a detailed discussion of safety precautions during system operation, refer to the Alinity s System Operations Manual, Section 8.

Reagent Handling

- Do not pool the calibrators.
- Do not freeze.
- For a detailed discussion of handling calibrators during system operation, refer to the Alinity s System Operations Manual, Section 7.

Reagent Storage

| | Storage Temperature | Maximum Storage Time | Additional Storage Instructions |
|----------|------------------------|--------------------------|---|
| Unopened | 2 to 8°C | Until expiration date | Store in an upright position. May be used immediately after removal from 2 to 8°C storage. |
| Onboard | System Temperature | 5 hours | |
| Opened | 2 to 8°C | 24 hours | Store tightly capped. Store in an upright position. Do not invert calibrators prior to loading them on the system. |



Indications of Deterioration

- Instability or deterioration should be suspected if there are precipitates, visible signs of leakage, turbidity, if calibration does not meet the appropriate package insert and/or Alinity s System Operations Manual criteria, or if controls do not meet the appropriate criteria.
- For troubleshooting information, refer to the Alinity s System Operations Manual, Section 10.

PROCEDURE

Materials Provided

• 06P0703 Alinity s HTLV I/II Calibrator Kit

Instructions for Use

- · Calibrator bottles are one-time use.
- For information on ordering calibrations and loading calibrators, refer to the Alinity s System Operations Manual, Section 5.

QUALITY CONTROL PROCEDURES

- Three replicates of Calibrator 1 are automatically tested by the system. The calibrator must be priority loaded.
- Once a calibration is accepted and stored, it may be used for 14 days. During this time, all subsequent samples may be tested without further calibration unless:
 - A reagent kit with a new lot number is used.
 - Daily quality control results are outside of quality control limits used to monitor and control system performance.
- This assay may require recalibration after maintenance to critical parts or subsystems or after service procedures have been performed.
- Refer to the Alinity s HTLV I/II Reagent Kit package insert and the Alinity s System Operations Manual for additional information.
- A single sample of each assay control must be tested to evaluate the calibration. For information on ordering controls, refer to the Alinity s System Operations Manual, Section 5.
 - Ensure that assay control values are within the ranges specified in the **RESULTS** section of the Alinity s HTLV I/II Assay Control Kit package insert.

BIBLIOGRAPHY

- US Department of Labor, Occupational Safety and Health Administration, 29 CFR Part 1910.1030, Bloodborne pathogens.
- US Department of Health and Human Services. *Biosafety in* Microbiological and Biomedical Laboratories. 5th ed. Washington, DC: US Government Printing Office; December 2009.
- World Health Organization. Laboratory Biosafety Manual. 3rd ed. Geneva: World Health Organization; 2004.
- Clinical and Laboratory Standards Institute (CLSI). Protection of Laboratory Workers from Occupationally Acquired Infections; Approved Guideline—Fourth Edition. CLSI Document M29-A4. Wayne, PA: CLSI; 2014.

Key to Symbols

| \bigcirc | Caution |
|---------------------------|--|
| ī | Consult instructions for use |
| | Manufacturer |
| X | Temperature limitation |
| | Use by/Expiration date |
| CAL 1 | Calibrator 1 |
| CN | Control Number |
| CONTAINS: AZIDE | Contains Sodium Azide. Contact with acids liberates very toxic gas. |
| DISTRIBUTED IN THE USA BY | Distributed in the USA by |
| INFORMATION FOR USA ONLY | Information needed for United States of America only <i>In vitro</i> Diagnostic Medical Device |
| LOT | Lot Number |
| PRODUCT OF GERMANY | Product of Germany |
| REF | List Number |
| SN | Serial Number |

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HTLV I/II Assay Control Kit

HTLV I/II REF 06P0720 G93468R02 C6P0S0

Instructions must be carefully followed. Reliability of assay results cannot be guaranteed if there are any deviations from these instructions.

NAME

Alinity s HTLV I/II Assay Control Kit

INTENDED USE

The Alinity s HTLV I/II Assay Controls are used to verify the calibration of the Alinity s System when it is used for the qualitative detection of antibodies to human T-lymphotropic virus Type I and/or human T-lymphotropic virus Type II (anti-HTLV I/HTLV II) in human serum and plasma.

REAGENTS

Kit Contents

 \sum 1 x 30 tests

CONTROL - 1 bottle of HTLV I/II Negative Control contains recalcified, human plasma. Preservatives: ProClin 950 and sodium azide.

CONTROL 1 1 bottle of HTLV I/II Positive Control 1 contains recalcified, inactivated, human plasma reactive for anti-HTLV I. Preservatives: ProClin 950 and sodium azide.

CONTROL + 2 1 bottle of HTLV I/II Positive Control 2 contains recalcified, inactivated, human plasma reactive for anti-HTLV II. Preservatives: ProClin 950 and sodium azide.

| Control | Quantity | Color | Minimum Activity (S/CO) |
|-------------|------------|-------------------|----------------------------|
| CONTROL - | 1 x 5.2 mL | None | Not applicable |
| CONTROL + 1 | 1 x 5.2 mL | Blue ^a | 1.18 |
| CONTROL + 2 | 1 x 5.2 mL | None | 1.35 |

S/CO = Sample to Cutoff

^a Dye: Acid Blue No. 9

Warnings and Precautions

IVD

For In Vitro Diagnostic Use

CAUTION: This product contains human-sourced and/or potentially infectious components. Refer to the **REAGENTS** section of this package insert. No known test method can offer complete assurance that products derived from human sources or inactivated microorganisms will not transmit infection. Therefore, all humansourced materials should be considered potentially infectious. It is recommended that these reagents and human specimens be handled in accordance with the OSHA Standard on Bloodborne Pathogens. Biosafety Level 2 or other appropriate biosafety practices should be used for materials that contain or are suspected of containing infectious agents.¹⁻⁴ The human plasma used in the negative control is nonreactive for anti-HTLV I/II, HBsAg, anti-HIV-1/HIV-2, anti-HCV, and HIV-1 RNA or HIV-1 Ag.

The human plasma used in the positive control 1 is reactive for anti-HTLV I and nonreactive for HBsAg, anti-HIV-1/HIV-2, anti-HCV, and HIV-1 RNA or HIV-1 Ag.

The human plasma used in the positive control 2 is reactive for anti-HTLV II and nonreactive for HBsAg, anti-HIV-1/HIV-2, anti-HCV, and HIV-1 RNA or HIV-1 Ag.

| • | ings and precautions apply to: |
|-------------------|--|
| CONTROL - , CONT | ROL + 1, and CONTROL + 2 |
| $\langle \rangle$ | |
| WARNING: | Contains methylisothiazolone and sodium azide. |
| H317 | May cause an allergic skin reaction. |
| EUH032 | Contact with acids liberates very toxic |
| | gas. |
| Prevention | |
| P261 | Avoid breathing mist / vapors / spray. |
| P272 | Contaminated work clothing should not be |
| | allowed out of the workplace. |
| P280 | Wear protective gloves / protective |
| | clothing / eye protection. |
| Response | · |
| P302+P352 | IF ON SKIN: Wash with plenty of water. |
| P333+P313 | If skin irritation or rash occurs: Get |
| | medical advice / attention. |
| P362+P364 | Take off contaminated clothing and wash |
| | it before reuse. |
| Disposal | |
| P501 | Dispose of contents / container in |
| | accordance with local regulations. |
| | |

Safety Data Sheets are available at www.transfusion.abbott or contact your local representative.

For a detailed discussion of safety precautions during system operation, refer to the Alinity s System Operations Manual, Section 8.

Reagent Handling

- Do not pool the assay controls.
- Do not freeze.
- For a detailed discussion of handling assay controls during system operation, refer to the Alinity s System Operations Manual, Section 7.



1

Reagent Storage

| | Storage Temperature | Maximum Storage Time | Additional Storage Instructions |
|----------|------------------------|--------------------------|---|
| Unopened | 2 to 8°C | Until expiration date | Store in an upright position. May be used immediately after removal from 2 to 8°C storage. |
| Onboard | System Temperature | 15 hours | |
| Opened | 2 to 8°C | 30 days | Store tightly capped. Return to refrigerated storage after use. Do not invert controls prior to loading on the system. |

Key to Symbols Caution Consult instructions for use Manufacturer Sufficient for Temperature limitation Use by/Expiration date CN Control Number Contains Sodium Azide. Contact CONTAINS: AZIDE with acids liberates very toxic gas. DISTRIBUTED IN THE USA BY INFORMATION FOR USA ONLY Negative Control CONTROL -Positive Control 1 CONTROL + 1

Indications of Deterioration

- Instability or deterioration should be suspected if there are precipitates, visible signs of leakage, turbidity, or if assay controls do not meet the appropriate package insert criteria.
- For troubleshooting information, refer to the Alinity s System Operations Manual, Section 10.

PROCEDURE

Materials Provided

• 06P0720 Alinity s HTLV I/II Assay Control Kit

Instructions for Use

For information on ordering assay controls, refer to the Alinity s System Operations Manual, Section 5.

QUALITY CONTROL PROCEDURES

- The Alinity s HTLV I/II Assay Controls must be tested once every 24 hours when the system is being used.
- Assay control values must be within the ranges specified in the RESULTS section of this package insert. When the assay control values are within range, sample results are generated, and a valid release control result is required to release test results. If an assay control value is not within range, sample results are not generated for in-process or scheduled samples. For troubleshooting information, refer to the Alinity s System Operations Manual, Section 10.
- Refer to the Alinity s HTLV I/II Reagent Kit package insert and the Alinity s System Operations Manual for additional information.

RESULTS

The following table details the acceptable Sample to Cutoff ratio (S/CO) specifications for the Alinity s HTLV I/II Assay Controls.

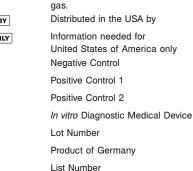
| Assay Control | S/CO Range |
|---------------|-------------|
| CONTROL - | ≤ 0.71 |
| CONTROL + 1 | 1.18 - 6.00 |
| CONTROL + 2 | 1.35 - 6.89 |

BIBLIOGRAPHY

- US Department of Labor, Occupational Safety and Health Administration, 29 CFR Part 1910.1030, Bloodborne pathogens.
- US Department of Health and Human Services. *Biosafety in* Microbiological and Biomedical Laboratories. 5th ed. Washington, DC: US Government Printing Office; December 2009.
- World Health Organization. Laboratory Biosafety Manual. 3rd ed. Geneva: World Health Organization; 2004.
- Clinical and Laboratory Standards Institute (CLSI). Protection of Laboratory Workers from Occupationally Acquired Infections; Approved Guideline—Fourth Edition. CLSI Document M29-A4. Wayne, PA: CLSI; 2014.



2



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CONTROL + 2

PRODUCT OF GERMANY

IVD

LOT

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SN

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Revised April 2019.

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HTLV I/II Release Control Kit

Revised April 2019.

Instructions must be carefully followed. Reliability of assay results cannot be guaranteed if there are any deviations from these instructions.

NAME

Alinity s HTLV I/II Release Control Kit

INTENDED USE

The Alinity s HTLV I/II Release Control is used to validate the Alinity s System functionality and release sample results when it is used for the qualitative detection of antibodies to human T-lymphotropic virus Type I and/or human T-lymphotropic virus Type II (anti-HTLV I/ HTLV II) in human serum and plasma.

REAGENTS

Kit Contents

 Σ 25 x 20 tests

RELEASE CONTROL 25 bottles of HTLV I/II Release Control contain recalcified, inactivated, human plasma reactive for anti-HTLV I. Preservatives: ProClin 950 and sodium azide.

| | | | Minimum Activity |
|-----------------|-------------|-------------------|------------------|
| Control | Quantity | Color | (S/CO) |
| RELEASE CONTROL | 25 x 4.1 mL | Blue ^a | 1.18 |

S/CO = Sample to Cutoff

^a Dye: Acid Blue No. 9

Warnings and Precautions

For In Vitro Diagnostic Use

CAUTION: This product contains human-sourced and/or potentially infectious components. Refer to the **REAGENTS** section of this package insert. No known test method can offer complete assurance that products derived from human sources or inactivated microorganisms will not transmit infection. Therefore, all humansourced materials should be considered potentially infectious. It is recommended that these reagents and human specimens be handled in accordance with the OSHA Standard on Bloodborne Pathogens. Biosafety Level 2 or other appropriate biosafety practices should be used for materials that contain or are suspected of containing infectious agents.¹⁻⁴

The human plasma used in the release control is reactive for anti-HTLV I and nonreactive for HBsAg, anti-HIV-1/HIV-2, anti-HCV, and HIV-1 RNA or HIV-1 Ag.

| The following warn | ings and precautions apply to: RELEASE CONTROL | |
|--------------------|--|--|
| \Diamond | | |
| WARNING: | Contains methylisothiazolone and sodium azide. | |
| H317 | May cause an allergic skin reaction. | |
| EUH032 | Contact with acids liberates very toxic gas. | |
| Prevention | | |
| P261 | Avoid breathing mist / vapors / spray. | |
| P272 | Contaminated work clothing should not be allowed out of the workplace. | |
| P280 | Wear protective gloves / protective clothing / eye protection. | |
| Response | · | |
| P302+P352 | IF ON SKIN: Wash with plenty of water. | |
| P333+P313 | If skin irritation or rash occurs: Get medical advice / attention. | |
| P362+P364 | Take off contaminated clothing and wash it before reuse. | |
| Disposal | | |
| P501 | Dispose of contents / container in accordance with local regulations. | |

Safety Data Sheets are available at www.transfusion.abbott or contact your local representative.

For a detailed discussion of safety precautions during system operation, refer to the Alinity s System Operations Manual, Section 8.

Reagent Handling

- Do not pool the release controls.
- Do not freeze.
- For a detailed discussion of handling the release control during system operation, refer to the Alinity s System Operations Manual, Section 7.

Reagent Storage

| | Storage Temperature | Maximum Storage Time | Additional Storage Instructions |
|----------|------------------------|--------------------------|---|
| Unopened | 2 to 8°C | Until expiration date | Store in an upright position. May be used immediately after removal from 2 to 8°C storage. |
| Onboard | System Temperature | 24 hours | |
| Opened | 2 to 8°C | 14 days | Store tightly capped. Return to refrigerated storage after use. Do not invert controls prior to loading on the system. |



Indications of Deterioration

- Instability or deterioration should be suspected if there are precipitates, visible signs of leakage, turbidity, or if the release control does not meet the appropriate package insert criteria.
- For troubleshooting information, refer to the Alinity s System Operations Manual, Section 10.

PROCEDURE

Materials Provided

06P0724 Alinity s HTLV I/II Release Control Kit

Instructions for Use

For information on ordering release controls, refer to the Alinity s System Operations Manual, Section 5.

QUALITY CONTROL PROCEDURES

- The Alinity s HTLV I/II Release Control must be tested in order to release test results. The release control is tested at user-defined intervals. For configuring the release control, refer to the Alinity s System Operations Manual, Section 2. For manually ordering the release control, refer to the Alinity s System Operations Manual, Section 5.
- The release control must meet the specifications defined in the RESULTS section of this package insert in order to validate the system functionality and release test results. If the release control does not meet specifications, refer to the Alinity s System Operations Manual, Section 10, for additional information.
- Refer to the Alinity s HTLV I/II Reagent Kit package insert and the Alinity s System Operations Manual for additional information.

RESULTS

The acceptable Sample to Cutoff ratio (S/CO) specification for the Alinity s HTLV I/II Release Control is shown below.

RELEASE CONTROL S/CO Range: 1.18 - 6.00

BIBLIOGRAPHY

- US Department of Labor, Occupational Safety and Health Administration, 29 CFR Part 1910.1030, Bloodborne pathogens.
- US Department of Health and Human Services. *Biosafety in* Microbiological and Biomedical Laboratories. 5th ed. Washington, DC: US Government Printing Office; December 2009.
- World Health Organization. Laboratory Biosafety Manual. 3rd ed. Geneva: World Health Organization; 2004.
- Clinical and Laboratory Standards Institute (CLSI). Protection of Laboratory Workers from Occupationally Acquired Infections; Approved Guideline—Fourth Edition. CLSI Document M29-A4. Wayne, PA: CLSI; 2014.

Key to Symbols

| \bigwedge | Caution |
|---------------------------|---|
| ī | Consult instructions for use |
| | Manufacturer |
| Σ | Sufficient for |
| | Temperature limitation |
| | Use by/Expiration date |
| CN | Control Number |
| CONTAINS: AZIDE | Contains Sodium Azide. Contact with acids liberates very toxic gas. |
| DISTRIBUTED IN THE USA BY | Distributed in the USA by |
| INFORMATION FOR USA ONLY | Information needed for United States of America only |
| IVD | In vitro Diagnostic Medical Device |
| LOT | Lot Number |
| PRODUCT OF GERMANY | Product of Germany |
| REF | List Number |
| RELEASE CONTROL | Release Control |
| SN | Serial Number |

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