INTENDED USE
ORTHO HCV Version 3.0 ELISA Test System is an enzyme-linked, immunosorbent assay for the qualitative detection of antibody to hepatitis C virus (anti-HCV) in human serum, plasma, and cadaveric specimens. This product is intended for use as a donor screening test to detect antibodies to hepatitis C virus in plasma and serum samples from individual human donors, including volunteer donors of Whole Blood, blood components, source plasma, and other living donors. It is also intended for use to screen organ donors when specimens are obtained while the donor’s heart is still beating, and in testing blood specimens to screen cadaveric (non-heart-beating) donors. This test is not intended for use on samples of cord blood.

SUMMARY AND EXPLANATION
ORTHO HCV Version 3.0 ELISA Test System is an enzyme-linked immunosorbent assay (ELISA) which utilizes microwells coated with recombinant hepatitis C virus encoded antigens as the solid phase. ELISA technology utilizes the principle that antigens or antibodies which become bound to the solid phase can be detected by complementary antibody or antigen which is labeled with an enzyme capable of acting on a chromogenic substrate. When enzyme substrate is applied, the presence of antigen or antibody can be detected by the development of a colored end product. Immunoassays of this type were first developed in the early 1970s. Since that time, ELISA technology has been extensively used for the detection of antigens and antibodies for a wide range of infectious diseases.

The hepatitis C virus (HCV) is now known to be the causative agent for most, if not all, blood-borne non-A, non-B hepatitis (NANBH). Studies throughout the world indicate that HCV is transmitted through contaminated blood and blood products, through blood transfusions or through other close, personal contacts. Currently, in the United States, greater than 90% of transfusion-associated hepatitis infections are considered to be NANBH infections. Worldwide, other forms of NANBH are recognized.

Three recombinant hepatitis C virus encoded antigens are used in ORTHO HCV Version 3.0 ELISA Test System. The three recombinant antigens, developed by Novartis Vaccines and Diagnostics, Inc., are c22-3, c200 and NS5.

A graphic representation of the putative HCV genome and recombinant proteins appears in Figure 1.

Figure 1
HCV Genome and Recombinant Proteins

HCV recombinant protein c22-3 is encoded by the putative core region of the HCV genome. Amino acid and nucleotide sequence comparisons of flaviviruses and pestiviruses with HCV suggest that c22-3 is derived from a structural region of the genome which encodes the RNA-binding nucleocapsid protein. Nucleocapsid proteins are thought to be involved in forming the viral core structure. Recent studies have indicated that antibodies which develop following infection with HCV are often reactive with c22-3. Moreover, studies performed using the CHIRON™ RIBA™ HCV 2.0 Strip Immunoblot Assay (SIA) for anti-HCV have shown that in many cases antibodies to c22-3 develop sooner following HCV infection than those to c100-3.
HCV recombinant protein c200 is encoded by the putative NS3 and NS4 regions of the HCV genome. Amino acid and nucleotide sequence comparisons of flaviviruses and pestiviruses with HCV suggest that c200 is derived from nonstructural regions of the genome. The c200 recombinant protein contains the c33c protein sequence genetically linked to the c100-3 protein sequence. c33c is also encoded by the putative NS3 portion of the HCV genome. Amino acid and nucleotide sequence comparisons of flaviviruses and pestiviruses with HCV suggest that the NS3 region encodes the viral helicase, an enzyme involved in the unwinding of RNA during replication of the viral genome by RNA-dependent RNA polymerase. Recent studies have indicated that antibodies which develop following infection with HCV are frequently reactive with c33c. Studies performed using the CHIRON™ RIBA™ HCV 2.0 SIA for anti-HCV have shown that antibodies reactive with c33c often develop sooner following HCV infection than do those to c100-3.11

HCV recombinant protein c100-3 is encoded by the putative NS4 region of the HCV genome. Amino acid and nucleotide sequence comparisons of flaviviruses and pestiviruses with HCV suggest that c100-3 is derived from a nonstructural region of the genome. At present, the function of this portion of the HCV genome is unknown. Antibodies which develop following infection with HCV are often reactive with c100-3.3

HCV recombinant protein NS5 is encoded by the putative NS5 region of the HCV genome. Amino acid and nucleotide sequence comparisons of flaviviruses and pestiviruses with HCV suggest that NS5 is derived from a nonstructural region of the genome that encodes the viral polymerase, an enzyme involved in replication of HCV. Recent studies have indicated that a significant proportion of persons infected with HCV develop antibodies to NS5.12,13

The use of HCV recombinant proteins derived from the core, NS3, NS4 and NS5 regions of the HCV genome has shown to be effective in identifying a greater number of diagnosed acute and chronic non-A, non-B hepatitis patients than single antigen (c100-3) assays.12,13 In addition, the use of these additional proteins allows for earlier detection of seroconversion following HCV infection. Although antibody responses to NS5 region-encoded antigens are not as prevalent in response to HCV infection as those to core and NS3 region-encoded antigens, the addition of NS5 to c22-3 and c200 recombinant proteins in ORTHO HCV Version 3.0 ELISA Test System affords antibody detection to a greater number of HCV-encoded epitopes.

The amino acid sequence of the three HCV recombinant proteins is as follows.

<table>
<thead>
<tr>
<th>Recombinant Protein</th>
<th>Polypeptide Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>c22-3</td>
<td>AA # 2-120</td>
</tr>
<tr>
<td>c200</td>
<td>AA # 1192-1931</td>
</tr>
<tr>
<td>NS5</td>
<td>AA # 2054-2995</td>
</tr>
</tbody>
</table>

The host organism for all three HCV recombinant proteins is *S. cerevisiae* (yeast).

The primary purpose of this assay is to screen blood donations so that units containing HCV antibody can be identified and eliminated from the blood supply. Although the presence of anti-HCV does not constitute a diagnosis of HCV infection, the determination of anti-HCV may be used as an aid in the diagnosis of hepatitis C and in the differential diagnosis of non-A, non-B hepatitis in conjunction with determination of liver enzymes, additional serological markers and clinical evaluation. The Hepatitis C Virus Encoded Antigen (Recombinant c22-3, c200 and NS5) used in the manufacture of ORTHO HCV Version 3.0 ELISA Test System is prepared under U.S. License by Novartis Vaccines and Diagnostics, Inc. under a shared manufacturing arrangement.

**PRINCIPLE OF THE PROCEDURE**

The assay procedure is a three-stage test carried out in a microwell coated with a combination of recombinant hepatitis C virus (rHCV) antigen (c22-3, c200 and NS5).

In the first stage, a diluted test specimen is incubated in the test well for a specified length of time. If antibody reactive to any of the three antigens is present in the specimen, antigen-antibody complexes will be formed on the microwell surface. If anti-HCV is not present, complexes will not be formed. In the subsequent washing step, unbound serum or plasma proteins will be removed.

In the second stage, murine monoclonal antibody conjugated to horseradish peroxidase is added to the microwell. The conjugate binds specifically to the human IgG portion of the antigen-antibody complexes. If antigen-antibody complexes are not present, the unbound conjugate will be removed by subsequent washing.

In the third stage, an enzyme detection system composed of o-phenylenediamine (OPD) and hydrogen peroxide is added to the test well. If bound conjugate is present, the OPD will be oxidized, resulting in a colored end product. In this reaction, peroxidase is divalently oxidized by hydrogen peroxide to form an intermediate compound, which is, in turn, reduced to its initial state by subsequent interaction with hydrogen ion donating OPD. The resulting oxidized form of OPD has an orange color. Sulfuric acid is then added to stop the reaction.

The color intensity is dependent upon the amount of bound conjugate and therefore is a function of the concentration of anti-HCV present in the specimen. The color intensity is measured with a microwell reader (photometer) designed to measure light absorbance in a microwell.
<table>
<thead>
<tr>
<th>Label Abbreviations</th>
<th>480 Test Kit Product Code 930740</th>
<th>2400 Test Kit Product Code 930750</th>
<th>Component Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCV</td>
<td>5 plates</td>
<td>25 plates</td>
<td>Hepatitis C Virus (HCV) Encoded Antigen (Recombinant c22-3, c200 and NS5) – Coated Microwell Plates (96 wells each) – c22-3, c200 and NS5 derived from yeast</td>
</tr>
<tr>
<td>CON</td>
<td>1 bottle (125 mL)</td>
<td>5 bottles (125 mL each)</td>
<td>Conjugate – Antibody to Human IgG (Murine Monoclonal) – anti-human IgG heavy chain (murine monoclonal) conjugated to horseradish peroxidase with bovine protein stabilizers Preservative: 1% ProClin™ 300</td>
</tr>
<tr>
<td>SD</td>
<td>1 bottle (190 mL)</td>
<td>4 bottles (190 mL each)</td>
<td>Specimen Diluent – phosphate-buffered saline with bovine protein stabilizers Preservative: 0.1% 2-chloroacetamide</td>
</tr>
<tr>
<td>PC</td>
<td>1 vial (1.0 mL)</td>
<td>4 vials (1.0 mL each)</td>
<td>Positive Control (Human) Source: Treated human serum or plasma containing anti-HCV and nonreactive for hepatitis B surface antigen (HBsAg) and antibody to human immunodeficiency virus type 1 (HIV-1) and type 2 (HIV-2). The anti-HCV serum or plasma has been treated to reduce the titer of potentially infectious virus. However, no test method can rule out the risk of potential infection; <strong>handle as if capable of transmitting infection</strong>. Preservatives: 0.2% sodium azide and 0.9% EDTA</td>
</tr>
<tr>
<td>NC</td>
<td>1 vial (1.5 mL)</td>
<td>5 vials (1.5 mL each)</td>
<td>Negative Control (Human) Source: Human serum or plasma nonreactive for HBsAg, antibody to HIV-1, antibody to HIV-2 and anti-HCV Preservatives: 0.2% sodium azide and 0.9% EDTA</td>
</tr>
<tr>
<td>OPD</td>
<td>1 vial (30 tablets)</td>
<td>5 vials (30 tablets each)</td>
<td>OPD Tablets – contains o-phenylenediamine•2HCl</td>
</tr>
<tr>
<td>SB</td>
<td>1 bottle (190 mL)</td>
<td>4 bottles (190 mL each)</td>
<td>Substrate Buffer-G – citrate-phosphate buffer with 0.02% hydrogen peroxide Preservative: 0.1% 2-chloroacetamide</td>
</tr>
<tr>
<td></td>
<td>21 84 Plate Sealers, disposable</td>
<td></td>
<td>Plate Sealers, disposable</td>
</tr>
</tbody>
</table>

**CAUTION: HANDLE AS IF CAPABLE OF TRANSMITTING INFECTIOUS AGENTS.**

**STORAGE REQUIREMENT**

Store unopened and opened components at 2 to 8°C

FOR IN VITRO DIAGNOSTIC USE

ORTHO HCV Version 3.0 ELISA Test System meets the FDA potency requirements.

**PRECAUTIONS**

1. **CAUTION:** Some components of this kit contain human blood derivatives. No known test method can offer complete assurance that products derived from human blood will not transmit infectious agents. Therefore, all blood derivatives should be handled as potentially infectious. It is recommended that these reagents and human specimens be handled using established good laboratory practices.14,15

2. Wear disposable gloves while handling kit reagents and specimens. Thoroughly wash hands afterward.

3. All specimens should be handled as potentially infectious agents.

4. Sodium azide is included as a preservative in the Positive Control and Negative Control. Sodium azide has been reported to form lead or copper azides in laboratory plumbing. These azides are potentially explosive. To prevent buildup, flush plumbing with a large volume of water while disposing of these solutions in the sink. Following are the Risk and Safety Requirements.16

   **R:** 22 – Harmful if swallowed
   **S:** 28 – After contact with skin, wash immediately with plenty of water.

5. Handle and dispose of all specimens and materials used to perform the test as if they contain infectious agents. Disposal of all specimens and materials should comply with all local, state and federal waste disposal requirements.17,18

6. 4N sulfuric acid (H2SO4) is a strong acid. Wipe up spills immediately. Flush the area of the spill with water. If the acid contacts the skin or eyes, flush with copious amounts of water and seek medical attention.

7. Handle OPD tablets with plastic or Teflon®-coated forceps only. Metal forceps may react with tablets and interfere with the test results. The vial cap may be used for counting and adding tablets.
8. Avoid contact of OPD with eyes, skin or clothing, as OPD may cause irritation or an allergic skin reaction. If OPD should come into contact with the skin, wash thoroughly with water. OPD is toxic for inhalation, ingestion and skin contact. In case of malaise, call a physician. Following are the Risk and Safety Requirements.16


Irritating to eyes. Limited evidence of a carcinogenic effect. May cause sensitization by skin contact.

Very toxic to aquatic organisms, may cause long-term adverse effects in the aquatic environment.

Possible risks of irreversible effects.

S: 26-36/37-45-60-61 – In case of contact with eyes, rinse immediately with plenty of water and seek medical advice. Wear suitable protective clothing and gloves. In case of accident or if you feel unwell, seek medical advice immediately (show the label where possible). This material and its container must be disposed of as hazardous waste. Avoid release to the environment. Refer to special instructions/Safety data sheets.

9. OPD tablets are light- and moisture-sensitive. Keep vial tightly closed when not in use. Bring vial to room temperature (15 to 30°C) before opening. The desiccant pouch must be retained in the vial at all times. Do not use tablets which are yellow, broken, or clumped.

10. Distilled or deionized water must be used for Wash Buffer preparation. Clinical laboratory reagent water Type I or Type II is acceptable.19 Store the water in nonmetallic containers.

11. Do not mix lot numbers of coated microwell plates, Specimen Diluent, Conjugate Reagent, Negative Control, or Positive Control from kits with different lot numbers. Any lot number of Substrate Buffer-G, OPD tablets, 4N sulfuric acid (H2SO4), and 20X Wash Buffer Concentrate may be used provided they are not used beyond the labeled expiration date.

12. All reagents and components must be at room temperature prior to use and kit components returned to 2 to 8°C after use.

13. The microwell strips are sealed in protective pouches with a humidity indicator desiccant. The desiccant, normally blue/purple in color, will turn pink if moisture is present in the pouch. If the desiccant is pink, the microwell strips should not be used.

14. Do not use reagents beyond their labeled expiration date.

15. Cross-contamination between reagents will invalidate the test results. Labeled, dedicated reservoirs for the appropriate reagents are recommended.

16. Ensure that specimen is added to the microwell. Failure to add specimen may produce an erroneous nonreactive result.

17. When using a single-channel micropipette for manual sample addition, use a new pipette tip for each specimen to be assayed. When using a multichannel micropipette, new tips are to be used for each reagent to be added.

18. Strict adherence to the specified wash procedure is crucial to ensure optimum assay performance (see Step 7 of Test Procedure).

19. Do not allow microwells to become dry once the assay has begun.

20. Do not touch the bottom exterior surface of the microwells. Fingerprints or scratches may interfere with reading the microwells. If necessary, wipe the bottom of the microwell strips carefully with a soft, lint-free, absorbent tissue to remove any moisture, dust or debris before reading.

21. Ensure that the microwell strips are level in the microwell strip holder during the test procedure.

22. Negative or positive control values which are not within the expected range (refer to Quality Control Procedures section) may indicate a technique problem or product deterioration.

23. Do not allow sodium hypochlorite fumes from chlorine bleach or other sources to contact the microwell strips during the assay as the color reaction may be inhibited.

24. All pipetting equipment should be used with care, calibrated regularly and maintained following the equipment manufacturer’s instructions.

25. The microwell reader should contain a reference filter with a setting at 620 nm or 630 nm. If an instrument without a reference filter is used, areas in the bottom of the microwells that are opaque, scratched or irregular may cause elevated readings.

26. ProClin™ 300 is included as a preservative in the Conjugate Reagent. Following are the Risk and Safety Requirements.16

R: 43 – May cause sensitization by skin contact.

S: 24-37 – Avoid contact with skin. Wear suitable gloves.

27. 2-chloroacetamide is included as a preservative in Specimen Diluent, 20X Wash Buffer Concentrate and Substrate Buffer-G.16

R: 43 – May cause sensitization by skin contact.

S: 24-37 – Avoid contact with skin. Wear suitable gloves.

28. Delays in plate processing may affect absorbance values.

29. Room temperature is defined as 15 to 30°C.

30. Visual inspections of the reagents should be performed prior to use to check for color change, cloudiness, and precipitates.

PREPARATION OF REAGENTS

1. Preparation of Wash Buffer (1X): Mix 1 part of 20X Wash Buffer Concentrate with 19 parts of distilled or deionized water (1 to 20 dilution). Wash Buffer (1X) is stable for 30 days when stored at room temperature. For longer storage (up to 60 days), keep at 2 to 8°C. Record the date the Wash Buffer (1X) is prepared and the expiration date on the container. Discard Wash Buffer (1X) if visibly contaminated.

NOTE: Any lot number of 20X Wash Buffer Concentrate may be used to prepare this reagent provided it is not used beyond its labeled expiration date.
2. Preparation of Substrate Solution: Clean glass or plastic vessels must be used. Prior to the end of the second incubation, transfer a sufficient amount of Substrate Buffer-G to a container and protect the contents from light. Completely dissolve the appropriate number of OPD tablets in Substrate Buffer-G prior to use. Each microwell plate requires at least 20 mL of Substrate Solution. More Substrate Solution may be needed depending upon the reagent dispenser used. See the instrument manufacturer’s instructions for additional reagent requirements. Below are guidelines for general use.

<table>
<thead>
<tr>
<th>Number of Wells</th>
<th>Number of Plates</th>
<th>Number of OPD Tablets</th>
<th>Substrate Buffer-G (mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>24</td>
<td>0.25</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>48</td>
<td>0.5</td>
<td>2</td>
<td>12</td>
</tr>
<tr>
<td>72</td>
<td>0.75</td>
<td>3</td>
<td>18</td>
</tr>
<tr>
<td>96</td>
<td>1</td>
<td>4</td>
<td>24</td>
</tr>
<tr>
<td>192</td>
<td>2</td>
<td>7</td>
<td>42</td>
</tr>
<tr>
<td>288</td>
<td>3</td>
<td>10</td>
<td>60</td>
</tr>
</tbody>
</table>

The Substrate Solution is stable for 60 minutes after the addition of OPD tablets when held at room temperature in the dark and should be colorless to very pale yellow when used. Record the time when the OPD tablets are added to the Substrate Buffer-G and when it will expire on the container. If it is noticeably yellow in color, discard and prepare more Substrate Solution as required. Do not use more than a single preparation of Substrate Solution per plate.

SPECIMEN COLLECTION, STORAGE, AND HANDLING

NOTE: Handle all specimens as if they are capable of transmitting infectious agents.

Living Donor Specimens
A. Blood specimens collected in glass, plastic, or serum-separator tubes may be used.
B. Plasma specimens collected in EDTA (glass and plastic tubes, plasma preparation tubes), lithium heparin, CPD, CP2D, CPDA-1, ACD, or 4% citrate anticoagulants may be used. Plasma collected with an improper ratio of specimen to anticoagulant should not be used.
C. Whole blood may be stored up to 25°C for 24 hours from time of draw and serum and EDTA plasma specimens may be stored up to 10 days from time of draw at 2-8°C prior to centrifugation. Do not freeze whole blood.
D. Specimens may be stored for up to 10 days from time of draw at 2-8°C following centrifugation, or up to 4 weeks at -20°C undergoing 5 freeze/thaw cycles. Store specimens in appropriately qualified freezers. Mix specimen thoroughly after thawing and before testing.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Whole Blood</th>
<th>Serum and Plasma</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>2-25°C</td>
<td>2-8°C</td>
</tr>
<tr>
<td>2</td>
<td></td>
<td>-20°C</td>
</tr>
<tr>
<td>Collection Time (days)</td>
<td>0 1 2 3 4 5 6 7 8 9 10 days</td>
<td>4 weeks</td>
</tr>
</tbody>
</table>

E. Studies have demonstrated that specimens may be shipped at ambient temperature (up to 37°C) for up to seven days or refrigerated (2 to 8°C) for up to seven days. Upon arrival, specimens should be stored at 2 to 8°C. For shipments requiring extensive transit times (greater than seven days), specimens should be kept frozen (-20°C or below).

F. If specimens are to be shipped, they must be packaged in compliance with International Air Transport Association (IATA) and other applicable guidelines and regulations.

G. No special preparation of the donor is required prior to specimen collection. Blood should be collected by approved medical techniques. Proper sample handling techniques should be employed to avoid microbial contamination.

H. Clear, non-hemolyzed samples are preferred. Precipitates in specimens should be removed by centrifugation.

I. No effect on reactivity was observed when 30 HCV reactive and 30 nonreactive specimens were treated with up to 800 mg/dL of hemoglobin and 30 mg/dL of bilirubin.

J. No effect on reactivity was observed for lipids when 30 HCV reactive and 30 nonreactive specimens were treated with up to 3000 mg/dL of triglyceride.

K. No effect on reactivity was observed in 20 nonreactive specimens containing ≥ 9.0 g/dL total protein.

L. No interference from human anti-mouse antibodies (HAMA) was observed in a 15 member commercially available HAMA panel. No interference from heterophilic antibodies was observed in a 15 member commercially available panel.

M. Do not use heat-treated specimens.

N. Specimens such as pleural fluids, saliva, urine, and nonhuman specimens have not been evaluated with this assay and should not be used.

Cadaveric Donor Specimens
O. Cadaveric specimens may be collected into serum, serum-separator tubes, or EDTA blood collection devices.

P. Cadaveric specimens may be stored for up to 10 days at 2 to 8°C and up to 4 weeks at -20°C undergoing 5 freeze/thaw cycles. Store specimens in appropriately qualified freezers. Mix specimen thoroughly after thawing and before testing.

Q. Studies have demonstrated that specimens may be shipped at ambient temperature (up to 37°C) for up to seven days or refrigerated (2 to 8°C) for up to seven days. Upon arrival, specimens should be stored at 2 to 8°C. For shipments requiring extensive transit times (greater than seven days), specimens should be kept frozen (-20°C or below).

R. If specimens are to be shipped, they must be packaged in compliance with International Air Transport Association (IATA) and other applicable guidelines and regulations.

S. Proper sample handling techniques should be employed to avoid microbial contamination.
T. Clear, non-hemolyzed samples are preferred. Precipitates in specimens should be removed by centrifugation.

U. No effect on reactivity was observed when the level of hemolysis in the cadaveric specimens ranged from 0 mg/dL to 800 mg/dL of hemoglobin.

Specimen Pooling
Testing of these specimens is not recommended. No data are available to interpret tests performed on pooled blood or processed plasma and products made from such pools.

PROCEDURE

Materials Provided
480 Test Kit (Product Code 930740)
2400 Test Kit (Product Code 930750)

Materials Required But Not Provided

1. Adjustable multichannel micropipette capable of delivering 50 µL and 200 µL with at least ±5% accuracy, or equivalent reagent dispenser.
2. Fixed or adjustable single-channel micropipettes capable of delivering 10 µL to 15 µL with at least ±10% accuracy, and 200 µL to 300 µL with at least ±5% accuracy or equivalent pipetter-dilutor.
3. 10 µL to 300 µL disposable pipette tips or equivalent
4. Appropriately sized serological pipette or graduated cylinder
5. Multichannel micropipette reservoir or equivalent reagent container
6. Multichannel aspirator-washer device capable of dispensing and aspirating 300 µL to 800 µL per well. (Consult the device operator's manual for additional technical information.)
7. Dual wavelength microwell reader capable of reading at 490 nm or 492 nm with a reference filter of 620 nm or 630 nm. If an instrument without a reference filter is used, areas in the bottom of the microwells that are opaque, scratched or irregular may cause erroneous readings. Linearity of the microwell reader must range from at least 0 to 2.5 absorbance units. Consult the instrument manufacturer's specifications.
8. 37°C ±1°C incubator (dry or humidified)
9. Distilled or deionized water, clinical laboratory reagent water Type I or Type II is acceptable (see PRECAUTIONS section)
10. 5.25% sodium hypochlorite (chlorine bleach)
11. 4N sulfuric acid (H₂SO₄) – available in the United States from Ortho-Clinical Diagnostics, Inc. (Product Code 933040) or equivalent. To determine suitability of another source of acid, prepare Substrate Solution as described under PREPARATION OF REAGENTS. Add 200 µL of Substrate Solution to three microwells, then add 50 µL of 4N sulfuric acid (H₂SO₄) to be tested to each microwell. Read the microwells at a wavelength of 490 nm or 492 nm with a reference filter of 620 nm or 630 nm at “0 time” and “60 minutes.” All absorbance values at each time interval must be less than or equal to 0.050.
12. White microwell strips (Product Code 50000312, Ortho-Clinical Diagnostics, Inc.) or equivalent uncoated microwells
13. 20X Wash Buffer Concentrate (Product Code 933730, 6 x 150 mL; Ortho-Clinical Diagnostics, Inc.) – phosphate buffer with sodium chloride and detergent. Preservative: 2% 2-chloroacetamide

Test Procedure

1. Approximately 30 minutes prior to the beginning of the procedure, bring kit components to room temperature (15 to 30°C). Invert liquid reagents gently several times, but avoid foaming. Check the incubator temperature; maintain at 37°C ±1°C.
2. Determine the total number of wells needed for the assay. In addition to specimens, one substrate blank, three negative controls and two positive controls must be included on each plate or partial plate. Unused wells should be stored at 2 to 8°C in the supplied foil pouch with desiccant, tightly sealed and used within 42 days of opening the foil pouch. Record the date the pouch is opened and the expiration date of the unused wells in the space provided on the pouch.

Performing the test on less than a full plate is permitted as long as the following conditions are met:
- Microwell strips from different plates can be mixed to assemble full or partial plates as long as they are from the same lot, within the open pouch expiration date and have come from plates that have previously demonstrated proper response to kit controls.
- When assembling a plate which contains strips from a newly opened, previously untested plate, one of these strips should be placed at the beginning of the plate and receive the full complement of kit controls.

CAUTION: Handle microwell strips with care. Do not touch the bottom exterior surface of the wells.
3. Assemble the microwell strips in the microwell strip holder, if necessary. Microwell strips must be level in the microwell strip holder. For incomplete plates, add white or uncoated microwell strips.
4. Prepare a record (plate map) identifying the placement of the controls and specimens in the microwells.

Arrange the assay control wells so that well 1A is the substrate blank. From well 1A arrange all controls in a row (horizontal) or column (vertical) configuration as follows. Configuration is dependent upon software.
5. Verify that any manual dispensing equipment is set to deliver the specified volumes as stated in the procedure, following the equipment manufacturer’s instructions.
Add controls and specimens to the microwells as follows.
Sample Addition:
a. Add 200 µL of Specimen Diluent to all wells except 1A.
b. Add 10 µL of the controls or specimen to the appropriate wells.
c. If the controls and specimens have been manually delivered, to ensure the complete addition of control, or specimen, mix the sample and Specimen Diluent in the well by flushing the pipette tip several times.
For Previously Diluted Sample Addition:
a. Add 300 µL of Specimen Diluent to a tube or container.
b. Add 15 µL of control or specimen to the tube. Mix thoroughly.
c. Transfer 210 µL of each previously diluted control or specimen to the appropriate well position.
6. For manual processing of microwell plates, cover the microwell strip holder with a plate sealer. When using an automated microplate processor for incubation, follow the instrument manufacturer’s recommendations with regard to microwell plate sealing. Incubate at 37°C ±1°C for 60 minutes ±5 minutes.
7. Level the strips in the microwell strip holder, if necessary. With an aspirator-washer device, aspirate and wash all wells five times with Wash Buffer (1X).
CAUTION: Strict adherence to the specified wash procedure is crucial to ensure optimum assay performance. Follow the steps specified in order to ensure thorough washing.
a. Aspirate the sample solutions from the microwells, then fill completely with Wash Buffer. Do not allow the wells to overflow. Allow approximately 20 seconds between the addition of Wash Buffer and subsequent aspiration.
b. Complete the aspirate/fill sequence four additional times (5 times total).
c. Completely aspirate wells. Invert the plate and firmly tap on a clean paper towel to remove excess Wash Buffer, if necessary.
8. Add 200 µL of Conjugate to all wells, except 1A.
9. For manual processing of microwell plates, cover the microwell strip holder with a new, unused plate sealer. When using an automated microplate processor for incubation, follow the instrument manufacturer’s recommendations with regard to microwell plate sealing. Incubate at 37°C ±1°C for 60 minutes ±5 minutes.
10. Prepare sufficient Substrate Solution prior to use in Step 12 to allow time for OPD tablets to dissolve completely. Refer to PREPARATION OF REAGENTS. Do not use more than a single preparation of Substrate Solution on a plate.
11. After the second incubation, wash the wells as described in Step 7.
12. Add 200 µL of Substrate Solution to all wells, including 1A using an adjustable multichannel micropipette or equivalent reagent dispenser capable of delivering 200 µL with at least ±5% accuracy. Incubate at room temperature (15 to 30°C) in the dark for 30 minutes ±1 minute.
13. Add 50 µL of 4N sulfuric acid (H₂SO₄) to all wells, including 1A using an adjustable multichannel micropipette or equivalent reagent dispenser capable of delivering 50 µL with at least ±5% accuracy. To ensure proper mixing, acid should be added forcibly in a steady stream. If necessary, gently tap the plate to mix the contents. Care should be taken to avoid splashing of the contents of the microwells. When using an automated microplate processor, follow the instrument manufacturer’s instructions with regard to mixing.
14. If necessary, wipe moisture from the bottom of the microwell strips carefully with a soft, lint-free, absorbent tissue before reading. If necessary, level the strips in the microwell strip holder. Read the microwell strip plate at a wavelength of 490 nm or 492 nm. For dual wavelength readers, set the reference wavelength at 620 nm or 630 nm. Blank the reader on well 1A according to the instrument manufacturer’s instructions.
For manual calculation, the user should ensure that the blank value (well 1A) has been subtracted from all control and specimen well values prior to applying the Quality Control criteria below.
NOTE: Microwell strip plates must be read within 60 minutes following the addition of 4N sulfuric acid (H₂SO₄). Plates must be stored in the dark until read.

Quality Control Procedures
1. **Substrate Blank Acceptance Criteria**
A plate is considered valid with respect to the substrate blank if the absorbance value of the substrate blank well (well 1A) is greater than or equal to 0.020 and less than or equal to 0.050. The plate is invalid if the substrate blank well is invalid.

2. **Negative Control Acceptance Criteria**
a. Individual negative control values must be less than or equal to 0.120 and greater than or equal to -0.005. Numbers which are between 0.000 and -0.005 inclusive are valid and should be rounded to 0.000 for calculation. If one of the three control values is outside either of these limits, recalculate the negative control mean (NCR) based upon the two acceptable control values. The plate is invalid and the test must be repeated if two or more of the three control values are outside either of the limits.
b. Determine the mean of the negative control values (NCR).
Example:

<table>
<thead>
<tr>
<th>Negative Control</th>
<th>Absorbance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.010</td>
</tr>
<tr>
<td>2</td>
<td>0.030</td>
</tr>
<tr>
<td>3</td>
<td>0.020</td>
</tr>
</tbody>
</table>

Total Absorbance = 0.060

\[
NCR = \frac{\text{Total Absorbance}}{3} = 0.020
\]
3. Positive Control Acceptance Criteria

The positive control is used to verify that the test kit components are capable of detecting a reactive specimen provided the test procedure has been strictly adhered to.

A plate is considered valid with respect to the positive control if both positive control values are greater than or equal to 0.800, within the linear range of the microwell reader and do not differ by more than 0.600.

**NOTE:** Results beyond the upper limit of the linear range of the microwell reader may appear as "OVER" or "***" or ">".

4. Calculation of the Cutoff Value

Cutoff Value = NCx + 0.600

Example:

<table>
<thead>
<tr>
<th>Negative Control</th>
<th>Absorbance</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.010</td>
</tr>
<tr>
<td>2</td>
<td>0.030</td>
</tr>
<tr>
<td>3</td>
<td>0.020</td>
</tr>
</tbody>
</table>

Total Absorbance = 0.060

NCx = \[ \frac{\text{Total Absorbance}}{3} \] = 0.020

Cutoff Value = 0.020 + 0.600 = 0.620

**INTERPRETATION OF RESULTS**

1. Specimens with absorbance values less than -0.025 should be retested in a single microwell. The specimen should be considered nonreactive if the retest absorbance value is less than the Cutoff Value, even if the retest absorbance value remains less than -0.025.

2. Specimens with absorbance values less than the Cutoff Value and greater than or equal to -0.025 are considered nonreactive. Further testing is not required.

3. Specimens with absorbance values greater than or equal to the Cutoff Value are considered initially reactive and should be retested in duplicate before final interpretation.

4. Upon retesting an initially reactive specimen, the specimen is considered repeatedly reactive for antibody to HCV if either or both duplicate determination(s) is (are) reactive, i.e., greater than or equal to the Cutoff Value.

5. After retesting an initially reactive specimen, the specimen is considered nonreactive for antibody to HCV if both duplicate determinations are negative, i.e., less than the Cutoff Value.

**LIMITATIONS OF THE PROCEDURE**

ORTHO HCV Version 3.0 ELISA Test System is limited to the detection of anti-HCV in human serum or plasma. The presence of anti-HCV does not constitute a diagnosis of hepatitis C, but may be indicative of recent and/or past infection by hepatitis C virus. A nonreactive test result does not exclude the possibility of exposure to hepatitis C virus. Levels of anti-HCV may be undetectable in early infection.

Data obtained from testing persons at increased risk or low risk for HCV infection suggest that repeatedly reactive specimens with high absorbance values are more likely to demonstrate the presence of anti-HCV in supplemental testing. Reactivity at or slightly above the Cutoff Value is more frequently nonspecific, especially in specimens obtained from persons at low risk for infection. However, the presence of anti-HCV in some of these specimens can be demonstrated by supplemental testing.

The positive control in the test kit is not to be used to quantitate assay sensitivity. The positive control is used to verify that the test kit components are capable of detecting a reactive specimen provided the test procedure has been strictly adhered to. When positive control values are beyond the linear range of the microwell reader, the positive control cannot be used to assess assay precision. Because true optical density values beyond the linear range are not known, differences between these values and values within the linear range cannot be quantitated.

**EXPECTED RESULTS**

In blood donor populations, the incidence of specimens found repeatedly reactive for anti-HCV by ORTHO HCV Version 3.0 ELISA Test System has typically been less than 1%.

**Reactivity in Blood Donors**

The specificity of ORTHO HCV Version 3.0 ELISA Test System in low-risk populations is based on a population of blood donors from four different test sites. A total of 33,025 specimens were tested. These consisted of 30,025 specimens from presumably healthy volunteer blood donors (sites 1, 2 and 3) and 3,000 specimens from commercial plasma donors (site 4). Rates of initial and repeat reactivities are shown in Table 1. Correlation between initial and repeat reactivity was 97.1%.

**Table 1. Prevalence of Anti-HCV Reactivity in Blood Donors**

<table>
<thead>
<tr>
<th>Site</th>
<th>Number Tested</th>
<th>Nonreactive</th>
<th>Initially Reactive</th>
<th>Repeatedly Reactive</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10,208</td>
<td>10,169</td>
<td>47 (0.46%)</td>
<td>39 (0.38%)</td>
</tr>
<tr>
<td>2</td>
<td>9,772</td>
<td>9,722</td>
<td>54 (0.55%)</td>
<td>50 (0.51%)</td>
</tr>
<tr>
<td>3</td>
<td>10,045</td>
<td>9,998</td>
<td>54 (0.54%)</td>
<td>47 (0.47%)</td>
</tr>
<tr>
<td>4</td>
<td>3,000</td>
<td>2,987</td>
<td>16 (0.53%)</td>
<td>13 (0.43%)</td>
</tr>
<tr>
<td>TOTAL</td>
<td>33,025</td>
<td>32,876</td>
<td>171 (0.52%)</td>
<td>149 (0.45%)</td>
</tr>
</tbody>
</table>
Specimens repeatedly reactive by ELISA were tested by supplemental strip immunoblot assay (SIA). Upon removal of specimens that were positive or indeterminate by SIA, specificity* in this low prevalence population was 99.95% (5 units per 10,000 donations).

*Specificity was calculated as follows:
Specificity = (TN/TN+FP)
Where TN = true negatives, that is, the number of specimens nonreactive by ELISA.
FP = false positives, that is, repeatedly reactive by ELISA and negative by SIA.

Detection of Anti-HCV Seroconversion in Transfusion Recipients with Transfusion-Associated NANBH (TA-NANBH)

Serial specimens collected from 21 patients with clinically documented TA-NANBH were tested at clinical (patient) site 2 by both ORTHO HCV Version 3.0 ELISA Test System and ORTHO HCV 2.0 ELISA Test System. HCV Version 3.0 detected seroconversion to anti-HCV in earlier bleeds in 5/21 (24%) patients. HCV 2.0 detected seroconversion to anti-HCV in an earlier bleed in 1/21 (5%) patients. In 15 patients, HCV Version 3.0 and HCV 2.0 detected anti-HCV in the same serial bleed.

Table 2. Difference in Detection of Anti-HCV in 21 TA-NANBH Patients

<table>
<thead>
<tr>
<th>Patient Number</th>
<th>Earlier Detection by HCV ELISA</th>
<th>Difference in Detection Bleeds (Days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>HCV Version 3.0</td>
<td>2 (20)</td>
</tr>
<tr>
<td>2</td>
<td>HCV Version 3.0</td>
<td>1 (34)</td>
</tr>
<tr>
<td>3</td>
<td>HCV Version 3.0</td>
<td>1 (27)</td>
</tr>
<tr>
<td>4</td>
<td>HCV Version 3.0</td>
<td>2 (20)</td>
</tr>
<tr>
<td>5</td>
<td>HCV Version 3.0</td>
<td>2 (29)</td>
</tr>
<tr>
<td>6</td>
<td>HCV 2.0</td>
<td>1 (35)</td>
</tr>
</tbody>
</table>

There was no difference in the time to detection of anti-HCV ELISA reactivity in the remaining 15 patients.

Reactivity in Patients with NANBH

NANBH patients from clinical (patient) site 3 were tested with ORTHO HCV Version 3.0 ELISA Test System. Patients were divided into acute and chronic duration disease based on alanine aminotransferase (ALT) levels and patterns. The frequency of anti-HCV detection was 75% in acute duration and 88% in chronic duration (Table 3).

Table 3. Detection of Anti-HCV in Patients with NANBH

<table>
<thead>
<tr>
<th>Category</th>
<th>Number of Patients</th>
<th>Repeatedly Reactive (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute</td>
<td>231</td>
<td>75.3</td>
</tr>
<tr>
<td>Chronic</td>
<td>59</td>
<td>88.1</td>
</tr>
</tbody>
</table>

* Diagnostic criteria for acute NANBH included: new onset of a symptomatic illness including a peak ALT of greater than 500 IU/L; absence of serologic markers for acute hepatitis A and B and no evidence of toxic (drug) induced hepatitis.

b Diagnostic criteria for chronic NANBH included: a negative test for hepatitis B surface antigen and abnormal ALT activity for more than 6 months.

Reactivity in High-Risk Populations

Reactivity in populations at risk for acquiring/transmitting NANBH was studied at two clinical (patient) sites. A total of 602 specimens from high-risk patients were tested with ORTHO HCV Version 3.0 ELISA Test System. Table 4 shows the prevalence of anti-HCV reactivity in three high-risk groups.

Table 4. Detection of Anti-HCV in High-Risk Populations

<table>
<thead>
<tr>
<th>Risk Group</th>
<th>Number of Specimens</th>
<th>Repeatedly Reactive (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemophiliacs</td>
<td>302</td>
<td>74.5</td>
</tr>
<tr>
<td>IV Drug Abusers</td>
<td>200</td>
<td>94.0</td>
</tr>
<tr>
<td>Renal Dialysis</td>
<td>100</td>
<td>19.0</td>
</tr>
</tbody>
</table>

SPECIFIC PERFORMANCE CHARACTERISTICS

Reproducibility

The interassay and intra-assay reproducibility of ORTHO HCV Version 3.0 ELISA Test System was assessed with a nonreactive specimen, a weakly reactive specimen (approximately 1.3 times the Cutoff Value), and a strongly reactive specimen (approximately 3.0 times the Cutoff Value). The specimens were tested with three different kit lots at two different test sites. Mean optical density (OD), standard deviation (SD) and coefficient of variation (%CV) results are shown in Table 5.

Table 5. Reproducibility of ORTHO HCV Version 3.0 ELISA Test System

<table>
<thead>
<tr>
<th>Parameter Measured</th>
<th>Nonreactive (Mean OD = 0.042) SD</th>
<th>%CV</th>
<th>Weakly Reactive (Mean OD = 0.773) SD</th>
<th>%CV</th>
<th>Strongly Reactive (Mean OD = 1.857) SD</th>
<th>%CV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interassay</td>
<td>0.003 NA*</td>
<td>0.079 10.3</td>
<td>0.168 9.0</td>
<td>0.086 11.1</td>
<td>0.158 8.5</td>
<td></td>
</tr>
<tr>
<td>Intra-assay</td>
<td>0.026 NA*</td>
<td>0.086 11.1</td>
<td>0.168 9.0</td>
<td>0.086 11.1</td>
<td>0.158 8.5</td>
<td></td>
</tr>
</tbody>
</table>

*% CVs are not meaningful when ODs approach zero.

Specificity

A total of 33,025 specimens from blood donors were tested, 32,876 of which were nonreactive. Upon removal of specimens positive or indeterminate in supplemental testing by SIA, specificity of ORTHO HCV Version 3.0 ELISA Test System in this low prevalence population was 99.95%.
Sensitivity
Sensitivity of ORTHO HCV Version 3.0 ELISA Test System was evaluated in two studies:

The time to detect anti-HCV seroconversion in TA-NANBH patients (compared to ORTHO HCV 2.0 ELISA Test System). See Table 2.

The frequency of detection of anti-HCV repeatedly reactive results in a NANBH patient population. See Table 3.

PERFORMANCE CHARACTERISTICS OF CADAVERIC SPECIMEN TESTING

Reproducibility
Reproducibility of ORTHO HCV Version 3.0 ELISA Test System was assessed using 20 cadaveric and 20 normal donor sera. These specimens were spiked with anti-HCV positive plasma to give reactivity near the assay cutoff. Each of the specimens was tested once on six different days on each of three lots of ORTHO HCV Version 3.0 ELISA Test System at one site. Reproducibility testing was performed by both manual and automated processing methods. For each processing method, cadaveric and living donor specimens were 100% reactive across kit lots and the %CVs were comparable for both specimen groups.

Kit Lot 1

<table>
<thead>
<tr>
<th>Number of Donors</th>
<th>Replicates</th>
<th>% Positive</th>
<th>Mean S/C</th>
<th>CV(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual Cadaveric</td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>1.828</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>1.706</td>
</tr>
<tr>
<td>Automated Cadaveric</td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>1.961</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>1.949</td>
</tr>
</tbody>
</table>

Kit Lot 2

<table>
<thead>
<tr>
<th>Number of Donors</th>
<th>Replicates</th>
<th>% Positive</th>
<th>Mean S/C</th>
<th>CV(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual Cadaveric</td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>1.981</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>1.851</td>
</tr>
<tr>
<td>Automated Cadaveric</td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>2.086</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>2.102</td>
</tr>
</tbody>
</table>

Kit Lot 3

<table>
<thead>
<tr>
<th>Number of Donors</th>
<th>Replicates</th>
<th>% Positive</th>
<th>Mean S/C</th>
<th>CV(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual Cadaveric</td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>2.070</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>1.943</td>
</tr>
<tr>
<td>Automated Cadaveric</td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>2.101</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>120</td>
<td>100</td>
<td>2.160</td>
</tr>
</tbody>
</table>

Specificity
Specificity was evaluated using 50 cadaveric specimens collected up to 23.7 hours after death and 50 normal donor specimens. Testing was performed across three lots of ORTHO HCV Version 3.0 ELISA Test System by both manual and automated processing methods. For the manual method, the mean signal to cutoff (S/C) ratio was 0.071 for the cadaveric specimens, and the mean S/C ratio was 0.030 for the normal donor specimens. For the automated method, the mean signal to cutoff (S/C) ratio was 0.064 for the cadaveric specimens, and the mean S/C ratio was 0.042 for the normal donor specimens. The results are presented in Table 6.

Table 6. Reactivity in the ORTHO HCV Version 3.0 ELISA Test System

<table>
<thead>
<tr>
<th>Population</th>
<th>Number of Specimens</th>
<th>Mean S/C</th>
<th>Nonreactive</th>
<th>Initially Reactive</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Manual Processing</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cadaveric</td>
<td>50</td>
<td>0.071</td>
<td>50 (100.0%)</td>
<td>0 (0.0%)</td>
</tr>
<tr>
<td>Normal Donor</td>
<td>50</td>
<td>0.030</td>
<td>50 (100.0%)</td>
<td>0 (0.0%)</td>
</tr>
<tr>
<td></td>
<td>Automated Processing</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cadaveric</td>
<td>50</td>
<td>0.064</td>
<td>50 (100.0%)</td>
<td>0 (0.0%)</td>
</tr>
<tr>
<td>Normal Donor</td>
<td>50</td>
<td>0.042</td>
<td>50 (100.0%)</td>
<td>0 (0.0%)</td>
</tr>
</tbody>
</table>

The ORTHO HCV Version 3.0 ELISA Test System has an estimated specificity in cadaveric specimens of 100.0% (50/50) with a 95% exact confidence interval of 92.9% to 100.0%.
Sensitivity

Sensitivity was evaluated using 50 cadaveric specimens collected up to 23.7 hours after death and 50 normal donor specimens. All specimens were pre-screened for anti-HCV and were found to be nonreactive. All specimens were spiked with anti-HCV positive plasma to give reactivity near the assay cutoff. Testing was performed approximately 47 hours after spiking using three lots of ORTHO HCV Version 3.0 ELISA Test System by both manual and automated processing methods. Since the specimens were spiked to be reactive, duplicate repeat testing was not performed for initially reactive specimens. For the manual method, the mean signal to cutoff (S/C) ratio was 1.904 for the cadaveric specimens, and the mean S/C ratio was 1.893 for the normal donor specimens. The calculated difference between the cadaveric specimens and the normal donor specimens tested by the manual method was 0.011 S/C, which was determined by the F-test not to be statistically significant (p=0.866). For the automated method, the mean signal to cutoff (S/C) ratio was 2.136 for the cadaveric specimens, and the mean S/C ratio was 2.362 for the normal donor specimens. The calculated difference between the cadaveric specimens and the normal donor specimens tested by the automated method was 0.226 S/C, which was determined by the F-test to be statistically significant (p=0.0003). However, all results for the cadaveric and normal donor specimens were reactive with the ORTHO HCV Version 3.0 ELISA Test System resulting in 100.0% reactivity. The results are presented in Table 7.

Table 7. Reactivity in the ORTHO HCV Version 3.0 ELISA Test System

<table>
<thead>
<tr>
<th>Population</th>
<th>Number of Specimens</th>
<th>Mean S/C</th>
<th>Nonreactive</th>
<th>Initially Reactive</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Manual Processing</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cadaveric</td>
<td>50</td>
<td>1.904</td>
<td>0 (0.0%)</td>
<td>50 (100.0%)</td>
</tr>
<tr>
<td>Normal Donor</td>
<td>50</td>
<td>1.893</td>
<td>0 (0.0%)</td>
<td>50 (100.0%)</td>
</tr>
<tr>
<td>Automated Processing</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cadaveric</td>
<td>50</td>
<td>2.136</td>
<td>0 (0.0%)</td>
<td>50 (100.0%)</td>
</tr>
<tr>
<td>Normal Donor</td>
<td>50</td>
<td>2.362</td>
<td>0 (0.0%)</td>
<td>50 (100.0%)</td>
</tr>
</tbody>
</table>

The ORTHO HCV Version 3.0 ELISA Test System has an estimated sensitivity in spiked cadaveric specimens of 100.0% (50/50) with a 95% exact confidence interval of 92.9% to 100.0%.

### SUMMARY OF REVISIONS

<table>
<thead>
<tr>
<th>Section</th>
<th>Revision</th>
</tr>
</thead>
<tbody>
<tr>
<td>INTENDED USE</td>
<td>Added the use of cadaveric specimens.</td>
</tr>
<tr>
<td>REAGENTS</td>
<td>Put 480 and 2400 Test Kit Component descriptions in table format.</td>
</tr>
<tr>
<td>STORAGE REQUIREMENT</td>
<td>Added “STORAGE REQUIREMENT” heading and revised the storage requirement information to “Store unopened and opened components at 2 to 8°C.”</td>
</tr>
<tr>
<td>PREPARATION OF REAGENTS</td>
<td>Step 1 – Changed first sentence to “Mix 1 part of 20X Wash Buffer Concentrate with 19 parts of distilled or deionized water (1 to 20 dilution).”</td>
</tr>
<tr>
<td>SPECIMEN COLLECTION, STORAGE, AND HANDLING</td>
<td>Step 2 – Added the statement “Do not use more than a single preparation of Substrate Solution per plate.” to the last sentence.</td>
</tr>
<tr>
<td></td>
<td>Changed section title to Specimen Collection, Storage, and Handling.</td>
</tr>
<tr>
<td></td>
<td>Reformatted section into three (3) subsections: Living Donor Specimens, Cadaveric Blood Specimens, and Specimen Pooling. Bulleted information format introduced.</td>
</tr>
<tr>
<td></td>
<td>Revised entire section to include updated specimen collection, preparation, shipping and storage information.</td>
</tr>
<tr>
<td></td>
<td>Added “Cadaveric specimens may be collected into serum, serum-separator tubes or EDTA blood collection devices.”</td>
</tr>
<tr>
<td></td>
<td>Added claims for total protein, human anti-mouse antibodies (HAMA) and heterophilic antibodies.</td>
</tr>
<tr>
<td></td>
<td>Revised statement on Specimen Pooling to read “Testing of these specimens is not recommended. No data are available to interpret tests performed on pooled blood or processed plasma and products made from such pools.”</td>
</tr>
<tr>
<td>MATERIALS PROVIDED</td>
<td>Deleted the words (“See REAGENTS for complete listing.”) Redundant information.</td>
</tr>
<tr>
<td>MATERIALS REQUIRED BUT NOT PROVIDED</td>
<td>Updated Product code for White microwell strips from 651200191 to 50000312.</td>
</tr>
<tr>
<td>QUALITY CONTROL PROCEDURES</td>
<td>Number 1 – Added the statement “The plate is invalid if the substrate blank well is invalid.”</td>
</tr>
<tr>
<td>PERFORMANCE CHARACTERISTICS</td>
<td>Added the reproducibility, specificity and sensitivity performance claims for cadaveric specimens.</td>
</tr>
<tr>
<td>BIBLIOGRAPHY</td>
<td>Added #23 for International Air Transportation Association (IATA): Dangerous Goods Regulations.</td>
</tr>
<tr>
<td>All Sections</td>
<td>Minor editorial clarifications and use of bold text to highlight critical information.</td>
</tr>
</tbody>
</table>
BIBLIOGRAPHY / BIBLIOGRAPHIE / BIBLIOGRAFIA

The following symbols may have been used in the labeling of this product. / Les symboles suivants ont pu être utilisés sur l’étiquette de ce produit. / Los siguientes símbolos pueden haber sido empleados en el etiquetado de este producto.

- **Do Not Reuse** / Ne pas réutiliser / No reutilizar
- **Use by or Expiration Date (Year-Month-Day)** / À utiliser avant la date de péremption (année-mois-jour) / Usar antes de o Fecha de caducidad (año-mes-día)
- **Lot Number** / Numéro de lot / Número de lote
- **Serial Number** / Numéro de série / Número de serie
- **Catalog Number or Product Code** / Référence catalogue ou code produit / Referencia de catálogo o Código del producto
- **Attention: See Instructions for Use** / Attention : consulter le feuillet technique / Atención: Consultar las instrucciones de uso
- **Manufacturer** / Fabricant / Fabricante
- **Authorized Representative in the European Community** / Mandataire dans l’Union européenne / Representante autorizado en la Unión Europea
- **Contains Sufficient for “n” Tests** / Suffisant pour << n >> dosages / Contiene suficiente para “n” ensayos

**KEY TO SYMBOLS / LÉGENDE DES SYMBOLES / CLAVE DE LOS SÍMBOLOS**

**IVD**
In vitro Diagnostic Medical Device / Pour diagnostic in vitro / Producto sanitario para diagnóstico in vitro

**Upper Limit of Temperature** / Conserver à une température égale ou inférieure à / Límite superior de temperatura

**Lower Limit of Temperature** / Conserver à une température égale ou supérieure à / Límite inferior de temperatura

Temperature Limitation / Conserver à une température comprise entre / Limitación de temperatura

**Consult Instructions for Use, “n” Version** / Consultez le feuillet technique << n >> version / Atención: ver las instrucciones de uso “n” versión

**Biological Risks** / Risques biologiques / Riesgos biológicos

**Do not use if damaged** / Ne pas utiliser si endommagé / No usar si está dañado

**Irritant** / Irritant / Irritante

**Harmful** / Nocif / Nocivo

**Toxic** / Toxique / Tóxico
KEY TO SYMBOLS / LÉGENDE DES SYMBOLES /
CLAVE DE LOS SÍMBOLOS

Continued / Suite / Continuación

Dangerous for the Environment / Dangereux pour l’environnement / Peligroso para el medio ambiente

Fragile, Handle with Care / Attention, fragile / Frágil; manipular con cuidado

Keep Dry / Tenir au sec / Mantener seco

This end up / Haut / Este lado hacía arriba

Positive Control / Contrôle positif / Control positivo

Negative Control / Contrôle négatif / Control negativo

Positive Calibrator / Calibrateur positif / Calibrador positivo

Negative Calibrator / Calibrateur négatif / Calibrador negativo

Confirmatory Control / Contrôle de confirmation / Control de confirmación

Recombinant Antigens Provided by / Antigènes recombinants fournis par / Antígenos recombinantes suministrados por

Antibody to Hepatitis B Surface Antigen / Antibodies to Hepatitis B Surface Antigen

Antibody to Hepatitis B Surface Antigen: Peroxidase Conjugate Concentrate

Der Grüne Punkt (the Green Dot). Manufacturer follows certain packaging material waste disposal management regulations. / Der Grüne Punkt (Point Vert). Le fabricant suit certaines règles de mise au rebut pour les déchets des matériaux d’emballage / Punto Verde (der grüne Punkt). El fabricante sigue la regulación sobre gestión de residuos de los embalajes