Hepatitis C virus RNA in Serum
NHANES 2013-2014

Laboratory Procedure Manual

Analyte: Hepatitis C virus RNA
Matrix: Serum
Method: AmpliPrep/COBAS® TaqMan® HCV Test
First Published: September 2013
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As performed by: Assay Development and Diagnostic Reference Laboratory
Laboratory Branch
Division of Viral Hepatitis
National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention

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Important Information for Users
The National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention periodically refines these laboratory methods. It is the responsibility of the user to contact the person listed on the title page of each write-up before using the analytical method to find out whether any changes have been made and what revisions, if any, have been incorporated.
This document details the Lab Protocol for testing the items listed in the following table:

<table>
<thead>
<tr>
<th>Lab Name</th>
<th>Variable Name</th>
<th>SAS Label</th>
</tr>
</thead>
<tbody>
<tr>
<td>HEPC_H</td>
<td>LBXHCR</td>
<td>Hepatitis C RNA (HCV-RNA)</td>
</tr>
</tbody>
</table>
1. SUMMARY OF TEST PRINCIPLE AND CLINICAL RELEVANCE

HCV infection is the most common chronic blood-borne infection in the United States with an estimated seroprevalence of 1.6%. An estimated 3.2 million Americans suffer from chronic HCV infection making it the leading cause of chronic liver disease. An estimated 85% of those with acute hepatitis C progress to chronic infection, of which 20%-25% will develop complications such as cirrhosis within 2 to 3 decades of its onset. Of the patients with cirrhosis, a smaller percentage will progress to decompensated liver disease, hepatocellular carcinoma, and death. In the United States, chronic hepatitis C (CHC) is responsible for an estimated 8,000 to 10,000 deaths per year and is the leading cause of liver transplantation. The burden of HCV-associated disease is likely to increase during the next 10 to 20 years as the infected cohort reaches an age at which complications of liver disease typically occur.

The COBAS AmpliPrep/COBAS TaqMan HCV Test is a nucleic acid amplification test for the quantitation of HCV RNA in human serum or plasma. Specimen preparation is automated using the COBAS AmpliPrep Instrument with amplification and detection automated using the COBAS TaqMan Analyzer or the COBAS TaqMan 48 Analyzer.

The COBAS AmpliPrep/COBAS TaqMan HCV Test is based on three major processes: (1) specimen preparation to isolate HCV RNA; (2) reverse transcription of the target RNA to generate complementary DNA (cDNA), and (3) simultaneous PCR amplification of target cDNA and detection of cleaved dual-labeled oligonucleotide detection probe specific to the target.

The COBAS AmpliPrep/COBAS TaqMan HCV Test permits automated specimen preparation followed by automated reverse transcription, PCR amplification and detection of HCV target RNA and HCV Quantitation Standard (QS) Armored RNA. The Master Mix reagent contains primers and probes specific for both HCV RNA and HCV QS Armored RNA. The Master Mix has been developed to ensure similar quantitation of HCV genotypes 1 through 6. The detection of amplified DNA is performed using a target-specific and a QS-specific dual-labeled oligonucleotide probe that permit independent identification of HCV amplicon and HCV QS amplicon.

The quantitation of HCV viral RNA is performed using the HCV QS. The HCV QS compensates for effects of inhibition and controls the preparation and amplification processes, allowing a more accurate quantitation of HCV RNA in each specimen. The HCV QS is a non-infectious Armored RNA construct that contains HCV sequences with identical primer binding sites as the HCV target RNA and a unique probe binding region that allows HCV QS amplicon to be distinguished from HCV target amplicon.

The HCV QS is added to each specimen at a known copy number and is carried through the specimen preparation, reverse transcription, PCR amplification and detection of cleaved dual-labeled oligonucleotide detection probes. The COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer calculates the HCV RNA concentration in the test specimens by comparing the HCV signal to the HCV QS signal for each specimen and control.
Hepatitis C virus RNA in Serum
NHANES 2013-2014

Target Selection
Selection of the target RNA sequence for HCV depends on identification of regions within the HCV genome that show maximum sequence conservation among the various HCV genotypes. Generic silica-based specimen preparation is used to capture the HCV RNA and HCV QS RNA and defined oligonucleotides are used as primers in amplification of the HCV RNA and HCV QS RNA. A target-specific and a QS-specific dual-labeled oligonucleotide probe permit independent identification of HCV amplicon and HCV QS amplicon. Accordingly, the appropriate selection of the primers and the dual-labeled oligonucleotide probe is critical to the ability of the test to amplify and detect the HCV genotypes. The COBAS AmpliPrep/COBAS TaqMan HCV Test uses reverse transcription and PCR amplification primers that define a sequence within the highly conserved region of the 5'-untranslated region of the HCV genome. The nucleotide sequence of the primers has been optimized to yield comparable amplification of six HCV genotypes.

Specimen Preparation
The COBAS AmpliPrep/COBAS TaqMan HCV Test utilizes automated specimen preparation on the COBAS AmpliPrep Instrument by a generic silica-based capture technique. The procedure processes 850 μL of plasma or serum. The HCV virus particles are lysed by incubation at elevated temperature with a protease and chaotrophic lysis/binding buffer that releases nucleic acids and protects the released HCV RNA from RNases in serum or plasma. Protease and a known number of HCV QS RNA molecules are introduced into each specimen along with the lysis reagent and magnetic glass particles. Subsequently, the mixture is incubated and the HCV RNA and HCV QS RNA are bound to the surface of the magnetic glass particles. Unbound substances, such as salts, proteins and other cellular impurities, are removed by washing the magnetic glass particles. After separating the beads and completing the washing steps, the adsorbed nucleic acids are eluted at elevated temperature with an aqueous solution. The processed specimen, containing the magnetic glass particles as well as released HCV RNA and HCV QS RNA, is added to the amplification mixture and transferred to the COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer. The HCV target RNA and the HCV QS RNA are then reverse transcribed, amplified and simultaneously detected by cleavage of a target-specific and a QS-specific dual-labeled oligonucleotide probe.

Reverse Transcription and PCR Amplification
The reverse transcription and PCR amplification reaction is performed with the thermostable recombinant enzyme Thermus specie DNA Polymerase (Z05). In the presence of manganese (Mn²⁺) and under the appropriate buffer conditions, Z05 has both reverse transcriptase and DNA polymerase activity. This allows both reverse transcription and PCR amplification to occur together with real-time detection of the amplicon.

Processed specimens are added to the amplification mixture in amplification tubes (K-tubes) in which both reverse transcription and PCR amplification occur. The reaction mixture is heated to allow a downstream primer to anneal specifically to the HCV target RNA and to the HCV QS RNA. In the presence of Mn²⁺ and excess deoxynucleotide triphosphates (dNTPs), including deoxyadenosine, deoxyguanosine, deoxycytidine and
Target Amplification
Following reverse transcription of the HCV target RNA and the HCV QS RNA, the Thermal Cycler in the COBAS TaqMan 48 Analyzer heats the reaction mixture to denature the RNA:cDNA hybrid and to expose the specific primer target sequences. As the mixture cools, the primers anneal to the target DNA. The thermostable *Thermus specie* Z05 DNA Polymerase (Z05) in the presence of Mn$^{2+}$ and excess deoxynucleotide triphosphates (dNTPs), including deoxyadenosine, deoxyguanosine, deoxycytidine and deoxyuridine (in place of thymidine) triphosphates, extends the annealed primers along the target template to produce a double-stranded DNA molecule termed an amplicon. The COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer automatically repeats this process for a designated number of cycles, with each cycle intended to double the amount of amplicon DNA. The required number of cycles is preprogrammed into the COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer. Amplification occurs only in the region of the HCV genome between the primers; the entire HCV genome is not amplified.

Selective Amplification
Selective amplification of target nucleic acid from the specimen is achieved in the COBAS AmpliPrep/COBAS TaqMan HCV Test by the use of AmpErase (uracil-N-glycosylase) enzyme and deoxyuridine triphosphate (dUTP). The AmpErase enzyme recognizes and catalyzes the destruction of DNA strands containing deoxyuridine but not DNA containing deoxythymidine. Deoxyuridine is not present in naturally occurring DNA, but is always present in amplicon due to the use of deoxyuridine triphosphate as one of the dNTPs in the Master Mix reagent; therefore, only amplicon contains deoxyuridine. Deoxyuridine renders contaminating amplicon susceptible to destruction by the AmpErase enzyme prior to amplification of the target DNA. Also, any nonspecific product formed after initial activation of the Master Mix by manganese is destroyed by the AmpErase enzyme. The AmpErase enzyme, which is included in the Master Mix reagent, catalyzes the cleavage of deoxyuridine-containing DNA at the deoxyuridine residues by opening the deoxyribose chain at the C1-position. When heated in the first thermal cycling step, the amplicon DNA chain breaks at the position of the deoxyuridine, thereby rendering the DNA non-amplifiable. The AmpErase enzyme remains inactive for a prolonged period of time once exposed to temperatures above 55°C, i.e., throughout the thermal cycling steps, and therefore does not destroy target amplicon formed after PCR reaction.
Detection of PCR Products in a COBAS TaqMan Test
The COBAS AmpliPrep/COBAS TaqMan HCV Test utilizes real-time PCR technology. The use of dual-labeled fluorescent probes allows for real-time detection of PCR product accumulation by monitoring of the emission intensity of fluorescent reporter dyes released during the amplification process. The probes consist of HCV and HCV QS-specific oligonucleotide probes with a reporter dye and a quencher dye. In the COBAS AmpliPrep/COBAS TaqMan HCV Test, the HCV and HCV QS probes are labeled with different fluorescent reporter dyes. When these probes are intact, the fluorescence of the reporter dye is suppressed by the proximity of the quencher dye due to Förster-type energy transfer effects. During PCR, the probe hybridizes to a target sequence and is cleaved by the 5'→3' nuclease activity of the thermostable Z05 DNA polymerase. Once the reporter and quencher dyes are released and separated, quenching no longer occurs, and the fluorescent activity of the reporter dye is increased. The amplification of HCV RNA and HCV QS RNA are measured independently at different wavelengths. This process is repeated for a designated number of cycles, each cycle effectively increasing the emission intensity of the individual reporter dyes, permitting independent identification of HCV RNA and HCV QS RNA. The PCR cycle where a growth curve starts exponential growth is related to the amount of starting material at the beginning of the PCR.

Fundamentals of COBAS TaqMan Test Quantitation
The COBAS AmpliPrep/COBAS TaqMan HCV Test is inherently quantitative over a wide dynamic range since the monitoring of amplicon is performed during the exponential phase of amplification. The higher the HCV titer of a specimen, the earlier the fluorescence of the reporter dye of the HCV probe rises above the baseline fluorescence level. Since the amount of HCV QS RNA is constant between all specimens, the fluorescence of the reporter dye of the HCV QS probe should appear at the same cycle for all specimens. In specimens, where the QS amplification and detection is affected by inhibition or poor specimen recovery, the appearance of fluorescence will be delayed, thereby enabling the calculated titer of HCV target RNA to be adjusted accordingly. The appearance of the specific fluorescent signals is reported as a critical threshold value (Ct). The Ct is defined as the fractional cycle number where reporter dye fluorescence exceeds a predetermined threshold (the Assigned Fluorescence Level), and starts the beginning of an exponential growth phase of this signal. A higher Ct value indicates a lower titer of initial HCV target material. A 2-fold increase in titer correlates with a decrease of 1 Ct for target HCV RNA; a 10-fold increase in titer correlates with a decrease of 3.3 Ct.

HCV RNA Quantitation
The COBAS AmpliPrep/COBAS TaqMan HCV Test quantitates HCV viral RNA by utilizing a second target sequence (HCV Quantitation Standard) that is added to each test specimen at a known concentration. The HCV QS is a non-infectious Armored RNA construct, containing fragments of HCV sequences with primer binding regions identical to those of the HCV target sequence. The HCV QS generates an amplification product of the same length and base composition as the HCV target RNA. The detection probe binding region of the HCV QS has been modified to differentiate HCV QS amplicon from HCV target amplicon.
Hepatitis C virus RNA in Serum  
NHANES 2013-2014

During the annealing phase of the PCR on the COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer, the specimens are illuminated and excited by filtered light and filtered emission fluorescence data are collected for each specimen. The readings from each specimen are then corrected for instrumental fluctuations. These fluorescence readings are sent by the instrument to the AMPLILINK software and stored in a database. Pre-Checks are used to determine if the HCV RNA and HCV QS RNA data represent sets that are valid, and flags are generated when the data lie outside the preset limits. After all Pre-Checks are completed and passed, the fluorescence readings are processed to generate Ct values for the HCV RNA and the HCV QS RNA. The lot-specific calibration constants provided with the COBAS AmpliPrep/COBAS TaqMan HCV Test are used to calculate the titer value for the specimens and controls based upon the HCV RNA and HCV QS RNA Ct values. The COBAS AmpliPrep/COBAS TaqMan HCV Test is standardized against the First WHO International Standard for Hepatitis C Virus RNA for Nucleic Acid Amplification Technology Assays (NIBSC code 96/790) and titer results are reported in International Units (IU/mL).

2. SAFETY PRECAUTIONS

Test kits contain components derived from human serum or plasma. Although various treatments in the manufacturing process are sufficient to inactivate most blood-borne pathogens, there is no assurance that these reagents are entirely noninfectious. Therefore, test kit components should be treated as though they are capable of transmitting HCV. Consider all serum specimens for analysis potentially positive for infectious agents including HIV, hepatitis B virus and HCV. Controls and samples should be handled as if infectious using safe laboratory procedures such as those outlined in *Biosafety in Microbiological and Biomedical Laboratories* and in the CLSI Document M29-A.

Observe universal precautions when performing the assay, thoroughly clean and disinfect all work surfaces with a freshly prepared solution of 0.5% sodium hypochlorite in deionized or distilled water, handle samples with extreme care to prevent sample contamination, use new, sterile aerosol barrier or positive displacement RNase-free pipette tips and sterile pipettes, wear personal protective apparel, disposable gloves and eyewear during all steps of this method to minimize both infectious and chemical contamination hazards.
Hepatitis C virus RNA in Serum
NHANES 2013-2014

Do not eat, drink, smoke, or apply cosmetics in areas where reagents or samples are handled. If skin or mucous membrane exposure occurs, immediately wash the area with copious amounts of water. Seek medical advice. Place all plastic and glassware contaminated with serum in a plastic autoclave bag for disposal. Do not use components beyond the expiration date on the kit. Alterations in the physical appearance of kit components may indicate instability or deterioration. Do not pool reagents from different lots or from different bottles of the same lot. Do not mix reagent cassettes or controls from different kits. Do not open COBAS AmpliPrep cassettes and exchange, mix, remove or add bottles. Store the kit away from any source of contaminating DNA, especially amplified nucleic acid. We recommend Biosafety Level 2 containment procedures as described in CDC/NIH publication #93-8395 be used by those handling test specimens and kit reagents. Use a Unidirectional work flow proceeding from the sample preparation to the amplification and detection steps. To help prevent laboratory areas from becoming contaminated with amplified RT-PCR product, maximize the physical separation of the pre- and post-amplification steps. Do not return samples, equipment, or reagents to the area where you performed the previous step. If you need to return to a previous work area, first perform the appropriate anti-contamination safeguards. Avoid microbial and RBase contamination of reagents.

Use all pipetting devices and instruments with care and follow the manufacturer’s instructions for calibration and quality control.

**HCV QS, CAP/CTM Mn** and **HCV MMX** contain sodium azide. Sodium azide may react with lead and copper plumbing to form highly explosive metal azides. While disposing of sodium azide containing solutions down laboratory sinks, flush the drains with a large volume of water to prevent azide buildup.

Do not allow **HCV CS2** and liquid waste from the COBAS AmpliPrep Instrument, which contain guanidine thiocyanate, to contact sodium hypochlorite (bleach) solution. These mixtures can produce a highly toxic gas.

When disposing of used COBAS AmpliPrep Sample Processing Units (SPUs), which contain guanidine thiocyanate, avoid any contact with sodium hypochlorite (bleach) solution. These mixtures can produce a highly toxic gas

Risk is minimal due to the small quantity of chemicals, the safety of packaging and the limited handling by the operators using the test kits.

3. COMPUTERIZATION; DATA MANAGEMENT SYSTEM

Data are stored in an Excel spreadsheet on the CDC LAN. Include with every run the HCV Negative control, HCV Low Positive control and HCV High Positive control as a part of the Quality Control.

For control orders, IU/mL value for the control must be within a specified range for the
Hepatitis C virus RNA in Serum
NHANES 2013-2014

run to be valid. After interpretation, format for reporting the results for controls is
Quantitative HCV RNA values in IU/mL and appropriate comments. Reporting is done by
manually updating previous reports sent to NCHS with an additional line for HCV RNA.
Electronically stored data are backed up routinely.

Finished data are reviewed by the laboratory supervisor and transmitted to the NCHS
along with other NHANES IV data.

Files stored on the CDC Local Area Network (LAN) are automatically backed up nightly.

Documentation for data system maintenance is maintained with printed copies of data
records for 2 years.

4. SPECIMEN COLLECTION, STORAGE, AND HANDLING PROCEDURES; CRITERIA
FOR SPECIMEN REJECTION

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

No special patient preparation before collection is necessary.

Blood should be collected in sterile collection tubes using red tops.

Collect one red top tube using standard venipuncture techniques. Follow the
manufacturer’s instructions for use of the collection tubes.

Separate serum from whole blood within 6 hours of collection by centrifugation at 1500
x g for 20 minutes at room temperature.

Transfer serum to a properly identified, sterile, screw-cap, polypropylene tube after
centrifugation.

Serum specimens may be stored at 2-8°C for up to 72 hours or frozen at -70°C or
colder for up to 6 weeks. Serum specimens may be frozen and thawed up to five times
without a loss of HCV RNA.

Specimens and controls should be handled as if infectious using safe laboratory
procedures such as those outlined in *Biosafety in Microbiological and Biomedical
Laboratories* and in the CLSI Document M29-A. Thoroughly clean and disinfect all work
surfaces with a freshly prepared solution of 0.5% sodium hypochlorite in deionized or
distilled water.

5. PROCEDURES FOR MICROSCOPIC EXAMINATIONS; CRITERIA FOR REJECTION
OF INADEQUATELY PREPARED SLIDES

Not applicable for this procedure.
6. PREPARATION OF REAGENTS, CALIBRATORS (STANDARDS), CONTROLS, AND ALL OTHER MATERIALS; EQUIPMENT AND INSTRUMENTATION

a. Instrumentation and Software

COBAS AmpliPrep Instrument

COBAS AmpliPrep TaqMan 48 Analyzer

AMPLILINK Software, Version 3.3.6 Series

Data Station for the AMPLILINK software, with printer

Multichannel pipettor (capacity 25 and 100 μL)

Aerosol barrier or positive displacement RNase-free tips (25 and 100 μL) and barrier-free tips

b. Reagents

Reagent Labeling and Preparation

All reagents are labeled by the manufacturer. The labeling includes contents, lot number, expiration date, and storage instructions.

All reagents are liquid, ready-to-use.

Reagent Storage and Use

Do not freeze reagents or controls.

Store HCV CS1, HCV CS2, HCV CS3 and HCV CS4 at 2-8ºC. Unused, these reagents are stable until the expiration date indicated. Once used, these reagents are stable for 28 days at 2-8ºC or until the expiration date, whichever comes first. HCV CS1, HCV CS2, HCV CS3 and HCV CS4 can be used for a maximum of 4 instrument cycles, up to a maximum of 64 hours cumulative on board the COBAS AmpliPrep Instrument. Reagents must be stored at 2-8°C between instrument cycles.

Store HCV H(+)+C, HCV L(+)C and CTM (−) C at 2-8°C. The controls are stable until the expiration date indicated. Once opened, any unused portion must be discarded.

Store Barcode clips [HCV H(+)C Clip, HCV L(+)C Clip and HCV (−) C Clip] at 2-30°C.

Store PG WR at 2-30°C. PG WR is stable until the expiration date indicated. Once opened, this reagent is stable for 28 days at 2-30°C or until the expiration date, whichever comes first.

Do not pool reagents from different lots or from different bottles of the same lot. Do not mix reagent cassettes or controls from different kits. Do not open COBAS AmpliPrep cassettes and exchange, mix, remove or add bottles. Do not pool reagents. Dispose of
unused reagents and waste in accordance with all local, country, state, and federal regulations.

Handle all reagents with caution and avoid contact with skin, eyes, or mouth. Refer to the package insert for any known toxicity.

HCV QS, CAP/CTM Mn\(^{2+}\) and HCV MMX contain sodium azide. Sodium azide may react with lead and copper plumbing to form highly explosive metal azides. While disposing of sodium azide containing solutions down laboratory sinks, flush the drains with a large volume of water to prevent azide buildup.

Do not allow HCV CS2 and liquid waste from the COBAS AmpliPrep Instrument, which contain guanidine thiocyanate, to contact sodium hypochlorite (bleach) solution. These mixtures can produce a highly toxic gas.

When disposing of used COBAS AmpliPrep Sample Processing Units (SPUs), which contain guanidine thiocyanate, avoid any contact with sodium hypochlorite (bleach) solution. These mixtures can produce a highly toxic gas.

These reagents carry a highly flammable warning: HCV CS1

These reagents are marked as harmful or irritants: HCV CS1, HCV CS2, and HCV CS3.

7. CALIBRATION AND CALIBRATION VERIFICATION PROCEDURES

a. Calibration Curve

No calibration curve is produced.

**AMPLILINK Software:**

Determines the Cycle Threshold value (Ct) for the HCV RNA and the HCV QS RNA.

Determines the HCV RNA concentration based upon the Ct values for the HCV RNA and HCV QS RNA and the lot-specific calibration coefficients provided on the cassette barcodes.

Determines that the calculated IU/mL for HCV L(+)C and HCV H(+)C fall within the lot specific assigned ranges encoded on the COBAS AmpliPrep/COBAS TaqMan HCV Test reagent cassette barcodes supplied with the kit.

b. Verification

Not Applicable
8. PROCEDURE OPERATING INSTRUCTIONS; CALCULATIONS; INTERPRETATION OF RESULTS

a. Preliminaries

Sample Preparation
If using frozen specimens, place the specimens at room temperature until completely thawed and vortex for 3-5 seconds before use. Controls should be removed from 2-8°C storage and equilibrated to ambient temperature before use.

b. Instrument Setup

Operation of Assay Procedure

Run Size and Workflow
Each kit contains reagents sufficient for 48 tests, which may be performed in batches of 12 to 24 tests. At least one replicate each of CTM (-) C, HCV L(+)C and HCV H(+)C must be included in each batch (see “Quality Control” section).

Workflow
The COBAS TaqMan 48 Analyzer run must be started within 120 minutes following completion of specimen and control preparation.

Note: Do not freeze or store processed specimens and controls at 2-8°C.

Part A. Maintenance and Priming

A1. The COBAS AmpliPrep Instrument is ready for operation in stand-by mode.

A2. Turn the Data Station for the AMPLILINK software ON. Prepare the Data Station as follows:
   a. Log onto Windows XP.
   b. Double click the AMPLILINK software icon.
   c. Log onto AMPLILINK software by entering the assigned User ID and password.

A3. Check the supply of PG WR using the Status Screen and replace if necessary.

A4. Perform all Maintenance that is listed in the Due Tab. The COBAS AmpliPrep Instrument will automatically prime the system.

Part B. Loading of Reagent Cassettes

Note: All reagent cassettes should be removed from 2-8°C storage, immediately loaded onto the COBAS AmpliPrep Instrument and allowed to equilibrate to ambient temperature on the instrument for at least 30 minutes before the first specimen is to be processed. Do not let reagent cassettes come to ambient temperature outside the instrument as condensation may form on the barcode.
Hepatitis C virus RNA in Serum
NHANES 2013-2014

labels. Do not wipe off condensation if it appears on the barcode labels.

B1. Place HCV CS1 onto a reagent rack. Place HCV CS2, HCV CS3 and HCV CS4 onto a separate reagent rack.

B2. Load the reagent rack containing HCV CS1 onto rack position A of the COBAS® AmpliPrep Instrument.

B3. Load the reagent rack containing HCV CS2, HCV CS3 and HCV CS4 onto rack position B, C, D or E of the COBAS® AmpliPrep Instrument (see Table 1 for additional information).

Part C. Loading of Disposables

Note: Determine the number of COBAS® AmpliPrep reagent cassettes, Sample Processing Units (SPUs), Input Sample tubes (S-tubes), K-tips and K-tubes needed. One SPU, one Input S-tube, one K-tip and one K-tube are needed for each specimen or control.

Multiple configurations for use of the COBAS AmpliPrep Instrument with the COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer are possible. For reference, see Table 1 below. Depending on the configuration used, load the appropriate number of reagent cassette racks, sample racks with Input S-tubes, SPU racks, K-tip racks, K-tube racks and K-carriers on K-carrier racks onto the respective rack positions of the COBAS AmpliPrep Instrument (see Table 1 for additional information).

C1. Place the SPUs in the SPU rack(s) and load the rack(s) onto rack position J, K or L of the COBAS® AmpliPrep Instrument.

C2. Depending on the configuration used, load full K-tube rack(s) onto rack position M, N, O or P of the COBAS® AmpliPrep Instrument.

C3. Load full K-tip rack(s) onto rack position M, N, O or P of the COBAS® AmpliPrep Instrument.

C4. For configurations 3 to 5 using the COBAS® TaqMan® 48 Analyzer, load K-carriers on K-carrier rack(s) onto rack position M, N, O or P of the COBAS® AmpliPrep Instrument.
### Table 1 Possible Configurations for using COBAS AmpliPrep Instrument with COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer

<table>
<thead>
<tr>
<th>Configuration</th>
<th>Transfer Mode to COBAS TaqMan Analyzer or COBAS TaqMan® 48 Analyzer</th>
<th>Racks, Carriers, and Disposables</th>
<th>Position on COBAS AmpliPrep Instrument</th>
</tr>
</thead>
<tbody>
<tr>
<td>3, 4</td>
<td>Manual transfer of K-carrier via K-carrier rack(s) onto COBAS TaqMan 48 Analyzer</td>
<td>K-tubes are placed on the sample rack</td>
<td>F-H</td>
</tr>
<tr>
<td></td>
<td></td>
<td>K-tips in full K-tip racks</td>
<td>M-P</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Input S-tubes on sample racks</td>
<td>F-H</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SPU in SPU rack</td>
<td>J-L</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CS1 on Cassette rack</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CS2, CS3, CS4 on Cassette rack</td>
<td>B-E</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Empty barcoded K-carrier on K-carrier rack</td>
<td>M-P</td>
</tr>
<tr>
<td></td>
<td></td>
<td>After specimen processing is finished: K-tubes in K-carrier on K-carrier rack</td>
<td>Same as above (M-P)</td>
</tr>
</tbody>
</table>
Part D. Ordering and Loading of Specimens

D1. Prepare sample racks as follows: Attach a barcode label clip to each sample rack position where a specimen (S-tube) is to be placed. Attach one of the specific barcode label clips for the controls [CTM (–) C, HCV L(+)C and HCV H(+)C] to each sample rack position where the controls (S-tube) are to be placed. The barcode label clips for controls should have the same control lot number as the lot number on the control vials in the kit. Take care in assigning the right control to the position with the appropriate control barcode clip. Place one Input S-tube into each position containing a barcode label clip.

D2. Using the AMPLILINK software, create specimen orders for each specimen and control in the Orders window Sample folder. Select the appropriate test file and complete by saving.

D3. Assign specimen and control orders to sample rack positions in the Orders window Sample Rack folder. The sample rack number must be for the rack prepared in Step D1.

D4. Print the Sample Rack Order report to use as a worksheet.

D5. Prepare specimen and control racks in the designated area for specimen and control addition as follows: Vortex each specimen and control [CTM (–) C, HCV L(+)C and HCV H(+)C] for 3 to 5 seconds. Avoid contaminating gloves when manipulating the specimens and controls.

D6. Transfer 1000 to 1050 μL of each specimen and control [CTM (–) C, HCV L(+)C and HCV H(+)C] to the appropriate barcode labeled Input S-tube using a micropipettor with an aerosol barrier or positive displacement RNase-free tip. Avoid transferring particulates and/or fibrin clots from the original specimen to the Input S-tube. Specimens and controls should be transferred to tube positions as assigned and recorded on the worksheet in step D4. The barcode label clips for controls should have the same control lot number as the lot number on the control vials in the kit. Assign the right control to the position with the appropriate control barcode clip. Avoid contaminating the upper part of the S-tubes with specimens or controls.

D7. For configurations 1 and 2, load the sample rack(s) filled with Input S-tubes onto rack positions F, G or H of the COBAS AmpliPrep Instrument.

D8. For configurations 3 to 5 using the COBAS TaqMan 48 Analyzer, load sample rack(s) with Input S-tubes and K-tubes (one for each Input S-tube, loaded in the right position adjacent to Input S-tubes) onto rack position F, G or H of the COBAS AmpliPrep Instrument.

Part E. Start of COBAS AmpliPrep Instrument Run

E1. Start the COBAS AmpliPrep Instrument using the AMPLILINK software.
Part F. End of COBAS AmpliPrep Instrument Run and Transfer to COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer (Only for Configurations 2–5)

F1. Check for flags or error messages in the system screen.

F2. Remove processed specimens and controls from the COBAS AmpliPrep Instrument on either sample racks (for COBAS TaqMan Analyzer without Docking Station) or K-carrier racks (for COBAS TaqMan 48 Analyzer), depending on the configuration (for further details see Part G).


*Note: All processed specimens and controls should not be exposed to light after completion of specimen and control preparation.*
Hepatitis C virus RNA in Serum
NHANES 2013-2014

Amplification and Detection

**COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer Set-up**

The COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer run must be started within 120 minutes following completion of specimen and control preparation.  

*Note: Do not freeze or store processed specimens and controls at 2-8°C.*

Part G. Loading Processed Specimens

G1. Depending on the instrument configuration, perform the appropriate steps to transfer the K-tubes to the COBAS TaqMan Analyzer or COBAS TaqMan 48 Analyzer:

- **Configuration 1:** Automated transfer of K-carrier via docking station to COBAS TaqMan Analyzer. Manual intervention is unnecessary.
- **Configuration 2 and 5:** Manual transfer of K-tubes in sample rack(s) to COBAS TaqMan Analyzer
- **Configuration 3, 4 and 5:** Manual transfer of K-carrier on K-carrier rack(s) to the COBAS TaqMan 48 Analyzer. Manual transfer of K-carriers into COBAS TaqMan 48 Analyzer using the K-carrier Transporter.

Part H. Start of COBAS TaqMan 48 Analyzer Run

H1. Start the COBAS TaqMan 48 Analyzer by one of the options below depending on the configuration used:

- **Configuration 1:** No intervention necessary.
- **Configuration 2 and 5:** Automatic start of the COBAS TaqMan Analyzer after insertion of sample rack(s).
- **Configuration 3, 4 and 5:** Fill K-carrier with empty K-tubes if fewer than 6 K-tubes on the K-carrier. Filling is guided by the AMPLILINK software. Open thermal cycler cover, load K-carrier into thermal cycler and close lid. Start the COBAS TaqMan 48 Analyzer run.

Part I. End of COBAS TaqMan 48 Analyzer Run

I1. At the completion of the COBAS TaqMan 48 Analyzer run, print Results Report. Check for flags or error messages in the Result section. Specimens with flags and comments are interpreted as described in the Results section. After acceptance, store data in archive.

I2. Remove used K-tubes from the COBAS TaqMan 48 Analyzer.
**Hepatitis C virus RNA in Serum**

**NHANES 2013-2014**

c. **Reporting results**

The COBAS TaqMan 48 Analyzer automatically determines the HCV RNA concentration for the specimens and controls. The HCV RNA concentration is expressed in International Units (IU)/mL.

**AMPLILINK Software:**
- Determines the Cycle Threshold value (Ct) for the HCV RNA and the HCV QS RNA.
- Determines the HCV RNA concentration based upon the Ct values for the HCV RNA and HCV QS RNA and the lot-specific calibration coefficients provided on the cassette barcodes.
- Determines that the calculated IU/mL for **HCV L(+)C** and **HCV H(+)C** fall within the lot specific assigned ranges encoded on the COBAS AmpliPrep/COBAS TaqMan HCV Test reagent cassette barcodes supplied with the kit.

Results are interpreted as follows:

<table>
<thead>
<tr>
<th>Titer Result</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target Not Detected</td>
<td>No Ct value for HCV obtained. Report results as &quot;HCV RNA not detected&quot;.</td>
</tr>
<tr>
<td>&lt;4.30E+01 IU/mL</td>
<td>Below 4.30E+01 IU/mL (lower limit of quantitation, LLoQ); HCV RNA is not quantifiable.</td>
</tr>
<tr>
<td>&gt;4.30E+01 IU/mL and &lt;6.90E+07 IU/mL</td>
<td>Results greater than or equal to 43 IU/mL and less than or equal to 6.90E+07 IU/mL are within the Linear Range of the assay.</td>
</tr>
<tr>
<td>&gt; 6.90E+07 IU/mL</td>
<td>Results are above the range of the assay. Report results as &quot;greater than 6.90E+07 HCV RNA IU/mL&quot;. If quantitative results are desired, the original specimen should be diluted 1:100 with HCV-negative human serum or EDTA plasma, depending on the matrix of the original specimen, and the test repeated. Multiply the reported result by the dilution factor.</td>
</tr>
</tbody>
</table>

d. **Recording of Data**

The Analyzer automatically determines the HCV RNA titer for the sample or control.

If **Negative Control, HCV Low Positive Control or HCV High Positive Control** is invalid then the entire run is invalid, repeat the entire run including control and sample preparation, reverse transcription, amplification and detection.
Hepatitis C virus RNA in Serum
NHANES 2013-2014

A valid run may include both valid and invalid sample results depending on whether
flags and/or comments are obtained for the individual samples.

e. Calculations

Not Applicable

f. Special Procedure Notes

Workflow in the laboratory must proceed unidirectional. It must begin in the Reagent
Preparation area, move to the Specimen Preparation area, and then move to the
Amplification/Detection area. Reagent preparation and specimen preparation are
performed in separate, segregated areas.

Supplies and equipment must be dedicated to each activity and not used for other
activities or moved between areas.

Gloves must be worn in each area and must be removed before leaving that area.
Supplies, equipment, and gloves used for the preparation activities must not be used
in the Amplification/Detection activities. Any amplification and detection supplies and
equipment must remain in that area at all times. All pipettors, pipettes, bulbs, pipette
tips, etc. must be dedicated to, and used only for, its individual PCR activity. It must
not be used for any non-PCR activity.

9. REPORTABLE RANGE OF RESULTS

Calculated results greater than or equal to 4.30 E+01 IU/mL and less than or equal to 6.90
E+07 IU/mL are within the linear range of the assay. For NHANES results are reported as
positive or negative (i.e., no Ct value for HCV obtained); quantitative values are not
reported.

10. QUALITY CONTROL (QC) PROCEDURES

One replicate each of the COBAS TaqMan Negative Control, the HCV Low Positive
Control and the HCV High Positive Control must be included in each test batch. The batch
is valid if no flags appear for any of the controls [HCV L(+), HCV H(+), and CTM (−)].
Based on results of a carry-over contamination study with alternating high positive HCV
samples and HCV negative samples, there are no requirements regarding the position of
the controls on the sample rack.
Check the batch printout for flags and comments to ensure that the batch is valid.

Negative Control
The CTM (−) C must yield a "Target Not Detected" result. If the CTM (−) C is flagged as
invalid, then the entire batch is invalid. Repeat the entire process (specimen and control
preparation, amplification and detection). If CTM (−) C is consistently invalid in multiple
Hepatitis C virus RNA in Serum
NHANES 2013-2014

Positive Controls
The assigned range for **HCV L(+)C** and **HCV H(+)C** is specific for each lot of reagents, and
is provided on the COBAS AmpliPrep/COBAS TaqMan HCV Test reagent cassette barcodes.

The HCV RNA IU/mL for **HCV L(+)C** and **HCV H(+)C** should fall within their assigned ranges. If one or both of the positive controls are flagged as invalid, then the entire batch is invalid. Repeat the entire process (specimen and control preparation, amplification and detection). If the HCV RNA titer of one or both of the positive controls is consistently outside the assigned ranges in multiple batches, contact your local Roche office for technical assistance.

Specimens and controls from separate specimen preparation batches may be amplified and detected at the same time. However, each separate specimen batch is validated individually by the set of controls included with the batch. Therefore, it is possible to reject one batch of specimens from a common amplification and/or detection run while accepting another batch upon the performance of the controls processed with those specimens.

All test specimens and controls prepared in the same batch should be amplified and detected in adjacent positions in the thermal cycler and on the detection plate. The exact order or placement of these specimens and controls in the thermal cycler or detection plate is not critical. Additional controls may be tested according to guidelines or requirements of local, state and/or federal regulations or accrediting organizations.

Control Material Preparation and Storage
Three controls are provided for use: HCV L(+)C, HCV H(+)C and CTM (–) C. The controls are liquid, ready-to-use. Store the controls at 2–8°C. The products are stable until the expiration date.

11. REMEDIAL ACTION IF CALIBRATION OR QC SYSTEMS FAIL TO MEET ACCEPTABLE CRITERIA

The entire run is considered to be invalid if one or more controls are not within specified limits.

Repeat the entire test process: specimen and control preparation, reverse transcription, amplification and detection.

*If one or more controls are consistently outside the specified limits, contact the Roche Response Center for technical assistance.*

Specimen Processing Lysis Control: Since the positive controls do not control for the lysis portion of Specimen Preparation, the user may consider a well-characterized, HCV RNA-positive specimen that is available in sufficient quantity to be included as an external control for the entire procedure. Additional external controls may be tested.
12. LIMITATIONS OF METHOD; INTERFERING SUBSTANCES AND CONDITIONS

Interfering Substances

Interfering substances include but are not limited to the following:
- Elevated levels of triglycerides, bilirubin, albumin, hemoglobin and human DNA in specimens as well as the presence of autoimmune diseases such as Systemic Lupus Erythematosus (SLE), Rheumatoid Arthritis (RA), and Antinuclear Antibody (ANA) have been shown not to interfere with the quantitation of HCV RNA by the COBAS AmpliPrep/COBAS TaqMan HCV Test.
- The following drug compounds tested at the Peak Plasma Level (Cmax) and at 3 times the Cmax have been shown not to interfere with the quantitation of HCV RNA by the COBAS AmpliPrep/COBAS TaqMan HCV Test:
  - **HIV Protease Inhibitors**: Indinavir Saquinavir Ritonavir Nelfinavir Amprenavir Lopinavir/Ritonavir
  - **Nucleoside HIV Reverse Transcriptase Inhibitors and DNA Polymerase Inhibitors**: Lamivudine Zidovudine Stavudine Abacavir Didanosine
  - **Non-nucleoside HIV Reverse Transcriptase Inhibitors**: Nevirapine Efavirenz
  - **HIV Fusion Inhibitors**: Enfurvitide
  - **Immune Modulators**: Interferon alfa-2a Interferon alfa-2b Peginterferon alfa-2a Peginterferon alfa-2a + Ribavirin Interferon alfa-2b+ Ribavirin
  - **Antidepressants**: Paroxetine HCl Fluoxetine Sertraline
  - **Compounds for Treatment of Herpes Viruses**: Ganciclovir Valganciclovir Acyclovir

Limitations of the Method

As with any test procedure, good laboratory technique is essential to the proper performance of this assay.

Procedural Limitations

This test has been validated for use with only human serum or plasma collected in EDTA anticoagulant. Testing of other specimen types may result in inaccurate results.

Quantitation of HCV RNA is dependent on the number of virus particles present in the specimen and may be affected by specimen collection methods, patient factors (e.g., age, presence of symptoms) and stage of infection.

Though rare, mutations within the highly conserved regions of the viral genome covered by the COBAS AmpliPrep/COBAS TaqMan HCV Test primers and/or probe may result in the under-quantitation of or failure to detect the presence of the virus in this circumstance.

Reliable results are dependent on adequate specimen collection, transport, storage and processing procedures.
The presence of AmpErase enzyme in the COBAS AmpliPrep/COBAS TaqMan HCV Master Mix reduces the risk of amplicon contamination. However, contamination from HCV positive controls and clinical specimens can be avoided only by good laboratory practices and careful adherence to the procedures specified in this Package Insert.

Use of this product should be limited to personnel trained in the techniques of PCR.

This product can only be used with the COBAS AmpliPrep Instrument and the COBAS® TaqMan Analyzer or COBAS TaqMan 48 Analyzer.

If another assay was initially used for quantitation of HCV viral RNA in order to assess treatment effect on the patient, it is recommended that prior to switching to the COBAS AmpliPrep/COBAS TaqMan HCV Test users perform method correlation studies in their laboratory to quantify assay differences.

13. REFERENCE RANGES (NORMAL VALUES)

All normal, noninfected humans should have negative values for HCV RNA.

14. CRITICAL CALL RESULTS ("PANIC VALUES”)

Not applicable.

15. SPECIMEN STORAGE AND HANDLING DURING TESTING

Specimens may remain at 20-25°C during preparation and testing only.

16. ALTERNATIVE METHODS FOR PERFORMING TEST OR STORING SPECIMENS IF TEST SYSTEM FAILS

Other FDA-licensed tests for HCV RNA may be substituted but must be accompanied by validation data to show substantial equivalence with this assay. Substitution of test methods may not be done without approval from the NCHS.

Alternate storage is not recommended.

17. TEST RESULT REPORTING SYSTEM; PROTOCOL FOR REPORTING CRITICAL CALLS (IF APPLICABLE)

Not applicable.

18. TRANSFER OR REFERRAL OF SPECIMENS; PROCEDURES FOR SPECIMEN ACCOUNTABILITY AND TRACKING

22
Hepatitis C virus RNA in Serum 
NHANES 2013-2014

Test results are documented through the lab management database (Section 3) for specimen tracking.

Specimens in long-term storage are arranged by study group. The storage location of each sample is listed with the test data. For NHANES, residual specimens are stored frozen and returned to the NCHS specimen bank after testing for each cycle has been completed.

REFERENCES


Hepatitis C virus RNA in Serum
NHANES 2013-2014