

Amplisure[®] HBV Quantitative Kit (Real Time PCR Kit)

IVD

Product Insert



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Index

	Page No.
Introduction	4
Product Description	4
Recommended Work Areas	4
General Precautions	5
Precautions while extracting Nucleic acid	5
Precautions while setting up a PCR reaction	5
Precautions for post PCR or equipment area/room	5
Precautions after completion of Real time PCR assay	5
Usage Limitations	6
Safety Precautions	6
Storage Conditions and Product Stability	6
Symbols	7
Kit Components	8
Materials required but not supplied	8
Quality Systems	9
Sample Type/Collection/Storage/Transport	9
Sample Type	9
Sample Collection, Storage and Transport	9
Assay Procedure	10
DNA Extraction	10
Use of Internal Control (IC)	10
Usage of Internal Control at the DNA extraction step	11
Usage of Internal Control at the Real time PCR step	11
qPCR Protocol	11
Preparation of Reaction Mastermix	11
PCR Programming	13
Plate Setup	13
Cycling conditions	14
Data Analysis	14
Setting the threshold for the qPCR Data analysis	14
Results	15
Interpretation	15
Troubleshoot	16

Assay Characteristics	17
Analytical Sensitivity	17
Analytical Sensitivity of qPCR assay	17
Analytical Sensitivity in consideration of Extraction Method	17
Linear Range	18
Conversion Factor (IU/ μ L to copies/ μ L)	18
Specificity	19
Detection of Different HBV genotypes using International Standard	20
Cross Reactivity Data	20
Precision	21
Intra-assay variability	21
Inter-assay variability	22
Abbreviations	23
References	24
Ordering Information	24

Introduction

Hepatitis B virus (HBV) is an enveloped DNA virus that belongs to the family Hepadnaviridae. Approximately 350 million people are chronically infected with HBV worldwide, placing them at high risk for developing cirrhosis, end-stage liver disease, and hepatocellular carcinoma. Detection and quantification of circulating HBV in plasma or serum play an important role in diagnosing and monitoring HBV infection as well as assessing response to therapy. Clinically, HBV DNA titers vary greatly, from levels as high as 10^{10} copies/mL during acute HBV infection to very low levels in HBe antigen- negative chronic carriers, in patients undergoing antiviral therapy and in those with occult HBV infection. Among the methods most commonly used for HBV DNA quantification, assays based on real-time PCR technology offer the great sensitivity and broad linear dynamic range.

Product Description

Amplisure[®] HBV Quantitative Kit is an *in-vitro* diagnostic kit for quantitation of Hepatitis B Virus in human plasma. The kit contains the necessary reagents for performing HBV (A-H genotypes) quantitation by Real – time PCR.

Pathogen detection by Real Time polymerase chain reaction (PCR) is based on the amplification of specific region (S gene) of the pathogen genome. The assay principle is based on Taqman probes which allow higher specificity and sensitivity.

In addition, the Amplisure[®] HBV Quantitative Kit contains a second amplification system to identify possible PCR inhibition by using an internal control (IC) without affecting the analytical sensitivity of the assay. External quantitation controls calibrated with WHO control are supplied, which allow the determination of the amount of Hepatitis B viral DNA.

Recommended Work areas

Molecular Diagnostics work area includes:

- a) Sample preparation area/room – for extraction of nucleic acids from clinical samples
- b) Pre-PCR area/room - for setting up PCR reaction
- c) PCR area/room – for performing PCR using the thermocyclers

As part of Good Laboratory Practices (GLP), it is recommended to have dedicated areas to avoid cross contamination.

General Precautions

Precautions while extracting Nucleic acid

Always wear proper attire (powder free gloves, facemask and Head cap) before starting the nucleic acid extraction procedure. During preparation of samples, compliance with good laboratory practices are essential to minimize the risk of cross-contamination between samples, and the inadvertent introduction of ribonucleases (RNases) into samples during and after the extraction procedure. Proper aseptic technique should always be used when working with RNA.

The Sample Preparation Area is dedicated to processing samples. All reagents used in the Sample Preparation Area should remain in this dedicated area at all times. Laboratory coats, pipettes, pipette tips and vortex mixer used in the Sample Preparation Area must remain in this area and not be moved to the Pre-PCR/PCR area. Discard the gloves before leaving this area. Do not bring amplified product into the Sample Preparation Area. Usage of filter tips is recommended while sample preparation and should be performed in a Biosafety cabinet.

Precautions while setting up a PCR reaction

PCR assay is sensitive and any accidental introduction of product from previous amplification reactions leads to incorrect results. Hence measures to reduce the risk of contamination in the laboratory should include physically separating the activities involved in performing PCR and complying with good laboratory practices.

It is recommended to have proper cleaning procedures to minimize the risk of cross contamination and carry over contamination (e.g. DNA OUT™, RNase OUT™, 0.1% Sodium Hypochlorite, Fumigation etc.).

It is recommended that areas should be defined in Pre-PCR room for preparation of mastermix and addition of templates. Laboratory coats and equipment used in the Pre-PCR Area must remain in this area and should not be moved to the Sample Preparation Area.

Precautions for post PCR or equipment area/room

The Real time PCR instrument/s should be kept in a separate segregated area away from Sample preparation area and Pre-PCR area.

Precautions after completion of Real time PCR assay

The reaction tubes or strips should be properly discarded without opening the caps, after the completion of run to avoid carry over contamination.

Usage Limitations

1. The kit and all its components are for *in-vitro* diagnostics only.
2. The product is to be used by personnel specially trained in the *in-vitro* diagnostics procedures only.
3. Follow the product insert strictly for optimal PCR results.
4. Do not use the kit beyond the expiry date mentioned on the kit box.
5. Follow the guidelines provided in product insert for sample collection, storage and transport.
6. For ideal performance, store the kit under recommended conditions only.











Safety Precautions

1. All patient specimens should be considered as potentially infectious and handled in a BSL2 biosafety hood with BSL3 practices.
2. Wear personal protective equipment, including gloves, head cap, face mask and lab coats when handling kit reagents/sample extraction. Wash hands thoroughly using detergents before and after performing the test.
3. Do not smoke, drink or eat in areas where kit reagents and/or human specimens are being used.
4. Dispose of unused kit reagents and human specimens as per regulatory guidelines.

Storage Conditions and Product Stability

1. All the kit reagents should be stored at -20°C. Replace all the kit components immediately at -20°C after usage.
2. Repeated thawing and freezing (more than 6 x) of all kit reagents should be avoided, as it reduces assay sensitivity. If needed, make aliquots of the kit reagents according to the volume used in the protocol prior to freezing.
3. Allow reagents to be thawed completely on Ice/4°C prior to use.
4. Kit reagents are stable through the end of the expiration date indicated on the box when stored at -20°C.

Symbols

Description of Symbol	Denotation
	<i>in-vitro</i> Diagnostic medical device
	Consult Instruction manual (Product Insert) for use
	Lot Number of the kit or Kit contents
	Catalogue number of Kit
	Contains sufficient for <N> reactions (Pack Size)
	Manufacturer
	Temperature limitation (Storage Condition)
	Use by MMM-YYYY (Expiry Date)
	Biological risk (handle carefully)
	Important Note

Kit Components

Color Coding (Caps)	Contents	Description	50 rxns (QT-HBV-50)	100 rxns (QT-HBV-100) 2 x 50 rxns
Yellow	RAS qDNA PCR Mix	DNA Amplification Reagent	1 x 750µL	2 x 750µL
Brown	RAS HBV PPM	HBV Primer-Probe Mix	1 x 110 µL	2 x 110 µL
Lavender	RAS HBQS1 (2 x 10 ⁴ IU/ µL)	HBV Quantitation Standards	1 x 60 µL	2 x 60 µL
Lavender	RAS HBQS2 (2 x 10 ³ IU/ µL)		1 x 60 µL	2 x 60 µL
Lavender	RAS HBQS3 (2 x 10 ² IU/ µL)		1 x 60 µL	2 x 60 µL
Lavender	RAS HBQS4 (2 x 10 ¹ IU/ µL)		1 x 60 µL	2 x 60 µL
Natural	RAS IC PCR Mix	Internal Control	1 x 50 µL	2 x 50 µL
Natural	RAS D-IC EX Mix		1 x 500 µL	2 x 500 µL
White	MBGW	Molecular Biology Grade water	1 x 1.00 mL	2 x 1.00 mL

Materials required but not supplied

The materials which are required but not supplied are listed below:

1. Viral DNA Extraction kit
2. Biosafety Cabinet
3. PCR Hood
4. Calibrated variable micropipettes
5. Sterile pipette filter tips (aerosol free)
6. Vortex mixer
7. Dry Bath
8. Benchtop centrifuge with rotor for 1.5 mL reaction tubes
9. Real Time PCR machine
10. Strip Tubes and Caps (0.2 mL) or PCR Tubes (0.2 mL) or 96 well plate
11. Cooling block (96 x 0.2 mL tubes)

12. 1.5 mL centrifuge tubes
13. 1.5 mL centrifuge tube stand
14. Cooling block (2 mL tubes)
15. Sterile powder free gloves
16. Facemask
17. Head cap
18. Lab coats

Quality Systems

In accordance with ISO-certified Quality Management System (9001:2008 and 13485: 2003) of RAS Lifesciences, each lot of Amplisure[®] HBV Quantitative PCR Kit is tested against predetermined specifications to ensure consistent product quality.

Sample Type/Collection/Storage/Transport

Sample Type

Plasma (K₂EDTA-Blood)

Heparinized Blood must not be used as they inhibit the PCR reaction

Sample Collection, Storage and Transport

3-5 mL of blood has to be drawn into a K₂EDTA vacutainer. Cap and swirl the tubes for uniform mixing of blood and K₂EDTA. When K₂EDTA is used, whole blood can be collected in tubes with or without a gel separator. The K₂EDTA blood samples are centrifuged (20 minutes at 1000–1500 x *g*) to separate plasma from cellular material and in the case of a non gel separator tube, the plasma should be removed to a secondary sterile tube within 4 hours of phlebotomy.

Collection and storage of unstabilized whole blood is not recommended for PCR analysis, because RNA degradation occurs in blood stored *ex vivo*. The sensitivity of the assay can be reduced if whole blood samples are frozen or stored for a longer period of time.

Plasma separated in a gel separator tube may be transported to the laboratory *in situ*. Plasma should be shipped at 2 to 8 °C and stored at -20°C as it is stable for up to five days at 2 to 8 °C and longer if frozen at -20°C or -70°C or lower. Do not store plasma samples in a “frost -free” freezer as the temperature is cycled several times per day on this type of freezer, causing degradation of nucleic acid targets.

Sample material should be transported in a leak proof, unbreakable transport container to avoid leakage of sample. The samples should be transported following the local and national instructions for the transport of pathogen material.

Assay Procedure

DNA Extraction

Amplisure® HBV Quantitative Kit has been validated using the following Viral DNA extraction kits:

- 1- Roche High Pure Viral Nucleic Acid kit (Cat. No. 11858874001)
- 2- QIAamp DNA Blood Mini Kit (Cat. No. 51104)

Follow the manufacturer's instructions mentioned in the manual for Viral DNA extraction. Different pack sizes of the above mentioned kits can be used. However the customer can also validate their own extraction process using other Viral DNA extraction Kits.

Recommended sample volume for extraction and elution are as follows:

Sl. No.	Name of the DNA Isolation Kit	Recommended Sample volume (<i>to be taken for DNA Extraction</i>)	Recommended Final Elution volume
1.	Roche High Pure Viral Nucleic Acid kit (Cat. No. 11858874001)	200 µL	100 µL
2.	QIAamp DNA Blood Mini Kit (Cat. No. 51104)	200 µL	100 µL

The analytical sensitivity of the assay in consideration of the purification was determined using the above defined volumes.

Use of Internal Control (IC)

Internal controls are supplied (RAS D-IC Ex Mix and RAS IC PCR Mix) along with Amplisure® HBV Quantitative Kit. This allows the user to control the Viral DNA isolation procedure as well as to check for possible PCR inhibition.



Internal control should only be used once, either at the Extraction step or at the PCR step

Usage of Internal Control at the DNA extraction step

If internal control (IC) is required to be added at the time of DNA extraction, add 20 µL of RAS *D-IC Ex mix* per isolation to the lysis buffer along with other components of kit used for lysis (as per kit instructions) and vortex for 5 seconds prior to usage.

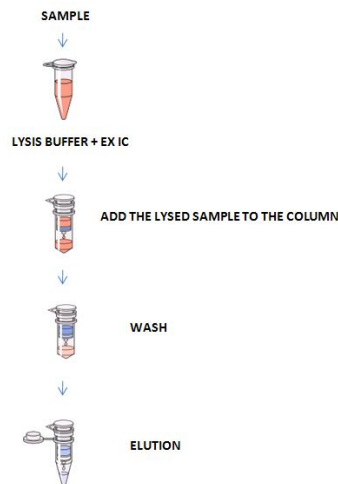


Fig1 Viral DNA Extraction Overview

Usage of Internal Control at the Real time PCR step

The internal control can optionally be used exclusively to check for possible PCR inhibition. For this application, add the internal control directly to the PCR master mix as described on Pg. No 15. Addition of Internal control should be done only once, either at the time of sample extraction or during PCR setup.

qPCR Protocol

Preparation of Reaction Mastermix

Quantitation procedure with Amplisure® HBV Quantitative Kit involves *1 step RT qPCR*. It is recommended that a minimum of three standards and a negative control (MBGW should be used as negative control) are required to be included in a single run for acquiring proper results.

Set up a real time single step RTPCR reaction as follows:

1- qPCR reaction mix composition without Internal Control (When D-IC Ex Mix is added during DNA extraction)

Components	Volume per reaction (µL) (for final vol. of 25 µL)
RAS qDNA PCR Mix	12.5
RAS HBV PPM	2.0
DNA/HBQS/ MBGW	5.0
MBGW	5.5

2- qPCR reaction mix composition with Internal Control (When D-IC Ex Mix is not added during DNA extraction)

Components	Volume per reaction (µL) (for final vol. of 25 µL)
RAS qDNA PCR Mix	12.5
RAS HBV PPM	2.0
DNA/HBQS/ MBGW	5.0
RAS IC PCR Mix	1.0
MBGW	4.5



- 1. Addition of IC PCR mix (if required) should be done at the time of preparation of master mix.**
- 2. The results may be inconsistent, if the IC PCR mix is added individually.**

Place the PCR plate/tubes/strips in real time thermocycler.

PCR Programming

The Amplisure HBV Quantitation kit is validated on the following instruments:

- Rotor-Gene™ 6000
- Rotor-Gene™ Q 5plex
- ABI 7500 DX Real-Time PCR System
- ABI 7300 Real-Time PCR System
- Eppendorf Realplex 4
- Bio-Rad™ CFX 96

Plate Setup

1. Program the plate setup by labeling the slots as per tube/strip/plate labels. The sequence of labeling of slots should be the same way as the tube/strip/plate is kept in the machine.
2. Select the type of sample (Unknown/Standard/NTC) for each slot.
3. When Standard is selected as sample type, mention the quantitation value in IU/μL (2E4, 2E3 etc)
4. Select the channel for acquisition (FAM/Yakima Yellow)

Sl. No.	Name of channel	Source wavelength (nm)	Detection wavelength (nm)
1.	FAM (Pathogen target)	470	510
2.	Yakima Yellow (Internal Control)	530	555

5. For background calibration in different instruments, follow the procedure described below:

Rotor-Gene™ 6000	- Perform 'Gain optimization'
Rotor-Gene™ Q 5plex	- Perform 'Gain optimization'
ABI 7500 DX Real-Time PCR System	- Select Passive Reference dye 'ROX'
ABI 7300 Real-Time PCR System	- Select Passive Reference dye 'ROX'
Eppendorf Realplex 4	- Select 'ROX' for background calibration
Bio-Rad™ CFX 96	- Select 'ROX' for background calibration



Preparation of reaction mastermix and cycling conditions are same for all the instruments listed in the product insert. For instrument specific protocols, please contact our technical support team at amplisure@raslifesciences.com

Cycling conditions

1. Configure the following program in the machine.

Steps	No. of cycles	Temperature (°C)	Time
1 (Initial denaturation)	1	95	10 min.
2 (PCR cycling)	40	95	15 sec.
		60*	60 sec
* Plate Read/Data Acquisition in FAM and Yakima Yellow channel			

2. Set the reaction volume as 25 μ L.
3. Plate read/Data Acquisition for FAM and Yakima Yellow channel should be incorporated in the third stage of step 2 (60°C/60 sec).
4. The ideal run time for the assay is 90 minutes. Note: *In case of Eppendorf Realplex 4, select RAMP rate as 35%.*



Preparation of reaction mastermix and cycling conditions are same for all the instruments listed in the product insert. For instrument specific protocols, please contact our technical support team at amplisure@raslifesciences.com

Data Analysis

Analyze the data after completion of the run. Check the $R_n/Cycle$ amplification plot and $\Delta R_n/Cycle$ amplification plot to observe the amplification signal generated by different samples in the run. Compare both the plots for data analysis. Also look for noisy signals, if observed as it might not give you a proper result.

Setting the threshold for the qPCR Data analysis

The threshold should be set either automatically (by the machine itself)/ or manually just above the background signal of the negative controls and negative samples by referring to $R_n/Cycle$ amplification plot. The mean threshold value calculated from these experiments will most likely work for the majority of future runs, but the user should nevertheless review the generated threshold value at regular intervals.

Result

The values for unknown samples would appear in the result column in *IU/μL* in FAM Channel. Samples showing no amplification in FAM channel should show amplification in VIC channel, and then only results should be considered. The negative control should not show any value in the result column.

Interpretation

Interpret the values for unknown samples, only if the Slope of Standards is between -3.1 to -3.6 and PCR efficiency is between 90%-110% (0.9 - 1.1) and there should be no amplification in negative control.

Observation	Interpretation	Conclusion
Amplification signal detected in HBV channel (FAM) and in Internal control channel (Yakima Yellow)	HBV DNA within quantitation range	Proceed for further Analysis
Amplification signal detected in HBV channel (FAM) but no signal in Internal Control channel (Yakima Yellow)	HBV DNA within quantitation range	
Amplification signal not detected in HBV channel (FAM) but detected in Internal control channel (Yakima Yellow)	HBV DNA below quantitation limit	
No Amplification signal detected in HBV channel (FAM) as well as Internal control channel (Yakima Yellow) in unknown samples	Possible inhibition of PCR	Dilute the DNA sample (1:100) and repeat the assay*

To convert the results from *IU/μL* to *IU/mL* use the following formula:

$\text{Result (IU/mL)} = \frac{\text{Result (IU/}\mu\text{L)} \times \text{Elution Volume (}\mu\text{L)}}{\text{Sample Volume (mL)}}$

i * For calculating the result of diluted sample (1:100); multiply the observed IU/mL value by dilution factor, 100

$\text{Result of 1:100 diluted sample (IU/mL)} = \text{Dilution Factor} \times \frac{\text{Result (IU/}\mu\text{L)} \times \text{Elution Volume (}\mu\text{L)}}{\text{Sample Volume (mL)}}$

Troubleshoot

Observation	Possible cause	Solution
No amplification signal for Standards in FAM channel	1. One of the components may not have been added. 2. Incorrect channel selection 3. Incorrect programming of the real time machine. 4. Instrument is not working properly	1. Repeat the assay by following the correct protocol and addition of reagents 2&3. Please recheck the PCR program 4. Contact manufacturer of thermocycler for technical support.
Weak amplification signal for standards (Signal below threshold) in FAM Channel	1. Improper PCR programming. 2. Inaccurate dispensing of reagents 3. Possible deterioration of kit components due to improper storage	1. Repeat the assay by following the correct protocol 2. Minimize Pipetting errors/Check for calibration status of pipettes
Identical/Similar Ct values observed in FAM channel	1. Possible contamination of Kit reagents / Standards/Work area.	1. Use fresh aliquots of Standards/Kit Reagents (if available) 2. Clean the PCR rack/Pipettes thoroughly as per GLP 3. Clean and Fumigate the work area overnight prior to use

i For any other technical query; please contact amplisure@raslifesciences.com

Assay Characteristics

The analytical sensitivity of PCR assay as well as the analytical sensitivity in consideration of the purification (DNA Extraction) was assessed for the Amplisure® HBV Quantitative Kit.

The analytical sensitivity in consideration of the purification is determined using WHO standards/ HBV positive clinical samples in combination with a particular extraction method.

Analytical Sensitivity of qPCR assay

The analytical sensitivity is determined independent from the selected extraction method, using an international standard of known concentration ie. 3rd WHO International Standard for Hepatitis B Virus for Nucleic acid amplification techniques (NIBSC code: 10/264).

To determine the analytical sensitivity of the Amplisure® HBV Quantitative Kit, a standard dilution series was set up from 1.05 IU/μL to 42 IU/μL and analyzed with Amplisure® HBV Real Time PCR Quantitation Kit.

Testing was carried out on 10 different days on 3 replicates. The results were analyzed by statistical analysis.

The analytical sensitivity of the Amplisure® HBV Quantitative Kit is 2.01 IU/μL.

Analytical Sensitivity in consideration of Extraction Method

The analytical sensitivity in consideration of the purification using Roche High Pure Viral Nucleic Acid kit (Cat. No .11858874001) and QIAamp DNA Blood Mini Kit (Cat. No. 51104) of the Amplisure® HBV Quantitative Kit was also determined using a dilution series of the 3rd WHO International Standard for Hepatitis B Virus for Nucleic acid amplification techniques (NIBSC code: 10/264) from 8.5 to 8500 IU/mL spiked in healthy plasma specimens (Negative for HBV by Real time PCR).

These were subjected to DNA extraction using the Roche High Pure Viral Nucleic Acid kit and QIAamp DNA Blood Mini Kit (Roche, extraction volume: 0.2 mL, elution volume: 100 μL and Qiagen, extraction volume: 0.2 mL, elution volume: 100 μL)

Each of the dilutions was analyzed with the Amplisure® HBV Quantitative Kit on 10 different days on 3 replicates. The results were analyzed by statistical analysis.

The analytical sensitivity in consideration of the purification of the Amplisure® HBV Quantitative Kit, is 85 IU/mL (with Roche High Pure Viral Nucleic Acid kit) and is 90 IU/mL (with QIAamp DNA Blood Mini Kit)

Linear Range

The linear range (analytical measurement) of the Amplisure® HBV Quantitative Kit was determined by analyzing a dilution series of HBV quantitation standards from 2×10^7 IU/ μ L to 2 IU/ μ L. The dilution series has been calibrated against the 3rd WHO International Standard for Hepatitis B Virus for Nucleic acid amplification techniques (NIBSC code: 10/264).

Each dilution was tested in replicates ($n = 3$) using the Amplisure® HBV Quantitative Kit.

The linear range of the Amplisure® HBV Quantitation Kit has been determined to cover concentrations from **20 IU/ μ L to 2×10^7 IU/ μ L.**

Conversion Factor (IU/ μ L to Copies/ μ L)

The 3rd WHO International Standard for Hepatitis B Virus for Nucleic acid amplification techniques (NIBSC code: 10/264) has been assigned an International Unit (IU) by the WHO as the number of genome equivalents.

The conversion factor value for Amplisure® HBV Quantitative kit has been assigned after calibration of the HBQS (HBV quantitation standards) with the 3rd WHO International Standard for Hepatitis B Virus for Nucleic acid amplification techniques (NIBSC code: 10/264). The Conversion Factor (IU/ μ L to Copies/ μ L) is entirely dependent on the current PCR assay kit. A Correlation and Regression analysis was performed to compare the two standard curves of HBV DNA values obtained by Amplisure® HBV Quantitative kit (copies/ μ L) and HBV NIBSC standards (IU/ μ L).

The correlation coefficient between the expected and estimated values was very good. Please see Fig. 2 below

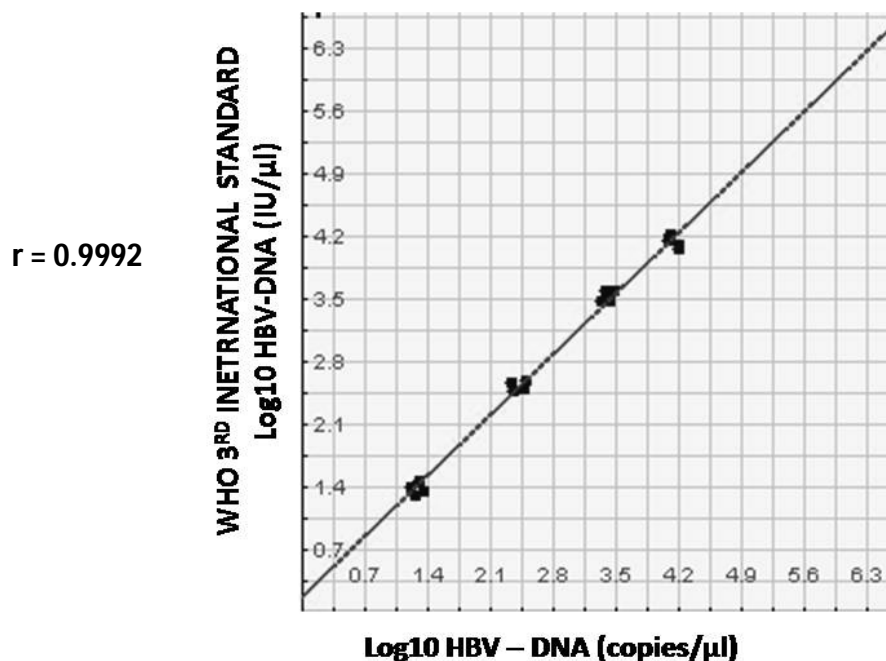


Fig. 2 Correlation and Regression plot of HBV NIBSC standards (IU/μL) and HBV DNA (copies/μL)

The conversion factor calculated for Amplisure® HBV PCR kit from IU/μL to Copies/ μL is 1.28, i.e. **1 IU/μL = 1.28 copies/μL**

To convert into copies/mL use the following formula:

$$\text{Result (copies/mL)} = \frac{\text{Calculated Value (copies/μL)} \times \text{Elution Volume (μL)}}{\text{Sample Volume (mL)}}$$

Specificity

The specificity of the Amplisure® HBV Quantitative Kit is ensured by the selection of the primers and probes, as well as the selection of stringent reaction conditions. The primers and probes were checked for possible homologies to all published sequences (Genbank) by BLAST analysis.

Detection of Different HBV genotypes (A-H) using International Standard

The detectability of all relevant genotypes (A-H) was ensured by a Database alignment and by a qPCR run with the following genotypes, using ≥ 400 positive HBV samples covering all genotypes (A-H) which were obtained from NABL accredited RAS Molecular Diagnostic Lab. The project was approved by ethical committee of RAS Lifesciences Pvt. Ltd. The entire clinical specimen was confirmed for genotypes by DNA sequencing.

Sl. No.	HBV Genotype Controls	Source of Controls	Result obtained with Amplisure® HBV Quantitative Kit
1.	Geno A	RAS*	Detected
2.	Geno B	RAS*	Detected
3.	Geno C	RAS*	Detected
4.	Geno D	RAS*	Detected
5.	Geno E	RAS*	Detected
6.	Geno F	RAS*	Detected
7.	Geno G	RAS*	Detected
8.	Geno H	RAS*	Detected

***NABL accredited RAS Molecular Diagnostic Lab**

Cross Reactivity Data

A potential cross-reactivity of the Amplisure® HBV Quantitative Kit was tested using the control group listed below. None of the tested pathogens has been reactive. No cross-reactivity appeared with mixed infections.

Moreover, the specificity was validated with 50 different healthy plasma specimens/other various sample types. These did not generate any signals with the HBV specific primers and probes, which are included in the Amplisure® HBV Quantitative Kit.

Pathogen Tested	Cross Reactivity with the HBV Primers/Probes
Adenovirus	-ve
Herpes Simplex Virus-1	-ve
Herpes Simplex Virus-2	-ve
Epstein Barr virus	-ve
Human Immunodeficiency Virus	-ve
Cytomegalo virus	-ve
Hepatitis C virus	-ve
Enterovirus	-ve
BK virus	-ve
MTB complex	-ve
Plasmodium spp.	-ve

Precision

The precision data of the Amplisure® HBV Quantitative Kit have been generated for HBV positive clinical specimens and HBV quantitation standards.

Intra-assay variability

The precision data consists of the *Intra-assay variability* which is variability of multiple results of samples of the same concentration within one experiment.

The data obtained were used to determine the standard deviation and the coefficient of variation for the pathogen specific PCR. Precision data of the Amplisure® HBV Quantitative Kit have been collected using all the quantitation standards (HBQS1- HBQS4). Also included were three different samples with different viral loads. Testing was performed with 3 replicates for 10 days.

Inter-assay variability

The precision data consists of the *Inter-assay variability* which is variability of multiple results of samples of the same concentration within different experiments.

The data obtained were used to determine the standard deviation and the coefficient of variation for the pathogen specific PCR. Precision data of the Amplisure® HBV Quantitative Kit have been collected using all the quantitation standards (HBQS1- HBQS4). Also included were three different samples with different viral loads. Testing was performed with 3 replicates for 10 days.

The precision data was calculated on basis of the CT values obtained.

Sample Type	Mean [Log Value (IU/ μ L)]	Variability Testing	Standard Deviation	Coefficient of Variation (%)
HBQS1	Log 4.3	Intra-assay variability	0.03	0.001
		Inter-assay variability	0.37	0.017
HBQS2	Log 3.3	Intra-assay variability	0.04	0.001
		Inter-assay variability	0.25	0.010
HBQS3	Log 2.3	Intra-assay variability	0.01	0.00
		Inter-assay variability	0.46	0.020
HBQS4	Log 1.3	Intra-assay variability	0.02	0.001
		Inter-assay variability	0.32	0.010
CONTROL 1	Log 9.15	Intra-assay variability	0.19	0.01
		Inter-assay variability	0.30	0.057
CONTROL 2	Log 7.30	Intra-assay variability	0.21	0.02
		Inter-assay variability	0.36	0.038
CONTROL 3	Log 4.48	Intra-assay variability	0.41	0.02
		Inter-assay variability	0.40	0.016

Abbreviations

<i>Abbreviation</i>	<i>Expansion</i>
HBV	Hepatitis B Virus
DNA	Deoxy Ribonucleic Acid
IC	Internal Control
DNases	Deoxyribonucleases
PCR	Polymerase Chain Reaction
BSL2	Bio Safety Level 2
BSL3	Bio Safety Level 3
mL	Milli Liters
µL	Micro Liters
K ₂ EDTA	Potassium Ethylene Diamine Tetra Acetate
g	Relative Centrifugal Force
qPCR Protocol	Quantitative PCR protocol
MBGW	Molecular Biology Grade Water
RT PCR	Real Time PCR
NTC	No Template Control
FAM	Carboxyfluorescein
ROX	Carboxy-X-rhodamine
NIBSC	National Institute for Biological Standards and Control
IU	International Units
WHO	World Health Organization
BLAST	Basic Local Alignment Search Tool
MTB complex	Mycobacterium tuberculosis complex
Sps	Species
Rxn	Reaction

References

1. Handbook "Sampling, Transportation, and Storage of Clinical Material for PCR Diagnostics", developed by Federal State Institution of Science Central Research Institute of Epidemiology of Federal Service for Surveillance on Consumers' Rights Protection and Human Well-Being, Moscow, 2008.
2. Liu Y, Hussain M, Wong S, Fung SK, Yim HJ, Lok AS. A genotype- independent real – time PCR assay for quantification of hepatitis B virus DNA. J Clin Microbiol. 2007; 45(2):553-558.

Ordering Information



QT-HBV-50	: 50 rxns
QT-HBV-100	: 100 rxns