# **Instructions for Use**

RealStar® HSV PCR Kit 1.0

06/2016 EN

# RealStar® HSV PCR Kit 1.0

For research use only!

(RUO)

REF

061003

 $\sum$ 

96

 $\prod$ 

06 2016



altona Diagnostics GmbH • Mörkenstr. 12 • D-22767 Hamburg

# Content

1.	Application	5
2.	Kit Components	5
3.	Storage	5
4.	Product Description	6
4.1	Real-Time PCR Instruments	8
5.	Procedure	9
5.1	Sample Preparation	9
5.2	Master Mix Setup	10
5.3	Reaction Setup	12
6.	Programming the Real-Time PCR Instrument	13
6.1	Settings	13
6.2	Fluorescence Detectors (Dyes)	13
6.3	Temperature Profile and Dye Acquisition	14
7.	Data Analysis	14
7.1	Interpretation of Results	15
7.1.1	Qualitative Analysis	15
7.1.2	Quantitative Analysis	15
8.	Technical Assistance	17
9.	Trademarks and Disclaimers	17
10.	Explanation of Symbols	18

# 1. Application

The RealStar® HSV PCR Kit 1.0 is a reagent system, based on real-time PCR technology, for the simultaneous detection and quantification of herpes simplex virus 1 (HSV-1) and herpes simplex virus 2 (HSV-2) specific DNA.

For research use only (RUO)! Not for use in diagnostic procedures.

# 2. Kit Components

Lid Color	Component	Number of Vials	Volume [µl/Vial]
Blue	Master A	8	60
Purple	Master B	8	180
Green	Internal Control	1	1000
Red	HSV-1 QS1-4*	4	250
Orange	HSV-2 QS1-4*	4	250
White	Water (PCR grade)	1	500

<sup>\*</sup> The RealStar® HSV PCR Kit 1.0 contains four HSV-1 Quantification Standards as well as four HSV-2 Quantification Standards.

# 3. Storage

- The RealStar® HSV PCR Kit 1.0 is shipped on dry ice. The components of the kit should arrive frozen. If one or more components are not frozen upon receipt, or if tubes have been compromised during shipment, contact altona Diagnostics GmbH for assistance.
- All components should be stored between -25°C and -15°C upon arrival.
- Repeated thawing and freezing of Master reagents (more than twice) should be avoided, as this might affect the performance of the assay. The reagents should be frozen in aliquots, if they are to be used intermittently.
- Storage between +2°C and +8°C should not exceed a period of two hours.
- · Protect Master A and Master B from light.

# 4. Product Description

The RealStar® HSV PCR Kit 1.0 is a reagent system, based on real-time PCR technology, for the simultaneous detection and quantification of herpes simplex virus 1 (HSV-1) and herpes simplex virus 2 (HSV-2) specific DNA. The assay includes a heterologous amplification system (Internal Control) to identify possible PCR inhibition and to confirm the integrity of the reagents of the kit.

Real-time PCR technology utilizes polymerase chain reaction (PCR) for the amplification of specific target sequences and target specific probes for the detection of the amplified DNA. The probes are labelled with fluorescent reporter and quencher dyes.

Probes specific for HSV-1 DNA are labelled with the fluorophore FAM<sup>TM</sup> whereas the probes specific for HSV-2 DNA are labelled with a fluorophore showing similar characteristics to  $Cy^{\otimes}5$ . The probe specific for Internal Control (IC) is labelled with the fluorophore  $JOE^{TM}$ .

Using probes linked to distinguishable dyes enables the parallel detection of HSV-1 and HSV-2 specific DNA as well as the detection of the Internal Control in corresponding detector channels of the real-time PCR instrument.

The test consists of two processes in a single tube assay:

- PCR amplification of target DNA and Internal Control
- Simultaneous detection of PCR amplicons by fluorescent dye labelled probes

The RealStar® HSV PCR Kit 1.0 consists of:

- Two Master reagents (Master A and Master B)
- Internal Control (IC)
- Four HSV-1 specific Quantification Standards (QS1-QS4)
- Four HSV-2 specific Quantification Standards (QS1-QS4)
- PCR grade water

Master A and Master B contain all components (buffer, enzymes, primers and probes) to allow PCR mediated amplification and target detection of HSV-1 and HSV-2 specific DNA and Internal Control in one reaction setup.

The Quantification Standards contain standardized concentrations of HSV-1 and HSV-2 specific DNA. The Quantification Standards can be used individually as positive controls, or together to generate a **standard curve**, which can be used to determine the concentration of HSV-1 and/or HSV-2 specific DNA in the sample.

The Quantification Standards have the following concentrations:

Quantification Standard	Concentration [copies/μΙ]
QS1	1.00E+04
QS2	1.00E+03
QS3	1.00E+02
QS4	1.00E+01

### 4.1 Real-Time PCR Instruments

The RealStar® HSV PCR Kit 1.0 can be used with the following real-time PCR instruments:

- m2000rt (Abbott Diagnostics)
- Mx 3005P™ QPCR System (Stratagene)
- VERSANT® kPCR Molecular System AD (Siemens)
- ABI Prism<sup>®</sup> 7500 SDS (Applied Biosystems)
- ABI Prism® 7500 Fast SDS (Applied Biosystems)
- Rotor-Gene® 6000 (Corbett Research)
- Rotor-Gene® Q5/6 plex Platform (QIAGEN)
- LightCycler® 480 Instrument II (Roche)
- CFX96™ Real-Time System (Bio-Rad)
- CFX96™ Deep Well Real-Time System (Bio-Rad)

### **NOTE**



Please ensure that all instruments used have been installed, calibrated, checked and maintained according to the manufacturer's instructions and recommendations.

### 5. Procedure

## 5.1 Sample Preparation

Extracted DNA is the starting material for the RealStar® HSV PCR Kit 1.0.

The quality of the extracted DNA has a profound impact on the performance of the entire test system. It has to be ensured that the system used for nucleic acid extraction is compatible with real-time PCR technology. The following kits and systems are suitable for nucleic acid extraction:

- QIAamp® DNA Mini Kit (QIAGEN)
- QIAsymphony® (QIAGEN)
- NucliSENS® easyMag® (bioMérieux)
- MagNa Pure 96 System (Roche)
- m2000sp (Abbott)
- Maxwell® 16IVD Instrument (Promega)
- VERSANT® kPCR Molecular System SP (Siemens Healthcare)

Alternative nucleic acid extraction systems and kits might also be appropriate.

If using a spin column based sample preparation procedure including washing buffers containing ethanol, it is highly recommended to perform an additional centrifugation step for 10 min at approximately 17000 x g ( $\sim$  13000 rpm), using a new collection tube, prior to the elution of the nucleic acid.

### **CAUTION**



If your sample preparation system is using washing buffers containing ethanol, make sure to eliminate any traces of ethanol prior to elution of the nucleic acid. Ethanol is a strong inhibitor of real-time PCR.

### **CAUTION**



The use of carrier RNA is crucial for extraction efficiency and stability of the extracted nucleic acid.

For additional information and technical support regarding pre-treatment and sample preparation please contact our Technical Support (see chapter 8. Technical Assistance).

# 5.2 Master Mix Setup

All reagents and samples should be thawed completely, mixed (by pipetting or gentle vortexing) and centrifuged briefly before use.

The RealStar® HSV PCR Kit 1.0 contains a heterologous Internal Control (IC), which can either be used as a PCR inhibition control or as a control of the sample preparation procedure (nucleic acid extraction) <u>and</u> as a PCR inhibition control.

► If the IC is used as a PCR inhibition control, but not as a control for the sample preparation procedure, set up the Master Mix according to the following pipetting scheme:

Number of Reactions (rxns)	1	12
Master A	5 μΙ	60 µl
Master B	15 µl	180 µl
Internal Control	1 µl	12 µl
Volume Master Mix	21 µl	252 μΙ

▶ If the IC is used as a control for the sample preparation procedure <u>and</u> as a PCR inhibition control, add the IC during the nucleic acid extraction procedure.

- ▶ No matter which method/system is used for nucleic acid extraction, the IC must not be added directly to the sample. The IC should always be added to the sample/lysis buffer mixture. The volume of the IC which has to be added, always and only depends on the elution volume. It represents 10% of the elution volume. For instance, if the nucleic acid is going to be eluted in 60 μl of elution buffer or water, 6 μl of IC per sample must be added into the sample/ lysis buffer mixture.
- ▶ If the IC was added during the sample preparation procedure, set up the Master Mix according to the following pipetting scheme:

Number of Reactions (rxns)	1	12
Master A	5 µl	60 µl
Master B	15 µl	180 µl
Volume Master Mix	20 μΙ	240 μl

### **CAUTION**



If the IC (Internal Control) was added during the sample preparation procedure, the Master Mix for the controls must be prepared including the IC.

### **CAUTION**



No matter which method/system is used for nucleic acid extraction, never add the IC directly to the sample.

# 5.3 Reaction Setup

- Pipette 20 μl of the Master Mix into each required well of an appropriate optical 96-well reaction plate or an appropriate optical reaction tube.
- Add 10 μl of the sample (eluate from the nucleic acid extraction) or 10 μl of the controls (Quantification Standard, Positive or Negative Control).

Reaction Setup				
Master Mix	20 μΙ			
Sample or Control	10 μΙ			
Total Volume	30 µl			

- ► Make sure that each Positive Control (QS) and at least one Negative Control is used per run.
- ► For quantification purposes all of each (HSV-1 and HSV-2) Quantification Standards (QS1 to QS4) should be used.
- ► Thoroughly mix the samples and controls with the Master Mix by pipetting up and down.
- ► Close the 96-well reaction plate with appropriate lids or optical adhesive film and the reaction tubes with appropriate lids.
- ► Centrifuge the 96-well reaction plate in a centrifuge with a microtiter plate rotor for 30 seconds at approximately 1000 x g (~ 3000 rpm).

# 6. Programming the Real-Time PCR Instrument

For basic information regarding the setup and programming of the different real-time PCR instruments, please refer to the user manual of the respective instrument. For detailed programming instructions regarding the use of the RealStar® HSV PCR Kit 1.0 on specific real-time PCR instruments please contact our Technical Support (see chapter 8. Technical Assistance).

# 6.1 Settings

Define the following settings:

Settings				
Reaction Volume	30 µl			
Ramp Rate	Default			
Passive Reference	ROX™			

# 6.2 Fluorescence Detectors (Dyes)

▶ Define the fluorescence detectors (dyes):

Target	Detector Name	Reporter	Quencher
HSV-1 specific DNA	HSV-1	FAM™	(None)
HSV-2 specific DNA	HSV-2	Cy®5	(None)
Internal Control	IC	JOE™	(None)

# 6.3 Temperature Profile and Dye Acquisition

▶ Define the temperature profile and dye acquisition:

	Stage	Cycle Repeats	Acquisition	Temperature [°C]	Time [min:sec]
Denaturation	Hold	1	-	95	10:00
Amplification	Cycling	45	-	95	00:15
Amplification	Cycling 45	yes	58	01:00	

# 7. Data Analysis

For basic information regarding data analysis on specific real-time PCR instruments, please refer to the user manual of the respective instrument.

For detailed instructions regarding the analysis of the data generated with the RealStar® HSV PCR Kit 1.0 on different real-time PCR instruments please contact our Technical Support (see chapter 8. Technical Assistance).

# 7.1 Interpretation of Results

# 7.1.1 Qualitative Analysis

Detection Channel		nel	Donald Intermediation
FAM™	Cy®5	JOE™	Result Interpretation
+	-	+*	HSV-1 specific DNA detected.
-	+	+*	HSV-2 specific DNA detected.
-	-	+	Neither HSV-1 nor HSV-2 specific DNA detected.  The sample does not contain detectable amounts of HSV-1 or HSV-2 specific DNA.
-	-	-	PCR inhibition or reagent failure. Repeat testing from original sample or collect and test a new sample.

<sup>\*</sup> Detection of the Internal Control in the JOE™ detection channel is not required for positive results either in the FAM™ detection channel or in the Cy®5 detection channel. High HSV-1 and/or HSV-2 DNA load/s in the sample can lead to reduced or absent Internal Control signals.

# 7.1.2 Quantitative Analysis

The RealStar® HSV PCR Kit 1.0 provides four Quantification Standards (QS) for HSV-1 and four Quantification Standards (QS) for HSV-2. In order to generate a **standard curve** for quantitative analysis, these have to be defined as **standards** with appropriate concentrations (chapter 4. Product Description). Using **standards** of known concentrations a standard curve for quantitative analysis can be generated.

$$C_t = m \cdot \log(N_0) + b$$

C, = Threshold Cycle

m = Slope

N<sub>o</sub> = Initial Concentration

b = Intercept

Derived from the standard curve positive samples of unknown concentrations can be quantified.

$$N_0 = 10^{\left(\frac{C_1 - b}{m}\right)}$$

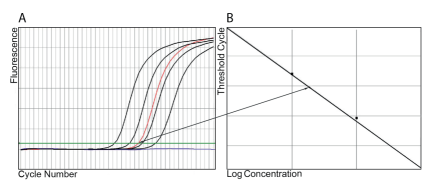


Figure 1: Quantification Standards (black), a positive (red) and a negative sample (blue) displayed in the Amplification Plot [A] and Standard Curve analysis [B]

To determine the **viral load of the original sample**, the following formula has to be applied:

### **NOTE**



The concentration of the "Sample" is displayed in copies/µl and refers to the concentration in the eluate.

# 8. Technical Assistance

For technical advice, please contact our Technical Support:

e-mail: support@altona-diagnostics.com

phone: +49-(0)40-5480676-0

### 9. Trademarks and Disclaimers

RealStar® (altona Diagnostics); Mx 3005P™ (Stratagene); ABI Prism® (Applied Biosystems); LightCycler® (Roche);Rotor-Gene®, QIAamp®, QIAsymphony® (QIAGEN); Maxwell® (Promega); NucliSENS®, easyMag® (bioMérieux); VERSANT® (Siemens Healthcare); CFX96™ (Bio-Rad); FAM™, JOE™ (Life Technologies); Cy® (GE Healthcare).

Registered names, trademarks, etc. used in this document, even if not specifically marked as such, are not to be considered unprotected by law.

For research use only! Not for use in diagnostic procedures.

© 2016 altona Diagnostics GmbH; all rights reserved.

# 10. Explanation of Symbols

RUO For research use only LOT Batch code CAP Cap color REF Product number CONT Content NUM Number COMP Component Version Consult instructions for use Contains sufficient for "n" tests/reactions (rxns) Temperature limit Use-by date Manufacturer Caution Note

