

IVD For *in Vitro* Diagnostic Use


For Professional Use Only

CMV/EBV/HHV6 Quant Real-TM

Handbook

Multiplex Real Time PCR Kit for quantitative detection and differentiation of Cytomegalovirus (CMV), Epstein Barr Virus (EBV) and Human Herpes 6 Virus (HHV6)

REF V48-100FRT

 **100**

NAME

CMV/EBV/HHV6 Quant Real - TM

INTENDED USE

The **CMV/EBV/HHV6 Quant Real-TM** is a “Real-Time Amplification” test for the quantitative detection and differentiation of Cytomegalovirus (CMV), Epstein Barr Virus (EBV) and Human Herpes 6 Virus (HHV6) in the biological materials. DNA is extracted from samples, amplified using real time amplification with fluorescent reporter dye probes specific for CMV/EBV/HHV6 and Internal Control (IC). Test contains an IC (β -globine gene) which allows controlling both PCR-analysis stages (DNA extraction and PCR amplification), material sampling, and storage conditions.

PRINCIPLE OF PCR DETECTION

CMV, EBV and HHV6 detection by polymerase chain reaction (PCR) with hybridization-fluorescence detection includes DNA extraction from clinical samples and PCR amplification of pathogen genome specific region with real-time hybridization-fluorescence detection. During DNA extraction from clinical material, human genomic DNA (endogenous internal control) is amplified. Endogenous internal control (IC Glob) allows controlling both PCR-analysis stages (DNA extraction and PCR amplification), material sampling, and storage adequacy. Then, the obtained samples are amplified using specific primers and polymerase (TaqF). In real-time PCR, the amplified product is detected using fluorescent dyes. These dyes are linked to oligonucleotide probes which bind specifically to the amplified product during thermocycling. The real-time monitoring of the fluorescence intensities during the real-time PCR allows the detection of accumulating product without re-opening the reaction tubes after the PCR run.

MATERIALS PROVIDED

<i>Reagent</i>	<i>Description</i>	<i>Volume, ml</i>	<i>Quantity</i>
PCR-mix-1-FRT EBV / CMV / HHV-6 / Glob	colorless clear liquid	0.6	2 tubes
PCR-mix-2-FRT	colorless clear liquid	0.3	2 tubes
Polymerase (TaqF)	colorless clear liquid	0.03	2 tubes
RNA-buffer	colorless clear liquid	0.6	1 tube
DNA calibrator KSG1	colorless clear liquid	0.2	1 tube
DNA calibrator KSG2	colorless clear liquid	0.2	1 tube
Negative Control (C-)*	colorless clear liquid	1.2	2 tubes
Positive Control DNA EBV / CMV / HHV-6 and human DNA**	colorless clear liquid	0.1	2 tubes

* *must be used in the extraction procedure as Negative Control of Extraction.*

** *must be used in the extraction procedure as Positive Control of Extraction (PCE).*

MATERIALS REQUIRED BUT NOT PROVIDED

- DNA extraction kit.
- Disposable powder-free gloves and laboratory coat.
- Automated pipettors (dosers) of variable volumes (from 5 to 20 μ l and from 20 to 200 μ l).
- Disposable tips with aerosol barriers (100 or 200 μ l) in tube racks.
- Tube racks
- Vortex mixer/desktop centrifuge.
- PCR box.
- Personal thermocyclers (for example, Rotor-Gene 3000 or Rotor-Gene 6000 (Corbett Research,); Rotor-Gene Q (Qiagen) iQ5 and iCycler iQ (Bio-Rad), Mx3000P (Stratagene) or equivalent).
- Disposable polypropylene microtubes for PCR or PCR-plate.
- Refrigerator for 2–8 °C.
- Deep-freezer for ≤ -16 °C.
- Waste bin for used tips.

STORAGE INSTRUCTIONS

All components of the **CMV/EBV/HHV6 Quant Real-TM** PCR kit (except for PCR-mix-1-FRT *EBV/CMV/HHV-6* /Glob, PCR-mix-2-FRT, and Polymerase (TaqF)) are to be stored at 2–8 °C when not in use. The kit can be shipped at 2-8°C but should be stored -20°C immediately on receipt. The shelf life of reagents before and after the first use is the same, unless otherwise stated.

STABILITY

CMV/EBV/HHV6 Quant Real-TM Test is stable up to the expiration date indicated on the kit label. The product will maintain performance through the control date printed on the label. Exposure to light, heat or humidity may affect the shelf life of some of the kit components and should be avoided. Repeated thawing and freezing of these reagents should be avoided, as this may reduce the sensitivity. Components stored under conditions other than those stated on the labels may not perform properly and may adversely affect the assay results.

WARNINGS AND PRECAUTIONS



***In Vitro* Diagnostic Medical Device**

For *In Vitro* Diagnostic Use Only

1. Wear disposable gloves, laboratory coats and eye protection when handling specimens and reagents. Thoroughly wash hands afterward.
2. Do not pipette by mouth.
3. Do not eat, drink, smoke, apply cosmetics, or handle contact lenses in laboratory work areas.
4. Do not use a kit after its expiration date.
5. Dispose of all specimens and unused reagents in accordance with local regulations.
6. Biosafety Level 2 should be used for materials that contain or are suspected of containing infectious agents.
7. Clean and disinfect all spills of specimens or reagents using a disinfectant such as 0,5% sodium hypochlorite, or other suitable disinfectant.
8. Avoid contact of specimens and reagents with the skin, eyes and mucous membranes. If these solutions come into contact, rinse immediately with water and seek medical advice immediately.
9. Material Safety Data Sheets (MSDS) are available on request.
10. Use of this product should be limited to personnel trained in the techniques of DNA amplification.
11. PCR reactions are sensitive to contamination. Measures to reduce the risk of contamination in the laboratory include physically separating the activities involved in performing PCR in compliance with good laboratory practice.
12. Workflow in the laboratory must proceed in a uni-directional manner, beginning in the Extraction Area and moving to the Amplification and Detection Area. Do not return samples, equipment and reagents in the area where you performed previous step.



Some components of this kit contain sodium azide as a preservative. Do not use metal tubing for reagent transfer.

PRODUCT USE LIMITATIONS

All reagents may exclusively be used in in vitro diagnostics. Use of this product should be limited to personnel trained in the techniques of DNA amplification (EN375). Strict compliance with the user manual is required for optimal PCR results. Attention should be paid to expiration dates printed on the box and labels of all components. Do not use a kit after its expiration date.

QUALITY CONTROL

In accordance with Sacace's ISO 13485-Certified Quality Management System, each lot is tested against predetermined specifications to ensure consistent product quality.

SAMPLE COLLECTION, STORAGE AND TRANSPORT

CMV/EBV/HHV6 Quant Real-TM can analyze DNA extracted from:

- *whole peripheral and umbilical cord blood* collected in either ACD or EDTA tubes;
- *buffy coat*;
- *tissue* homogenized with mechanical homogenizer and dissolved in PBS sterile;
- *urine (sediment)*;
- *swabs*: insert the swab into the nuclease-free 1,5 ml tube and add 0,2 mL of Transport medium. Vigorously agitate swabs in medium for 15-20 sec.

It is recommended to process samples immediately after collection. Store samples at 2–8 °C for no longer than 24 hours, or freeze at –20/80°C. Transportation of clinical specimens must comply with country, federal, state and local regulations for the transport of etiologic agents.

DNA ISOLATION

The following isolation kit is recommended:

⇒ **DNA-Sorb-B** (Sacace, REF K-1-1/B)

⇒ **DNA/RNA-Prep** (Sacace, REF K-2-9)



Extract DNA according to the manufacturer's instructions.



Transfer **100 µl** of **Negative Control** to the tube labeled C-. Transfer **90 µl** of **Negative Control** and **10 µl** of **Positive Control DNA EBV/CMV/HHV-6 and human DNA** to the tube labeled PCE.

PROTOCOL (Reaction volume 25 µl):

1. Prepare in the new sterile tube for each sample **10*N µl** of **PCR-mix-1 "CMV/EBV/HHV6/IC"**, **5,0*N** of **PCR-Buffer-FRT** and **0,5*N** of **TaqF DNA Polymerase**. Vortex and centrifuge for 2-3 sec.

2. Prepare required quantity of reaction tubes for samples and controls and add **15 µl** of **Reaction Mix** and **10 µl** of **extracted DNA** sample to appropriate tube. Mix by pipetting.

(Re-centrifuge all the tubes with extracted DNA for 2 min at maximum speed (12000-16000 g) and take carefully supernatant. N.B. don't disturb the pellet, sorbent inhibit reaction!).

3. **For qualitative analysis:**

NCA - Add **10 µl** of **RNA-buffer** to the tube labeled NCA (Negative Control of Amplification).

C+ - Add **10 µl** of DNA calibrator **KSG2** to the tube labeled C+ (Positive Control of Amplification).

4. **For quantitative analysis:**

NCA	- Add 10 µl of RNA-buffer to the tube labeled NCA (Negative Control of Amplification).
Calibrators KSG1 and KSG2	- Add 10 µl of KSG1 to two tubes and add 10 µl of KSG2 to other two tubes

Close tubes and transfer them into the instrument in this order: samples, negative controls, positive control, Standards.

Amplification program for rotor-type instruments¹

Step	Temperature, °C	Time	Fluorescence detection	Cycles
Hold	95	15 min	–	1
Cycling 1	95	5 s	–	5
	60	20 s	–	
	72	15 s	–	
Cycling 2	95	5 s	–	40
	60	20 s	FAM/Green, JOE/Yellow, ROX/Orange, Cy5/Red	
	72	15 s	–	

Amplification program for plate-type and modular type instruments²

Step	Temperature, °C	Time	Fluorescence detection	Cycles
1	95	15 min	–	1
2	95	5 s	–	5
	60	20 s	–	
	72	15 s	–	
3	95	5 s	–	40
	60	30 s	FAM, JOE/HEX/Cy3, ROX/TexasRed, Cy5	
	72	15 s	–	

¹ For example Rotor-Gene™ 6000/Q (Qiagen)

² For example, SaCycler-96™ (Sacace), iQ5™ (BioRad); Mx3005P™ (Agilent Technologies), ABI® 7500 Real Time PCR (Applied Biosystems), SmartCycler® (Cepheid)

RESULTS ANALYSIS

β -Globin gene DNA (IC) is detected in the FAM/Green channel, EBV DNA is detected in the JOE/HEX/Cy3/Yellow channel, CMV DNA is detected in the ROX/TexasRed/Orange channel, and HHV6 DNA is detected in the Cy5/Red channel.

Interpretation of results

The results are interpreted by the software of the used Instrument by the crossing (or not) of the fluorescence curve with the threshold line.

1. The sample is considered to be **positive** for **EBV DNA** if its Ct value in the results grid on the JOE/HEX/Cy3/Yellow channel is detected and does not exceed the threshold value of positive result.
2. The sample is considered to be **positive** for **CMV DNA** if its Ct value in the results grid on the ROX/Orange/TexasRed channel is defined and does not exceed the threshold value of positive result.
3. The sample is considered to be **positive** for **HHV6 DNA** if its Ct value in the results grid on the Cy5/Red channel is defined and does not exceed the threshold value of positive result.
4. For qualitative analysis, the sample is considered to be **negative** if its Ct value in the results grid in the FAM/Green channel does not exceed the Ct value indicated in the **Product Data Sheet**.
5. For quantitative analysis, the quantity of IC Glob DNA should be greater than 2000 copies per reaction for whole blood, white blood cells, viscera biopsy material or more than 500 copies per reaction for saliva and oropharyngeal swabs.



*For cerebrospinal fluid (liquor), the Ct value can be greater than the Ct value indicated in the **Product Data Card** in the results grid in the FAM/Green channel or the quantity of IC Glob DNA can be less than 500 copies per reaction in case of quantitative analysis because the cerebrospinal fluid samples may contain a very small number of cells.*

6. For **qualitative** analysis, the result of analysis is considered to be **invalid** if the Ct value is not detected in the results grid (the fluorescence curve does not cross the threshold line) or if it is greater than the threshold value in the JOE/HEX/Yellow, ROX/Orange, or Cy5/Red channel and the Ct value in the results grid in the FAM/Green channel exceeds the Ct value indicated in the **Product Data Sheet**.
7. For **quantitative** analysis, the analysis result is considered to be **invalid** if the Ct value is not detected in the results grid (the fluorescence curve does not cross the threshold line) or if it is greater than the boundary value in the JOE/HEX/Yellow, ROX/Orange, or Cy5/Red channel and the quantity of IC Glob DNA is less than 2000 copies per reaction for whole blood, white blood cells, viscera biopsy material or if it is less than 500 copies per reaction for saliva and oropharyngeal swabs. In such cases, PCR analysis of the sample should be repeated.
8. For qualitative analysis, results of analysis are considered reliable only if the results obtained for both Positive and Negative Controls of amplification as well as for the Negative Control of extraction are correct. For quantitative analysis, results on C+ should fall in range of concentrations indicated in the **Product Data Sheet**.

Table 1. Results for controls

Control	Stage for control	Ct in channel				Interpretation
		FAM/Green	JOE/HEX/ Cy3/Yellow	ROX/Orange/ TexasRed	Cy5/Red	
NCE	DNA extraction, PCR	Neg	Neg	Neg	Neg	OK
NCA	PCR	Neg	Neg	Neg	Neg	OK
C+	PCR	POS	POS	POS	POS	OK
QS1 QS2	PCR	Pos (see Data Sheet)	Pos (see Data Sheet)	Pos (see Data Sheet)	Pos (see Data Sheet)	OK

Quantitative results

In quantitative analysis, if total DNA is extracted from **human whole blood, white blood cells and biopsy material**, the concentration in log of DNA copies per standard cell quantity (10^5) in control and test samples is calculated by the following formula:

For *CMV*:

$$\log \left\{ \frac{\text{CMV DNA copies in PCR sample}}{\text{Glob DNA copies in PCR sample}} \times 2 \cdot 10^5 \right\} = \log \{ \text{CMV DNA copies} / 10^5 \text{ of cells} \}.$$

For *EBV*:

$$\log \left\{ \frac{\text{EBV DNA copies in PCR sample}}{\text{Glob DNA copies in PCR sample}} \times 2 \cdot 10^5 \right\} = \log \{ \text{EBV DNA copies} / 10^5 \text{ of cells} \}.$$

For *HHV6*:

$$\log \left\{ \frac{\text{HHV6 DNA copies in PCR sample}}{\text{Glob DNA copies in PCR sample}} \times 2 \cdot 10^5 \right\} = \log \{ \text{HHV6 DNA copies} / 10^5 \text{ of cells} \}.$$

The results can be calculated manually or using Excel tables. To do this copy the names of the samples and insert them in the first column (Column A). Copy the concentrations of EBV DNA from the channel Joe(Yellow)/HEX/Cy3 and paste in the second column of Excel table (Column B). Copy the concentrations of IC Glob from the channel Fam(Green) and paste in the third column of Excel table (Column C). Insert in the column D the formula D=LOG (B/C*200000): log values will appear.

Name	Calc Conc (copies/reaction) Joe(Yellow)/HEX/Cy3	Calc Conc (copies/reaction) Fam(Green)	log EBV/10 ⁵ cells
A	B	C	D
1	8742	125640	4,1
2	253	87787	2,8
3		65765	
4	648	16354	3,9
5		76865	
QS1	9962	9793	
QS1	10011	10143	
QS2	98	103	
QS2	102	97	
Neg PCR			

Use the same procedure for calculation of CMV (ROX/Orange/TexasRed channel) and HHV6 (Cy5/Red channel) log quantity inserting in the column B the relative results.

If total DNA is extracted from **saliva, oropharyngeal swabs and cerebrospinal fluid (liquor)**, the concentration of DNA per ml of sample (Conc DNA) is calculated by the following formula:

$$\text{Conc DNA} = \text{C DNA} \times 100 \text{ (copies/ml)}$$

C DNA is the number of *EBV* DNA copies, or the number of *CMV* DNA copies, or the number of *HHV6* DNA copies in DNA sample.

Table 2. Example of Qualitative Analysis

Ct limits			
IC	EBV	CMV	HHV6
28	35	35	35

No.	Description	Fam (IC)	Joe (EBV)	Rox (CMV)	Cy5 (HHV6)	Result	EBV	CMV	HHV6
	Name	Ct	Ct	Ct	Ct				
1	344	27,18			28	HHV6	-	-	+
2	445	26,41		34,12	32,1	CMV, HHV6	-	+	+
3	451	29,81				Invalid	?	?	?
4	456	23,3	28,48		27,7	EBV, HHV6	+	-	+
5	461	29,02		35,08		Invalid-?, (low CMV)	?	low	?
6	472	24,83	33,28			EBV	+	-	-
7	477	17,51	24,06		34,95	EBV, HHV6	+	-	+
8	489	21,32	21,85		27,2	EBV, HHV6	+	-	+
9	491	23,47	28,15			EBV	+	-	-
10	494	29,88				Invalid	?	?	?
11	497	16,29	31,06		34,18	EBV, HHV6	+	-	+
12	501	18,5		32,64		CMV	-	+	-
13	C+	27,23	30,18	28,47	27,25	OK			
14	C+	26,06	30,45	27,95	26,58	OK			
15	C+	26,37	30,8	28,17	26,73	OK			
16	C- (Neg. Control)					OK			
17	C- (DNA-buffer)					OK			
18	C- (DNA-buffer)					OK			

QUALITY CONTROL PROCEDURE

CMV/EBV/HHV6 Quant Real-TM PCR contains the Internal Control IC (human beta-globine gene), which allows to control the presence of cellular material in the sample. If the sample is not correctly prepared or it is an insufficient quantity of epithelial cells the Internal Control will not be detected.

A negative control of extraction (NCE), negative amplification control (NCA), positive amplification control (C+) are required for every run to verify that the specimen preparation, the amplification and the detection steps are performed correctly.

If the controls are out of their expected range (see table Results for Controls), all of the specimens and controls from that run must be processed beginning from the sample preparation step.

TROUBLESHOOTING

Results of analysis are not taken into account in the following cases:

1. The presence of any Ct value on JOE/Yellow/HEX, FAM/Green, ROX/Orange and Cy5/Red channels in the results grid for the Negative Control of Amplification (NCA) and for the Neg. Control of Extraction (C-) indicates contamination of reagents or samples. In this case, PCR analysis should be repeated for all samples in which pathogen DNA was detected starting from the DNA extraction stage.
2. For qualitative analysis, if the Ct value in the results grid for the Positive Control of PCR on the JOE/Yellow/HEX, FAM/Green, ROX/Orange, or Cy5/Red channels is absent, it is necessary to repeat amplification for all samples where pathogen DNA was not detected.
3. If the Ct value for the sample is not detected on JOE/Yellow/HEX/Cy3, ROX/Orange/TexasRed, Cy5/Red channel or it exceeds the boundary Ct value specified in the Data Sheet and the Ct value for the sample is greater than the maximum Ct value for IC in the FAM/Green channel, analysis should be repeated starting from the DNA extraction stage. This error may be caused by incorrect treatment of clinical material, which resulted in the loss of DNA, or by the presence of PCR inhibitors.
4. If the Ct value for the sample is detected in JOE/Yellow/HEX/Cy3, ROX/Orange/TexasRed or Cy5/Red channel and it is greater than the boundary Ct value specified in the Data Sheet, the result is considered to be equivocal. It is necessary to repeat analysis of such sample in duplicate. If a reproducible positive Ct value is obtained, the result is considered to be positive; otherwise, the result is considered to be equivocal.

PERFORMANCE CHARACTERISTICS

Sensitivity

The analytical sensitivity of **CMV/EBV/HHV6 Quant Real-TM** PCR kit is specified in the table below.

Type of clinical material	Nucleic acid extraction kit	Sensitivity
Cerebrospinal fluid (liquor), saliva, oropharyngeal swabs, and lavages	DNA/RNA-Prep	400 copies/ml
Whole human blood, white blood cells, viscera biopsy material	DNA/RNA-Prep	5 DNA copies per 10 ⁵ cells

Specificity

CMV/EBV/HHV6 Quant Real-TM PCR kit is intended for *Epstein-Barr virus (EBV)* DNA, *Human Herpes Virus type 6 (HHV6)* DNA and *human cytomegalovirus (CMV)* DNA detection. Specific activity of **CMV/EBV/HHV6 Quant Real-TM** PCR kit was confirmed by analysis of reference *CMV* strain AD 169, QCMD panel for *Epstein-Barr virus*, as well as by analysis of clinical material with subsequent confirmation of results by sequencing the amplified fragments. The activity of the PCR kit components with respect to DNA of other viruses (herpes simplex virus types 1 and 2, human herpes virus type 8, Varicella Zoster Virus, Parvovirus B19, and others), bacterial pathogens (*Staphylococcus aureus*, *Streptococcus pyogenes*, *Streptococcus agalactiae*, and others) and human DNA was absent. The clinical specificity of **CMV/EBV/HHV6 Quant Real-TM** PCR kit was confirmed in laboratory clinical trials.

Target region: CMV – MIE, EBV – LMP, HHV6 – pol gene

REFERENCES

- PCR detection of cytomegalovirus DNA in serum as a diagnostic test for congenital cytomegalovirus infection. C T Nelson, A S Ista, M K Wilkerson, and G J Demmler. *J Clin Microbiol.* 1995 December; 33(12): 3317–3318.
- Detection of Cytomegalovirus DNA in Peripheral Blood of Patients Infected with Human Immunodeficiency Virus. D. Shibata, W. John Martin, Maria D. Appleman, Dennis M. Causey, J. M. Leedom, N. Arnheim. *J Infect Dis.* (1988) 158 (6): 1185-1192.
- Multiplex PCR for six herpesviruses after hematopoietic stem cell transplantation. Sawada A, Koyama-Sato M, Yasui M, Kondo O, Ishihara T, Takeshita Y, Okamura T, Nishikawa M, Inoue M, Kawa *Pediatr Int.* 2011 Aug 2. doi: 10.1111/j.1442-200X.2011.03437.
- Cytomegalovirus Infections in Non-immunocompromised and Immunocompromised Patients in the Intensive Care Unit. Florescu DF, Kalil AC. *Infect Disord Drug Targets.* 2011 Jun 16.
- Comparison of PCR, Antigenemia Assay, and Rapid Blood Culture for Detection and Prevention of Cytomegalovirus Disease after Lung Transplantation. Adriana Weinberg, Tony N. Hodges, Shaobing Li, Guanyung Cai, M. R. Zamora. *Journal of Clinical Microbiology*, February 2000, p. 768-772, Vol. 38, No. 2
- Optimization of Quantitative Detection of Cytomegalovirus **DNA in Plasma by Real-Time PCR**. Michael Boeckh, MeeiLi Huang, James Ferrenberg, Terry Stevens-Ayers, Laurence Stensland, W. Garrett Nichols, and Lawrence Corey. *Journal of Clinical Microbiology*, March 2004, p. 1142-1148, Vol. 42, No. 3
- Quantification of Human Cytomegalovirus DNA by Real-Time PCR. Elyanne Gault, Yanne Michel, Axelle Dehé, Chahrazed Belabani, Jean-Claude Nicolas, Antoine Garbarg-Chenon. *J Clin Microbiol.* 2001 February; 39(2): 772–775
- Definitions of Cytomegalovirus Infection and Disease in Transplant Recipients. Per Ljungman, Paul Griffiths, Carlos Paya, ...*Clin Infect Dis.* (2002) 34 (8): 1094-1097

KEY TO SYMBOLS USED



List Number



Lot Number



For *in Vitro* Diagnostic Use



Store at



Manufacturer



Consult instructions for use



Expiration Date



Caution!



Contains sufficient
for <n> tests



Version

NCA

Negative Control of
Amplification

C-

Negative control of
Extraction

C+

Positive Control of
Amplification

IC

Internal Control

*iQ5™ is a registered trademark of Bio-Rad Laboratories

*Rotor-Gene™ Technology is a registered trademark of Qiagen

*MX3005P® is a registered trademark of Agilent Technologies

*ABI® is a registered trademark of Applied Biosystems

*SaCycler™ is a registered trademark of Sacace Biotechnologies

NOTE



Sacace Biotechnologies Srl

*via Scalabrini, 44 – 22100 – Como – Italy Tel +390314892927 Fax +390314892926
mail: info@sacace.com web: www.sacace.com*