Minimum detection limit of an in-house nested-PCR assay for herpes simplex virus and varicella zoster virus


Introduction: Herpes simplex virus (HSV) and varicella zoster virus (VZV) are responsible for a variety of human diseases, including central nervous system diseases. The use of polymerase chain reaction (PCR) techniques on cerebrospinal fluid samples has allowed the detection of viral DNA with high sensitivity and specificity. Methods: Serial dilutions of quantified commercial controls of each virus were subjected to an in-house nested-PCR technique. Results: The minimum detection limits for HSV and VZV were 5 and 10 copies/µL, respectively. Conclusions: The detection limit of nested-PCR for HSV and VZV in this study was similar to the limits found in previous studies.

Keywords: Herpes simplex virus. Varicella zoster virus. Polymerase chain reaction.

Herpes simplex virus type 1 and type 2 (HSV-1, HSV-2) and varicella zoster virus (VZV), which belong to the Herpesviridae family, are morphologically very similar, with large double-stranded linear deoxyribonucleic acid (DNA) genomes. These viruses are associated with a variety of clinical symptoms, including central nervous system (CNS) diseases such as meningitis, severe encephalitis, skin and mucosal infections and keratoconjunctivitis. In immunocompromised patients, the viruses may also trigger severe clinical conditions, including pneumonia and disseminated infection.

The prevalence rates of HSV-1 and 2 vary widely between and within countries; however, it is well known that HSV is highly prevalent in the general population. Both viruses are transmitted through contact with infected skin during viral reactivations. Most commonly, HSV-1, or oral herpes, causes sores around the mouth and lips (also called fever blisters or cold sores) and can infect the CNS. Although HSV-1 can cause genital herpes, most cases of genital herpes are caused by HSV-2. Unlike HSV-1 infection, which usually establishes latency in the trigeminal ganglion, typically causing an encephalitic illness, HSV-2 usually establishes latency in the sacral sensory ganglia and typically causes a meningitic illness. HSV-2 sometimes causes sores around the genitals or rectum. Although HSV-2 sores may occur in other locations, they are usually found below the waist.

Primary VZV infection causes varicella or chicken pox, a frequent illness affecting health professionals and the general population alike. Varicella is considered to be a highly infectious disease with high incidence among children. VZV can also reactivate from its latent state to cause herpes zoster or shingles, which is more common in the elderly. CNS complications can follow both primary infection and reactivation of VZV as meningitis and encephalitis. The more severe manifestations arise when VZV invades the spinal cord or cerebral arteries after viral reactivation, causing diseases such as myelitis and focal vasculopathies.

Before the introduction of molecular techniques, laboratory diagnosis of viral infections of the CNS, such as meningitis caused by HSV or VZV, relied on virus isolation in cell culture, detection of specific antibody production in cerebrospinal fluid (CSF) or, for encephalitis caused by HSV, viral antigen detection in brain biopsy specimens. The clinical usefulness of viral culture of CSF for the detection of HSV and VZV is limited because of poor sensitivity. Serological techniques have no role in the diagnosis of active HSV or VZV diseases such as genital infection or encephalitis. Polymerase chain reaction (PCR) protocols (commercial or in-house assays) have emerged as a tool for the rapid and accurate diagnosis of CNS infections. The sensitivity of PCR for the diagnosis of CNS disease is high for HSV encephalitis and enteroviral meningitis, increasing our understanding of the etiological roles of viruses in CNS disease.

Many molecular biology laboratories use in-house PCR assays – techniques specifically developed by laboratories for the diagnosis of infectious diseases. However, the lack of standardization and the poor reproducibility of in-house techniques have been argued as limitations against their routine use as diagnostic methods. When developing in-house PCR assays, one of the parameters to be analyzed as part of standardization studies is the minimum detection limit.

In this context, the purpose of this study was to determine the detection threshold of a duplex nested-PCR technique for...
HSV and VZV that was standardized by the Molecular Biology Laboratory of the Hospital de Clínicas de Porto Alegre.

Serial dilutions of viral DNA of known concentrations (1.3 x 10⁴ and 1.7 x 10⁴ copies/µL) were prepared from quantified HSV and VZV commercial controls (Advanced Biotechnologies®, Columbia, USA), respectively. Nine-fold dilutions of each virus (1,000, 500, 250, 100, 75, 50, 10, 5 and 1 copy/µL) were subjected to nested-PCR. Extraction and purification of viral nucleic acids were performed using a QIAamp Viral RNA Mini Kit (Qiagen®, Valencia, USA) according to the manufacturer’s instructions. The dilutions were then submitted to an in-house nested-PCR assay. This technique uses two stages of consecutive amplification, with the products of the first reactions (413bp for HSV and 272bp for VZV) serving as the targets for the second (280bp for HSV and 208bp for VZV). That is, in this assay, second amplification primers lie (are nested) within the first amplification primers. HSV and VZV external (HSV: 5' ATC CGA ACG CAG CCC CGC TG and 5' TCC GGS GGC AGC AGG GTG CT; VZV: 5' ACG GGT CTT GCC GGA GCT GTT and 5' AAT GCC GTG AGG GTG CT; HSV: 5' GCG CCG TCA GCG AGG ATA AC and 5' AGC TGT ATA SGG CGA CGG TG; VZV: 5' ACC TTA AAA CTC ACT ACC AGT and 5' CTA ATC CAA GGC GGG TGC AT) and internal primers (HSV: 5' GCG CCG TCA GCG AGG ATA AC and 5' AGC TGT ATA SGG CGA CGG TG; VZV: 5' ACC TTA AAA CTC ACT ACC AGT and 5' CTA ATC CAA GGC GGG TGC AT) amplify fragments of the D glycoprotein gene and gene 29 that are present in HSV and VZV genomes, respectively. The first-stage PCR reaction contained 16mM (NH₄)₂SO₄, 67mM Tris-HCl (pH 8.8, 25°C), 1.5 mM MgCl₂, 0.01% (w/v) Tween-20 (Southern Cross Biotechnology Ltd., Cape Town, South Africa), 0.25mM of each dNTP, 0.2μM primers and 2.5U DNA Super-Therm polymerase (Southern Cross Biotechnology Ltd., Cape Town, South Africa). The final volume of the reaction mixture was 50μL and contained 20μL of DNA extracted from the commercial controls. Amplification was conducted in a Techne Thermocycler TC414 (Techne®, New Jersey, USA). The reaction began with denaturation at 94°C for 45s, followed by 33 cycles at 94°C for 30s, annealing at 60°C for 30s and polymerization at 72°C for 30s. In the second amplification reaction, the reaction mixture was identical to the first; however, half of the individual volumes of the reagents were used, and 2μL of the amplicon obtained in the first reaction was added. Additionally, the internal instead of the external primers were used. The reaction began with denaturation at 94°C for 45s, followed by 33 cycles at 94°C for 20s, annealing at 60°C for 30s and polymerization at 72°C. The final products of these reactions were electrophoresed on 2% agarose gels stained with 0.5μg/mL ethidium bromide. All testing was performed with 100- to 1,000-bp molecular weight markers (Invitrogen®, Carlsbad, USA). As a negative control, a reaction containing 20μL of sterile water was used instead of the commercial control dilutions. The results were qualitatively expressed as positive (amplified DNA product) or negative.

For the statistical analysis, WinPepi® (version 10.5) was used to calculate the number of PCR assay repetitions. For a sensitivity of 90%, with a 10% error margin (assuming a significance level of 95%), it was necessary to perform the PCR assay at least 33 times for each dilution. The lower limit of detection was defined as the lowest concentration of viral DNA yielding 100% positivity in all repetitions. For testing categorical variables with PCR results, Fisher’s exact test was used with a 95% confidence interval.

First, all nine dilutions (1 to 1,000 copies/µL) were tested for HSV to determine the lowest range of dilution for further testing. Because amplification of the target product was observed with both smaller and larger dilutions, it was established that the assay’s minimum detection limit would most likely be between 1 and 10 copies/µL. Therefore, at least 33 repetitions of the nested PCR assay were performed for each virus within a range of higher dilutions, with 20μL of extracted DNA. The lower limits of detection were determined to be 5 copies/µL for HSV and 10 copies/µL for VZV (Table 1). To verify whether the PCR protocol was specific for HSV and

<table>
<thead>
<tr>
<th>Viral DNA dilutions</th>
<th>PCR repetitions</th>
<th>positive amplifications n</th>
<th>%</th>
<th>PCR repetitions</th>
<th>positive amplifications n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 copy/µL</td>
<td>33</td>
<td>26</td>
<td>78.7</td>
<td>33</td>
<td>11</td>
<td>33.3</td>
</tr>
<tr>
<td>5 copies/µL</td>
<td>33</td>
<td>33</td>
<td>100.0</td>
<td>33</td>
<td>24</td>
<td>72.7</td>
</tr>
<tr>
<td>10 copies/µL</td>
<td>36</td>
<td>36</td>
<td>100.0</td>
<td>33</td>
<td>33</td>
<td>100.0</td>
</tr>
<tr>
<td>50 copies/µL</td>
<td>9</td>
<td>9</td>
<td>100.0</td>
<td>1</td>
<td>1</td>
<td>100.0</td>
</tr>
<tr>
<td>75 copies/µL</td>
<td>9</td>
<td>9</td>
<td>100.0</td>
<td>1</td>
<td>1</td>
<td>100.0</td>
</tr>
<tr>
<td>100 copies/µL</td>
<td>16</td>
<td>16</td>
<td>100.0</td>
<td>1</td>
<td>1</td>
<td>100.0</td>
</tr>
<tr>
<td>250 copies/µL</td>
<td>7</td>
<td>7</td>
<td>100.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>500 copies/µL</td>
<td>3</td>
<td>3</td>
<td>100.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>1,000 copies/µL</td>
<td>3</td>
<td>3</td>
<td>100.0</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

HSV: herpes simplex virus; VZV: varicella zoster virus; DNA: deoxyribonucleic acid; PCR: polymerase chain reaction.
VZV, we performed a single PCR assay using DNA extracted from HSV and VZV and from pathogens commonly isolated in CSF samples that may be involved in CNS infections, including *Mycobacterium tuberculosis*, cytomegalovirus, Epstein-Barr virus, human herpes virus 6, polyomavirus JC and BK, adenovirus, enterovirus and *Toxoplasma gondii*. The HSV and VZV primers used yielded specific fragments, confirming the specificity of the assay. HSV- (280bp) and VZV-specific bands (208bp) were compared with a molecular weight marker (MWM), as illustrated in Figure 1.

The application of molecular biology techniques has led to a revolution in the laboratory diagnosis of many infectious diseases of the CNS, as these techniques have high sensitivity and specificity. PCR has been extensively used in the diagnosis of CNS infections for its speed and capacity to detect small amounts of target DNA in the CSF. Molecular analysis using PCR methods can be performed using commercial assays or in-house methods – in other words, lower-cost, laboratory-standardized alternatives. Assays can be developed in-house by creating a multiplex panel containing multiple pairs of primers directed at various target genes, improving screening. However, there is wide intra-laboratory variability in PCR assays, resulting from the use of different brands of reagents (enzymes and primers) and laboratory equipment (pipettes and cyclers) and inter-operator variability. The standardization of the technique is crucial, including the determination of the lower limit of detection.

In this study, we observed that the lower limit of detection for the proposed PCR assay was 5 copies/µL for HSV and 10 copies/µL for VZV, indicating high analytical sensitivity. In a previous study comparing the detection limits of nested-PCR and real-time polymerase chain reaction (RT-PCR) for HSV, 100% detection was reported for both methods for dilutions between 55 and 60 viral copies per reaction (2.75 - 3 copies/µL). In addition, for reactions with dilutions between 2.5 and 12 copies, 50% were positive by RT-PCR, and 12.5% were positive by nested-PCR. Furthermore, Tanaka et al. developed a multiplex-PCR assay for rapid and simultaneous detection of six types of human herpes viruses, including HSV and VZV. In that study, the PCR assay had a lower detection limit of approximately 20 viral copies per reaction (0.8 copies/µL) for each type of virus. In this regard, there is a small variation of sensitivity in different studies, which can be explained by several factors, such as the sequence of primers chosen, enzyme activity, DNA extraction method and the detection method used.

The introduction of molecular techniques, such as PCR, and the adoption of optimized protocols for the detection of infectious diseases are fundamental measures for the diagnosis of CNS infections. Comparing the results obtained in the literature with the results obtained in this study, it is possible to affirm that a lower limit of detection was detected for the proposed PCR assay for HSV and VZV, demonstrating that the protocol is highly sensitive. Determining the lower limits of detection of PCR assays and following quality assurance guidelines are key to ensuring the reliability of the results.

**CONFLICT OF INTEREST**

The authors declare that there is no conflict of interest.

**FINANCIAL SUPPORT**

We thank the Research Incentive Fund (*Fundo de Incentivo à Pesquisa e Eventos* - FIPE), the Graduate Research Group (GPPG) and Hospital de Clínicas de Porto Alegre.
# REFERENCES


