Traumatic Lumbar Puncture is Unlikely to Reduce Modern Molecular Detection of HSV Encephalitis (HSVE)

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Abstract

Polymerase chain reaction (PCR) is the gold-standard for diagnosing HSVE, but may be inhibited by blood contamination. We mimicked traumatic lumbar puncture and measured its effect on the molecular detection of HSV-1 in CSF. Clinically meaningful reductions in sensitivity were not observed. Pre-PCR processing allows sensitive detection of HSVE despite traumatic lumbar puncture.

Keywords: Traumatic lumbar puncture; Traumatic tap; HSV; HSV encephalitis; PCR inhibition

Introduction

The detection of HSV DNA in the cerebrospinal fluid (CSF) by PCR is the gold standard for diagnosing HSVE [1,2], which can produce very small quantities of DNA. Therefore, PCR detection must be extremely sensitive. Overall sensitivities and specificities of CSF PCR in neonatal HSV disease have ranged from 71-100% [1]. Even higher sensitivities and specificities (98 and 94%, respectively) of the assay have been reported in adults with HSVE when compared to brain biopsy [2], and sensitivity of the assay is maintained up to a week after initiating antiviral therapy [2,3].

The sensitivity of real-time PCR is affected by the quality of target nucleic acids, reaction conditions, and the presence of extraneous matter [4]. Amplification can be inhibited by contaminants present in blood, including haemoglobin, lactoferrin, and immunoglobulin's [5,6]. Taq and AmpliTaq Gold DNA polymerases (utilized widely in Hot Start/Fast Start "master mix" real-time PCR protocols) are inhibited by as little as 14,000 RBC/mm³ [6]. Heme, particularly the degenerated heme complex, is considered a universal PCR inhibitor. Release of iron ions from heme contaminants affects ion balance and disturbance of DNA polymerase activity and primer/probe annealing [6]. Hemin is a competitive inhibitor of target DNA, and immunoglobulin G may bind directly to ssDNA, further inhibiting nucleic acid amplification [5].

Modern real-time PCR molecular diagnostic assays typically attempt to avoid these inhibitory effects by processing techniques applied prior to amplification, detection and quantification of the PCR product [4]. Specimen processing techniques include cell separation and removal of visible RBC's from CSF by centrifugation. Extraction techniques concentrate nucleic acids and remove/neutralize PCR inhibitors. Column-based solid-phase (DNA manual mini-kits) or automated (QIAGEN EZ-1 or NucliSENS EasyMAG) magnetic resin/silica DNA extraction and purification methods are effective and commonly employed [6]. Furthermore, PCR chemistry has been optimized through utilization of specific buffers, pH, salt conditions, and amplification facilitators (bovine serum albumin is used to bind heme; DMSO is used to destabilize DNA) [6].

Traumatic lumbar punctures occur frequently, approaching 14-40% [7-9]. The visible threshold of detection (CSF visibly pink tinged) is defined as >400 RBC/mm³. Published ranges of blood within the CSF after traumatic LPs include 1,000-23,000 RBCs/mm³ (median 3,000-4,000 RBCs/mm³) [9]. Viral-induced haemorrhage into the CSF contributes to this contamination [10].

Newer molecular diagnostic techniques including pre-PCR processing may reduce PCR inhibition by blood products. Within the CSF, the clinically applicable limitations of sensitivity by blood products on real-time PCR have not been evaluated. We mimicked traumatic lumbar puncture and studied the effects of various clinically-relevant concentrations of human blood on the quantitative real-time PCR detection of HSV-1 in human CSF.

Materials and Methods

Human CSF was obtained and pooled from patients undergoing routine ventriculoperitoneal shunt placement. It was determined free of RBC's and detectable HSV virus within the clinical laboratory by automated CBC technology and real-time PCR.

Sample treatment techniques prior to Real-time PCR

Normal pooled human CSF was spiked with clinically-relevant HSV-1 concentrations (Figure 1) derived from human fetal lung fibroblast (MRC-5) tissue culture. For experimental condition A (EC-A), whole blood collected in EDTA and serially diluted in blood buffer (CELL-DYN Sapphire, Germany) was then added to this spiked CSF to produce final concentrations of 200,000 to 0.2 RBC/mm³ (Figure
For experimental condition B (EC-B), samples were prepared as EC-A but followed by 30 s of mini-centrifugation and removal of supernatant. For experimental condition C (EC-C), whole blood was haemolysed (frozen on dry ice for 10 min followed by 30 s of vortexing and repeated three times). This haemolysed blood was then added to spike CSF followed by 30 s of mini-centrifugation for each prepared specimen (similar to EC-B). A positive control included pooled human CSF spiked with HSV-1 and RBC diluent blood buffer. Negative controls included: (1) pooled human CSF, and (2) RBC diluent blood buffer.

![Figure 1: The solid bars represent viral load data from patients with HSV-1 induced encephalitis diagnosed from 2004-2013. The solid line is the computed normalized distribution of the viral loads. The maximum lowering of measured viral load (0.53 log10 copies/mL) contributed experimentally by 2,000 RBC/mm$^3$ (inset*) could potentially render a false negative test in only 1-2 of 28 (3.6-7.1%) of our patient population with HSVE. *Inset: pooled CSF containing various experimentally-added concentrations of non-haemolysed whole blood (RBC/mm$^3$).](image)

HSV-1 viral DNA was extracted from 200 μL of these processed samples utilizing the automated Biorobot EZ-1 instrument/Virus Mini Kit v2.0 (QIAGEN, Germantown, Maryland, USA) and eluted to 60 μL of sterile water. The concentration within the CSF of HSV-1 spiked into samples was chosen to mimic the mean viral load in clinical patients with HSV-1 encephalitis, generated by the same real-time PCR utilized for this experimental study.

Detection and quantification of HSV-1 DNA by Real-time PCR

Detection of extracted HSV-1 DNA pol conserved gene segment was carried out by the Light Cycler 2.0 (Roche, Indianapolis, IN, USA) via real-time PCR utilizing a home-brew assay based on hybridization probe technology. As previously described [11], forward and reverse primer HSV polymerases were used to amplify a 140 base-pair product. Real-time detection and melting temperature differentiation of HSV-1 and 2 was determined with a pair of fluorescence-labelled probes. The reaction mixture was prepared in a controlled access reagent preparation room and consisted of 15 μL of PCR master mix and 5 μL of DNA extract per CSF sample. Cycling conditions and melting curve analysis after amplification were then determined [11]. Samples were run in duplicate with a negative and positive control, and were considered positive if both replicates had Ct values <40 and negative if there was no amplification within 40 amplification cycles.

Quantification was based on external standard curves made from highly-concentrated, purified and deactivated whole HSV virus isolated from cell culture and quantified via pico green (Zeptometrix, Buffalo, NY, USA).

Statistical methods

After testing for normality of distribution, the Wilcoxon Rank Sum (Mann-Whitney U) test compared median viral load (log10 copies/mL) between experimental groups and control (CSF + HSV, no blood). Two-tailed p values <0.05 were considered statistically significant.

Human subjects protection

IRB approval was obtained for all studies as appropriate.

Results

Despite the addition of widely differing concentrations of haemolysed and non-haemolysed blood and the application of different processing techniques (the absence of processing v/s sample centrifugation prior to extraction), no blood concentration significantly affected PCR detection of HSV-1 DNA in CSF compared to control (CSF+HSV without blood) [Mean 4.60 log10 copies/mL (CI: 4.04-5.17)], p all >0.05].

A trend toward reduction in the quantitative detection of HSV-1 occurred in the HSV spiked CSF containing 2,000 RBC/mm$^3$ of haemolysed blood (EC-C) (Figure insert*). However, this did not reach statistical significance [maximum lowering of measured viral load was 0.53 log10 copies/mL of HSV DNA, (p all >0.05)]. When applied to a calculated normalized distribution of the clinical mean CSF viral load from our 28 patients with HSVIE quantitatively detected by real-time PCR in our laboratory from 2004-2013 (Figure 1), this maximum lowering of measured viral load could potentially render a false negative test in only 1-2 of 28 (3.6-7.1%) patients with HSVE.

Discussion

Traumatic LP occurs often [8,9], and haemorrhagic CSF, secondary to viral-induced necrosis in patients with HSVE, is well described [10]. Most real-time PCR assays, including ours, utilize commercially available TaqDNA polymerases, which are sensitive to heme-induced inhibition [4-6]. Thus, blood contamination interferes with PCR amplification. Whether this phenomenon occurs to a clinically meaningful extent using modern PCR processing techniques and PCR assays has not been evaluated. We demonstrated that despite the phenomenon of PCR inhibition by blood contaminants cited by manufacturers, modern molecular diagnostic pre-PCR processing techniques, most importantly improved extraction capabilities and optimal PCR chemistry, likely overcome the previously-observed PCR inhibition by blood products. This allows the sensitive detection of HSVE despite traumatic lumbar puncture.

In an effort to optimize sensitivity of molecular diagnostic assays, sample-processing techniques of visibly bloody CSF specimens, including centrifugation prior to extraction, are widely practiced. But some degree of limitation of generalizability of these results warrants discussion. Performance of PCR varies amongst laboratories given the lack of inter-laboratory quantitative and non-quantitative standards and the use of single-lab ("home-brew") developed assays [6]. Reagent
buffers and amplification facilitators utilized to provide optimal PCR chemistry may also vary. However, results of our study should not undermine the importance of modern extraction procedures, in addition to PCR chemistry, employed to reduce such theoretical inhibition.

Modern extraction techniques are widely utilized [6] and are equally effective [12]. Manual column-based solid-phase mini-kits or the automated biorobots, which use magnetic resin/silica for DNA extraction and purification, have replaced older Chelex 100 resin or organic phenol chloroform extraction techniques, providing concentrated and more highly purified DNA given their ability to more effectively adsorb water soluble complexes of ferric heme and serum proteins [6]. Modern pre-PCR processing techniques may not be standardized, but optimal chemistry and extraction is generally utilized in commercial kits and assays, allowing sensitive detection of HSV in CSF despite the presence of blood products.

The morbidity and mortality of HSVE remains unacceptably high. Up to 69% of neonates and 44-62% of children and adults will suffer long-term neurological impairment despite treatment (acyclovir) [10,13]. Early initiation of acyclovir and treatment until the cerebrospinal fluid viral load becomes undetectable is prudent as rapid reduction and suppression of viral replication is necessary to improve clinical outcomes in patients with HSVE [1,14]. Quantitative real-time polymerase chain reaction (PCR) molecular techniques enable earlier detection and initiation of therapy for HSVE before widespread dissemination of virus and allow the clinician to potentially monitor response to treatment. Our study has demonstrated that despite traumatic lumbar puncture, the essential sensitivity of PCR in CSF is maintained in patients with HSVE.

Conclusions

Advanced pre-PCR processing techniques including sample treatment (extraction) and optimal PCR chemistry appear to overcome the previously observed PCR inhibition by blood products allowing sensitive detection of HSVE despite traumatic lumbar puncture.

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References