Enteroviral and Herpes Simplex Virus Central Nervous System Infections in Infants <90 Days Old: A Paediatric Investigators’ Collaborative Network on Infections in Canada (PICNIC) Study

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Research article
Abstract

Background: The relative contribution of viruses to central nervous system (CNS) infections in young infants is not clear. For viral CNS infections, there are limited data on features that suggest HSV etiology or on predictors of unfavorable outcome.

Methods: In this cross-sectional retrospective study, seven centers from the Pediatric Investigators Collaborative Network on Infections in Canada identified infants <90 days of age with CNS infection proven to be due to enterovirus (EV) or herpes simplex virus (HSV) January 1, 2013 through December 31, 2014.

Results: Of 174 CNS infections with a proven etiology, EV accounted for 103 (59%) and HSV for 7 (4%). All HSV cases and 41 (40%) EV cases presented before 21 days of age. Four HSV cases (57%) and 5 EV cases (5%) had seizures. Three (43%) HSV and 23 (23%) EV cases lacked cerebrospinal fluid (CSF) pleocytosis. HSV cases were more likely to require ICU admission (p=0.010), present with seizures (p=0.031) and have extra-CNS disease (p<0.001). Unfavorable outcome occurred in 12 cases (11% of all EV and HSV infections) but was more likely following HSV than EV infection (4 (57%) versus 8 (8%); p=0.002).

Conclusions: Viruses accounted for approximately two-thirds of proven CNS infections in the first 90 days of life. Empiric therapy for HSV should be considered in suspected CNS infections in the first 21 days even in the absence of CSF pleocytosis unless CSF parameters are suggestive of bacterial meningitis. Neurodevelopmental follow-up should be considered in infants whose course of illness is complicated by seizures.

Background

The prevention of bacterial meningitis by conjugate vaccines has resulted in viruses accounting for an increasing proportion of central nervous system (CNS) disease in childhood. Improvement in viral diagnostics has made this trend more apparent. Previous studies of viral CNS disease were limited by small sample size, included cases where the etiology was not proven or did not focus on infants. The most common viruses associated with CNS infections are enteroviruses (EV), which most frequently manifest as self-limited aseptic meningitis with no recognized long-term sequelae. By contrast, herpes simplex virus (HSV) CNS infections result in significant morbidity and mortality, especially if acyclovir therapy is delayed. It is therefore vital that clinicians know what clinical and laboratory features should prompt them to start empiric acyclovir.

This was a cross-sectional analysis to identify infants less than 90 days of age with proven CNS infections. This age range was selected as diagnosis of CNS infections is particularly challenging in young infants. We sought to a) determine the relative contribution of HSV and EV to microbiologically-confirmed CNS infections, b) provide a comparative analysis of the epidemiology and outcome of HSV
and EV CNS infection, c) describe factors associated with HSV aetiology and d) identify factors associated with unfavorable outcome.

**Methods**

**Study Population and Design:**

Seven paediatric academic centres within the Paediatric Investigators Collaborative Network on Infections in Canada (PICNIC) retrospectively enrolled hospitalized infants <90 days of age with microbiologically-confirmed CNS infection January 1, 2013 through December 31, 2014. Cases were identified using appropriate discharge diagnostic codes from the International Statistical Classification of Diseases and Related Health Problems, Tenth Revision (ICD 10) (Appendix A) and charts were then reviewed. A previous publication described cases of bacterial CNS infection as proven if bacteria were detected from cerebrospinal fluid (CSF) or brain abscess by means of culture or PCR or probable if CSF pleocytosis was present, along with bacterial growth from another sterile site. For the purposes of this study, we included all proven cases of HSV or EV CNS infection based on the identification of a virus in the CSF by polymerase chain reaction (PCR) during life or in the brain tissue using PCR at autopsy. All study centres offered routine PCR testing for HSV and EV. None used multiplex PCR. Only two centres offered HPeV testing during the study period so it was not possible to compare HPeV cases to other viral cases. Cases with coinfection were excluded unless the investigator deemed that a virus was the main pathogen. There were no other exclusion criteria.

Ethics board approval was obtained from all participating centres with the primary approval coming from the Health Research Ethics Board of the University of Alberta (Study number PRO00055909).

**Study Definitions:**

1) Case classification: a) early onset if diagnosis was made within the first 6 days of life, b) late onset if diagnosis was made day 7 through 29 of life and c) very late onset if diagnosis was made day 30 through 90 of life.

2) Infants were considered to have extra-CNS disease if there was microbiological, clinical or other laboratory findings consistent with viral disease at other sites.

3) Infants who had i) seizures or ii) head imaging suggesting parenchymal involvement were presumed to have meningoencephalitis. All other cases were deemed to have meningitis.

4) Unfavourable outcomes were defined as:

- Neurodevelopmental sequelae (any one of hearing loss, visual impairment, other neurological sequelae such as extensive intracranial haemorrhage or hydrocephalus, or developmental delay noted at follow-up) OR
- Death
Data Collection and Analysis:

Demographic, clinical, microbiological, head imaging reports, treatment, outcome and any available follow-up data were extracted from medical records and entered into Research Electronic Data Capture (REDCap) by each participating center. Follow-up data were collected at variable time points depending upon local protocols and parental compliance with follow-up. Follow-up data were not available if the neonatal follow-up program was not in the institution where the infant was admitted.

Two separate although related comparative analyses were undertaken comparing clinical features and outcome by etiology (HSV versus EV). Descriptive analysis was conducted. Chi-square or Fisher’s exact test was used to compare categorical variables and non-parametric tests were used to compare continuous variables (Mann–Whitney U test). Exploratory analysis was conducted using univariate analyses and where sample size allowed, multivariate analyses to identify clinical, laboratory or outcome differences between EV and HSV cases were conducted using factors identified as significant in univariate analysis. Additionally, we used univariate analysis to explore potential factors associated with an unfavorable outcome overall. We adjusted for multiple comparisons using Bonferroni correction. Epi-info version 7 (Centers for Disease Control and Prevention) was used for statistical analysis.

Results

Relative contribution of viruses to microbiologically-confirmed CNS infections

There were 174 cases of proven CNS infections in infants <90 days old, of which 111 (64%) were viral in origin. One case was excluded due to coinfection with group B streptococcus and EV. The most common identified viral pathogen was EV (N=103; 93%) followed by HSV (N=7; 6%) and human parechovirus (HPeV) (N=1; 1%). The HSV cases included 3 with HSV1 (1 with isolated CNS disease, and 2 with disseminated disease) and 4 with HSV2 (1 with isolated CNS disease, and 3 with disseminated disease).

Descriptive analysis of EV and HSV CNS infections

Demographics: The median birth weight was 3343g (range 1670-4900g) and median gestational age was 37 weeks (range 29-40 weeks). Sixteen infants were preterm (15%). Infants presented at a median age of 22.5 days (range 3-84 days), with 5 cases occurring during the birth hospitalization (all were EV infection on day 3 to day 21 of life in infants born at 31 to 35 weeks GA). HSV cases presented earlier than EV cases (median 14 days versus 25 days of life; p=0.02) (Table 1). Fifty-two (50%) of EV cases and 3(43%) of HSV cases presented August through October (Fig 1).

Maternal History: Among HSV cases, three (38%) were born to mothers with active genital lesions documented at or within 7 days of delivery (Table 1). Data on the reasons for the mode of delivery were not collected. One of the three mothers had recurrent HSV1 disease and was not compliant with acyclovir prophylaxis; her infant presented with HSV1 on day 4 of life after vaginal delivery. The other two mothers had first clinical episode of genital HSV within 7 days of delivery. Their infants received no screening or
empiric treatment and presented with HSV2 on days 6 and 9 after caesarean and vaginal delivery, respectively; the duration of rupture of membranes was not available. For EV infection, 3 mothers had documented illness compatible with EV within 10 days prior to delivery; their infants presented on days 5, 6 and 8.

**Timing of Presentation:** Early-onset infection occurred in 10 of the 110 infants (8 EV and 2 HSV). The EV cases all presented after day 2 of life and 3 of the 8 had severe disease including: fatal myocarditis, shock with coagulopathy and meningoencephalitis. The two early-onset cases with HSV infection presented on days 4 and 6.

For late-onset disease, there were 63 cases of EV meningitis and 5 cases of HSV meningoencephalitis. All infants (N=32) with very late onset infection had EV. All HSV cases presented before 21 days of age.

**Clinical Features:** There were 9 (8%) cases with seizures. Eight had seizures during the admission for the CNS infection (5 with EV and 3 with HSV) and the ninth developed seizures after discharge coinciding with CNS HSV relapse. For the 4 HSV cases, 2 had seizures only within the first 72 hours following diagnosis, 1 after 72 hours but prior to hospital discharge and as mentioned previously, one case only after discharge. Five of 103 infants with EV had seizures (5%), with 3 presenting in the first 72 hours following diagnosis and 2 presenting after 72 hours but prior to hospital discharge. The age at diagnosis of CNS infection for these 5 cases was 5, 10, 10, 14 and 84 days.

There were 14 (12%) infants with extra-CNS involvement. Five infants had extra-CNS HSV infection, consisting of vesicular lesions without other extra-CNS involvement (N=1), transaminitis and pneumonitis (N=1), transaminitis and vesicles (N=1) and transaminitis, pneumonitis and coagulopathy (N=2). Coagulopathy was complicated by spontaneous intracranial haemorrhages (intraventricular and parenchymal) in one of these two infants. Extra-CNS manifestations in EV cases included rash (N=2), pneumonia (N=2), shock with coagulopathy (N=2), myocarditis (N=2) and transaminitis (N=1). The median age of onset of the 7 cases with organ involvement (omitting the 2 with skin involvement) was 9 days (range 5-73 days). Extra-CNS involvement was more likely in HSV than EV cases (p <0.001), even if skin involvement was not considered (4 (57%) versus 5 (6%); p=0.001) (Table 1).

**Microbiology:** All cases were diagnosed using PCR analysis of CSF. HSV PCR testing was also positive on skin lesions in two infants and from the conjunctiva of one (in the absence of ophthalmological abnormalities). EV typing was not available. Suspected urinary tract coinfections occurred in 4 infants with EV infection (Table 1). Systemic candidiasis complicated the course of one infant with HSV meningoencephalitis with liver failure, coagulopathy and intraventricular haemorrhages requiring external ventricular drain (EVD) placement. *Candida albicans* was isolated from blood and CSF obtained from EVD just prior to demise. Newborn screen, immunoglobulin assay and flow cytometry failed to identify an underlying immunodeficiency in this fatal case.
**CSF Findings:** The median values for cell count, glucose and protein on the initial CSF were not significantly different between HSV and EV (Table 2). Thirty-six (33%) infants (4 with HSV and 32 with EV) had CSF white blood cell (WBC) counts less than 30 $\times 10^6$/L. Notably, 5 (5%) of infants with EV infections had CSF WBC $>2000$ $\times 10^6$/L.

**Head Imaging:** Thirty-three (30%) of the 111 infants had head imaging performed (HSV (N=7) and EV (N=26)). Among the cases of HSV, magnetic resonance imaging was abnormal in 4/6 (67%) and appeared consistent with infection; the seventh case had only a head ultrasound which was normal. Among the cases of EV, 7/26 (27%) had abnormalities detected on imaging but only 5 (19%) of these were attributed to infection (diffusion restriction abnormalities).

**Meningoencephalitis:** Thirteen infants (HSV=6; EV=7) fulfilled the study criteria for meningoencephalitis. The EV cases presented at a median of 10 days of age (range 5 to 84 days). One case of disseminated HSV2 infection did not meet our definition of meningoencephalitis as the infant did not have documented seizures and only had a normal head ultrasound documented, but did not have MRI or CT imaging performed. Infants with meningoencephalitis were younger (p=0.012), more likely to require ICU admission (p<0.001), more likely to have disseminated disease (p=0.007) and more likely to die or have developmental delay (8 (62%) vs 4 (4%); P<0.001) than those without meningoencephalitis. Poor long term outcome in survivors with meningoencephalitis was equally likely whether the cause was HSV (3/5; 60%) or EV (4/7; 57%) (p=1.0). Adjusting for multiple comparisons, these associations remained significant.

**Antiviral Treatment and Prophylaxis:** All HSV cases received acyclovir treatment for a median of 21 days (range 21-51 days). One infant received acyclovir until demise on day 42 of acyclovir therapy. Acyclovir resistance was first tested for on a sample just prior to death and was proven to be present. Another infant did not have documented CSF clearance until 51 days of therapy. For the other 5 cases, repeat testing done between 19 and 22 days of treatment confirmed successful clearance of HSV from CSF. Three (50%) surviving infants were documented to have been discharged on oral acyclovir as prophylaxis for minimum 6 months.

**Outcome:** There were 2 deaths (2%), one from disseminated EV (a 6-day old infant with myocarditis who required extracorporeal membrane oxygenation) and one from HSV2 (the infant with systemic candidiasis and with persistent HSV detection in CSF until death at day 48 of life). Autopsies were not performed. Virologically-proven recurrence of HSV1 meningoencephalitis presenting as infantile spasms occurred in 1 (33%) of the 3 infants who received oral acyclovir until 6 months of life; this occurred 2 weeks after oral acyclovir was discontinued. Ten (9%) of the 108 surviving infants had neurodevelopmental sequelae documented at discharge or follow-up (Table 1). Neurodevelopmental outcomes were not available for infants who had HSV persistence documented in CSF as the single survivor was lost to follow-up. All 3 of the HSV (2 HSV2; 1 HSV) and 1 of the EV survivors with neurodevelopmental sequelae developed seizure disorders requiring anticonvulsant therapy. Overall, unfavorable outcome occurred in 12 cases (11% of all EV and HSV infections) but was more likely
following HSV than EV infection (4 (57%) versus 8 (8%); p=0.002) (Table 1). Three (75%) of four HSV cases with unfavorable outcome were caused by HSV2. One of 3 (33%) cases of HSV1 had poor outcome compared to 3 of 4 (75%) cases with HSV2. All cases of EV meningoencephalitis survived. There were no differences by pathogen in the incidence of poor neurodevelopmental outcomes in surviving infants with presumed encephalitis, (3/5 (60%) HSV vs 4/7 (57%) EV; p=1.0. Eight (62%) of 13 infants (HSV=4; EV=4) with meningoencephalitis had unfavorable outcome.

**Factors associated with HSV aetiology:** In univariate analysis, HSV cases were more likely than EV cases to require intensive care unit (ICU) admission (p=0.010), have seizures at any time (p=0.001), have extra-CNS disease (p<0.001) and have unfavorable outcome (p<0.001) (Table 1). The latter three remained significant after correcting for multiple comparisons. Seizures (p=0.005) and extra-CNS disease (p=0.002) remained significant after controlling for ICU admission.

Among infants <30 days of age (N=78), the presence of seizures or extra-CNS disease was more likely in HSV than in EV CNS infection (6 of 7; (86%) versus. 10 of 71; 14%); p<0.001).

**Factors associated with unfavorable outcome:**

In the univariate analysis, several factors were identified (Table 3). After adjusting for multiple comparisons, the factors associated with unfavorable outcome included younger age (p=0.003), HSV etiology (p=0.002), seizures (p<0.001), ICU admission (p<0.001) and meningoencephalitis (p<0.001) (Table 3). The latter 3 remained significant when analysis was limited to the subgroup of infants with EV CNS infections (Table 4 – The sample size limited multivariate analysis).

**Discussion**

Viral infections accounted for about two-thirds of CNS infections in the first 90 days of life where a CSF pathogen was detected in the current study. Trends in Canada are not clear but in a population-based United Kingdom (UK) study, the authors document a dramatic rise in admissions for viral meningitis in infants between 2005 and 2011. They show a major increase in the proportion of cases of viral meningitis recognized to be due to EV over time, from 90 (3%) of 2770 admissions for viral meningitis in 1968-1985 to 811 (47%) of 1716 viral meningitis admissions in 2007-2011. These changing trends probably reflect the UK adoption of molecular diagnostic screening for viral meningitis resulting in increased detection over conventional viral culture methods which were not consistently applied in earlier years. Further, molecular testing has facilitated the detection of viruses like HPeV that are missed by viral isolation techniques.

Consistent with prior literature, CNS infection with HSV was much more likely than infection with EV to lead to meningoencephalitis and long-term neurodevelopmental morbidity or death. However, among the subgroup of EV cases with meningoencephalitis, outcomes were comparable to cases of HSV...
meningoencephalitis. Identifying clinical or laboratory markers that distinguish HSV from non-HSV viral infections is vital to ensure that empiric acyclovir is started at presentation in all HSV cases. We identified younger age, seizures, ICU admission and the presence of extra-CNS features as factors associated with HSV infection; however, only seizures and extra-CNS disease remained significant in the multivariate analysis and 3 of 7 infants with HSV CNS disease (43%) did not have seizures. Most if not all HSV meningoencephalitis in the neonatal period comes from perinatal transmission, and in our study all presented by day 21 of life. There should be limited use of empiric acyclovir beyond the first month of life. However, HSV meningoencephalitis can present at any age and in a 2018 study of 46 cases up to 60 days of age, the IQR was 9 to 24 days.

Most genital HSV infections are subclinical. A small percentage of neonatal HSV cases may arise from post-natal transmission from saliva. Therefore, all infants should be assumed to be at risk of HSV infection irrespective of maternal history. In addition, as demonstrated in 4 of the 7 infants with CNS HSV in our cohort, the absence of CSF pleocytosis does not exclude CNS HSV infection. Furthermore, one case in our cohort had HSV detected by PCR on CSF analysis from day 5 of illness after a negative PCR on day 2 of illness. Thus, if the clinical picture is suggestive of HSV infection and initial CSF HSV testing returns negative, acyclovir should be continued until another CSF sample is retested to ensure that the original sample was not falsely negative. The potential value of repeating CSF analysis towards the end of treatment course is exemplified by the two cases with persistent detection of HSV in CSF, although there are not studies to prove that continuing intravenous acyclovir beyond the usual 21-day course improve prognosis.

As noted in our fatal HSV2 case, the possibility of acyclovir resistance should be considered in children with persistent detection of the virus in CSF. While rare, the possibility of acyclovir resistance needs to be kept in mind as alternative therapy including foscarnet or vidarabine may be of benefit.

A major limitation of our study was the retrospective design. Searching laboratory records rather than discharge codes might have identified more cases but was not practical at all sites. A surveillance program would be required to detect suspected in addition to proven cases. The lack of HPeV testing at most sites precluded study of this virus. It is likely that other viral etiologies of meningitis or meningoencephalitis will eventually be identified. Methods of molecular testing varied by study center. Emerging molecular diagnostic panels may eventually improve diagnosis of CNS infections. Infants with mild viral meningitis are not always recognized to have CNS infection. The total number of infants investigated at the 7 centers to yield the 174 with proven CNS infections is not known. The small sample size would have precluded detecting all differences in clinical presentation and outcome for EV versus HSV. Application of the International Encephalitis Consortium definition of encephalitis to infants is problematic as it can be difficult to determine if they have altered level of consciousness or focal signs and they are less likely than older children to manifest fever or CSF pleocytosis. An EEG is not always performed. Therefore, we used a simplified definition for meningoencephalitis; this definition was highly dependent upon the decision to perform and the interpretation of head imaging (which was not always
obtained) and recognition of seizures so could have missed or over-diagnosed cases. Rarely, aseptic meningitis can also result in seizures and head imaging abnormalities. Infants who had coagulopathy or were too systemically ill to have CSF obtained or who died before they had a diagnosis would have been missed. Molecular testing for HSV (and presumably for other viruses) can be falsely negative early in the course of infection. However, the inclusion of only proven cases was deemed to yield the most accurate data. There was inconsistent access to data on neurodevelopment follow-up and the timing and nature of this follow-up was not standardized between centers. Study results may not be applicable to resource poor settings.

**Conclusions**

Proven viral CNS infections appear to be more common than proven bacterial infections in the first 90 days of life. Age <21 days and presence of seizures or extra-CNS involvement are clues to HSV infection, even in the absence of CSF pleocytosis. However, not all infants with CNS HSV have seizures. Although most infants with EV CNS infections have good outcomes, the subset who have seizures and/or abnormal head imaging may have outcomes similar to those of infants with HSV meningoencephalitis and require neurodevelopmental follow-up. Further studies should address the contribution of HPeV to viral CNS infections and explore predictors of long-term morbidity.

**Abbreviations**

CNS – central nervous system

CSF – cerebrospinal fluid

EV – enterovirus

HSV – herpes simplex virus

IQR – interquartile range

WBC – white blood cell count

**Declarations**

- Ethics approval and consent to participate: Ethics approval was obtained at each site for conduct of this study with the primary approval coming from the Health Research Ethics Board of the University of Alberta (Study number PRO00055909). Parental consent was waived as it was a retrospective chart review.
- Consent for publication: Not applicable
- Availability of data and material: All data are stored in REDCap. An anonymized version is available from the corresponding author upon reasonable requests.
• Competing interests: Joseph Ting is an Associate Editor for *BMC Pediatrics*.

• Funding: No funding was obtained for this study.

• Authors' contributions: JR and MB wrote the first draft of the protocol, designed the case report form and finalized the manuscript. MB performed the data analysis. DP wrote the first draft of the manuscript. CR, LO, JB1, JB2, SK, AB, JM, AB, JT and AR provided input into the protocol, case report form or manuscript and organized data collection. All authors approved the final version.

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Tables

Table 1: Comparison of demographic, clinical and outcome features in infants with HSV and EV meningitis by univariate analysis
| Characteristics | HSV N=7 | EV N=103 | P-Value$^A$ |
|-----------------|--------|----------|-------------|
| Male gender, n (%) | 3 (43) | 54 (52) | 0.71 |
| Age at onset (d), median (IQR) | 14 (6-19) | 25 (12-33) | 0.02 |
| Overall subset fulfilling meningoencephalitis criteria [Proportion (%) age <28d] | 16 (6-19) | 10 (6-27) | 1.0 |
| [6/6 (100)] | [6/7 (86)] | 1.0 |
| Gestation (wks), median (IQR) | 37 (37-38) | 37 (29-38) | 0.31 |
| ICU admission, n (%) | 4 (57) | 12/98 (12) | 0.010 |
| Seizures, n (%) | 4 (57) | 5 (5) | .001* |
| Had at least one seizure at any time | 2 (29) | 3 (3) | 0.03 |
| 3(43) | 5 (5) | 0.008 |
| Had a seizure in first 72 hours of | 1(16) | 0 (0) | 0.06 |
| admission | Had at least one seizure during the admission | Had seizures only after discharge |
|-----------|---------------------------------------------|----------------------------------|
| Multisystemic infection** | 5 (71) | 8 (8) | <0.001* |
| Coinfection with bacteria or fungus, n (%) | 1D (14%) | 4E (4) | 0.28 |
| Abnormal head imaging**, n (%) | 4/7 (57) | 5/26 (19) | 0.068 |
| Meningoencephalitis, n (%) | 6/7 (86) | 7 (7)G | <0.001* |
| stay | Length of stay (d), median (IQR) | 25 (21-43) | 3 (3-5) | <0.001* |
| p | Follow-up (mo), median (IQR) | 16 (10-24) | 6 (1-12) | 0.03 |
| Neurodevelopmental | 3/6 (50%) | 7/102 | 0.01, n (%) |
| Abnormalities at discharge or follow-up<sup>H</sup> | | |
|---|---|---|
| Death<sup>I</sup> or neurological complications<sup>J</sup> or neurodevelopmental abnormalities, n (%) | 4/7 (57) | 8 (8) | 0.002* |
| All infants | 4/6 (67) | 4/7 (57) | 1.00 |
| Infants with encephalitis | | | |

Legend: CSF – cerebrospinal fluid; EV- enterovirus; HSV – herpes simplex virus; IQR – interquartile range; mo-months

For comparison of proportions, Fishers exact test (2-sided) was used; for comparison of medians, Mann-Whitney test was used.

These were identified as independent risk factors after controlling for age and ICU admission, respectively.

All infants with long-term seizures had neurodevelopmental delay (range mild to profound).

This infant had multiple blood and CSF samples positive for *C. albicans*, which developed in the latter part of therapy for disseminated HSV.

Infants received antimicrobial therapy for urinary tract infections (*E. coli* (N=2) and *E. faecalis* (N=2); the case of group B streptococcus coinfected with EV is not included.)
This comparison was limited to those abnormalities that were consistent with CNS infection.

All 7 cases with abnormal imaging or seizures had EV

4 of the 10 infants with neurodevelopmental abnormalities (3 with HSV and 1 with EV) required ongoing anticonvulsant therapy

One death was due to EV and one was due to HSV

After adjusting for multiple comparisons (Bonferroni correction), these variables remained significant.

Table 2: Comparison of initial cerebrospinal fluid findings in infants with HSV and EV central nervous system infections by univariate analysis
| CSF white blood cell (WBC) (x10^6/L) at diagnosis, median (IQR) | HSV N=7       | EV N=100^A | P-Value^B |
|---------------------------------------------------------------|---------------|------------|-----------|
| CSF WBC, n (%)                                                | 26 (2,146)    | 153 (17.5,422) | 0.08      |
| >9 x10^6/L                                                   | 4 (57)        | 79 (79)    | 0.18      |
| >15 x10^6/L                                                  | 4 (57)        | 76 (76)    | 0.37      |
| >100 x10^6/L                                                 | 3 (43)        | 57 (57)    | 0.70      |
| >1000 x10^6/L                                                | 0             | 22 (22)    | 0.34      |
| >1000 x10^6/L                                                | 0             | 10 (10)    | 1.00      |
| Median percentage polymorphonuclear contribution to CSF cell count | 6 (3,14)     | 25 (6,51)  | 0.04      |
| CSF pleocytosis^E                                            | 4 (57)        | 77 (77)    | 0.36      |
|                                | 0/7 (0)   | 23/86 (27) | 0.19 |
|--------------------------------|-----------|------------|------|
| CSF cell count with polymorphonuclear dominance (>50%), n (%) |           |            |      |
| CSF red blood cell count (x10^6/L), median (IQR) | 26 (1-6950) | 23 (2-138) | 0.66 |
| CSF glucose (mmol/L), median (IQR) | 2.4 (2.1-2.8) | 2.4 (2.2-2.7) | 0.90 |
| CSF glucose <2.1 mmol/L, n (%) | 0/7 (0) | 10 (10) | 0.91 |
| CSF protein (g/L), median (IQR) | 0.76 (0.53-0.90) | 0.79 (0.59-1.02) | 0.86 |
| Protein >1 g/L, n (%) | 1 (14) | 25/98 (25) | 0.68 |

Legend: CSF – cerebrospinal fluid; HSV – herpes simplex virus; IQR – interquartile range; WBC – white blood cell count

A Three of the EV cases had CSF sent only for microbiological analysis; so only 100 cases had CSF analysis that included a cell count, protein or glucose level; 2/7 (29%) of the EV cases that were classified as meningoencephalitis had CSF WBC <30 x10^6/L.
For comparison of proportions, Fishers exact test (2-sided) was used; for comparison of medians, Mann-Whitney test was used.

EV cases were more likely than HSV cases to have to have one or more of the parameters (cell count $>1000 \times 10^6/L$, Glucose $<2.0$ mmol/L and CSF Protein $>1.0$ g/L $\geq 1$) that suggested bacterial meningitis (65 (64%) versus 1 (13%); $p=0.006$).

EV cases with CSF WBC $>2000 \times 10^6/L$ had median CSF WBC of 2630 (range 2020-6400) $\times 10^6/L$

CSF pleocytosis was defined as CSF white cell count $>15 \times 10^6/L$ for infants 0-28 days of age and $>9 \times 10^6/L$ for infants beyond neonatal period. Of 74 neonates with spinal taps 22 (30%) neonates had CSF WBC $<15 \times 10^6/L$ and 4/33 (12%) infants $>28$d old had CSF wbc $<9 \times 10^6/L$

Table 3: Demographic, clinical and laboratory factors associated with unfavorable outcome following viral CNS infection by univariate analysis
|                                | Unfavorable Outcome N=12 | Favorable outcome N=98 | P-Value\(^A\) |
|--------------------------------|--------------------------|------------------------|---------------|
| n (%)                          | 9/12 (75)                | 45/98 (45)             | 0.07          |
| \(\text{IQR}\) of seizure onset (d), set, n (%) | 9 (5.5-18.5)             | 25 (14-33)             | 0.003*        |
| Of seizure or during hospitalisation, n (%) | 7/12 (58)                | 2/98 (2)               | <0.001*       |
| NMR imaging, n                 | 7/11 (64)                | 4/22 (18)              | 0.02          |
| Encephalitis                   | 8/12 (67)                | 6/98 (6)               | <0.001*       |
| Infection care mission, n      | 7/11 (64)                | 9/95 (9)               | <0.001*       |
| Meningitis virus, n            | 4/12 (33)                | 3/98 (3)               | 0.002*        |
|                              | 1/12 (8)                 | 2/98 (2)               | 0.29          |
|                              | 3/12 (25)                | 1/98 (1)               | 0.004         |
|                              | 8/12 (67)                | 95/98 (97)             | 0.002         |
| Te blood test                 | 104 (27.5-762)           | 147 (11-358)           | 0.75          |
| Parameter                      | EV Median (IQR) | HSV Median (IQR) | Significance |
|-------------------------------|-----------------|------------------|--------------|
| Protein (g/L)                 | 1.0 (0.64-1.27) | 0.74 (0.56-0.95) | 0.12         |
| Glucose (mmol/L)              | 2.1 (2.0-2.45)  | 2.4 (2.2-2.75)   | 0.03         |
| N. System (%)                 | 5/12 (42)       | 8/99 (8)         | 0.005        |

Legend: CSF – cerebrospinal fluid; HSV – herpes simplex virus

For comparison of proportions, Fishers exact test (2-sided) was used; for comparison of medians, Mann-Whitney test was used.

The presence of one or more of parameters suggestive of bacterial meningitis (cell count >1000 x10^6/L, Glucose <2.0 mmol/L and CSF Protein >1.0 g/L) in infants with EV or HSV infection were not associated with unfavorable outcome.

*These variables remained significant at a p value < 0.004 after Bonferroni correction applied for multiple comparisons.
Table 4: Demographic Clinical and Laboratory Factors Associated with Unfavorable Outcome Following Enteroviral CNS Infection

| Demographic Clinical and Laboratory Factors | Unfavorable Outcome |
|--------------------------------------------|---------------------|

...
|                | Unfavorable Outcome N=8 | Good outcome N=95 | P-Value<sup>A</sup> |
|----------------|-------------------------|-------------------|---------------------|
| Gender, n      | 6/8 (75)                | 43/95 (45)        | 0.15                |
| Onset (d), n (IQR) | 9 (5.5-19.5)          | 25 (14-34)        | 0.02                |
| y of seizure or during treatment, n (%) | 3/8 (38)              | 2/95 (2)          | 0.003*              |
| mal imaging, n (%) | 3/7 (43)              | 4/19 (21)         | 0.34                |
| goencephalitis, % | 4 (50)                 | 3 (95)            | <0.001*             |
| ive care unit sion, n (%) | 4/7 (57)              | 8/91 (9)          | 0.004*              |
| hite blood unt x10<sup>6</sup> /L<sup>B</sup>, n (IQR) | 180 (41-1271)          | 153 (15.5-393.5)  | 0.36                |
| otein (g/L), n (IQR) | 1.10 (0.73-1.27)       | 0.76 (0.59-0.99)  | 0.13                |
| Parameter                        | Group A (N=8) | Group B (N=90) | P-value |
|---------------------------------|---------------|----------------|---------|
| Protein over 1 g/L              | 5/8 (63%)     | 20/90 (22%)    | 0.02    |
| Glucose (mmol/L)                | 2.1 (1.95-2.29) | 2.45 (2.20-2.75) | 0.01    |
| CSF protein over 1 g/L          |               |                |         |
| CSF glucose < 2.0 mmol/L        |               |                |         |
| CNS disease, 1st CSF meter      | 2/8 (25%)     | 6/95 (6%)      | 0.12    |

Legend: CNS – central nervous system; CSF – cerebrospinal fluid

A For comparison of proportions, Fishers exact test (2-sided) was used; for comparison of medians, Mann-Whitney test was used.

B The presence of one or more of parameters suggestive of bacterial meningitis (cell count >1000 x10⁶/L, Glucose <2.0 mmol/L and CSF Protein >1.0 g/L) in infants with EV
was not associated with unfavorable outcome (8 (100%) infants with EV had unfavorable outcome versus vs 57 (62%) with favorable outcome; p=0.05)*These variables remained significant at a p value of < 0.005 after Bonferroni correction applied for multiple comparisons

Appendix

Appendix 1 - ICD10CA codes used to identify potential cases

A170 Tuberculous meningitis
A203 Plague meningitis
A321 Listerial meningitis and meningoencephalitis
A390 Meningococcal meningitis
A870 Enteroviral meningitis
A871 Adenoviral meningitis
A872 Lymphocytic choriomeningitis
A878 Other viral meningitis
A879 Viral meningitis, unspecified
B003 Herpesviral meningitis
B010 Varicella meningitis
B021 Zoster meningitis
B051 Measles complicated by meningitis
B261 Mumps meningitis
B375 Candidal meningitis
B384 Coccidioidomycosis meningitis
G000 Haemophilus meningitis
G001 Pneumococcal meningitis
G002  Streptococcal meningitis
G003  Staphylococcal meningitis
G008  Other bacterial meningitis
G009  Bacterial meningitis, unspecified G00 Bacterial meningitis, not elsewhere classified
G01   Meningitis in bacterial diseases classified elsewhere
G020* Meningitis in viral diseases classified elsewhere
G021* Meningitis in mycoses
G028* Meningitis in other specified infectious and parasitic diseases classified elsewhere
G030  Nonpyogenic meningitis
G031  Chronic meningitis
G032  Benign recurrent meningitis [Mollaret]
G038  Meningitis due to other specified causes
G039  Meningitis, unspecified
A811  Subacute sclerosing panencephalitis
A830  Japanese encephalitis
A831  Western equine encephalitis
A832  Eastern equine encephalitis
A833  St Louis encephalitis
A834  Australian encephalitis
A835  California encephalitis
A838  Other mosquito-borne viral encephalitis
A839  Mosquito-borne viral encephalitis, unspecified
A840  Far Eastern tick-borne encephalitis [Russian spring-summer encephalitis]
A841  Central European tick-borne encephalitis
A848  Other tick-borne viral encephalitis
A849  Tick-borne viral encephalitis, unspecified
A850  Enteroviral encephalitis
A851  Adenoviral encephalitis
A852  Arthropod-borne viral encephalitis, unspecified
A858  Other specified viral encephalitis
A86  Unspecified viral encephalitis
A922  Venezuelan equine fever
A923  West Nile virus infection
B004  Herpesviral encephalitis
B011  Varicella encephalitis
B020  Zoster encephalitis
B050  Measles complicated by encephalitis
B262  Mumps encephalitis
B582  Toxoplasma meningoencephalitis
G040  Acute disseminated encephalitis
G042  Bacterial meningoencephalitis and meningomyelitis, not elsewhere classified
G048  Other encephalitis, myelitis and encephalomyelitis
G049  Encephalitis, myelitis and encephalomyelitis, unspecified
G050*  Encephalitis, myelitis and encephalomyelitis in bacterial diseases classified elsewhere
G05.1*  Encephalitis, myelitis and encephalomyelitis in viral diseases classified elsewhere
G052*  Encephalitis, myelitis and encephalomyelitis in other infectious and parasitic diseases classified elsewhere
G058*  Encephalitis, myelitis and encephalomyelitis in other diseases classified elsewhere
Figure 1

Seasonality of HSV and enteroviral CNS infections in infants < 90 days of age