There are few conditions more devastating than severe encephalitis, in which a previously healthy child presents, within days, from a nonspecific and apparently benign febrile illness to coma, seizure and irreversible brain injury. The current article focuses on key aspects of the etiology and management of acute and subacute childhood encephalitis beyond the neonatal period in the Canadian context. Several recent publications provide comprehensive reviews on the diagnostic approach to suspected encephalitis in both children and adults (1,2).

Strictly speaking, a diagnosis of encephalitis requires proof of brain inflammation. However, in practice, the diagnosis is usually a clinical one, based on the presence of symptoms and signs including fever, headache, seizure or altered consciousness, and indirect evidence of brain inflammation such as cerebrospinal fluid (CSF) pleocytosis, elevated CSF protein level or suggestive neuroimaging abnormalities. Reduced consciousness – the hallmark of encephalopathy – is present in most cases of encephalitis, but can also be observed with other infections, such as bacterial meningitis, sepsis or malaria, and with numerous noninfectious disorders including metabolic, immunological, toxic and drug-related conditions. A case definition for encephalitis has been proposed both as an aid to diagnosis and to enable future research studies to be more comparable with one another (Table 1) (1).

The incidence of encephalitis in developed countries ranges from 0.7 to 13.8 per 100,000 population (3). In a United Kingdom study, incidence rates varied from 1.5 per 100,000 overall to 2.8 per 100,000 for children and 5.7 per 100,000 for infants (4). The incidence of encephalitis due to specific pathogens is dynamic, influenced by a variety of factors including vaccination rates for diseases such as measles, mumps and varicella; geographical spread of vector-borne pathogens due to climate change or inadvertent air/ship transport, as occurred with the introduction of West Nile virus into North America; and the emergence of previously unknown zoonotic pathogens (eg, Nipah virus).

Pathogenesis varies according to etiology. Classic viral encephalitis due to herpes simplex virus (HSV), enteroviruses and arboviruses is caused by direct infection of the central nervous system (CNS) and necrosis of neurons or other cells. Measles, mumps and Mycoplasma pneumoniae can cause both acute encephalitis due to direct invasion of the CNS and postinfectious immune-mediated encephalitis. Influenza and other respiratory virus-associated encephalitides may involve excessive production of proinflammatory cytokines, mitochondrial dysregulation or endothelial dysfunction (3). Acute disseminated encephalomyelitis (ADEM) is likely a consequence of autoimmunity, although the precise mechanisms are not fully understood. Development of antibodies directed against the N-methyl-D-aspartate receptor (NMDAR) is responsible for anti-NMDAR encephalitis. Post-HSV encephalitis ‘neurological relapse’ (typically characterized by choreoathetoid movements), in which HSV CSF polymerase chain reaction (PCR) testing is negative, has been attributed to the production of anti-NMDAR antibodies or, less frequently, anti-dopamine-2 receptor antibodies (5,6).

A confirmed or probable etiology can be identified in 30% to 60% of acute childhood encephalitis cases. The most common infectious causes in otherwise healthy Canadian children are HSV, varicella zoster virus (VZV), Epstein Barr virus (EBV), M pneumoniae and influenza, adenovirus and other respiratory viruses (Table 2). Arboviruses endemic to North America and enterovirus/parechoviruses are important considerations during the summer and fall months (7-10), whereas influenza and other respiratory viruses are more typically observed during the winter months. M pneumoniae encephalitis can be observed year-round, with increased incidence coinciding with epidemic respiratory disease every three to seven years (11). Less commonly encountered pathogens include measles, mumps, rubies, Bartonella henselae, Baylisascaris procyoni and travel-related arboviruses, Rickettsia and free-living amoebae. ADEM and anti-NMDAR encephalitis are the predominant noninfectious causes of encephalitis in children, together accounting for approximately 15% to 25% of all encephalitis cases, respectively (3,12). A review of cases involving patients <30 years of age referred to the California Encephalitis Project suggested that the frequency of anti-NMDAR encephalitis was higher than any single viral etiology (12). The main causes of subacute encephalitis in children are shown in Table 3.

In the immunocompromised host, herpesvirus groups (including HSV, VZV, EBV, cytomegalovirus and human herpes virus-6 [HHV-6]), enteroviruses and polyoma JC virus (progressive multifocal leukoencephalopathy) are the predominant causes of encephalitis (13,14). Despite a confirmed or probable etiology, many cases of encephalitis remain unexplained (12). A review of cases involving patients <30 years of age referred to the California Encephalitis Project suggested that the frequency of anti-NMDAR encephalitis was higher than any single viral etiology (12). The main causes of subacute encephalitis in children are shown in Table 3.

### TABLE 1

| Major criterion (required) | Minor criteria (two for possible encephalitis, ≥3 for probable or confirmed encephalitis) |
|---------------------------|------------------------------------------------------------------------------------------|
| Altered mental status, defined as decreased or altered level of consciousness, lethargy, or personality change lasting ≥24 h with no alternative cause identified | Documented fever ≥38.0°C (100.4°F) within 72 h before or after presentation |
| Generalized or partial seizures not attributable to a pre-existing seizure disorder | Abnormality of brain parenchyma on neuroimaging suggestive of encephalitis |
| New onset of focal neurological findings | Abnormality on electroencephalography that is consistent with encephalitis |
| Cerebrospinal fluid pleocytosis (leukocyte count ≥5×10⁹/L) | Abnormality on electroencephalography that is consistent with encephalitis and not attributable to another cause |
| Abnormality of brain parenchyma on neuroimaging suggestive of encephalitis that is either new from previous studies or appears to be acute in onset | *Adapted from reference 1; †Confirmed encephalitis requires: pathological confirmation of brain inflammation consistent with encephalitis; pathological, microbiological or serological evidence of acute infection with a microorganism that is strongly associated with encephalitis from an appropriate clinical specimen; or laboratory evidence of an autoimmune condition that is strongly associated with encephalitis* |

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Can J Infect Dis Med Microbiol Vol 26 No 2 March/April 2015 69
leukoencephalopathy) are important considerations. Of note, the absence of rash does not preclude VZV as a cause of encephalitis in severely immunocompromised individuals; elevated liver enzyme levels can serve as a clinical indicator in this circumstance. Measles inclusion body encephalitis is typically observed within one year of measles virus infection or measles vaccination in children with primary or secondary immune deficiency (13,14). Encephalitis due to West Nile virus and rabies has been observed in adults following organ transplantation from an infected donor (15,16).

Travel-related viral encephalitides, not endemic to Canada, should be considered if symptom onset began within three to four weeks of leaving the region (1-3,17). Japanese encephalitis virus, endemic to Japan, Korea, Taiwan, China, Southeast Asia, India, Nepal and northern Australia, is a predominant travel-related pathogen (18). Rural travel during the summer months to central or northern Europe or Russia, including Siberia, should lead to consideration of tick-borne encephalitis virus. Nipah virus infection is acquired following close contact with swine, bats or bat roosting sites, or infected

### TABLE 2

**Selected clinical, microbiological and treatment aspects of infectious causes of childhood encephalitis acquired in Canada**

| Cause                  | Comments |
|------------------------|----------|
| HSV (5,6,22)           | PCR of CSF can be negative during the first 72 h of illness; repeat lumbar puncture if clinical picture is consistent with HSV encephalitis. Treatment: intravenous acyclovir 30–60 mg/kg/day for children three months to 12 years of age and 30 mg/kg/day for children ≥12 years of age in three divided doses for three weeks. Testing for anti-NMDAR antibodies in CSF and serum is warranted for new-onset postencephalitis neurological symptoms. Corticosteroids may be of benefit for post-HSV encephalitis symptomatic relapse if CSF tests negative for HSV by PCR and there are no new necrotic lesions on MRI. |
| EBV (24)              | Serology is the mainstay of diagnosis (monospot; IgM and IgG VCA; IgG EA, VCA and EBNA). Detection of EBV by PCR in CSF may represent true disease or virus DNA within passenger lymphocytes. Treatment: intravenous ganciclovir 10 mg/kg/day in two divided doses for 2–3 weeks should be considered for severe cases; complete resolution without treatment is common |
| VZV (25)              | Concurrent or recent chickenpox in majority. CSF PCR is negative in 75% of children. Treatment: intravenous acyclovir – for children ≥2 years of age, 80 mg/kg per day in four divided doses (max 3200 mg/day) if ≥40 kg, 3200 mg in four divided doses if >40 kg for 1–3 weeks |
| HHV-6                 | Restricted to immunocompromised subjects. Detection of HHV-6 in CSF by PCR may represent true infection or latent virus within lymphocytes or DNA integration related to vertical germline transmission. Treatment: intravenous ganciclovir 10 mg/kg/day in two divided doses for 2–6 weeks should be considered (foscarnet is an alternative agent if ganciclovir is contraindicated). |
| HHV-7 (26)            | Rare; mainly teenagers and young adults. Majority of positive CSF PCR test results associated with latent infection (eg, likely false positive with respect to encephalitis causation); microbiological diagnosis requires confirmation of seroconversion (not available commercially). No data regarding treatment are available |
| Enterovirus           | Parechovirus disease generally restricted to those <3 months of age. Most children with parechovirus encephalitis do not demonstrate pleocytosis. CSF PCR for enteroviruses relatively insensitive; testing of respiratory samples and stool recommended. Treatment: IVIG may be considered for severe cases, but evidence is inconclusive; pleconaril has activity against some serotypes of enterovirus but is currently not available |
| West Nile virus (7)   | Serology is the mainstay of diagnosis (CSF and serum); may be negative early in course or positive from remote infection. CSF and blood PCR have poor sensitivity after neurosurgical symptom onset |
| Other arboviruses (8,10,18,27) | |
| Rabies (28)           | Diagnostic tests include serology on serum and CSF, RT-PCR or virus isolation from saliva, detection of rabies antigen in cutaneous nerves. Treatment: intravenous acyclovir 30–60 mg/kg/day for children three months to 12 years of age and 30 mg/kg/day for children ≥12 years of age in three divided doses for three weeks. Testing for anti-NMDAR antibodies in CSF and serum is warranted for new-onset postencephalitis neurological symptoms. Corticosteroids may be of benefit for post-Rabies encephalitis symptomatic relapse if CSF tests negative for Rabies by PCR and there are no new necrotic lesions on MRI. |
| Influenza A and B (5) | Abrupt onset before or concurrent with respiratory symptom onset in some cases. Acute necrotizing encephalopathy is a severe form of disease including bilateral symmetric necrosis of the thalamus, putamina and cerebellar white matter on MRI. PCR on CSF is almost uniformly negative. Treatment: oseltamivir according to weight for 5–10 days |
| Measles               | Occurs in children who are incompletely immunized. Cough, corony, conjunctivitis, erythematous maculopapular rash, Koplik spots. Serology and direct detection by PCR in CSF, nasopharyngeal and urine samples |
| Mumps                 | Occurs in children who are incompletely immunized. Parotitis, testicular pain. Serology and direct detection by PCR in CSF, saliva |
| Mycoplasma pneumoniae (11,23) | Absence of a respiratory prodrome does not preclude as a cause. Both direct and immune-mediated pathogenesis hypothesized. PCR of CSF and respiratory samples plus serology in peripheral blood recommended for diagnosis; due to poor specificity and predictive value, serology should not be relied upon in isolation for diagnosis. Treatment: A 10–14 day course of antymycoplasmal antibiotic therapy may be considered; role of corticosteroids uncertain |
| Bartonella henselae   | Consider in those with exposure to cats. Serology is the mainstay of microbiological diagnosis. PCR available, but utility in diagnosis of encephalopathy/encephalitis unknown. Treatment: doxycycline or cotrimoxazole plus rifampin for 4–6 weeks may be considered |
| Baylisascaris procyonis (19) | Consider in toddlers or cognitively impaired children with exposure to soil potentially contaminated with raccoon feces. Peripheral blood and stool examination for Baylisascaris DNA integration related to vertical germline transmission. Treatment: intravenous ganciclovir 10 mg/kg/day in two divided doses for 2–3 weeks. |
| CMV                   | Consider in those who are immunocompromised. Serological diagnosis (requires acute and convalescent serum); consider in summer and fall |
| Parainfluenza virus   | Consider in those with exposure to cats. Serology is the mainstay of microbiological diagnosis. PCR available, but utility in diagnosis of encephalopathy/encephalitis unknown. Treatment: doxycycline or cotrimoxazole plus rifampin for 4–6 weeks may be considered |

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**Notes:**

1. The 2012 Red Book recommends a dose of 30–45 mg/kg/day in three divided doses for children three months to 12 years of age. However, some experts prefer to use the upper limit of the approved dose range (60 mg/kg/day in three divided doses) for children <12 years of age; close monitoring of renal function is required.
2. In adults, herpes zoster-associated encephalitis predominates (mainly in immunocompromised hosts). Some experts prefer to use 1500 mg/m²/day in three divided doses, the dose usually used to treat varicella in immunocompromised hosts; duration of therapy has not been well established, and close monitoring of renal function is required.
3. Eastern equine and Powassan encephalitis viruses are a consideration primarily in the eastern provinces; Western equine encephalitis predominately in the western and prairie provinces and into Ontario; St Louis and the California serogroup viruses (Snowshoe hare and Jamestown Canyon) encephalitis viruses are more ubiquitous. Other respiratory viruses, including adenovirus, parainfluenza viruses and respiratory syncytial virus, are occasionally associated with encephalitis or infection-associated encephalopathy. CSF Cerebrospinal fluid; EA Early antigen; EBNA Epstein Barr nuclear antigen; EBV Epstein Barr virus; HHV-6 Human herpes virus-6; HHV-7 Human herpes virus-7; HSV Herpes simplex virus; Ig Immunoglobulin; IVIG Intravenous immunoglobulin; MRI Magnetic resonance imaging; NMDAR N-methyl-D-aspartate receptor; PCR Polymerase chain reaction; RT-PCR Reverse transcriptase PCR; VCA Viral capsid antigen; VZV Varicella zoster virus
humans in south Asia. Recent travel to Australia or New Guinea suggests the possibility of encephalitis due to Murray Valley virus or Hendra virus, the latter being associated with contact with horses. Rift Valley fever virus can be acquired from mosquito bites, or direct or indirect contact with animals or animal products (including carcasses) in sub-Saharan Africa or the Arabian Peninsula. Dengue and Chikungunya viruses are rare causes of encephalitis that can be acquired in many tropical and subtropical regions, including the Caribbean. Other considerations include herpes B virus in cases with a history of bite or scratch by a macaque monkey (old world), Toscana virus transmitted by sandflies in Mediterranean countries, Taunya virus in Russia and other parts of Europe, and Venezuelan equine encephalitis virus endemic to central and South America.

The microbiological investigation of the previously well child with acute encephalitis need not be exhaustive and should be tailored to the history and findings on physical examination (1-3). PCR testing of CSF for HSV, EBV, entero-/parechoviruses and M pneumoniae and of respiratory samples for M pneumoniae and entero-/parechoviruses is recommended for all cases. Stool should be tested for enteroviruses and adenovirus. Serology on peripheral blood for acute EBV and M pneumoniae infection should also be performed, and an acute serum sample stored for subsequent testing for less commonly encountered pathogens (1). Serological testing of serum and CSF for West Nile virus and other endemic arboviruses should be performed when disease onset is during the summer and fall months. In the presence of respiratory symptoms, respiratory samples should be tested for influenza A and B, adenoviruses, parainfluenza viruses, respiratory syncytial virus and rhinoviruses. A vesicular rash should prompt testing of lesion samples for HSV, VZV and selectively for enteroviruses by PCR (or culture). Serology and PCR testing of CSF as well as respiratory or salivary samples for measles and mumps is warranted in children with suggestive clinical findings. Beyslisascaris procyonis serology is appropriate for toddlers or autistic/developmentally delayed children with peripheral or CSF eosinophilia, particularly if there is a history of pica or playing in soil frequented by raccoons (19). In case of travel, a detailed history that includes precise travel dates, specific regions visited (rural versus urban), any documented exposures to arbovirus vectors, animals (including bites or scratches), or swimming in lakes and brackish water should direct testing for travel-related infections (1-3).

Treatment of acute childhood encephalitis begins with meticulous supportive care that includes management of fluid and electrolytes, respiratory status, seizures and increased intracranial pressure. Intravenous acyclovir should be given to all children pending microbiological test results (Table 2). A full three-week course of therapy should be administered if HSV disease is confirmed or if the clinical picture is most compatible with HSV encephalitis regardless of PCR results, particularly if the lumbar puncture was performed within 72 h of symptom onset. Treatment directed at other pathogens should be considered, although efficacy is less well established (Table 2). First-line therapy for ADEM and anti-NMDAR encephalitis consists of high-dose corticosteroids; other treatment modalities include intravenous immunoglobulin, plasma exchange and, for refractory cases of anti-NMDAR encephalitis, rituximab (20). Excision of ovarian teratoma or other tumours associated with anti-NMDAR encephalitis speeds up clinical response and reduces relapses (21).

The risk for death and permanent neurological sequelae including cognitive impairment, psychiatric or behavioural disorders, epilepsy, motor dysfunction, cranial nerve deficits or movement disorders vary according to etiology. Rubies encephalitis, subacute sclerosing panencephalitis, measles inclusion-body encephalitis, rubella panencephalitis and non-HIV-associated progressive multifocal leuкоencephalopathy are almost uniformly fatal. Although mortality is now uncommon, neurological sequelae occur in 50% to 60% or more of children with HSV encephalitis despite acyclovir treatment (22). The outcome of M pneumoniae-associated encephalitis is similarly poor, with 50% to 60% of children experiencing sequelae (23). Death or neurological sequelae following arboviral encephalitis varies from 50% to 70% for Eastern equine and Powassan encephalitis to 10% to 30% for Japanese, St Louis equine and La Crosse encephalitis (8-10,18). Most children with ADEM respond to immune-modulation, but approximately 20% experience relapses, some of whom are eventually diagnosed with multiple sclerosis. With appropriate treatment, 85% of children with anti-NMDAR encephalitis demonstrate either complete recovery (60%) or minimal residual deficits (25%) (6).

Table 3: Selected causes, clinical features and diagnostic tests for subacute childhood encephalitis

| Condition                  | Selected clinical features and diagnostic methods                                                                 |
|----------------------------|---------------------------------------------------------------------------------------------------------------|
| Anti-NMDAR encephalitis†   | Female preponderance (70%)                                                                                   |
|                            | Psychiatric or behavioural symptoms, seizures, movement disorder, catatonia, hallucinations                   |
|                            | Anti-NMDAR antibodies present in CSF and selectively in serum                                               |
|                            | Teratoma, other tumours should be sought, but are uncommon in children <12 years of age (regardless of sex) and in boys of any age |
| Subacute sclerosing panencephalitis | Onset typically between 5–15 years of age                                                                    |
|                            | Cognitive decline, personality change, strange behaviour, myoclonic jerks, seizures, dementia                 |
|                            | Electroencephalography showing periodic complexes                                                           |
|                            | Elevated CSF, serum anti-measles titers (CSF/serum titre ratio typically <200)                              |
| Measles inclusion body encephalitis | Immunocompromised hosts only; most associated with hematological malignancies                                |
|                            | Onset within one year of measles infection (or vaccination)                                                 |
|                            | Altered mentality, focal afebrile seizures                                                                  |
|                            | Electroencephalography often show epilepsia partialis                                                     |
|                            | Brain biopsy required for diagnosis (RT-PCR, electron microscopy)                                            |
| JC virus (PML)             | Predisposed to by significant cell-mediated immune deficiency state                                          |
|                            | Cognitive dysfunction, focal neurological deficits                                                          |
|                            | Non-enhancing confluent subcortical white matter                                                          |
|                            | Hyperintensities on T2/fluid attenuation inversion                                                          |
|                            | Recovery magnetic resonance imaging images                                                                 |
|                            | Microbiological diagnosis with CSF PCR                                                                    |
| Rubella panencephalitis    | Onset typically between 10–20 years of age                                                                  |
|                            | Other clinical features of congenital rubella present                                                      |
|                            | Anti-rubella IgG and IgA in CSF may be elevated                                                            |

*Other infections that can rarely present with a subacute/chronic encephalitic picture in childhood include HIV, Mycobacterium tuberculosis, congenital syphilis and variant Creutzfeldt-Jakob disease; †Encephalitis due to other autoantibodies to neuronal surface γ-aminobutyric acid-A receptor, γ-aminobutyric acid-B receptor, γ-aminobutyric acid-D receptor, γ-aminobutyric acid-E receptor, γ-aminobutyric acid-F receptor, γ-aminobutyric acid-G receptor, γ-aminobutyric acid-H receptor, γ-aminobutyric acid-I receptor, γ-aminobutyric acid-J receptor, γ-aminobutyric acid-K receptor, γ-aminobutyric acid-L receptor, γ-aminobutyric acid-M receptor, γ-aminobutyric acid-N receptor, γ-aminobutyric acid-O receptor, γ-aminobutyric acid-P receptor, γ-aminobutyric acid-Q receptor, γ-aminobutyric acid-R receptor, γ-aminobutyric acid-S receptor, γ-aminobutyric acid-T receptor, γ-aminobutyric acid-U receptor, γ-aminobutyric acid-V receptor, γ-aminobutyric acid-W receptor, γ-aminobutyric acid-X receptor, γ-aminobutyric acid-Y receptor, γ-aminobutyric acid-Z receptor.
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