WHAT IS ENCEPHALOPATHY?

Encephalopathy is a clinical state characterised by an alteration of consciousness, behaviour and/or cognition. It can present with lethargy and drowsiness, or conversely with a heightened state of agitation and confusion. It arises from diffuse brain dysfunction that can result from myriad underlying conditions, ranging from primary neurologic to systemic disorders. While encephalopathy more commonly stems from a global brain insult, it can present with a range of focal neurological manifestations such as seizures, visual disturbances, speech abnormalities, motor weakness, and sensory and autonomic deficits. Encephalitis refers specifically to brain inflammation, usually associated with fever, with evidence of cerebrospinal inflammation (pleocytosis and raised protein levels) and/or neuroimaging findings of encephalitis.

HOW RELEVANT IS THIS TO MY PRACTICE?

Altered behaviour is common in children and has many causes. In the general clinic setting, it is important to pick up common clinical mimics of encephalopathy that can be managed without immediate referral to the hospital. These include post-ictal drowsiness from simple febrile seizures, and metabolic derangements such as hypoglycaemia. Conversely, it is also important to identify red flags in the history and physical examination that necessitate either activation of emergency services or an urgent referral to the neurologist for further workup.

In particular, encephalitis accounts for a significant proportion of encephalopathy cases that persist beyond 24 hours, and can result in significant morbidity and mortality. The worldwide incidence of encephalitis ranges from 3.3 to 10.7 per 100,000 children per year.1-8 Arriving at an accurate diagnosis for the underlying cause is critical because many causes are reversible, and early specific treatment confers a better prognosis.

WHAT CAN I DO IN MY PRACTICE?

In this primary care approach to encephalopathy in paediatric patients, we propose looking at the main differential diagnoses by age group and time course of onset. A comprehensive history is vital, especially from the caregiver, as patient accounts may be limited due to their age and clinical status. Key features in the physical examination should be specifically looked out for. These provide a framework to guide investigations and management (Table I).

Pertinent features in history taking

Careful attention needs to be paid to the onset, duration and progression of symptoms (Table II). A rapidly dropping Glasgow Coma Scale (GCS) score (usually taken as a drop of 2 or more) with headache and vomiting suggests raised intracranial pressure and necessitates immediate evacuation to the nearest hospital. In the presence of major trauma (e.g. fall from greater than 3 body height/3 m, road traffic accident), intracranial bleeds must always be suspected. There should also be a high index of suspicion for non-accidental injury in infants under one year of age, especially with concerning features in the history or physical examination. These include unexplained bruises (of varying ages, in areas that are usually clothed, or suspicious circular, square, tramline or herringbone patterns); a history that is vague, inconsistent or implausible; or delay in seeking medical attention. In the absence of trauma, intracranial bleeds can also be the first manifestation of an underlying arteriovenous malformation. Evolution of symptoms over hours to days is most commonly a result of either infectious or infection-associated immune encephalitis. Some atypical infections such as tuberculosis (TB) and human immunodeficiency virus (HIV) may present more insidiously over weeks. Other considerations for more chronic presentation include underlying inflammatory, neurometabolic and neoplastic processes.

In children, functionality and deviation from baseline behaviour is critical in ascertaining the presence, and degree, of encephalopathy. It is thus important to elicit this information from the main caregiver specifically, if possible; this may not necessarily be the parent who brought the child but a domestic helper or grandparent. The caregiver’s reports of irritability, lethargy or drowsiness should always be taken seriously, especially if these symptoms are disproportionate to any fever spikes. In younger patients, aversion to feeds, disinterest in toys and play, and decreased activity may be the only manifestations of encephalopathy. Infants with poor intake (usually considered less than 50% of normal) and dehydration (increased thirst,
Table I. Recommended approach to history and physical examination.

<table>
<thead>
<tr>
<th>Item</th>
<th>Characteristic</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Initial approach</strong></td>
<td></td>
</tr>
</tbody>
</table>
| Is the patient stable?            | • Reduced Glasgow Coma Scale score  
• Abnormal vitals  
• Ongoing seizure                                                                                  |
| **History taking**                |                                                                                                                                               |
| Characterisation of episode       | • Drowsiness, lethargy  
• Confusion, agitation  
• Personality change  
• Hallucinations  
• Speech disturbances  
• Abnormal movements or gait                                                                       |
| Time course                       | • Hyperacute (over minutes)  
• Acute (over days)  
• Subacute (over weeks)  
• Chronic (over months)                                                                                 |
| Baseline function                 | • Reduced intake/output  
• Decreased activity  
• Disinterest in play/poor engagement                                                                       |
| History of seizures               | • Generalised vs. focal  
• First episode vs. recurrent  
• Febrile vs. afebrile  
• Duration                                                                                             |
| Associated symptoms               | • Fever, headache, vomiting  
• Vision abnormalities  
• Diarrhoea/abdominal pain  
• Cough/nasal congestion/sore throat  
• Systemic features: rash, joint pain, oral ulcers  
• Malaise, loss of appetite/weight                                                                        |
| Significant exposures             | • Trauma  
• Toxins and drugs  
• Travel  
• Animal and sick contacts                                                                             |
| Past medical history              | • Perinatal history  
• Epilepsy  
• Developmental milestones  
• Neurogenetic diagnoses  
• Psychiatric disorders  
• Other systemic conditions                                                                               |
| Developmental/academic history    | • Appropriate vs. delayed vs. regression  
• Issues raised by teachers in school                                                                 |
| Family history                    | • Epilepsy  
• Neurogenetic, metabolic, developmental disorders  
• Autoimmune diseases                                                                                   |
| **Physical examination**          |                                                                                                                                               |
| General inspection                | • Lethargy, confusion, agitation  
• Mood and affect  
• Dysmorphism  
• Poor interaction with environment and caregiver  
• Speech abnormalities: quantity (excessive, aphasia), quality (pressured, echolalia, volume), articulation (slurred)  
• Abnormal posture/gait  
• Presence of involuntary movements                                                                        |
| Head                              | • Abnormal occipitofrontal circumference: crossing centiles, macrocephaly, microcephaly  
• Bulging or tense anterior fontanelles                                                                    |

(Contd...)
Table I. (Contd...)  

<table>
<thead>
<tr>
<th>Item</th>
<th>Characteristic</th>
</tr>
</thead>
</table>
| Eyes                      | • Pupillary reaction, anisocoria  
|                           | • Eye movements (sustained deviation, nystagmus, opsoconius)  
|                           | • Ophthalmoplegia  
|                           | • Visual acuity and fields  
|                           | • Fundoscopy: papilloedema, pale discs  
| Neurological examination  | • Focal neurological deficits  
|                           | • Persistence of primitive reflexes  
|                           | • Neurocutaneous stigmata  
| Focused examination (guided by history) | • Injuries/bruises  
|                           | • Rashes, mouth ulcers, joint swelling  
|                           | • Lymphadenopathy  
|                           | • Hepatosplenomegaly  
| Investigations            | Hypoglycaemia, hyperglycaemia  

Table II. Primary neurological conditions causing encephalopathy.  

<table>
<thead>
<tr>
<th>Patient group</th>
<th>Hyperacute (minutes to hours)</th>
<th>Acute (hours to days)</th>
<th>Subacute (days to weeks)</th>
<th>Chronic (weeks to months)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Neonate/infants</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
|                                   | • Seizures                    | • Infection (bacterial, viral)  
|                                   | • Non-accidental injury (SDH, ICH) | • Vascular (stroke, cerebral venous thrombosis)  
|                                   | • Hypoxic-ischaemic injury    | • Hyperammonaemia (defects in urea or amino acid metabolism)  
|                                   | • Metabolic derangements      |                       | • Hydrocephalus            | • Neurometabolic disorders  
|                                   | (hypoglycaemia, hyperglycaemia, hyponatraemia) | | • Infection (TB, fungal, parasitic)  
|                                   |                              |                       |                          | • Genetic syndromes  
|                                   |                              |                       |                          | • Epileptic encephalopathies  
|                                   |                              |                       |                          | • Neoplasms (primary brain tumours, metastasis)  
| **Preschoolers/older children**  |                              |                       |                          |                           |
|                                   | • Seizures                    | • Infection (bacterial, viral)  
|                                   | • Trauma                      | • Inflammation (post-infectious, autoimmune encephalitis, most commonly anti-NMDAR encephalitis, acute demyelinating conditions)  
|                                   | • Vascular (ICH, ischaemic stroke) |                       | • Infection (TB, fungal, parasitic)  
|                                   | • Hypoglycaemia, DKA          |                       | • Inflammation (post-infectious, autoimmune encephalitis, most commonly anti-NMDAR encephalitis, acute demyelinating conditions)  
|                                   | • Drugs/toxins                |                       | • Neurometabolic disorders  
|                                   |                              |                       | • Neoplasm (primary brain tumours, metastasis)  
|                                   |                              |                       | • Infections (TB, HIV)  
| **Adolescents**                   |                              |                       |                          |                           |
|                                   | • Seizures                    | • Infection (bacterial, viral)  
|                                   | • Trauma                      | • Inflammation (post-infectious, autoimmune encephalitis, most commonly anti-NMDAR encephalitis, acute demyelinating conditions)  
|                                   | • Vascular (ICH, ischaemic stroke) |                       | • Infection (TB, fungal, parasitic)  
|                                   | • Drugs/toxins ingestion      |                       | • Inflammation (post-infectious, autoimmune encephalitis, most commonly anti-NMDAR encephalitis, acute demyelinating conditions)  
|                                   | • Posterior reversible encephalopathy syndrome |                       | • Neurometabolic disorders  
|                                   | • Migraine                    |                       | • Neoplasm (primary brain tumours, metastasis)  
|                                   |                              |                       | • Neurometabolic disorders  
|                                   |                              |                       | • Infections (TB, HIV)  

CNS: central nervous system; DKA: diabetic ketoacidosis; HIV: human immunodeficiency virus; ICH: intracranial haemorrhage; NMDAR: N-methyl-D-aspartate receptor; SDH: subdural haemorrhage; SLE: systemic lupus erythematosus; TB: tuberculosis

reduced urinary output) due to an otherwise simple viral illness are nevertheless predisposed to hypoglycaemia or cerebral venous thrombosis, both of which can also result in altered mentation. Even if the child appears well during the point of clinical contact, there should be a high index of suspicion for delirium, which presents with fluctuating levels of alertness. It may be prudent in these cases to advise admission for monitoring.

Seizures are sometimes the first presentation of encephalopathy but are far more commonly caused by simple febrile seizures or epilepsy. As a general rule, a simple febrile fit in a child older than 18 months, who has no red flags (i.e. normal physical/ neurological examination, duration < 15 minutes, no recurrence within 24 hours, no focal lateralising signs, normal developmental milestones, no prior head trauma/central nervous system
infections) and has returned to baseline function, can be managed in the primary care setting; all other patients should be referred to the emergency department. It should also be noted that neonatal seizures can be subtle and are usually not generalised in view of immature cerebral motor pathways.\(^4\) However, if episodes are recurrent and stereotypical, clinicians need to have a high index of suspicion for prompt referral. Caregivers should be encouraged to take videos of suspected episodes, as reviewing them is helpful in characterising the semiology and directing further investigation.

The presence of fever suggests an infective or infection-associated encephalitis, which is often preceded or accompanied by respiratory or gastrointestinal symptoms. Correspondingly, the commonest pathogens detected with encephalitis are *Streptococcus*, influenza virus, *Mycoplasma pneumoniae*, *Escherichia coli*, enterovirus and rotavirus.\(^5\) Travel and contact history (including animals) is important in screening for more atypical infections. In older teens, systemic features (photosensitive rash, joint pain, ulcers) should be sought, as autoimmune diseases such as systemic lupus erythematosus tend to present in this age group.

In previously healthy children, an acute to subacute onset of neuropsychiatric symptoms (personality change, auditory hallucinations) accompanied by a suggestive clinical phenotype that includes movement abnormalities, speech impairment, seizures, mood lability and sleep dysregulation raises the suspicion of autoimmune encephalitis.\(^6,7\) In the last decade, autoimmune encephalitis has been shown to rival the frequency of viral aetiologies\(^8\) and should be considered early in the presentation of encephalopathy. Many are seronegative, but the most common autoantibody in children targets the N-methyl-D-aspartate receptor (NMDAR). A timely diagnosis will avoid unnecessary diagnostic and treatment costs and permit earlier directed therapy.\(^9\)

Finally, environmental risk factors should be explored. In the younger child, caregivers should be questioned on the possibility of accidental toxin exposure (e.g. child left unattended or medications/toxins within reach). In the adolescent, the possibility of alcohol use, drug use or intentional overdose should not be overlooked.

**Clinical examination**

The first priority of the clinical examination is to ensure the child is haemodynamically and neurologically stable, with a normal, age-appropriate heart rate, respiratory rate, oxygen saturations and blood pressure (Table III). State of consciousness should be assessed, with particular attention on the pattern of breathing, pupillary size and reactivity, and GCS score. This assessment tool is quick to use and focuses on three domains – eye movements, verbal responses and motor responses – with a total score ranging from 3 to 15 (Table IV). A GCS of ≤ 8, or a rapidly dropping GCS, indicates more extensive obtundation and is considered a neurological emergency. There should be immediate activation of services to the nearest emergency department.

It is important to note that while the GCS score is helpful, it should not be the only measure of an altered mental state. Encephalopathy can be present even with a GCS score of 15, as the score does not take into consideration other behavioural or cognitive changes such as mood lability and higher-level executive functioning. The inclusion of a neurocognitive test such as the Mini Mental State Examination for the older paediatric patient can reveal subtle cognitive dysfunction that would be missed at first impression (Box 1). Attention should also be paid to the quality and quantity of verbal responses; expressive aphasia or dysarthria can be secondary to an underlying stroke or tumour; verbal reduction or pressured speech are observed in patients with anti-NMDAR encephalitis.

As the clinical history is being obtained from the caregiver, an observant clinician can simultaneously glean valuable information from watching the patient in the caregiver’s lap or at play. This is especially helpful in younger children who may be fretful and refuse more direct examination. The child’s resting posture, movement and interaction should be observed, noting poor tone, asymmetry of limb movement, abnormal posturing, lack of interaction or poor eye contact with caregivers. In an older patient, gait and movement abnormalities should be specifically looked for, even before performing a formal neurological examination. Choreoathetoid and other dyskinetic movements of the face, mouth, tongue and limbs and/or catatonic features (staring, minimal responsiveness or excessive hyperactivity, rigid posturing) are typical presentations for autoimmune encephalitis if drug ingestion has been ruled out.\(^6,7\)

A detailed examination of the eyes and head can reveal important clues. Small or large pupils that are sluggish to light in the absence of a dropping GCS score may suggest drug ingestion. Persistent gaze deviation may be a sign of ongoing seizure. An isolated sixth nerve palsy and/or papilloedema points to raised intracranial pressure, while ophthalmoplegia

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**Table III. Normal vital parameters by age.**

<table>
<thead>
<tr>
<th>Age group</th>
<th>Heart rate (bpm)</th>
<th>Respiratory rate (min)</th>
<th>SBP* (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neonate</td>
<td>120–180</td>
<td>40–60</td>
<td>60–80*</td>
</tr>
<tr>
<td>Infant (1 mth–1 yr)</td>
<td>110–160</td>
<td>30–40</td>
<td>70–90</td>
</tr>
<tr>
<td>Toddler (1–2 yr)</td>
<td>100–150</td>
<td>25–35</td>
<td>80–95</td>
</tr>
<tr>
<td>Young child (2–7 yr)</td>
<td>95–140</td>
<td>25–30</td>
<td>90–110</td>
</tr>
<tr>
<td>Older child (7–12 yr)</td>
<td>80–120</td>
<td>20–25</td>
<td>100–120</td>
</tr>
</tbody>
</table>

\(*\) Hypotension is defined as SBP < 60 mmHg in a neonate, < 70 mmHg in infants below 12 mth, < 70 mmHg + [2 × (age in yr)] in children 1–10 yr, < 90 mmHg in children above 10 yr. Mean arterial pressure is calculated as 50 + [2 × (age in yr)] mmHg. SBP: systolic blood pressure.
should prompt careful examination of the other cranial nerves for better localisation of the pathology. Reduced visual acuity with red desaturation is highly suggestive of optic neuritis in the setting of a demyelinating disorder, such as neuromyelitis optica or multiple sclerosis.

Occipitofrontal circumference (OFC) should be routinely measured in all children aged two years and under. Serial OFCs that rapidly increase and cross centiles raise the suspicion of hydrocephalus or a space-occupying lesion, while microcephaly (OFC < third centile) or macrocephaly (OFC > 97th centile) could be clues of underlying perinatal insults or neurogenetic disorders. A tense or bulging anterior fontanelle in a non-crying infant is also indicative of raised intracranial pressure.

A developmental assessment together with the full neurological examination should be performed in infants and younger children, especially those with more chronic symptoms, as this may be a manifestation of underlying malignancy or neurometabolic disorder. Developmental regression, defined as a plateauing or loss of milestones already acquired, is a red flag for early specialist referral.

**Investigations**

In the primary care setting, a quick hypocount would exclude any reversible glucose derangement that can account for the altered mental status. Hypoglycaemia is generally defined as serum glucose less than 3.5 mmol/L. It is a particularly important diagnosis to exclude in neonates and infants, as delayed treatment can result in neuronal injury in a vulnerable young brain. Hypoglycaemia can also be the first manifestation of an underlying metabolic or endocrine disorder. Hyperglycaemia in the setting of acidosis is suspicious for diabetic ketoacidosis (serum glucose > 11 mmol/L, pH < 7.3, HCO$_3$ < 1.5 mmol/L), which can present with lethargy, drowsiness and Kussmaul’s breathing.

**Management**

Stable patients (i.e. GCS 15 and normal vitals with no focal neurological deficits) with more chronic behavioural change or developmental delay/regression may be referred for an early neurology consult. Otherwise, most patients need to be referred to an emergency department.

While there may be limited resuscitative ability in the general clinic, some simple measures can be instituted while awaiting transfer. Hypoglycaemia can be easily treated with oral feeds (simple sugars, e.g. sweets, juice or a glucose gel) or an intravenous 2 mL/kg bolus of 10% dextrose if the child is symptomatic or unable to tolerate orally. There is often concomitant dehydration in diabetic ketoacidosis, but fluid boluses should be avoided unless there is haemodynamic shock, as rapid rehydration increases the risk of cerebral oedema. Ongoing seizure lasting more than five minutes can be aborted with either rectal diazepam 0.4 mg/kg (max dose 10 mg) or intravenous lorazepam 0.1 mg/kg (max dose 4 mg). Each dose can be repeated once with a maximum of two doses of benzodiazepines using either route. The child should be placed in a lateral position on a level surface, preferably in a well-ventilated area, and oxygen can be administered to keep saturations above 95%. These interim measures help to stabilise the patient before the transfer.

**Further management in the tertiary setting**

Once the child has been referred and stabilised appropriately, more detailed investigations and subsequent management can be undertaken in the tertiary setting. Diagnostic studies in hospitals are generally approached in two stages. A first-line screen includes basic haematology, biochemistry and pathogen identification in blood, cerebrospinal fluid and other specific sites dictated by the initial assessment. Autoimmune and metabolic screening is added if there was developmental delay, systemic symptoms or a significant family history at presentation. If there

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**Table IV. Glasgow Coma Scale (GCS).**

<table>
<thead>
<tr>
<th>Score</th>
<th>Modified GCS for infants</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spontaneous</td>
<td>Spontaneous</td>
</tr>
<tr>
<td>To speech</td>
<td>To speech</td>
</tr>
<tr>
<td>To pain</td>
<td>To pain</td>
</tr>
<tr>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Oriented</td>
<td>Coos/babbles</td>
</tr>
<tr>
<td>Confused</td>
<td>Irritable</td>
</tr>
<tr>
<td>Inappropriate words</td>
<td>Cries to pain</td>
</tr>
<tr>
<td>Incomprehensible sounds</td>
<td>Moans to pain</td>
</tr>
<tr>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>-directed</td>
<td>Normal spontaneous</td>
</tr>
<tr>
<td>Withdraws to touch</td>
<td>Withdraws to pain</td>
</tr>
<tr>
<td>Abnormal flexion</td>
<td>Abnormal flexion</td>
</tr>
<tr>
<td>Abnormal extension</td>
<td>Abnormal extension</td>
</tr>
<tr>
<td>None</td>
<td>None</td>
</tr>
</tbody>
</table>

**Box 1. Mini Mental State Examination:**

<table>
<thead>
<tr>
<th>Orientation</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Date and place of birth</td>
</tr>
<tr>
<td>• Country and place</td>
</tr>
<tr>
<td>• Current time, day and date</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Registration</th>
</tr>
</thead>
<tbody>
<tr>
<td>• ‘I will name three objects for you to remember;’ (e.g. watch, pen, apple)</td>
</tr>
<tr>
<td>• ‘Repeat them after me.’</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Attention and Calculation</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Five serial 7s (serially subtract 7 from 100) or for younger children, five serial 3s (serially subtract 3 from 20)</td>
</tr>
<tr>
<td>• Spelling WORLD backwards</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Recall</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Ask for names of objects learnt earlier</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Language</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Name two objects or body parts</td>
</tr>
<tr>
<td>• Obey a three-stage command (e.g. take this paper with your left hand, fold it and put it on the table)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Copying</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Copy a pair of intersecting pentagons</td>
</tr>
</tbody>
</table>
are persistent symptoms or clinical deterioration, investigations are extended to include a wider search for atypical organisms, such as TB, and more specialised tests for central nervous system inflammation, such as anti-NMDAR receptor antibodies.

Computed tomography of the brain is quick and detects intracranial bleeds and focal lesions, but magnetic resonance imaging is the preferred imaging modality for accurate and sensitive identification and characterisation of various pathology with no risk of ionising radiation. Demyelinating disorders and an increasing number of viral-associated encephalitis now have recognisable clinico-radiological phenotypes, which facilitates early diagnosis and treatment. Magnetic resonance spectroscopy and magnetic resonance angiography are also helpful in neurometabolic and vascular aetiologies. Electroencephalography can be utilised to detect subclinical seizures, assess the degree of encephalopathy and localise areas of pathology. Genetic testing is also increasingly employed to aid in diagnosis, especially in epileptic encephalopathies and neurometabolic disorders.

Specific treatment will subsequently be guided by the likely aetiology. In acute-onset encephalopathy, early broad-spectrum antimicrobials may be initiated while awaiting diagnostic studies. First-line therapy consists of intravenous ceftriaxone and acyclovir, with the addition of ciprofloxacin if mycoplasma infection is suspected. Immunotherapy with intravenous methylprednisolone and/or immunoglobulin is given for probable autoimmune encephalitis. General neuroprotective measures are instituted to prevent secondary cerebral injury. More chronic encephalopathy will require detailed workup and neurorehabilitative intervention to optimise the patient’s functional status.

**TAKE HOME MESSAGES**

1. Encephalopathy is a broad term characterised by impaired awareness, cognition and behaviour, and can be caused by a wide range of conditions from systemic to primary neurologic disorders.

2. Common clinical mimics of encephalopathy include post-ictal drowsiness, hypoglycaemia and toxin ingestion. A good history and physical examination should be able to identify causes.

3. A useful approach to narrowing the differential diagnoses is determining the onset and evolution of symptoms in specific age groups, followed by thorough and directed history taking and examination that includes GCS scoring and the Mini Mental State Examination.

4. As neonatal encephalopathy and seizures can present subty, a high index of suspicion is required. Video-recording of episodes can be very useful in characterising their semiology.

5. A simple hypocount or point-of-care test can be performed in the primary care clinic setting to rule out hypoglycaemia and diabetic ketoacidosis.

6. Acute seizure management involves placing the patient in a lateral position with supplemental oxygen in a well-ventilated room. If the seizure persists beyond five minutes, rectal diazepam or intravenous lorazepam should be administered.

You called Brian’s main caregiver, his grandmother, and found out that his temperature had been spiking since the day before. He was also taking less than half his usual share of food, and was lethargic and minimally interactive for most of the day, even during the times when he was afebrile. In view of the above, you quickly sent Brian via ambulance to the children’s emergency department after ensuring that he was otherwise stable. Two months later, his grandmother thanked you for your prompt assessment and transfer. It meant that Brian received timely treatment for his viral encephalitis and subsequently had a full recovery. He suffered no long-term complications and was back to his usual cheerful self.

**REFERENCES**


SINGAPORE MEDICAL COUNCIL CATEGORY 3B CME PROGRAMME
(Code SMJ 202012A)

1. Encephalopathy describes an alteration of awareness and behaviour, and can present not only as drowsiness but also agitation and confusion.
2. All children with altered behaviour must be referred to the hospital.
3. A simple febrile fit is a common cause of encephalopathy.
4. Neonatal seizures are most often focal and can be difficult to distinguish from other non-epileptic movements.
5. An 18-month-old child seemed lethargic, only opening her eyes when called. On examination, she was fretful and reached out spontaneously for her mother. She only cried during the consult and did not say any words or babble, although her mother said she was usually able to say ‘yes’, ‘no’ or ‘Mama’. Her Glasgow Coma Scale (GCS) score is 14.
6. A developmental assessment should always be done, as stagnation or regression of milestones can accompany encephalopathy in more chronic cases.
7. Encephalopathy can be the first presenting symptom of systemic lupus erythematosus.
8. Accidental toxin ingestion is a possible cause of encephalopathy in a previously well three-year-old who seemed lethargic during the past few hours.
9. Hypoglycaemia can mimic encephalopathy and should be checked, especially in children aged under one year with a history of poor feeding and lethargy.
10. An 11-year-old girl presented with a one-week history of auditory hallucinations and decreased verbal output. Her mother said that she was sticking out her tongue and licking her lips frequently in the past three days. You should refer her for a psychiatric consult.
11. Infection is an important cause of encephalitis to exclude in children of all age groups.
12. A drop in GCS score of > 2 with tachycardia and elevated blood pressure is a red flag for immediate transfer to the emergency department.
13. All causes of encephalopathy are attributed to neurological conditions.
14. A teenager who has a GCS score of 15 but is unable to recall objects and obey a three-stage command on the Mini Mental State Examination should be suspected to be encephalopathic.
15. If a child looks lethargic and the hypocount shows a blood sugar level of 3 mmol/L, a cup of juice or a sweet should be offered and a repeat hypocount check done in half an hour.
16. Electroencephalography is a diagnostic test for epilepsy.
17. A two-year-old child presented with a fever of 38.5°C and a generalised seizure that had been going on for five minutes. Are these statements true or false?
   (a) The child should be placed in a prone position.
   (b) She requires supplemental oxygen.
   (c) You should give her rectal diazepam 0.4 mg/kg, which can be repeated once if the seizure does not stop.
   (d) Refer the patient to a general neurology clinic if the seizure has aborted and she is well.

Doctor’s particulars:
Name in full: ____________________________________ MCR no.: __________________________
Specialty: ____________________________________ Email: __________________________________

SUBMISSION INSTRUCTIONS:
Visit the SMJ website: http://www.smj.org.sg/current-issue and select the appropriate quiz. You will be redirected to the SMA login page.
For SMA member: (1) Log in with your username and password (if you do not know your password, please click on ‘Forgot your password?’). (2) Select your answers for each quiz and click ‘Submit’.
For non-SMA member: (1) Create an SMJ CME account, or log in with your SMJ CME username and password (for returning users). (2) Make payment of SGD 21.40 (inclusive of 7% GST) via PayPal to access this month’s quizzes. (3) Select your answers for each quiz and click ‘Submit’.

RESULTS:
(1) Answers will be published online in the SMJ February 2021 issue. (2) The MCR numbers of successful candidates will be posted online at the SMJ website by 8 February 2021. (3) Passing mark is 60%. No mark will be deducted for incorrect answers. (4) The SMJ editorial office will submit the list of successful candidates to the Singapore Medical Council. (5) One CME point is awarded for successful candidates. (6) SMC credits CME points according to the month of publication of the CME article (i.e. points awarded for a quiz published in the December 2020 issue will be credited for the month of December 2020, even if the deadline is in February 2021).