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# Toxic-metabolic encephalopathy in adults: critical discussion and pragmatical diagnostic approach.

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### Abstract:

Toxic-metabolic encephalopathy (TME) results from an acute cerebral dysfunction due to different metabolic disturbances including medications or illicit-drugs. It can lead to altered consciousness, going from delirium to coma, which may require intensive care and invasive mechanical ventilation. Even if it is a life-threatening condition, TME might have an excellent prognosis if its etiology is rapidly identified and treated adequately. This review summarizes the main etiologies, their differential diagnosis, and diagnostic strategy and management of TME with a critical discussion on the definition of TME.

**Keywords:** toxic-metabolic encephalopathy, metabolic encephalopathy, ICU, inborn error of metabolism, disorders of consciousness, coma, delirium, encephalopathy, blood-brain barrier.

# **Abbreviations**

CSF: cerebrospinal fluid

CT: computerized tomography EEG: electroencephalogram

cEEG: continuous electroencephalogram FLAIR: fluid-attenuated inversion recovery

HE: hepatic encephalopathy ICU: intensive care unit

IEM: inborn error of metabolism

MELAS: mitochondrial encephalopathy with lactic acidosis and stroke-like episodes

MERRF: myoclonic epilepsy with ragged red fibers

TME: toxic-metabolic encephalopathy

#### Introduction

Among the many causes of altered consciousness, toxic-metabolic encephalopathy (TME) is probably the most common evoked diagnosis. However, whereas specific causes are well defined, the definition of TME is complex and a generally accepted definition is still lacking. Metabolic encephalopathy, and the term toxic frequently added, is nowadays however a commonly used term in medicine. We should keep in mind that its delineation is blurred and its existence as a nosological entity can be questioned. Indeed, whereas metabolic encephalopathy has initially been proposed as the cerebral consequences of one major organ dysfunction, it has progressively been applied to all hemispheric neurological symptoms without any focal sign [1]. This probably explains some concurrence between a purely symptomatic description of neurological status compared to a more pathophysiological description that could be, imperfectly, expressed by toxic-metabolic encephalopathy [2]. One must nevertheless keep in mind the fact that while the pathophysiology of some metabolic encephalopathies is relatively well understood, the pathophysiology of most of others remains largely unknown. The absence of a precise nosological description has probably hindered pathophysiological studies in the past.

Despite these shortcomings, in the common acceptation of TME, the term refers to a potentially reversible pathological process within the brain, arising from an extracerebral origin, secondary to different metabolic disturbances including the effect of medications/drugs and their withdrawal.

The clinical presentation of TME is nonspecific and implies altered consciousness, without, in general, any focal and/or asymmetrical neurological signs. Abnormal movements such as asterixis or myoclonus can be found. This life-threatening condition could account for 15% of all causes of coma in the intensive care unit

(ICU), with the limitations that have been raised previously [3][4]. Thus, an accurate diagnostic approach is critical for proper management.

This review will successively discuss the following points:

- (1) How is TME most commonly defined and what are the main causes?
- (2) What are the clinical presentations of TME?
- (3) What are the main differential diagnoses to rule out prior to evoking TME?
- (4) What etiological workup should be performed?
- (5) How should TME be managed and what is the possible outcome?
- (6) Some perspectives

# 1. How is TME most commonly defined and what are the main causes?

# 1.1 Commonly accepted definition and pathophysiology

According to the recent nomenclature updated by an expert consensus, the term acute encephalopathy should be defined as a rapidly developing brain process which is expressed clinically as delirium or coma and may have additional features, such as seizures or extrapyramidal signs [2] (Supplemental Table 1). Thus, despite its initial limitation to only cerebral consequences of organ dysfunction, TME could be defined as a developing brain process secondary to a metabolic disturbance but is often described as multifactorial in origin [5][1][6]. This metabolic disturbance can be either innate, very rare, or acquired, most frequently, and is secondary to an organ failure, to altered homeostasis, the deficit or the accumulation of some endogenous or exogenous substances, including medications, illicit drugs or any neurotoxic substance.

This process could be responsible for compromised cellular functioning of neurons but probably also astrocytes, and for long-lasting compromised signal transmission

between different brain regions. Moreover, some brain areas such as the frontal regions could be particularly susceptible to metabolic disturbance [7]. Some circumstances associate multiple homeostasis disturbances, for instance as in hepatic encephalopathy (HE), where neurological symptoms can be secondary to hyperammonemia and systemic and cerebral inflammation, as well as intracerebral accumulation of bile acids and medications [8]. The implication of increased permeability of the blood-brain barrier (BBB) and the expression of specific transporters - ATP binding cassette transporter or efflux pumps - on it has been proposed [9][10] and could explain differences in the severity of the neurological signs despite very similar triggering events. Indeed, patients with previous cerebral alterations – elderly patients, patients with prior neurological diseases or patients without prior neurological diseases but with altered BBB permeability like cirrhotic patients for example – are more prone to develop encephalopathy: [11][12][13][14]. For instance, 20% of patients with liver cirrhosis display hepatic encephalopathy (HE) at diagnosis, and approximately 70% will develop HE during the disease course [8]. Increased permeability of the BBB has been recently shown in this condition [10][15] as as have the accumulation of efflux pump substrates in the cerebrospinal fluid (CSF). Most of the patients with encephalopathy in ICU have many medications due to their intensive care condition, sedatives, antimicrobial agents for nosocomial infection, or due to the cause of their admission to ICU [16][17]. Therefore, they might be more vulnerable to metabolic disturbance. Some polymorphisms of P-glycoprotein (P-gp) have been found to be associated with an increased risk of ICU delirium [18]. The number of prescribed medications could thus constitute a risk factor mainly through possible drug competition on BBB efflux pumps [19][10][20]. One illustrative example is the concomitant use of over-the-counter loperamide with another P-gp

substrate, frequently verapamil, to deliberately induce neurological toxic effects by loperamide, a "peripheral" opioid. Indeed, when taken concomitantly with verapamil that will be effluxed by P-gp, loperamide will accumulate inside the brain as attested by cerebral scintigraphy and positron emission tomography [19] and lead to alteration of consciousness.

### 1.2 Main causes of TME

Some etiologies of TME, such as anoxic-ischemic encephalopathy and hypercapnic encephalopathy, are easily recognized by physicians as they often occur in an obvious context of organ dysfunction: heart failure or cardiac arrest and respiratory failure, respectively. Hepatic encephalopathy or uremic encephalopathy are frequently associated with other causes of TME due to the patient's condition, especially septic-associated encephalopathy. Some toxic etiologies, for instance carbon monoxide poisoning, are more difficult to evoke if not in their common medical scenarios, particularly if the victim was alone at home without other victims. Illicit drug abuse is another example. Table 1 summarizes the main causes of acquired TME. Among TMEs due to deficiencies, hypoglycemia and Gayet-Wernicke's encephalopathy should be rapidly ruled out since these life-threatening conditions require rapid supplementation to preclude definitive neurological segualae. Gayet-Wernicke's encephalopathy is observed after exhaustion of thiamine (vitamin B1) reserves mainly in patients with chronic alcohol consumption [21]. Nowadays, it can also be encountered following obesity surgery, with poor or insufficient postoperative supplementation or following certain chemotherapies, which may interfere with thiamine absorption. Anecdotally, among rare causes of TME, endocrinological etiologies such as hypothyroidism or adrenal insufficiency can be cited, although the

cerebral hypometabolism observed in these conditions is simply the expression of a global body hypometabolism. Among TMEs secondary to the accumulation of a substance, either an endogenous or an exogenous substance, hyperammonemia and drug-induced encephalopathy should be promptly evoked as well [6][22]. A list of drugs and medication commonly associated with TME is shown in Table 2. It should be noticed that some drugs are able to induce hyperammonemia as well. In the ICU, medications responsible for encephalopathy are usually analgesics, antiepileptic medications and antimicrobial drugs, but neuroleptics, antidepressants, and immunosuppressive agents can also be responsible [23]. Another mechanism of encephalopathy, which is neither due to a deficiency nor an accumulation of a substance, we can cite rapid osmolar shift, responsible for osmotic demyelination. Usually, this entity is attributed to an overly rapid correction of hyponatremia [24]. The implication of this osmotic shift in cirrhotic patients with hyponatremia has been proposed to be implicated or associated with HE [25][26]. In a patient with a history of alcoholism or illicit drug abuse, acute alcoholic or another drug withdrawal must be evoked. Some specific causes such as vasculitis or specific TME, e.g. Chasing the *Dragon* in heroin users [27], could be looked for.

Finally, encephalopathy can be an acute manifestation of an inborn error of metabolism (IEM). IEMs are rare conditions mainly diagnosed in childhood, but those can occur in adulthood with an acute presentation and few or no symptoms beforehand [28]. IEMs involve brain functioning via three main mechanisms: *i*) intoxication with an endogenous substance due to its rapid accumulation; *ii*) an energy deficit due to deficiency in energetic metabolism processes; *iii*) cellular organelle dysfunction leading to a synthesis deficit regarding a crucial molecule or catabolism abnormalities, particularly in enzymes involved in the degradation of

complex molecules [29][30][28]. Table 3 summarizes IEMs, with their diagnostic investigations. Even though these diseases are rare, specific experience is needed for both diagnosis and management. In most developed countries, dedicated networks exist and it is helpful to identify the local contact upstream (see in France: http://www.filiere-g2m.fr or http://www.sfeim.org).

Due to its heterogeneous non-specific clinical presentation with symptom fluctuations, onset of encephalopathy requires an appropriate neurological examination and an accurate diagnostic approach, essential for tailored patient management.

# 2. What are the clinical presentations of TME?

Although not specific, subacute delirium with myoclonus in the absence of focal neurological sign should suggest TME [31].

#### 2.1 Delirium

The term *delirium* (Supplemental Table 1) is defined by a change from baseline, with disturbance in attention, awareness, and cognition (memory deficit, disorientation, language, visuospatial or perception disturbance) over a short period of time, with a trend to fluctuation in severity over time. To fulfilled *delirium* definition, these disturbances should not be explained by a pre-existing neurological disorder [2] [32]. The inability to pay attention is a subtle sign that appears precociously and should constitute a red flag. Altered awareness ranges from drowsiness to deep coma, and can be assessed using the Glasgow Coma Scale or the FOUR score (full outline of unresponsiveness) [33] [34]. It can lead to the loss of airway reflexes with a risk of airway obstruction, aspiration pneumonia or bradypnea, which may require ICU

admission and mechanical invasive ventilation. In the ICU, delirium can be scored using the Confusion Assessment Method for the ICU (CAM-ICU)[35]. This scale can be used even in mechanically ventilated patients and is a validated tool to assess delirium.

# 2.2 Sleep disorders

Reversal of circadian rhythm, insomnia, hypersomnia, and rapid eye movement sleep behavior disorder can be observed in TME, sometimes independently from delirium, and should be sought.

# 2.3 Myoclonus and asterixis

*Myoclonus*, a movement disorder characterized by brief and involuntary muscular jerks, is common in TME. Frequently visible in the upper limbs or on the face, it can sometimes be observed over the whole body. TME can also be revealed by *asterixis*, a sudden postural fall which is called negative myoclonus [36][37]. It is characterized by irregular and sustained intermittent muscle relaxation (distal negative myoclonus), and revealed classically in the upper limbs by the Barré test. It is quite common for these myoclonus to be misinterpreted as epileptic activity, both clinically and at EEG, and treated as such, thereby worsening TME and associated consciousness disorders [38].

### 2.4 Cranial nerve palsy

Even though a differential diagnosis should be ruled out first prior to evoking TME in the diagnostic approach of a cranial nerve palsy, this presentation is possible. It is an expression of brain stem dysfunction, with cranial nerve signs (oculomotor disturbances, nystagmus, or changes in pupil size), pathological brain stem reflexes, and bulbar syndrome [39].

# 2.5 Extrapyramidal signs and movement disorders

According to the TME etiology, extrapyramidal signs can be observed as well as other movement disorders [39]. These symptoms are classically observed during the accumulation of certain substances, such as excess copper during Wilson's disease or in manganese intoxication [40][41]. Extrapyramidal signs can also be observed in overt HE [42]. These are typically reversible after liver transplantation and have been associated with the accumulation of manganese ([43]). However, these symptoms are rarely seen in other causes of TME.

The diversity of clinical manifestations might be explained by the heterogeneity of brain regions involved depending on the related TME etiology.

# 3. What are the main differential diagnoses to rule out prior to evoking TME?

Table 4 summarizes the main differential diagnoses that should at least be discussed, clinical symptoms to look for and associated complementary examinations. Briefly, if a neurological localization sign is observed or if an acute coma occurs, the first urgent diagnosis to rule out is a vascular cause or a space occupying process, by brain imaging [44]. Note that post-traumatic brain injury is not considered as a TME. Bacterial meningitis has to be ruled out in a patient with fever and associated meningism [45]. Skin purpuric lesions, which in this context must lead to suspect an invasive meningococcal disease, should be sought. Similarly, fever associated with a focal neurological sign or seizures should raise the hypothesis of an ongoing brain infectious process (viral encephalitis, rhombencephalitis, brain

abscess or septic embolism from an endocarditis). Posterior reversible encephalopathy syndrome can also mimic a TME (see for review [46]). Immune-mediated encephalitis can be evoked in a patient with alteration of consciousness, focal neurologic signs, movement disorders and seizures (see for review [47]). However, it should be recalled that apart from very common biochemical or immunological tests, most of the biological work-up requires several weeks to obtain definitive results. The consequence is that the diagnosis of TME is generally retained before some results are known, based on the clinical context and minimal negative etiological work-up.

# 4. Which etiological work-up should be performed?

# 4.1 First-line etiological work-up

This first-line etiological work-up has as objectives to confirm TME, to determine its etiology and to rule out differential diagnoses (Table 4). Whereas EEG can confirm TME, brain imaging will mainly rule out differential diagnoses. Blood sampling will bring clues in favor of TME but will inconsistently enable definitive confirmation of TME.

#### Blood sampling

Capillary glucose level must be determined whenever there is an acute neurological symptom. Confirmation on a blood sample is required, but a low capillary glucose level should be treated rapidly by glucose administration. Blood chemistry with sodium and calcium levels and other blood tests – urea, creatinine, liver enzymes and prothrombin time – enables evoking the most common causes of TME: electrolyte disorder, kidney or liver failure. Blood gases can show evidence of

hypercarbia associated with respiratory failure [5]. Depending on the context, other tests including vitamin B1 assay before any supplementation, search for illicit-drugs or medications – especially, opioids, benzodiazepines, neuroleptics, amphetamines, and cannabinoids – in urine and blood samples, as well as blood tests for thyroid hormones, cortisol, vitamin B9, B12, C or PP can be ordered [48]. It can be helpful to store blood and urine samples, drawn before any supplementation, for further unplanned analyses.

In the absence of any definite diagnosis in the hours after symptom onset, ammonemia should be determined without delay and without waiting for the results of all tests. Indeed, even when the context is not suggestive of acute liver failure or chronic liver disease, hyperammonemia is an important finding since it could reveal hepatic encephalopathy associated with an undiagnosed liver disease, or with a porto-systemic shunt without liver disease; drug-associated hyperammonemia (sodium valproate, 5-fluorouracile or asparaginase for the most frequent ones) [49] is another possibility, or more rarely an undiagnosed IEM, especially urea cycle defect [28][50][30]. Some miscellaneous causes of hyperammonemia are also possible [51][52]. In the specific context of liver disease, neurological symptoms suggestive of HE with a normal ammonia level should prompt search for a differential diagnosis such aseptic or drug-associated encephalopathy. For example, using metabolomics our group recently showed that about 60% of cirrhotic patients presenting with encephalopathy had significant levels of antimicrobial agents in their CSF [53]. On the other hand, ammonia levels above 150 µmol/l, but the cut-off is still discussed, would be suggestive of IEM that can be associated with atypical liver presentation. It should however be recalled that some technical issues persist with ammonia assays [54].

# **Brain imaging**

TME diagnosis is based on history taking and physical examination, however, if there is a doubt or an atypical presentation, brain imaging may be needed to rule out major differential diagnoses and to exclude brain lesions (Table 2). If brain imaging is performed, magnetic resonance imaging should be preferred to computed tomography (CT) as it is more sensitive [55]. However, if unavailable, or cumbersome to perform as in the ICU, a CT-scan should at least be performed if no positive diagnosis has already been established. In most cases, brain imaging is normal in TME. Nevertheless, unfrequently, it can show either diffuse or focal edema or specific lesions as in Gayet-Wernicke's encephalopathy or metronidazole-associated TME. For instance, bilateral T2 and FLAIR hyperintensity of the dentate nucleus, the cerebellar peduncles and the inferior colliculus associated with olivary hypertrophy is evocative of a metronidazole-induced encephalopathy [56]. Similarly, T2 and FLAIR hyperintensity of the splenium of the corpus callosum have been described in patients treated with carbamazepine [57].

Magnetic resonance spectroscopy can also help the investigation. For instance, a classical spectroscopic profile of hepatic encephalopathy shows an increased peak of glutamine/glutamate and a decreased peak of choline and myoinositol [58].

# Electroencephalogram (EEG)

EEG is a primary tool for the positive diagnosis as it can detect and monitor TME in the ICU and because of its availability at bedside, its non-invasiveness, and its excellent temporal resolution. In case of TME, it will typically show the presence of slow background activity with fluctuating triphasic waves, maximal over anterior

regions, and an inconstant reactivity to stimulations (Fig. 1). However, an unreactive EEG pattern must raise the possibility of a differential diagnosis, such as status epilepticus or severe hypoxic ischemic encephalopathy. The degree of EEG disturbance correlates fairy well with the level of consciousness, and allows to graduate TME severity [59][60][61]. More specific patterns can sometimes be found such as unreactive pseudo periodic triphasic waves in cefepime-induced encephalopathy, an increase in frequency of the posterior dominant rhythm in hyperthyroidism, or frontal intermittent rhythmic delta activity in hyperglycemia or hyponatremia [62][63]. Several drugs including baclofen, barbiturates or propofol can also cause burst-suppression on EEG.

EEG can also detect interictal patterns, non-convulsive seizures or electrical status epilepticus that represent frequent differential diagnoses and possible complications of TME [38][64] (Fig. 2). More recently, several studies demonstrated the value of continuous electroencephalographic monitoring (cEEG) in the ICU. cEEG provides continuous monitoring of brain function and is recommended to identify non-convulsive seizures in patients with an altered mental status without known acute brain injury. cEEG may also provide some prognostic elements in hypoxemic ischemic encephalopathy [65] [66] [67] [68] [69] [70]. The place of cEEG in TME has however to be clarified in the future even though most neuro-ICUs use regularly it in this condition.

### Lumbar puncture

Lumbar puncture is normal in TME or IEM, outside specific techniques not routinely available [10] and may be performed mainly to rule out a differential diagnosis, especially in the case of altered consciousness with fever suggesting

meningoencephalitis. Quiet recently, two studies proposed that the concentrations of drugs or the level of neurofilaments in the CSF could be prognostic in TME [71] [72]. This has however to be confirmed.

# 4.2 If the first-line etiological work-up doesn't lead to a diagnosis

The most important points are probably to review all the tests that have been performed, and sometimes redo the doubtful ones, and go back to the medical history. Indeed, having a detailed history of symptom onset, of personal and familial history will enable to orient the second-line etiological work-up.

If an IEM is suspected, additional explorations – thoraco-abdominopelvic CT scan, ophthalmologic examination – should be proposed to identify frequently associated non-symptomatic lesions in other organs, or in the spine or the peripheral or central nervous system. Spine magnetic resonance imaging, electroneuromyography and audiogram are sometimes needed. In some case, histological analysis is performed on a muscle or brain biopsy [73] [74].

Evidence of neurosensorial impairment (deafness, blindness), diabetes mellitus, family history in association with TME and myoclonus will lead to evoke mitochondrial encephalopathy and will need genetic testing for MELAS and MERF and muscular biopsy. An acute encephalopathy onset in a context of fasting, high protein intake, pregnancy, previous episodes of altered consciousness, unexplained rhabdomyolysis or metabolic acidosis will lead to evoke an IEM. If so, an expert center should be rapidly contacted, and preventive measures urgently taken. Specific dosages or genetic analysis (one or many specific genomic regions with next generation

sequencing, or whole exome sequencing) are needed but are guided by the expert's advice [75].

Broader diffusion of metabolomic techniques, liquid chromatography coupled to mass spectroscopy, now also enables posteriori identification of thousands of toxic substances based on hair analyses [76].

# 5. How should TME be managed and what is the possible outcome?

Management of TME includes symptomatic management including neuroprotective measures and etiological management. Whatever the etiology of TME, management consists in avoiding fever, controlling mean arterial blood pressure, glucose and sodium levels, avoiding hypercarbia but also severe hypocarbia that has been associated with cerebral ischemia. Several studies recently suggested that hyperoxia could be deleterious, thus systemic oxygen administration should probably be avoided and oxygen delivered only if oxygen level is low [77]. In the ICU, some simple environmental measures are frequently added and recommended: quiet ambiance with low intensity lights and night switch off, ear plugs and eye masks, displaying clocks in each room [78]. If psychomotor agitation is present, haloperidol and dexmedetomidine are the preferred drugs [79]. In case of a non-rapidly reversible comatose state, mechanical ventilation can be mandatory. Seizure management have no specificities in TME outside the frequent pitfall of misdiagnosing myoclonus or triphasic waves as seizures or epileptic activity [38]. Some authors prefer to avoid antiepileptic drugs that can be responsible for TME in this condition and suggest avoiding sodium valproate in patients with previously altered brains.

The etiological management of TME is of major importance. Electrolyte disorders must be corrected, glucose be given in case of hypoglycemia, B1 supplementation prescribed in Gayet-Wernicke's encephalopathy, invasive or non-invasive mechanical ventilation be started in case of hypercapnic encephalopathy, the cause of sepsis be identified and antimicrobial agents be given. If encephalopathy onset is secondary to medications, simplifying the prescription is of major importance. Depending on the associated causes of TME and the possible complex interaction between the different drugs on efflux pumps, stopping every non-indispensable molecule is an easy practical advice. If one specific compound is clearly suspected, cefepime for example, its replacement by another equivalent drugs is indicated. Whereas in the case of TME associated for example to lithium or phenytoin intoxication, renal replacement therapy can be discussed [80], some others associated with toxic alcohol, methotrexate or ifosfamide specific antidotes can be proposed [81] [82] [83]. IEM treatment should be done in collaboration with the local expert center. In the first hours, it is important to avoid further degradation by applying a preventive strategy. In case of urea cycle defects, protein restriction during the first 24 hours should be done only in association with the administration of glucose and lipids in order to prevent endogenous protein catabolism [28]. Whatever the etiology of TME, any precipitant factor should be treated; infection being probably the most frequent one.

Very few data on outcome of TME are available. The large range of etiologies preclude definitive answers. However, due to the theorical reversible nature of TME, it appears, by experience, that the prognosis of TME is good. This is an important point especially for intensivists and has two major implications. First, patients with suspected TME should be admitted in the ICU and second it is important to give the

patients with TME some time to improve before discussing care withdrawal or with-holding. In some circumstances, where the disturbance has lasted for a long time and management has been delayed, definitive neurological sequalae can be present.

This reinforces the need for early recognition and early management/treatment [84].

# 6. Perspectives

We previously discussed shortcomings of the term toxic-metabolic encephalopathy. Thus, how to deal with it? Quiet recently, it has been shown that the use of the term encephalopathy or delirium in original articles in Pubmed were largely dependent on the specialty of the authors [2]. Whereas internists, neurologists mainly use encephalopathy; anesthesiologists, intensivists, psychiatrics, and geriatrics use delirium. Furthermore, the authors showed that the corresponding published literature is highly segregated. A panel of experts using a modified Delphi method proposed the definition presented in **Supplemental Table 1** for encephalopathy and delirium and discarded the use of some other terms. Should a similar approach be applied to TME? Indeed, the imperfect shape precludes for more detailed studies on epidemiology, pharmacology, pharmacoepidemiology or pathophysiology. Only such an approach could improve our knowledge and provide strong evidence for better management.

### 7. Conclusion

TME is a frequently encountered condition in the emergency room, the ICU and/or in patients with chronic diseases. Even if most cases are reversible and have good outcome, delayed diagnosis and management can be associated with worse

outcome and neurological sequalae. This reinforces the need for an urgent diagnosis and management. Clinically, the presentation is unspecific with altered consciousness, without focal signs classically, and can be associated to asterixis or myoclonus. Apart the medical history and the context, EEG is of major importance to make a positive diagnostic of TME whereas brain imaging is mainly important to rule out differential diagnosis. In the absence of any definite diagnosis, ammonia should be measured. Past medical history of sensory disturbance, neuropathy, epilepsy in childhood, consanguineous marriage in the family or an unknow neurological disease in patient's relative can raise the possibility of an IEM. TME management encompass symptomatic measures aimed to avoid neurological worsening and the treatment of the cause of TME. Some technical evolution, metabolomics or improved brain imaging, will probably provide a more rapid diagnosis of TME in a near future [22] [53] [85] [86].

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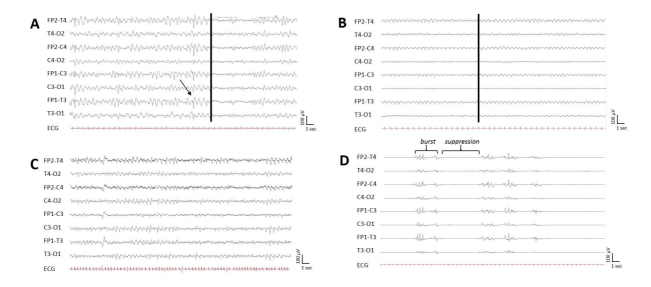
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# Legends

Figure 1: Various representative EEG patterns of TME.



# EEGs are in bipolar montage

A: hepatic encephalopathy pattern, with slow activity (2-3Hz) and fluctuating triphasic waves (shown by the arrow), maximal over anterior regions, with reactivity to stimulation (shown by the vertical line)

B: more severe hepatic encephalopathy pattern, with a continuous slow background activity, with no reactivity to stimulation (shown by the vertical line)

C: pseudo-periodic discharges with triphasic morphology, over the frontocentral area, in cefepime-induced encephalopathy

D: burst-suppression pattern: alternating episodes of isoelectric EEG and bursting slow waves, induced by thiopental

Abbreviations: EEG: electroencephalography; TME: toxic-metabolic encephalopathy

Altered consciousness Identified non-modifiable risk +/- myoclonus factors of TME: STEP 1: (Perform capillary blood glucose) -Elderly -Prior neurological disease -Cirrhosis/Kidney disease -Focal neurological sign -Polymedication Brain imaging YES -Meningism, fever +/- lumbar puncture -Visual field loss Seizures **NORMAL** YES Possible TME First-line biological assessment : EEG: TME pattern Metabolic disturbance or Positive toxic test YES YES NO **Confirmed TME** YES Second line assessment NB: keep in mind that TME can be associated to an other

Figure 2: Diagnostic algorithm for suspected toxic-metabolic encephalopathy

Abbreviations: EEG: electroencephalography; TME: toxic-metabolic encephalopathy; IEM: inborn error of metabolism

Metabolic disturbance/IEM

Table 1: Main causes of acquired TME.

Metabolic encephalopathy			
Hepatic encephalopathy	Acute or chronic liver failure		
	Portosystemic shunt		
Hypercapnic encephalopathy	End-stage COPD		
	Acute respiratory failure		
	Diaphragmatic dysfunction		
Uremic encephalopathy	Kidney failure		
Electrolyte disturbances	Hyper/hyponatremia		
	Hyper/hypocalcemia		
Endocrine disorders	Hyper/hypoglycemia		
	Hyper/hypothyroidism		
	Hyper/hypocortisolism		
Deficiency-related encephalopathy	Thiamin (vitamin B1) deficiency: Gayet-Wernicke's		
	encephalopathy		
	Niacin (vitamin B3 or PP) deficiency: pellagrous		
	encephalopathy		
	Pyridoxine (vitamin B6) deficiency		
	Folate (vitamin B9) deficiency		
	Cyanocobalamin (vitamin B12) deficiency		
	Ascorbic acid (vitamin C) deficiency: scurvy		
	Vitamin E deficiency (cerebellum)		
Septic-associated encephalopathy	Sepsis, septic shock		
	Acute pancreatitis, ARDS		
Anoxic-ischemic encephalopathy	Cardiac arrest		
	Shock		
Toxic encephalopathy			
Drug-induced encephalopathy	see Table 3		
Alcohol abuse			
Illicit drug abuse	Methamphetamine, heroin, cocaine, synthetic cathinones		
Other toxic encephalopathy	Heavy metal (lead, manganese, mercury)		
	Environmental agents (methanol, ethylene glycol, carbon		
	monoxide)		
	Organophosphate		
Withdrawal syndrome	Alcohol		
	Benzodiazepine		

Abbreviation: ARDS, Acute Respiratory Distress Syndrome; COPD, Chronic obstructive Pulmonary Disease

Table 2: Main drugs that could lead to toxic metabolic encephalopathy.

Antiepileptics	Benzodiazepines	
	Valproic acid*	
	Barbiturates*	
	Phenytoin	
	Gabapentin	
	Lacosamide	
	Carbamazepine#	
Psychiatrics	Tricyclic antidepressants	
	Selective serotonin reuptake inhibitors	
	Neuroleptics	
	Lithium	
Oncologic	Methotrexate*	
	L-asparaginase*	
	5-fluoro-uracil*	
	Ifosfamide	
Immunosuppressants	Calcineurin inhibitors	
	Tacrolimus	
Antimicrobial agents	Betalactams (including carbapenems, cefepime)	
	Fluoroquinolone	
	Metrodinazole#	
	Linezolid	
	Foscavir, aciclovir	
	Interferon alpha	
	Fluconazole	
Miscellaneous	Dopamine agonists	
	Levodopa	
	Opioids	
	Proton pump inhibitors	
	Baclofen	
	Loperamide	

<sup>\*</sup>can be associated with hyperammonemia; if associated with a P-gp inhibitor such as verapamil or proton pumps inhibitor #brain magnetic resonance imaging can be abnormal

Table 3: Categories of inborn errors of metabolism in adults, with their main diagnostic investigations.

Intoxication by an endogenous substance				
Amino acid disorders	Amino acids chromatography in blood and urine, homocysteine assay			
homocystinuria, leucinosis				
Organic acidurias	Organic acids chromatography in blood and urine, acylcarnitine profile			
1	onic acidemia (cobalamin disorders), methylmalonic acidemia hylmalonic acidemia (methylmalonyl-CoA mutase)			
Urea cycle defects	Ammonemia, amino acids chromatography, urine orotic acid			
OTC, CPS, ASA				
Metal storage disorders	Serum copper, ceruloplasmin, iron, ferritin, transferrin assay, hepatic biopsy, genetic testing			
Wilson disease, hemochromatosis, neurodegeneration with brain iron accumulation				
Porphyrias	5-aminolevulinic acid, urinary porphobilinogen assay			
Energy deficiency				
Mitochondrial disorders	Lactate/ pyruvate in blood and CSF, stress test, ENMG, muscle biopsy, genomic sequencing, molecular analyses, extra-neurological evaluations			
MERRF, MELAS, POLG, Leigh, Ke	earn-Sayre			
Glycogen storage diseases	Hepatic/muscle biopsy, lactate level, genetic testing			
Disorders of fatty acid oxidation	Organic acids chromatography, acylcarnitine profile			
Carnitine uptake defect/carnitine tra	ansport defect			
Deficiency of complex molecules	s synthesis or catabolism			
Peroxisomal disorders	Phytanic acid, pristanic acid assay, serum very-long-chain fatty acid			
Refsum disease	1			
Lysosomal storage diseases	Enzyme assay, genomic sequencing			
Gaucher disease, Fabry disease				
Congenital disorders of glycosylation	Carbohydrate deficient transferrin and protein-linked glycan analysis, genomic sequencing			
grycosyration	genomic sequencing			

Abbreviations: CPS, carbamyl phosphate synthetase; MERRF: myoclonic epilepsy with ragged red fibers, MELAS: Mitochondrial Encephalomyopathy, Lactic Acidosis and Stroke-Like Episodes, OTC, ornithine transcarbamylase; CSF: cerebrospinal fluid ENMG: electroneuromyogram.

Table 4: Differential diagnosis of TME, with clinical manifestations and complementary examinations.

Differential diagnosis	Clinical manifestations	Complementary examinations
- Infectious meningoencephalitis	Fever Headache Meningismus Subacute onset Skin purpuric lesion	Lumbar puncture Brain imaging
Autoimmune encephalitis	Neuropsychiatric manifestations Seizures Movement disorders Autonomic dysfunction	Brain imaging Lumbar puncture Blood/CSF antibodies - neuronal cell-surface - synaptic receptors EEG
Posterior reversible encephalopathy syndrome	Headaches High blood pressure Elevated plasmatic creatinine	Brain imaging EEG
Convulsive or non-convulsive status epilepticus	Generalized or unilateral Myoclonus Seizures	EEG / cEEG
Psychiatric or cognitive disorders	Normal neurological examination	Psychiatric evaluation
Cerebrovascular disease - Ischemic stroke - Intracerebral hemorrhage - Subarachnoid hemorrhage - Cerebral venous thrombosis - Sub/extra-dural hematoma	Neurological focal sign Sudden onset Headache Possible intracranial Hypertension syndrome	Brain imaging
Space occupying brain lesion - Tumor - Abscess - Inflammatory lesion / ADEM	Neurological localization sign Chronic or subacute onset Intracranial hypertension syndrome	Brain imaging ± Lumbar puncture ± Brain biopsy

Abbreviations: ADEM: acute disseminated encephalomyelitis; CSF: cerebrospinal fluid; EEG: electroencephalogram; cEEG: continuous EEG.

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