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## Review

# Pediatric refractory and super-refractory status epilepticus



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# ABSTRACT

*Purpose*: To summarize the available evidence related to pediatric refractory status epilepticus (RSE) and super-refractory status epilepticus (SRSE), with emphasis on epidemiology, etiologies, therapeutic approaches, and clinical outcomes.

Methods: Narrative review of the medical literature using MEDLINE database.

Results: RSE is defined as status epilepticus (SE) that fails to respond to adequately used first- and second-line antiepileptic drugs. SRSE occurs when SE persist for 24 h or more after administration of anesthesia, or recurs after its withdrawal. RSE and SRSE represent complex neurological emergencies associated with long-term neurological dysfunction and high mortality. Challenges in management arise as the underlying etiology is not always promptly recognized and therapeutic options become limited with prolonged seizures. Treatment decisions mainly rely on case series or experts' opinions. The comparative effectiveness of different treatment strategies has not been evaluated in large prospective series or randomized clinical trials. Continuous infusion of anesthetic agents is the most common treatment for RSE and SRSE, although many questions on optimal dosing and rate of administration remain unanswered. The use of non-pharmacological therapies is documented in case series or reports with low level of evidence. In addition to neurological complications resulting from prolonged seizures, children with RSE/SRSE often develop systemic complications associated with polypharmacy and prolonged hospital stay.

*Conclusion:* RSE and SRSE are neurological emergencies with limited therapeutic options. Multi-national collaborative efforts are desirable to evaluate the safety and efficacy of current RSE/SRSE therapies, and potentially impact patients' outcomes.

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## 1. Introduction

Refractory status epilepticus (RSE) is a life-threatening neurological emergency associated with significant morbidity and mortality. RSE is defined as seizure activity that persists after administration of a first-line benzodiazepine (BZD) and a second-line antiseizure drug (ASD) [1]. Primarily encountered and treated

Abbreviations: ASD, Anti-seizure drug; BZD, Benzodiazepine; CEEG, Continuous Encephalography; CI, Continuous infusions; CNS, Central Nervous System; FIRES, Febrile illness-related epilepsy syndrome; GCSE, Generalized convulsive status epilepticus; ICU, Intensive Care Unit; KD, Ketogenic diet; NCS, Neurocritical Care Society; NCSE, Non convulsive status epilepticus; NORSE, New onset refractory status epilepticus; RSE, Refractory status epilepticus; RCSE, Refractory convulsive status epilepticus; SRSE, Super-refractory status epilepticus; TBI, Traumatic brain injury.

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in the intensive care unit (ICU), RSE patients usually receive additional boluses of second-line ASDs (e.g. fosphenytoin, levetiracetam, and valproate) or are placed in a medically induced coma with intravenous (IV) continuous infusions (CI) of an anesthetic (e.g. midazolam, propofol, barbiturates) for seizure control. Nevertheless, continuous or intermittent seizures may persist for 24 h or more following the administration of general anesthesia or recur after its withdrawal. The resulting condition is known as super-refractory status epilepticus (SRSE) [2]. Identifying the underlying etiology of RSE and SRSE can be challenging. Treatment decisions mainly rely on case series or experts' opinions. The comparative effectiveness of heterogeneous treatment strategies has not been systematically evaluated in large prospective series or randomized clinical trials.

The annual incidence of status epilepticus (SE) is estimated to be 17–23 episodes per 100,000 children [3,4]. Of these SE patients, between 10% and 40% develop RSE [5–7] with a mortality rate of 16–43.5% [8–10]. The few epidemiological studies on SRSE are based mainly in adult population [11] indicating that 10–15% of

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RSE cases progress to SRSE [12,13], and approximately one-third of RSE and SRSE patients die [12–14]. In children, a retrospective study showed that of 602 convulsive SE episodes, SRSE occurred in 7.14% [15]. In the United States, most data come from case series or case reports. Without large series, estimating the exact incidence and mortality of SRSE is challenging. In this review we present existing evidence on pediatric RSE/SRSE including clinical presentations and etiologies. We address current diagnostic and treatment approaches, as well as clinical outcomes of RSE/SRSE in children.

## 1.1. Clinical presentation

Patients who develop RSE/SRSE present clinically in different ways. The most common presentation is generalized convulsive SE (GCSE), the major seizure type for pediatric SE [16–18], or focal SE with impaired consciousness after failure to initial treatment. Other common clinical presentations are subtler and thus, are suggestive of non-convulsive SE (NCSE). In the latter clinical scenario, patients usually present with stupor/coma after a GCSE or after an acute brain insult [19,20]. Although convulsive SE often evolves to NCSE, the distinction between these two phenotypes is crucial as the therapeutic approaches and outcomes are different, being NCSE more often associated with medical refractoriness due to delayed recognition [20].

## 1.2. Etiology

The etiologies for pediatric RSE and SRSE reported in the literature vary. The most commonly reported etiologies include: acute symptomatic causes (e.g. presumed infectious or immune mediated encephalitis, central nervous system (CNS) infections, traumatic brain injury (TBI), brain ischemia), remote symptomatic with acute precipitant causes (e.g. CNS lymphoproliferative disease, human immunodeficiency virus (HIV) infection, hypoxic-ischemic encephalopathy, developmental delay, epilepsy), remote symptomatic and progressive encephalopathies (e.g. Alpers disease, metabolic diseases such as medium chain acyl-CoA dehydrogenase deficiency, epileptic encephalopathies), febrile SE (excluding CNS infections), and unknown etiologies (e.g. cryptogenic) [3,8,21,22]. Nonetheless, previous studies show that etiology varies according to age groups and geographic location. In a study of 151 refractory convulsive SE (RCSE) episodes, the most common etiology was acute symptomatic (28.5%) in neonates and infants; prolonged febrile convulsions (33.8%) in children 1-5 years; and remote symptomatic etiologies in 40% of patients between 5 and 10 years old, and in 36.8% patients between 10 and 16 years old [23]. In contrast, the etiologies contributing to RSE and SRSE development in the adult population include acute brain injury (e.g. cerebrovascular disease, CNS infections, brain tumor, traumatic brain injury), intoxication/withdrawal syndromes, low levels of antiepileptic drugs, metabolic distrubances, and systemic infections [19]. In developing countries, acute symptomatic etiology remains the most frequent etiology in children, followed by remote symptomatic and unknown etiologies. Within the acute symptomatic etiology, CNS infections (e.g. herpes simplex virus (HSV), HIV, neurocysticercosis, malaria, tuberculosis), viral/autoimmune encephalitis and meningitis are common causes [7,24-30].

New Onset RSE (NORSE), on the other hand, is a clinical presentation described in patients without epilepsy or a relevant preexisting neurological disorder, who present with RSE without an identifiable acute cause or active structural, toxic or metabolic cause [31]. A subcategory of NORSE known as febrile infection-related epilepsy syndrome (FIRES) commonly presents in

previously healthy school-aged children [32,33]. These patients usually have a preceding febrile illness, with fever starting between 2 weeks and 24h prior to the onset of SE, and can present with or without fever at the onset of SE [31]. The etiology of this syndrome remains unknown, and the seizures in this patient population are notoriously difficult to control in both the acute and chronic settings. While NORSE and FIRES are relatively rare epilepsy syndromes, they are frequent in the SRSE population. In a small study of ketogenic diet (KD) usage for SRSE, the most common etiologies included encephalitis/FIRES (55.55%), followed by FIRES (22.22%), epileptic encephalopathy (11.11%), and central nervous system-hemophagocytic lymphohistiocytosis (11.11%) [34]. Another study showed that 40% patients with SRSE had a diagnosis of immune-mediated encephalitis (Rasmussen's syndrome, post-infectious mycoplasma encephalitis and Anti-Nmethyl-D-aspartate (NMDA) receptor encephalitis), followed by FIRES (20%), genetic epilepsies (PCDH19 and GABRG2 mutations) (20%), epilepsy of a known metabolic etiology (10%) and NORSE (10%) [35]. Therefore, when initial diagnostic workup reveals negative results, these epilepsy syndromes should be considered in the differential diagnosis.

# 1.3. Pathophysiology

In RSE and particularly SRSE, mechanisms responsible for seizure termination fail and additional pathophysiologic processes develop leading to persistence of SE. At a cellular level, SE intensifies the internalization of synaptic y-amino butyric acid type A (GABA-A) receptors whereas the function of extra synaptic receptors is preserved. This synaptic "receptor trafficking" leads to an overall reduction of the inhibitory activity of GABA, playing a key role in the development of pharmacoresistance [36]. Additionally, increased number of glutaminergic receptors at the neuronal surface may contribute to seizure perpetuation due to changes in concentrations of ions, like chloride, in the cellular environment. Furthermore, the persistence of seizures and development of SRSE may be explained by sensitivity to NMDAmediated neuronal stimulation [37], mitochondrial failure [38], blood-brain barrier damage, and neuro inflammation (i.e. proinflammatory cytokines, autoantibodies to neural elements) [2]. All of these factors result in excitotoxicity [39], which is directly responsible for neuronal injury and cell loss, and ultimately poor clinical outcomes. Finally, previous studies demonstrate the important role of time from seizure onset to treatment administration on seizure duration [40,41]. If treatment is delayed or inadequate, seizures can rapidly become self-sustained and fail to respond to the intrinsic mechanisms normally involved in seizure termination [42].

# 2. Diagnosis & treatment

## 2.1. Diagnostic approach

Approximately 15% of patients with a prolonged convulsive seizure episode (>5 min) may achieve seizure cessation without medical intervention [6]; however, the majority of patients experience a seizure lasting more than 30 min. Therefore, RSE recognition is crucial and requires prompt diagnostic evaluation and treatment in order to prevent long-term sequelae. At this point in time, SE patients may have already completed a battery of testing [43–45], yet efforts should remain focused on identifying the etiology of RSE/SRSE (Table 1) [22,43–45]. Laboratory investigations for inflammatory and immune-related etiologies (e.g. serum and CSF-autoantibodies, IgG index) should be tiered according to the disease phenotype [45,46]. With more readily available genetic testing, evaluations for genetic epilepsy

# **Table 1**Recommended diagnostic workup for pediatric RSE/SRSE.

#### Always recommended

Finger stick blood glucose

Monitor vital signs

CT/MRI (almost always appropriate except in epileptic patients with a prior normal neuroimaging or with a generalized seizure syndrome and generalized seizures) Serum electrolytes including calcium and magnesium

cEEG monitoring

#### Specific circumstances

## Known epilepsy patient

ASD levels Consider CT/MRI

Consider Electrolytes

\*Decision making largely dependent on the patient's seizure history and associated comorbidities.

#### Febrile patient

### SE with fever (presumed Febrile SE) in a patient <5 years, improved clinical state and SE resolving (no concerns for CNS infection)

Identification of primary source of fever

# SE with fever in a patient>5 years, improved clinical state and SE resolving

Identification of primary source of fever

CT/MRI consider giving IV contrast if possible

### SE with fever of unknown etiology and no improvement of clinical state

CBC

Lumbar puncture with CSF investigation of infectious etiologies

CT/MRI consider giving IV contrast if possible

#### Suspected non-infectious encephalitis (immune/inflammatory)

CRP

ESR

Auto-antibodies including ANA, anti-dsDNA, ANCA, APS & ENA panel

Serum anti-neuronal antibodies including anti-NMDAR, -AMPA & -VGKC, -GABA

Lumbar puncture with oligoclonal bands, and CSF anti-neuronal antibodies (as above)

Paraneoplastic evaluation if appropriate

# Suspected genetic syndrome

Genetic consultation

Tiered genetic testing per age, clinical exam and seizure phenotype

#### Additional considerations

Toxicology screen

Consider medication side effect (chemotherapeutics, immune-modulators, etc.)

In rheumatologic disease consider: CRP, ESR, CMP, ANA, ANCA, APS panel, ENA panel

Abbreviations: CT: computed tomography, MRI: magnetic resonance imaging, cEEG: continuous electroencephalogram, ASD: anti-seizure drug, SE: status epilepticus, CNS: central nervous system, IV: intravenous, CBC: complete blood count, CSF: cerebrospinal fluid, CRP: C-reactive protein, ESR: erythrocyte sedimentation rate, ANA: anti-nuclear antibody, ANCA: anti-neutrophil cytoplasmic antibody, APS: anti-phospholipid syndrome (lupus anticoagulant, anti- $\beta$ 2-glycoprotein,-cardiolipin), ENA: extractable nuclear antigen (anti-Smith, -RNP,-Ro,-La), NMDAR (N-methyl-D-aspartate), AMPA ( $\alpha$ -amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid), VGKC (voltage gated potassium channel), GABA ( $\gamma$ -Amino butyric acid)

syndromes should be considered in patients in whom an etiology is not established with the initial testing. These should be suspected particularly in patients that present at young ages, or with a history of dysmorphic features, developmental delay or a family history of epilepsy.

Lastly, continuous electroencephalography (cEEG) monitoring is essential for the diagnosis and management of RSE/SRSE [43,47,48]. Monitoring with cEEG allows clinicians to target electrographic seizure cessation or achievement of burst-suppression as the case requires [43,47,48]. It is helpful to guide the induction of pharmacological coma as well as the identification of subsequent electrographic seizures, often recurring during titration or weaning of anesthetic agents [49]. Furthermore, there is evidence that approximately 1/3 of the children with convulsive SE may develop electrographic seizures and of these, almost half evolve to electrographic SE [50]. There is certainly an increased awareness of the valuable aid of cEEG in the initial treatment and diagnosis of RSE/SRSE [51,52]; however, not all of the available pediatric studies include it in their RSE/SRSE management. This highly limits the generalizability of results and comparison of clinical outcomes across studies, particularly in NCSE.

# 2.2. Treatment

The therapeutic approach in RSE/SRSE aims to achieve seizure control with prevention of excitotoxicity, neuroprotection and avoidance of systemic complications [53]. Coma induction with

anesthetic agents is the most common treatment after failure of first- and second-line ASDs. The choice of anesthetic agent is often individualized. Unfortunately, due to the lack of randomized clinical trials to guide clinical practice, the goal of pharmacological induced coma (termination of seizures, burst-suppression or complete suppression of EEG activity [54]), duration, and weaning parameters remain unclear. In general, CIs should be titrated to achieve electrographic seizure cessation or burst-suppression; which should be maintained for at least 24-48h before withdrawal of CI agents [43]. If seizures recur after the weaning period, CIs are usually up-titrated until cessation is re-attained and for additional 24–48h. In addition to coma induction with anesthetic agents, the use of adjunctive therapies such as immunomodulation, ketogenic diet, hypothermia, electroconvulsive therapy and vagus nerve stimulation might be considered despite their low level of evidence. The treatment approaches for RSE/SRSE are heterogeneous and usually employed in a trial and error fashion until a response is achieved. Table 2 summarizes the most common pharmacological and non-pharmacological therapies for the treatment of RSE/SRSE [43,55,56] and their corresponding level of evidence.

# 2.2.1. Anesthetics and continuous infusions

The most commonly used agents for coma induction are midazolam [15,26,57–59], barbiturates and propofol [53,60]. Midazolam is a fast-acting BZD that enhances the action of GABA on the GABA-A receptors in the CNS [61,62]. It has a short duration

**Table 2**Pharmacological and non-pharmacological therapies for the treatment of RSE/SRSE.

	Mechanism of action	Dose	Adverse Events	Clinical Considerations	Level of evidence	References
Pharmacological therap	vies					
Benzodiazepines Midazolam	Positive allosteric modulation of GABA-A receptors, Increases	Loading dose: 0.2 mg/kg;	Hypotension, respiratory depression	Prolonged use may cause tachyphylaxis and drug	Class IIA, Level B	[43]
Wanashai ama	frequency of CI channel opening	administer at an infusion rate of 2 mg/min Infusion rate: 0.05–2 mg/kg/h Breakthrough SE: 0.1–0.2 mg/kg bolus, increase rate by 0.05–0.1 mg/kg/h. every 3–4 h		accumulation	Class IV	[9,153]
IV anesthetic agents Barbiturates Pentobarbital	Activation of GABA receptors- increase mean CI channel	Loading dose: 5- 15 mg/kg; infusion	Hypotension, cardiac and respiratory depression,	Long half-life (15–50 h) Requires mechanical	Class IIB, Level B	[43]
PEHLODAI DILAI	opening duration, inhibition of NMDA receptors, alteration in conductance of Cl <sup>-</sup> , K <sup>*</sup> , Ca <sup>2*</sup> ion channels. Same as Pentobarbital	0, 0	paralytic ileus, infection	ventilation. Can exacerbate porphyria Hepatic enzyme inducer Drug accumulation with prolonged use	Class IV	[10,65].
Thiopental	Same as the mechanism described above	2–7 mg/kg, infusion rate ≤ 50 mg/min Infusion/ maintenance rate: 0.5–5 mg/kg/h Breakthrough SE:1– 2 mg/kg bolus, titrate by 0.5–1 mg/ kg/h. every 12 h.	Hypotension, cardiac and respiratory depression	Requires mechanical ventilation, titrate infusion rates to EEG burst-suppression	Class IV	[66]
Propofol	Chloride channel conductance, enhances GABA-A receptor	Initial loading dose: 1–2 mg/kg	PRIS, hypotension,	Requires mechanical ventilation	Class IIB, Level B	[43]
	Children Grish A receptor	Initial infusion rate 20 mcg/kg/min titrated by 5– 10 mcg/kg/min Use with caution with doses > 65 mcg/kg/min Breakthrough SE: Increase infusion rate by 5–10 mcg/ kg/min every 5 min	cardiac and respiratory depression	Prolonged infusion of propofol is a relative contraindication in children (due to risk of PRIS) and in patients with metabolic acidosis, mitochondrial disorders or hypertriglyceridemia Reduces ICP Caution with concomitant use of steroid or catecholamine therapy	Class IV	[66]
Ketamine	reduces neuronal excitability	0.5–3 mg/kg Infusion rate: 1–10 mg/kg/h	Tachycardia, hypertension, ICP elevation	Relative contraindication in patients with ICP. Ketamine is an enzyme inducer and inhibitor (CYP2C9)	Class IV	[73,154]
Inhalational anesthes Isoflurane	<b>ia</b> Enhancement of GABA-A	Concentration 1–5%	Hypotension requiring	High seizure recurrence	Class IV	[83,84,87]
	receptors, noncompetitive antagonist of NMDA receptor	Titrate to achieve burst-suppression on EEG	use of vasopressors, atelectasis, paralytic ileus, infection, deep vein thrombosis	rate		
Immunomodulatory therapy						
IVIG	Alteration of IgG-specific receptors (FcγR) expression and function (decreases cytokine production), attenuation of complement mediated cell damage	1–2 g/kg divided over 3–5 days	Hypersensitivity reactions, transfusion related acute lung injury, thromboembolic events, renal dysfunction with concentrated solutions, aseptic meningitis	Immunomodulatory therapies may be considered in patients with cryptogenic, autoimmune etiologies of RSE/SRSE.	Class IV	[33,93,94,155]
Corticosteroids:    Methyl    prednisolone  Prednisone	Inhibition of inflammation- associated proteins (e.g. cytokines, chemokines) and immunosuppressive action Same as the mechanism described above	1 g/day for 3–5 days 60 mg daily	Glucose intolerance, psychiatric disturbances, altered immune function, adrenal suppression		Class IV	[91–94,156,157]

Table 2 (Continued)

	Mechanism of action	Dose	Adverse Events	Clinical Considerations	Level of evidence	References
Plasmapheresis	Removal of circulating autoantibodies, immune factors or high weight proteins that may participate in inflammatory process	5 exchanges over 5 days			Class IV	[33,94,96,140,158– 160]
Non-pharmacologica	al alternatives					
Ketogenic diet	Ketosis mediated decreased glycolysis, increase in free and polyunsaturated fatty acids, anti-inflammatory action, stabilization of neuronal membrane	4:1 (ratio fat to carbohydrates and proteins)	Hypoglycemia, hyperlipidemia, weight loss, acute pancreatitis, metabolic acidosis	Contraindicated in pyruvate carboxylase deficiency, disorders of fatty acid oxidation and metabolism, or porphyria	Class IV	[34,35,101,107–115]
Hypothermia	Reduction of Na <sup>+</sup> exchange, decreased K <sup>+</sup> conductance, regulation of glutamatergic synaptic transmission, disruption of synchronized discharges	32–35°C x 24h Rewarming $\leq$ 0.5 °C/ h	Deep venous thrombosis, infections, cardiac arrhythmias, electrolyte disturbances, acute intestinal ischemia, coagulation disorders	Requires EEG monitoring	Class IV	[123,124,126,127,161– 163]
Electroconvulsive therapy	Enhancement of GABA neurotransmission, increase of seizure threshold and reduction of neural metabolic activity	Variable protocols	May induce seizures and non-convulsive SE after treatment, amnesia, headache, cognitive impairment	Relative contraindication in patients with cardiovascular conditions Requires EEG monitoring	Class IV	[134–136,164]
Vagus nerve stimulation	Modulation of the locus coeruleus, thalamus and limbic circuit through noradrenergic and serotoninergic projections, elevation of GABA levels in brainstem	Surgical implantation	Hoarseness, surgical infection, rarely asystole or bradycardia		Class IV	[137–142]

Abbreviations: RSE: Refractory status epilepticus; SRSE, Super-refractory status epilepticus GABA-A,  $\Upsilon$ -amino-butyric acid type A; mg, milligrams; MCG, micrograms; kg, kilograms; PRIS: Propofol related infusion syndrome; NMDA, N-methyl-p-aspartate; EEG: Electroencephalogram.

of action and is generally administered as an initial loading dose of 0.2 mg/kg followed by an infusion rate of 0.05-2 mg/kg/h. It is used to achieve electrographic/clinical seizure cessation or burstsuppression [43,55]. A randomized, open label study showed that the efficacy of continuous midazolam in controlling RSE was similar to infusion of diazepam (86% vs 89% with diazepam). Midazolam however, was associated with a higher recurrence rate (57% vs. 16% with diazepam) [43,63]. Similarly, a study of 27 children with refractory generalized convulsive SE, demonstrated that midazolam infusion (0.2 mg/kg as bolus followed by 5 mcg/kg/ min as CI) was effective in the control of RSE in 26 (96%) children within 65 min, with no adverse events [28]. Even though the use of CI midazolam may cause cardiorespiratory depression and hypotension; the risk is still low compared to other anesthetics [64]. Another important aspect to consider is the risk of tachyphylaxis, which may occur after prolonged midazolam use. Thus, requiring constant monitoring and adjustment of the dosing [43].

If midazolam fails to control SE, clinicians typically resort to using barbiturates as a second agent. Pentobarbital and thiopental are both barbiturates that act similar to midazolam, through the enhancement of GABA activity. Additionally, they inhibit glutamate NMDA receptors and alter the ion conductance in the axonal membrane. Pentobarbital is administered with an initial bolus of 5-15 mg/kg (may give additional 5-10 mg/kg) followed by an infusion rate of 0.5-5 mg/kg/h. to achieve electrographic/clinical seizure cessation or burst-suppression. A case series of 26 children treated with pentobarbital showed that 75% of patients achieved burst-suppression pattern on EEG, with a relapse rate of 22% upon its weaning [10]. Another study of 30 patients presenting RSE showed that 33% achieved burst-suppression with pentobarbital without relapse; while 66.7% required titration of pentobarbital to reachieve burst-suppression. The authors also found that patients older than five years and those who achieved burst-suppression within one day of pentobarbital initiation were more likely to have

positive outcomes [65]. Thiopental is less commonly used, but can be administered with an initial bolus of 2-7 mg/kg followed by an infusion rate of 0.5-5 mg/kg/h. [43]. Its efficacy in terminating seizures showed to be lower (55%; 11/20 patients) than propofol (64%; 14/22 patients) in a retrospective study [66]. A major drawback of barbiturates is the long half-life, which leads to a delayed recovery time. In addition, barbiturates are associated with a high rate of side effects including hypotension, respiratory depression, infections, anemia and prolonged length of ICU stay [64.67]. Reported effectiveness in SE termination with barbiturates varies between 64% and 69% with an estimated seizure recurrence rate of 22% [10.64.65.67]. Because the majority of patients initiated on pentobarbital infusions have failed to stop ongoing seizure activity with midazolam, showing the potential for a more severe form of SRSE, the higher seizure recurrence rate associated with pentobarbital infusions should be interpreted cautiously.

The use of propofol is reported in the treatment of adult RSE [68]. Although the mechanism of action is similar to midazolam and barbiturates (GABA-A receptor agonist), its use in children is limited. A previous report showed that children have increased risk of propofol infusion syndrome (PRIS), a life-threatening condition characterized by metabolic acidosis, rhabdomyolysis, arrhythmias, myocardial and renal failure, that results from administering a dose higher than 4 mg/kg/h. during 48 or more hours [69]. Thus, propofol should be used with caution in pediatric patients, particularly with doses greater than 65 mcg/kg/min.

Ketamine is an alternative therapy for control of RSE [70]. It acts as a noncompetitive antagonist of NMDA receptor and has been postulated to reduce epileptiform burst discharges and prevent glutamate-mediated neuro-toxicity [71,72]. Since, in late stages of SE, there is a decrease in the number of active GABA-A receptors and up-regulated glutamate NMDA receptors, ketamine emerged as a promising treatment. A retrospective multicenter study including 46 adults and 12 children reported that permanent SE control was likely attributed to ketamine in 32% (19/60) of patients, and transient

control in 13% (8/60) when used early. When ketamine was administered as third- or fourth-line treatment, SE control was achieved in 60% (6/10) of patients [73]. Similarly, a systematic review of 162 adults and 52 children, showed that ketamine was effective in 56.5% and 36.5%, of the adults and children, respectively. From the pediatric studies included in the review, those who reported ketamine dosing indicated either an initial bolus followed by CI, isolated CI or oral administration. The initial bolus dosing ranged between 2 and 3 mg/kg, followed by an infusion rate of 7.5 mcg/kg/h. to 10 mg/kg/h. When ketamine was administered as isolated CI, the dose ranged between 7 and 60 mcg/kg/min [74]. Despite the sympathomimetic properties, ketamine has a relative safe profile [74]. Compared to other anesthetics, it has the advantage of avoiding endotracheal intubation due to lack of respiratory compromise [75]. There is a current ongoing multicenter, randomized, controlled trial [76] evaluating the efficacy of ketamine in pediatric RSE. We anticipate that the results will be essential to stratify anesthetic utilization for the treatment of RSE/SRSE. They could help to overcome current limitations of retrospective data particularly, heterogeneity of medications prior to ketamine use, dosing and timing of administration.

One of the last resorts of the SE/RSE treatment protocols is inhalational anesthesia. The antiepileptic effects of inhaled anesthetics likely involve potentiation of GABA-A receptors and inhibition of glutamate NMDA receptors [77,78]. Isoflurane and desflurane are commonly used for the treatment of RSE and have shown to be effective in inducing burst-suppression that is easily titratable [79,80]. In the children, the most common inhalational anesthetic used is isoflurane [81–86]. It is administered with endtidal concentrations of 0.5-2.3% [82.83] through an anesthetic machine. This poses a potential logistical limitation in the ICU [81,82]. A clinical series of 11 RSE episodes in 9 patients (4 adults, 5 children) demonstrated achievement of seizure cessation (EEG burst-suppression pattern) with isoflurane in all patients. Nonetheless, 8/11 (72.7%) episodes relapsed upon discontinuation of isoflurane and 6/9 (66.7%) patients died [83]. Another series of 5 adults and 2 children presenting RSE, showed that isoflurane and desflurane stopped seizures in 100% of cases, with sustained EEG burst-suppression pattern [87]. Overall, the efficacy of inhalational anesthetics seems to be transient and thus, should be considered as a temporary measure while additional workup is done to establish the etiology and/or adjunctive therapy is administered. The most commonly reported side effect of inhalational anesthetics is hypotension, requiring use of vasopressors. Other common side effects include atelectasis, deep vein thrombosis, infections and paralytic ileus [2], all of which are limiting factors for their use.

## 2.2.2. Immune modulators

In cases of RSE of a presumed autoimmune/inflammatory etiology or in cryptogenic NORSE, clinicians frequently use immunomodulatory therapies as trial in attempts to control seizures. In this scenario, their use is supported by recent discoveries on immunologic (antibodies against neural receptors such as voltage-gated potassium channels and NMDA receptors) and inflammatory (activation of inflammatory signaling pathways such as Interleukin-1 receptor/toll-like receptor pathway) processes that may contribute to their underlying pathophysiology [53,88,89]. The most commonly used therapies include corticosteroids, IV immunoglobulin (IVIG) and plasmapheresis [33,90-94]; however, their efficacy remains controversial. A series of 5 young adults with new onset SRSE, showed that early administration of immunotherapy (steroids, IVIG, and/or azathioprine) achieved seizure control in 3 patients with subsequent AEDs, and was associated with good outcomes [95]. Nonetheless, a review including 21 RSE/SRSE cases treated with adjunctive immunomodulatory therapy showed adequate seizure control in only 5% of cases [68].

Plasmapheresis is often used in parallel to other immunotherapies, particularly in RSE/SRSE etiologies such as FIRES, anti-NMDA encephalitis, and autoimmune paraneoplastic encephalitis. A previous series showed the potential benefit of early administration of plasmapheresis in FIRES. In this series, 4 patients received plasmapheresis at days 11, 12, 20 and 30, respectively, after seizure onset. The two patients that received plasmapheresis at days 11 and 12 presented better clinical outcomes than those who received it later [96]. On the contrary, a retrospective case series showed that plasmapheresis was no efficacious in two patients with FIRES [94].

There are less conventional immune therapies that target proinflammatory cytokines, which are considered to play an important role in the etiopathology of resistant epilepsies [97]. Recently, the use of Anakinra has gained attention for the treatment of FIRES [98]. Anakinra is an antagonist of the interleukin (IL) 1 receptor type 1, which inhibits the biological actions of IL-1 $\beta$  [99]. IL-1 $\beta$  is a pro-inflammatory cytokine with ictogenic properties. In animal models with refractory seizures, microglia and astrocytes exhibit overexpression of IL-1 $\beta$  [100], which makes of Anakinra a potential therapeutic approach for new onset RSE/SRSE.

Similarly, the potential involvement of toll-like receptors and IL receptors in the role of innate immunity as a precipitating factor of seizures makes inflammatory mediators appealing therapeutic targets. Drugs such as Pralnacasan (inhibitor of IL-1 $\beta$  converting enzyme), Belnacasan (selective inhibitor of the interleukin converting enzyme/caspase-1 family), VX765 (selective inhibitor of interleukin converting enzyme), Resveratrol (suppressor of nuclear factor  $\kappa B$  induced by toll like receptors) and Ifenprodil (sensitive blocker of NR2B-contaning NMDA receptors) are experimental inflammatory modulating therapies that deserve further exploration for the treatment of RSE/SRSE [97].

# 2.2.3. Other therapies

Ketogenic diet (KD) is a high-fat, low-carbohydrate, and adequate-protein diet considered a safe and effective optional therapy for patients with drug-resistant epilepsy [101]. It has gained significant attention in recent years for the treatment of SE [102–104] as an adjunctive therapy due to its anti-inflammatory and antiseizure properties. In the pediatric population, small series report a collective efficacy rate of approximately 54% [34,35,105-107]. In a series of 9 children with FIRES, KD was administered with a ratio of fat to combined protein and carbohydrate of 4:1 between days 4 and 55 of seizure onset. Seizure cessation was achieved in 7 of 8 (87.5%) patients that reached ketosis, within 2-4 days of ketonuria. Six of the 7 patients were maintained on the diet and re-experienced mild seizures (~2 seizures per week) only after few months. The remaining patient returned to RSE after termination of KD, dying 10 days later [108]. Overall, there is heterogeneity regarding timing of implementation of KD, time to ketosis and clinical outcomes [34,35,101,107–115]. Moreover, the simultaneous use of pharmacologic and non-pharmacologic therapies challenges the full understanding of the impact of KD efficacy in RSE/SRSE as well as the optimal parameters of administration. An important consideration is the involvement of a dietitian in the multidisciplinary team, as they will be crucial for achieving and maintaining ketosis in these patients. In children who are unable to take enteral feeds, implementation of KD can become complicated. Thus, IV KD can be used in these patients as a temporary measure [34,105]. Contraindications to KD implementation include carnitine deficiencies, beta-oxidation metabolic defects, pyruvate carboxylase deficiency and porphyria [116]. In the absence of a contraindication, KD could be considered in earlier stages of RSE and SRSE management.

The use of neurosteroids (e.g. allopregnanolone) in RSE/SRSE was initially supported by their action on extrasynaptic GABA-A receptors. These are different to BZDs' receptors as they do not

undergo internalization with prolonged seizures [117], and thus, represented a promising therapy for RSE/SRSE. Their use was reported in adults [118] and two pediatric patients [119], in whom the neurosteroids allowed the general anesthetic infusions to be successfully weaned. Nonetheless, a recent randomized controlled trial in adults and children failed to demonstrate the efficacy of IV allopregnanolone (brexanolone) compared to placebo (43.9% vs. 42.4%; p = 0.877) in the resolution of SRSE, when it was added to the standard care [120]. These results highlight the importance of data derived from randomized clinical trials to inform and guide current clinical RSE/SRSE management.

Therapeutic hypothermia is described mainly for the treatment of TBI. Currently, it is considered as an adjunctive therapy for RSE/SRSE due to its neuroprotective and antiepileptic properties demonstrated in animal studies. Moreover, it has the capacity to reduce cerebral metabolic rate, cerebral edema, inflammation, oxidative stress, and glutaminergic drive [121,122]. In children, multiple case reports describe resolution of RSE with adjunctive hypothermia [123-126]. In a series of 5 children with RSE, mild hypothermia (32-35 °C) was beneficial in reducing seizure burden, and prevented RSE relapse in all the patients [127]. Nonetheless, more robust evidence of its efficacy emerged with a multicenter randomized clinical trial comparing therapeutic hypothermia to the standard medical treatment. This study included 270 patients with convulsive SE and concluded that hypothermia was not associated with a lower rate of progression to RSE or SRSE, or improved clinical outcomes [128]. Important considerations before its implementation include the possible interaction with anesthetics and ASDs clearance [121], as well as awareness of common side effects: deep venous thrombosis. infections, cardiac arrhythmias, electrolyte disturbances, acute intestinal ischemia and coagulation disorders [55,129].

Electroconvulsive therapy (ECT) is also documented for the treatment of SRSE [130] in case series and reports. ECT increases GABA levels, leading to reduction of neuronal metabolic activity and interruption of seizures through the induction of a refractory period [131–133]. It is considered in children with focal and generalized RSE, with variable etiology (e.g. structural, intractable epilepsy) in whom more than five ASDs have been administered prior to ECT [134–136]. In these case reports the patients' outcomes are variable ranging from transient response to ECT and mild improvement of seizure frequency [135] to no clinical improvement followed by death [134]. Administering acute ECT is logistically challenging because its availability in some centers is limited. The side effects reported in the literature include transient lethargy or amnesic events [135], though most of the studies fail to document this information.

Another non-pharmacological option includes the vagus nerve stimulation [137–142]. Similar to ECT, there is certainly no evidence that suggest consistent improvement in seizure control based on the current data.

Emergency epilepsy surgery should be considered for RSE and SRSE treatment in two scenarios: failure to maximal medical therapy for at least two weeks or when a structural abnormality is identified [143]. Although surgery is generally contemplated in the late course of RSE, patients may undergo surgery as early as eight days after SE onset [84]. Since typical pre-operative diagnostic tools such as cortical mapping are challenging to obtain in an ICU setting, emergency surgery must be carefully weighed. The initial diagnostic approach includes the identification of the epileptogenic focus on EEG. Prior literature in children with RSE treated with surgery showed that common EEG features include generalized or non-focal lateralized discharges [144]. However, in some cases these findings can be hindered by the use of barbiturates and BZDs. If that is the case, ancillary testing such as ictal single-photon emission computerized tomography (SPECT), MRI and possibly

magneto-encephalogram (MEG) should be utilized. If a structural lesion is identified, the extent of the lesion should be further characterized through the mapping of the eloquent cortex and definition of the epileptogenic zone [143].

### 3. Outcomes of RSE and SRSE in children

#### 3.1. Clinical outcomes

The nature of RSE/SRSE is highly heterogeneous and several factors may contribute to the patients' prognosis. Mortality estimates in pediatric RSE are 13.7–43.5% [8–10,24], related to etiology, patients' age (<3 years old), and initial EEG findings (i.e. multifocal or generalized abnormalities) [8]. The underlying etiology is usually recognized as a primary predictor of outcomes [6,7,24,145,146]. Patients presenting with RSE due to acute symptomatic etiologies are less likely to return to baseline neurological function and are at higher risk of developing drugresistant epilepsy [145]. Similarly, existing evidence shows that patients with treatment delays [147], longer RSE duration [41,148], and those who present with non-convulsive SE have worse clinical outcomes [149].

Not only are patients with RSE/SRSE at risk for higher mortality, they are also at risk for neurological and systemic complications as a consequence of the prolonged length of ICU stay and anesthetic use. While the data for SRSE is scarce, a series of 10 children with SRSE showed a median ICU and hospital length stay of 27 and 62 days, respectively; and a mortality rate of 11.1% [35]. Another study identified 43 SRSE patients out of 305 children with SE. The patients in the SRSE cohort presented more often with adverse events due to medications. This cohort had a case fatality rate of 21.3% when compared to the 5.1% in all the analyzed SE children [15]. ICU related comorbidities include hypotension, prolonged respiratory failure, sepsis, pneumonia, urinary tract infections and prolonged immobility [53]. Moreover, as these patients are often on multiple therapies, systemic complications may arise from interactions between medications or their adverse effects. A study of 171 adult patients with SE evaluated the outcome of patients who were treated with IV anesthetic drugs (37%). After controlling for confounders, the authors found that the use of CIs was associated with increased frequency of infections during SE as well as a 2.9-fold relative risk for death. Additionally, these patients had higher rates of intubation, severe hypotension and poor functional outcome on the Glasgow Outcome Score at long-term follow-up [150].

Similarly, there is evidence in children with febrile RSE that induction of therapeutic burst-suppression may lead to increased risk of hemodynamic instability and poor outcomes as compared to electrographic seizure control [151]. Two retrospective case series showed that recurrence of SE after initiation of pentobarbital was associated with worse neurologic outcomes, suggesting that it should be considered as a poor prognostic factor [10,152]. The direct association between CIs use and clinical outcomes in RSE/ SRSE warrants further investigation. It remains unclear whether worse outcomes are a result of the administration of CIs and their side effects, the natural progression of a more severe case of RSE/ SRSE or a combination of these two. We consider urgent that future studies address the adequate approach to CI administration regarding dosing, up titration versus substituting for another anesthetic infusion, as well as the treatment goals in view of a potential impact on clinical outcomes.

## 4. Conclusion

RSE is a neurological emergency in the pediatric population. Literature on SRSE in children is limited despite the morbidity associated with this disorder. Current clinical practice is challenged by the heterogeneous etiologies and multiple factors involved in the progression from SE to RSE and SRSE. Moreover, there is a need to understand how aggressively RSE/SRSE patients should be treated initially, as the intrinsic risks of treatment and their effect on clinical outcomes should be taken into consideration. A multicenter and multinational collaborative effort is desirable to evaluate epidemiological data on pediatric RSE/SRSE, prevention strategies and available therapeutic options in order to provide more definitive evidence for their efficacy and safety in RSE and SRSE.

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#### References

- Falco-Walter J.J., Bleck T. Treatment of established status epilepticus. J Clin Med 2016;5:49.
- [2] Shorvon S. Super-refractory status epilepticus: an approach to therapy in this difficult clinical situation. Epilepsia 2011;52:.
- [3] Chin RF, Neville BG, Peckham C, Bedford H, Wade A, Scott RC. Incidence, cause, and short-term outcome of convulsive status epilepticus in childhood: prospective population-based study. Lancet 2006;368:222–9.
- [4] Raspall-Chaure M, Chin RF, Neville BG, Bedford H, Scott RC. The epidemiology of convulsive status epilepticus in children: a critical review. Epilepsia 2007;48:1652–63.
- [5] Lewena S, Young S. When benzodiazepines fail: how effective is second line therapy for status epilepticus in children. Emerg Med Australas 2006;18:45– 50.
- [6] Eriksson K, Metsaranta P, Huhtala H, Auvinen A, Kuusela AL, Koivikko M. Treatment delay and the risk of prolonged status epilepticus. Neurology 2005:65:1316–8.
- [7] Barzegar M, Mahdavi M, Galegolab Behbehani A, Tabrizi A. Refractory convulsive status epilepticus in children: etiology, associated risk factors and outcome. Iran | Child Neurol 2015;9:24–31.
- [8] Sahin M, Menache CC, Holmes GL, Riviello JJ. Outcome of severe refractory status epilepticus in children. Epilepsia 2001;42:1461–7.
- [9] Gilbert DL, Gartside PS, Glauser TA. Efficacy and mortality in treatment of refractory generalized convulsive status epilepticus in children: a metaanalysis. J Child Neurol 1999;14:602–9.
- [10] Kim SJ, Lee DY, Kim JS. Neurologic outcomes of pediatric epileptic patients with pentobarbital coma. Pediatr Neurol 2001;25:217–20.
- [11] Chateauneuf AL, Moyer JD, Jacq G, Cavelot S, Bedos JP, Legriel S. Super-refractory status epilepticus: epidemiology, early predictors, and outcomes. Intensive Care Med 2017;43:1532–4.
- [12] Novy J, Logroscino G, Rossetti AO. Refractory status epilepticus: a prospective observational study. Epilepsia 2010;51(2):251–6.
- [13] Ferlisi M, Shorvon S. The outcome of therapies in refractory and superrefractory convulsive status epilepticus and recommendations for therapy. Brain 2012;135(8):2314–28.
- [14] Hocker SE, Britton JW, Mandrekar JN, Wijdicks EFM, Rabinstein AA. Predictors of outcome in refractory status epilepticus. JAMA Neurol 2013;70(1):72–7.
- [15] Kravljanac R, Djuric M, Jankovic B, Pekmezovic T. Etiology, clinical course and response to the treatment of status epilepticus in children: a 16-year singlecenter experience based on 602 episodes of status epilepticus. Eur J Paediatr Neurol 2015;19:584–90.
- [16] Gross-Tsur V, Shinnar S. Convulsive status epilepticus in children. Epilepsia 1993;34(Suppl. 1):S12–20.
- [17] Treiman DM. Generalized convulsive status epilepticus. In: Jerome Engel JTAP, editor. Epilepsy: A Comprehensive Textbook. Lippincott Williams & Wilkins; 2008. p. 665–76.
- [18] Shorvon S. Tonic clonic status epilepticus. J Neurol Neurosurg Psychiatry 1993;56:125–34.
- [19] Hocker S, Tatum WO, LaRoche S, Freeman WD. Refractory and superrefractory status epilepticus—an update. Curr Neurol Neurosci Rep 2014;14:452.
- [20] Holtkamp M, Meierkord H. Nonconvulsive status epilepticus: a diagnostic and therapeutic challenge in the intensive care setting. Ther Adv Neurol Disord 2011;4:169–81.
- [21] Hussain N, Appleton R, Thorburn K. Aetiology, course and outcome of children admitted to paediatric intensive care with convulsive status epilepticus: a retrospective 5-year review. Seizure 2007;16:305–12.
- [22] Freilich ER, Schreiber JM, Zelleke T, Gaillard WD. Pediatric status epilepticus: identification and evaluation. Curr Opin Pediatr 2014;26:655–61.
- [23] Tully I, Draper ES, Lamming CR, Mattison D, Thomas C, Martland T, et al. Admissions to paediatric intensive care units (PICU) with refractory convulsive status epilepticus (RCSE): a two-year multi-centre study. Seizure 2015;29:153–61.

- [24] Lingappa L, Konanki R, Patel R, Vooturi S, Jayalakshmi S. Clinical profile and outcome of refractory convulsive status epilepticus in older children from a developing country. Seizure 2016;36:31–5.
- [25] Li Y, Tian L, Zeng T, Chen J, Chen L, Zhou D. Clinical features and outcome of super-refractory status epilepticus: a retrospective analysis in West China. Seizure 2014;23:722–7.
- [26] Saz EU, Karapinar B, Ozcetin M, Polat M, Tosun A, Serdaroglu G, et al. Convulsive status epilepticus in children: etiology, treatment protocol and outcome. Seizure 2011;20:115–8.
- [27] Gupta N, Jain P, Singh K, Bhattacharya S. Super-refractory status epilepticus with hemophagocytic syndrome in a child with HIV infection. J Trop Pediatr 2017;63:414–6.
- [28] Ozdemir D, Gulez P, Uran N, Yendur G, Kavakli T, Aydin A. Efficacy of continuous midazolam infusion and mortality in childhood refractory generalized convulsive status epilepticus. Seizure 2005;14:129–32.
- [29] Aroor S, Shravan K, Mundkur SC, Jayakrishnan C, Rao SS. Super-refractory status epilepticus: a therapeutic challenge in paediatrics. J Clin Diagn Res 2017;11:Sr01–sr04.
- [30] Reddy Y, Balakrishna Y, Mubaiwa L. Convulsive status epilepticus in a quaternary hospital paediatric intensive care unit (PICU) in South Africa: an 8 year review. Seizure 2017;51:55–60.
- [31] Hirsch LJ, Gaspard N, van Baalen A, Nabbout R, Demeret S, Loddenkemper T, et al. Proposed consensus definitions for new-onset refractory status epilepticus (NORSE), febrile infection-related epilepsy syndrome (FIRES), and related conditions. Epilepsia 2018;59:739–44.
- [32] Fox K, Wells ME, Tennison M, Vaughn B. Febrile infection-related epilepsy syndrome (FIRES): a literature review and case study. Neurodiagn J 2017;57:224– 33.
- [33] van Baalen A, Hausler M, Boor R, Rohr A, Sperner J, Kurlemann G, et al. Febrile infection-related epilepsy syndrome (FIRES): a nonencephalitic encephalopathy in childhood. Epilepsia 2010;51:1323–8.
- [34] Farias-Moeller R, Bartolini L, Pasupuleti A. Brittany cines RD, kao A, carpenter JL: a practical approach to ketogenic diet in the pediatric intensive care unit for super-refractory status epilepticus. Neurocrit Care 2017;26:267–72.
- [35] Appavu B, Vanatta L, Condie J, Kerrigan JF, Jarrar R. Ketogenic diet treatment for pediatric super-refractory status epilepticus. Seizure 2016;41:62–5.
- [36] Macdonald RL, Kapur J. Acute cellular alterations in the hippocampus after status epilepticus. Epilepsia 1999;40(Supp. 1)S9–20 discussion S21-2.
- [37] Borris DJ, Bertram EH, Kapur J. Ketamine controls prolonged status epilepticus. Epilepsy Res 2000;42:117–22.
- [38] Cock H. The role of mitochondria in status epilepticus. Epilepsia 2007;48 (Suppl. 8):24–7.
- [39] Fujikawa DG. Prolonged seizures and cellular injury: understanding the connection. Epilepsy Behav 2005;7(Suppl. 3):S3–11.
- [40] Waterhouse EJ, Garnett LK, Towne AR, Morton LD, Barnes T, Ko D, et al. Prospective population-based study of intermittent and continuous convulsive status epilepticus in Richmond, Virginia. Epilepsia 1999;40:752–8.
- [41] Sanchez Fernandez I, Abend NS, Agadi S, An S, Arya R, Brenton JN, et al. Time from convulsive status epilepticus onset to anticonvulsant administration in children. Neurology 2015;84:2304–11.
- [42] Lado FA, Moshe SL. How do seizures stop. Epilepsia 2008;49:1651-64.
- [43] Brophy GM, Bell R, Claassen J, Alldredge B, Bleck TP, Glauser T. Guidelines for the evaluation and management of status epilepticus. Neurocrit Care 2012;17:.
- [44] Glauser T, Shinnar S, Gloss D, Alldredge B, Arya R, Bainbridge J, et al. Evidence-based guideline: treatment of convulsive status epilepticus in children and adults: report of the guideline committee of the american epilepsy society. Epilepsy Curr 2016;16:48–61.
- [45] Van Mater H. Pediatric inflammatory brain diseases: a diagnostic approach. Curr Opin Rheumatol 2014:26:553–61.
- [46] Graus F, Titulaer MJ, Balu R, Benseler S, Bien CG, Cellucci T, et al. A clinical approach to diagnosis of autoimmune encephalitis. Lancet Neurol 2016; 15:391–404.
- [47] Herman ST, Abend NS, Bleck TP, Chapman KE, Drislane FW, Emerson RG, et al. Consensus statement on continuous EEG in critically ill adults and children, part I: indications. J Clin Neurophysiol 2015;32:87–95.
- [48] Herman ST, Abend NS, Bleck TP, Chapman KE, Drislane FW, Emerson RG, et al. Consensus statement on continuous EEG in critically ill adults and children, part II: personnel, technical specifications, and clinical practice. J Clin Neurophysiol 2015;32:96–108.
- [49] Abend NS, Topjian AA, Gutierrez-Colina AM, Donnelly M, Clancy RR, Dlugos DJ. Impact of continuous EEG monitoring on clinical management in critically ill children. Neurocritical care 2011;15:70–5.
- [50] Sanchez Fernandez I, Abend NS, Arndt DH, Carpenter JL, Chapman KE, Cornett KM, et al. Electrographic seizures after convulsive status epilepticus in children and young adults: a retrospective multicenter study. J Pediatr 2014;164:339–46 e1-2.
- [51] Cardoso I, Acevedo K, Hernandez M, Santin J, Moya P, Godoy J, et al. [Refractory status epilepticus in children: characterisation of epilepsies, continuous electroencephalographic monitoring and response to treatment]. Rev Neurol 2013:56:401–8.
- [52] Sanchez SM, Arndt DH, Carpenter JL, Chapman KE, Cornett KM, Dlugos DJ, et al. Electroencephalography monitoring in critically ill children: current practice and implications for future study design. Epilepsia 2013;54:1419–27.
- [53] Shorvon S, Ferlisi M. The treatment of super-refractory status epilepticus: a critical review of available therapies and a clinical treatment protocol. Brain 2011;134:.

- [54] Abend NS, Bearden D, Helbig I, McGuire J, Narula S, Panzer JA, et al. Status epilepticus and refractory status epilepticus management. Semin Pediatr Neurol 2014;21:263–74.
- [55] Bayrlee A, Ganeshalingam N, Kurczewski L, Brophy GM. Treatment of superrefractory status epilepticus. Curr Neurol Neurosci Rep 2015;15:66.
- [56] Loddenkemper T, Goodkin HP. Treatment of pediatric status epilepticus. Curr Treat Options Neurol 2011;13:560–73.
- [57] Tasker RC, Goodkin HP, Sanchez Fernandez I, Chapman KE, Abend NS, Arya R, et al. Pediatric status epilepticus research G. refractory status epilepticus in children: intention to treat with continuous infusions of midazolam and pentobarbital. Pediatr Crit Care Med 2016;17:968–75.
- [58] Patten W, Naqvi SZ, Raszynski A, Totapally BR. Complications during the management of pediatric refractory status epilepticus with benzodiazepine and pentobarbital infusions. Indian J Crit Care Med 2015;19:275–7.
- [59] Hayashi K, Osawa M, Aihara M, Izumi T, Ohtsuka Y, Haginoya K, et al. Research Committee on Clinical Evidence of Medical Treatment for Status Epilepticus in C. Efficacy of intravenous midazolam for status epilepticus in childhood. Pediatr Neurol 2007;36:366–72.
- [60] Tasker RC, Goodkin HP, Sanchez Fernandez I, Chapman KE, Abend NS, Arya R, et al. Refractory status epilepticus in children: intention to treat with continuous infusions of midazolam and pentobarbital. Pediatr Crit Care Med 2016;17:968–75.
- [61] Kaye AD, Fox CJ, Padnos IW, Ehrhardt Jr. KP, Diaz JH, Cornett EM, et al. Pharmacologic considerations of anesthetic agents in pediatric patients: a comprehensive review. Anesthesiol Clin 2017;35:e73–94.
- [62] Macdonald RL, McLean MJ. Mechanisms of anticonvulsant drug action. Electroencephalogr Clin Neurophysiol Suppl 1987;39:200–8.
- [63] Singhi S, Murthy A, Singhi P, Jayashree M. Continuous midazolam versus diazepam infusion for refractory convulsive status epilepticus. J Child Neurol 2002;17:106–10.
- [64] Wilkes R, Tasker RC. Intensive care treatment of uncontrolled status epilepticus in children: systematic literature search of midazolam and anesthetic therapies\*. Pediatr Crit Care Med 2014;15:632–9.
- [65] Barberio M, Reiter PD, Kaufman J, Knupp K, Dobyns EL. Continuous infusion pentobarbital for refractory status epilepticus in children. J Child Neurol 2012;27:721–6.
- [66] van Gestel JP, Blusse van Oud-Alblas HJ, Malingre M, Ververs FF, Braun KP, van Nieuwenhuizen O. Propofol and thiopental for refractory status epilepticus in children. Neurology 2005;65:591–2.
- [67] Bellante F, Legros B, Depondt C, Creteur J, Taccone FS, Gaspard N. Midazolam and thiopental for the treatment of refractory status epilepticus: a retrospective comparison of efficacy and safety. J Neurol 2016;263:799–806.
- [68] Ferlisi M, Shorvon S. The outcome of therapies in refractory and superrefractory convulsive status epilepticus and recommendations for therapy. Brain 2012;135:2314–28.
- [69] Fodale V, La Monaca E. Propofol infusion syndrome: an overview of a perplexing disease. Drug Saf 2008;31:293–303.
- [70] Keros S, Buraniqi E, Alex B, Antonetty A, Fialho H, Hafeez B, et al. Increasing ketamine use for refractory status epilepticus in US pediatric hospitals. J Child Neurol 2017:32:638–46.
- [71] Wasterlain CG, Mechanistic Chen JW. pharmacologic aspects of status epilepticus and its treatment with new antiepileptic drugs. Epilepsia 2008;49(Suppl. 9):63–73.
- [72] Aram JA, Martin D, Tomczyk M, Zeman S, Millar J, Pohler G, et al. Neocortical epileptogenesis in vitro: studies with N-methyl-D-aspartate, phencyclidine, sigma and dextromethorphan receptor ligands. J Pharmacol Exp Ther 1989;248:320–8.
- [73] Gaspard N, Foreman B, Judd LM, Brenton JN, Nathan BR, McCoy BM, et al. From the Critical Care EEGMRC. Intravenous ketamine for the treatment of refractory status epilepticus: a retrospective multi-center study. Epilepsia 2013;54:1498-503.
- [74] Zeiler FA, Teitelbaum J, Gillman LM, West M. NMDA antagonists for refractory seizures. Neurocrit Care 2014;20:502–13.
- [75] Ilvento L, Rosati A, Marini C, L'Erario M, Mirabile L, Guerrini R. Ketamine in refractory convulsive status epilepticus in children avoids endotracheal intubation. Epilepsy Behav 2015;49:343–6.
- [76] Rosati A, Ilvento L, L'Erario M, De Masi S, Biggeri A, Fabbro G, et al. Efficacy of ketamine in refractory convulsive status epilepticus in children: a protocol for a sequential design, multicentre, randomised, controlled, open-label, non-profit trial (KETASER01). BMJ Open 2016;6:e011565.
- [77] Kudo M, Aono M, Lee Y, Massey G, Pearlstein RD, Warner DS. Effects of volatile anesthetics on N-methyl-D-aspartate excitotoxicity in primary rat neuronalglial cultures. Anesthesiology 2001;95:756–65.
- [78] Solt K, Eger El. 2nd, Raines DE. Differential modulation of human N-methyl-D-aspartate receptors by structurally diverse general anesthetics. Anesth Analg 2006;102:1407-11.
- [79] Savard M, Dupré N, Turgeon A, Desbiens R, Langevin S, Brunet D. POLG mitochondrial disorder heralded by propofol infusion syndrome: A case report. 2013.
- [80] Tasker RC, Vitali SH. Continuous infusion, general anesthesia and other intensive care treatment for uncontrolled status epilepticus. Curr Opin Pediatr 2014;26:682–9.
- [81] Tobias JD. Therapeutic applications and uses of inhalational anesthesia in the pediatric intensive care unit. Pediatr Crit Care Med 2008;9:169–79.
- [82] Wheless JW. Treatment of refractory convulsive status epilepticus in children: other therapies. Semin Pediatr Neurol 2010;17:190–4.

- [83] Kofke WA, Young RS, Davis P, Woelfel SK, Gray L, Johnson D, et al. Isoflurane for refractory status epilepticus: a clinical series. Anesthesiology 1989;71:653–9.
- [84] Alexopoulos A, Lachhwani DK, Gupta A, Kotagal P, Harrison AM, Bingaman W, et al. Resective surgery to treat refractory status epilepticus in children with focal epileptogenesis. Neurology 2005:64:567–70.
- focal epileptogenesis. Neurology 2005;64:567–70.

  [85] Fugate JE, Burns JD, Wijdicks EF, Warner DO, Jankowski CJ, Rabinstein AA. Prolonged high-dose isoflurane for refractory status epilepticus: is it safe. Anesth Analg 2010;111:1520–4.
- [86] Lippert MM. Isoflurane anaesthesia for status epilepticus. S Afr Med J 1989;75:350-1.
- [87] Mirsattari SM, Sharpe MD, Young GB. Treatment of refractory status epilepticus with inhalational anesthetic agents isoflurane and desflurane. Arch Neurol 2004;61:1254–9.
- [88] Vezzani A, Ruegg S. The pivotal role of immunity and inflammatory processes in epilepsy is increasingly recognized: introduction. Epilepsia 2011;52(Suppl. 3):1–4.
- [89] Vezzani A, Balosso S, Aronica E, Ravizza T. Basic mechanisms of status epilepticus due to infection and inflammation. Epilepsia 2009;50(Suppl. 12):56-7.
- [90] Ferlisi M, Hocker S, Grade M, Trinka E, Shorvon S, International Steering Committee of the StEp A. Preliminary results of the global audit of treatment of refractory status epilepticus. Epilepsy Behav 2015;49:318–24.
- [91] Almaabdi KH, Alshehri RO, Althubiti AA, Alsharef ZH, Mulla SN, Alshaer DS, et al. Intravenous methylprednisolone for intractable childhood epilepsy. Pediatr Neurol 2014;50:334–6.
- [92] Erol I, Alehan F, Yalcin K. Refractory status epilepticus owing to human parvovirus B19 encephalitis in a child. J Child Neurol 2006;21:820-2.
- [93] Moreno-Medinilla EE, Negrillo-Ruano R, Calvo-Medina R, Mora-Ramirez MD, Martinez-Anton JL. Status epilepticus in paediatrics: a retrospective study and review of the literature. Rev Neurol 2015;60:394–400.
- [94] Kramer U, Chi CS, Lin KL, Specchio N, Sahin M, Olson H, et al. Febrile infectionrelated epilepsy syndrome (FIRES): pathogenesis, treatment, and outcome: a multicenter study on 77 children. Epilepsia 2011;52:1956–65.
- [95] Gall CR, Jumma O, Mohanraj R. Five cases of new onset refractory status epilepticus (NORSE) syndrome: outcomes with early immunotherapy. Seizure 2013;22:217–20.
- [96] Chevret L, Husson B, Nguefack S, Nehlig A, Bouilleret V. Prolonged refractory status epilepticus with early and persistent restricted hippocampal signal MRI abnormality. J Neurol 2008;255:112–6.
- [97] Matin N, Tabatabaie O, Falsaperla R, Lubrano R, Pavone P, Mahmood F, et al. Epilepsy and innate immune system: a possible immunogenic predisposition and related therapeutic implications. Human Vaccines & Immunotherapeutics 2015;11:2021–9.
- [98] Kenney-Jung DL, Vezzani A, Kahoud RJ, LaFrance-Corey RG, Ho ML, Muskardin TW, et al. Febrile infection-related epilepsy syndrome treated with anakinra. Ann Neurol 2016;80:939–45.
- [99] Dinarello CA, Simon A, van der Meer JW. Treating inflammation by blocking interleukin-1 in a broad spectrum of diseases. Nat Rev Drug Discov 2012;11:633–52.
- [100] Dube C, Vezzani A, Behrens M, Bartfai T, Baram TZ. Interleukin-1beta contributes to the generation of experimental febrile seizures. Ann Neurol 2005:57:152–5.
- [101] Caraballo RH, Flesler S, Armeno M, Fortini S, Agustinho A, Mestre G, et al. Ketogenic diet in pediatric patients with refractory focal status epilepticus. Epilepsy Res 2014;108:1912–6.
- [102] Freeman JM, Vining EP, Kossoff EH, Pyzik PL, Ye X, Goodman SN. A blinded, crossover study of the efficacy of the ketogenic diet. Epilepsia 2009;50:322–5.
- [103] Bodenant M, Moreau C, Sejourne C, Auvin S, Delval A, Cuisset JM, et al. Interest of the ketogenic diet in a refractory status epilepticus in adults. Rev Neurol (Paris) 2008:164:194–9.
- [104] Kossoff EH, Nabbout R. Use of dietary therapy for status epilepticus. J Child Neurol 2013;28:1049–51.
- [105] Chiusolo F, Diamanti A, Bianchi R, Fusco L, Elia M, Capriati T, et al. From intravenous to enteral ketogenic diet in PICU: a potential treatment strategy for refractory status epilepticus. Eur J Paediatr Neurol 2016;20:843–7.
- [106] Caraballo R, Noli D, Cachia P. Epilepsy of infancy with migrating focal seizures: three patients treated with the ketogenic diet. Epileptic Disord 2015;17:194–7.
- [107] Fung EL, Chang SK, Yam KK, Yau PY. Ketogenic diet as a therapeutic option in super-refractory status epilepticus. Pediatr Neonatol 2015;56:429–31.
- [108] Nabbout R, Mazzuca M, Hubert P, Peudennier S, Allaire C, Flurin V, et al. Efficacy of ketogenic diet in severe refractory status epilepticus initiating fever induced refractory epileptic encephalopathy in school age children (FIRES). Epilepsia 2010;51:2033–7.
- [109] Caraballo RH, Valenzuela GR, Armeno M, Fortini S, Mestre G, Cresta A. The ketogenic diet in two paediatric patients with refractory myoclonic status epilepticus. Epileptic Disord 2015;17:491–5.
- [110] Amer S, Shah P, Kommineni V. Refractory status epilepticus from NMDA receptor encephalitis successfully treated with an adjunctive ketogenic diet. Ann Indian Acad Neurol 2015;18:256–7.
- [111] Lin JJ, Lin KL, Chan OW, Hsia SH, Wang HS. Intravenous ketogenic diet therapy for treatment of the acute stage of super-refractory status epilepticus in a pediatric patient. Pediatr Neurol 2015;52:442–5.
- [112] Singh RK, Joshi SM, Potter DM, Leber SM, Carlson MD, Shellhaas RA. Cognitive outcomes in febrile infection-related epilepsy syndrome treated with the ketogenic diet. Pediatrics 2014;134:e1431–5.

- [113] Cobo NH, Sankar R, Murata KK, Sewak SL, Kezele MA, Matsumoto JH. The ketogenic diet as broad-spectrum treatment for super-refractory pediatric status epilepticus: challenges in implementation in the pediatric and neonatal intensive care units. J Child Neurol 2015;30:259–66.
- [114] O'Connor SE, Ream MA, Richardson C, Mikati MA, Trescher WH, Byler DL, et al. The ketogenic diet for the treatment of pediatric status epilepticus. Pediatr Neurol 2014;50:101–3.
- [115] Sort R, Born AP, Pedersen KN, Fonsmark L, Uldall P. Ketogenic diet in 3 cases of childhood refractory status epilepticus. Eur J Paediatr Neurol 2013;17:531–6.
- [116] Kossoff EH, Zupec-Kania BA, Amark PE, Ballaban-Gil KR, Christina Bergqvist AG, Blackford R, et al. Optimal clinical management of children receiving the ketogenic diet: recommendations of the International Ketogenic Diet Study Group. Epilepsia 2009;50:304–17.
- [117] Trinka E, Brigo F, Shorvon S. Recent advances in status epilepticus. Curr Opin Neurol 2016;29:189–98.
- [118] Vaitkevicius H, Husain AM, Rosenthal ES, Rosand J, Bobb W, Reddy K, et al. First-in-man allopregnanolone use in super-refractory status epilepticus. Ann Clin Transl Neurol 2017;4:411–4.
- [119] Broomall E, Natale JE, Grimason M, Goldstein J, Smith CM, Chang C, et al. Pediatric super-refractory status epilepticus treated with allopregnanolone. Ann Neurol 2014;76:911–5.
- [120] Rosenthal ESW, Mark. A Study With SAGE-547 for Super-Refractory Status Epilepticus. 2017.
- [121] Corry JJ, Dhar R, Murphy T, Diringer MN. Hypothermia for refractory status epilepticus. Neurocrit Care 2008;9:189–97.
- [122] Motamedi GK, Lesser RP, Vicini S. Therapeutic brain hypothermia, its mechanisms of action, and its prospects as a treatment for epilepsy. Epilepsia 2013:54:959–70.
- [123] Lin JJ, Lin KL, Hsia SH, Wang HS. Therapeutic hypothermia for febrile infection-related epilepsy syndrome in two patients. Pediatr Neurol 2012;47:448–50.
- [124] Shein SL, Reynolds TQ, Gedela S, Kochanek PM, Bell MJ. Therapeutic hypothermia for refractory status epilepticus in a child with malignant migrating partial seizures of infancy and SCN1A mutation: a case report. Ther Hypothermia Temp Manage 2012;2:144–9.
- [125] Elting JW, Naalt J, Fock JM. Mild hypothermia for refractory focal status epilepticus in an infant with hemimegalencephaly. Eur J Paediatr Neurol 2010;14:452–5.
- [126] Orlowski JP, Erenberg G, Lueders H, Cruse RP. Hypothermia and barbiturate coma for refractory status epilepticus. Crit Care Med 1984;12:367–72.
- [127] Guilliams K, Rosen M, Buttram S, Zempel J, Pineda J, Miller B, et al. Hypothermia for pediatric refractory status epilepticus. Epilepsia 2013;54:1586–94.
- [128] Legriel S, Lemiale V, Schenck M, Chelly J, Laurent V, Daviaud F, et al. Hypothermia for neuroprotection in convulsive status epilepticus. N Engl J Med 2016;375:2457–67.
- [129] Zeiler FA, Zeiler KJ, Teitelbaum J, Gillman LM, West M. Therapeutic hypothermia for refractory status epilepticus. Can J Neurol Sci 2015:42:221–9.
- [130] Ahmed J, Metrick M, Gilbert A, Glasson A, Singh R, Ambrous W, et al. Electroconvulsive therapy for super-refractory status epilepticus. J ECT 2017.
- [131] Sackeim HA. The anticonvulsant hypothesis of the mechanisms of action of ECT: current status. J ECT 1999;15:5–26.
- [132] Yang X, Wang X. Potential mechanisms and clinical applications of mild hypothermia and electroconvulsive therapy on refractory status epilepticus. Expert Rev Neurother 2015;15:135–44.
- [133] Duthie AC, Perrin JS, Bennett DM, Currie J, Reid IC. Anticonvulsant mechanisms of electroconvulsive therapy and relation to therapeutic efficacy. J ECT 2015;31:173–8.
- [134] Morales OG, Henry ME, Nobler MS, Wassermann EM, Lisanby SH. Electroconvulsive therapy and repetitive transcranial magnetic stimulation in children and adolescents: a review and report of two cases of epilepsia partialis continua. Child Adolesc Psychiatr Clin N Am 2005;14:193–210 viiiix.
- [135] Griesemer DA, Kellner CH, Beale MD, Smith GM. Electroconvulsive therapy for treatment of intractable seizures: initial findings in two children. Neurology 1997:49:1389–92.
- [136] Shin HW, O'Donovan CA, Boggs JG, Grefe A, Harper A, Bell WL, et al. Successful ECT treatment for medically refractory nonconvulsive status epilepticus in pediatric patient. Seizure 2011;20:433–6.
- [137] Yamazoe T, Okanishi T, Yamamoto A, Yamada T, Nishimura M, Fujimoto A, et al. New-onset refractory status epilepticus treated with vagus nerve stimulation: a case report. Seizure 2017;47:1–4.
- [138] De Herdt V, Waterschoot L, Vonck K, Dermaut B, Verhelst H, Van Coster R, et al. Vagus nerve stimulation for refractory status epilepticus. Eur J Paediatr Neurol 2009;13:286–9.

- [139] De Benedictis A, Freri E, Rizzi M, Franzini A, Ragona F, Specchio N, et al. Vagus nerve stimulation for drug-resistant Epilepsia Partialis Continua: report of four cases. Epilepsy Res 2013;107:163–71.
- [140] Howell KB, Katanyuwong K, Mackay MT, Bailey CA, Scheffer IE, Freeman JL, et al. Long-term follow-up of febrile infection-related epilepsy syndrome. Epilepsia 2012;53:101–10.
- [141] Winston KR, Levisohn P, Miller BR, Freeman J. Vagal nerve stimulation for status epilepticus. Pediatr Neurosurg 2001;34:190-2.
- [142] Zamponi N, Rychlicki F, Corpaci L, Cesaroni E, Trignani R. Vagus nerve stimulation (VNS) is effective in treating catastrophic 1 epilepsy in very young children. Neurosurg Rev 2008;31:291–7.
- [143] Bhatia S, Ahmad F, Miller I, Ragheb J, Morrison G, Jayakar P, et al. Surgical treatment of refractory status epilepticus in children: clinical article. J Neurosurg Pediatr 2013;12:360-6.
- [144] Vendrame M, Loddenkemper T. Surgical treatment of refractory status epilepticus in children: candidate selection and outcome. Semin Pediatr Neurol 2010;17:182–9.
- [145] Sahin M, Menache CC, Holmes GL, Riviello Jr. JJ. Prolonged treatment for acute symptomatic refractory status epilepticus: outcome in children. Neurology 2003;61:398–401.
- [146] Mayer SA, Claassen J, Lokin J, Mendelsohn F, Dennis LJ, Fitzsimmons BF. Refractory status epilepticus: frequency, risk factors, and impact on outcome. Arch Neurol 2002;59(2):205–2010.
- [147] Gaínza-Lein M, Sánchez Fernández I, Jackson M, et al. Association of time to treatment with short-term outcomes for pediatric patients with refractory convulsive status epilepticus. JAMA Neurol. 2018;75:410–8.
- [148] Claassen J, Lokin JK, Fitzsimmons BF, Mendelsohn FA, Mayer SA. Predictors of functional disability and mortality after status epilepticus. Neurology 2002:58:139–42.
- [149] Abend NS, Wusthoff CJ, Goldberg EM, Dlugos DJ. Electrographic seizures and status epilepticus in critically ill children and neonates with encephalopathy. Lancet Neurol 2013;12:1170–9.
- [150] Sutter R, Marsch S, Fuhr P, Kaplan PW, Rüeg S. Anesthetic drugs in status epilepticus: risk or rescue? A 6-year cohort study. Neurology 2014;82:.
- [151] Lin J-J, Chou C-C, Lan S-Y, Hsiao H-J, Wang Y, Chan O-W, et al. Therapeutic burst-suppression coma in pediatric febrile refractory status epilepticus. Brain Dev 2018;39:693–702.
- [152] Claassen J, Hirsch LJ, Emerson RG, Mayer SA. Treatment of refractory status epilepticus with pentobarbital, propofol, or midazolam: a systematic review. Epilepsia 2002;43:.
- [153] Hayashi K, Osawa M, Aihara M, Izumi T, Ohtsuka Y, Haginoya K, et al. Efficacy of intravenous midazolam for status epilepticus in childhood. Pediatr Neurol 2007;36:366–72.
- [154] Rosati A, L'Erario M, Ilvento L, Cecchi C, Pisano T, Mirabile L, et al. Efficacy and safety of ketamine in refractory status epilepticus in children. Neurology 2012;79:2355–8.
- [155] Gaspard N, Foreman BP, Alvarez V, Cabrera Kang C, Probasco JC, Jongeling AC, et al. New-onset refractory status epilepticus: etiology, clinical features, and outcome. Neurology 2015;85:1604–13.
- [156] Illingworth MA, Hanrahan D, Anderson CE, O'Kane K, Anderson J, Casey M, et al. Elevated VGKC-complex antibodies in a boy with fever-induced refractory epileptic encephalopathy in school-age children (FIRES). Dev Med Child Neurol 2011;53:1053-7.
- [157] Caputo D, Iorio R, Vigevano F, Fusco L. Febrile infection-related epilepsy syndrome (FIRES) with super-refractory status epilepticus revealing autoimmune encephalitis due to GABAAR antibodies. Eur J Paediatr Neurol 2018:22:182-5.
- [158] Caraballo RH, Reyes G, Avaria MF, Buompadre MC, Gonzalez M, Fortini S, et al. Febrile infection-related epilepsy syndrome: a study of 12 patients. Seizure 2013:22:553–9.
- [159] Finne X, Sindic C, van Pesch V, El Sankari S, de Tourtchaninoff M, Denays R, et al. Anti-N-methyl-D-aspartate receptor encephalitis with favorable outcome despite prolonged status epilepticus, Neurocrit Care 2013;18:89–92.
- [160] Lousa M, Sanchez-Alonso S, Rodriguez-Diaz R, Dalmau J. Status epilepticus with neuron-reactive serum antibodies: response to plasma exchange. Neurology 2000;54:2163–5.
- [161] Vastola EF, Homan R, Rosen A. Inhibition of focal seizures by moderate hypothermia: a clinical and experimental study. Arch Neurol 1969;20:430–9.
- [162] Cereda C, Berger MM, Rossetti AO. Bowel ischemia: a rare complication of thiopental treatment for status epilepticus. Neurocrit Care 2009;10:355–8.
- [163] Miras Veiga A, Moreno DC, Menendez AI, Siscart IM, Fernandez MD, Sanchez EC, et al. Effectiveness of electroconvulsive therapy for refractory status epilepticus in febrile infection-related epilepsy syndrome. Neuropediatrics 2017;48:45–8.
- [164] Viparelli U, Viparelli G. ECT and grand mal epilepsy. Convuls Ther 1992;8:39-