

# Pharmacotherapy for Status Epilepticus

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**Abstract** Status epilepticus (SE) represents the most severe form of epilepsy. It is one of the most common neurologic emergencies, with an incidence of up to 61 per 100,000 per year and an estimated mortality of 20 %. Clinically, tonic-clonic convulsive SE is divided into four subsequent stages: early, established, refractory, and super-refractory. Pharmacotherapy of status epilepticus, especially of its later stages, represents an “evidence-free zone,” due to a lack of high-quality, controlled trials to inform clinical decisions. This comprehensive narrative review focuses on the pharmacotherapy of SE, presented according to the four-staged approach outlined above, and providing pharmacological properties and efficacy/safety data for each antiepileptic drug according to the strength of scientific evidence from the available literature. Data sources included MEDLINE and back-tracking of references in pertinent studies. Intravenous lorazepam or

intramuscular midazolam effectively control early SE in approximately 63–73 % of patients. Despite a suboptimal safety profile, intravenous phenytoin or phenobarbital are widely used treatments for established SE; alternatives include valproate, levetiracetam, and lacosamide. Anesthetics are widely used in refractory and super-refractory SE, despite the current lack of trials in this field. Data on alternative treatments in the later stages are limited. Valproate and levetiracetam represent safe and effective alternatives to phenobarbital and phenytoin for treatment of established SE persisting despite first-line treatment with benzodiazepines. To date there are no class I data to support recommendations for most antiepileptic drugs for established, refractory, and super-refractory SE. Limiting the methodologic heterogeneity across studies is required and high-class randomized, controlled trials to inform clinicians about the best treatment in established and refractory status are needed.

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## Key Points

Initial treatment of early status epilepticus (SE) with intravenous lorazepam or intramuscular midazolam is able to control seizures in 63–73 %; buccal midazolam may be an alternative whenever intravenous or intramuscular application of other benzodiazepines is not possible.

In established SE, intravenous antiepileptic drugs (phenytoin/fosphenytoin, valproate, levetiracetam, phenobarbital) are most commonly used, but there is no class I evidence for choosing one over the other; valproate and levetiracetam represent safe and effective alternatives to phenobarbital and phenytoin; lacosamide is another potential alternative to phenytoin and phenobarbital, but current evidence is too sparse to give recommendations.

Refractory and super-refractory SE is treated with anesthetics (propofol, midazolam, thiopental/pentobarbital) with lower success rates and a high morbidity and mortality. Potential drugs to be considered in super-refractory SE are ketamine, magnesium, and immunomodulatory treatments, as well other cause-directed and non-medical treatments.

Other drugs which might be useful in the treatment of SE, such as clonazepam, paraldehyde, chlormethiazole (clomethiazole), or lidocaine, have a long history, but there is no higher-class evidence to support their use other than as second or third alternatives in refractory cases.

## 1 Introduction

Status epilepticus (SE) can be regarded as the most severe and extreme form of an epileptic seizure. Tonic-clonic SE (i.e., convulsive SE, CSE) can be defined as ongoing convulsive seizure activity or repeated convulsive seizures, without regaining consciousness between seizures, for more than 5 min [1]. Non-convulsive SE (NCSE) can be defined as an “enduring epileptic condition with reduced or altered consciousness, behavioral and vegetative abnormalities, or merely subjective symptoms like auras, but without major convulsive movements for more than 30 min” [2, 3]. A Task Force of the International League Against Epilepsy (ILAE) recently defined SE as “a condition resulting either from the

failure of the mechanisms responsible for seizure termination or from the initiation of mechanisms, which lead to abnormally prolonged seizures (after time point t1)... which can have long-term consequences (after time point t2), including neuronal death, neuronal injury, and alteration of neuronal networks, depending on the type and duration of seizures”. [4]. The time limits for t1 were set at 5 min for generalized convulsive SE, and 10 min for focal SE with impaired consciousness (formerly complex-partial SE). In the new classification NCSE is divided into those patients with and without coma following two broad clinical categories: while the former are “ictally comatose”, often seen as a progression of CSE, the “walking wounded” with aura continua, absence status, or focal SE with impaired consciousness have a less severe prognosis and do usually not need the full armamentarium of emergency treatment as described below.

SE is most prevalent in the population with structural brain damage. In patients with epilepsy, SE can be precipitated by drug withdrawal, intercurrent illness, or metabolic disturbance. The mortality of SE is around 20 %, but may be as high as 40 % in the elderly with acute symptomatic SE [5–9] and many co-morbidities [10]. The annual incidence has been estimated to be approximately 18–28 cases per 100,000 per year, but may be as high as 61 per 100,000 per year, depending on the population studied [11–16]. The incidence is highest in the elderly and has a second peak in the neonatal period [17–22].

Although the first descriptions go back to Babylonian Times (Sakikku-Board, 718–614 BC) [23] and recognition of absence status was evident in the 16th century [24], detailed descriptions of the clinical picture and first pathophysiology considerations occurred in the 19th and 20th centuries. In their seminal work, Clark and Prout recognized three phases of CSE [25–27]:

- (a) In patients with epilepsy, an early phase can be characterized, where frequency and severity of seizures increases in a crescendo pattern. Synonyms are premonitory status, impending status, and heraldic status. In patients without pre-existing epilepsy, the phase with a crescendo-like increase in seizure frequency and severity is missing, and SE starts abruptly. Ongoing convulsive epileptic activity for more than 5 min is now often called early SE.
- (b) Established SE designates continuous seizure activity with convulsions or intermittent seizures without regaining consciousness between the seizures. For more than 10 and up to 30 min, or failure of initial treatment (usually benzodiazepines) of early SE.
- (c) With increasing duration, a decrease in motor activity (electromechanical dissociation) occurs while the patient remains in a coma. This phase is called

advanced SE or refractory SE, referring to the failed treatment (usually with antiepileptic drugs, AEDs) of early and established SE. Other terms are subtle SE or stuporous SE.

- (d) At the third London-Innsbruck Colloquium on Status Epilepticus [28], the fourth stage of SE, called super-refractory SE, was characterized. At this stage seizures continue despite maximal treatment with intravenous (IV) anesthetics for more than 24 h in an intensive care unit. These patients have ictal EEG discharges when anesthesia is lessened. This stage has also been termed malignant SE [29] (Fig. 1).

It has to be acknowledged that there is no clear definition of the stages and one might merge into the other. While Clark and Prout described the stages of SE in mostly untreated patients, clinical practice now defines the first stage with a time frame (5 min of convulsive and 10 min of focal non-convulsive), and the later stages by treatment response: it has now been commonly accepted to designate established SE as “benzodiazepine-resistant SE,” while the term refractory SE is used, when treatment with benzodiazepines and one or more IV AEDs have failed. This also implies that the timeframes given above, which are used by most clinicians, may vary considerably with treatment. By nature this lack of clear definitions leads to a high degree of variability in the current literature.

In 2007, at the First London-Innsbruck Colloquium on Status Epilepticus, a workshop was held with the purpose of outlining the options of optimal pharmacotherapy of the various forms of SE. A consensus was reached and a treatment protocol published, which followed the conventional pattern of tonic-clonic SE established [30, see Flow

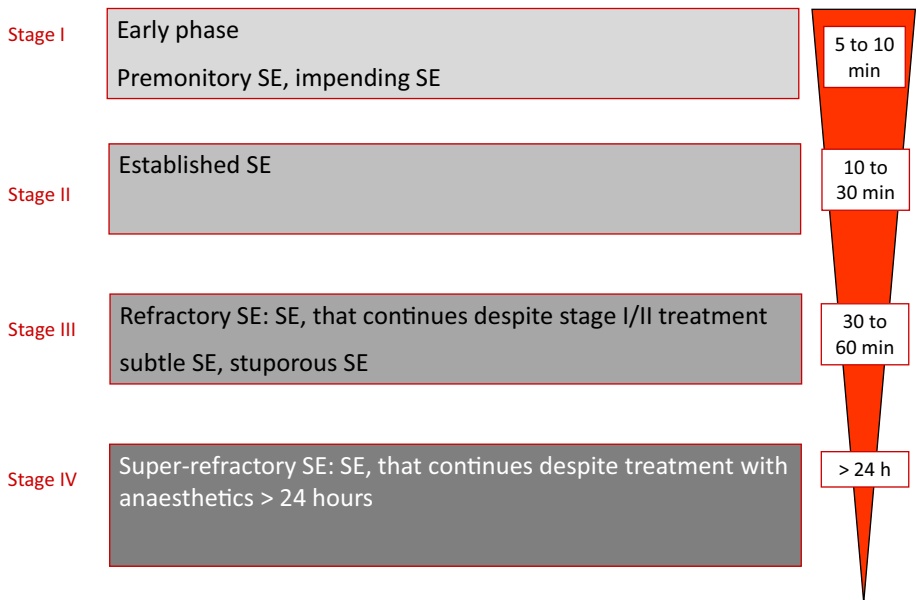
chart in “Appendix”]. The European Federation of Neurological Societies and other groups have also published similar recommendations [31, 32]. Recent reviews [33, 34] covered the history of pharmacotherapy of SE outlining the enormous range of therapies that have been advocated since the 19th century.

Data on the pharmacotherapy of SE are most often observational, having a high degree of heterogeneity and high-class randomized, controlled trials are only available for the early stages of SE. Therefore we discuss the pharmacotherapy of SE in a narrative, rather than in a systematic review. In this article we will review the data following the same principles of a staged approach as outlined above.

Treatment of SE, especially of its later stages, the pharmacological management of which represents a terra incognita [28], is almost an “evidence-free zone,” due to a lack of adequate numbers of high-quality, controlled trials to inform clinical decisions, especially in the later stages of the disorder. In most clinical trials performed in this area, often burdened by severe methodologic limitations including excessive clinical heterogeneity, investigators use different definitions of SE (and its stages), adopt inappropriate comparators, or use unclear methods of data presentation [35–37], so that reaching definite evidence is an extremely challenging task.

Given this serious limitation, in this narrative review we presented the most relevant studies on this topic (Table 1) taking into account the “evidence-pyramid” [38]: whenever available, data from controlled clinical trials (randomized/not randomized) were preferred over uncontrolled trials or case series, unless reporting relevant clinical results in terms of efficacy or tolerability. Similarly,

**Fig. 1** Clinical course of convulsive status epilepticus (SE)



**Table 1** Main randomized, controlled clinical trials conducted in different stages of status epilepticus (SE)

Study	Country	Definition of SE	Participants and age	Interventions	Clinical seizure cessation	Adverse effects
<b>Early SE: Stage I</b>						
Remy et al. [53]	France	Seizures >20 min or 2 GTCS within 20 min	Adults 16–65 years	Rectal DZP 30 mg vs. rectal DZP 20 mg	DZP 30 mg 13/18 DZP 20 mg 6/21	DZP 30 mg 10/18 DZP 20 mg 13/21
Dreifuss et al. [52]	USA	Acute repetitive seizures	Adults >18 years	Rectal DZP 0.2 mg/kg vs. placebo	DZP 31/46 Placebo 14/49	DZP 19/46 Placebo 13/49
Cereghino et al. [50]	USA	Multiple seizures within 12–24 h	Adults 18–76 years	Rectal DZP 0.2 mg/kg vs. placebo	DZP 22/31 Placebo 11/39	DZP 10/31 Placebo 9/39
Shaner et al. [117]	California	GTCS >30 min or 3 GTCS within 1 h or GTCS >5 min	Adults >15 years	IV DZP 2–20 mg + IV PHT 6–18 mg/kg based on initial drug levels vs. IV PB 10–30 mg/kg IV	DZP + PHT 10/18 PB 16/18	DZP + PHT 9/18 PB 9/18
Allredge et al. [40]	USA	Seizures >5 min	Adults >18 years	IV DZP 5 mg or LZP 2 mg vs. placebo	LZP 39/66 DZP 29/68	LZP 7/66 DZP 7/68
Chamberlain et al. [43]	USA	Seizures >5 min	Children 3 months–18 years	IV DZP 0.2 mg/kg vs. IV LZP 0.1 mg/kg	Placebo 15/71 DZP 101/140 LZP 97/133	Placebo 16/71 DZP 157** LZP 155**
Leppik et al. [41]	USA	≥3 GTCS in 1 h; confusional state with ongoing EEG abnormalities	Adults >18 years	IV LZP 2 mg vs. IV DZP 5 mg	LZP 33/37 DZP 25/32	LZP 0/37 DZP 4/32
Treiman et al. [81]	USA	Seizures >10 min or 2 GTCS within 10 min or subtle generalized convulsive SE (coma and ictal discharges on EEG)	Adults >18 years	IV LZP vs. IV PB vs. IV DZP + PHT vs. IV PHT	LZP 63/97 PB 53/91 DZP + PHT 53/95	LZP 42/97 PB 46/91 DZP + PHT 48/95
Arya et al. [48]	India	Seizures at arrival at emergency department	Children 6–14 years	IN LZP 0.1 mg/kg vs. IV LZP 0.1 mg/kg	PHT 44/101 IN LZP 59/71 IV LZP 56/70	PHT 44/101 IN LZP 0/71 IV LZP 0/70
Gathwala et al. [58]	India	Seizures at arrival at emergency department	Children 6 months–14 years	IV MDZ 0.1 mg/kg vs. IV DZP 0.3 mg/kg vs. IV LZP 0.1 mg/kg	MDZ 36/40 DZP 29/40 LZP 38/40	MDZ 4/40 DZP 24/40 LZP 4/40

Table 1 continued

Study	Country	Definition of SE	Participants and age	Interventions	Clinical seizure cessation	Adverse effects
Lahat et al. [59]	Israel	Febrile seizures >10 min	Children 6 months–5 years	IN MDZ 0.2 mg/kg vs. IV DZP 0.3 mg/kg	MDZ 23/26 DZP 24/26	MDZ 0/26 DZP 0/26
Fisgin et al. [62]	Turkey	Seizures at arrival at emergency department	Children 1 month–13 years	IN MDZ 0.2 mg/kg vs. rectal DZP 0.3 mg/kg	MDZ 20/23 DZP 13/22	MDZ 2/23 DZP 0/22
Mahmoudian and Zadeh [60]	Iran	Seizures at arrival at emergency department	Children 2 months–15 years	IN MDZ 5 mg/ml vs. IV DZP 0.2 mg/kg	MDZ 35/35 DZP 35/35	MDZ 0/35 DZP 0/35
Thakker and Shanbag [61]	India	Seizures >10 min	Children 1 month–12 years	IN MDZ 0.2 mg/kg vs. IV DZP 0.3 mg/kg	MDZ 18/27 DZP 15/23	MDZ 0/27 DZP 1/23
Scott et al. [65]	UK	Seizures at arrival of paramedics	Children/adults 5–22 years	Buccal MDZ 10 mg vs. rectal DZP 10 mg	MDZ 30/40* DZP 23/39	MDZ: 0/40 DZP 0/40
McIntyre et al. [63]	UK	Seizures at arrival at emergency department	Children 7 months–15 years	Buccal MDZ 0.5 mg/kg vs. rectal DZP 0.5 mg/kg	MDZ 71/109* DZP 45/110	MDZ 5/109 DZP 7/110
Mpimbaza et al. [66]	Uganda	Seizures at arrival at emergency department or >5 min	Children 3 months–12 years	Buccal MDZ 0.5 mg/kg vs. rectal DZP 0.5 mg/kg	MDZ 125/165 DZP 114/165	MDZ 1/165 DZP 0/165
Talukdar and Chakrabarty [64]	India	Seizures at arrival at emergency department	Children Birth to 12 years	Buccal MDZ 0.2 mg/kg vs. IV DZP 0.3 mg/kg	MDZ 51/60 DZP 56/60	MDZ 0/60 DZP 0/60
Chamberlain et al. [217]	USA	Seizures >10 min	Children Birth to 18 years	IM MDZ 0.2 mg/kg vs. IV DZP 0.3 mg/kg	MDZ 12/13 DZP 10/11	MDZ 1/13 DZP 1/11
Shah and Deshmukh [67]	India	Seizures at arrival at emergency department	Children 1 month–12 years	IM MDZ 0.2 mg/kg vs. IV DZP 0.2 mg/kg	MDZ 45/50 DZP 54/65	MDZ 0/50 DZP 7/65
Silbergleit et al. [44]	USA	Seizures >5 min	Children and adults 0–102 years	IM MDZ 5–10 mg (according to body weight) vs. IV LZP 2–4 mg (according to body weight)	MDZ 329/448 LZP 282/445	MDZ 75/448 LZP 77/445
Misra et al. [82]	India	Seizures >10 min	Children/adults 1–85 years	IV VPA 30 mg/kg bolus vs. IV PHT 18 mg/kg	VPA 23/35 PHT 14/33	VPA 4/35 PHT 6/33

Table 1 continued

Study	Country	Definition of SE	Participants and age	Interventions	Clinical seizure cessation	Adverse effects
Glad et al. [84]	Israel	Seizures >30 min	Adults >18 years	IV VPA 30 mg/kg bolus vs. IV PHT 18 mg/kg bolus	VPA 13/18	VPA 0/18
Misra et al. [116]	India	Seizures >5 min	Children/adults 1–75 years	IV LEV 20 mg/kg over 15 min vs. IV LEP 0.1 mg/kg over 2–4 min	PHT 7/9 LEV 29/38 LZP 31/41	PHT 2/9 LEV 62** LZP 94**
Established SE: stage II						
Shaner et al. [117]	California	GTCS >30 min or 3 GTCS within 1 h or GTCS >5 min	Adults >15 years	IV DZP 2–20 mg + IV PHT 6–18 mg/kg based on initial drug levels vs. IV PB 10–30 mg/kg	DZP + PHT 10/18	DZP + PHT 9/18
Treiman et al. [81]	USA	Seizures >10 min or 2 GTCS within 10 min or subtle generalized convulsive SE (coma and ictal discharges on EEG)	Adults >18 years	IV LZP vs. IV PB vs. IV DZP + PHT vs. IV PHT	PB 16/18 LZP 63/97 PB 53/91 DZP + PHT 53/95	PB 9/18 LZP 42/97 PB 46/91 DZP + PHT 48/95
Agarwal et al. [83]	India	Seizures >5 min refractory to IV DZP	Children/adults >2 years	IV VPA 20 mg/kg bolus vs. IV PHT 20 mg/kg	PHT 44/101 VPA 44/50 PHT 42/50	PHT 44/101 VPA 4/50 PHT 8/50
Chen et al. [218]	China	Seizures >5 min refractory to IV DZP	Adults 15–99 years	IV VPA 30 mg/kg bolus followed by infusion at 1–2 mg/kg vs. IV DZP 0.2 mg/kg bolus followed by infusion at 4 mg/h for 3 min and then increased every 3 min by 1 µg/min until seizure cessation or maximal duration (1 h) reached	VPA 15/30 DZP 20/36	VPA 5/30 DZP 4/36
Malamiri et al. [219]	Iran	Seizures >5 min, not controlled by DZP	Children 3–16 years	IV VPA 20 mg/kg bolus vs. IV PB 20 mg/kg bolus	VPA 27/30 PB 23/30	7/30 22/30
Refractory SE: stage III						
Singhi et al. [127]	India	Motor seizures uncontrolled after 2 doses of DZP and PHT infusion	Children 2–12 years	IV MDZ 0.2 mg/kg bolus followed by 2–10 µg/kg/min infusion vs. DZP 0.01–0.1 mg/kg/min infusion	MDZ 18/21 DZP 17/19	MDZ 8/21 DZP 9/19
Mehta et al. [220]	India	Seizures >30 min, not controlled by DZP and PHT	Children 5 months–12 years	IV VPA 30 mg/kg bolus vs. IV DZP 10 µg/Kg/min increased by 10 µg/kg/h every 5 min	VPA 16/20 DZP 17/20	VPA 0** DZP 22**
Rossetti et al. [135]	Switzerland, USA	Seizures >30 min, not controlled by benzodiazepine and PHT or VPA or PB or LEV	Adults 16–87 years	IV PRO 2 mg/kg bolus then titrated toward burst-suppression or 2 mg/kg/h vs. IV PTB 5 mg/kg bolus then titrated toward burst-suppression or 2 mg/kg/h or IV THP 2 mg/kg bolus then titrated toward burst-suppression or 4 mg/kg/h	PRO 6/14 PTB or THP 2/9	PRO15** PTB or THP 12**
Super-refractory SE: stage IV						
No randomized, controlled clinical trials available to inform clinical decisions						

Table 1 continued

Study	Country	Definition of SE	Participants and age	Interventions	Clinical seizure cessation	Adverse effects
Studies not explicitly reporting a definition of SE						
Appleton et al. [42]	UK	SE not defined	Children	LZP 0.0.5–0.1 mg/kg IV vs. DZP 0.3–0.4 mg/kg IV	LZP 26/27 DZP 29/34	LZP 0/27 DZP 0/34
McCormick et al. [221]	USA	SE not defined	Children 1 month–15 years	MDZ 0.2 mg/kg IV vs. LZP 0.1/kg IV	MDZ 14/15 LZP 8/12	MDZ 1/15 LZP 2/12
<i>DZP</i> diazepam, <i>GTCS</i> generalized tonic-clonic seizures, <i>IM</i> intramuscular, <i>IN</i> intranasal, <i>IV</i> intravenous, <i>LZP</i> lorazepam, <i>mo</i> months, <i>PB</i> phenobarbital, <i>PHT</i> phenytoin, <i>PRO</i> propofol, <i>PTB</i> pentobarbital, <i>SE</i> status epilepticus, <i>THP</i> thiopental, <i>VPA</i> valproate						

\* Expressed as number of episodes

\*\* Expressed as number of episodes; some patients experienced more than one adverse effects

cumulative data obtained from high-quality systematic literature reviews were reported as the best available evidence on this topic.

## 2 Early Status Epilepticus: Stage I

All AEDs commonly used as first-line treatment in SE are benzodiazepines. These drugs bind to the gamma-aminobutyric acid (GABA)-A receptors, increasing channel opening frequency at the receptor, with subsequent increased chloride conductance and neuronal hyperpolarization, leading to enhanced inhibitory neurotransmission and antiepileptic action [39].

### 2.1 Lorazepam (Intravenous (IV) and Intranasal (IN))

Lorazepam can be administered either intravenously or intranasally, although to date most evidence in the treatment of SE refers to its IV use. Although it has a longer initial duration of action than diazepam, lorazepam administered intravenously is usually preferred as initial treatment of early SE, because it is less lipid-soluble and consequently does not undergo the rapid redistribution into peripheral tissues seen with diazepam. This pharmacologic advantage has been clinically substantiated in randomized, controlled trials comparing IV lorazepam with placebo [40], IV diazepam [41–43], and IM midazolam [44]. In a meta-analysis, lorazepam was better than placebo for risk of non-cessation of seizures (relative risk (RR) 0.52; 95 % confidence interval (CI) 0.38–0.71), better than diazepam for reducing risk of non-cessation of seizures (RR 0.64; 95 % CI 0.45–0.90), and had a lower risk for continuation of SE requiring a different drug or general anesthesia (RR 0.63; 95 % CI 0.45–0.88) [36, 45]. There was no statistically significant difference between lorazepam and diazepam administered intravenously in terms of respiratory failure/depression, or hypotension [36, 45].

IM midazolam was non-inferior to IV lorazepam in a landmark study in early SE [44] (see Sect. 2.4 for details). Recently an intranasal (IN) administration of lorazepam has been proposed as an alternative, non-invasive delivery route for this drug, considering the favorable pharmacokinetics with rapid absorption from the IN route leading to rapid blood concentrations required for seizure termination [46, 47]. The favorable pharmacokinetics of IN lorazepam in relation to standard (IV) administration have been confirmed in one randomized, open-label non-inferiority trial conducted in 141 consecutive children aged 6–14 years who presented with convulsions to the emergency room, showing that IN lorazepam was not inferior to IV lorazepam in terms of clinical seizure remission within



10 min of drug administration [48]. It has to be noted that this study included not only children in SE, but also those who had a seizure in the emergency room, which can explain the high rate of treatment success, potentially leading to a bias towards non-inferiority.

## 2.2 Diazepam (IV, Rectal)

Diazepam is a highly lipophilic benzodiazepine, which rapidly enters into the brain but subsequently is rapidly redistributed into peripheral tissues [39]. This pharmacokinetic property is responsible for its fast anticonvulsant effect in spite of its longer elimination half-life. Diazepam can be administered either intravenously or rectally, with demonstrated significantly higher efficacy over placebo in terms of controlling acute repetitive convulsive seizures in adults and children for both methods of administration [40, 49–52]. Diazepam 30 mg intrarectal gel was found to have higher efficacy than 20 mg in seizure cessation without any statistically significant increase in adverse effects [53].

A meta-analysis of the literature indicates that, compared with placebo, after diazepam administration there is a lower risk of requirement for ventilator support and continuation of SE requiring a different drug or general anesthesia with diazepam (304 patients included overall) [36]. In a recent double-blind, randomized, controlled, superiority trial IV diazepam was compared to IV lorazepam in pediatric SE [43]. 273 children aged 3 months to <18 years were randomized to either 0.2 mg/kg diazepam ( $n = 140$ ) or 0.1 mg/kg lorazepam ( $n = 133$ ). The rates for cessation of SE within 10 min and without recurrence over 30 min were 72.1 % (101/140) in the diazepam group and 72.9 % (97/133) in the lorazepam group. There were also no differences in all secondary outcomes (e.g., requirement of assisted ventilation), except that patients in the lorazepam group were more often sedated (66.9 vs. 50 %).

## 2.3 Clonazepam (IV)

Clonazepam is more lipophilic than lorazepam, but less lipophilic than diazepam, making it therefore less prone to redistribution. Its long half-life of 17–55 hs and rapid onset of action makes it an attractive agent for emergency treatment of seizures and SE. To date, there is limited evidence to support the use of IV clonazepam in the treatment of early SE. In one uncontrolled case series (17 children) with SE treated with this drug, seizure cessation was reported in all patients after administration of doses between 0.25 and 0.75 mg, with no adverse effects reported [54]. In a subsequent uncontrolled, open-label trial (24 patients), the administration of an IV bolus injection of 1–2 mg clonazepam led to complete control of 100 % (7/7)

petit mal, 50 % (7/14) grand mal, and 66 % of partial complex cases of SE (mean time to clinical seizure cessation after administration was 1.75 min) [55]. Adverse effects consisted exclusively of transient mild to moderate drowsiness occurring in 40 % of the patients. One study comparing IV clonazepam alone to clonazepam followed by levetiracetam in generalized CSE was reported to recruit in 2011, but final results have not been published yet [56].

Quite surprisingly, despite these favorable preliminary data, no further controlled trials has been conducted to evaluate efficacy and tolerability of this drug in the treatment of SE. Despite this lack of evidence, clonazepam is extensively used in France, The Netherlands, Belgium, and other European countries.

## 2.4 Midazolam (IV, Intramuscular (IM), Intranasal, Buccal)

Midazolam is a benzodiazepine with the advantage of multiple routes of administration, due to its water solubility. At physiologic pH the ring structure of midazolam closes and it becomes highly lipophilic, crossing the blood-brain barrier rapidly [57].

Midazolam administered intravenously was found to be similar in terms of seizure recurrence to IV diazepam or IV lorazepam in a pediatric non-randomized, controlled trial, with no significant differences in mean duration to clinical seizure cessation [58].

A recent double-blind, randomized, non-inferiority trial compared the efficacy of IM midazolam with that of IV lorazepam for children and adults with CSE treated by paramedics before admission to hospital [44]. Midazolam was found to be at least as safe and effective as IV lorazepam: at the time of arrival in the emergency department, seizures were absent without rescue therapy in 73.4 % (329/448) in the IM-midazolam group and in 63.4 % (282/445) in the IV-lorazepam group. The two treatment groups were similar with regard to the need for endotracheal intubation (14.1 % of subjects with midazolam and 14.4 % with lorazepam) and recurrence of seizures (11.4 % and 10.6 %, respectively). Among subjects whose seizures ceased before arrival in the emergency department, the median times to active treatment were significantly lower in the midazolam group, although the onset of action (i.e., seizure cessation) occurred sooner after IV lorazepam administration, and adverse-event rates were similar in the two groups. Overall, these findings indicate that IM midazolam is a practical, safe, and effective alternative to IV lorazepam for the treatment of prolonged convulsive seizures in prehospital settings.

In general, IN and buccal routes of administration are more convenient than IV administration for the treatment of SE, because these formulations deliver the medication



non-invasively and more rapidly than by the IV route, and may be used also by paramedics. In three randomized, controlled trials comparing IN midazolam with IV diazepam, IN midazolam was equally effective as IV diazepam, with a lower mean time to control of seizures in the midazolam group than in the diazepam group, and no significant side effects observed in either group [59–61]. In addition, IN midazolam was found to be more effective than rectal diazepam in children with prolonged convulsive seizures, without serious complications [62].

In a prospective randomized trial, buccal midazolam was found to be more effective than rectal diazepam in children with convulsive febrile seizures [63]. No statistically significant differences in terms of efficacy were found in other studies comparing buccal midazolam with IV diazepam [64] or rectal diazepam [65, 66]. These findings support treatment protocols recommending its use as first-line treatment of acute tonic-clonic seizures in childhood including CSE where IV access is difficult or not available [31]. Time to obtain IV access may be relevant, and may explain a shorter time for controlling the convulsive episodes in patients receiving buccal midazolam compared with patients treated with IV [64] or rectal [65] diazepam. Similarly, in a study in children comparing IM midazolam and IV diazepam, mean interval to cessation of convulsions with IM midazolam was found to be significantly lower than in the diazepam group without prior IV access [67].

### 3 Established Status Epilepticus: Stage II

#### 3.1 Phenytoin/Fosphenytoin (IV)

Phenytoin has a pKa of 8.3 and is highly lipid soluble but insoluble in water. To keep phenytoin in solution it has to be prepared in a highly alkaline solvent with pH values of around 12 [68]. It has been used extensively over the past 50 years in the treatment of SE [68], but it took almost 20 years to recognize the appropriate doses of phenytoin to be effective in SE [69]. Due to its slow rate of infusion (maximum 50 mg/min) and its delayed onset of action, it should not be used in early SE [30–32, 70]. The recommended dose is 18–20 mg/kg for adults and 15 mg/kg in the elderly (>65 years). Though phenytoin is not sedative, hypotension (28–50 %) and cardiac arrhythmias (2 %) may complicate the treatment [71, 72]. Patients over the age of 50 years and with pre-existing cardiac disease are at special risk for cardiovascular complications of phenytoin [71]. Phenytoin needs polypropylene glycol in an alkaline solution to prevent precipitation of the substance, which may result in local irritation, thrombophlebitis, compartment syndrome, and ‘purple glove syndrome,’ as well as tissue necrosis with extravasation [73–76].

Despite its long-standing use in SE there are only nine studies including four randomized, controlled studies in adults and children with various forms and stages of SE [69, 77–84]. In a randomized, controlled, double-blind study for initial treatment of tonic clonic status, IV phenytoin alone was significantly less effective (defined as cessation of all clinical and electrographic seizure activity within 20 min after the start of infusion and absence of seizure relapse 60 min after treatment onset) than lorazepam (43.6 vs. 64.9 %,  $p = 0.002$ ). In three other studies, two in early SE [82, 84] and one in established SE [83], phenytoin was compared with valproate. These studies were assessed in a recent systematic review, showing that there was no detectable difference between valproate and phenytoin in clinical seizure cessation (RR 1.31, 95 % CI 0.93–1.84), seizure freedom at 24 h (RR 0.96; 95 % CI 0.88–1.06), but significantly more side effects with phenytoin (RR 0.31; 95 % CI 0.12–0.85) [37]. The overall success rate with IV phenytoin has ranged from 44 % in a randomized, controlled study to 90 % in the uncontrolled studies [85]. It is important to consider that many of the patients in these studies had ineffective pretreatment with benzodiazepines, paraldehyde, or phenobarbital, making an estimate of the effect of phenytoin difficult to assess.

Fosphenytoin is a water-soluble precursor which is rapidly transformed to phenytoin. Advantages are a faster rate of infusion, up to 150 mg/min and a better local tolerability [86, 87]. Because of the 15-min time to conversion from fosphenytoin to phenytoin a similar delayed action is expected [88]. Due to its very high costs, fosphenytoin is not on the formulary of most hospitals in Europe.

#### 3.2 Valproic Acid (IV)

Valproic acid (or the sodium salt of it, i.e., valproate) is worldwide the most often prescribed AED [89]. It has a broad spectrum of efficacy, against all seizure types, and a well-known risk and adverse-event profile [90–94]. Several studies on the pharmacokinetics of IV valproate in healthy probands and patients with epilepsies are available [95–97]. Maximum plasma concentrations were reached within minutes, and onset of action is quick. Valproic acid is highly bound to plasma protein (>90 %) and is extensively metabolized in the liver (glucuronidation and beta-oxidation). The terminal half-life is 12 h. Experimental data demonstrated a rapid and reliable onset of action in several SE models [98, 99]. Since the introduction of IV valproate, experiences with this agent in 860 patients with various forms of SE have been reported, including six randomized, controlled trials, four non-randomized, controlled trials, and 20 uncontrolled trials (eight prospective, 12 retrospective) [100]. The overall response rate to

abrogate SE was 70.9 % (601/848; 95 % CI 67.8–73.9) [100]. The most commonly used effective dose was 15–45 mg/kg as a bolus (6–10 mg/kg/min) followed by 1–3 mg/kg/h infusion [85, 100–104].

Valproic acid is non-sedating and has been used in critically ill patients with rapid infusions up to 40 mg/kg/min with good tolerability [102, 105]. Safety studies of IV valproic acid in patients with SE showed a low incidence of adverse events overall (<10 %), mainly dizziness, thrombocytopenia, and mild hypotension, which was independent of infusion rates [100]. Of note is the good cardiovascular and respiratory tolerability. The most serious concern relates to the possibility of acute encephalopathy, which is sometimes related to hepatic abnormalities or hyperammonaemia [89, 93, 106, 107].

### 3.3 Levetiracetam (IV)

Levetiracetam is an efficacious and well tolerated drug with a broad spectrum of efficacy against all seizure types and a low potential for interactions due to minimal hepatic metabolism and low plasma protein binding (<10 %) [108–110]. IV levetiracetam has been available since 2006. In animal experiments a rapid antiepileptic activity has been demonstrated, especially when used in combination with diazepam [111]. Retrospective case series and prospective safety studies on more than 1000 patients including more than 500 with various forms of SE have been reported [85, 112, 113]. The safety profile of levetiracetam is advantageous, with a very low rate of adverse effects reported (most often somnolence and sedation, and rarely agitation and thrombocytopenia) [113].

In a meta-analysis including ten studies (seven retrospective observational, two prospective observational, one prospective randomized control) reporting on 234 patients, efficacy ranged from 44 to 94 %, with a higher efficacy reported in the retrospective studies [114]. A recent meta-analysis on five different treatment options including levetiracetam in benzodiazepine-resistant SE identified 27 studies (798 patients with CSE) [115]. The relative effectiveness of levetiracetam was 68.5 % (95 % CI 56.2–78.7), compared with phenobarbital 73.6 % (95 % CI 58.3–84.8 %), phenytoin 50.2 % (95 % CI 34.2–66.1), and valproate 75.7 % (95 % CI 63.7–84.8). The authors concluded that levetiracetam, valproate, or phenobarbital, but not phenytoin, should be used as first-line drugs in benzodiazepine-resistant SE [115].

Interestingly, although levetiracetam is considered a drug for the treatment of established SE, a recent randomized, open-labeled study compared its use with iIV lorazepam in 79 patients with early (stage I) SE [116]. Both drugs were equally effective in reaching clinical seizure cessation within 10 min of administration, with

significantly higher respiratory failure requiring artificial ventilation among patients receiving lorazepam. This pilot study suggests that levetiracetam is also effective in early-stage SE, where it might represent an alternative to lorazepam.

### 3.4 Phenobarbital (IV, IM)

In a randomized, controlled trial on CSE, IV phenobarbital was at least as effective as a combination of diazepam and phenytoin [117]. In the Veterans Affairs-study phenobarbital was not inferior to lorazepam in the initial treatment of SE [81]. The central depressive effect of phenobarbital, especially following the use of benzodiazepines, limits its clinical utility, when alternatives are available. However, over the years wide experience has been gained in adults and children, as well as in the newborn. The main disadvantages are sedation, respiratory depression, and hypotension. The usual recommended adult IV loading dose of phenobarbital is 10 mg/kg (doses up to 20 mg/kg have been used and recommended) given at the rate of 100 mg/min, up to a total amount of 700 mg in 7 min. In the acute setting respiration and blood pressure have to be monitored and the patient has to be under clinical observation.

## 4 Refractory Status Epilepticus: Stage III

After failure of stage I treatment (with benzodiazepines) and failure of stage II treatment (with phenytoin, levetiracetam, or valproic acid), 31–43 % of patients enter the refractory stage [118, 119]. In refractory and super-refractory SE, IV anesthetic drugs (thiopental/pentobarbital, midazolam, or propofol) are commonly used [28]. Treatment recommendations at this stage depend on retrospective case series and uncontrolled studies [120–122]. In two systemic reviews, none of the treatments currently available was superior to another [123, 124]. Propofol may be associated with an infusion syndrome, characterized by metabolic acidosis, rhabdomyolysis, renal failure, and heart failure [125]. Treatment duration of less than 48 h and doses of no more than 5 mg/kg/h are recommended. The rate of cardiovascular and metabolic complications seemed to be lowest with midazolam and highest with barbiturates, which cause a severe immunosuppression, and often lead to infections in the compromised patient [123, 124]. The use of anesthetics in refractory and super-refractory SE was associated with more infections during SE (43 % vs. 11 %;  $p = 0.0001$ ) and a 2.9-fold relative risk for death (2.88; 95 % CI 1.45–5.73) in a 6-year prospective cohort study including 171 patients (63 of them received IV anesthetics) [126].

#### 4.1 Midazolam (Continuous IV Infusion)

One study compared continuous IV midazolam infusion versus continuous IV diazepam infusion in patients with refractory SE defined as seizures not controlled by two bolus doses of diazepam (0.3 mg/kg) and phenytoin infusion (20 mg/kg in normal saline infusion over 20 min) followed by a repeat dose of benzodiazepine [127]. Continuous midazolam and diazepam infusions were equally effective for cessation of SE, although midazolam was associated with a higher seizure recurrence rate and mortality. About half of the patients required mechanical ventilation and 40 % of patients in both groups had hypotension. In a recent study, high-dose continuous IV midazolam ( $n = 100$ ) was compared with a historical control of the same center treated with a lower dose ( $n = 29$ ) [128]. The median maximum dose was 0.4 mg/kg/h (interquartile range [IQR] 0.2, 1.0) for the high-dose group and 0.2 mg/kg/h (IQR 0.1, 0.3) for the low-dose [128] group ( $p < 0.001$ ), with a similar duration of infusion. Withdrawal seizures, occurring within 48 h of discontinuation of continuous midazolam infusion, were less frequent in the high-dose group (15 vs. 64 %; odds ratio (OR) 0.10; 95 % CI 0.03–0.27) and discharge mortality was lower in the high-dose group (40 vs. 62 %; OR 0.34; 95 % CI 0.13–0.92) compared with the low-dose group.

#### 4.2 Propofol

Propofol is an anesthetic agent, acting as an *N*-methyl-D-aspartate (NMDA) antagonist in vitro [129], with a shorter duration of action and lower tendency to accumulate in the body than barbiturates. It may cause hypotension, but reduces intracranial pressure and brain metabolic requirements [130], and may also have immunomodulatory effects [131]. Its prolonged use has been reported to cause the so-called “propofol infusion syndrome,” which includes potentially fatal myocardial failure with lactic acidosis, hypertriglyceridemia, and rhabdomyolysis, with an observed incidence of around 1 % [132–134].

Propofol has been directly compared with thiopental sodium (see Sect. 4.3) in a small (only 24 patients recruited of the 150 needed; 14 subjects received propofol, nine thiopental), single-blind, multicenter trial evaluating adult patients with refractory SE [135]. This study showed a wide CI, suggesting that the drugs may differ in efficacy up to more than twofold. No difference was found between the drugs with respect to control of seizure activity and return to baseline clinical conditions at 3 months. Infections and arterial hypotension did not differ between groups, although thiopental use was associated with significantly longer mechanical ventilation.

To date, robust evidence on the role of propofol based on large, randomized, controlled trials for this serious condition is still lacking [136].

#### 4.3 Thiopental, Pentobarbital

Thiopental sodium and pentobarbital are barbiturates acting as GABA-A agonists, with enhanced inhibitory neurotransmission and antiepileptic action. Both drugs have a prolonged duration of action, mainly due to their accumulation in the body, resulting in a long recovery time [137]. Furthermore, they may cause hypotension and cardiorespiratory depression, which may require the use of additional drugs to control pressure and breathing, as well as immunosuppression [138–140].

As reported earlier (Sect. 4.2), compared with propofol, the use of thiopental seems to be associated with longer mechanical ventilation, with no difference with respect to control of seizure activity, infections, and arterial hypotension [135].

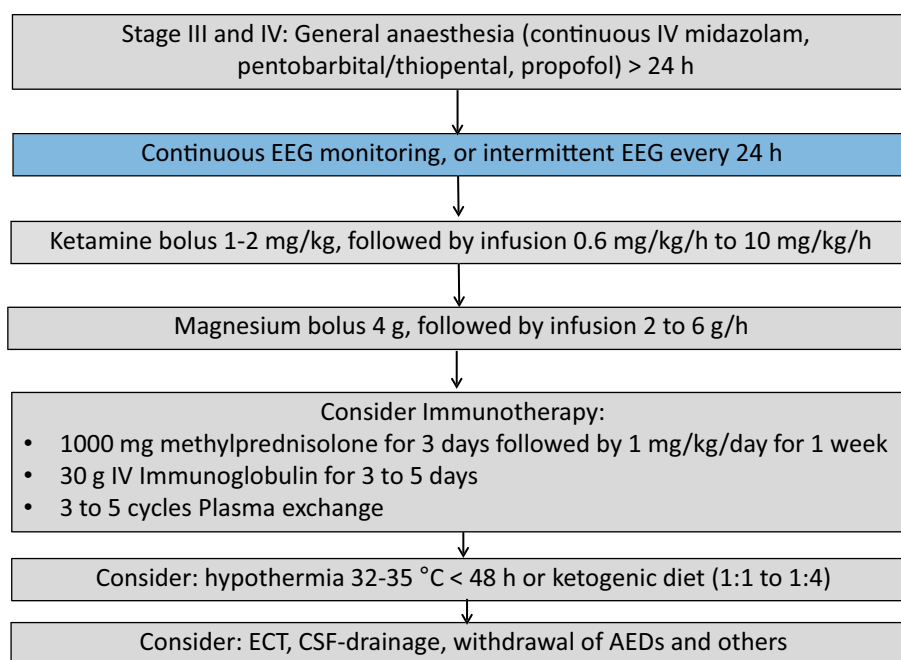
A systematic review evaluated the outcomes of anesthetics (thiopental/pentobarbital,  $n = 192$ ; midazolam,  $n = 585$ ; propofol,  $n = 143$ ) for terminating refractory and super-refractory SE. Barbiturate treatment achieved seizure control in 64 % (midazolam 78 %, propofol 68 %), but was associated with death rate during treatment in 19 % (midazolam 2 %, propofol 8 %). The authors concluded that a treatment protocol for stages III and IV of SE divided into first-line, second-line, and third-line therapy is suggested on the basis of their outcome evaluation [124]. Thus, barbiturates should be restricted to the most severe forms of refractory SE.

Despite the wide use of barbiturate anesthesia for the treatment of refractory SE, evidence derived from comparative randomized, controlled trials on the efficacy of barbiturates on which to base a choice is still lacking.

#### 4.4 Isoflurane (Inhalation)

Isoflurane is an inhalational anesthetic acting by enhancing GABA activity, with subsequent increased inhibitory neurotransmission. In a case series, this drug (used for up to 55 h) led to seizure cessation in nine patients, although with a mortality rate of 67 % [141]. A subsequent study reported its use in seven patients with refractory SE (defined as “continued seizures after failure of two or three antiepileptic drugs”) [142]: isoflurane consistently stopped epileptic discharges with adequate, sustained suppression of electroencephalographic bursts within minutes of administration. Complications were common: all patients developed severe hypotension requiring vasopressors and atelectasis; paralytic ileus occurred in three patients; and the outcome was fatal in three patients. No renal or hepatic

**Fig. 2** Example of a treatment protocol for super-refractory status epilepticus (SE). Modified after Meierkord et al. [32]. *AEDs* antiepileptic drugs, *CSF* cerebrospinal fluid, *ECT* electroconvulsive therapy, *EEG* electroencephalogram, *IV* intravenous



dysfunction occurred. Traditionally the use of inhalational anesthetics needed a vaporizer and a cart, which was not practical in the crowded environment of most intensive care units. The recently developed small self-contained vaporizers have become more widely available and will make the use of inhalational anesthetics in intensive care units more feasible.

#### 4.5 Etomidate (IV)

Etomidate is an anesthetic for which the exact mechanism of action is unknown. It has a favorable safety profile with regard to cardiovascular side effects, and may induce sedation rapidly [143]. However, its use is limited by a concern for reversible cortisol inhibition occurring after the administration of this drug, which requires hormonal substitution during treatment with etomidate [144]. In a case series (eight patients), rapid control of seizure activity was obtained in all cases, with hypotension occurring in five subjects [145].

### 5 Super-Refractory Status Epilepticus: Stage IV

When treatment with an IV anesthetic for more than 24 h is not successful in controlling SE, the condition can be termed super-refractory SE or malignant SE [28]. There have been no studies of this stage, and evidence is extremely sparse. A new initiative of a global audit (<http://www.status-epilepticus.net>) is aimed at gaining more information on the treatments used and outcomes achieved at this stage of SE. Recommendations are based exclusively on small

case series, including several treatment options, such as ketamine, corticosteroids, magnesium, IV immunoglobulin, a ketogenic diet, and neurostimulation, and in lesional cases, also surgery. It is advised to establish a protocol-driven treatment approach (Fig. 2).

#### 5.1 Ketamine (IV infusion)

Ketamine has a strong antagonistic effect on the NMDA-glutamate receptor. It has a half-life of 2–3 h and is extensively metabolized by the hepatic cytochrome P450 pathway to its active metabolite, norketamine [146]. Animal models (hippocampal electrical stimulation or pilocarpine animal models) have demonstrated the efficacy of ketamine in refractory SE in rats, even in late stages [146–148].

To date, there are two retrospective case series [149, 150] and nine single case reports, reporting 80 episodes of refractory SE in adults treated with ketamine [151–159].

One retrospective case series included 46 adults and 12 children, with 60 episodes of refractory SE. The outcome data for adults has not been separated from children [150].

In total 23 CSE, 13 NCSE, five focal SE, 38 focal NCSE, and one SE of infantile spasms were treated with ketamine. The overall success rate was 56 % after treatment with ketamine. The bolus dose ranged from 0.5 to 2 mg/kg [151, 157]. In three case reports no bolus was administered [155, 159, 160].

In the case series the loading dose varied from 1 to 2 mg/kg [149, 150], and the subsequent infusion rate ranged from 0.6 mg/kg/h to a maximal 10 mg/kg/h [150, 153]. The treatment duration of ketamine was at least 2 h [152]

and at most 27 days [150]. The time of SE until ketamine administration was started ranged from a few hours [152] up to 140 days [155]. A median of five medications (min four, max nine) prior to ketamine were used.

With regard to adverse events, one patient suffered from dysarthria, drooling, and appendicular ataxia after treatment with ketamine; upon magnetic resonance imaging a cerebellar atrophy was described. The authors concluded this to be a neurotoxic effect of ketamine. In comparison with other reports they used a higher loading—and a higher infusion dose of ketamine (2 mg/kg and a maximum dose of 7.5 mg/kg/h). After a follow-up of 21 months there was a slight clinical improvement [157].

One patient suffered from hypertension, with systolic blood pressure >220 mmHg for 10 min, after the initial bolus of 0.5 mg/kg was administered [154]. In one case series a patient had symptoms like propofol-infusion syndrome after 4 days of high-dose ketamine (4.5 mg/kg/h) and midazolam, and recovered after discontinuation. Another two patients developed supraventricular tachycardia [150]. A systematic review evaluating efficacy in pediatrics and adults showed that currently there exists Oxford level 4, Grade C evidence to support the use of ketamine for refractory SE. But they believe that there is a potential benefit with low adverse effects of NMDA antagonists—further prospective studies of early ketamine administration are needed [159].

## 5.2 Magnesium (IV)

Magnesium sulphate probably has an antiepileptic action through blocking the NMDA receptor. Magnesium sulphate has been used in SE since 1901, but has not gained much attention until now. It is currently used as the drug of choice in treating seizures occurring in eclampsia [161]. The evidence on its use in patients with SE is based on single case reports, with some evidence of benefit [162, 163]. To date, no comparative study has been conducted to assess its role in SE management. However, its infusion is safe and without significant adverse events. It has been suggested to administer magnesium as an initial IV bolus, followed by a continuous infusion at a dose that increases the serum level to ~3.5 mmol/L [163].

## 5.3 Topiramate (Enteral)

Topiramate is a broad-spectrum AED with several mechanisms of action, including blockade of the ionotropic glutamatergic AMPA receptor [164]. There is no commercially available IV formulation, but it can be administered enterally.

Evidence on the use of topiramate in super-refractory SE is based on 95 patients reported to date in the literature

[124, 165]. The dose of topiramate used in studies ranged between 2 and 25 mg/kg/day in children and up to 1600 mg/day in adults, leading to clinical seizure cessation in 62/95 (65 %) of patients. Metabolic acidosis was the most frequently reported side effect with its use.

## 5.4 Immunotherapy: Corticosteroids, IV Immunoglobulins, Plasma Exchange

The recent discovery that super-refractory SE may be caused by antibodies against neural cell receptors (voltage-gated potassium and *N*-methyl-D-aspartate (NMDA)) and evidence on the role of inflammation in epileptogenesis [166–168] led to the increasing use of immunotherapy in this stage of SE, even in the absence of any defined immunologic disease. However, to date no single comparative study has been conducted to evaluate the efficacy and tolerability of immunotherapies in super-refractory SE, and evidence is based on more than 50 patients treated so far [28]. In all reported cases, other therapies were also introduced concurrently, so that it is extremely difficult to definitively dissect the antiepileptic efficacy of immunotherapy from that of other treatments.

The rationale for the use of immunotherapies is that in some cases super-refractory SE without a clear underlying cause might be due to occult immunologic diseases with antibodies (not yet identified) directed against neural elements, which may also explain the persistence of the SE. Consequently, even in the absence of an immunologic underlying cause for the SE, a trial of high-dose steroids (1 g of IV prednisolone/day for 3 days followed by 1 mg/kg/day for ~1 week) can be given. In case of no efficacy within 2 days, either IV immunoglobulins (at a dose of 0.4 g/kg over 5 days) or plasma exchange can be tried in addition. If there is a response, treatment is continued with long-term corticosteroids, repeated courses of IV immunoglobulins and, later, other immunomodulatory agents such as cyclophosphamide or rituximab [28].

## 6 Other Drugs Used in Status Epilepticus

### 6.1 Lacosamide (IV)

Lacosamide has been available since 2008 in the European Union for the treatment of focal epilepsies [169–171]. Bioequivalence studies in healthy probands and adults with epilepsy report a good tolerability of the IV solution [172, 173]. In animal models on SE, lacosamide demonstrated a good efficacy [174]. First case reports and retrospective case series on lacosamide were published shortly after availability of the IV formulation [175–178]. There are in total 19 studies (ten single case reports and nine case



series), reporting a total of 136 episodes of refractory SE (50 % NCSE, 31 % focal SE, and 19 % CSE) treated with lacosamide [176]. All retrospective case series included patients with various forms of SE in different stages. The most commonly used bolus dose was 400 mg, followed by a daily dose of 200–400 mg lacosamide. The overall success rate was 56 % (76/136). Adverse events were reported in 25 % (34/136) of patients: mild sedation in 25 cases, one patient with possible angioedema, two with allergic skin reactions, four with hypotension, and one with pruritus. One patient developed a third-degree atrioventricular (AV) block and paroxysmal asystole [176]. There are also two single case reports (one in a patient with NCSE, one with neuropathy), reporting on AV-conduction abnormalities associated with an application of the drug [179, 180]. There was a small increase in PR interval at the end of the infusion reported in the bioequivalence studies, but this was judged to be clinically not relevant [172, 173].

## 6.2 Paraldehyde (IM, Rectal)

Paraldehyde is a drug with proven anticonvulsant properties both in animal models [181] and in humans [182]. It acts through a mechanism not yet identified, and appears to be safe with regard to cardiovascular tolerability [143]. The rectal route, which is less painful and carries no risk of sterile abscess, has largely replaced the traditional IM injection [182].

In one pediatric study rectal paraldehyde was found to be effective in terminating over 60 % of acute and prolonged convulsive seizure episodes within 10 min of its administration and without any documented adverse side effects [182]. A similar finding (clinical seizure cessation within 10 min of its administration) was found in a subsequent open-label randomized, controlled clinical trial comparing IN lorazepam and IM paraldehyde in children with prolonged (>5 min) tonic-clonic seizures; no clinically important cardiorespiratory events were reported with IM paraldehyde [183]. Of note, in this study IN lorazepam was more effective (i.e., less likely to require additional drugs to terminate seizure), significantly safer, and cheaper than IM paraldehyde in the treatment of acute tonic-clonic seizures.

## 6.3 Lidocaine (IV)

Lidocaine acts as a local anesthetic drug by inhibiting ionic currents through voltage-gated sodium channels during abnormal membrane depolarization [184]. Interestingly, IV lidocaine has been widely used in Japan for the treatment of CSE, although this drug has no official approval for the management of this condition. Most studies assessing its role in SE were performed in Japan [185, 186]. To date, there are no large, double-blind, placebo-controlled studies

evaluating lidocaine in SE, although numerous case reports and case series support its use [185–188]. Most of the available data derive from patients refractory to multiple AEDs. Furthermore, additional data supporting the use of lidocaine in SE come from pediatric uncontrolled studies, where this drug proved effective in controlling SE in neonates not responding to barbiturates [187, 188].

Overall effectiveness of lidocaine in cessation of CSE ranges between 35.8 % and 53 % [187–190]. Furthermore, unlike other AEDs used for the treatment of SE carrying the risk of respiratory depression, lidocaine has been reported to reduce the rate of mechanical ventilation [187].

## 6.4 Chlormethiazole (IV)

Chlormethiazole (clomethiazole) is a thiazole derivative acting by enhancing GABA activity, with subsequent increased inhibitory neurotransmission. No controlled studies have been conducted to evaluate its effectiveness and tolerability in SE. To date, only a few reports of patients with SE (mostly children) treated with IV chlormethiazole are available [191–194], so that the evidence supporting its use in SE is scarce. Several adverse effects related to its use have been reported including thrombophlebitis and respiratory depression [195].

## 6.5 Carbamazepine (Rectal, IV)

Carbamazepine has ILAE level A evidence for treatment of adults with focal epilepsy. This drug is characterized by poor water solubility, which represents an obstacle in the development of a parenteral formulation. IV carbamazepine has been tested in animal models [196, 197], showing rapid onset of anticonvulsant action and lack of respiratory or cardiovascular adverse effects, and very recently also in humans [198–200]. Novel IV formulations with favorable pharmacokinetics have been developed [201], but their use in clinical practice needs to be definitively assessed.

To date, there is no study reporting on patients with SE treated with IV carbamazepine. However, there is a case series reporting on seven patients presenting with cluster seizures or generalized tonic-clonic SE, who received a rectally administered oral syrup of carbamazepine with subsequent measurement of serum levels (the goal was to achieve minimum therapeutic levels of total carbamazepine >20 µmol/L) [202]. In all patients seizures had terminated prior to administration of carbamazepine, but no patient had a seizure recurrence after carbamazepine administration. This study therefore indicates the potential long-acting efficacy of rectal carbamazepine in achieving maintenance of freedom from seizures in patients with SE, but does not support its use in terminating SE.



## 7 Conclusions

In this article, we reviewed the pharmacotherapy of SE according to a staged approach based on pragmatic definitions. Morbidity and mortality correlate with duration of epileptic activity, the rapid identification of the cause of SE, age, and comorbidity of the patients [9, 10, 203–206]. Indeed, if left untreated, SE is potentially fatal or can lead to irreversible brain damage.

There is general agreement that treatment of status should follow a staged treatment protocol, so that patients may receive prompt and appropriate treatment [30]. Despite conflicting data [207], clinical seizure cessation seems to be clearly associated with adherence to treatment protocols [208–210]. Yet many hospitals do not have these or do not review and update them regularly. Randomized, controlled trials show that IV lorazepam [40, 41, 81] or IM midazolam [44, 211, 212] are the most efficient drugs in early-stage SE.

Midazolam buccal administration is a widely used alternative first-line treatment for emergency management of acute seizure disorders by non-medical personnel, prior to admission to hospital. The IN route may also be used for early SE, particularly in children, but its value compared with the buccal or IM routes is less established. Compared to lorazepam or midazolam, diazepam is more widely available throughout the world, representing a valuable alternative to lorazepam for early SE. However, compared to lorazepam, diazepam undergoes a rapid redistribution with subsequent shorter duration of action, and higher risk of accumulation after repeated administration. Rectal diazepam administration is a valuable therapeutic alternative to IV diazepam or lorazepam when this method of administration is not feasible, as it often occurs in pre-hospital settings without skilled health-care personnel. However, buccal or IN midazolam represent alternatives that are more socially accepted than rectal diazepam.

Approximately 30–40 % of all patients do not respond to benzodiazepines administered as first-line drugs, needing further treatment with IV AEDs. Even with their use, a high proportion of patients remain refractory, requiring the use of anesthetics [81, 85, 213, 214]. IV phenytoin or phenobarbital are widely used treatments for established SE. However, phenytoin and phenobarbital are not efficacious in some epilepsy syndromes and there are several concerns related to their safety profile, as both drugs may cause cardiac arrhythmias, hypotension, and respiratory depression, the latter being aggravated by co-administration with benzodiazepines.

Hence, IV formulations of other AEDs, including valproate, levetiracetam, and lacosamide have generated considerable interest as possible treatments for established SE after failure of benzodiazepines [100, 112, 113, 176–178, 215].

Classic randomized clinical trials satisfying contemporary standards are extremely challenging to perform in the setting of established SE. To date there are no class I data to support treatment recommendations for most AEDs that may be used for established, refractory, and super-refractory SE. The only partial exception concerns the evidence for use of IV valproate in the treatment of established SE, which has been shown to have a similar efficacy but a better tolerability than phenytoin in a meta-analysis [37, 100]. A recent systematic review compared the relative efficacy of five AEDs in treatment of stage II, benzodiazepine-resistant CSE [115]. Valproate was effective in controlling SE in 75.7 % (95 % CI 63.7–84.8), levetiracetam in 68.5 %, (95 % CI 56.2–78.7), phenobarbital in 73.6 % (95 % CI 58.3–84.8), and phenytoin in 50.2 % (95 % CI 34.2–66.1). Considered overall, cumulative data from the literature are therefore consistent with valproate and levetiracetam being a safe and effective therapeutic alternative to phenobarbital and phenytoin for treatment of established SE persisting despite conventional first-line treatment with benzodiazepines. A high-class randomized, controlled trial comparing valproate, levetiracetam, and phenytoin for established SE (ESET Trial) is underway, and the first results can be expected in the next few years [216]. Limiting the methodological heterogeneity across studies will be of the utmost importance in further research in the treatment of SE: investigators should adopt uniform definitions of SE and its different stages, provide individual patient data, and report their results clearly and explicitly.

### Compliance with Ethical Standards

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**Conflicts of interest** Eugen Trinka has acted as a paid consultant to Eisai, Ever Neuropharma, Biogen Idec, Medtronic, Bial, and UCB and has received speakers' honoraria from Bial, Eisai, GL Pharma, GlaxoSmithKline, Boehringer, Viropharma, Actavis, and UCB Pharma in the past 3 years. Eugen Trinka has received research funding from UCB Pharma, Biogen Idec, Red Bull, Merck, the European Union, FWF Österreichischer Fond zur Wissenschaftsförderung, and Bundesministerium für Wissenschaft und Forschung. Eugen Trinka is also part of the investigators planning the ESET-Trial and member of the Task Force on Classification of Status Epilepticus of the ILAE. Julia Höfler has received speakers' honoraria from UCB Pharma and travel support from Eisai and GL Pharma. Markus Leitinger has received travel grants from Medtronic. Francesco Brigo has received speaker's honoraria from UCB Pharma.

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## Appendix

### Early Status Epilepticus: Stage I

Intravenous (IV) route available:

- *Lorazepam* 0.07 mg/kg (usually 4 mg) IV bolus (maximum rate 2 mg/kg); if necessary can be repeated once.

OR

- *Diazepam* 5–10 mg IV bolus (maximum rate 5 mg/min); if necessary can be repeated once up to 20 mg.

OR

- *Clonazepam* 1 mg IV bolus (maximum rate 0.5 mg/min); if necessary can be repeated once after 5 min.

If IV route is difficult or not possible:

- *Midazolam* 10 mg buccal (5 mg in the elderly or in patients <50 kg); if necessary can be repeated once after 10 min. Alternatively, use 10 mg/2 mL injection via buccal route.

OR

- *Midazolam* 10 mg intramuscularly (5 mg in the elderly or in patients <50 kg); if necessary can be repeated once after 10 min.

OR

- *Diazepam* 10 mg rectal (5 mg in the elderly or in patients <50 kg); if necessary can be repeated once after 10 min.

### Established Status Epilepticus: Stage II

- *Phenobarbital* 10 mg/kg (range 10–20) IV bolus infusion at a max. rate of 100 mg/min.

OR

- *Phenytoin* 18 mg/kg (range 15–20) IV bolus infusion at max. rate of 50 mg/min.

OR

- *Fosphenytoin* 15 mg PE/kg (range 15–20) IV bolus infusion at max. rate of 100 mg PE/min.

OR

- *Valproate* 30 mg/kg (range 15–30) IV bolus infusion at 3–6 mg/kg/min.

OR

- *Levetiracetam*\* 30 mg/kg (range 30–60) IV bolus infused over 10 min.

OR

- *Lacosamide*\* 200–400 mg IV bolus infused over 3–5 min.

### Refractory Status Epilepticus: Stage III

The following drugs need to be administered by a neurointensivist/experienced anaesthetist in an intensive care unit setting:

- *Propofol* 2 mg/kg IV bolus infusion, repeated if necessary, and then followed by a continuous infusion of 5–10 mg/kg/h initially, reducing to a dose sufficient to maintain a burst-suppression pattern on the EEG (usually 1–3 mg/kg/h).

OR

- *Thiopental* 100–250 mg IV bolus infusion given over 20 s with further 50-mg boluses every 2–3 min until seizure control, followed by a continuous IV infusion at a dose sufficient to maintain a burst-suppression pattern on the EEG (usually 3–5 mg/kg/h).

OR

- *Pentobarbital* 5–15 mg/kg IV bolus, followed by a continuous IV infusion at a dose sufficient to maintain a burst-suppression pattern on the EEG (usually 0.5–3 mg/kg/h).

OR

- *Midazolam* 0.1–0.3 mg/kg IV bolus infusion at max. rate of 4 mg/min initially, followed by a continuous IV infusion at a dose sufficient to maintain a burst-suppression pattern on the EEG (usually 0.05–0.4 mg/kg/h).

After seizure control has been achieved for at least 12 h, the drug dosage should be slowly reduced over a further 12 h. If seizures recur, the anesthetic agent should be administered again for another 12 h, and then withdrawal attempted again. This cycle may need to be repeated every 24 h until achievement of seizure control.

Continuous EEG monitoring is indicated to assess level of anesthesia (burst-suppression pattern) and abolition of ictal discharges after drug withdrawal

### Super-Refractory Status Epilepticus: Stage IV

No randomized, controlled trials are available in the literature to inform about the use of any drug in the treatment of super-refractory status epilepticus.

Maintain the use of anesthetic drugs used in stage III.

Consider the use of:

- *Ketamine*\* 1–3 mg/kg IV bolus, followed by a continuous IV infusion at a dose sufficient to maintain a burst-suppression pattern on the EEG (usually up to 5 mg/kg/h).

As second-line therapy [124] consider:

- *Hypothermia*\* levels of hypothermia uncertain, usually target temperatures between 32 and 35 °C continued in the first instance for 24–48 h;
- *Magnesium infusion*\* dose of 2–6 g/h to obtain a serum level of 3.5 mmol/L;
- *Pyridoxine infusion*\* (in young children): 180–300 mg;
- *Immunologic therapy*\* high-dose steroids (1 g/day in adults) over 3 days and continued at lower doses (1 mg/kg/day) over 1 week; in addition, course of IV immunoglobulin (0.4 g/kg/day) over 5 days or plasma exchange;
- Ketogenic diet\*;
- Emergency neurosurgery\* (including focal resection, multiple subpial transection, corpus callosotomy, and hemispherectomy, even in combination).

As third-line therapy [124] consider:

- Electroconvulsive therapy\*;
- Cerebrospinal fluid drainage\*.

\*No randomized, controlled trials available in the literature to inform about the use of this drug/procedure in this stage of status epilepticus.

Ferlisi and Shorvon [124].

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