**Status Epilepticus Management: A Short Review**

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**About the Author**

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**Summary**

Status epilepticus is a common neurological emergency, with high morbidity and mortality, now defined as 5 minutes or more of (1) continuous clinical and/or electrographic seizure activity or (2) recurrent seizure activity without recovery (returning to baseline) between seizures. In observational studies, many disparities exist in management among different medical teams. Here, the author reviews evidence-based medicine data on this management, with emphasis on the use of drugs and electroencephalography. Benzodiazepine (either IV lorazepam or IM midazolam) are the recognized first line of treatment. When a patient is still seizing thereafter, the second line is usually either IV phenytoin or IV valproate, where available. A persisting status epilepticus should then be considered as refractory and managed with anaesthetic drug, keeping in mind that the natural history of that state is to evolve to non-convulsive status epilepticus, where an emergent EEG is the only way to diagnose that condition with certainty.

**Résumé**

Le status epilepticus est une urgence neurologique fréquente, associée à une haute morbidité et mortalité, maintenant définie comme 5 minutes ou plus (1) d’activité ictale continue Clinique et/ou électrographique ou (2) d’activité ictale récurrente sans résolution (retour au niveau de base) entre les crises. Dans des études observationnelles, de nombreuses disparités existent dans sa prise en charge entre les différentes équipes médicales. Nous allons ici réviser les données de la médecine basée sur l’évidence à propos de cette prise en charge, avec une emphase particulière sur les médications et l’utilisation de l’EEG. Les benzodiazépines (soit le lorazépam IV ou le midazolam IM) constituent la première ligne de traitement reconnue. Quand un patient demeure toujours en crise par la suite, la phénytoine IV ou l’acide valproïque IV (lorsque disponible) sont la seconde ligne de traitement. Un status epilepticus persistant devra par la suite être considéré comme étant réfractaire et traité à l’aide de médication anesthésiante, tout en gardant en tête que l’histoire naturelle de cette condition est d’évoluer en status épilepticus non convulsif, où un EEG d’urgence est la seule façon d’en permettre le diagnostic avec certitude.
Status epilepticus (SE) is one of the more common neurological emergencies, with a 9–21% mortality associated at hospital discharge.1 Among important factors influencing mortality, SE duration is of major importance: mortality goes from 2.7% to 32% if SE has, respectively, a length of less than 1 hour or more than 1 hour.2 Moreover, the natural history of SE is similar to that for cardiac arrest since a patient with SE will eventually evolve to an “electromechanical dissociation” state, called nonconvulsive SE. At that point, diagnostic delay has been proven to be the major factor influencing mortality;3 and only an electroencephalogram (EEG) can be used to accurately diagnose that state. This article therefore concentrates on the major aspects concerning the timely management of SE, its definition, and its appropriate pharmacological management.

The definition of SE used to involve a patient seizing consecutively for 30 minutes, since irreversible lesions appeared in the brain in a monkey model at that point.4 New guidelines no longer support that definition.1 Since all class 1 studies available considered shorter definitions (from 5 to 10 minutes5–7) and new data supported the idea that even a few minutes of evolution may have an important impact on SE resolution,7 the most recent Neurocritical Care Society (NCS) guidelines1 state that SE can be “defined as 5 minutes or more of (i) continuous clinical and/or electrographic seizure activity or (ii) recurrent seizure activity without recovery (returning to baseline) between seizures.” This definition also implies that most of the usual tonic-clonic seizures, which have a length of approximately 1 minute and almost always less than 2 minutes,8 do not usually necessitate an acute drug treatment. Eventually, 30% of the SE will evolve to a refractory state (often nonconvulsive), defined as a non-response to a benzodiazepine and a second acceptable drug in appropriate dosing, with an even higher mortality of 23–61%.1

Three principal class 1 studies (prospective, multicentric, double-blind, and randomized) influence the way we manage SE pharmacologically. The two first are paramedic studies, and the last one is an emergency room (ER) study.

The first to consider here is the San Francisco Emergency Medical Service study.4 Paramedics there were treating adults with 5 minutes or more of consecutive clinical seizure activity with intravenous (IV) diazepam 5 mg, IV lorazepam 2 mg, or placebo. Placebo was considered ethical in this case since San Francisco paramedics had not been treating SE with IV drugs previously and there was some concern that IV benzodiazepine in that setting may cause respiratory distress and morbidity. The primary outcome was the presence or not of a persisting SE at arrival in the emergency department. A total of 205 patients were enrolled in that study. A significant statistical difference favoured the use of lorazepam compared with placebo (40.9% versus 78.9%, \( p = .001 \)), whereas a non-statistically significant difference was found when compared with diazepam (57.4%). Interestingly, the group having more respiratory complications was the placebo group (22.5%, versus 10.6% for lorazepam and 10.3% for diazepam; \( p = .08 \)), supporting the notion that seizing is bad for airway protection.

The second is the most recent one, the Rampart study,7 which had the objective to prove that intramuscular (IM) midazolam (5–10 mg) is not inferior to IV lorazepam (2–4 mg). This objective was motivated by the point that an IV access may be tricky to find in a seizing patient, and moreover because the IV lorazepam solution has to be refrigerated. The study finally demonstrated a superiority of IM midazolam, with 73.4% of patients being seizure free when entering the emergency room compared with 63.4% of those treated with IV lorazepam (\( p < .001 \)). Both drugs were considered safe. It has been suggested that the difference between the two groups was secondary to the fact that it took less time to use the IM treatment (1.2 minutes) than the IV treatment (4.8 minutes).

The third study is known as the Veteran study.5 Four different blinded treatments were evaluated in the emergency room for patients seizing for at least 10 minutes. The only statistical difference was between IV lorazepam (64.9% efficacy) and IV phenytoin (43.6% efficacy), with intermediary results for phenobarbital and a phenytoin-plus-diazepam combination. The lorazepam dosage used in this study was 0.1 mg/kg. A very interesting point is that the investigators also blindly evaluated the second- and third-line efficacy of these drugs, with a collective result of 7.3% and 2.0%, respectively. These data are the basis of the refractory SE definition discussed above.

In parallel with treatment, a diagnosis workup has to be made. The NCS guidelines1 suggest the following: (1) vital sign monitoring; (2) finger-stick glucose; (3) brain computed tomography (CT) scan (for most cases); (4) basic blood tests (including calcium, magnesium, and anticonvulsant levels); and (5) continuous EEG monitoring. Based on the clinical presentation, a lumbar puncture, magnetic resonance imaging (MRI), and a toxicology screen can be considered. EEG monitoring remains the most difficult recommendation to follow, but it is motivated by the possibility of a nonconvulsive seizure evolution, which is present in 14% of controlled convulsive SE.9 The “15 minute rule” can be used here; this means that even if a patient is still comatose, there must be a clinical improvement within 15 minutes of the last seizure. If not, if available, an emergent EEG must be considered to rule out nonconvulsive SE; if this is unavailable, an intubation associated with an anaesthetic agent (preferably propofol) should be instituted and then stopped after 12–24 hours of treatment. If the patient does not improve
in the 15–30 minutes after the end of the propofol perfusion, he or she must be transferred to a hospital where EEG (preferably continuous EEG) is available. If needed, supportive treatment must also include ventilation assistance, pressors, and fluid resuscitation (especially keeping in mind the acute tubular necrosis risk secondary to rhabdomyolysis). Hyperthermia should be treated.

When benzodiazepines are not working, IV phenytoin or valproic acid must be considered; but, as discussed above, most patients will remain refractory at this point. Of interest, two prospective comparative studies\(^{10,11}\) have since compared IV phenytoin and valproate, favouring valproate in each case. However, valproate is still underused and therefore has low general clinical experience. As a consequence, IV phenytoin and valproate are usually considered equal as second line choices.

IV anaesthetics are to be considered thereafter, particularly midazolam (bolus 0.2 mg/kg, perfusion 0.1–2.0 mg/kg/h) or propofol (bolus 2–5 mg/kg, perfusion 1.0–4.5 mg/kg/h; avoid in children and for perfusions lasting >48 hours because of the risk of deadly propofol infusion syndrome\(^12\)). If the patient is still seizing during perfusion with midazolam or propofol, additional options are barbiturate, isoflurane, or ketamine. Finally, anaesthetic treatment has to be stopped every 24–48 hours. Other treatments may be tried in parallel at this point; there is some anecdotal evidence for the use of topiramate, levetiracetam, magnesium, or lacosamide, or even surgery and electroconvulsive therapy.\(^13\) Since many good outcomes have been reported, even when a patient has been in SE for weeks, if the prognosis is otherwise not bad (e.g., the SE is not caused by a malignant tumour), the key point is to be very patient.

In conclusion, SE is a neurological emergency in which the decisions taken during the first few hours have a great impact on the outcome, and where evidence-based medicine can guide our initial decisions. A suggested treatment algorithm is provided in Figure 1. Refractory SE is a less well-known entity, but remains a treatable state. However, with no evidence-based guidelines available at this time, its management is more art than science.

**Figure 1.** Suggested algorithm in treating status epilepticus (see text for details). EEG = electroencephalogram; NCSE = nonconvulsive status epilepticus; r/o = rule out.

**References**