Merrf Syndrome and Refractory Status Epilepticus

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Abstract: MERRF syndrome (or Myoclonic Epilepsy with Ragged Red Fibers) is a mitochondrial disease maternally-inherited. In this progressive disorder, status epilepticus is common and highly resistant to treatment including generalized anesthesia. We report two cases with Merrf syndrome (sister and brother), presented with status epilepticus refractory even to Thiopental, Propofol and Midazolam. In the same family is and another girl, diagnosed with Merrf syndrome. Given that it wasn’t possible to stop the refractory status in these cases, we suggest Ketamine as an alternative to general anesthesia as for blocking seizure activity, as for the potential neuroprotective effect.

Keywords: Merrf syndrome, status epilepticus, ketamine.

1. Introduction

MERRF syndrome (or Myoclonic Epilepsy with Ragged Red Fibers) is a mitochondrial disease, caused by a maternally-inherited mutation at position 8344 in the mitochondrial genome. This point mutation disrupts the mitochondrial gene for tRNA-Lys and so disrupts synthesis of proteins essential for oxidative phosphorylation.(1) The characteristic symptom of MERRF syndrome are myoclonic seizures that are usually sudden, brief, jerking, spasms that can affect the limbs or the entire body.(2) Status epilepticus can be severe and refractory even to Midazolam, Thiopental and propofol. In such cases we have failure of GABA-ergic anesthetics, so failure of inhibition mechanisms. On the other hand glutamate activation of NMDA receptors give excitatory effects and lead to neurologic injury with cell death. With the development of GABA_A resistance, SE excitatory effects and lead to neurologic injury with cell death. Mortality of SE is about 20% but varies widely, primarily on the basis of age, etiology, and duration of SE.(3) In a recent series, 11 out of 26 patients with refractory even him to midazolam, phenytoine, thiopental and propofol. Asking carefully the children's relatives, we learn that an uncle of the children has died young, and the genetic tree of the child's mother had other cases of new age deaths, without a clear cause. By the help of our geneticist doctor we made the diagnosis of Merrf syndrome, but we couldn’t help even him for the refractory status in which he presented. (He died after some days)

In the same family is and another girl, which has been too diagnosed for Merrf syndrome (mutation A8344G - analyzing the DNA of lymphocytes), and presents epilepsy.

2. Case Reports

We report three cases of Merrf syndrome, all from a family. First child, ten years old, had the first presentation in hospital with a refractory myoclonic status epilepticus. According to the family, it was the first time that the child posed such a state. The status was refractory to Diazepam (two times with 0.3 mg/kg IV), Phenytoine (20 mg/kg IV) and Phenobarbital ( loading dose 20mg/kg IV). No changes were observed at the CT of the head, performed within a few hours of presentation. Naturally the girl has been intubated and maintained in mechanical ventilation. Despite continuous treatment with Thiopental (5 mg/kg loading dose, 5 mg/kg/h maintenance dose), Midazolam (0.3 mg/kg loading dose, 0.3 μg/kg/h maintenance dose) and Propofol (2 mg/kg loading dose, 5 mg/kg/h maintenance dose), the status didn’t stop for any moment). The girl died three days after admission. At that moment we haven’t done a checkup for the other siblings, to could diagnose and maybe decide about the treatment with Levetiracetam or another antiepileptic which is of choice in such cases. In this progressive disorder, SE is common and highly resistant to treatment including generalized anesthesia(1), refractory even to Thiopental and propofol such in our case reports. In a recent series, 11 out of 26 patients with mitochondrial disease died, and in over 80% malignant SE was the recorded cause of death. (4)

Refractory status epilepticus (RSE) is an important and serious clinical problem that typically requires prolonged and high-level intensive care, and is often associated with poor outcome such in our cases. (2,5,6,7) The overall mortality of SE is about 20% but varies widely, primarily on the basis of age, etiology, and duration of SE.(3) In a systemic review of Claassen at al. (8) for refractory status from 193 patients, forty-eight percent of patients died. Even with current best practice, mortality in patients who experience refractory generalized convulsive SE is about
50% and only the minority return to their premorbid functional baseline.(9)

Several intravenous agents have been used for RSE; however, problems occur with their toxicity and/or effectiveness.(7,10,11,12) The most common treatment protocols for status epilepticus specify intravenous benzodiazepine (either lorazepam or diazepam) as initial antiepileptic drug (AED) therapy, followed by phentoin or fosphenytoin and then phenobarbital if seizures continue (according these protocol we treat and our children). (2,7) Most patients respond to the first or second AED. However, when the patient fails to respond to this standard, initial treatment, the patient is considered to be “refractory” and requires additional, more aggressive treatment.

Neuronal injury is probably mediated by excess excitation via activation of the N-methyl-D-aspartate (NMDA) subtype of glutamate receptors and consequent elevated intracellular calcium that causes acute necrosis and delayed apoptotic cell death.(13)

During RSE glutamate activation of NMDA receptors give excitatory effects and lead also to neurologic injury with cell death. With the development of GABA\(_A\) resistance (failure of inhibition mechanisms in cases refractory to GABA-ergic anesthetics), SE increase the sensitivity to NMDA antagonists. Therefore, new treatment options are urgently needed. The ideal new drug for refractory RSE would be one that has the ability to stop seizures more effectively and safely than current drugs, and that has neuroprotective properties to prevent the brain damage and neurologic morbidity caused by RSE.(9)

A few cases suggest strong anticonvulsant properties of ketamine even after failure of GABA-ergic anesthetics.(2,10,11,12,14,15,16) Ketamine inhibits NMDA (N-methyl-D-aspartate) receptors by binding to the phencyclidine site on the NMDA-receptor protein, blocking seizure activity, and having a potential neuroprotective effect. In addition it can increase blood pressure due to its sympathomimetic properties.(6,10) These characteristics make it a reasonable potential choice as adjunctive therapy in RSE.

In an abstract, Bleck and colleagues(11,17) described their experience with the use of ketamine in seven critically ill patients with RSE. Their main observation was that ketamine produced electrographic seizure control in over half the patients without causing hemodynamic instability. A case report by Sheth and Gidal(15) also provides evidence for the potential utility of ketamine in extremely refractory cases.

The general (dissociative) anesthetic dose is 1-5 mg/kg, with infusion of 1-5 mg/kg/hour (20-80 mcg/kg/minute).(10) It is recommended that it should be administered with a benzodiazepine in an attempt to decrease later psychiatric side effects.(16) Naturally, not only to give hope to this mother, but also for other similar cases with Merrf syndrome, given the RSE, even to phentoin, thiopental and propofol, after failure of GABAergic anesthetics, we suggest to incorporate ketamine into the therapeutic protocol for the strong anticonvulsant properties due to increased NMDA receptor expression with ongoing seizure activity.

4. Conclusion

Future treatment of refractory SE should involve both GABA-ergic anesthetics and NMDA antagonists to stop seizure activity and prevent acute and delayed neuronal injury.

For patients experiencing prolonged status epilepticus, refractory even to Thiopental and propofol, ketamine may be an alternative to general anesthesia.

References