Interictal Ionized Magnesium/Total Serum Magnesium Ratio in Serbian Population with Drug Resistant Epilepsy—Whether is Severe Epilepsy in Fact Brain Injury?

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Abstract

Introduction
We hypothesized that our patients with drug resistant epilepsy have low interictal ionized magnesium/total serum magnesium ratio to indicate on the protracted brain injury.

Objective
The aim of this study was to examine interictal ionized magnesium/total serum magnesium ratio in patients with drug resistant epilepsy and to consider illness and AEDs related predictors of the possible brain injury.

Methods
Patients with drug resistant epilepsy of unknown cause were tested for interictal total serum magnesium concentrations and serum ionized magnesium concentrations at the endpoint visit 14 years later. Ionized magnesium/total serum magnesium ratio cut off point was 0.60. Groups were monitored in relation to the: seizures types, seizures frequency, duration of epilepsy, appearance of status epilepticus (SE), and longest used first line antiepileptic drugs (AEDs).

Results
According to our results, 60.6% (N=63) of the patients with drug resistant epilepsy of unknown cause had lower interictal ionized magnesium/total serum magnesium ratio (mean ratio 0.53 ± 0.05). Odds Ratio was 29.19 (95% CI 10.94 to 77.90%). In addition we have found significant differences among groups in: age (p=0.001), the length of suffering from epilepsy (p=0.017), the seizure frequency (p=0.000), the experiencing of the SE (p=0.000), and the longest used first line AEDs (p=0.012) with 54% of the phenobarbital in the study group.

Conclusion
Patients with drug resistant epilepsy of unknown cause have low interictal ionized magnesium/total serum magnesium ratio which can indicate on the protracted brain injury. More evidence is necessary to proclaim ionized magnesium as marker of the protracted brain injury induced by drug resistant epilepsy. Predictors for the low interictal ionized magnesium/total serum magnesium ratio in this group of patients are: status epilepticus, older age in relation with longer suffering from epilepsy, more frequent seizures, and longest total intake of the phenobarbital.

Keywords
Drug resistant epilepsy, Ionized magnesium, Brain injury, Status epilepticus, Seizure frequency, Phenobarbital, Interictal
Research
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Introduction

Proportion of the intractable seizures among people with epilepsy (PWE) is about 15.6%, and cumulative risk for developing inadequate seizure control is about 30%. Epilepsy with intractable seizures or drug resistant epilepsy, according to International League against Epilepsy (ILAE) is defined as “failure of adequate trials of two tolerated, appropriately chosen and used antiepileptics drugs schedules (whether as monotherapies or in combination) to achieve sustained seizure freedom” [1].

Although the seizures are more frequently observed in patients with electrolyte disorders (hyponatremia, hypocalcemia, and hypomagnesemia), those kinds of seizures are classified as acute symptomatic seizures and they are unrelated to epilepsy and intractable seizures [2].

Magnesium can potentially modulate seizure activity because of its role as antagonist of the excitatory calcium influx through N-methyl-D-aspartate (NMDA) receptors [3]. Accordingly, lower serum magnesium concentrations were found in PWE than in controls [4,5]. Reducing in serum magnesium concentration was more pronounced in PWE who had status epilepticus and severe epilepsy than in those with mild epilepsy [6]. But, hypomagnesemia can be induced by a variety of reasons; from starvation, alcohol intake, diarrhea and vomiting, renal tubular defects, to intake of antimicrobials or even antiepileptic drugs (AEDs) [7-9]. Therefore, numerous studies has proved that serum ionized magnesium monitoring is more precise clinical marker for brain condition [10]. Decrease of serum or plasma ionized magnesium levels associated with brain injury were found in experimental and clinical studies on child traumatic brain injury (TBI), severe close TBI in adults, open TBI, stroke, blast trauma, combined neurotrauma models, and epilepsy [11-17]. Serum ionized magnesium was significantly lower in PWE than in healthy controls, and levels were lower postictal (within 24hour after seizure) than interictal [18,19].

We hypothesized that our patients with drug resistant epilepsy (DRE) have low interictal ionized magnesium/ total serum magnesium ratio to indicate on the epilepsy induced protracted brain injury.

Objective

The aim of this study was to examine Interictal ionized magnesium/ total serum magnesium ratio in patients with drug resistant epilepsy and to consider illness and AEDs related factors that may contributed to the possible epilepsy induced brain injury.

Methods

The study was designed as a clinical cohort study which included patients who met criteria for DRE of unknown cause (epilepsies of unknown cause are epilepsies that in the past were termed “cryptogenic” including those that were considered “undetermined”, and even partially those that were considered idiopathic, but without precise genetic defect in which seizures are the core symptom of the disorder) [20]. The cohorts were defined in regards to presence or absence of decreased serum Interictal ionized magnesium/total serum magnesium ratio on the endpoint visit, 14 years after baseline visit.

This study is a part of a cohort study on the new onset interictal psychiatric disorders in patients with drug resistant epilepsy of unknown cause, and influence of the seizure frequency, illness duration, low interictal ionized magnesium/ total serum magnesium ratio, and experiencing status epilepticus on their occurrence.

We have prescreened for the study patients from authors’ clinical database of PWE, and enroll them in the study during the year 2001. Study was completed in December 2015.

Patients were prescreened for the study according to inclusion criteria of the ILAE standards for drug resistant epilepsy of unknown cause (current terminology) i.e. idiopathic, cryptogenic and undetermined DRE, with exclusion of genetic epileptic syndromes (terminology used at the study start time) [21]. Exclusion criteria at screening visit and at endpoint visit was hypomagnesemia of any etiology, and additional exclusion criteria at screening visit was low Interictal ionized magnesium/ total serum magnesium ratio.

We have enrolled 124 female and male outpatients between the age of 18 and 50 years (less than 65 years at the time of endpoint visit), treated in Neurology Department, Institute for Neuropsychiatric Disorders “Dr Laza Lazarevic”(current name Clinic for Psychiatric Disorders „Dr Laza Lazarevic”). The study was completed after 14 years by 104 patients (Figure 1).

All patients gave their informed consent prior to screening. The study was approved by the ethical committee of the Clinic for Psychiatric Disorders.
Prescreened from data base of PWE (n=871)
Inclusion criteria: Drug resistant epilepsy - ILAE standards
Idiopathic, cryptogenic or undetermined epilepsy

Excluded: genetic epileptic syndromes

Screened (n=147)

Excluded (n=23):
- Low serum magnesium (n=2)
- Low ionized magnesium (n=18)
- IC withdraw (N=2)
- Other (n=1)

Enrolled (n=124)

Study period 14 years - Follow up (monthly visits)

Discontinued (n=19):
- Referral to another institution (n=9)
- Lack of compliance (n=7)
- Moving to another country (n=3)

No more drug resistant, still in the study (n=23)

Excluded (n=1):
- Low serum magnesium (n=1)

End point visit (n=104)

Excluded: genetic epileptic syndromes

Analysis (n=104)
Exclusion from the analysis (n=0)

Figure 1: Patients' flow chart.
**Statistical Analysis**

All collected data were organized and evaluated using dedicated software (IBM SPSS 24.0, USA) and were analyzed by descriptive statistical parameters and regression models. Descriptive statistical methods were represented by measures of central tendency (mean and median), variability (standard deviation and variation interval) and were expressed in number of patients and percentages. We used Medcalc for calculating odds ratio and diagnostic test evaluation calculator. For testing data of different categories (gender, age, illness related factors and therapy), Pearson’s $\chi^2$ and ANOVA tests were applied. We used linear regression models to examine illness and AEDs related predictors of the potential brain injury in patients with DRE of unknown cause. Level of statistical significance was set at $p<0.05$.

**Results**

According to our results, 60.6% (N=63) of the patients with drug resistant epilepsy of unknown cause had lower interictal ionized magnesium/total serum magnesium ratio, while 39.4% (N=41) had normal ratio. Odds ratio was 29.19 (95% CI 10.94 to 77.90), and Relative risk for having low interictal ionized magnesium ratio in DRE was 3.07 (95% CI 2.36 to 4.00). Mean interictal Ionized magnesium/total serum magnesium ratio in the study group was $0.53 \pm 0.05$, while in the control group was $0.70 \pm 0.06$ (Table 1).

Distribution of demographic characteristics in the observed groups shows that there were no gender differences between the groups ($p=0.452$), while we found statistical significant age differences among groups ($p=0.001$), with odds ratio for having low ionized magnesium ratio after the age of 50 years of 2.93 (95%CI 1.29 to 6.65) (Table 2).

The significant differences between groups were present compared to length of suffering from epilepsy ($p=0.017$) with odds ratio for longer illness duration (more than 35 years) of 7.06 (95%CI 2.73 to 18.2), and seizure frequency ($p=0.000$), with odds ratio for more frequent seizures of 20.46 (95% CI 7.033 to 59.53%). There was no significant difference between groups relative to seizure types ($p=0.530$), with odds ratio for generalized seizures of 0.95 (95% CI 0.43 to 2.09) (Table 3).

We found high statistical significance for the difference between two groups in the occurrence of seizures, and significant differences in illness duration, seizure frequency and seizure types.
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of the status epilepticus (p=0.000) with 95.2% of the patients who had the SE in study group, and relative risk for having low interictal ionized magnesium/total serum magnesium ratio was 14.67 (Table 4). Linear diagram of the interictal ionized magnesium/total serum magnesium ratio in regards to status epilepticus is in Figure 2.

Significant differences between the groups were also found in relation to the longest used first line AEDs (p=0.012), and study group AEDs profile was: 54% phenobarbital, 22.3% carbamazepine, 14.3% valproate, and 9.5% lamotrigine use. There was no AEDs pattern in control group where AEDs were presented in similar percentage (29.3% phenobarbital, 26.8% carbamazepine, 24.4% valproate, 19.5% lamotrigine).

Linear regression analysis marked older age (p=0.01), longer suffering from epilepsy (p=0.011), more frequent seizures (p=0.000), experiencing of the status epilepticus (p=0.000), and longest use of phenobarbital (p=0.037), as predictors of the lower ionized magnesium/total serum magnesium ratio and potential brain injury.

**Discussion**

Numerous experimental and clinical studies have showed that lower ionized magnesium levels can be considered as a significant predictor of the brain injury and even of long-term neurobehavioral outcome [24]. As magnesium is a potential modulator of seizure activity because of its ability to antagonize the excitatory calcium influx through the NMDA receptor, it was expected to find lower ionized magnesium and higher ionized calcium levels during the seizures [18], and even to link low ionized magnesium levels to sudden unexpected death in epilepsy [25], or to consider magnesium supplementation in the overall management of the people with refractory epilepsy [26]. This is the first paper considering interictal ionized magnesium/total serum magnesium ratio in DRE.

Based on a present knowledge, we hypothesized that patients with drug resistant epilepsy have low interictal ionized magnesium/total serum magnesium ratio to indicate on the epilepsy induced protracted brain injury. We found significantly lower interictal ionized magnesium/total serum magnesium ratio in 60, 6% of patients with DRE of unknown cause, thereby supporting the evidence on protracted brain injury in refractory seizures [18,19,26].

Additional evidence for this claim was found in this long term study, in which we have excluded patients with low interictal ionized magnesium/total serum magnesium ratio at screening visit, or patients with hypomagnesemia of any etiology.
at screening and endpoint visit. In this way, we have excluded even a least likely possibility that our patients had chronic magnesium deprivation such as in TRPM6 gene mutation, which lowers seizure threshold [27].

By examining the illness and AEDs related factors which can lead to a decrease in Interictal ionized magnesium/total serum magnesium ratio, thus contributing to the possible brain injury, we have not found significant gender differences among study and control group. Previous studies in healthy adults have shown that ionized magnesium and total magnesium concentrations in men were not different from those in young women or in menopausal women [28], and similar pattern is present in our patients with DRE, but cannot be used to compare with our results.

We have not found enough results from literature to compare low ionized magnesium and types of seizures. We already know that decreased ionized magnesium levels provoke generalized seizures, so it is expected that we have more generalized than partial seizures. But in this “chicken or egg” situation where longer suffering from epilepsy-more seizures-more frequent seizures induced lower Interictal ionized magnesium/total serum magnesium ratio, and low magnesium induced more seizures (mostly generalized), situation is not so clear. We have more generalized seizures but odds ratio was 0.95 for 95% CI 0.43 to 2.09, and differences were not significant. We assume that DRE of unknown cause is by itself a risk factor, and that the status epilepticus, more frequent seizures and longer suffering from epilepsy are more dominant for developing protracted brain injury than type of seizures.

Age, duration of epilepsy, and seizure frequency were significantly different among study and control group. There are not enough literature data in this field to compare with our results, but

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**Figure 2:** Interictal ionized magnesium/total serum magnesium ratio and status epilepticus.

1Interictal ionizes magnesium/total serum magnesium ratio
there are some studies on total magnesium, or ionized magnesium in pediatric population where age, seizure frequency and duration of treatment did not influence the plasma magnesium levels [29,30]. Our results indicate that lower interictal ionized magnesium/ total serum magnesium ratio is associated with older age (53.08 ± 9.78), longer suffering from epilepsy (38.44 ± 10.11), and more frequent seizures, which places our study on the side of other studies proving that intractable epilepsy damage the brain [31-33]. For more precise results, we have excluded 20 patients at the screening due to lower ionized magnesium or low total serum magnesium (Figure 1). Suppose that we didn’t exclude them from the study at the screening visit, our results on the “age dependent - longer suffering from epilepsy “model, and “longer suffering from epilepsy- more seizures” model would not have such importance.

Maybe the most intriguing, severe, and the previously least explored illness related factor which we monitored in our study, is occurrence of status epilepticus. There is a lot of evidence that magnesium is efficient in treatment of the SE [34], less evidence on lower serum magnesium levels in SE [35], but there is almost no evidence on the state of ionized magnesium during the SE or after it. More than 15% of the PWE had SE at least once during the illness, and patients with drug resistant epilepsy is even more at risk [36]. Status epilepticus in humans and animal models results in significant cerebral damage and in increased risk of subsequent seizures, associated with a characteristic pattern of neuronal loss particularly affecting the hippocampus [37-41]. Exactly as we knew that SE could cause brain damage, and that the low ionized magnesium levels are could be a clinical marker of the brain injury, we have anticipated that low ionized magnesium/total serum magnesium ratio would be in significant correlation with SE. This proved to be true because 95, 2% of the patients in the study group had SE, and that there is significant difference between the groups in the occurrence of the SE, with relative risk to have low interictal ionized magnesium/total serum magnesium ratio of 14.67 if you experienced status epilepticus. This proves that the appearance of status epilepticus is the most accurate factor for lower interictal ionized magnesium/total serum magnesium ratio and therefore for the protracted brain injury.

Some data in the literature show that some AEDs, in the first place phenobarbital, could cause brain damage, and another one claiming that other AEDs doesn’t cause additional brain damage [42,43]. Our results indicate that there was a significant difference among groups in relation to the longest used first line AEDs, and the study group AEDs profile was with the domination of the phenobarbital and the minimum of the valproate and lamotrigine, while at the same time the control group had no specific AEDs pattern. Our results support the evidence on the phenobarbital’s potential for brain damage.

**Conclusion**

Patients with drug resistant epilepsy of unknown cause have low interictal ionized magnesium/ total serum magnesium ratio which can indicate on the protracted brain injury. We assume that DRE of unknown cause is by itself a risk factor for the low interictal ionized magnesium/total serum magnesium ratio, but more evidence are necessary to proclaim ionized magnesium as marker of the protracted brain injury induced by drug resistant epilepsy. Predictors for the low interictal ionized magnesium/total serum magnesium ratio in this group of patients are: status epilepticus, older age in relation with longer suffering from epilepsy, more frequent seizures, and longest total intake of the phenobarbital.

**References**


