Randomized Controlled Trial between Levetiracetam and Phenobarbital in the Treatment of Neonatal Seizure due to Perinatal Asphyxia

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Abstract:

Background: Seizure occurs more frequently in neonatal period and incidence of seizure is 50%-68% in perinatal asphyxia. At present phenobarbital is the drug of choice for treating neonatal seizure, which has some adverse effects on neurodevelopment status. Levetiracetam is a novel antiepileptic agent well-tolerated and effective in focal, generalized and neonatal seizure as well and lacks the adverse effects like phenobarbital. The present study was undertaken to compare the safety and efficacy of levetiracetam to phenobarbital in the treatment of neonatal seizure due to perinatal asphyxia.

Methodology: This interventional study (Randomized Controlled Trial) was conducted in Department of Neonatology, Bangabandhu Sheikh Mujib Medical University (BSMMU), Dhaka and Dhaka Medical College Hospital, Dhaka, Bangladesh from 1st January’ 2014 to 30th June’ 2015. Intravenous levetiracetam injection, 50 mg/kg loading followed by 10 mg/kg 8 hourly maintenance was used and phenobarbital intravenous 20-40 mg/kg loading and 2.5 mg/kg/dose 12 hourly maintenance was given as per institutional protocol.

Results: Sixty-nine term asphyxiated neonates (intention to treat population) provided analyzable data. Seizure control was found significantly higher (p = 0.011) higher in levetiracetam group in comparison to phenobarbital group (71% vs 40%). Need for more than one drug was significantly lower in levetiracetam group (p=0.011). Adverse effects were found significantly (p=0.001) lower in levetiracetam group (9% vs 43%). No serious adverse effect was observed in any group and most common adverse effect was somnolence in both group followed by irritability. Restlessness, sedation and shallow breathing were found only in phenobarbital group.

Conclusion: Levetiracetam is more effective and safe in comparison to phenobarbital in the treatment of neonatal seizure due to perinatal asphyxia.

Key words: Seizure, Perinatal Asphyxia, Levetiracetam, Phenobarbitone.

Introduction:
Seizure occurs more frequently in neonatal period than at any other time of life and incidence of neonatal seizure ranges from 0.95 to 3.5 per thousand live birth.¹ According to Brown et al and Finer et al, incidence of seizure is 50%-68% in perinatal asphyxia.² At present phenobarbital is the drug of choice for treating neonatal seizure, which has some adverse effects on neurodevelopment.³ In addition, the known risk of cognitive impairment of phenobarbital in infants and toddlers should be considered.³ Levetiracetam is a novel antiepileptic agent well-tolerated and effective in focal, generalized and neonatal seizure as well and currently licensed as adjunctive therapy in the treatment of children and...
infants with epilepsy starting from 1 month of age in partial onset seizures, and already licensed in children aged 4 years or above for other indications. Studies showed its efficacy 71% - 86%\textsuperscript{4,5} whereas efficacy was found only 29%-85% in case of Phenobarbitone\textsuperscript{6,7} More over levetiracetam is rapidly and completely absorbed after oral administration.\textsuperscript{8} In a study by Maitre et al Levetiracetam showed better neurodevelopmental outcome in comparison to phenobarbitone\textsuperscript{9} levetiracetam has not been found to increase neuronal apoptosis in animal models.\textsuperscript{10} Prospective studies with small patient groups in infants and very young children revealed similar results.\textsuperscript{11,12} Levetiracetam is increasingly being used as an antiepileptic drug in the neonatal period. Use of levetiracetam in term and preterm neonates with rarely observed adverse effects were found in an analysis of surveys from neonatologists and pediatric neurologists.\textsuperscript{13} Levetiracetam may be an alternative for hypoxia induced neonatal seizure. No such study was done comparing safety and efficacy of levetiracetam with phenobarbitone in seizure due to perinatal asphyxia. So this study is intended to evaluate the efficacy and side effects of levetiracetam which may open a new frontier of neonatal seizure management and may help to tailor new guideline.

Materials and Methods
It was a randomized controlled trial conducted in the Department of Neonatology, Bangabandhu Sheikh Mujib Medical University (BSMMU), Dhaka and Dhaka Medical College Hospital, Dhaka, Bangladesh from 1st January, 2014 to 30st June, 2015 with prior institutional review board (IRB) approval. Term neonates, aged up to 48 hours, admitted in Neonatal Intensive Care Unit with history of perinatal asphyxia and clinical seizure (both before and after admission) diagnosed by doctors and those have given written consent were included in the study. Convulsion management was given according to institutional protocol (Maintenance of temperature, airway, breathing, circulation, correction of hypoglycemia & hypocalcaemia) except the anticonvulsants. For seizure control in neonates with active convulsion either levetiracetam or phenobarbital was allocated according to the randomization. Levetiracetam receivers were in intervention group and phenobarbital receivers were in control group. Levetiracetam was administered intravenous as a loading dose of 50 mg/kg. Maintenance was administered intravenous at a dose of 10 mg/kg 8 hourly. All bolus and maintenance doses were diluted with normal saline to a concentration of 20 mg/ml and boluses were administered over 15 minutes. Phenobarbital loading was given at a dose of 20-40 mg/kg intravenously and maintenance doses were given 2.5 mg/kg/dose 12 hourly according to institutional protocol. When convulsion stopped completely and not recurred within 48 hours then it was considered as convulsion control and maintenance was followed up upto 48 hours. Adverse effects like life threatening conditions as well as somnolence, irritability, insomnia, hypersomnia, increased frequency of convulsion, respiratory depression, rash, and any other side effects were noted. When one drug either Phenobarbital or levetiracitum failed to control convulsion alone in a patient then that patient was treated as institutional protocol (Phenobarbital Fosphenytoin Midazolam and so on). Drugs were purchased from local market and all the levetiracetam were from same manufacturer and which is also same in case of Phenobarbital. For statistical analysis Independent sample t-test and chi-square test were used and p value<0.05 was considered significant. Statistical software used for analysis of data was SPSS version 20.

Results:
In this study 69 patients reached randomization and out of them 34 were allocated to intervention group and 35 were allocated for control group. Finally, 34 from intervention group and 35 from control group was analyzed (Fig.-1). Most of the patients were normal in birth weight category and it was 94% and 86% in intervention and control group respectively. Most of the patients were delivered by normal vaginal delivery in both groups. It was 68% and 77% in intervention and control group respectively. Seizure developed in most of the patients within 12 hours of birth. In intervention group 74% and in control group 71% neonates developed seizure within 12 hours (Fig.-2). Most of the patients got admitted within 12 hours of age, among them 62% were in intervention group and 63% in control group. Among the four basic types of seizure clonic type was predominant in both groups. It was 56% and 49% in intervention and control group respectively followed by subtle type which was 32% in intervention group and 37% in control group. There was no myoclonic type seizure in any of the group (Fig.-3).
Fig.- 1: Diagram of flow of participants in each stage of the study.

Fig.- 2: Age at seizure onset

Fig.- 3: Seizure type in patients
Seizure control was significantly higher (p = 0.011) in intervention group (71%), but it was 40% in control group and need for more than one drug was significantly lower in intervention group (p = 0.011). Adverse effects were significantly lower in intervention group (p = 0.001), which was 9% in intervention group and 43% in control group and no life threatening adverse effect was observed in both the groups. There was no significant difference in terms of hospital stay in both the groups but mean hospital stay was less in intervention group in comparison to control group (Table-II). After predefined period of follow up (48 hours), finally 68% patients of intervention group and 69% patients of control group were discharged with advice.

Discussion:
Neonatal seizure has wide variety of causes but in this study we focused on neonatal seizure due to perinatal asphyxia, the most common cause.

In baseline data, male sex was found predominant in comparison to female in both groups. It is probably due the social custom of our country where still male offspring is more preferred and taken care of by the parents and family members and they are less interested to spend time and money for female offspring by taking them to distant higher level hospitals. In this study it was found that with a comparable baseline characteristics levetiracetam has significantly higher efficacy in comparison to phenobarbital (71% vs 40%) in controlling neonatal seizure due to perinatal asphyxia. Furwentsches et al. on their pilot study on 6 neonates with seizure used levetiracetam 10 mg/kg/day loading increasing to 30 mg/kg/day over 3 days. All 6 patients became seizure free within 6 days. They used much lower loading dose than us, and used additional doses of phenobarbital during titration of levetiracetam dose and needed several doses and days to control seizure though they had 100% seizure control over 6 days, it is not clear whether this seizure control was attributed by phenobarbital or not. Whereas we used single loading dose of 50 mg/kg followed by 10 mg/kg maintenance 8 hourly and only immediate seizure control was taken into account. Khan et al in their retrospective study with levetiracetam as first or second line anticonvulsant reported 86% seizure control within one hour but some of them recurred.
so the actual control would be lower and for that we think that current study finding is comparable. Khan et al. again in their retrospective study reported 82% seizure control in preterm neonates over 24 hours. It would be lesser if they only take into account those cases with immediate seizure control like this current study and would be comparable. Khan et al retrospectively reviewed neonates treated with 10-50 mg/kg loading of levetiracetam for seizure found 86% seizure control within one hour and 100% within 72 hours and we think our finding would be comparable if we take seizure control over one hour into account. Pharmacokinetic studies have established a benign safety profile for levetiracetam. Li et al. used initial loading doses of 15 to 40 mg/kg and demonstrated that levetiracetam was well-tolerated in 18 neonates with seizures. The only adverse event present was somnolence. They also found linear kinetics, minimal protein binding and no hepatic metabolism with levetiracetam use in children and neonates. In our study a dose of 50 mg/kg was used as an initial loading dose and were maintained on 10 mg/kg every 8 hours and patients tolerated well. There are no clear dosing recommendations available currently in the literatures where doses range from 10 to 100 mg/kg were used. In terms of adverse effects levetiracetam was found much safer in comparison to phenobarbital. Adverse effects, were found in 9% cases of levetiracetam group and 43% in phenobarbital group. Most of the study with levetiracetam found no significant adverse effect and like us they also found somnolence as comparatively commoner insignificant side effect.

Conclusion: Levetiracetam is more effective and safe in comparison to phenobarbital in the treatment of neonatal seizure due to perinatal asphyxia. Multicenter study with larger sample size, use of EEG, monitoring drug concentration and renal functions are recommended.

References


