Neonatal Seizure Management: Is the Timing of Treatment Critical?

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Objective To assess the impact of the time to treatment of the first electrographic seizure on subsequent seizure burden and describe overall seizure management in a large neonatal cohort.

Study design Newborns (36-44 weeks of gestation) requiring electroencephalographic (EEG) monitoring recruited to 2 multicenter European studies were included. Infants who received antiseizure medication exclusively after electrographic seizure onset were grouped based on the time to treatment of the first seizure: antiseizure medication within 1 hour, between 1 and 2 hours, and after 2 hours. Outcomes measured were seizure burden, maximum seizure burden, status epilepticus, number of seizures, and antiseizure medication dose over the first 24 hours after seizure onset.

Results Out of 472 newborns recruited, 154 (32.6%) had confirmed electrographic seizures. Sixty-nine infants received antiseizure medication exclusively after the onset of electrographic seizure, including 21 infants within 1 hour of seizure onset, 15 between 1 and 2 hours after seizure onset, and 33 at >2 hours after seizure onset. Significantly lower seizure burden and fewer seizures were noted in the infants treated with antiseizure medication within 1 hour of seizure onset (P = .029 and .035, respectively). Overall, 258 of 472 infants (54.7%) received antiseizure medication during the study period, of whom 40 without electrographic seizures received treatment exclusively during EEG monitoring and 11 with electrographic seizures received no treatment.

Conclusions Treatment of neonatal seizures may be time-critical, but more research is needed to confirm this. Improvements in neonatal seizure diagnosis and treatment are also needed. (J Pediatr 2022;243:61-8).

See editorial, p 7

O wing to the unique physiologic properties of the immature brain,1 seizures are common in newborn infants, with an incidence of 1-3.5/1000 live births in term infants.2,4 Although a wide variety of causes have been reported, the leading cause remains hypoxic ischemic encephalopathy (HIE), despite the introduction of therapeutic hypothermia.3,5 Seizure recognition is challenging in newborns because many seizures are subclinical or have subtle clinical manifestations; in addition, treatment of seizures can cause an “uncoupling” of clinical and electroencephalography (EEG) features.6

There has been an increase in the use of EEG in neonatal units for diagnosing seizures.5,7 Amplitude-integrated EEG (aEEG) is still preferred by some neonatologists, because it is easy to perform and not dependent on 24/7 neurophysiology support.8,9 International guidelines recommend the use of continuous conventional electroencephalography (cEEG) for a minimum of 24 hours as the gold standard for seizure diagnosis in the newborn.10-12

There is increasing evidence that neonatal seizures are associated with poor neurodevelopmental outcome, and that untreated seizures might add to the initial brain injury.13,14 Responsiveness to antiseizure medication may decrease with recurrent and prolonged seizures.15-17 The hypothesis that earlier treatment leads to better response is supported by animal work in mice and rats which has shown a progressive increase in intracellular chloride with recurrent seizures, increasing the likelihood of more seizures and decreasing the responsiveness to treatment.17 Current published guidelines for management of all neonatal seizures recommend initiation of antiseizure medication as soon as possible.

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aEEG Amplitude-integrated electroencephalography

cc EEG Conventional electroencephalography

EEG Electroencephalography

HIE Hypoxic ischemic encephalopathy
following seizure recognition, but there are no recommendations on specific target times for treatment. There remains significant variability in how seizures are diagnosed and managed, and a consensus is needed.

Our group previously showed that only 11% of “seizure episodes” were treated within the first hour after onset. Another study investigating the time from onset of aEEG seizure to treatment with antiseizure medication found that 32.1% of patients were treated within 1 hour of onset, 19.8% were treated within 1-2 hours of onset, and the majority (48.1%) were treated at >2 hours after onset. The primary aim of our present analysis was to assess whether the time to treatment of first electrographic seizure has an impact on subsequent seizure burden. To achieve this goal, we used a large multicenter European neonatal cohort. Our secondary aim was to describe the initial seizure management for all infants included in this cohort.

**Methods**

The present study is a secondary data analysis of 2 European multicenter cohort studies that recruited newborns across 8 European tertiary neonatal intensive care units between January 2011 and February 2017 (ClinicalTrials.gov identifiers NCT02160171 and NCT02431780). Both studies examined the feasibility and efficacy of cEEG monitoring and a new automated neonatal seizure detection algorithm (ANSeR algorithm) and included infants of 36-44 weeks corrected gestational age requiring EEG monitoring for suspected seizures. Infants were excluded only if parental/guardian written consent was refused or they were at corrected gestational age <36 weeks. Relevant information regarding delivery and neonatal course were recorded in study-designed electronic databases. To assess the effect of treatment timing of the first electrographic seizure (main aim), we included all infants who received antiseizure medication exclusively after electrographic seizure onset and who had at least 24 hours of cEEG recording after electrographic seizure onset. The secondary aim was to describe the initiation of seizure management for all infants included in the 2 studies.

Ethical approval was granted for both studies by national and local Ethics Committees specific to each participating center.

**EEG Monitoring**

All infants underwent prolonged video cEEG monitoring as clinically indicated, using a 10:20 EEG electrode modified neonatal system with disposable electrodes placed at F3, F4, C3, C4, Cz, T3, T4, O1/P3, and O2/P4. Three different EEG machines were used for monitoring: Neurofax EEG-1200 (Nihon Kohden), NicoletOne ICU Monitor (Natus), and XLTek EEG (Natus). Teams at each site were trained in EEG electrode application and maintenance of good quality recordings. The clinical teams at each site had the aEEG signals from F3-C3 and F4-C4, 8 raw EEG channels, and electrocardiography and respiratory traces displayed on the EEG monitors and available to review. No standard EEG review protocol was imposed during the study period, and the clinical teams reviewed the monitoring data as recommended by the local guidelines.

**Seizure Analysis**

This analysis included all the EEG monitoring performed for each infant, regardless of the time of study enrollment. Electrographic seizures were annotated by 1 of 4 neonatal neurophysiologists blinded to the infant’s medical history and outcome. A standard EEG review protocol for seizure annotation was used. Electrographic seizures were defined as a minimum 10 seconds of sudden, repetitive, and evolving stereotyped waveforms involving at least 1 EEG channel. An infant was considered to have seizures if at least 1 electrographic seizure was annotated. The following summary measures of seizures were calculated: seizure period (time [in hours] from the beginning of the first electrographic seizure to the end of the last electrographic seizure), total seizure burden (duration [in minutes] of all seizures occurring during the entire monitoring period), maximum hourly seizure burden (maximum seizure burden within 1 hour [in minutes/hour]), status epilepticus (seizure burden in a minimum of 30 minutes within 1 hour), and number of seizures during the entire monitoring period. Seizure characteristics were described for all infants who experienced seizures during the EEG monitoring period.

**Antiseizure Medication Treatment**

EEG recordings were reviewed by local teams from each site in real time, and all seizures detected were managed according to local protocols. For each infant, antiseizure medication treatment was recorded in the study database (ie, type of drug, dose, and time of administration). The antiseizure medication was chosen according to local protocols and at the clinician’s discretion.

**Treatment Timing Analysis**

For the purpose of treatment timing analysis, we considered the first antiseizure medication dose administered after the first electrographic seizure. This analysis included only infants with anti-seizure medication given exclusively after electrographic seizure onset and with a minimum of 24 hours of EEG. Infants who received antiseizure medication before their first electrographic seizure and infants who did not receive antiseizure medication treatment throughout the study period were excluded from this analysis, as this could be a bias for seizure diagnosis and management by the clinical teams. The treatment timing cohort was divided into 3 groups: treatment with antiseizure medication within 1 hour of electrographic seizure onset, between 1 and 2 hours after electrographic seizure onset, and at >2 hours after electrographic seizure onset. The primary outcome was seizure burden, and the secondary outcomes were maximum seizure burden, presence of status epilepticus, number of seizures, and total number of antiseizure medication doses. The American Clinical Neurophysiology Society guidelines recommend at least 24 hours of cEEG monitoring for neonates at risk of seizures and when seizures are confirmed, at least 24 hours of seizure-free cEEG; therefore, all outcomes were calculated over the first 24 hours after electrographic seizure onset. A post hoc analysis was performed in the subgroup of infants.
diagnosed with HIE, from which infants with encephalopathy caused by factors other than HIE injury were excluded.

Statistical Analyses
Data are reported as median and IQR for continuous variables and as frequency and percentage for categorical variables. Differences in outcomes between the treatment groups were investigated based on type of data and normality. The Mann-Whitney U test and Kruskal-Wallis test were used for continuous outcome variables, and the χ² test was used for categorical outcome variables. Post hoc pairwise comparisons with a Bonferroni correction were performed if the omnibus test was significant. All tests were 2-sided, and a \( P \) value < .05 was considered statistically significant. Statistical analysis was performed using SPSS version 25.0 (IBM).

Results
The 2 studies included a total of 472 infants: 318 (67.4%) without seizures and 154 (32.6%) with seizures (Figure 1). The neonatal characteristics for the whole cohort are presented in Table I (available at www.jpeds.com). The percentages of infants with moderate and severe HIE, stroke, and metabolic/genetic disorders were higher in the seizure group compared with nonseizure group, but otherwise the 2 groups were similar.

Seizure Characteristics and Treatment
Out of 154 infants with evidence of electrographic seizures, 31 (20.1%) had no antiseizure medication given after electrographic seizure onset, including 11 who received no antiseizure medication at all and 20 who received antiseizure medication only before electrographic seizure onset (Table II). Based on the timing of antiseizure medication administration after electrographic seizure onset, 26 infants (16.9%) received treatment within 1 hour, 23 (14.9%) received treatment between 1 and 2 hours, and 74 (48.1%) received treatment after 2 hours.

The median age of seizure onset for all infants was 22 (IQR, 14-54) hours after birth, with a median total seizure burden of 69 (IQR, 23-154) minutes. Compared with infants in the other groups, those in the group treated with antiseizure medication within 1 hour had an earlier onset of seizure, a longer duration of first electrographic seizure, and a higher seizure burden in the first hour after seizure onset (Table II).

Primary Aim: Treatment Timing Analysis
This analysis included infants who received antiseizure medication exclusively at any time point after the first electrographic seizure (n = 69 infants) (Table III). We excluded infants who did not receive any antiseizure medication during the study period (n = 11), infants who received antiseizure medication exclusively before electrographic seizure onset (n = 20), infants who received antiseizure medication both before and after electrographic seizure onset (n = 52), and infants with <24 hours of EEG monitoring after their first electrographic seizure (n = 2).

Seizure burden and the number of seizures differed significantly between the antiseizure medication treatment groups (\( P = .029 \) and .035, respectively). The pairwise analysis showed that from the onset of first electrographic seizure, the seizure burden calculated over the subsequent 24 hours was significantly lower in the <1 hour antiseizure medication group compared with the >2 hour antiseizure medication group (median seizure burden 36 [IQR, 15-70] minutes vs 75 [IQR, 30-152] minutes; \( P = .025 \)) (Figure 2; available at www.jpeds.com). The number of seizures was also significantly lower in the <1 hour group compared with the >2 hour group (median number of seizures, 10 [IQR, 2-24] vs 28 [IQR, 11-50]; \( P = .032 \)).

Figure 1. Study flow diagram. ASM, anti-seizure medication; EEG, electroencephalography.
We investigated etiology, therapeutic hypothermia status, age at start of EEG monitoring, and age at first electrographic seizure as potential confounding variables and found that they were not associated with the timing of antiseizure medication administration. The background etiologies were not statistically significantly different among the antiseizure medication treatment groups (<1 hour group: moderate HIE, n = 7 [33.3%]; severe HIE, n = 5 [28.6%]; 1-2 hour group: moderate HIE, n = 6 [28.6%]; severe HIE, n = 5 [23.8%]; stroke, n = 3 [14.3%]; other, n = 6 [28.6%]; >2 hours group: moderate HIE, n = 12 [36.4%]; severe HIE, n = 8 [24.2%]; stroke, n = 7 [21.2%]; other, n = 6 [18.2%]; P = .928). Therapeutic hypothermia was provided to 12 infants (57.1%) in the <1 hour group, 9 infants (60%) in the 1-2 hours group, and 18 infants (54.5%) in the >2 hours group (P = .937).

The median age at the start of EEG monitoring 6.9 (IQR, 3.4-38.6) hours of life in the <1 hour group, 9.0 (IQR, 3.0-41.6) hours of life in the 1-2 hours group, and 6.9 (IQR, 3.6-33.1) hours of life in the >2 hours group (P = .976). The median age at the first electrographic seizure was 14.1 (IQR, 8.6-39.6) hours of life in the <1 hour group, 16.2 (IQR, 9-42.3) hours of life in the 1-2 hours group, and 14.6 (9.9-48.2) hours of life in the >2 hours group (P = .851).

Comparing the group of infants who received antiseizure medication exclusively after their first electrographic seizure (n = 69) with the group who received antiseizure medication before and after their first seizure (n = 52), there were no significant differences in terms of total seizure burden (median, 74 [IQR, 32-167] minutes vs 79 [IQR, 21-158] minutes; P = .582), maximum seizure burden (median, 24 [IQR, 13-34] minutes vs 19 [IQR, 8-29] minutes; P = .071), number of seizures (median, 25 [IQR, 10-50] vs 33 [IQR, 12-58]; P = .449), and presence of status epilepticus (24 [34.8%] vs 12 [23.1%]; P = .163).

Post hoc analysis also was performed in the subgroup of infants who had a diagnosis of HIE (n = 42), and the seizure burden was significantly different among the antiseizure medication treatment groups (P = .009). The pairwise comparison showed a significantly lower seizure burden in the <1 hour group compared with the >2 hours group (median 41 [IQR, 24-67] minutes vs 86 [IQR, 68-168] minutes; P = .007) (Table IV). For the HIE cohort, therapeutic hypothermia status and age at the start of EEG monitoring were investigated as potential confounders and were not associated with antiseizure medication timing (P = .834 and = .984, respectively).

Secondary Aim: Overall Description of Antiseizure Medication Treatment

Among the 472 infants, 258 (54.7%) received at least 1 dose of antiseizure medication before or during EEG monitoring.

### Table II. Seizure characteristics* of all infants with electrographic seizures and for infants by treatment timing group after electrographic seizure onset: descriptive analysis

| Characteristics                              | All (n = 154) | Antiseizure medication treatment at <1 h (n = 26) | Antiseizure medication treatment at 1-2 h (n = 23) | Antiseizure medication treatment at >2 h (n = 74) | No antiseizure medication treatment (n = 31) |
|---------------------------------------------|--------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|-----------------------------------------------|
| General characteristics                     |              |                                               |                                               |                                               |                                               |
| Seizure period, h, median (IQR)             | 16.5 (6.7-40.2) | 18.1 (0.7-57.4) | 16.6 (8.3-54.5) | 26.6 (10.0-48.2) | 8.1 (2.7-14.8) |
| Total seizure burden, min, median (IQR)     | 69 (23-154) | 45 (21-127) | 104 (34-167) | 75 (27-162) | 30 (4-106) |
| Number of seizures, median (IQR)            | 21 (9-52) | 23 (5-33) | 28 (12-52) | 32 (12-60) | 10 (2-32) |
| Median seizure duration, s, median (IQR)    | 104 (65-189) | 89 (56-495) | 105 (75-191) | 108 (64-160) | 98 (45-164) |
| Maximum seizure burden, min/h, median (IQR)| 22 (10-32) | 22 (12-36) | 24 (16-35) | 22 (10-31) | 14 (2-28) |
| Age at first electrographic seizure, h, median (IQR) | 35 (19-63) | 21 (11-49) | 32 (19-80) | 36 (20-68) | 38 (28-59) |
| Status epilepticus (yes), n (%)             | 43 (27.9) | 8 (30.8) | 8 (34.8) | 20 (27.0) | 7 (22.6) |
| Characteristics related to first electrographic seizure |              |                                               |                                               |                                               |                                               |
| Age at first electrographic seizure, h, median (IQR) | 22 (14-54) | 14 (9-39) | 20 (15-57) | 21 (14-56) | 36 (19-59) |
| Duration of first seizure, s, median (IQR)  | 93 (46-273) | 268 (65-1385) | 104 (37-681) | 75 (43-161) | 114 (45-287) |
| Seizure burden in the first hour of seizure period, min, median (IQR) | 6.0 (2.3-15.1) | 15.7 (10.0-30.8) | 9.7 (2.6-22.1) | 4.4 (1.8-10.2) | 3.9 (1.9-11.7) |
| Number of seizures in the first hour of seizure period, median (IQR) | 2 (1-4) | 2 (1-6) | 3 (1-4) | 2 (1-4) | 2 (1-3) |

*Seizure characteristics were calculated based on complete EEG monitoring throughout the study period.
The infants included in this analysis were those who received no antiseizure medication before their first electrographic seizure, with at least 1 antiseizure medication dose given after the first electrographic seizure and with at least 24 hours of EEG monitoring after the first electrographic seizure.

Similarly, the post hoc analysis investigating treatment who received antiseizure medication after 1 hour of seizure burden in the first hour in this group) compared with infants the subsequent following 24 hours (despite a higher seizure onset had the lowest seizure burden and fewer seizures over 32.1% of the infants received antiseizure medication starting within 1 hour of aEEG seizure onset.22 The higher proportion of early treatment in that study compared with our cohort could be explained by their use of a seizure detection algorithm and a single expert site. Previous studies have shown that a high seizure burden is independently associated with worse brain injury detected on magnetic resonance imaging timing in infants with a diagnosis of HIE showed a lower seizure burden in the early (within 1 hour) treatment group. Current international guidelines recommend that treatment for neonatal seizures should be administered as soon as possible but do not specify an optimal treatment target.18 The ANSeR phase 1 study cohort reported by Rennie et al was included in this analysis, together with a neonatal cohort (ANSeR phase 2 study) recruited for a randomized controlled trial of a seizure detection algorithm.3,23 Rennie et al showed that only 11% of “seizure episodes” (clusters of seizures separated by <2 hours) were treated within 1 hour. In the present study, we analyzed only the treatment of first electrographic seizures and found that among all the infants with electrographic seizures, 26 (16.9%) received treatment for their first electrographic seizure within the first hour; however, only 21 received antiseizure medication exclusively after the onset of electrographic seizure (5 infants received antiseizure medication before the start of EEG monitoring due to clinical seizures). In another study, 32.1% of the infants received antiseizure medication starting within 1 hour of aEEG seizure onset.22 The findings of this study demonstrate that the group of infants with electrographic seizures treated within 1 hour of seizure onset had the lowest seizure burden and fewer seizures over the subsequent following 24 hours (despite a higher seizure burden in the first hour in this group) compared with infants who received antiseizure medication after 1 hour of seizure onset. Similarly, the post hoc analysis investigating treatment

### Discussion

The infants included in this analysis were those who received no antiseizure medication before their first electrographic seizure, with at least 1 antiseizure medication dose given after the first electrographic seizure and with at least 24 hours of EEG monitoring after the first electrographic seizure.

*P values were from the Kruskal-Wallis test unless indicated otherwise. P < .05 was considered to indicate statistical significance.

†P value from the χ² test.

### Table III. Antiseizure treatment group analysis (n = 69)

| Outcomes                                      | Treatment at <1 h (n = 21) | Treatment at 1-2 h (n = 15) | Treatment at >2 h (n = 33) | P value* | Pairwise comparison |
|-----------------------------------------------|-----------------------------|-----------------------------|-----------------------------|----------|---------------------|
| Seizure burden within 24 h, min, median (IQR) | 36 (15-70)                  | 71 (32-112)                 | 75 (30-152)                 | .029     | <1 h vs >2 h        |
| Maximum seizure burden within 24 h, min/h, median (IQR) | 16 (11-24)                  | 20 (12-40)                  | 27 (12-35)                  | .224     |                     |
| Number of seizures within 24 h, median (IQR)  | 10 (2-24)                   | 18 (6-32)                   | 28 (11-50)                  | .035     | <1 h vs >2 h        |
| Status epilepticus within 24 h (yes), n (%)   | 3 (14.3)                    | 4 (26.7)                    | 14 (42.4)                   | .089†    |                     |
| Total doses of antiseizure medication within 24 h, median (IQR) | 2 (1-4)                     | 2 (1-3)                     | 2 (1-3)                     | .712     |                     |

The infants included in this analysis were those who received no antiseizure medication before their first electrographic seizure, with at least 1 antiseizure medication dose given after the first electrographic seizure and with at least 24 hours of EEG monitoring after the first electrographic seizure. *P values were from the Kruskal-Wallis test unless indicated otherwise. P < .05 was considered to indicate statistical significance.

†P value from the χ² test.

### Table IV. Antiseizure treatment group analysis for infants with HIE (n = 42)

| Outcomes                                      | Treatment at <1 h (n = 12) | Treatment at 1-2 h (n = 10) | Treatment at >2 h (n = 20) | P value* | Pairwise comparison |
|-----------------------------------------------|-----------------------------|-----------------------------|-----------------------------|----------|---------------------|
| Seizure burden within 24 h, min, median (IQR) | 41 (24-67)                  | 87 (34-113)                 | 86 (68-168)                 | .009     | <1 h vs >2 h        |
| Maximum seizure burden within 24 h, min/h, median (IQR) | 21 (13-35)                  | 24 (11-41)                  | 32 (23-40)                  | .132     |                     |
| Number of seizures within 24 h, median (IQR)  | 10 (1-23)                   | 15 (6-34)                   | 24 (11-58)                  | .056     |                     |
| Status epilepticus within 24 h (yes), n (%)   | 3 (25.0)                    | 3 (30.0)                    | 12 (60.0)                   | .098†    |                     |
| Total doses of antiseizure medication within 24 h, median (IQR) | 2 (1-4)                     | 3 (2-3)                     | 2 (1-4)                     | .909     |                     |

The infants included in this analysis were those who received no antiseizure medication before their first electrographic seizure, with at least 1 antiseizure medication dose given after the first electrographic seizure and with at least 24 hours of EEG monitoring after the first electrographic seizure.

*P values were from the Kruskal-Wallis test unless indicated otherwise. P < .05 was considered to indicate statistical significance.

†P value from the χ² test.

Neonatal Seizure Management: Is the Timing of Treatment Critical?
and worse neurodevelopmental outcomes, suggesting that early antiseizure medication administration resulting in a reduction of seizure burden might lead to a decrease in magnetic resonance imaging–detected brain injury and better long-term outcomes.13,25-27 A randomized controlled trial netic resonance imaging–detected brain injury and better reduction of seizure burden might lead to a decrease in mag-

early antiseizure medication administration resulting in a and worse neurodevelopmental outcomes, suggesting that

THE JOURNAL OF PEDIATRICS

studies have documented that exposure to antiseizure medica-

tor hypothermia status, age at initiation of EEG moni-

toring, and age at first electrographic seizure as possible confounders for this analysis but noted no significant differ-

ences among the treatment groups. Our results are supported by previous animal and human work demonstrating that the sooner the treatment, the better the response.17,29

For our analysis of treatment timing, we had a large multi-
center European neonatal cohort from eight tertiary neonatal intensive care units available, and thus we also described the overall seizure management in this cohort. Out of 472 new-

borns, 258 (54.7%) received at least 1 dose of antiseizure medication during the study period. Seventy-eight infants received antiseizure medication before EEG monitoring was started for suspected clinical seizures, among whom 58 infants had no evidence of further electrographic seizures. We can only assume that in these infants, the seizures responded to the initial antiseizure medication treatment, or that what was diagnosed as a clinical seizure was likely part of a neonatal movement disorder rather than a true seizure, as demonstrated previously by our group.30

The majority of infants who received antiseizure medication exclusively during EEG monitoring had demonstrated electro-

graphic seizures; however, 40 infants received antiseizure medica-
tion exclusively during EEG monitoring and had no evidence of electrographic seizures. Most of the infants who did not receive antiseizure medication had no electrographic seizures throughout the duration of EEG monitoring (203 infants); how-
ever, 11 infants with electrographic seizures had no treatment. Among these 11 infants with electrographic seizures, 6 had a to-
tal seizure burden of >20 minutes and 2 had status epilepticus. These results demonstrate that recognition of seizures remains a major challenge for neonatologists and inappropriate antisei-

ure medication treatment (undertreatment and overtreat-
ment) remains an ongoing concern. Animal and human studies have documented that exposure to antiseizure medica-
can lead to neuronal apoptosis, poor brain development, and later cognitive impairments; therefore, it is important to administer antiseizure medication appropriately to infants who may already have some degree of brain injury.31-35 These re-

results illustrate the diagnostic difficulties that clinicians face, even when the gold standard diagnostic tool (cEEG) is used. Current guidelines recommend the use of cEEG monitoring for at least 24 hours to detect seizures in neonates, but there are no clear protocols for EEG review, and neurophysiologic support is limited even in tertiary neonatal centers.10 Implementation of EEG monitoring and seizure management protocols and spec-
ific EEG review and interpretation training have been shown to be beneficial.29,36

Several limitations must be considered when interpreting our present results. This is a secondary analysis of infants recruited to 2 European studies of cEEG monitoring, not a prospective study investigating treatment of neonatal seizures. Given the sample sizes, only large differences between antiseizure medication groups could be detected across the outcomes investigated. When seizures were detected, treatment was initiated in accord-

cance with local clinical guidelines, and because there is no consensus on the optimal target time for treatment, the treat-
ment cutoffs used for this analysis were selected based on previ-

s, International League Against Epilepsy definition of neonatal seizure does not include a minimum duration of 10 seconds as long as an evolution in frequency and morphology is demon-

strated.37 In our cohort, the mean duration of the first electro-

graphic seizure was 93 (IQR, 46-273) seconds, and only 9% of the infants had a first electrographic seizure of <30 seconds. Although we cannot definitely rule out that this did not influ-

ence the overall results, we believe that it is unlikely, given that the majority of seizures in our cohort were >10 seconds. How-
ever, we do believe that a more detailed analysis of shorter-
duration discharges using the International League Against Ep-
ilepsy definition could provide valuable insight into the impact of electrographic seizures on the developing brain and is a priori-

ity area for our future research.

For our treatment timing analysis, infants who received anti-

seizure medication before electrographic seizure onset (including previous EEG start) were excluded to minimize the cumulative effect of antiseizure medication on the electro-

graphic seizure burden. When comparing the group of infants with antiseizure medication given exclusively after the first elec-

trographic seizure (n = 69 infants) with the group of infants with antiseizure medication given before and after the first electro-

graphic seizure (n = 52 infants), we found no significant differ-

ences in terms of total seizure burden, maximum seizure burden, number of seizures, or presence of status epilepticus. In addition, infants who did not receive antiseizure medication after the emergence of electrographic seizure were excluded, because electrographic seizures were not recognized and not treated by the cotside clinical teams. As a result, we could include
only 45% (69 of 154) of all infants with electrographic seizures in the treatment timing analysis. Comparing the group of infants included in the treatment timing analysis (n = 69) with the group of infants excluded (n = 85), we noted that more than one-half of the excluded infants were born outside of the recruiting hospitals, and thus the EEG monitoring was started significantly later in this group. All participating centers were tertiary referral hospitals, and almost one-half of the infants with electrographic seizures were outborn. This may explain the high proportion of infants who received antiseizure medication before EEG commencement.

This analysis looked at short-term outcomes, derived from the seizure burden, and our 2-year developmental follow-up analysis is not yet completed. Our findings suggest that there may be an association between treatment timing and subsequent seizure burden and that treatment of neonatal seizures might be time-critical; however, this needs to be confirmed in a large prospective study. Inappropriate treatment remains an ongoing concern; many infants continue to be treated who do not require treatment and in those who do, delayed onset of treatment remains problematic. Recognition of electrographic seizures remains a major challenge for neonatologists. With the increasing use of prolonged neonatal cEEG monitoring and insufficient neonatal neurophysiology expertise, additional support from automated seizure detection algorithms might be the solution.23,38,39

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Data Statement

Data sharing statement available at www.jpeds.com.

References

1. Jensen FE. Neonatal seizures: an update on mechanisms and management. Clin Perinatol 2009;36:881-900. vii.
2. Pisani F, Facini C, Bianchi E, Giussani G, Piccolo B, Beghi E. Incidence of neonatal seizures, perinatal risk factors for epilepsy and mortality after neonatal seizures in the province of Parma, Italy. Epilepsia 2018;59:1764-73.
3. Rennie JM, de Vries LS, Blennow M, Foran A, Shah DK, Livingstone V, et al. Characterisation of neonatal seizures and their treatment using continuous EEG monitoring: a multicentre experience. Arch Dis Child Fetal Neonatal Ed 2019;104:F493-501.
4. Glass HC, Pham TN, Danielsen B, Towner D, Glidden D, Wu YW. Antenatal and intrapartum risk factors for seizures in term newborns: a population-based study, California 1998-2002. J Pediatr 2009;154: 24-8.e1.
5. Glass HC, Shellhaas RA, Wusthoff CJ, Chang T, Abend NS, Chu CJ, et al. Contemporary profile of seizures in neonates: a prospective cohort study. J Pediatr 2016;174:98-103.e1.
6. Malone A, Ryan CA, Fitzgerald A, Burgoyne L, Connolly S, Boylan GB. Interobserver agreement in neonatal seizure identification. Epilepsia 2009;50:2097-101.
7. Boylan G, Burgoyne L, Moore C, O’Flaherty B, Rennie J. An international survey of EEG use in the neonatal intensive care unit. Acta Paediatr 2016;99:1150-5.
8. Battie SG, Sall E, Webster R, Varin M, Lemery B, Hahn C, et al. Continuous electroencephalography monitoring for critically ill neonates: a Canadian perspective. Can J Neurol Sci 2019;46:394-402.
9. Wang Z, Zhang P, Zhou W, Zhou X, Shi Y, Cheng X, et al. Electroencephalography monitoring in the neonatal intensive care unit: a Chinese perspective. Transl Pediatr 2021;10:552-9.
10. Shellhaas RA, Chang T, Tsuchida T, Scher MS, Rivielo JJ, Abend NS, et al. The American Clinical Neurophysiology Society’s guideline on continuous electroencephalography monitoring in neonates. J Clin Neurophysiol 2011;28:611-7.
11. Rennie JM, Chorley G, Boylan GB, Pressler R, Nguyen Y, Hooper R. Non-expert use of the cerebral function monitor for neonatal seizure detection. Arch Dis Child Fetal Neonatal Ed 2004;89:F37-40.
12. Pellegrin S, Munoz FM, Padula M, Heath PT, Meller L, Top K, et al. Neonatal seizures: case definition & guidelines for data collection, analysis, and presentation of immunization safety data. Vaccine 2019;37:7596-609.
13. Kharoshankaya L, Stevenson NJ, Livingstone V, Murray DM, Murphy BP, Ahearne CE, et al. Seizure burden and neurodevelopmental outcome in neonates with hypoxic-ischemic encephalopathy. Dev Med Child Neonol 2016;58:1242-8.
14. De Haan TR, Langeslag I, van der Lee JH, van Kaa AH. A systematic review comparing neurodevelopmental outcome in term infants with hypoxic and vascular brain injury with and without seizures. BMC Pediatr 2018;18:147.
15. Glass HC, Wirrell E. Controversies in neonatal seizure management. J Child Neurol 2009;24:591-9.
16. Ronen GM, Buckley D, Penney S, Streiner DL. Long-term prognosis in children with neonatal seizures: a population-based study. Neurology 2007;69:1816-22.
17. Dhital VI, Kuchibhotla KV, Glykys JC, Kahle TK, Swiercz WB, Feng G, et al. Progressive NRKCCI-dependent neuronal chloride accumulation during neonatal seizures. J Neurosci 2010;30:11745-61.
18. World Health Organisation. Guidelines on neonatal seizures. 2011. Accessed November 10, 2020, https://apps.who.int/iris/bitstream/handle/10665/77756/9789241548304_eng.pdf;jsessionid=72779185D616A91411E068F22276B7F?sequence=1
19. Gossling L, Alik JIP, Stavroulakis T, Hart AR. Investigating and managing neonatal seizures in the UK: an explanatory sequential mixed methods approach. BMC Pediatr 2020;20:36.
20. McNally MA, Hartman AL. Variability in preferred management of electrographic seizures in neonatal hypoxic ischemic encephalopathy. Pediatr Neurol 2017;77:37-41.
21. Glass HC, Glidden D, Jeremy RJ, Barkovich AJ, Ferriero DM, Miller SP. Clinical neonatal seizures are independently associated with outcome in infants at risk for hypoxic-ischemic brain injury. J Pediatr 2009;155:318-23.
22. Apers WMJ, de Vries LS, Groenendaal F, Toet MC, Weeke LC. Delay in treatment of neonatal seizures: a retrospective cohort study. Neonatology 2020;117:599-605.
23. Pavel AM, Rennie JM, de Vries LS, Blennow M, Foran A, Shah DK, et al. A machine-learning algorithm for neonatal seizure recognition: a multicentre, randomised, controlled trial. Lancet Child Adolesc Health 2020;4:740-9.
24. Clancy RR, Legido A. The exact ictal and interictal duration of electroencephalographic neonatal seizures. Epilepsia 1987;28:537-41.
25. Shah DK, Wusthoff CJ, Clarke P, Wyatt JS, Ramaiah SM, Dias RJ, et al. Electrographic seizures are associated with brain injury in newborns undergoing therapeutic hypothermia. Arch Dis Child Fetal Neonatal Ed 2014;49:F219-24.
26. Srinivasakumar P, Zempel J, Trivedi S, Wallendorf M, Rao R, Smith B, et al. Treating EEG seizures in hypoxic ischemic encephalopathy: a randomized controlled trial. Pediatrics 2015;136:e1302-9.
Newborn Screening for Sickle Cell Disease: A Promise (Un)Fulfilled
Huntsman RG, Metters JS, Yawson GI. The diagnosis of sickle cell disease in the newborn infant. J Pediatr 1972;80:279-81.

Sickle cell disease (SCD) is the most common inherited blood cell disorder, affecting more than 100,000 people in the US. The sickle cell trait (SCT) occurs in more than 1.5 million Americans and 8% of African Americans. Current worldwide estimates indicate that there are 300,000 SCD annual births and 100 million people living with SCT.

In 1949, the discovery by Linus Pauling that sickle hemoglobin has a higher isoelectric point enabled the diagnostic use of electrophoretic techniques to diagnose SCD and SCT. Challenges, however, remained as the predominant hemoglobin at birth is fetal hemoglobin (approximately 90%), with only a fraction being sickle hemoglobin. Fifty years ago, Huntsman et al compared multiple diagnostic techniques demonstrating that agar gel electrophoresis could clearly identify SCD and SCT in cord blood. This discovery led to the establishment of newborn screening (NBS) programs for SCD in 1975, which are now implemented universally in the US. Mortality in US children with SCD has been virtually eliminated owing partly to universal NBS (at present using an advanced high-throughput methodology) allowing for timely penicillin prophylaxis and early initiation of comprehensive care. In stark contrast, in low-resource settings such as sub-Saharan Africa, where there is lack of uniform NBS, it is estimated that up to 90% of children with SCD will die before adulthood; one-half these children will not survive until their fifth birthday.

Despite these successes in the US, there are wide variations in states’ follow-up practices and mandates. The number of neonates who receive follow-up for positive SCD NBS results are not tracked. Additionally, although NBS identifies newborns who have SCT, there are no standardized practices for follow-up of carriers and screening of family members. This point is important not just for reproductive decision-making, but also owing to accumulating evidence that SCT is not entirely benign. Although pediatricians take on these tasks despite a lack of adequate funding and structured policy, it is also our moral obligation to advocate for an improved national and international response.

Other rare diseases such as cystic fibrosis and hemophilia, with much lower prevalences, have garnered far greater funding for research and comprehensive clinical care. These racial inequities directly impact the health outcomes of this vulnerable population.

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Figure 2. Seizure burden within 24 hours from electrographic seizure onset by treatment groups (boxplot). ASM, anti-seizure medication.
Table I. Neonatal characteristics

| Characteristics | All infants | Seizure group | Non-seizure group | P value* |
|-----------------|------------|---------------|-------------------|---------|
|                 | Number     | Value         | Number            | Value               | Number     | Value               | Number     | Value               |         |
| Gestational age at birth, wk, median (IQR) | 472 | 40 (39-41) | 154 | 40 (39-41) | 318 | 40 (38-41) | .117 |
| Birth weight, g, median (IQR) | 472 | 3440 (3061-3800) | 154 | 3400 (3088-3793) | 318 | 3450 (3056-3818) | .954 |
| Male sex, n (%) | 472 | 292 (61.9) | 154 | 91 (59.1) | 318 | 201 (63.2) | .388 |
| Mode of delivery, n (%) | 470 | 154 | 316 | .372 |
| Spontaneous vaginal delivery | 171 (36.4) | 57 (37.0) | 114 (36.1) | |
| Instrumental vaginal delivery | 126 (26.8) | 35 (22.7) | 91 (28.8) | |
| Elective cesarean delivery | 43 (9.1) | 18 (11.7) | 25 (8.0) | |
| Emergency cesarean delivery | 130 (27.7) | 44 (28.6) | 86 (27.1) | |
| Place of birth, n (%) | 472 | 154 | 318 | .648 |
| Inborn in recruiting hospital | 255 (54.0) | 81 (52.6) | 174 (54.7) | |
| Outborn home/prehospital | 19 (4.0) | 8 (5.2) | 11 (3.5) | |
| Outborn other hospital | 198 (42.0) | 65 (42.2) | 133 (41.8) | |
| Apgar score at 1 min, median (IQR) | 447 | 3 (1-7) | 148 | 2 (1-8) | 299 | 3 (1-7) | .094 |
| Apgar score at 5 min, median (IQR) | 451 | 6 (4-9) | 148 | 6 (3-9) | 303 | 6 (4-9) | .185 |
| Assisted ventilation at 10 min of life (yes), n (%) | 461 | 216 (46.9) | 153 | 76 (49.7) | 308 | 140 (45.5) | .393 |
| Lowest cord pH, median (IQR) | 339 | 7.11 (6.93-7.23) | 104 | 7.10 (6.90-7.20) | 235 | 7.12 (6.94-7.24) | .182 |
| Therapeutic hypothermia (yes), n (%) | 472 | 234 (49.6) | 154 | 84 (54.5) | 318 | 150 (47.2) | .133 |
| Diagnosis before discharge, n (%) | 472 | 318 | <.001 |
| Mild HIE | 82 (17.4) | 0 (0) | 82 (26.0) | |
| Moderate HIE | 125 (26.5) | 50 (32.5) | 75 (23.6) | |
| Severe HIE | 56 (11.9) | 40 (26.0) | 16 (5.0) | |
| Stroke | 50 (10.5) | 26 (16.9) | 24 (7.5) | |
| Metabolic/genetic disorder | 43 (9.1) | 27 (17.5) | 16 (5.0) | |
| Suspected seizures, unconfirmed | 31 (6.6) | 0 (0) | 31 (9.7) | |
| Perinatal asphyxia without encephalopathy | 22 (4.7) | 0 (0) | 22 (7.0) | |
| Sepsis/meningitis | 20 (4.2) | 3 (2.0) | 17 (5.3) | |
| Intracranial hemorrhage | 10 (2.1) | 1 (0.6) | 9 (2.8) | |
| Other† | 33 (7.0) | 7 (4.5) | 26 (8.2) | |
| Age at start of EEG monitoring, h, median (IQR) | 472 | 14.5 (5.6-44.4) | 154 | 14.5 (5.7-41.0) | 318 | 14.5 (5.5-49.1) | .576 |
| Duration of EEG monitoring, h, median (IQR) | 472 | 65.7 (29.6-92.8) | 154 | 86.4 (52.2-102.6) | 318 | 47.6 (23.5-88.0) | <.001 |

*P values were from the Mann-Whitney U test for continuous data and the χ² test for categorical data unless indicated otherwise. P < .05 was considered to indicate statistical significance.
†n = 8 seizures of unknown origin; n = 7 transient metabolic deficit; n = 5 congenital brain malformation; n = 3 for each of postnatal cardiorespiratory arrest and respiratory distress; n = 2 for each of neonatal drug withdrawal syndrome and congenital cardiac anomaly; n = 1 for each of congenital anemia, meconium aspiration syndrome, tracheoesophageal atresia with cystic periventricular leukomalacia.