# **Original Article**



# Evaluating Levetiracetam Weight-Based Dosing in Benzodiazepine-Refractory Status Epilepticus

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**Abstract:** *Background:* Levetiracetam (LEV) is an antiseizure medication (ASM) used as a second line after benzodiazepines for status epilepticus treatment. Current literature lacks direct head-to-head comparisons between different LEV loading dose strategies, leading to uncertainty about superior dosing methods and thus clinical practice variations. *Methods:* A retrospective cohort study was designed to compare efficacy and safety of low (<30 mg/kg) versus high ( $\geq$ 30 mg/kg) weight-based LEV loading doses in adults with benzodiazepine-refractory status epilepticus (BRSE). The primary outcome of this study was termination of BRSE. No requirement for additional ASM after LEV was a surrogate for BRSE termination. Secondary endpoints included endotracheal intubation, intensive care unit (ICU) admission, 30-day all-cause mortality and adverse drug reactions. Statistical analysis included discrete and inferential statistics, including logistic regression and win-ratio analysis, to control for potential confounding variables. *Results:* Of the 106 patients included in this study, 54 (51%) did not require additional ASM after LEV, thereby achieving seizure termination. There was a higher proportion of patients with seizure termination in the higher weight-based dosing group as compared to the lower weight-based group (66% vs 40%, respectively; aOR 3.07; 95% CI: 1.36–7.21). There were lower rates for endotracheal intubation, ICU admission and all-cause mortality in the higher dosing group. Adverse events were comparable between the both groups. *Conclusion:* LEV's high weight-based loading dose strategy ( $\geq$ 30 mg/kg) is more effective in the termination of BRSE as compared to the lower weight-based loading dose strategy ( $\geq$ 30 mg/kg) is more effective in the termination of BRSE as compared to the lower weight-based loading dose strategy ( $\geq$ 30 mg/kg).

**RÉSUMÉ**: Évaluation du dosage de lévétiracétam selon le poids en ce qui regarde l'état de mal épileptique réfractaire aux benzodiazépines. Contexte : Après les benzodiazépines, le lévétiracétam est un médicament anticonvulsivant de deuxième intention (second line) pour le traitement de l'état de mal épileptique (EME). La littérature actuelle manque de comparaisons directes entre les différentes stratégies de dose de charge (loading dose) du lévétiracétam, ce qui entraîne une incertitude quant aux méthodes de dosage supérieures et donc des variations dans la pratique clinique. Méthodes : Une étude de cohorte rétrospective a été conçue pour comparer l'efficacité et l'innocuité de faibles doses de charge (< 30 mg/kg) de lévétiracétam par rapport à des doses élevées (≥ 30 mg/kg), et ce, selon le poids d'adultes présentant un EME réfractaire aux benzodiazépines. Le principal résultat de cette étude était l'arrêt d'un EME réfractaire aux benzodiazépines. Le fait de ne pas recourir à un traitement anticonvulsivant supplémentaire après l'administration de lévétiracétam s'est avéré un critère de substitution de la fin d'un EME réfractaire aux benzodiazépines. Les critères secondaires d'évaluation comprenaient l'intubation endotrachéale ; l'admission aux soins intensifs ; la mortalité, toutes causes confondues, au bout de 30 jours ; et finalement les effets indésirables des médicaments. L'analyse statistique comprenait des statistiques discrètes et inférentielles, y compris la régression logistique et l'analyse des rapports de gain, afin de contrôler les variables de confusion potentielles. Résultats : Sur les 106 patients inclus dans cette étude, 54 (51%) n'ont pas eu besoin d'un traitement anticonvulsivant supplémentaire après le lévétiracétam, ce qui a permis de mettre fin aux crises. La proportion de patients dont les crises ont pris fin était plus élevée dans le groupe où les doses de charge basées sur le poids étaient plus élevées en comparaison avec le groupe où ces mêmes doses étaient plus faibles (respectivement 66 % contre 40 % ; RC ajusté 3,07 ; IC 95 % : 1,36 - 7,21). À noter aussi que les taux d'intubation endotrachéale, d'admission aux soins intensifs et de mortalité, toutes causes confondues, étaient plus faibles dans le groupe ayant bénéficié de doses de charge plus élevées. Enfin, les événements indésirables se sont révélés comparables entre les deux groupes. Conclusion : Une stratégie de doses de charge élevées de lévétiracétam (≥ 30 mg/kg) est plus efficace pour mettre fin à un EME réfractaire aux benzodiazépines qu'une stratégie de doses de charge plus faibles (< 30 mg/kg).

Keywords: Antiseizure; antiepileptic; levetiracetam; seizure; refractory

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# Highlights

- Intravenous levetiracetam (LEV) is used as the second-line treatment option for status epilepticus, but there is still uncertainty regarding the optimal loading dose strategy.
- Weight-based dosing ≥30 mg/kg of LEV seems to result in more favorable outcomes.
- Future prospective studies are required to definitively guide optimal dosing.

#### Introduction

Status epilepticus (SE) is a critical neurological emergency where seizures persist and fail to terminate spontaneously. Prolonged seizures, lasting a minimum of 5 minutes, can lead to continuous seizure activity with potential for irreversible neuronal damage and fatality if not appropriately treated.<sup>1</sup> First-line treatment with benzodiazepines (BZD) is commonly administered to abort seizure activity. However, benzodiazepine-refractory SE (BRSE) can occur and is associated with mortality rates ranging from 35% to 60%.<sup>2</sup> As a result, an effective second-line antiseizure medication (ASM) such as levetiracetam (LEV) is essential in managing BRSE. A loading dose is recommended in emergency situations to critically achieve adequate seizure control.<sup>3</sup> It can be administered as a weight-based dose ranging from 20 to 60 mg/kg or a fixed dose of 1000 to 3000 mg.<sup>4</sup>

Both fixed and weight-based LEV dosing strategies are commonly used in clinical practice. The Neurocritical Care Society Guidelines recommend a fixed LEV dosing strategy for SE based on observational data.<sup>5,6</sup> A study by Eue *et al.* demonstrated a 44.2% response rate (primary outcome of no BRSE) with up to 2000 mg IV LEV loading dose.<sup>6</sup> However, international guidelines recommend weight-based loading dose in patients with BRSE based on the results from the ESETT trial, which showed a LEV loading dose of 60 mg/kg to a maximum of 4500 mg effectively controlled seizures in 47% of refractory SE cases with minimal adverse events.<sup>7,8,9</sup> A recent systematic review suggested that 20–60 mg/kg weight-based LEV dosing is efficacious in 46.9–81.8% of cases;<sup>10</sup> however, it lacks a direct head-to-head comparison of the safety and efficacy between different dosing strategies, leading to clinical ambiguity about the optimal LEV loading dose strategy for SE.

Two prior retrospective analyses have explored the efficacy and safety of different LEV loading doses in BRSE. In a 2023 study of 202 patients, a dosing strategy of 35 mg/kg was not associated with worse clinical outcomes.<sup>11</sup> Similarly, a 2024 analysis of 218 patients found no significant differences in seizure termination or recurrence rates among LEV dosing groups of <20 mg/kg, 21-39 mg/kg and ≥40 mg/kg.12 However, these studies yielded negative results, contributing to ongoing uncertainty about the optimal LEV loading dose strategy. In this retrospective cohort study, we aimed to address this clinical question by comparing rates of successful BRSE termination in patients receiving low (<30 mg/kg) versus high (≥30 mg/kg) LEV weight-based dosing. The decision to use 30 mg/kg as a threshold to distinguish low versus high LEV dosing was arbitrary and reflects commonly observed practices in clinical settings (i.e., a fixed 2000 mg loading dose for a 70 kg patient correlates to an approximately 30 mg/kg weight-based load). The low weight-based dosing threshold would thus ideally capture most common fixed LEV doses seen in clinical practice.5

# Methods

This was a single-center, retrospective cohort study of patients admitted to the emergency department of Vancouver General Hospital (VGH) between November 2022 and December 2023. The study included patients who were  $\geq 18$  years old with convulsive SE lasting at least 5 minutes despite receiving at least one dose of BZD. The use of electroencephalogram (EEG) monitoring was not considered as part of the study's inclusion criteria. No EEG monitoring was involved. Patients were excluded if they did not receive BZD prior to LEV, received loading doses of other parenteral ASM prior to LEV, were intubated or admitted under critical care services prior to SE event, were pregnant or had traumatic brain injuries or toxicological seizures due to overdoses or alcohol withdrawal. The primary outcome of this study was to compare successful BRSE termination between the two groups. The need for additional ASM after receiving a LEV load was used as a surrogate marker for BRSE termination. This surrogate endpoint was selected as the best available objective measure to confirm clinical resolution of BRSE. The secondary outcomes were the proportion of patients who required endotracheal intubation, intensive care unit (ICU) admission or experienced adverse drug reactions in each group. Also 30-day all-cause mortality was captured in both groups. Ethics and operational approval for this study were obtained from the University of British Columbia Clinical Research Ethics Board and Vancouver Coastal Health Research Institute, respectively.

### Data collection and assessment of outcomes

All patients presenting to the VGH Emergency Department between November 2022 and December 2023 who received intravenous LEV were screened for inclusion in the study. A total of 106 patients met the inclusion criteria. The electronic medical records of the included patients were reviewed for baseline demographics including age, weight, serum creatinine and estimated glomerular filtration rate (eGFR). Additionally, risk factors for seizures such as history and type of epilepsy, seizure etiology and electrolyte imbalances were collected for each patient. Electrolyte imbalances were defined for hyponatremia (serum sodium levels below 125 mmol/L), hypomagnesemia (serum magnesium levels below 0.8 mmol/L) and hypocalcemia (serum calcium levels below 1.9 mmol/L).<sup>13</sup> Finally, patient charts were reviewed for clinical outcome data including termination of BRSE, endotracheal intubation, ICU admission, 30day all-cause mortality and adverse drug events. The majority of data were collected from consultation notes and electronic administration records. Details such as dosage and route of BZD administered, LEV loading dose, drug and dose of additional ASM(s), documented reasons for death and documented adverse drug events were manually extracted from each health record. All data were stored in a secure database using Research Electronic Data Capture (REDCap).<sup>14</sup> LEV loading doses administered were dichotomized using a built-in REDCap calculator that divided the administered dose by the patient's documented weight, yielding a mg/kg dosage. This was further categorized as <30 mg/kg and  $\geq 30 \text{ mg/kg}$ .

# Sample size

The sample size was calculated based on data from the study by Eue *et al.*<sup>6</sup> and ESETT trial,<sup>9</sup> assuming a 1:1 recruitment ratio. With an expected moderate effect size of 0.5, power of 80% and an alpha of 0.05, a total of 314 patient encounters would be required.

#### **Statistical analysis**

Descriptive statistics were used to summarize patient characteristics. Normally distributed variables were reported as mean and

#### Table 1. Baseline characteristics

Characteristics	Total (N = 106)	≥30 mg/kg (n = 44)	<30 mg/kg (n = 62)	
Age – year (median, IQR)	63.5 (43,77)	66 (38,79)	62.5 (44,74)	
Weight – kg (mean, SD)	69.7 ± 16.8	62.6 ± 13.6	$74.6 \pm 17.1$	
Serum creatinine – µmol/L (median, IQR)	73.0 (58.8, 91.2)	70.0 (58.0, 88.3)	75.5 (64.5, 92.3)	
Estimated GFR – mL/min/1.73 m <sup>2</sup> (mean, SD)	83.2 ± 27.7	82.8 ± 27.2	84.3 ± 27.3	
History of epilepsy – no. (%)	38 (36%)	12 (27%)	26 (42%)	
Generalized	21 (55%)	6 (50%)	15 (57%)	
Focal	12 (32%)	4 (33%)	8 (31%)	
Generalized and focal	4 (10%)	2 (17%)	2 (8%)	
Non-convulsive	1 (3%)	0	1 (4%)	
Underlying etiology				
Structural – no. (%)	74 (70%)	29 (65%)	45 (72%)	
Infectious – no. (%)	5 (5%)	3 (7%)	2 (3%)	
Genetic – no. (%)	5 (5%)	2 (5%)	3 (5%)	
Biochemical – no. (%)	4 (4%)	2 (5%)	2 (3%)	
Other* – no. (%)	9 (8%)	3 (7%)	6 (10%)	
Unknown – no. (%)	9 (8%)	5 (11%)	4 (7%)	
Electrolytes				
Hyponatremia <sup>¶</sup> – no. (%)	2 (2%)	1 (2%)	1 (2%)	
Hypomagnesemia <sup>†</sup> – no. (%)	31 (29%)	16 (36%)	15 (24%)	
Hypocalcemia <sup>‡</sup> – no. (%)	4 (4%)	1 (2%)	3 (5%)	
Index seizure episode (median, IQR)				
Lorazepam initial dose (mg) (n = 84)	2.0 (1.0, 2.0)	2.0 (1.3, 2.0) (n = 34)	2.0 (1.0, 2.0) (n = 50)	
Midazolam initial dose (mg) (n = 22)	5.0 (2.2, 5.0)	5.0 (3.1, 5.0) (n = 10)	3.3 (1.8, 5.0) (n = 12)	
LEV dose (mg/kg)	28.0 (19.3, 38.0)	39.0 (36.0, 43.5)	20.5 (16.0, 26.0)	
LEV dose (mg)	2000 (1500, 2000)	2500 (2000, 3000)	1500 (1000, 2000)	

GFR = glomerular filtration rate; LEV = levetiracetam.

\*Medication noncompliance, metabolic.

¶ Na < 125;  $^{\dagger}$  Mg < 0.8;  $^{\ddagger}$  Ca < 1.9.

Table 2. Primary outcome (adjusted odds ratio\* for termination of BRSE)

	Total (N = 106)	≥30 mg/kg (n = 44)	<30 mg/kg (n = 62)	Adjusted odds ratio	95% confidence interval
Termination of BRSE	54 (51%)	29 (66%)	25 (40%)	3.07	1.36 - 7.21

BRSE = benzodiazepine-refractory status epilepticus.

\*Adjusted for age and estimated glomerular filtration rate.

standard deviation, and non-normally distributed variables were reported as median and interquartile range, based on the Shapiro-Wilk test. A multiple logistic regression was utilized to compare the proportion of patients with BRSE termination between the two dosing strategies. An adjusted odds ratio (aOR) with its 95% confidence interval for the association between type of LEV dose and the termination of BRSE was calculated. Age, history of

#### Table 3. Secondary endpoints

	Total (N = 106)	≥30 mg/kg (n = 44)	<30 mg/kg (n = 62)
Endotracheal intubation	20 (19%)	5 (11%)	15 (24%)
ICU admission	23 (22%)	7 (16%)	16 (26%)
30-day all-cause mortality	21 (20%)	5 (11%) <sup>†</sup>	16 (26%)*

ICU = intensive care unit.

 $^{\dagger}$ Documented reasons in electronic records: refractory SE (n = 2), hydrocephalus (n = 1), end-stage cancer (n = 1), unknown (n = 1).

\*Documented reasons in electronic records: end-stage cancer (n = 5), hemorrhagic stroke (n = 4), respiratory failure (n = 2), cognitive decline (n = 2), sepsis (n = 2), refractory SE (n = 1).

 Table 4.
 Safety outcomes

Adverse drug reaction	≥30 mg/kg (n = 44)	<30 mg/kg (n = 62)
Agitation	2 (5%)	3 (5%)
Aggressive behavior	1 (2%)	1 (2%)
Delirium	1 (2%)	0
Rash	1 (2%)	0
Sedation	0	1 (2%)

epilepsy and eGFR were considered as potential adjustment variables. Final adjustment variables were selected by performing a likelihood ratio test. A few missing eGFR variables were addressed and imputed with their median. The secondary outcomes were examined using descriptive statistics to describe exploratory trends and observations. In addition, patients were randomly split into two groups and conducted win-ratio analyses based on a composite outcome consisting in order of mortality, ICU admission, endotracheal intubation and BRSE termination. A win ratio was calculated for both the unstratified sample and groups stratified by age of 65 years old and seizure history.<sup>15</sup> All analyses were performed in R 4.1.2.<sup>16</sup>

#### Results

A total of 106 patients who presented to the VGH Emergency Department and were treated with LEV for BRSE met the inclusion criteria. Patient baseline characteristics and relevant BRSE risk factors are presented in Table 1.

The median age of patients was 63.5 years, with an average weight of about 70 kg. Thirty-eight patients (36%) had a history of epilepsy, and the seizure etiology was mainly structural (70%). Patients were initially treated with lorazepam (79%) or midazolam (21%), with median initial doses presented in Table 1. BZD was given either intravenously (90%), intramuscularly (7%) or sublingually (3%). The median LEV loading dose was 28.0 mg/kg (or median dose of 2000 mg).

Out of 106 patients, 54 (51%) had termination of BRSE, not requiring additional ASM (Table 2). There was a larger proportion of patients with termination of BRSE in the higher ( $\geq$ 30 mg/kg) compared to the lower (<30 mg/kg) weight-based dosing group (66% vs 40%, respectively). The unadjusted OR was 2.86 (95% CI: 1.30–6.52). The significance of potential adjustment variables was assessed, and age and eGFR were included in the final model. eGFR was identified as a variable influencing patient response to each loading dose strategy (specifically, patients with better renal function were more likely to respond favorably to LEV). The aOR was 3.07 (95% CI: 1.36–7.2). There were lower rates for endotracheal intubation, ICU admission and all-cause mortality in the higher weight-based dosing group (Table 3). There was a total of 21 deaths (20%) recorded in this study, with higher rates observed in the lower weight-based dosing group. Among the reported causes, two patients in the higher weight-based dosing group died due to refractory SE, compared to one patient in the lower weight-based dosing group. The majority of these deaths were non-SE-related (Table 3). LEV was well tolerated in both groups, with adverse drug reactions being comparable between both dosing strategies (Table 4).

Win ratio analysis revealed differences between higher and lower weight-based dosing strategies. In the overall analysis, the win ratio was 2.78 (95% CI: 2.70–2.86), reflecting a greater likelihood of favorable outcomes with higher weight-based dosing. Stratified analyses showed consistent patterns. When stratified by age ( $\geq$ 65 years), the win ratio was 1.91 (95% CI: 1.85–1.97), and when stratified by seizure history, the win ratio was 1.92 (95% CI: 1.86–1.97).

# Discussion

In this retrospective cohort study, the use of a LEV loading dose of  $\geq$  30 mg/kg was more effective in terminating BRSE compared to the lower weight-based loading dose of <30 mg/kg. Despite existing studies with negative results, our study remains the first to demonstrate a positive association between a specific LEV dosing threshold ( $\geq$ 30 mg/kg) and improved clinical outcomes in BRSE. Notably, higher weight-based loading doses were linked to increased rates of seizure termination, lower rates of endotracheal intubation, ICU admission and all-cause mortality, while maintaining a tolerable safety profile. In contrast to the studies by Schowe et al. and Kuffer et al.,<sup>11,12</sup> our study employed a different statistical approach, utilizing regression analysis to calculate aORs for the primary outcome and a win-ratio analysis for hierarchical comparisons of the primary outcome and select secondary outcomes (Figure S1). These methodological differences may account for the discrepancy in findings and further highlight the importance of dosing strategies in optimizing BRSE management.

The observed association between higher weight-based LEV loading doses and BRSE termination is consistent with findings from other contemporary trials such as ESETT, where clinical termination of SE was achieved with high weight-based LEV dosing (60 mg/kg to a maximum of 4500 mg). Given linear kinetics, low protein binding (<10%) and relatively high volume of distribution (0.5–0.7 L/kg in adults),<sup>17</sup> the pharmacokinetic profile of LEV supports the use of higher initial doses to rapidly achieve therapeutic serum levels for effective acute seizure control. Current guidelines offer reference ranges for LEV trough level concentrations, but there is a lack of evidence to associate specific levels with seizure control.<sup>18,19</sup> A recent meta-regression and pharmacokinetic modeling analysis suggests that a weight-based loading dose of 40 mg/kg, up to 4500 mg, may achieve optimal LEV serum concentrations in refractory SE.<sup>20</sup> This underscores the importance of weight-based dosing to achieve adequate drug concentrations after administration. This approach may mitigate the risk of underdosing, which could necessitate additional ASM to manage refractory SE, as observed in this study with the lower weightbased dosing group.

While the median LEV weight-based loading dose in this study population was 28.0 mg/kg (or median dose of 2000 mg), the median weight-based loading dose was 20.5 mg/kg (or median dose of 1500 mg) in the lower dosing group and 39.0 mg/kg (or median dose of 2500 mg) in the higher dosing group. The maximum loading dose given in this study did not exceed the recommended maximum dose of 4500 mg (maximum weightbased loading dose observed was 62.5 mg/kg (or dose of 4300 mg)). Based on existing research and observational findings from this study, it can be suggested that initiating LEV treatment with a loading dose of at least 30 mg/kg, up to 60 mg/kg with a maximum of 4500 mg, might be an effective practice approach for improving BRSE management outcomes. The decision to adopt this threshold would require replication in research to further validate and establish optimal LEV weight-based dosing guidelines for managing BRSE effectively.

Lower rates for endotracheal intubation and ICU admission in the higher weight-based dosing group suggest that adequate loading doses of LEV are crucial for managing seizures effectively, as evidenced by the results of our win-ratio analyses. By achieving therapeutic concentrations promptly, LEV can terminate seizures and prevent invasive interventions such as intubation and admission to the ICU. Overall, there were more non-SE-related deaths compared to SE-related deaths in this patient population (Table 3). This study was not powered to compare mortality rates between the two dosing strategies; however, the persistent benefit seen in the win-ratio analysis showcases the benefits in other clinically important outcomes seen with higher weight-based LEV dosing. The adverse drug reactions observed in this study were consistent with the limited side effects reported in previous LEV studies, supporting its overall favorable safety profile.

Lorazepam was the most commonly administered BZD in both dosing groups, with a recommended dose of 4 mg for initial treatment of SE. The median initial lorazepam dose in both groups was subtherapeutic, although it was relatively closer to the recommended dose in the lower weight-based LEV dosing group, suggesting that despite a conventionally recommended BZD dose, the lower weight-based LEV dose group fared worse. The observation that patients received various initial doses of BZD may reflect clinical practice variability or individual patient factors such as the severity of seizure presentation. This variability limits the assessment of adequate dosing with BZD. While optimal BZD dosing should ideally be confirmed before considering LEV, the results of this study highlight the importance of a systematic approach to medication dosing in SE and underscore the equal importance of adequate LEV dosing in managing BRSE to ensure optimal patient outcomes.

There are several limitations to this study. First, the retrospective design limits the ability to control for unknown confounders and the resultant selection bias, although efforts were made to correct for this limitation by performing a regression analysis. Second, future studies with larger sample sizes would be beneficial to confirm these results and ensure that the observed effects are robust and applicable to a broader population. Third, data collection was limited to what was available and documented in the electronic health record. Given the lack of objective data to confirm clinical resolution of BRSE, additional ASM after LEV load was used as a surrogate endpoint for BRSE termination. There may be a possibility that patients who did not return to neurological baseline (such as patients with non-convulsive SE) were missed and not captured in the study.

# Conclusion

The study demonstrated that patients who received  $\geq$  30mg/kg of LEV as a loading dose had greater odds of BRSE termination compared to

those who received <30mg/kg. It also showed lower rates for clinically important outcomes of endotracheal intubation, ICU admission and 30-day all-cause mortality in the higher LEV dosing group. Dosing guidance provided by this study may help improve successful seizure control and patient outcomes in clinical practice. Prospective studies are warranted to further explore the role of higher weight-based dosing and treatment of BRSE in this clinical setting.

**Supplementary material.** The supplementary material for this article can be found at https://doi.org/10.1017/cjn.2025.56.

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#### References

- Trinka E, Cock H, Hesdorffer D, et al. A definition and classification of status epilepticus-report of the ILAE task force on classification of status epilepticus. *Epilepsia*. 2015;56:1515–1523. DOI: 10.1111/epi.13121.
- Wylie T, Sandhu DS, Murr N. Status epilepticus, *StatPearls*. Treasure Island (FL): StatPearls Publishing; 2023.
- Wheless JW, Clarke D, Hovinga CA, et al. Rapid infusion of a loading dose of intravenous levetiracetam with minimal dilution: a safety study. J Child Neurol. 2009;24:946–951. DOI: 10.1177/0883073808331351.
- Brophy GM, Bell R, Claassen J, et al. Guidelines for the evaluation and management of status epilepticus. *Neurocrit Care*. 2012;17:3–23. DOI: 10. 1007/s12028-012-9695-z.
- Uges JW, van Huizen MD, Engelsman J, et al. Safety and pharmacokinetics of intravenous levetiracetam infusion as add-on in status epilepticus. *Epilepsia*. 2009;50:415–421. DOI: 10.1111/j.1528-1167.2008.01889.x.
- Eue S, Grumbt M, Müller M, Schulze A. Two years of experience in the treatment of status epilepticus with intravenous levetiracetam. *Epilepsy Behav.* 2009;15:467–469. DOI: 10.1016/j.yebeh.2009.05.020.
- 7. Vossler DG, Bainbridge JL, Boggs JG, et al. Treatment of refractory convulsive status epilepticus: a comprehensive review by the American

epilepsy society treatments committee. *Epilepsy Curr*. 2020;20:245–264. DOI: 10.1177/1535759720928269.

- Glauser T, Shinnar S, Gloss D, et al. Evidence-based guideline: treatment of convulsive status epilepticus in children and adults: report of the guideline committee of the American Epilepsy Society. *Epilepsy Curr.* 2016;16:48–61. DOI: 10.5698/1535-7597-16.1.48.
- Chamberlain JM, Kapur J, Shinnar S, et al. Efficacy of levetiracetam, fosphenytoin, and valproate for established status epilepticus by age group (ESETT): a double-blind, responsive-adaptive, randomised controlled trial [published correction appears in Lancet. [published correction appears in Lancet. 2023 May 6;401(10387):1498]. Lancet. 2020;395(10231):1217-1224. DOI: 10.1016/S0140-6736(20)30611-5020.
- Webb CA, Wanbon R, Otto ED. Levetiracetam for status epilepticus in adults: a systematic review. *Can J Hosp Pharm.* 2022;75:46–53. DOI: 10.4212/cjhp.v75i1.3254.
- Kuffer I, Novy J, Rossetti AO. Status epilepticus prognosis following levetiracetam administration: analysis of loading doses. *Eur J Neurol.* 2023;30:1957–1962. DOI: 10.1111/ene.15791. Epub 2023 Apr 2.
- Schowe C, Frick CD, Weitkamp LU, Jarboe L. Evaluation of levetiracetam loading dose in adult patients with benzodiazepine-refractory status epilepticus. *Am J Emerg Med.* 2024;85:148–152. DOI: 10.1016/j.ajem. 2024.09.007.
- Nardone R, Brigo F, Trinka E. Acute symptomatic seizures caused by electrolyte disturbances. J Clin Neurol. 2016;12:21–33. DOI: 10.3988/jcn. 2016.12.1.21. PMID: 26754778; PMCID: PMC4712283.
- Harris PA, Taylor R, Thielke R, Payne J, Gonzalez N, Conde JG. Research electronic data capture (REDCap)-a metadata-driven methodology and workflow process for providing translational research informatics support. J Biomed Inform. 2009;42:377–381. DOI: 10.1016/j.jbi.2008. 08.010.
- Pocock SJ, Ariti CA, Collier TJ, Wang D. The win ratio: a new approach to the analysis of composite endpoints in clinical trials based on clinical priorities. *Eur Heart J.* 2012;33:176–182. DOI: 10.1093/eurheartj/ehr352.
- 16. R Core Team. R: a language and environment for statistical computing [Computer software]. Vienna: R Foundation for Statistical Computing; 2021.
- Wright C, Downing J, Mungall D, et al. Clinical pharmacology and pharmacokinetics of levetiracetam. *Front Neurol.* 2013;4:192. DOI: 10.3389/ fneur.2013.00192.
- Patsalos PN, Berry DJ, Bourgeois BF, et al. Antiepileptic drugs-best practice guidelines for therapeutic drug monitoring: a position paper by the subcommission on therapeutic drug monitoring ILAE commission on therapeutic strategies. *Epilepsia*. 2008;49:1239–1276. DOI: 10.1111/j.1528-1167.2008.01561.x.
- Hiemke C, Bergemann N, Clement H, et al. Consensus guidelines for therapeutic drug monitoring in neuropsychopharmacology: update 2017. *Pharmacopsychiatry*. 2018;51:9–62. DOI: 10.1055/s-0037-1600991.
- Lau A, Haag H, Maharaj A. A simulation-based assessment of levetiracetam concentrations following fixed and weight-based loading doses: a metaregression and pharmacokinetic modeling analysis. J Clin Pharmacol. 2024;64:1173–1180. DOI: 10.1002/jcph.2449. Epub ahead of print.