Efficacy, Tolerability, and Safety of Concentrated Intranasal Midazolam Spray as Emergency Medication in Epilepsy Patients During Video-EEG Monitoring

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Abstract

Background An efficient, well tolerated, and safe emergency treatment with a rapid onset of action is needed to prevent seizure clusters and to terminate prolonged seizures and status epilepticus.

Objectives This study aimed to examine the efficacy, tolerability, and safety of intranasal midazolam (in-MDZ) spray in clinical practice.

Methods In this retrospective, multicenter observational study, we evaluated all patients with peri-ictal application of in-MDZ during video-EEG monitoring at the epilepsy centers in Frankfurt and Marburg between 2014 and 2017. For every patient, we analyzed the recurrence of any seizure or generalized tonic–clonic seizures after index seizures with and without in-MDZ administration. Treatment-emergent adverse events (TEAEs) were also evaluated.

Results In-MDZ was used in 243 patients with epilepsy (mean age 35.5 years; range 5–76 years; 46.5% female) for treatment of 459 seizures. A median dose of in-MDZ 5 mg (i.e., two puffs; range 2.5–15 mg) was administered within a median time from EEG seizure onset until in-MDZ application of 1.18 min [interquartile range (IQR) 1.27], while median time from clinical seizure onset until in-MDZ administration was 1.08 min (IQR 1.19). In-MDZ was given within 1 min after EEG seizure onset in 171 seizures. An intraindividual comparison of seizures with and without application of in-MDZ was feasible in 171 patients, demonstrating that in-MDZ reduced the occurrence of any (Cox proportional-hazard model \( p < 0.001 \)) and generalized tonic–clonic seizure (Cox proportional-hazard model \( p = 0.0167 \)) over a period of 24 h. The seizure-free timespan was doubled from a median of 5.0 h in controls to a median of 10.67 h after in-MDZ administration. We additionally clustered in-MDZ administrations for the 119 patients who received in-MDZ more than once, comparing them with the index cases without in-MDZ. Even when considering subsequent seizures with in-MDZ administration, a patient receiving in-MDZ is still half as likely to incur another seizure in the upcoming 24 h as compared with when the same patient does not receive in-MDZ (hazard ratio 0.50; 95% CI 0.42–0.60; \( p < 0.01 \)). In-MDZ was well tolerated without major adverse events. The most common side effects were irritation of the nasal mucosa [37 cases (8.1%)], prolonged sedation [26 cases (5.7%)], and nausea and vomiting [12 cases (2.6%)]. A decline in oxygen saturation was measured after 78 seizures (17%).

Conclusion We conclude that in-MDZ is a safe and efficient treatment option to prevent short-term recurrence of seizures. In-MDZ can be administered very quickly by trained staff within 1–2 min after seizure onset. No major cardiocirculatory or respiratory adverse events were observed.

1 Introduction

The emergency treatment of acute seizures is required to be rapid and efficient, as ongoing epileptic activity results in increased morbidity and mortality [1]. The ideal anticonvulsant agent is safe, easily administered, and has a long-lasting antiseizure effect [2]. Rescue medication should prevent seizure recurrence as well as the occurrence of prolonged seizures or status epilepticus [3–5]. Benzodiazepines such as lorazepam, diazepam, and midazolam are established first-line drugs for acute seizures [6], with available delivery routes that include intravenous, rectal, intramuscular, buccal, and intranasal [5–11]. Difficulties with achieving intravenous access may lead to a delay in drug...
Key Points

Use of intranasal midazolam spray was analyzed in a large cohort (n = 243; 459 seizures) of patients with epilepsy.

In general, intranasal midazolam spray reduced the occurrence of any (Cox proportional-hazard model \( p < 0.001 \)) and generalized tonic–clonic seizure (\( p = 0.0167 \)) over a period of 24 h. The seizure-free timespan was doubled from a median of 5.0 h in controls to a median of 10.67 h after intranasal midazolam administration.

Intranasal midazolam spray can be administered easily and quickly after seizure onset by hospital staff. We did not observe any clinically relevant treatment-emergent effects.

administration, rendering the development of suitable non-parenteral routes vital, as responsiveness to benzodiazepines is time-dependent [12]. Jaw clenching [9], hypersalivation, and uncontrollable swallowing constitute some limitations inherent to the buccal route, making it difficult to minimize variability in pharmacodynamics [13], which might also prove problematic for intramuscular injections [8]. Rectal administration is becoming less popular due to the social distress and sense of shame it imposes on both patients and caregivers [14].

In comparison, intranasal midazolam (in-MDZ) may be a more favorable option as it can be administered in a significantly shorter interval without the need for an intravenous route [15] and has been shown to be reliable and efficient [8, 14–19]. Furthermore, as compared with rectal diazepam, it is at least as effective in its anticonvulsive action [14, 20, 21] and is preferred by caregivers [20, 22]. Additionally, in-MDZ can be easily administered, rendering drug delivery more convenient in cases where patients are cognitively impaired and gaining their cooperation is more difficult [8]. However, study populations have so far been small and data on adult patients remain relatively scarce [23, 24].

The aim of this study was to ascertain the efficacy, tolerability, and safety of in-MDZ during video-EEG monitoring.

2 Methods

In this retrospective, multicenter observational study, we examined the medical records of all patients with an epilepsy diagnosis who received in-MDZ during video-EEG monitoring between September 2014 and August 2017 at the university hospitals in Marburg and Frankfurt. Patients were admitted for presurgical evaluation or evaluation of their epilepsy syndrome, both usually scheduled for 120 h and requiring withdrawal of antiepileptic drugs. In both units, each patient is prescribed in-MDZ as a pro re nata emergency medication in case of frequent or prolonged seizures or generalized convulsive seizures; a dedicated emergency protocol is not utilized. Patients treated with in-MDZ between 2008 and 2014 who were previously analyzed and reported in a pilot study by Kay et al. [17] were not included.

All patients who received in-MDZ were identifiable through the narcotic drug documentation, which is mandatory for the use of concentrated midazolam nasal spray under Article III of German Narcotic Law. This is due to the fact that the midazolam content in one nasal spray device exceeds 15 mg. We included all patients with epilepsy and administration of in-MDZ. Patients in whom in-MDZ was administered for reasons other than seizure treatment (e.g., periprocedural sedation), or erroneously for treatment for psychogenic non-epileptic seizures, were excluded from analysis. The analysis was approved by the ethics committee of the Goethe-University Frankfurt and adhered to the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement [27]. The informed consent was waived by the ethics committees because of the retrospective nature of the study. This study was not sponsored or funded by any company.

Briefly, EEG, electrocardiogram (ECG), and video data were reviewed in patients who underwent video-EEG monitoring while in-MDZ was administered. Patients were monitored with scalp electrodes according to the 10–20 international system [25]. Sphenoidal electrodes or intracranial depth or subdural electrodes were placed in selected patients when clinically indicated. Heart rate analysis was based on the coregistered ECG tracing [29]. If available, the saturation of peripheral oxygen was measured interictally and immediately after a seizure using a pulse oximeter.

The classification of seizure types and epilepsies was performed based on the latest definitions proposed by the International League against Epilepsy (ILAE) [26, 28]. A standardized questionnaire was used to gather information concerning etiology, concomitant diseases, frequency of seizures, anticonvulsant treatment, and, if applicable, its discontinuation during the video-EEG monitoring.

EEG seizure onset and EEG seizure endpoint, and timing of in-MDZ treatment as well as seizure type were recorded. The following 24 h were examined for the occurrence of seizures. For every administration of midazolam, treatment-emergent adverse events (TEAEs) were recorded from the medical reports and the annotations to the video and EEG data if applicable, including a deterioration in oxygenation, nasal irritation, headache, cough, nausea and vomiting, and prolonged sedation. As proposed by Bancke et al. [30], we defined a drop of peripheral oxygen saturation below 90% to be a TEAE. Measurements of peripheral oxygen saturation were available in 310 out of 459 in-MDZ applications.

The concentrated midazolam spray was prepared by the central pharmacies of the involved university hospitals in Marburg.
and Frankfurt, based on a formula that was previously published [17, 31]. The clear and colorless solution contained midazolam hydrochloride buffered to a pH of 3.3 with a mixture of water and 1 N of hydrochloride acid. As an antimicrobial preservative, we used benzalkonium chloride and sodium ethylenediaminetetraacetic acid. Stored at room temperature and under light protection, the spray lasts up to 3 months. Medical personnel applying the nasal spray were trained to deliver the drug, ensuring standardized administration. To ensure equal dosage, nasal spray devices (Zscheile & Klinger, Hamburg, Germany) were used and a dose of 2.5 mg of midazolam per puff (140 µL) was administered.

Intraindividual comparisons comparing index seizures with and without midazolam were performed to examine the potential effects of in-MDZ on increasing the seizure-free interval within the subsequent 24 h and the semiology of the upcoming seizure. For each patient, the timespan between seizures both with and without midazolam application and the respective following seizures were compared. To prevent potential bias, the first seizure with midazolam administration was defined as the in-MDZ index seizure and compared with the control index seizure without in-MDZ. The control index seizure was either the first seizure of the patient recorded via monitoring or, if no seizure prior to midazolam application was available, the first seizure after midazolam application without in-MDZ. If a patient received midazolam during more than one seizure, every application was recorded and assessed for tolerability and safety. Where patients received in-MDZ on several occasions, we additionally clustered in-MDZ administrations for every patient and compared them with the patients’ respective index seizure without in-MDZ.

Statistic evaluations were executed using the SPSS version 25.0 software program (IBM Corp., Armonk, NY, USA) and R (R Foundation for Statistical Computing, Vienna, Austria). McNemar’s test was used to determine if there are differences between the paired in-MDZ and control groups. The seizure-free interval and heart rate were non-normally distributed and groups were compared using a Wilcoxon paired sample test. Seizure recurrence was depicted in survival curves and test comparisons from a Cox proportional-hazard model for both the in-MDZ and control groups. Seizure recurrence was normally distributed and groups were compared using a Wilcoxon paired sample test. Seizure recurrence was depicted in survival curves and test comparisons from a Cox proportional-hazard model for both the in-MDZ and control groups. Seizure recurrence was normally distributed and groups were compared using a Wilcoxon paired sample test.

### 3 Results

#### 3.1 Patient Characteristics

In total, 243 patients [113 female (46.5%)] received in-MDZ during the study period with sufficient EEG and video data for analysis (including 167 in Frankfurt and 76 in Marburg). The mean ± standard deviation (SD) age was 35.5 ± 14.5 years [range 5–76 years, 19 patients below the age of 18 years (7.8%)], with a mean ± SD age of epilepsy onset of 18.1 ± 14.7 years (range 0–64 years). The overall mean ± SD duration of epilepsy to the time of admission was 16.9 ± 13.3 years (range 0–60 years). Patients reported a mean ± SD of 19.4 ± 99 seizures per month (median 4 seizures; range 0–1500 seizures), including 17.5 ± 98.7 focal seizures (median 2 seizures; range 0–1500 seizures) and 1.2 ± 3.9 generalized tonic–clonic seizures (GTCS) [median 0.1 seizures; range 0–30 seizures; 197 (81%) patients had GTCS in their lifetime].

Patients were found to be taking a mean of 2.2 ± 0.9 antiseizure drugs (ASDs) (range 0–4), with 50 (20.6%) of them taking monotherapy, 190 (78.2%) taking polytherapy, and three (1.2%) being currently without any medication. The patients had failed a mean ± SD of 2.6 ± 2.5 ASDs in the past (median 2 ASDs; range 0–15 ASDs; current medications not included). In total, 224 patients (92.2%) had drug-resistant epilepsy according to the definition of the ILAE [32]. Fourteen (5.8%) patients had undergone neurosurgery for the implantation of electrodes prior to the monitoring.

The most prevalent epilepsy type was focal (91.0%), followed by generalized epilepsy (7.0%), while the epilepsy syndrome remained unknown in 2.1%. A total of 141 patients (58.0%) had additional medical conditions and, in 40 cases (16.5%), neurological impairments were present. For a more detailed description of patient characteristics, see Table 1.

#### 3.2 Characteristics, Tolerability, and Safety of in-MDZ Application During Video-EEG Monitoring

Video-EEG monitoring was performed for a mean ± SD of 113.9 ± 34.1 h (median 107; range 47–254). In the majority (n = 232; 95.5%) of patients, antiseizure medication was reduced or discontinued during video-EEG monitoring, while, in nine patients (3.7%), medication use was unchanged and two patients (0.8%) did not receive ASD treatment. Overall, midazolam was used 459 times (cases), with 119 patients receiving in-MDZ more than once. Patients received in-MDZ during or after a GTCS in 184 cases (40.1%), 261 times due to focal seizures (56.9%), and in 14 cases (3%) in-MDZ was given due to unstable EEG patterns. A median dose of in-MDZ 5 mg (i.e., two puffs; SD 1.2 mg; range 2.5–15 mg) was administered; the median was lower (3.75 mg) in those aged 12 years and younger, and 5 mg in the elderly aged 60 years and older.

The median time from EEG seizure onset until in-MDZ application was 1.18 min [interquartile range (IQR) 1.27; range 0.1–52.4], while median time from clinical seizure onset until in-MDZ administration was 1.08 min (IQR 1.19);
range – 0.25 to 52.3); in one case, in-MDZ was administered before clinical seizure onset. In-MDZ was given within 1 min after EEG seizure onset in 171 cases.

We observed some kind of TEAE in 108 (44.4%) patients and 136 cases (29.6%). TEAEs included irritation of the nasal mucosa (n = 32 patients, 13.2%; 37 cases, 8.1%), headache (n = 4 patients, 1.6%; 4 cases, 0.9%), cough (n = 8 patients, 3.3%; 9 cases, 2.0%), nausea and vomiting (n = 10 patients, 4.1%; 12 cases, 2.6%), and prolonged sedation (n = 22 patients, 9.1%; 26 cases, 5.7%). One patient had a nosebleed following several attempts to administer the drug despite excessive turning of the head.

Heart rate analysis showed a comparable increase of heart rate 10 min after GTCS (29.3 bpm in in-MDZ; 25.8 bpm in controls) and 10 min after seizures without generalization (5.3 bpm in in-MDZ, 4.0 bpm in controls) with and without in-MDZ.

In 310 seizures, the measurement of oxygen saturation was feasible. Oxygen levels were measured directly after seizure end, which occurred a median of 3.43 min after application of in-MDZ. The mean ± SD oxygen saturation immediately postictally being 92.3% ± 6% (range 51–99). There was no intubation necessary in any patient. A drop in peripheral oxygen saturation below 90% occurred 78 times (17%) in 67 patients (27.6%). Such a desaturation below 90% was observed 36 times after seizures without generalization and in 42 GTCS. Among the latter, one patient had respiratory insufficiency with a minimum oxygen concentration of 51% after having suffered a GTCS. Oxygenation rose above 90% within 42 s under administration of oxygen.

3.3 Difference in Number of Recurring Seizures with and without Administration of in-MDZ

To evaluate the difference in number of recurring seizures for a postictal period of 24 h, we compared pairs of seizures in each single patient, with and without administration of in-MDZ. Overall, 171 pairs of seizures with in-MDZ application and seizures without treatment were suitable for statistic comparison. Regarding generalization, we observed a higher proportion of GTCS for index seizures treated with in-MDZ (60/171; 35.1%) than for controls (16/171; 9.4%; p < 0.001). In 72 patients, there was no index seizure without in-MDZ available because the patients either received in-MDZ in every seizure (n = 69 patients; 28.4%); were treated with in-MDZ due to unstable EEG patterns (n = 2 patients; 0.8%); or the EEG pattern could not be clearly distinguished in the index seizure without in-MDZ (n = 1). These patients were included in the study for tolerability assessment only.

Among 116 patients, the index seizure without in-MDZ took place a median of 17.28 h before the in-MDZ seizure (IQR 25.34; range 0.15–209.12), whereas in 52 patients, the index seizure without in-MDZ took place a median of

| Table 1 Clinical characteristics of patients treated with intranasal midazolam (in-MDZ) |
|---------------------------------------------------------------|
| Patients treated with in-MDZ (n = 243) % (n) |                     |
| Age in years (mean ± SD) | 35.5 ± 14.5          |
| Age at epilepsy onset (mean ± SD) | 18.1 ± 14.7          |
| Epilepsy duration in years (mean ± SD) | 16.9 ± 13.3          |
| Epilepsy type |                     |
| Focal epilepsy | 86.4 (210)          |
| Right hemisphere | 34.2 (83)          |
| Left hemisphere | 39.9 (97)            |
| Bilateral | 12.3 (30)            |
| Genetic (idiopathic) generalized epilepsy | 6.2 (15) |
| Unknown | 7.4 (18)             |
| History of status epilepticus |                     |
| Yes | 4.1 (10)             |
| No | 95.9 (233)           |
| History of febrile convulsions |                     |
| Yes | 9.9 (24)             |
| No | 84.8 (206)           |
| Unknown | 5.3 (13)          |
| Imaging findings |                     |
| No appreciable disease | 37.2 (96) |
| Cavernoma | 3.5 (9)              |
| Cortical dysplasia | 14.0 (36)          |
| Hippocampal sclerosis | 22.1 (57)      |
| Tumor | 9.7 (25)            |
| Other | 7.0 (18)            |
| No imaging available | 3.5 (9)       |
| Previous neurosurgery | 3.1 (8)    |
| Comorbidities (n = 141) |                     |
| Arterial hypertension | 15.6 (22)    |
| Diabetes | 6.4 (9)            |
| Adipositas | 5.7 (8)            |
| Depression | 19.1 (27)        |
| Psychiatric conditions | 11.3 (16)  |
| Endocrine disease | 9.9 (14)         |
| Abuse of alcohol | 4.3 (6)       |
| Abuse of nicotine | 17.7 (25)       |
| Other | 9.9 (14)            |
| Antiseizure drugs |                     |
| Levetiracetam | 60.9 (148)      |
| Lamotrigine | 42.8 (104)        |
| Lacosamide | 18.5 (45)         |
| Oxcarbazepine | 17.3 (42)        |
| Zonisamide | 16.0 (39)         |
| Valproate | 12.8 (31)         |
| Carbamazepine | 10.7 (26)       |
| Topiramate | 7.4 (18)          |
| Other | 30.0 (73)           |
Intranasal Midazolam Spray During Video-EEG Monitoring

9.03 h after the in-MDZ index seizure (IQR 16.85; range 0.69–143.95) ($p < 0.001$). The seizure-free interval as per intention-to-treat analysis is depicted in a survival curve in Fig. 1 ($p < 0.001$). Further, the median timespan until the occurrence of the next seizure was 10.67 h [range 0.03–24, 95% confidence interval (CI) 7.28–14.77] after the application of in-MDZ, which was a significantly longer period than that after the index seizure without in-MDZ (5.00 h; range 0.02–24 h, 95% CI 4.12–6.55).

In-MDZ had a significant effect on seizure recurrence within the upcoming 24 h. Of the total cohort, 93.6% of patients remained seizure-free for 60 min following in-MDZ (vs 85.4% without in-MDZ), 60.8% remained seizure-free for 6 h (vs 43.9% without in-MDZ), and 29.2% did not experience another seizure in the following 24 h (vs 14% without in-MDZ).

The rate of GTCS in the following 24-hour interval was lower in the in-MDZ group than that in the control group for the same time period (see Table 2). The recurrence of GTCS within 24 h is depicted as a survival curve as per intention-to-treat analysis in Fig. 2 ($p = 0.0167$).

To increase the validity of our findings, we additionally clustered in-MDZ administrations for the 119 patients who received in-MDZ more than once, comparing them to the index cases without in-MDZ. We found that, even when considering subsequent seizures with in-MDZ administration, a patient receiving in-MDZ is still half as likely to incur another seizure in the upcoming 24 h as compared with when the same patient does not receive in-MDZ (hazard ratio 0.50; 95% CI 0.42–0.60). This finding was statistically significant with $p < 0.01$.

### 4 Discussion

Our study indicates that in-MDZ can be used as an effective anticonvulsant agent in emergency seizure treatment for epilepsy patients, in that it reduces seizure recurrence within the upcoming 24 h and does not cause any serious adverse events.

Previous research in children and smaller cohorts has already suggested in-MDZ to be a recommendable alternative to rectal diazepam [18, 33], intravenous diazepam [24], and intravenous lorazepam both in terms of comfort of drug delivery and effect. In their comparison of in-MDZ with intravenous diazepam, Lahat et al. [24] showed intravenous diazepam to have a more rapid effect, but the time until drug delivery was longer and thus in-MDZ was more favorable in terms of time until seizure cessation. Fışgin et al. [16] concluded that intranasal midazolam was more effective than rectal diazepam in terminating seizures, ending 87%

### Table 2 Occurrence of any seizure and of generalized tonic–clonic seizure (GTCS) in patients with and without application of intranasal midazolam (in-MDZ)

| Time (h) | Index seizure with in-MDZ treatment | Index seizure without in-MDZ treatment | $p$ value* |
|---------|------------------------------------|--------------------------------------|------------|
|         | No seizure recurrence (no. of patients) | Seizure recurrence (no. of patients) | Patients with GTCS | No seizure recurrence (no. of patients) | Seizure recurrence (no. of patients) | Patients with GTCS |
| 1       | 160 | 11 | 0 | 146 | 25 | 3 | < 0.001 |
| 2       | 143 | 28 | 1 | 121 | 50 | 6 | < 0.001 |
| 3       | 134 | 37 | 2 | 110 | 61 | 8 | < 0.001 |
| 4       | 126 | 45 | 3 | 91 | 80 | 11 | 0.002 |
| 5       | 115 | 56 | 5 | 82 | 89 | 14 | 0.080 |
| 6       | 104 | 67 | 8 | 75 | 96 | 16 | 0.621 |
| 9       | 90 | 81 | 10 | 64 | 107 | 19 | 0.254 |
| 12      | 80 | 91 | 10 | 52 | 119 | 21 | 0.007 |
| 24      | 50 | 121 | 18 | 24 | 147 | 25 | < 0.001 |

*McNemar’s test
As opposed to 60% of seizures within 10 min (p < 0.05). With special consideration of seizure prevention, Kay et al. [17] examined the medical records of 75 patients with an overall 110 applications of in-MDZ. Comparing 63 pairs of seizures intra-individually with and without in-MDZ, they found seizure recurrence to have been successfully prevented by in-MDZ and the seizure-free time span to have been increased significantly to a median of 5.8 h as compared with 2.4 h in patients without in-MDZ.

Our findings are supported by Detyniecki et al. [34], who conducted an open-label extension study involving 161 patients that examined the safety profile and seizure-related outcomes of in-MDZ in an outpatient setting. Seizure recurrence in their trial was 31% within 6 h, comparable to 39.5% in our study. Regarding the 24-h interval, these authors found seizure recurrence to be 36.9%, which was considerably less than our result of 70.1%. This may be due to the fact that they did not discontinue ASDs, as well as a result of limited surveillance since, in their study, follow-up observation was carried out by caregivers only. In our study design, only hospitalized patients were included so as to be able to ascertain the availability of the time of seizure onset, delivery of the study drug, and seizure termination. Thus, patients were under constant video-EEG monitoring and seizure recurrence was thus much more likely to be detected.

In-MDZ seems to be a safe treatment measure, but, as with any drug, it has weaknesses. Detyniecki et al. [34] detected TEAEs in 57.1% of patients (n = 161), mainly nasal irritation (12.4%) and somnolence (9.3%). Kay et al. [17] (n = 75) observed some side effects in four patients (5.3%), with three showing nasal irritation and one patient delaying the application of the study drug due to per-ictal movement of the head. In a study of 21 patients, De Haan et al. [33] reported 68% of in-MDZ administrations (n = 59) to have caused drowsiness, while 29% led to nasal irritation. In our study and in accordance with the findings mentioned above, we observed various side effects in 108 (44.4%) patients. The main TEAEs included a drop in peripheral oxygen levels, nasal irritation, prolonged sedation, and nausea and vomiting. One patient had respiratory depression, but he recovered quickly after oxygen was applied and no intubation was necessary. Oxygen levels of all other patients recovered within seconds after seizure cessation. No major adverse events were detected that required additional medical assistance. Midazolam and benzodiazepines in general are known for their potential for respiratory depression [35]. In our study, desaturation occurred 78 times (17%). The number of patients with postictically reduced oxygen saturation is similar or rather low when compared with results of dedicated studies on the effect of seizures on breathing. Bateman et al. demonstrated that pulse oximetry will show oxygen desaturations below 90% in 33.2% of all seizures, and such data was well demonstrated in several recent studies, pointing especially at GTCS that lead to a temporary decline in breathing efforts and reduced peripheral oxygenation [32–39]. Overall, postictal respiratory depression may be caused by multiple reasons not limited to anticonvulsive treatment [9]. Additionally, an inherent weakness to our study was that there was no specific timeframe in which oxygen levels were to be measured, rendering a more specific statement with regard to post-in-MDZ oxygenation difficult. Data may also have been altered due to inadequate positioning of the pulse oximeter or displacement during ictal movement due to body position changes [40, 41]. Overall, we did not observe a need for assisted breathing in any patient.

Other than oxygenation, the most common adverse event patients encountered was nasal irritation. This is caused by a need for the midazolam formulation to be acidic in order to be water soluble [42], but may also be independent of the medication given that it may stem from the application of larger volumes of liquids into the nasal cavity [43]. It could thus be observed in most patients but many may be oblivious to it because of the reduced vigilance during and after a seizure [33]. Its occurrence may be lessened by the addition of lidocaine to the solution as suggested by Humphries.
et al. [6], although this approach requires further investigation. Recently, a ready-made formulation of in-MDZ was approved by the United States Food and Drug Administration (FDA) (Nayzilam®; UCB, Brussels, Belgium) to treat acute repetitive seizures [44]. Nayzilam offers 5 mg of midazolam per puff. The new formulation solved the problem of the low pH value by use of other solvents, resulting in a pH range between 5.0 and 9.0, and therefore the side effect of nasal irritation should not occur. Furthermore, use of buccal midazolam can avoid this problem of nasal irritation; however, buccal midazolam is only licensed to children and adolescents below the age of 18 years. Another alternative is the recently FDA-approved intranasal diazepam spray (Valtoco®, Neurelis, Inc., San Diego, CA, USA) [45].

Due to its retrospective design, this study has inherent limitations, as we were not able to randomize or blind the administration of in-MDZ and we did not screen for all possible adverse effects. Although we protocollled in detail the delivery of in-MDZ and the administration of other ASDs, we did not systematically obtain serum levels. However, this is offset by the detailed video and EEG analysis. Furthermore, the study results are not generalizable to individuals with first seizure as the analyzed patients suffered from known epilepsy, an analysis for different etiologies was also not possible due to limited numbers.

In search of new delivery routes, Dhir et al. [46] suggested that intrapulmonal midazolam might be even more efficient as the lung is more vascularized than the nasal mucosa and the drug reaches the arterial blood system directly via this approach. Intrapulmonal delivery is currently under investigation in phase II trials [47]. It must be noted that anatomical differences in patients may render intrapulmonal drug delivery more difficult in children or geriatric patients [47], and seizure-related respiratory changes such as hypopnea, insufficient depth of breathing, or secretion may alter the rate of absorption [36]. To date, in-MDZ remains a preferable alternative, allowing for the easy administration of emergency treatment and protection against damaging subsequent seizures.

**5 Conclusion**

In-MDZ is a reliable and safe anticonvulsive agent in epilepsy patients that can lower the likelihood of seizure occurrence within the upcoming 24 h following its administration. We did not observe any serious TEAEs and there was no need for assisted breathing after treatment with in-MDZ. In-MDZ can be administered very quickly by trained staff within 1–2 min after seizure onset. Better formulation of in-MDZ will help to reduce irritation of the nasal mucosa and thus increase patient comfort and use of this emergency medication.

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**Compliance with Ethical Standards**

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**Ethical approval** The study was approved by the ethics committee of the Goethe-University Frankfurt in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki and its later amendments.

**Informed consent** Informed consent was waived by the ethics committee of the Goethe-University Frankfurt due to the retrospective nature of this study.

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**References**

1. Physicians Emergency, Committee Policies. Clinical policies subcommittee. Clinical policy. Ann Emerg Med. 2004;43(2):278–90. https://doi.org/10.1016/j.annemergmed.2003.10.045.

2. Daniel H, Lowenstein M, Aldredge B. Status epilepticus. New Engl J Med. 1998;338(14):970–6.
3. Poukas VS, Pollard JR, Anderson CT. Rescue therapies for seizures. Curr Neurol Neurosci Rep. 2011;11(4):418–22. https://doi.org/10.1007/s11910-011-0207-x.

4. Strzelczyk A, Ansorge S, Hapfelmeier J, Bonthapally V, Erder MH, Rosenow F. Costs, length of stay, and mortality of super-refractory status epilepticus: A population-based study from Germany. Epilepsia. 2017;58:1533–41. https://doi.org/10.1111/epi.13837.

5. Schubert-Bast S, Zöllner JP, Ansorge S, et al. Burden and epidemiology of status epilepticus in infants, children, and adolescents: A population-based study on German health insurance data. Epilepsia. 2019;60:911–20. https://doi.org/10.1111/epi.14729.

6. Humphries LK, Eiland LS. Treatment of acute seizures: is intranasal midazolam a viable option? J Pediatr Pharmacol Ther. 2013;18(2):79–87. https://doi.org/10.5863/1551-6776.18.2.79.

7. Elisabeth R, Tréluyer JM, Pons G. Pharmacokinetic optimisation of benzodiazepine therapy for acute seizures. Focus on delivery routes. Clin Pharmacokinet. 1999;36(6):409–24. https://doi.org/10.2165/00003495-199936060-00003.

8. Anderson GD, Saneto RP. Current oral and non-oral routes of antiepileptic drug delivery. Adv Drug Deliv Rev. 2012;64(10):911–8. https://doi.org/10.1016/j.addr.2012.01.017.

9. McIntyre J, Norris E, Appleton R, et al. Safety and efficacy of intranasal midazolam versus rectal diazepam for emergency treatment of seizures in children: a randomised controlled trial. Lancet. 2005;366:205–10.

10. Galvin GM, Jelinek GA. Midazolam: an effective intravenous agent for seizure control. Arch Emerg Med. 1987;4(3):169–72. https://doi.org/10.1136/emermed.4.3.169.

11. Kadel J, Bauer S, Hermsen AM, et al. Use of emergency medication in adult patients with epilepsy: a multicentre cohort study from Germany. CNS Drugs. 2018;32(8):771–81. https://doi.org/10.1007/s40263-018-0544-2.

12. Kapur J, Macdonald RL. Rapid seizure-induced reduction of benzodiazepine and Zn⁺ sensitivity of hippocampal dentate granule cell GABAA receptors. J Neurosci. 1997;17(19):7532–7540. http://www.ncbi.nlm.nih.gov/pubmed/9295398).

13. Scott RC, Besag FMC, Boyd SG, Berry D, Neville BGR. Bucal absorption of midazolam: pharmacokinetics and EEG pharmacodynamics. Epilepsia. 1998;39(3):290–4.

14. Bhattacharyya M, Kalra V, Gulati S. Intranasal midazolam vs rectal diazepam in acute childhood seizures. Pediatr Neurol. 2006;34(5):355–9. https://doi.org/10.1016/j.pediatrneuro.2005.09.006.

15. Thakker A, Shanbag P. A randomized controlled trial of intranasal midazolam versus intravenous-diazepam for emergency treatment of seizures in children. J Neurol. 2013;260(2):470–4. https://doi.org/10.1007/s00415-012-6659-3.

16. Fisgin T, Gurer Y, Tezic T, et al. Effects of intranasal midazolam and rectal diazepam on acute convulsions in children: prospective randomized study. J Child Neurol. 2002;17(2):123–6. https://doi.org/10.1177/08830730201700206.

17. Lahat E, et al. Comparison of intranasal midazolam with intravenous-diazepam for acute childhood seizures. J Neurol. 2013;260(2):470–4. https://doi.org/10.1007/s00415-012-6659-3.

18. Holsti M, Sill BL, Firth SD, Filloux FM, Joyce SM, Furnival RA. Prehospital intranasal midazolam for the treatment of pediatric seizures. Pediatr Emerg Care. 2007;23(3):148–53. https://doi.org/10.1097/PEC.0b013e3180328c92.

19. Kay L, Merkel N, Von Blomberg A, et al. Intranasal midazolam as first-line in-hospital treatment for status epilepticus: a pharmaco-EEG cohort study. Ann Clin Transl Neurol. 2019;6:2413–25. https://doi.org/10.1002/acn3.50932.

20. Holsti M, Dudley N, Schunk J, et al. Intranasal midazolam vs rectal diazepam for the home treatment of acute seizures in pediatric patients with epilepsy. Arch Pediatr Adolesc Med. 2010;164(8):747–53. https://doi.org/10.1001/archpediatrics.2010.130.

21. McMullan J, Sasson C, Pancioli A, Silbergleit R. Midazolam versus Diazepam for the treatment of status Epilepticus in children and young adults: a meta-analysis. Acad Emerg Med. 2010;17(6):575–82. https://doi.org/10.1111/j.1553-2712.2010.00751.x.

22. Harbord MG, Kyrkou NE, Kay D, Coulthard KP. Use of intranasal midazolam to treat acute seizures in paediatric community settings. J Paediatr Child Health. 2004;40(9–10):556–8.

23. Mula M. The safety and tolerability of intranasal midazolam in epilepsy. Expert Rev Neurother. 2014;14(7):735–40.

24. Lahat E, et al. Comparison of intranasal midazolam with intravenous-diazepam for treating febrile seizures in children: prospective randomised study. BJM. 2000;321(7523):83–6. https://doi.org/10.1136/bmj.321.7253.83.

25. Klem GH, Lüders HO, Jasper HH, Elger C. The ten-twenty electrode system of the International Federation. Electroencephalogr Clin Neurophysiol. 1999;10(2):371–5. https://doi.org/10.1016/S0161-488X(98)00053-1.

26. Fisher RS, Cross JH, French JA, et al. Operational classification of seizure types by the International League Against Epilepsy: position paper of the ILAE Commission for Classification and Terminology. Z Epileptol. 2018;31(4):272–81. https://doi.org/10.1007/s00228-018-0216-8.

27. von Elm E, Altman DG, Egger M, Pocock SJ, Gotzsche PC, Vandenbroucke JP. The strengthening the reporting of observational studies in epidemiology (STROBE) statement: guidelines for reporting observational studies. PLoS Medicine. 2007;4:e296. https://doi.org/10.1371/journal.pmed.0040296.

28. Scheffer E, Berkovic S, Capovilla G, et al. Classification delle epilepsie della Internationale League Against Epilepsy: position paper della Commissione ILAE per la Classificazione e la Terminologia. Epilepsia. 2017;58(4):512–21. https://doi.org/10.1111/epi.13709.

29. Strzelczyk A, Adjei P, Scott CA, et al. Postictal increase in T-wave alternans after generalized tonic-clonic seizures. Epilepsia. 2011;52:2112–7. https://doi.org/10.1111/j.1528-1167.2011.03266.x.

30. Bancke LL, Dworak HA, Rodvold KA, Halvorsen MB, Gidal BE. Pharmacokinetics, pharmacodynamics, and safety of USL261, a midazolam formulation optimized for intranasal delivery, in a randomized study with healthy volunteers. Epilepsia. 2015;56(11):1723–31. https://doi.org/10.1111/epi.13131.

31. Knoester PD, Jonker DM, Van Der Hoeven RTM, et al. Pharmacokinetics and pharmacodynamics of midazolam administered as a concentrated intranasal spray. A study in healthy volunteers. Br J Clin Pharmacol. 2002;53(5):501–7. https://doi.org/10.1046/j.1365-2125.2002.01588.x.

32. Kwan P, Arzimanoglou A, Berg AT, et al. Definition of drug resistant epilepsy: consensus proposal by the ad hoc Task Force of the ILAE Commission on Therapeutic Strategies. Epilepsia. 2010;51(6):1069–77. https://doi.org/10.1111/j.1528-1167.2009.02397.x.

33. De Haan G, Van Der Geest P, Doelman G, Bertram E, Edelbrock P. A comparison of midazolam nasal spray and diazepam rectal solution for the residential treatment of seizure exacerbations. Epilepsia. 2010;51(3):478–82. https://doi.org/10.1111/j.1528-1167.2009.02333.x.

34. Detynecki K, Van Ess PJ, Sequeira DJ, Whless JW, Meng TC, Pullman WE. Safety and efficacy of midazolam nasal spray in the outpatient treatment of patients with seizure clusters—a
randomized, double-blind, placebo-controlled trial. Epilepsia. 2019;60:1797–808. https://doi.org/10.1111/epi.15159.

35. Gonzalez Castro LN, Mehta JH, Brayanov JB, Mullen GJ. Quantification of respiratory depression during pre-operative administration of midazolam using a non-invasive respiratory volume monitor. PLoS One. 2017;12(2):1–11. https://doi.org/10.1371/journal.pone.0172750.

36. Devinsky O. Effects of seizures on autonomic and cardiovascular function. Epilepsy Curr. 2004;4(2):43–6.

37. Vilella L, Lacuey N, Hampson JP, et al. Incidence, recurrence, and risk factors for peri-ictal central apnea and sudden unexpected death in epilepsy. Front Neurol. 2019;10:1–13. https://doi.org/10.3389/fneur.2019.00166.

38. Lacuey N, Zonji B, Hampson JP, et al. The incidence and significance of pericentral apnea in epileptic seizures. Epilepsia. 2018;59(3):573–82. https://doi.org/10.1111/epi.14006.

39. Bateman LM, Li CS, Seyer M. Ictal hypoxemia in localization-related epilepsy: analysis of incidence, severity and risk factors. Brain. 2008;131(12):3239–45. https://doi.org/10.1093/brain/awn277.

40. Chan ED, Chan MM, Chan MM. Pulse oximetry: understanding its basic principles facilitates appreciation of its limitations. Respir Med. 2013;107(6):789–99. https://doi.org/10.1016/j.resmed.2013.02.004.

41. Mahr K, Bergmann MP, Kay L, et al. Prone, lateral, or supine positioning at seizure onset determines the postictal body position: A multicenter video-EEG monitoring cohort study. Seizure. 2020;76:173–8. https://doi.org/10.1016/j.seizure.2020.02.008.

42. Reves J, Fragen R, Vinik H, Greenblatt D. Midazolam: pharmacology and uses. Anesthesiology. 1985;62(3):310–24.

43. Bailey AM, Baum RA, Horn K, et al. Review of intranasally administered medications for use in the emergency department. J Emerg Med. 2017;53(1):38–48. https://doi.org/10.1016/j.jemermed.2017.01.020.

44. U.S. Food and Drug Administration. NAYZILAM® (midazolam) nasal spray CIV US Prescribing Information. 2019;1–23. https://www.accessdata.fda.gov/drugsatfda_docs/label/2019/211321s000lbl.pdf.

45. U.S. Food and Drug Administration. VALTOCO® (diazepam nasal spray), CIV US Prescribing Information. 2020;1–28. https://www.accessdata.fda.gov/drugsatfda_docs/label/2020/211635s000lbl.pdf.

46. Dhiri A, Zolowska D, Rogawski MA. Seizure protection by intrapulmonary delivery of midazolam in mice. Neuropharmacology. 2013;73:425–31. https://doi.org/10.1016/j.neuropharm.2013.06.002.

47. Maglalang PD, Rautiola D, Siegel RA, et al. Rescue therapies for seizure emergencies: new modes of administration. Epilepsia. 2018;59:207–15. https://doi.org/10.1111/epi.14479.

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