Title: Therapeutic drug monitoring newer antiepileptic drugs: a randomised trial for dosage adjustment.

Running head: Antiepileptic drugs therapeutic drug monitoring

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ABSTRACT

Objective: Therapeutic drug monitoring (TDM) of antiepileptic drugs (AEDs) is widely established for older generation AEDs, whereas there is limited evidence about newer AEDs. Our aim is to assess the benefit of newer generation AEDs TDM in epilepsy.

Methods: We performed a randomised, controlled trial comparing systematic with rescue TDM of lamotrigine, levetiracetam, oxcarbazepine, topiramate, brivaracetam, zonisamide, or pregabalin. Participants were adults with epilepsy, in whom treatment with newer generation AEDs was initiated or needed adjustment. In the systematic TDM arm, AED plasma levels were available at each appointment, whereas in the rescue TDM arm, levels were known only if a study endpoint was reached (inefficacy or adverse events). The primary outcome was the proportion of participants followed over one year without reaching one of the predefined endpoint.

Results: 151 participants were enrolled; global retention in the study was similar in both arms (56% overall, 58% in the systematic and 53% in rescue TDM arm, p=0.6 Cox regression). There was no difference in term of outcome regarding treatment efficacy or tolerability. Partial adherence of clinicians to TDM (adjusting or not AED dosage based on blood levels) did not explain this lack of benefit.

Interpretation: This study provides Class A evidence that systematic drug level monitoring of newer generation AEDs does not bring tangible benefits in the management of patients with epilepsy. Poor correlation between clinical effects and drug levels likely accounts for this finding. However, TDM is useful in several situations, such as pregnancy, as well as compliance issues.

Key words: efficacy, adverse events, antiepileptic drug blood levels, pharmacokinetic, prospective.
INTRODUCTION:

In clinical practice, therapeutic drug monitoring (TDM) is useful to adjust medication dosages of treatments that have large inter-individual and low intra-individual pharmacokinetic variability, show a good correlation between plasma concentrations and clinical effects, and have a narrow therapeutic index. Therapeutic drug monitoring (TDM) of older generation antiepileptic drugs (AEDs) such as phenytoin (PHT), carbamazepine, phenobarbital (PB) or valproate, has been widely implemented since 1960. The relationship between AEDs plasma levels and clinical effect has been well established for those agents, allowing to define reference ranges which are widely accepted.

The use of older generation AEDs is decreasing as newer generation AEDs are being increasingly prescribed, mostly because of their better tolerability profile. In this evolving situation, the International League Against Epilepsy (ILAE) published recommendations for the use of TDM. Evidence about TDM in newer generation AEDs is however very scarce. Observational studies showed a correlation between plasma concentration and clinical response for lamotrigine (LTG), levetiracetam (LEV), and topiramate (TPM), whereas for others (brivaracetam (BRV), lacosamide (LCM), pregabalin (PGB), zonisamide (ZNS), perampanel (PER), oxcarbazepine (OXC)) this relationship remains controversial. Some of these AEDs, such as LTG, LCM, ZNS, or felbamate, show significant inter-individual pharmacokinetic variability due to interactions or metabolism polymorphisms, as they are substrates of hepatic cytochromes or glucuronyltransferases. Other newer generation AEDs have less pharmacokinetic variability (LEV, TPM, OXC), but still show significant changes in their bioavailability with certain co-medications or under specific physiological modifications, such as pregnancy. At times, those pharmacokinetic changes may exert dramatic consequences.

TDM utility for newer generation AEDs was never assessed in a controlled trial. Its usefulness tends however to be accepted and its use is even recommended in specific situations, such as pregnancy.
The aim of this study was to assess whether the systematic monitoring of newer generation AEDs plasma levels provides a tangible benefit in the care of patients with epilepsy.

METHODS:

We set up a randomised controlled two-arm clinical trial comparing systematic versus rescue TDM (ration 1:1). In the rescue TDM arm, medication plasma levels were available only when one of the specific predefined endpoint for failure was reached (status epilepticus, ≥2 seizures with loss of awareness during one year of follow-up, need to add-on a further AED or to discontinue the study drug because of predefined inefficacy or poor tolerability).

The study was approved by the ethical committee of our institution (2015-00079) and it was registered in the clinicalTrials.gov database (NCT02739282). The study was independently monitored, and funded by the Swiss National Scientific Foundation (grant 320030_163430).

Participants

Participants were consenting patients older than 18 years of age, diagnosed with epilepsy and followed in our outpatient clinic. They were treated with newer generation AEDs on mono- or poly-therapy (LTG, LEV, LCM, OXC, TPM, ZNS, BRV, PER), either starting a treatment or requiring dosage adjustment because of inefficacy or adverse events. Pregnant women, in whom systematic TDM is recommended, were excluded. The study was presented by JN, AOR, AFH, and PR to the potential participants.

Protocol

Participants were randomized 1:1 (following a predefined randomisation list) either into the systematic or rescue TDM arm. Allocation was in a list covered with paint, which had to be scratched. The randomisation list was organised in cluster and done by the pharmacists (T.B. and P.A.). Each patient was followed for a studied period of one year with visits set as required clinically, usually 3 to 4 times per year. Participants in both arms had blood samples taken at each visit. Time interval between visits was established by the treating physician according to clinical needs. The participants were instructed
to take their medication at least 6 hours before or after the visit to avoid peak level at the sampling time. All blood samples were collected at steady state after the last dosage adjustment.

A pharmacist specialised in TDM (P.A.) and a neurologist (I.A.R.) assessed all plasma levels to ascertain if concentrations were within the reference ranges, taking into account the time of the last dosing and the medication pharmacokinetics. ILAE recommended reference ranges were used (table 2) \(^{10}\). In the systematic TDM arm, results were systematically communicated to the clinician in charge of the patient within 24 hours. We did not provide target levels to clinicians in the systematic TDM arm, but we compared drug levels (after extrapolation of trough levels when needed) with ILAE recommended reference ranges (table 2) \(^{10}\). Clinicians were then free to adjust medication using these results. In the rescue TDM arm, drug levels were blinded to the physicians during the study follow-up, AED serum levels were communicated to the treating physician only if a study endpoint (see below) was reached. Otherwise, the results became available only at the end of the follow-up period. Therefore, in systematic TDM arm, drug levels were available for the clinician at each, whereas in the rescue TDM arm, drug levels were available only if the treatment failed and needed to be rescued. The rescue TDM arm assessed therefore the management of patients without drug levels available; once an endpoint was reached, drug levels would be available in that arm at the same time the patient would be excluded from the study (drug levels and dosage adjustment performed thereafter would not be considered as part of the outcome). In both arms, clinicians in charge were free to adjust the treatment dosage based on their clinical judgement (for example, if the patient suffered from adverse events AED doses were decreased or AED was stopped and changed), but, as designed, knowledge of drug level was available only for patients in the systematic TDM arm. The inclusion period lasted 18 months and the follow-up 1 year.

At the first visit, demographic data, epilepsy characteristics, current and previous treatments were recorded. At later visits, adverse events and seizure frequency, as well as treatment changes were recorded. The final visit occurred at the end of the 1-year follow-up or when an endpoint was reached.
A combined endpoint (representing treatment failure) was used as the primary outcome of the trial, accounting for both efficacy and adverse events. It was defined by the occurrence of any of: \( \geq 2 \) seizures with impaired awareness (with generalised tonic-clonic seizures), status epilepticus (defined as any seizure lasting >5 minutes), need of an add-on AED, or need to discontinue the studied drug (for either lack of efficacy or adverse reactions); the last two criteria were left to the clinician’s judgement. Severe adverse events were defined as hospitalisation or urgent medical visit, death, life-threatening condition, or condition leading to a persistent disability. Upon occurrence of an endpoint, the participant would be taken off study with drug levels being made available to the clinician in charge of the patient. We recorded the retention of participants, defined as time of follow-up in the study without reaching any of the previous endpoints. In the rescue group, once the patient got an endpoint and the AED levels were available, if there was AED levels equal 0 or very low, then the patient was contacted and asked by the compliance. When the compliance was bad the AED doses were maintained, and when the compliance was good we suspected a fast inducer metabolization and the dose was increased.

**Determination of plasma AED levels**

Plasma levels were determined in the clinical pharmacology laboratory of our institution, which developed an Ultra-performance liquid Chromatography coupled to tandem Mass Spectrometry (UPLC-MS/MS) method, requiring 100 µL of plasma for simultaneous quantification within 7 min of the AEDs. Quality controls/certification in external laboratories were regularly performed.

**Statistical analysis**

Our inclusion target was 150 participants to have sufficient power to demonstrate a 20% difference in treatment failures.

Retention in the study without reaching a predefined endpoint was assessed using survival analysis. *Systematic TDM* and *rescue TDM* were compared using Cox Regression. The outcome was analysed taking into account all patients included (“intention to monitor”) as well as considering only patients who followed the protocol (“per protocol”). Subgroup analysis (drug
resistant epilepsy, focal or generalized epilepsy, patients on monotherapy or polytherapy, patients included due to dosage adjustment or to start of a new AED, patients treated with LTG) were analysed as secondary outcomes. Pearson Chi squared and Mann Whitney U tests were used in univariable analyses. \( P \) values < 0.05 were considered as significant. All analyses were performed using SPSS version 25 (IBM Inc).

RESULTS

Patients

We enrolled 151 patients between June 2016 and December 2017, the overall flow of the trial is shown in figure 1; demographic details of all participants are shown in table 1. Both groups were comparable with respect to age, sex, origin, type of epilepsy, drug resistant epilepsy, epilepsy duration and number of previously tried AEDs. Most had a focal epilepsy (75.5%) and half of the patients had drug resistant epilepsy, according to ILAE’s definition of two adequate AEDs failing to fully control epilepsy 30. Most prescribed AEDs were LTG (66 participants) and LEV (31), followed by ZNS (21), TPM (9), LCM (6) and OXC (4). Half (55%) of the participants were treated with monotherapy. Considering inclusion criteria, 87 (57%) patients were enrolled because of dosage adjustments and 64 (42%) because of introduction of a new AED, of which 17 (26%) were drug naïve.

Outcome

Global retention rate was 56%: 69 patients completed the one-year follow-up without reaching an endpoint. In the “intention to monitor” analysis, both arms had similar retention rates (58% in the systematic TDM arm versus 53% in the rescue TDM arm, \( p=0.6 \), Cox regression, figure 2). Among the 61 participants with endpoints, 34 (55.7%) had ≥2 seizures with loss of awareness, 18 (29.5%) reported adverse events requiring treatment changes, 8 (13.1%) required treatment changes because of inefficacy, and 1 (0.01%) had focal status epilepticus.

In the “per protocol” analysis (figure 1), the overall retention was 56%, without significant difference between systematic (58%) and rescue (55%) TDM arms (\( p=0.7 \), Cox Regression). Retention rates between the two arms were not
different in subgroups (systematic vs rescue arm): drug-resistant epilepsy (60% in systematic vs 47% in rescue arm, p=0.3); focal epilepsy (58% in systematic vs 57% in rescue arm, p=0.9); generalized epilepsy (58% in systematic vs 49% in rescue arm, p=0.6); participants on mono-therapy (82% in systematic vs 70% in rescue arm, p=0.3), or polytherapy (36% in systematic vs 35.5% in rescue arm, p=0.9); participants included for dosage adjustment (52% in systematic vs 49% in rescue arm, p=0.7); participants included for starting an AED (73% in systematic vs 65% in rescue arm, p=0.6), drug naïve patients (85% in systematic vs 90.5% in rescue arm, p=0.3); and participants treated with LTG (53% in systematic vs 56% in rescue arm, p=0.7).

**Plasma drug levels**

All AED blood were measured at least 6 hours after the last dosing, 166 (out of 400, 41.5%) were trough levels.

In the “per protocol” analysis, every participant had at least one AED plasma level. Considering all plasma levels of all AEDs in each participant over the follow-up period, the median proportion of drug levels within the reference ranges per participant was 0.5 (range: 0-1). When stratifying all participants according to their proportion of plasma levels within the reference ranges into high (≥ 50%) or low (<50%), there was no difference of retention between participants with a high proportion of drug levels within the range (52%) versus those with a low proportion (59%) (p=0.5, Cox Regression).

In order to analyse the impact of TDM in the systematic TDM arm, we calculated the proportion of prescribed dosage changes for each participant with the aim of bringing serum levels within the reference range (typically, increasing dosage when the drug level was below the reference range), as opposed to when those changes were not prescribed, or when they were not aimed at bringing drug levels within the ranges. Taking into consideration all changes made for all medications over the follow-up period, the median proportion of changes aimed at bringing drug level in the reference ranges was 0.5 (range: 0-1). When comparing retention rate of participants with a high (≥ 50%) proportion of changes targeting the references ranges and those with a low proportion (<50%), there was no significant difference (58% vs 70%, p=0.7)
We assessed the proportion of medication changes prescribed by physicians in each arm. After the initial change which was the inclusion criterion, 43 dosages adjustments were prescribed in the systematic arm during the follow-up in 32 participants (out of 58, 55%) and 38 adjustments were made in the rescue arm in 31 participants (out of 59, 53%). There was no significant difference between both arms (p=0.3 chi squared).

We classified adjustments made at each visits in three of the different categories: treatment increase when medication(s) dosage was only increased; treatment decrease when medication(s) dosage was only decreased and balanced changes when some of the medications dosage were increased whereas others were decreased. In a total 81 changes during the follow-up in both arms: treatment increase accounted for 39.1% of changes in the systematic arm, 40.6% to the rescue arm, treatment decrease for 17.2% in the systematic arm, 11.8% in the rescue arm; balanced changes for 43.1% in the systematic arm and 47.4% in the rescue. There was no difference between both arms (p=0.7, Chi Squared).

The median number of visits (and TDM for systematic arm) was 3 in both arms. The median interval between TDM/visits in both the systematic and the rescue arm was 91.3 days (range 10.5-186 in the systematic arm, range 9.5-130.6 in the rescue arm, p=0.9).

**Efficacy**

When all inefficacy endpoints were being considered together (occurrence of seizures with loss of awareness, need to change treatment because of insufficient efficacy, occurrence of status epilepticus), there was no retention difference between systematic (65%) and rescue TDM arms (66%, p= 0.9) in the per protocol analysis. When considering the most commonly prescribed AED (LTG, n=49), last plasma levels in participants who had an inefficacy endpoint (median: 3.5 mg/L, range: 0.8-12.2) were not different to last plasma levels in participants without such endpoints in both arms (2.2 mg/l, 0-17.9, p=0.3, Mann Whitney). Similarly, there was no difference for participants on LEV (n=20; median: 10.9 mg/L, range 0-20.3 vs 7.9 mg/L, 1.2-33.6, p=0.6, Mann Whitney).
**Adverse events**

In the per protocol analysis, 45% of participants reported an adverse event at least once (leading to an endpoint or not) during the follow-up period. Most common features were tiredness (25%), psychiatric symptoms (depression, aggressiveness, 23%), weight changes (17%), and unsteadiness (13%). The retention in both arms when considering only patients having toxicity endpoints (18) was similar in systematic (90%) and in rescue TDM (85%) (p=0.4). The occurrence of serious adverse events was also similar in both arms (3.9% in systematic vs 1.3% in rescue TDM, p= 0.6 Chi Square) over the entire follow-up. There was no difference in last LTG plasma levels in participants who reported adverse events endpoints (median: 1.7 mg/L, range: 0-17.9) when compared with last plasma levels of participants without this endpoint, in both arms (2.7 mg/L, 0.7-12.5, p=0.8, Mann Whitney). There was also no difference between arms for participants on LEV (n=20; 4.4 mg/L, range 1.2-7.7 mg/L vs 8.9, range 0-20.3 mg/L, p=0.2, Mann Whitney).

**DISCUSSION**

This trial does not show any significant benefit of systematic TDM of newer generation AED in an outpatients setting compared to rescue TDM, representing TDM as it is widely used in clinical practice. Retention was similar in both groups (59% in the systematic TDM group versus 56%). These results suggest that continuously monitoring drug levels provides at most a modest benefit in the outcome of patient with epilepsy. There was no particular subgroup benefitting of systematic TDM.

Retention in the monotherapy group (76%) and in drug naive participants (90.9%) were greater than the global retention as a favourable outcome (at least in term of seizure control) is more likely in these patients\(^\text{31}\), with no difference between both arms. We also failed to find a difference between both arms in participants with drug resistant epilepsy, a group in which one might have expected that finer medication dosage adjustment might have been more useful given the greater prevalence of polytherapy and their risk of drug-drug interactions \(^\text{32}\). There was also no benefit of systematic TDM in term of efficacy or tolerability, respectively.
The overall retention in the study (56%) was comparable with a similar clinical trial assessing the benefit of TDM of older generation AEDs 33, and with retention studies assessing add-on newer generation AEDs 34,35, supporting the generalizability of our observations. Our methodology was comparable to the two previous clinical trial of older generation AED TDM 33,36. In those studies, however, the treating physician was invited to modify the dosage of patients in the systematic TDM arm in order to achieve concentrations within the reference ranges. We chose a more pragmatic approach letting the clinician decide whether to adjust dosages, taking into account the overall context of the patient with the knowledge of drug level compared to the reference range.

Despite the negative results of the previous clinical trials, older generation AEDs TDM is widely established in clinical practice, at least for PHT 3,4,37–39. TDM usefulness for CBZ and VPA is not clear 5,7,33,36,40. Several limitations of previous as well as our TDM trial may explain these negative outcomes. The adherence of clinicians to adjust dosage according to AED plasma levels was questioned 41. Despite being invited to do so, several authors pointed out that a substantial proportion of drug levels fell below or above the reference range in the previous trials 10, suggesting adjustments after TDM may not have been performed systematically. Another retrospective study suggested that modifying dosage (for PHT and PB) according to plasma levels is associated with a better outcome in term of seizure frequency and adverse events 41; interestingly, TDM adherence (adjusting AED doses based on TDM) by physicians was not a determinant of the outcome when comparing high versus low adherence in our study. Physicians might also have learned from TDM in clinical practice and published studies to anticipate AED pharmacokinetic changes, and they may adjust AED dosage reasonably well without knowing actual drug levels. Another more likely explanation for our negative findings on newer AEDs is the poor correlation between clinical effects and AEDs levels 20,21. We did indeed not find any differences of drug levels (for LTG and LEV), in participants reporting treatment failure, either in term of efficacy or tolerability. Maximal (ceiling) AED levels associated with remission 42,43 might have provided better references in regards of the endpoints of the trial. It was indeed shown that patients may reach optimal seizure control with AED plasma levels often below reference ranges 44–46. Individual reference ranges (referring to drug levels associated
with a response in an individual patient) might be also more useful in clinical practice \(^{10,39,47-49}\), but assessing this aspect went beyond our aims.

Our trial assessed the general systematic use of TDM for dosage adjustment, there are however situations in which TDM might be potentially useful: when toxicity is suspected in patients with intellectual disability in whom assessment of adverse effects is difficult; in lack of efficacy despite appropriate dosages; if compliance issues or drug interactions are suspected; in pregnancy; or in case of renal or hepatic dysfunction. When initiating therapy, individual variability also makes targeting a drug level more desirable than a dosage, there is however that this translates in clinical benefit \(^{10}\).

Our trial has limitations. Due to regulatory issues, its design excluded patients unable to consent, such as those with intellectual disability or cognitive decline, which might represent important groups for TDM, as those patients are less able to report adverse events \(^{10}\). Intervals between visits were also irregular, as we chose to remain as pragmatic as possible to maximise adherence with trial protocol. Reflecting clinical practice, blood samples were drawn during the time the participants spent in clinic, drug levels were therefore not for half of them trough concentrations. Participants were however instructed not to take their medication when coming at the morning clinic, blood samples were drawn at least 6 hours after dosing, and were then interpreted taking into account last dosing and medication pharmacokinetics. The endpoints we chose were also more stringent than those used in the previous trials, particularly occurrence of ≥2 seizures with loss of awareness in patients with drug resistant epilepsy. In previous trials, endpoints were typically the need to switch or to add another AED because of toxicity or inefficacy without clearer criteria \(^{33,36}\). The endpoints we chose seem however more likely to reflect on a real benefit in patients’ quality of life, such as the return of the ability to drive.

The sample size may also limit the ability to show a small significant difference between both arms; but the intervention effect is likely to be small. The trial included all newer generation AEDs available at the time of study, making it a heterogeneous group of drugs to assess. Some of these compounds are likely to be better candidate for TDM, such as LTG. We did however not find any obvious difference when analysing individually LTG and LEV, although the sample size was not large enough to drive firm conclusions.
We could not analyse specific value of systematic TDM for topiramate, oxcarbazepine or lacosamide, due to the small sample size, so usefulness of TDM of these AEDs is unclear. Another important limitation is that the treating physicians were experienced neurologist specialised in epilepsy, our finding could therefore not be applied to patients treated by general neurologists and primary care physicians. Our study also assessed the usefulness of TDM to prevent treatment failure (as defined by our endpoints), but it did not assessed TDM potential benefit after the treatment failed. Our trial assessed the general systematic use of TDM for dosage adjustment, there are however situations in which TDM might be useful; when toxicity is suspected in patients with intellectual disability in whom assessment of adverse effects is difficult; in lack of efficacy despite appropriate dosages; if compliance issues or drug interactions are suspected; in pregnancy; or in case of renal or hepatic dysfunction. When initiating therapy, individual variability also makes targeting a drug level more desirable than a dosage, there is however no evidence that this translates in clinical benefit.

In summary, systematic TDM of newer generation AEDs in patients with epilepsy appears unlikely to bring any tangible benefit in clinical practice; systematic monitoring of newer generation is therefore not justified. TDM should be reserved for selected situations such as pregnancy/pharmacokinetic changes, drug interactions or renal or hepatic failure, suspicion of non-compliance, suspicion of toxicity, lack of effect in spite of correct dosages\textsuperscript{2,10,47,48,50}.

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Author contributions:


Potential conflict of interest:
The authors have nothing to report relevant for the study.

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Figure legends:

Figure 1: Flow diagram of the study. The single death was not related to the study (brain tumour progression).

Figure 2: Overall retention in both arms (“intention to monitor” analysis). There was no difference between the systematic TDM arm (58%) and the rescue TDM arm (53%, p = 0.5 Cox regression).
Table 1: Demographic characteristics of participants according to their randomization. *Combination of all inefficacy endpoints: patients (≥2) with loss of awareness, status epilepticus and need to add a treatment because of inefficacy.

<table>
<thead>
<tr>
<th></th>
<th>Total (151)</th>
<th>Systematic TDM arm (76)</th>
<th>Rescue TDM arm (75)</th>
<th>P value</th>
<th>Test used</th>
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</thead>
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<tr>
<td>Sex (female)</td>
<td>55.6%</td>
<td>55.3%</td>
<td>56%</td>
<td>0.9</td>
<td>Pearson’s Chi Square</td>
</tr>
<tr>
<td>Age, median, (range)</td>
<td>37(18-82)</td>
<td>38 (18-82)</td>
<td>36 (18-76)</td>
<td>0.5</td>
<td>U Mann Whitney</td>
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<tr>
<td>Focal epilepsy</td>
<td>75.5%</td>
<td>76.3%</td>
<td>74.7%</td>
<td>0.8</td>
<td>Pearson’s Chi Square</td>
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<tr>
<td>Drug-resistant epilepsy</td>
<td>48.7%</td>
<td>46.7%</td>
<td>50.7%</td>
<td>0.6</td>
<td>Pearson’s Chi Square</td>
</tr>
<tr>
<td>Epilepsy duration-years</td>
<td>7 (0-47)</td>
<td>8 (0-44)</td>
<td>7 (0-47)</td>
<td>0.7</td>
<td>U Mann Whitney</td>
</tr>
<tr>
<td>Number of previously</td>
<td>1 (0-9)</td>
<td>1 (0-7)</td>
<td>1 (0-9)</td>
<td>0.3</td>
<td>U Mann Whitney</td>
</tr>
<tr>
<td>tried AEDs median, (range)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>All inefficacy endpoints*</td>
<td>43 (28.5%)</td>
<td>24 (31.6%)</td>
<td>19 (25.3%)</td>
<td>0.4</td>
<td>Pearson’s Chi Square</td>
</tr>
<tr>
<td>Adverse event endpoints</td>
<td>18 (12%)</td>
<td>7 (9.2%)</td>
<td>11(14.6%)</td>
<td>0.6</td>
<td>Pearson’s Chi Square</td>
</tr>
</tbody>
</table>
Table 2: AED and ILAE recommended reference ranges (Patsalos, Epilepsia 2008, Patsalos Seeizure 2014, Patsalos Therapeutic drug monitoring 2018)

<table>
<thead>
<tr>
<th>Antiepileptic drug</th>
<th>Reference range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brivaracetam (BRV)</td>
<td>0.4-1.2 mg/L</td>
</tr>
<tr>
<td>Lacosamide (LCM)</td>
<td>10-20 mg/L</td>
</tr>
<tr>
<td>Lamotrigine (LTG)</td>
<td>2.5-15 mg/L</td>
</tr>
<tr>
<td>Levetiracetam (LEV)</td>
<td>12-46 mg/L</td>
</tr>
<tr>
<td>Oxcarbazepine (OXC)</td>
<td>3-35 mg/L</td>
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<tr>
<td>Perampanel (PER)</td>
<td>180-980 μg/L</td>
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<tr>
<td>Pregabalin (PGB)</td>
<td>2.8-10 mg/L</td>
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<tr>
<td>Topiramate (TPM)</td>
<td>5-20 mg/L</td>
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<tr>
<td>Zonisamide (ZNS)</td>
<td>10-40 mg/L</td>
</tr>
</tbody>
</table>
Figure 1: Flow-diagram of the study. The single death was not related to the study (brain tumour progression).
Figure 2: Retention (56%) without endpoints, taking into account all participants ("intention to monitor" analysis).
Figure 3: Overall retention in both arms ("intention to monitor" analysis). There was no difference between the systematic TDM arm (58%) and the rescue TDM arm (53%, p= 0.5 Cox regression).