Therapeutic monitoring of anti-seizure medications in low- and middle-income countries: a systematic review

Abstract

Background

The treatment gap for epilepsy is large in low- and middle-income countries (LMICs) and the effectiveness and safety of the available anti-seizure medication (ASMs) is not fully understood. We systematically reviewed available evidence on therapeutic drug monitoring (TDM) of ASM in LMIC.

Methods

We searched four main databases (PubMed, Psych-Info, CINAHL and Embase) up to 31st December 2020, with eligible articles screened using a PRISMA checklist and a set of exclusion and inclusion criteria. Full texts were examined to evaluate the extent and practice of TDM in LMICs. Analyses were performed using Stata 13 and descriptive statistics were used to pool median distribution of TDM across studies.

Results

Of the 6,309 articles identified in the initial search, 65 (1.0%) met the eligibility criteria. TDM of ASMs was mostly done to assess toxicity (42.8%), but rarely to monitor adherence (9.0%). TDM differed by economic status and infrastructural status with majority of the studies...
coming from Europe (53.8%) and upper-middle-income countries (87.6%). First generation ASMs (82.3%) were more likely to be monitored than second generation ASMs (17.6%) and carbamazepine was the most frequently monitored drug. Fluorescence Polarization Immunoassay (FPIA) was the most common technique used for TDM (41.5%) followed by High-Performance Liquid Chromatography (HPLC) (16.9%). In addition, FPIA was the cheapest method of TDM based on approximated costs ($1000, TDx system). Assay validation and quality control were reported variably, and reference ranges used during TDM of ASMs were relatively uniform.

Conclusions

TDM is mostly done to evaluate ASM toxicity, but rarely for other reasons such as evaluating adherence or assessing clinical efficacy. There is a need for more investment in comprehensive and targeted TDM in LMICs when initiating treatment, switching therapies, adding or removing ASM and evaluating treatment response and safety of both first generation and second generation ASMs.

Plain Language Summary

The number of people with epilepsy who do not have access to treatment is high in low- and middle-income countries (LMICs) and the effectiveness and safety of the available medication for epilepsy is not fully understood. We systematically reviewed available evidence on therapeutic drug monitoring (TDM), i.e. measuring medication levels to ensure they are within the recommended ranges in a LMIC.

We searched four main databases (PubMed, Psych-Info, CINAHL and Embase) up to 31st December 2020, with eligible articles screened using a PRISMA checklist and a set of criteria tailored to our study objectives. Full texts were examined to evaluate the extent and practice of TDM in LMICs. Analyses were performed using Stata 13 and we used statistical methods to describe the distribution of TDM across studies. Of the 6,309 articles identified in the initial search, 65 (1.0%) met the set criteria for inclusion. Measurement of medication levels was mostly done to check for side effects (42.8%), but rarely to ensure if patients were taking their medication as prescribed (9.0%). Distribution differed by economic status with the majority of the studies coming from Europe (53.8%) and upper-middle-income countries (87.6%). Older medications for epilepsy (82.3%) were more likely to be monitored than newer drugs (17.6%), with carbamazepine being the most frequently measured drug. A laboratory method called Fluorescence Polarization Immunoassay was the most common (41.5%) and affordable (costing about $1000). Reference ranges for medication levels used during TDM were relatively uniform.

We concluded that TDM was rarely done when evaluating medication adherence or clinical efficacy. Therefore, there is a need for more investment in comprehensive and targeted TDM in LMICs when initiating treatment, switching therapies, adding or removing
medications and evaluating treatment response and safety of old and newer medications for epilepsy.

**Keywords**
Therapeutic drug monitoring, TDM, anti-seizure medication, ASM, low- and middle-income countries, LMICs, resource-limited settings

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**Introduction**

Epilepsy is a serious neurological disorder characterized by an enduring predisposition to experience unprovoked seizures and often results in neurobehavioural, psychosocial consequences (Fisher et al., 2005) and premature mortality (Levira et al., 2017). Approximately 70 million people worldwide have epilepsy, more than 85% of whom live in low- and middle-income countries (LMICs) (Ngugi et al., 2011; World Health Organization, 2018). Appropriate treatment schedules of anti-seizure medication (ASMs), the main-stay of therapy for seizure disorders, results in complete control of seizures in up to 70% of patients yet the treatment gap is still large in LMICs (Mbuba et al., 2012; Meyer et al., 2010). In these resource-poor settings, health-seeking behavior depends on availability and accessibility of health care facilities, cultural beliefs and financial resources (Newton & Garcia, 2012), resulting in people seeking treatment only once the disease is already severe (Kariki et al., 2015).

The ASMs such as carbamazepine, phenobarbital, phenytoin and valproic acid are listed in the Health Organization (WHO) essential medicines list as first line anti-seizure medications for the management of epilepsy (WHO Model List of Essential Medicines, 2020). Lamotrigine and ethosuximide are recommended for the management of focal and focal to bilateral tonic-clonic seizures and absence seizures, respectively. Benzodiazepines such as clonazepam and diazepam are commonly used in resource-limited countries for acute management of acute seizures including status epilepticus (World Health Organization, 2004). Phenobarbital, phenytoin, carbamazepine and sodium valproate are widely available in public health facilities in LMICs (Ibinda et al., 2017), while second-generation ASMs such as ethosuximide, gabapentin, lamotrigine, topiramate, levetiracetam, pregabalin and clobazam are only available in private facilities or tertiary centres, not within the reach of the poor populations (Chin, 2012).

ASMs therapy requires a careful balance of controlling seizures while minimizing toxicity. Routine monitoring allows the prescriber to evaluate treatment response while identifying adverse effects and drug interactions. First generation ASM which are commonly used in low-resource settings display highly variable pharmacokinetic profiles which may affect the concentrations in blood following mono- or polytherapy (Knezevic & Marzinke, 2018). For instance, carbamazepine has a narrow therapeutic index which means that a small change in the administered dose can result in sub-clinical response or toxicity while auto-induction in the first two weeks of treatment may result in delayed response. Likewise, phenytoin displays zero-order pharmacokinetics at high doses which can lead to accumulation and toxicity (Hwang & Tsai, 2004). These variations in ASM concentrations can be empirically assessed clinically but the extent can only be validated through monitoring of plasma levels.

Despite great efficacy in controlling seizures in people with epilepsy, majority of ASM have been linked with several side effects including feelings of tiredness, stomach upset, dizziness, blurred vision, cognitive and behavioural problems (Kwan et al., 2013; Mutanana et al., 2020). These adverse effects require monitoring but TDM is rarely done in LMICs due to limited infrastructure, resources and lack of technical expertise to operate equipment for mass spectrometry and immunoassays (Tomson et al., 2007). In view of the gaps in availability, access, and effects of use of ASMs in LMIC highlighted above, and because of limited resources, we need a criterion to guide implementation of targeted TDM in LMIC. Therefore, we conducted a review to examine available evidence on TDM, purposes, methods and ASM reference ranges used during monitoring in these settings.

**Methods**

**Protocol registration**

The protocol for this review was published on PROSPERO under the registration number CRD42018116937.

**Search strategy**

The main search terms “monitoring”, “antiepileptic drugs” and “low- and middle-income countries” were developed, connected with a Boolean operator and then entered into the major databases (Table 1). There were no restrictions based on date, language or publication format during the search to allow us to retrieve as many relevant articles as possible. Although all the search terms were in English, we postulated that many peer-reviewed non-English articles would have English-translated abstracts therefore, non-English articles would be picked up during the search. The initial database search was carried out

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Table 1. Search strategy.

| ((Monitoring) OR (Measurement) OR (Evaluation) OR (Examination) OR (Observation) OR (Concentration) OR (Levels)) AND (Antiepileptic drugs) OR (AED) OR (Antiepileptic agents) OR (Antiseizure medications) OR (ASM) OR (Anticonvulsants) OR (Carbamazepine) OR (Phenobarbital) OR (Phenytoin) OR (Sodium Valproate) OR (Ethosuximide) OR (Levetiracetam) OR (Gabapentin) OR (Lamotrigine) OR (Topiramate) OR (Pregabalin) OR (Clobazam) OR (Diazepam) OR (Clonazepam)) AND ((LMICs) OR (Low-income countries) OR (Lower-middle-income countries) OR (Upper-middle-income countries) OR (Sub-Saharan Africa) OR (Africa) OR (East Asia) OR (South Asia) OR (Middle East) OR (South America) OR (Tajikistan) OR (Ukraine) OR (Georgia) OR (Moldova) OR (Kosovo) OR (Turkey) OR (Serbia) OR (Russia) OR (Kazakhstan) OR (Bulgaria) OR (Belarus) OR (Armenia) OR (Albania)) |
on 8th November 2018 and later updated to 31st December 2020. The results in different databases were as follows: PubMed (N= 2,964), Psych-Info (N= 86), CINAHL (N= 39) and Embase (N= 3,821).

Inclusion and exclusion criteria for selecting studies
This review focused on studies reporting monitoring of both detectable and optimal levels of all ASMs in human serum, plasma and other body fluids. We only included primary studies originating from low-, lower-middle or upper-middle-income countries using the World Bank GNI per capita cut-offs for the 2021 fiscal year (World Bank, 2020). The studies included were either cross-sectional, retrospective or prospective cohort studies published in peer-reviewed journals. Although we did not impose language restrictions during the search, only articles written in English or those that had English translations were included due to the language expertise of the reviewers. There was no year of publication restriction. Reviews, case studies, conference abstracts, poster sessions, editorials and case reports were excluded from this review.

Data screening, appraisal and extraction
Screening of eligible studies was done using Microsoft Excel (Microsoft Office 365 ProPlus ; Version 2002) by MA assisted by SK; first by title and abstract and then by full text. Hand-searching of reference lists of eligible studies and relevant reviews was also conducted. For inaccessible articles, personal requests for full texts were made to the authors via email. Any discrepancies arising during the screening stage were resolved through consensus and in consultation with CN. A standardized form was used to extract data relevant to the objectives including study setting, methodology, interventions and outcome measures such as ASMs monitored, purpose and methods of TDM.

The quality of each eligible article was assessed using the Newcastle-Ottawa Scale (NOS) for observational studies (http://www.ohri.ca/programs/clinical_epidemiology/nos_manual.pdf), with a focus on selection- case and control definitions and representativeness (having the outcome of interest); comparability (criteria used to decide on the cases and comparison group and matching), if any; and ascertainment of exposure and non-response rate. We used the adapted and published NOS for cross-sectional and case-control studies and assessment was done at the study level. Eligible studies were scored out of 10 as per the quality appraisal scale. Those that scored between zero to three were considered poor quality, four to six were rated average quality and studies that scored seven and above were rated good quality. Study selection and data extraction were conducted by MA, assisted by SK who independently reviewed 30% of all eligible full-text articles. Any discrepancies arising during the screening and data extraction were resolved through consensus and in consultation with CN.

Outcome measures included the proportion of TDM by country, continent, WHO region, income level and classification as a percentage and the mean and median reference ranges used during monitoring by ASM. We extracted data on the regional distribution of the included articles, type of ASM used, method of conducting TDM, validation procedures and references ranges.

Statistical analysis
Analyses were performed using Stata (StataCorp. 2013. Stata Statistical Software: Release 13. College Station, TX: StataCorp LP). Frequencies between studies were compared using Pearson’s chi-squared test and Fisher’s exact test, when the observations were few. Continuous measures were compared using Man-Whitney U test because most did not follow a Gaussian distribution. Descriptive statistics were reported using measures of central tendency. We had planned to conduct a meta-analysis, but a narrative synthesis was conducted to summarize the outcomes of interest due to significant variability in the reporting of outcomes.

Results
Search results
Following a search of four major databases and reference lists of included studies and relevant reviews, 6,039 studies were retrieved. After screening by title and abstract, 198 studies were selected for full text retrieval as shown by the PRISMA flow diagram (Figure 1). Of the 198 studies, only 65 (32.8%) were eligible for qualitative and quantitative analyses (Table 2).

Study characteristics
In terms of WHO regional groupings, the European region contributed a bulk of the studies (53.8%), while the Americas contributed the least (1.5%). Nearly a third (27.7%) of the studies were from the African region while 13.8% were from the Western pacific region (Table 3). Turkey and South Africa were well-represented with each country contributing 36.9% and 20.2% of the studies, respectively while China and Serbia each contributed 13.0% of the included studies. Those originating from the remaining nine countries each contributed less than 5% of the overall proportion. Majority of the studies were from upper-middle-income countries (87.6%) with studies from low- and lower-middle-income countries contributing less than 10% each (Table 3).

This review included studies published from 1977 to 2020, with majority (43.0%) being from the last 10 years (2010 to 2020) and cohorts were the most popular study design (76.9%). Fifty-eight (89.2%) of the studies were hospital-based and a similar proportion of the studies was from urban and semi-urban areas (Table 3). Fifty-three (81.5%) of the included studies were of average quality, 12 (18.4%) studies were classified as high quality and no studies had a score of less than 4 based on the Newcastle-Ottawa scale rating. The age group of participants was not specified in 14 (21.5%) studies but in studies that specified age groups, 41.5% of the studies involved adolescents and children while 36.9% involved adults.
Figure 1. A PRISMA flow diagram used to summarize the number of studies identified, screened, eligible and included and excluded at each stage.

Table 2. List of included articles.

| Author          | Year | Study Title                                                                 |
|-----------------|------|------------------------------------------------------------------------------|
| Abaci A, et al. | 2009 | Metabolic alterations during valproic acid treatment: a prospective study    |
| Akin R, et al.  | 1998 | Evaluation of bone mineral density in children receiving antiepileptic drugs |
| Aldemir E, et al.| 2012 | Valproate-associated reproductive hormone abnormalities: do bipolar men have the same risk as epileptic men? |
| Altunbasak S, et al. | 1997 | Asymptomatic Hyperammonemia in Children Treated With Valproic Acid          |
| Atugonza R, et al. | 2016 | Multiple anti-epileptic drug use in children with epilepsy in Mulago hospital, Uganda: a cross sectional study |
| Brzakovic B, et al. | 1997 | Cerebrospinal Fluid and Plasma Pharmacokinetics of Phenobarbital after Intravenous Administration to Patients with Status Epilepticus |
| Brzakovic BB, et al. | 2012 | Impact of age, weight and concomitant treatment on lamotrigine pharmacokinetics |
| Buchanan N, et al. | 1977 | Serum levels of anticonvulsants and haematological sequelae in Black patients |
| Cai MW, et al.  | 1993 | Free Phenytoin Monitoring in Serum and Saliva of Epileptic Patients in China |
| Chang Y, et al. | 2014 | Correlation of the UGTIA4 Gene Polymorphism With Serum Concentration and Therapeutic Efficacy of Lamotrigine in Han Chinese of Northern China |
| Chbili C, et al. | 2016 | Effects of EPHX1 and CYP3A4*22 genetic polymorphisms on carbamazepine metabolism and drug response among Tunisian epileptic patients |
| Chen Z, et al.  | 2011 | Simultaneous determination of valproic acid and 2-propyl-4-pentenoic acid for the prediction of clinical adverse effects in Chinese patients with epilepsy |
| Cicek NP, et al. | 2018 | The effects of valproate and topiramate use on serum insulin, leptin, neuropeptide y and ghrelin levels in epileptic children |
| Author            | Year | Study Title                                                                                           |
|-------------------|------|--------------------------------------------------------------------------------------------------------|
| Daci A, et al.    | 2015 | Polymorphic Variants of SCN1A and EPHX1 Influence Plasma Carbamazepine Concentration, Metabolism and   |
|                   |      | Pharmacoresistance in a Population of Kosovar Albanian Epileptic Patients                              |
| Davies VA, et al. | 1985 | Precursor Prothrombin Status In Patients Receiving Anticonvulsant Drugs                                 |
| Deda G, et al.    | 2003 | Effect of long-term carbamazepine therapy on serum lipids, vitamin B12 and folic acid levels in children |
| Djordjevic N, et al. | 2016 | CYP1A2 genotype affects carbamazepine pharmacokinetics in children with epilepsy                      |
| Dowse and Futter | 1991 | Outpatient compliance with theophylline and phenytoin therapy                                         |
| Ecevit C, et al.  | 2004 | Effect of carbamazepine and valproate on bone mineral density                                         |
| Erdal A, et al.   | 2013 | The evaluation of sexual functions and sex hormones in male and female epilepsy patients taking         |
|                   |      | valproic acid and carbamazepine monotherapy                                                          |
| Geda G, et al.    | 2002 | Serum lipids, vitamin B12 and folic acid levels in children receiving long-term valproate therapy      |
| Gulcebi MI, et al.| 2018 | The effect of serum levetiracetam concentrations on therapeutic response and IL1-beta concentration   |
|                   |      | in patients with epilepsy                                                                             |
| Gulcebi MI, et al.| 2011 | The relationship between UGT1A4 polymorphism and serum concentrations of lamotrigine in patients with |
|                   |      | epilepsy                                                                                                |
| Hemingway C, et al.| 1999 | The effect of carbamazepine and sodium valproate on the blood and serum values of children from a      |
|                   |      | third-world environment                                                                               |
| Hundt HKL, et al. | 1983 | Carbamazepine and its major metabolites in plasma: a summary of eight years of therapeutic drug         |
|                   |      | monitoring                                                                                             |
| Ibinda F, et al.  | 2017 | Magnitude and Factors Associated with Nonadherence to Antiepileptic Drug Treatment in Africa: A        |
|                   |      | cross-sectional multi-site study                                                                       |
| Ibinda F, et al.  | 2014 | Evaluation of Kilifi Epilepsy Education Programme: A randomized controlled trial                        |
| Ilic V, et al.    | 2016 | Duration of valproic acid monotherapy correlates with subclinical thyroid dysfunction in children with  |
|                   |      | epilepsy                                                                                                |
| Incecik F, et al. | 2007 | Effects of valproic acid on hearing in epileptic patients                                              |
| Jakovljevic MB, et al.| 2008 | Inverse correlation of valproic acid serum concentrations and quality of life in adolescents with epilepsy |
| Joubert PH, et al.| 1977 | Epilepsy and IgA Deficiency-the Effect of Sodium Valproate                                             |
| Jovanovic M, et al.| 2014 | Effect of Long-term Topiramate Therapy on Serum Bicarbonate and Potassium Levels in Adult Epileptic     |
|                   |      | Patients                                                                                                |
| Kurekci AE, et al.| 1995 | Plasma trace element, plasma glutathione peroxidase, and superoxide dismutase levels in epileptic       |
|                   |      | children receiving antiepileptic drug therapy                                                          |
| Lai M             | 1985 | Steady state serum levels of anticonvulsant drugs in Chinese epileptic patients living in Taiwan        |
| Lalic M et al.    | 2009 | Lamotrigine and valproate pharmacokinetics interactions in epileptic patients                          |
| Leary PM, et al.  | 1989 | The prevalence of adverse reactions to anticonvulsants in children with epilepsy                       |
| Lu Y, et al.      | 2017 | Effects of UGT1A9 genetic polymorphisms on monohydroxylated derivative of oxcarbazepine concentrations |
|                   |      | and oxcarbazepine monotherapeutic efficacy in Chinese patients with epilepsy                           |
| Ma et al.         | 2019 | Pharmacist impact on adherence of valproic acid therapy in pediatric patients with epilepsy using active |
|                   |      | education techniques                                                                                   |
| Majola MP, et al. | 2000 | Factors influencing phenytoin-induced gingival enlargement                                              |
| Mbuba CK, et al.  | 2012 | Risk factors associated with the epilepsy treatment gap in Kilifi, Kenya: a cross-sectional study        |
| McFayden ML, et al.| 1990 | The relevance of a First-World therapeutic drug monitoring service to the treatment of epilepsy in Third-|
|                   |      | World conditions                                                                                        |
| McFayden ML, et al.| 2007 | Therapeutic Drug Monitoring and Clinical Outcomes in Epileptic Egyptian Patients: A gene Polymorphism  |
|                   |      | Perspective Study                                                                                        |
| Milovanovic DD, et al.| 2016 | The influence of CYP2C8*3 on carbamazepine serum concentrations in epileptic paediatric patients         |
Anti-seizure medications monitored

First generation ASMs (carbamazepine, phenobarbital, phenytoin and valproic acid) were monitored exclusively in 56 (82.3%) of the studies, while second generation ASMs (including lamotrigine, levetiracetam, oxcarbazepine and topiramate) were monitored in 12 (17.6%) of the included studies. Anticonvulsant benzodiazepines were only monitored in one study (Table 4). There were 108 instances where TDM of ASMs was reported with some studies reporting monitoring of more than one agent. Valproic acid (30.5%) and carbamazepine (27.7%) contributed to nearly a third of the overall proportion of ASMs monitored with phenytoin and phenobarbital accounting for 17.1% and 13.8%, respectively.

The remaining ASMs including lamotrigine, topiramate and oxcarbazepine and benzodiazepines accounted for less than 1.0–5.5%.

There were significant differences across the various WHO regions for TDM of first generation ASMs (p=0.037), with studies from the Americas and Eastern Mediterranean regions not reporting any monitoring of second generation ASMs or anticonvulsant benzodiazepines (Table 4). A similar trend for first generation ASMs was observed in the various continents with significant differences in the proportions (p=0.016). However, studies from Asia had an equal distribution (44.4%) of studies reporting monitoring of first and second generation ASMs.
ASMs (Table 4). There were no significant differences in monitoring of either first or second generation ASMs or benzodiazepines across studies from different income levels or urban and rural settings.

Purpose of Therapeutic Drug Monitoring (TDM)
Overall, there were 77 instances where the reasons for conducting TDM were reported with some studies reporting multiple purposes for TDM. Monitoring to assess side effects constituted nearly half (42.8%) of the reported instances while monitoring with the aim of evaluating compliance and to examine effect of genetics on ASM levels accounted for 12 (15.5%) and 11 (14.2%) of the instances. Other reasons for TDM that were reported accounted 2.5 to 10.3% of the instances, including monitoring to assess clinical efficacy, to determine drug interactions or for the purpose of dose titration (Table 5).

There were significant differences across the various WHO regions of monitoring to assess unwanted ASM side effects (p<0.05) accounting for 62.8% and 43.4% of the instances in the European and African regions, respectively (Table 5). Across the regions, there were also significant differences in TDM conducted to examine the effect of genetic variations on ASM levels (p= 0.025), making up a third (31.2% and 33.3%) of the proportions in the Western Pacific and Eastern Mediterranean regions. In addition, there were significant differences across the regions of TDM to assess adherence (p=0.012) and clinical efficacy (p=0.023) and a similar trend was observed across the continents. There were no significant differences in purposes of TDM across various income levels (Table 5). A similar trend was observed when urban and rural areas were compared except for TDM conducted to assess compliance which accounted for nearly half (42.8%) of the TDM in rural areas but only 6.7% in urban areas (p=0.019).

Methods of TDM
Twelve different TDM techniques were reported in 50 (76.3%) of the included studies with some studies reporting more than one method (53 instances). Fluorescence polarization immunoassay (FPIA) was reported in nearly half (41.5%) of the instances (Table 6), while High Performance Liquid Chromatography (HPLC) and HPLC with ultraviolet detection (HPLC-UV) were used in 9 (16.9%) and 5 (9.4%) of the instances. The remaining techniques including Liquid Chromatography-Mass Spectrometry (LC-MS) and Gas Chromatography (GC) accounted for 1.8 to 5.6% of the overall proportion (Table 6).

FPIA was the most common method of TDM in the African (64.2%) and European regions (42.3%) with a trend toward significance for differences across the regions (p=0.075). HPLC was popular in studies from the Eastern Mediterranean (50.0%) and Western Pacific (36.3%) regions with a significant difference across the regions (p=0.035). A similar trend was observed across the continents, but with no significant differences while the TDM method used in the single study from South America (Americas region) was not reported. The trend remained same across different income levels and in urban and rural areas with FPIA being the most popular TDM method in low- (50.0%), lower-middle- (60.0%) and upper-middle-income (41.3%) countries and urban (40.0%) and rural (75.0%) areas. There were no significant differences across the studies across different income levels or when comparing studies from urban and rural areas.

The overall cost of the various techniques was unspecified in all the included studies, therefore, the cost of purchase of equipment was used to allude to the likely cost per test. LC-MS equipment was the most expensive based on commercially quoted prices, costing about $149,000 (TSQ Quantum Access from Thermo-Fischer), while FPIA equipment (TDx from Abbot Technologies) was the least expensive, costing about $1,000 (Table 6).
### Table 4. Distribution different ASM classes across WHO regions, continents, income and urbanisation status.

| ASM Classification                  | First generation ASM N (%) | Second generation ASM N (%) | Benzodiazepine N (%) |
|-------------------------------------|----------------------------|-----------------------------|----------------------|
| Overall (N=68)                      | 56 (82.3)                  | 12 (17.6)                   | 1 (1.4)              |
| **WHO Region**                      |                            |                             |                      |
| African (N=19)                      | 18 (94.7)                  | -                           | 1 (5.2)              |
| Americas (N=1)                      | 1 (100.0)                  | -                           | -                    |
| Eastern Mediterranean (N=2)         | 2 (100.0)                  | -                           | -                    |
| European (N=35)                     | 28 (80.0)                  | 5 (14.2)                    | 2 (5.7)              |
| Western Pacific (N=9)               | 4 (44.4)                   | 5 (55.5)                    | -                    |
| **P value**                         | **0.037**                  | **0.005**                   | 1.000                |
| **Continent**                       |                            |                             |                      |
| Africa (N=21)                       | 20 (95.0)                  | -                           | 1 (5.0)              |
| Asia (N=9)                          | 4 (44.4)                   | 4 (44.4)                    | 1 (11.2)             |
| Europe (N=35)                       | 28 (80.0)                  | 5 (14.2)                    | 2 (5.7)              |
| South America (N=1)                 | 1 (100.0)                  | -                           | -                    |
| **P value**                         | **0.016**                  | **0.010**                   | 0.636                |
| **Income status**                   |                            |                             |                      |
| Low-income (N=3)                    | 3 (100.0)                  | -                           | -                    |
| Lower-middle-income (N=8)           | 8 (100.0)                  | -                           | -                    |
| Upper-middle-income (N= 62)        | 49 (79.0)                  | 12 (19.3)                   | 1 (1.6)              |
| **P value**                         | **0.389**                  | **0.475**                   | 1.000                |
| **Urbanisation**                    |                            |                             |                      |
| Rural (N=7)                         | 7 (100.0)                  | -                           | -                    |
| Urban (N=65)                        | 52 (80.0)                  | 12 (18.4)                   | 1 (1.5)              |
| **P value**                         | **0.232**                  | **0.262**                   | 0.903                |

### Assay validation and quality control

Fourteen (23%) of the included studies described validation of the various assays used in TDM. Nine (64%) of these studies reported either overall coefficients of variation (CV), intra- and interassay CV, percentage accuracy, percentage precision and intra- and interday measures of accuracy including reproducibility and relative standard deviation. This significant variability in reported outcomes made it difficult to pool these outcomes.

Among studies that reported intra- and interassay CVs, one study using LC-MS/MS to assay phenobarbital, valproic acid and carbamazepine in dried blood spots reported 3.8% and 4.1% as inter- and intraassay CV, respectively. Accuracy and precision in this study was reported as below 13% and 10%, respectively (Velghe et al., 2020). Two studies using FPIA to assay phenytoin reported the mean intra- and interassay CVs were 4.2% (interquartile range [IQR] 2.1-6.4) and 4.4% (IQR 3.4-5.4) respectively (Cai et al., 1993; Smit et al., 1999). Among studies that reported intra- and interday measures, one study that used both HPLC and GC to assay valproic acid reported the intra- and interday reproducibility ranging from 1.54 to 5.14% and 1.59 to 4.97%, respectively for HPLC with a precision within 5.1% and from 0.59 to 2.04% and 2.67 to 5.2%, respectively for GC with the precision within 5.5% (Ma et al., 2019). A different study that used HPLC to assay 10-hydroxycarbazepine reported the intra and interday precision as ranging from 0.58 to 16.67% (Yang et al., 2019) while another study that used HPLC-UV to assay carbamazepine (CBZ) reported the intra- and inter-day precision of CBZ and carbamazepine-epoxide (CBZ-E) as ranging from 0.56% to 6.14% (Wang et al., 2015a).
The inter- and intraday relative standard deviations of lamotrigine assays using HPLC reported in a different study were lower than 15% for all measured analytes while for valproic acid, the assay using GC had a coefficient of variation (C.V.) lower than 4% (Wang et al., 2015b). One study reported the inter- and intraday CV for lamotrigine assayed using HPLC as 6.57 ± 2.01% and 4.95 ± 1.87% precision (Brzaković et al., 2012) while another reported similar CV being less than 8% and less than 12% respectively, for phenobarbital in both cerebrospinal fluid and plasma assayed using reversed-phase HPLC-UV (Brzaković et al., 1997). In terms of quality control, duplicate determinations were the most commonly used quality control methods and were reported in four (28%) of these studies, two studies on lamotrigine and two on phenytoin.

Reference ASM ranges used during TDM
Eighteen (27.6%) of the included studies reported the ASM reference ranges used during TDM. Majority of these studies were from the African (38.8%) and European regions (30.0%) with the other regions contributing between 5 to 16.6% of the proportion. Overall, uniform reference ranges as defined by ILAE were used for phenytoin (10.0–20.0µg/ml), valproic acid (50.0–100.0µg/ml), lamotrigine (3.0–14.0µg/ml)

| Purpose of TDM | Side Effects N (%) | Adherence N (%) | Clinical Efficacy N (%) | Drug Interactions N (%) | Dose Titration N (%) | Method Validation N (%) | Genetic Variations N (%) |
|----------------|---------------------|----------------|------------------------|------------------------|---------------------|------------------------|------------------------|
| Overall (N=77) | 33 (42.8)           | 7 (9.0)        | 12 (15.5)              | 8 (10.3)               | 4 (5.1)             | 2 (2.5)                | 11 (14.2)              |
| Region
| Americas (N=1) | -                   | -               | 1 (100.0)              | -                      | -                   | -                      | -                      |
| Western Pacific (N=16) | 1 (6.2)          | 1 (6.2)        | 3 (18.7)               | 3 (18.7)               | 1 (6.2)             | 2 (12.5)               | 5 (31.2)               |
| Europe (N=35) | 22 (62.8)          | -               | 3 (8.5)                | 4 (11.4)               | 1 (2.8)             | -                      | 5 (14.2)               |
| Eastern Mediterranean (N=3) | -               | -               | 2 (66.6)              | -                      | -                   | -                      | 1 (33.3)               |
| African (N=23) | 10 (43.4)          | 6 (26.0)       | 3 (13.0)               | 1 (4.3)                | 2 (8.6)             | 1 (4.3)                | -                      |
| P value | >0.05               | 0.012           | 0.023                  | 0.528                  | 0.635               | 0.215                  | 0.025                  |
| Continent
| South America (N=1) | -               | -               | 1 (100.0)              | -                      | -                   | -                      | -                      |
| Asia (N=17) | 2 (11.7)           | 1 (5.8)        | 3 (17.6)               | 3 (17.6)               | 1 (5.8)             | 2 (11.7)               | 5 (29.4)               |
| Europe (N=35) | 22 (62.8)          | -               | 3 (8.5)                | 4 (11.4)               | 1 (2.8)             | -                      | 5 (14.2)               |
| Africa (N=25) | 10 (40.0)          | 6 (24.0)       | 5 (20.0)               | 1 (4.0)                | 2 (8.0)             | 1 (4.0)                | -                      |
| P value | 0.002               | 0.007           | 0.108                  | 0.416                  | 0.823               | 0.122                  | 0.020                  |
| Income
| Low-income (N=4) | 1 (25.0)          | 1 (25.0)        | 1 (25.0)               | -                      | -                   | 1 (25.0)               | -                      |
| Lower-middle-income (N=5) | -             | 1 (20.0)       | 2 (40.0)               | -                      | -                   | -                      | 2 (40.0)               |
| Upper-middle-income (N=69) | 32 (46.3)    | 5 (7.2)        | 9 (13.0)               | 8 (11.5)               | 4 (5.7)             | 2 (2.8)                | 9 (13.0)               |
| P value | 0.097               | 0.183           | 0.184                  | 1.000                  | 1.000               | 0.157                  | 0.226                  |
| Urbanization
| Urban (N=74) | 33 (44.5)          | 5 (6.7)        | 11 (14.8)              | 8 (10.8)               | 4 (5.4)             | 2 (2.7)                | 11 (14.8)              |
| Rural (N=7) | 1 (14.2)           | 3 (42.8)       | 2 (28.5)               | -                      | -                   | 1 (14.2)               | -                      |
| P value | 0.123               | 0.019           | 0.372                  | 0.469                  | 0.692               | 0.240                  | 0.345                  |
and phenobarbital (10.0–40.0 µg/ml) while the median reference range used for carbamazepine (4.0–11.8 µg/ml) varied from the set limits. Across the regions, similar findings were observed for carbamazepine while phenobarbital median reference ranges also varied (Table 7).

**Discussion**

To the best of our knowledge, this is the first systematic review to summarize evidence on therapeutic drug monitoring of ASMs in LMICs, including the methods used, reference ranges and purposes of conducting TDM. Majority of TDM was conducted in Europe compared to the rest of the world and in upper middle-income countries compared to low- and lower-middle-income countries, perhaps because of logistical burden involved. TDM for first generation ASMs was commonly reported overall while monitoring to assess unwanted ASM-related side effects was the most popular reason for conducting TDM. FPIA was the most common method used for TDM, consistent with its the lowest commercial quote compared to other equipment. Assay validation and quality control were reported in a small proportion of the studies with variability in the reporting of outcomes. Majority of the studies used the International League Against Epilepsy (ILAE)-recommended reference ranges (Patsalos et al., 2008) during TDM.

TDM by geographic regions and economic blocks

TDM appeared to be influenced by economic status and infrastructural status. For example, more than half (53%) of TDM was in Europe (as a WHO region or continent), where countries can afford TDM apparatus and machines compared to the rest of the world. Similarly, Turkey and Serbia, which are upper middle-income countries from Europe, were leading in TDM compared to other individual countries. Africa contributed a substantial proportion of TDM (27%), but economically developed countries in Africa were over-represented, for instance, South Africa, which was leading in Africa, contributed nearly 20% of all TDM. Indeed, over 87% of all TDM was in upper-middle-income countries, which are wealthier than low- and lower-middle-income countries. Therefore, there is need to increase investment on TDM in LMICs.

**Methods and techniques used in TDM**

FPIA was the most common technique used for TDM overall and in Africa and Europe compared to others such as HPLC. In Africa, this could be due to the relatively low cost of equipment (TDx system) based on commercially quoted

| Method of TDM (N = 50 studies) | Total N = 53 (%) | Estimated Cost of Equipment |
|--------------------------------|-----------------|----------------------------|
| Auto-analyzer                  | 3 (5.6)         | $14,000                    |
| Chemiluminescent Immunoassay  | 2 (3.7)         | Unknown                    |
| Enzyme Multiplied Immunoassay Technique | 3 (5.6) | $1600 (Cobas), $11,500 (Syva) |
| Fluorescence Polarization Immunoassay | 22 (41.5) | $1000 (TDx) |
| Gas Chromatography             | 2 (3.7)         | $79,500                    |
| High Performance Liquid Chromatography (HPLC) | 9 (16.9) | $21,500                    |
| HPLC with fluorescence detection | 1 (1.8) | $1799.99                   |
| HPLC with Ultraviolet detection (HPLC-UV) | 5 (9.4) | $2,695                     |
| Liquid Chromatography-Mass Spectrometry LC-MS | 3 (5.6) | $149,000                   |
| Reversed phase HPLC with UV detection | 1 (1.8) | $1100                      |
| Thin Layer Chromatography (TLC) | 1 (1.8)         | Unknown                    |
| TLC with fluorescence detection | 1 (1.8)         | Unknown                    |
Table 7. Median reference ranges used during TDM across the regions.

| ASM | Carbamazepine | Phenobarbital | Phenytoin | Valproic acid | Lamotrigine |
|-----|---------------|---------------|-----------|---------------|-------------|
|     | Lower limit (IQR) | Upper limit (IQR) | Lower limit (IQR) | Upper limit (IQR) | Lower limit (IQR) | Upper limit (IQR) | Lower limit (IQR) | Upper limit (IQR) | Lower limit (IQR) | Upper limit (IQR) |
| Overall N=18 | 4.0 (4.0-4.6) | 11.81 (10.0-12.0) | 10.0 (9.2-10.0) | 40.0 (32.4-40.0) | 10.0 (10.0-10.0) | 20.0 (20.0-20.0) | 50 (50.0-50.2) | 100.0 (100.0-100.0) | - | - |
| WHO Region | | | | | | | | | | |
| African N=7 | 4.0 (4.0-4.0) | 12 (11.8-12.0) | 10 (9.2-10.0) | 40 (25.5-40.0) | 10.0 (10.0-10.0) | 20.0 (20.0-20.0) | 50 (50.0-50.2) | 100.0 (100.0-100.0) | - | - |
| European N=6 | 4.0 (4.0-4.0) | 10.0 (10.0-10.0) | 12.5 (10.0-15.0) | 40.0 (40.0-40.0) | 10.0 (10.0-10.0) | 20.0 (20.0-20.0) | 50.0 (50.0-50.0) | 100.0 (100.0-100.0) | 3.0 (3.0-3.0) | 14.0 (14.0-14.0) |
| Western Pacific N=3 | - | - | - | - | 10.0 (10.0-10.0) | 20.0 (20.0-20.0) | 50.0 (50.0-50.0) | 100.0 (100.0-100.0) | - | - |
| Eastern Mediterranean N=1 | - | - | - | - | 10.0 (10.0-10.0) | 20.0 (20.0-20.0) | - | - | - | - |
| Americas N=1 | 4.6 (4.6-4.6) | 11.6 (11.6-11.6) | 9.2 (9.2-9.2) | 39.4 (39.4-39.4) | - | - | - | - | - | - |
prices and low sample volumes (Kang et al., 2012). Surprisingly, FPIA was equally popular in Europe, implying possible knowledge exchange and collaboration between the continents. HPLC, which is slightly more expensive in terms of equipment, was popular in Asian countries probably informed by efficiency or production and assemblage of these machines in those settings. The WHO recommends an estimation of cost per test based on equipment, reagents and consumables, personnel, facility and quality management but only commercially quoted equipment prices were available for comparison (World Health Organization, 2019). For the studies that reported assay validation and quality control during TDM, authors reported compliance with either international or national bioanalytical guidelines, but this could not be validated.

**Recommended therapeutic ranges of ASMs during TDM**

In 2008, a best practice guideline was developed by ILAE (Patsalos et al., 2008) to guide therapeutic drug monitoring of anti-seizure medications and interpretation of the ASM levels. Only a very small proportion of the studies reported the reference range used during the assays, mostly from Africa and Europe (the regions that formed the bulk of the included studies). Among the studies, all the reported reference ranges were within the ILAE recommendations except for variations in ranges used for phenobarbital and carbamazepine. These variations could be attribute to ranges used in three studies published before the publication of the guidelines in 2008. Notably, ILAE recommends use of the reference ranges as a benchmark upon which individual concentrations should be established to achieve the best clinical outcomes, and this should be emphasized in future studies in LMIC.

**Strengths and limitations**

This review outlines findings on TDM that are generalizable to LMICs, including middle-income countries in Europe, which may have easily been overlooked. The review followed standard procedures such as a PRISMA flow-diagram used to clearly elucidate the inclusion and exclusion criteria. For references ranges used during TDM of the various agents, we used the recommended ILAE guidelines to determine appropriateness.

However, this review has several limitations. Significant between-study variation was observed across the outcomes of interest and can be attributed to different study designs, methods of TDM, sample sizes, types of ASMs measured and the different populations with varying ethnicities in the included studies. Majority of the studies were from Eastern Europe, specifically Turkey and Serbia, which may not be representative of the poorer continents. In Africa, majority of the studies were from South Africa, an upper-middle-income country, which may not represent most low- and lower middle-income groups. Some studies were based on very small sample sizes that may not offer acceptable precision for comparing the outcomes of interest. The estimated cost of the various TDM methods was extrapolated from the cost of equipment as data required for laboratory costing was not reported in any of the studies. We excluded articles that were not written in English or did not have English translations meaning relevant articles and their outcomes may have been missed. Finally, it is difficult to rule out the possibility that there was some publication bias as a result of the likely lack of publication of negative results as well as other biases such as inherent methodological differences that we could not evaluate in the present analysis.

**Conclusions and public health implications**

There is evidence of TDM in LMICs, especially in upper-middle-income countries, possibly due to availability of resources and technical expertise. First generation ASMs were the most commonly monitored and TDM was mostly conducted for the purpose of assessing unwanted side effects and toxicities related to ASMs. FPIA was the most common method of TDM and likely to be the most affordable.

TDM is important in complementing best clinical practice by ensuring that prescribed ASM are safe and effective. This will likely improve adherence to treatment and encourage people with epilepsy to seek care thus, reducing the epilepsy treatment gap in LMIC. TDM would be most useful when initiating new treatment, switching therapies, adding on ASM, investigating toxicity related to ASM and in cases of refractory epilepsy. A targeted approach for implementation of monitoring would be most useful in resource-limited settings because of cost implications. Development of standard operating procedures for guiding TDM in LMIC and training of primary health care providers on appropriate prescription of ASMs in epilepsy should also be prioritised.

**Data availability**

**Underlying data**

All data underlying the results (except for individual participant data) are available as part of the article and no additional source data are required. For data pertaining to the individual participants in the included articles, this is deemed to be licensed under the authors of the primary studies.

**Reporting guidelines**

Harvard Dataverse: PRISMA checklist for ‘Therapeutic monitoring of anti-seizure medications in low- and middle-income countries: a systematic review’, https://doi.org/10.7910/DVN/JMBK4R (Odhiambo et al., 2021)

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC BY 4.0).
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Version 2

Reviewer Report 14 July 2023

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✔ Bridget MacDonald

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I think this article is fine now and deserves to be indexed.

I retain my concerns about what appears to be a belief that it's good to monitor drugs to see if there are side effects, as opposed to clinically assessing patients adequately and listening to them.

There is no comment on whether tests are done more if the system for reimbursing doctors rewards doctors to do more tests. I think this does influence therapeutic drug monitoring in some countries.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Epidemiology, epilepsy, general neurology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 25 January 2022

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Thank you for doing this study which is interesting and has value. I am commenting from the perspective of a neurologist with an interest in epilepsy and epidemiology with some experience of advising colleagues in low income countries. I am not in any way an expert in how therapeutic drug monitoring is carried out although I may use its results in clinical practice. Hence I will not comment on that aspect at all.

There are, for me, several issues that need to be clarified.

Most importantly this study cannot be read as a study of where and how much TDM is used in different places of the world. It is an account of who has written articles about TDM. It would be wrong to conflate these two separate things. The article should be altered to reflect this.

More specific issues: (I leave the citation from the paper in italics, my comments in normal text)

○ For instance, phenobarbital is associated with adverse effects including risk of withdrawal seizures, hyperactivity in children, risk of drug abuse and suicidal ideation (Nimaga et al., 2002), but it is the only ASM that is easily available and affordable in many LMICs (Chin, 2012). These adverse effects require monitoring but TDM is rarely done in LMICs due to limited infrastructure, resources and lack of technical expertise to operate equipment for mass spectrometry and immunoassays (Tomson et al., 2007).

This paragraph is somewhat cherry picking – using a citation to support the notion that phenobarbital is “bad”, when some evidence is contrary to that view e.g. when Deb Pal (Randomised controlled trial to assess acceptability of phenobarbital for childhood epilepsy in rural India. Lancet Volume 351, Issue 9095, 3 January 1998, Pages 19-23)1 looked into this in a well designed study in poor village children in India the outcomes are far more nuanced.

○ Routine monitoring allows the prescriber to check treatment efficacy while identifying adverse effects and drug interactions. Drug interactions are often complicated by pharmacogenetics, medication for comorbidities and patient adherence (Knezovic & Marzinke, 2018).

It seems to me that although this view is often heard what one needs is to discuss whether there is evidence demonstrating that therapeutic drug monitoring achieves the aim of better therapeutic effect with fewer side-effects better than good clinical practice without TDM. UK’s NICE guidelines, for example, does not advocate their use except under a limited number of circumstances because there is little evidence to support it. There is marked variance in the use of TDM across Europe which is better explained by tradition than research.

○ In Methods – what did you do to ascertain that articles written in languages other than English were picked up and then analysed? All search terms were in English and only 8 were identified as eligible but not analysed because they were not in English. One might, for example, expect American literature to have Spanish and Portuguese articles. Overall the non-English group seems low to me and so more comment on this needs to be made.
There is a need for more investment in comprehensive TDM in LMICs that incorporates dose titration of ASM using pharmacokinetics and pharmacodynamics modelling, for both first generation and second generation ASMs.

I am unsure that there is evidence that TBM achieves better clinical outcomes, it could be argued that the shortfall in treatment for people with epilepsy in resource poor settings would be made worse if unnecessary and expensive monitoring were advocated. It seems there is a better rationale for monitoring when introducing (novel) medications to new populations where the risk of marked differences in drug metabolism might be suspected (be that due to diet or genetics) as part of an "introduction of a new drug to new population" study rather than a monitoring system.

In all, this article is interesting and well written but I find the framing of the study not quite right. Additionally, it purports to measure something it does not. The text could be altered to reflect this without altering the value of the data and work that has gone into it.

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Are the rationale for, and objectives of, the Systematic Review clearly stated?
No

Are sufficient details of the methods and analysis provided to allow replication by others?
Yes

Is the statistical analysis and its interpretation appropriate?
Partly

Are the conclusions drawn adequately supported by the results presented in the review?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: epidemiology, epilepsy, general neurology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 09 May 2023
Mercy Odhiambo

Thank you for taking the time to review our article. Please see below our responses to your comments. Most importantly this study cannot be read as a study of where and how much...
TDM is used in different places of the world. It is an account of who has written articles about TDM. It would be wrong to conflate these two separate things. The article should be altered to reflect this.

Response: Thank you for your comment. We think that lack of published articles may not necessarily mean that TDM is not being conducted in these settings as TDM may be conducted in clinics and hospitals but not published if not conducted systematically or in the context of research. However, we have reworded the objective to emphasise this that this review was conducted to examine available evidence on TDM, purposes, and methods and ASM reference ranges used during monitoring in these settings.

More specific issues: (I leave the citation from the paper in italics, my comments in normal text) For instance, phenobarbital is associated with adverse effects including risk of withdrawal seizures, hyperactivity in children, risk of drug abuse and suicidal ideation (Nimaga et al., 2002), but it is the only ASM that is easily available and affordable in many LMICs (Chin, 2012). These adverse effects require monitoring but TDM is rarely done in LMICs due to limited infrastructure, resources and lack of technical expertise to operate equipment for mass spectrometry and immunoassays (Tomson et al., 2007). This paragraph is somewhat cherry picking – using a citation to support the notion that phenobarbital is “bad”, when some evidence is contrary to that view e.g. when Deb Pal (Randomised controlled trial to assess acceptability of phenobarbital for childhood epilepsy in rural India. Lancet Volume 351, Issue 9095, 3 January 1998, Pages 19-23) looked into this in a well designed study in poor village children in India the outcomes are far more nuanced.

Response: Thank you for the observation. Indeed, some studies show benefit while other show some adverse effects of most anti-seizure medication in use. We have re-written this section to comment on adverse events related to anti-seizure medications in general such as feelings of tiredness, stomach upset, dizziness, blurred vision, cognitive and behavioural problems (Kwan et al., 2013; Mutanana, Tsvere, & Chiweshe, 2020) as this remains a concern and justification for TDM.

Routine monitoring allows the prescriber to check treatment efficacy while identifying adverse effects and drug interactions. Drug interactions are often complicated by pharmacogenetics, medication for comorbidities and patient adherence (Knezevic & Marzinke, 2018). It seems to me that although this view is often heard what one needs is to discuss whether there is evidence demonstrating that therapeutic drug monitoring achieves the aim of better therapeutic effect with fewer side-effects better than good clinical practice without TDM. UK’s NICE guidelines, for example, does not advocate their use except under a limited number of circumstances because there is little evidence to support it. There is marked variance in the use of TDM across Europe which is better explained by tradition than research.

Response: Thank you for the observation. We have elaborated on the benefit of monitoring; emphasising the highly variable pharmacokinetic profiles of these ASM. For instance, phenytoin displays zero-order pharmacokinetics meaning it can easily accumulate and cause toxicity if not monitored while carbamazepine has a narrow therapeutic index meaning a slight change in its dose may be the difference between sub-clinical response and toxicity. These variations may be empirically assessed clinically but can only be ascertained through TDM.
In Methods – what did you do to ascertain that articles written in languages other than English were picked up and then analysed? All search terms were in English and only 8 were identified as eligible but not analysed because they were not in English. One might, for example, expect American literature to have Spanish and Portuguese articles. Overall the non-English group seems low to me and so more comment on this needs to be made.

**Response:** This is a good observation. Even though our entire search strategy was in English, we postulate that most non-English articles will have English translated titles and abstracts, so these would be picked up during the search. We have added this explanation in the search strategy section and also noted this as a limitation.

There is a need for more investment in comprehensive TDM in LMICs that incorporates dose titration of ASM using pharmacokinetics and pharmacodynamics modelling, for both first generation and second generation ASMs I am unsure that there is evidence that TBM achieves better clinical outcomes, it could be argued that the shortfall in treatment for people with epilepsy in resource poor settings would be made worse if unnecessary and expensive monitoring were advocated. It seems there is a better rationale for monitoring when introducing (novel) medications to new populations where the risk of marked differences in drug metabolism might be suspected (be that due to diet or genetics) as part of an "introduction of a new drug to new population" study rather than a monitoring system.

**Response:** Thank you for this observation. Indeed, monitoring can be cost-intensive, and we propose introducing monitoring using a targeted approach. It would be useful in establishing the efficacy and side effect profile of new drugs as suggested. Additionally, in cases of suspected drug-related adverse effects for patients on polytherapy, for instance, it would be difficult to identify the offending drug at first glance but TDM would offer a fairly quick resolution. We have re-framed the conclusion to reflect situations where TDM would be most useful in low-income settings and play a role in improving patient outcomes.

**Competing Interests:** No competing interests were disclosed.
Generally, the title and their work are interesting but I have some comments that will be seen by authors.

1. Title: There is a difference between seizure and epilepsy but you write antiseizure medication in your title but you are talking about epilepsy in your background and other sections so please revise your title and the terms antiseizure medications in your manuscript.

2. Abstract section: Method: Please add the search period and the software used for your analysis

3. In your main manuscript, you put the protocol registration before the result section. because you followed PRISMA guidelines, the protocol registration should be written just after method and before inclusion and exclusion criteria.

4. The quality of observational studies is assessed based on the Newcastle Ottawa quality assessment scale adapted for crossectional studies. So it is advisable to use that adapted assessment scale rather than adapting by yourself.

5. Where are your outcome measurements?

6. In your strength and limitation part of paragraph 2 line 16, you mentioned as there is publication bias. Where is the publication bias and how you assessed it?

Are the rationale for, and objectives of, the Systematic Review clearly stated?
Yes

Are sufficient details of the methods and analysis provided to allow replication by others?
Yes

Is the statistical analysis and its interpretation appropriate?
Yes

Are the conclusions drawn adequately supported by the results presented in the review?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: cancer

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
Mercy Odhiambo

Thank you for taking the time to review our article. Please see below our responses to your comments.

Title: There is a difference between seizure and epilepsy but you write antiseizure medication in your title but you are talking about epilepsy in your background and other sections so please revise your title and the terms antiseizure medications in your manuscript.

Response: anti-seizure medications (ASM) is the current recommended term, having been changed from the previously used term; anti-epileptic drugs. The reason for this change is because these ASM are known to suppress the seizure activity and do not affect the underlying disease i.e. that is epilepsy. This will allow communication between patients and physicians that symptomatic treatments such as ASM do not cure the disease and have to be taken at regular intervals to prevent recurrent seizures.

Abstract section: Method: Please add the search period and the software used for your analysis

Response: We have added the search period (initially on 8th November 2018 and updated on 31st December 2020) and the software used for analysis (Stata 13) in the abstract.

In your main manuscript, you put the protocol registration before the result section. because you followed PRISMA guidelines, the protocol registration should be written just after method and before inclusion and exclusion criteria.

Response: We have moved the protocol registration section to immediately after the title of the methods section.

The quality of observational studies is assessed based on the Newcastle Ottawa quality assessment scale adapted for cross sectional studies. So, it is advisable to use that adapted assessment scale rather than adapting by yourself.

Response: We used all the quality appraisal tools as obtained from the source and we did not adapt any by ourselves, but rather applied it to the aspects available in the searched studies. We have re-phrased the wording to reflect that we used the published NOS that has been adapted for cross-sectional and case-control studies.

Where are your outcome measurements?

Response: Thank you for this observation. We extracted data on the regional distribution of the included articles, type of ASM used, method of conducting TDM, validation procedures and references ranges. We have added the outcome measurements in the methods and they include the proportion of TDM by country, continent, WHO region and income level as a percentage, the proportions of ASM monitored by country, continent, WHO region, income level and classification as a percentage and the mean and median reference ranges used during monitoring by ASM.
In your strength and limitation part of paragraph 2 line 16, you mentioned that there is publication bias. Where is the publication bias and how you assessed it? **Response:** We postulate that they may be publication bias due to lack of publication of negative results as well as other biases due to methodological differences that we could not evaluate in the present analysis. We have added this statement to the text to clarify the same.

**Competing Interests:** No competing interests were disclosed.