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***Observational Study***

**Chronic antiepileptic drug use and functional network efficiency: A functional magnetic resonance imaging study**

van Veenendaal TM *et al*. Antiepileptic drugs and functional network efficiency

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# Abstract

***AIM***

To increase our insight in the neuronal mechanisms underlying cognitive side-effects of antiepileptic drug (AED) treatment.

***METHODS***

The relation between functional magnetic resonance-acquired brain network measures, AED use, and cognitive function was investigated. Three groups of patients with epilepsy with a different risk profile for developing cognitive side effects were included: A “low risk” category (lamotrigine or levetiracetam, *n* = 16), an “intermediate risk” category (carbamazepine, oxcarbazepine, phenytoin, or valproate, *n* = 34) and a “high risk” category (topiramate, *n* = 5). Brain connectivity was assessed using resting state functional magnetic resonance imaging and graph theoretical network analysis. The Computerized Visual Searching Task was used to measure central information processing speed, a common cognitive side effect of AED treatment.

***RESULTS***

Central information processing speed was lower in patients taking AEDs from the intermediate and high risk categories, compared with patients from the low risk category. The effect of risk category on global efficiency was significant (*P* < 0.05, ANCOVA), with a significantly higher global efficiency for patient from the low category compared with the high risk category (*P* < 0.05, *post-hoc* test). Risk category had no significant effect on the clustering coefficient (ANCOVA, *P* > 0.2). Also no significant associations between information processing speed and global efficiency or the clustering coefficient (linear regression analysis, *P* > 0.15) were observed.

***CONCLUSION***

Only the four patients taking topiramate show aberrant network measures, suggesting that alterations in functional brain network organization may be only subtle and measureable in patients with more severe cognitive side effects.

**Key words:** Antiepileptic drugs; Cognitive side effects; Brain networks; Resting state; Functional magnetic resonance imaging; Graph analysis

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**Core tip:** Slowed information processing is a commonly observed cognitive side-effect of antiepileptic drug (AED) treatment. We aimed to increase our insight in the neuronal mechanisms underlying this side-effect. Therefore, the relation between functional MR-acquired brain network measures, AED use, and cognitive function was investigated. No associations were found between information processing speed and graph measures, and only the four patients taking topiramate (with a high risk on cognitive side effects) showed aberrant network measures. The results suggest that alterations in functional brain network organization may be only subtle and measureable in patients with more severe cognitive side effects.

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# INTRODUCTION

Epilepsy is generally treated with antiepileptic drugs (AEDs). A persistent problem in AED treatment is the occurrence of adverse events among which cognitive side effects are commonly seen[1,2]. The cognitive side effects account for a high percentage of the disease burden[3] and lead to early drug discontinuation[4]. The prevalence and severity of the cognitive side effects varies among different AEDs. Several AEDs, such as topiramate, are associated with cognitive problems such as language deficit (anomia), while other AEDs such as lamotrigine seem to induce less cognitive side effects or even have activating effects[5]. Despite specific differences, a decreased central information processing speed is commonly observed among the different AEDs to some extent[2].

AEDs control epileptic seizures *via* several distinct mechanisms, such as enhancement of GABAergic inhibition, reduction of glutamatergic neurotransmission, or modulation of the voltage-gated ion channels[6]. Changes in brain metabolic processes also affect healthy brain activity, and are likely to be responsible for cognitive side effects[1]. Functional magnetic resonance imaging (fMRI) enables assessment of this brain activity, and can be employed to measure combined effects of different mechanism of action of AEDs[7]. Several fMRI studies have shown altered brain activity patterns in healthy participants[8] or patients with epilepsy[9,10] treated with AEDs. For instance, altered brain activity patterns appeared to be associated with language impairments when taking topiramate[11-13].

Cognitive functions are mediated by the concerted action of multiple and distributed brain regions. These brain regions show correlations of their fMRI time signals, which is commonly interpreted as functional connectivity. Collectively, these functional connections form a brain network, which can be analyzed and characterized using graph theoretical analysis. Brain networks appear to be efficient networks, characterized by a high functional segregation and integration, *i.e*., different brain regions form densely interconnected groups, enabling specialized information processing, and also rapid communication between distributed brain regions. Several graph measures are available to quantify these characteristics[14].

Cognitive performance has been associated with the efficiency of functional brain networks[15,16], while impaired functional brain networks have been associated with cognitive decline in epilepsy[17,18]. Furthermore, associations between drug load, cognition and graph measures were shown in one of these studies, although this was not the main focus of the current study[17]. Another study associated the use of carbamazepine with altered graph measures when compared with other AEDs, but did not investigate the relation with cognitive effects[19]. In the current study, we aim to test whether chronic use of AEDs, associated with a high risk for cognitive side-effects, affects functional resting-state network measures differently than long-term use of AEDs associated with milder cognitive side-effects. Furthermore, we will test whether functional resting-state network measures are associated with impaired cognitive functioning.

**MATERIALS AND METHODS**

## *Patients*

Three groups of patients with epilepsy were compared in this observational, cross-sectional study[20]. These groups were subdivided based on the AEDs that were being used, in accordance to Samarasekera *et al*[21]. The first group, the low risk category, consisted of patients using lamotrigine or levetiracetam. Patients taking carbamazepine, oxcarbazepine, phenytoin, or valproate were included in the intermediate risk category, while the high risk category comprised patients taking topiramate. Patients on polytherapy took at most two different AEDs and were categorized according to their AED associated with the greatest cognitive risk. By including patients with AEDs from the three risk groups, a range in slowing of information processing speed is set out for.

All patients were clinically diagnosed with localization-related epilepsy and aged between 18 and 70 years. The patients were recruited from our tertiary epilepsy referral center. Participants not eligible for MRI, because of metal implants, claustrophobia, or pregnancy, were excluded from this study. Furthermore, patients did not experience seizures at least 12 h prior to MRI. This study was approved by the local Medical Ethical Committee and all participants provided written informed consent.

## *Neuropsychological investigation*

Cognitive functioning was assessed by two neuropsychological tasks. The Computerized Visual Searching Task (CVST) was used to measure visual (complex) information processing speed[22]. Slowing of this central information processing speed is a common side effect of AEDs[2], and therefore the CVST is considered to be sensitive for treatment effects[23]. With the CVST, a centered grid is shown surrounded by 24 other grid patterns. Participants have to find the (only) grid identical to the centered one as fast as possible.

The Raven Standard Progressive Matrices was administered to assess global cognitive performance. This is a non-verbal reasoning test which gives an indication of fluid intelligence[24]. Previous studies suggested that intelligence stays relatively unaffected by AEDs[23].

## *Epilepsy severity*

As several epilepsy related characteristics might affect functional brain networks[25], a score was composed to account for these effects. This epilepsy severity score was assessed in all patients and compared between the different risk categories. Epilepsy severity was characterized using a summarized score between zero and seven, composed by the sum of subscores for seizure type (tonic-clonic: 1, other: 0), previous occurrence of status epilepticus (yes: 1, no: 0), seizure-related injury (yes: 1, no: 0) and seizure frequency (seizure free: 0, yearly: 1, monthly: 2, weekly: 3, daily: 4).

## *MRI data acquisition*

MRI data were acquired on a 3.0T MRI scanner equipped with an 8-channel head coil (Philips Achieva, Philips Medical Systems, Best, The Netherlands). The scanning protocol included resting-state functional MRI and a T1-weighted scan. Functional MRI data were acquired using whole-brain single-shot multi-slice echo planar imaging (EPI) sequence sensitive to the blood-oxygen-level-dependent (BOLD) effect (195 volumes, 32 slices, in-plane resolution 2 mm × 2 mm, 4 mm thick slices, repetition time 2000 ms, echo time 35 ms, flip angle: 90°, acquisition time: 7 min). A 3D T1-weighted scan was acquired for anatomic reference (voxel size 1 mm × 1 mm × 1 mm, repetition time 8.3 ms, echo time 4.8 ms, inversion time 1022 ms, 180 slices, flip angle 8°, acquisition time 6 min).

## *Data preprocessing*

Preprocessing of the functional images was performed using SPM8 (Wellcome Department of Cognitive Neurology, London, United Kingdom). The functional images were corrected for differences in slice timing and head movement, coregistered to the T1 image and spatially (FWHM 6 mm) and temporally filtered (band pass 0.01-0.1 Hz). The BOLD signal originating from the white matter and ventricles, which is assumed to reflect physiological noise[26], and the six translation and rotation parameters obtained from the motion correction were deregressed from the BOLD signal.

The T1-weighted scan was parcellated into 82 cortical and subcortical brain regions using FreeSurfer v5.1.0 (The General Hospital Corporation, Boston MA, United States). Subsequently, a connectivity matrix was created by calculating the Pearson’s correlation coefficient between the average (deregressed) BOLD time signal of each combination of two regions. Negative correlations were set to zero. The correlation values were thresholded, based on the average connectivity matrix, to obtain connectivity matrices with only the strongest connections. The number of included connections was varied, with sparsity levels ranging from 0 to 0.9 (0 is fully connected, whereas 1 indicates no connections).

## *Data analysis*

The Brain Connectivity Toolbox[14] was employed to compute graph measures for each individual connectivity matrix. The clustering coefficient and the characteristic path length are commonly used to characterize the functional segregation and integration, respectively. The clustering coefficient quantifies the fraction of a node’s neighbor that is also connected to each other. The characteristic path length is defined as the average shortest distance (the inverse correlation coefficient) between all pairs of nodes. As, in sparse networks, a single weak connection can result in a large, or even infinite average path lengths, global efficiency was computed instead of characteristic path length, which avoids this effect by using inverse path lengths[27]. One hundred null models of the connectivity matrices were computed by randomizing the connections of the original matrices, while preserving the degree and weight distribution[28]. The graph measures were divided by the mean global efficiency and clustering coefficient of these null models, providing a normalized global efficiency (*Eg*) and clustering coefficient (γ).

## *Statistical analysis*

To test whether the clustering coefficient and global efficiency differed between the risk categories, an analysis of covariance (ANCOVA) was applied with the graph measures as outcome, cognitive risk category as fixed factor and age as covariate. Associations with cognition were assessed with linear regression analysis, with CVST time as outcome, and *Eg* or γ, age, and the percentage corrects answers in the Raven test as independent variables. To assess whether these results were affected by confounders, these analyses were repeated with gender, epilepsy severity score, or drug load (ratio of prescribed daily dose to defined daily dose[29]) added to the regression analyses as additional covariates. All statistical analyses were performed in MATLAB (version R2012b). *P*-values lower than 0.05 were considered significant.

**RESULTS**

## *Patient characteristics*

In total, 58 patients were included in this study. Three of these patients did not finish the procedures due to claustrophobia, resulting in 16 patients taking AEDs from the low risk category, 34 taking AEDs from the intermediate risk category, and 5 taking high risk AEDs. The age and drug load were significantly higher in the intermediate risk category than in the low risk category (Table 1). Also the number of patients on polytherapy was significantly higher in the intermediate risk category compared with the low risk category, while the high and low risk categories significantly differed in number of patients on polytherapy. The risk categories did not differ in gender distribution, educational level, or epilepsy severity.

## *Neuropsychological assessment*

The results of the CVST and the Raven task are summarized in Table 2. The CVST reaction time was slower than the normal range (range: 7.3 to 30.8 s, while the mean ± SD was 10.3 ± 4.1 s in normal population[31]). A significant effect of risk category on CVST reaction time was observed, which remained significant when controlling for age, gender, and global cognitive level (*P* = 0.009, ANCOVA). Post-hoc tests showed significant differences in CVST between the low and intermediate risk category (*P* = 0.035, estimated adjusted mean difference 3.5 s), and between the low and high risk category (*P* = 0.004, adjusted mean difference 7.8 s). No significant differences were found between the percentage correct answers Raven scores of the different risk categories.

## *Network topology*

Of the 55 included patients, seven were excluded from further analysis: one patient was excluded because of excessive head motion (maximum head movement of 8.0 mm, while the maximum head movement was below 1.5 mm in all other patients), one because of a deeper large lesion mass, and five patients were excluded because of a failure to automatically parcellate the cortex, due to cortical abnormalities. The analysis was therefore performed on 48 patients: 15 patients taking AEDs from the low-risk category, 29 patients taking AEDs from the intermediate risk category and 4 patients taking the high risk medication. The maximum head displacement did not differ between the three risk categories.

The functional networks were fully connected and showed small-world characteristics within the sparsity range 0.32-0.66 (which was defined as γ/λ significantly larger than one, with γ the normalized clustering coefficient, and λ the normalized characteristic path length). Only the sparsity levels within this range were considered for further analyses. The ANCOVA test revealed significant effects of risk category on *Eg* at most sparsities within this sparsity range (Figure 1). *Post-hoc* tests showed a significantly higher *Eg* for patients from the low category (*n* = 14) compared with the high risk category (*n* = 4), and for patients from the intermediate category (*n* = 29) compared with the high risk category (*n* = 4). *Eg* or γ did not differ significantly between patients from the low and intermediate risk categories (*P* > 0.2 at all sparsity levels), and no significant associations were observed between γ or *Eg* and CVST time (*P* > 0.15 at all sparsity levels). Gender, epilepsy severity score, or drug load were not significantly associated with the γ, *Eg*, or CVST reaction time, and the results of these adjusted analyses were consistent with the results of the analyses without these additional covariates (< 10% change in effect size of the variable of interest).

# DISCUSSION

The current study investigated whether patients taking AEDs with a different risk for cognitive side-effects have different functional brain topologies. To this end, we included epilepsy patients with chronic AED treatment with different risk profiles, *i.e.*, a low risk category, intermediate-risk, and high risk category. Furthermore, we assessed whether cognitive problems, in terms of a decreased central information processing speed, could be associated with the functional brain organization.

A higher global efficiency was shown in patients taking TPM (*n* = 4, the high risk category), compared with patients taking the low (*n* = 14) and intermediate risk AEDs (*n* = 29). The directionality of this difference is strikingly, as this result seems to contradict the cognitive side effects of TPM. The global efficiency is suggested to be particularly important for more complex cognitive tasks, for which different brain areas are involved[32]. The “better” global efficiency in TPM users might however be interpreted as a compensatory mechanism, or could be explained by a “survivor effect”. As patients with side effects are more likely to switch to other AEDs, it is likely that these patients are less vulnerable for cognitive problems. The higher global efficiency in the high-risk group might therefore reflect a lower susceptibility for cognitive side effects of these patients[33,34]. However, these patients did have a lower processing speed compared with the other patients, which argues against this explanation and in favor of a compensatory mechanism.

No differences in graph measures were observed between the patients groups taking AEDs from the low and from the intermediate risk category. It is possible that the effects of TPM on brain organization are more pronounced compared with effects of other AEDs, but TPM can also have distinctive effects on brain organization. TPM is suggested to have a unique cognitive profile, with specific effects on verbal fluency. Moreover, it has multiple mechanisms of action, and both these mechanisms and its chemical structure differ from other AEDs[35].

Furthermore, no associations were found between processing speed and graph measures, in contrast to a previous study that showed not only associations between intellectual decline and a lowered clustering coefficient in patients with epilepsy, but also with increasing drug load[17]. The latter suggests that the intellectual decline (which was based on intelligence tests) was a side effect of the AED treatment, but this could also result from differences in epilepsy characteristics. That study included more patients with a high drug load (15% of the patients had a drug load higher than 3) than the current study (no drug loads higher than 3 in the included patients), thus it is possible that the effects on graph measures are only measureable in patients with higher drug loads or AEDs with high risks on cognitive complaints.

The measured information processing speed covered the whole range from normal to a clearly affected processing speed, and patients taking AEDs known to induce cognitive side effects, showed lower processing speeds than patients with lower risk AEDs. These results could therefore not explain the lack of associations between graph measures and information processing speed, or the lack of differences in graph measures between the low and intermediate risk category. Also no trends were shown, while the total number of participants (48), and the number of patients in the low (16) and intermediate risk categories (34) were relatively large, making it unlikely that this lack of findings were due to limited power.

All included patients in the current study were diagnosed with localization-related epilepsy. Epilepsy is associated with a decreased global efficiency and increased clustering coefficient, although some studies showed a decreased clustering coefficient in patients with epilepsy[36]. It is therefore plausible that the functional brain networks of all three groups of patients in this study were already altered compared with healthy participants, irrespective of AED treatment.

This study has several limitations. Although we tried to include comparable patient groups, the risk categories differed in age and drug load, suggesting that our study population is biased. Therefore, the analyses were corrected for these characteristics by including age and drug load as covariates. Besides these characteristics, also other factors could have confounded our results, such as the location of the epileptic focus or effects of AEDs on the neurovascular coupling, which should be assessed in separate studies[37]. Finally, no information is available about changes over time and causality due to the cross-sectional design.

No differences in functional network graph measures could be detected between patients with epilepsy after chronic use of AEDs with a different risks on cognitive side effects. Only the four patients taking TPM, which has a high risk for developing cognitive side effects, showed a more efficient brain network topology, which might be a compensatory mechanism. Also no associations were found between the graph measures and the measured cognitive impairments, specifically slowing of central information processing. Alterations in functional brain network organization may be only subtle and measureable in patients with more severe cognitive side-effects.

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**COMMENTS**

## *Background*

A persistent problem in antiepileptic drug (AED) treatment is the occurrence of adverse events among which cognitive side effects are commonly seen. The prevalence and severity of the cognitive side effects varies among different AEDs, but a decreased central information processing speed is commonly observed among the different AEDs to some extent.

## *Research frontiers*

Functional brain network analysis is being applied to study cognitive processes and cognitive problems.

## *Innovations and breakthroughs*

This is the first study that specifically assessed the relation between functional brain network measures and cognitive problems in patients taking different types of AEDs.

## *Applications*

To summarize the practical applications of your research findings, so that readers may understand the perspectives by which this study will affect the field and future research.

## *Terminology*

AED: Antiepileptic drug; Clustering coefficient: Network measure; indication of segregation of a network; Global efficiency: Network measure; indication of integration of a network; CVST: Computerized Visual Searching Task; measures visual (complex) information processing speed.

***Peer-review***

The manuscript is well writtern.

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**Table 1 Patient characteristics for the three risk categories1**

|  |  |  |  |
| --- | --- | --- | --- |
|  | Low risk (*n* = 16) | Intermediate risk(*n* = 34) | High risk(*n* = 5) |
| GeneralMale/female Age (yr)2Educational level3 | 5/11 (31%/69%)39.5 ± 13.45 (range 2-6) | 16/18 (47%/53%)50.7 ± 12.5a5 (range 2-7) | 0/5 (0%/100%)42.4 ± 15.85 (range 4-6) |
| Epilepsy-relatedSymptomatic/non-symptomatic epilepsySeizure frequencyWeeklyMonthlyYearlySeizure freeYears since epilepsy onset2Epilepsy severityscore2 | 2/14 (13/88%)04 (25%)2 (13%)10 (63%)22.7 ± 11.71.4 ± 0.8 | 15/19 (44/56%)1 (3%)3 (9%)6 (18%)24 (71%)30.4 ± 13.41.2 ± 1.0 | 0/5002 (40%)3 (60%)26.8 ± 23.31.0 ± 0.7 |
| AED-relatedMono-/polytherapyMedication typeCBZLEVLTGOXCPHTTPMVPADrug load2,4  | 16/007 (44%)9 (56%)00001.3 ± 0.6 | 8/26 (24/77%)a17 (50%)6 (18%)10 (29%)4 (12%)16 (47%)07 (21%)1.8 ± 0.7a | 3/2 (60/40%)c1 (20%)01 (20%)005 (100%)1 (20%)1.2 ± 1.0 |

Differences between the risk groups were tested using a Fisher’s exact test (gender, symptomatic epilepsy, number of different AEDs), a Mann-Whitney test (educational level, seizure frequency, epilepsy severity score), or a student’s *t*-test (all remaining variables). aIndicates significant differences between the low and intermediate risk category (*P* < 0.05); cIndicates differences between the low and high risk category (*P* < 0.05). 1Low risk: Lamotrigine (LTG), levetiracetam (LEV); Intermediate risk: Valproate (VPA), carbamazepine (CBZ), oxcarbazepine (OXC) and phenytoin (PHT); High risk: Topiramate (TPM); 2Mean ± SD; 3Median (range). Scores are according to Verhage[30], range 1 (did not finish primary school) to 7 (Master’s degree); 4The drug load is defined as the ratio of the prescribed daily dose to the defined daily dose[29]. AED: Antiepileptic drug.

**Table 2 Results of the neuropsychological investigation, represented as mean ± SD for each risk category**

|  |  |
| --- | --- |
|  | Cognitive test results |
| CVST (s)1 | Raven2 |
| Risk categoryLow riskIntermediate riskHigh risk | 11.5 ± 2.915.7 ± 6.420.2 ± 6.7 | 71.7% ± 10.3%73.2% ± 10.1%71.7% ­± 3.1% |
| *P*-value3 | 0.008 | 0.85 |

1Mean reaction time on the Computerized Visual Searching Task (CVST)[22]; 2Percentage correct answers on the Raven Standard Progressive Matrices[24]; 3Tested with ANOVA.



**Figure 1 Mean clustering coefficient (A** and **C) and global efficiency (B** and **D) for each risk category.** Both clustering coefficient and global efficiency are normalized, *i.e*., the measures are divided by the clustering coefficient and global efficiency of random networks. A and B show the graph measures as a function of sparsity, while B and D show the results at a single sparsity level. Error bars show standard deviations, while the asterisks indicate significant differences between the risk categories (*P* < 0.05, with age included as covariate).