Localization of Epileptogenic Zone With the Correction of Pathological Networks

Chuanzuo Yang¹, Guoming Luan²,³,⁴, Qian Wang²,³, Zhao Liu²,³, Feng Zhai²,³ and Qingyun Wang¹*

¹Department of Dynamics and Control, Beihang University, Beijing, China, ²Beijing Key Laboratory of Epilepsy, Sanbo Brain Hospital, Capital Medical University, Beijing, China, ³Department of Neurosurgery, Epilepsy Center, Sanbo Brain Hospital, Capital Medical University, Beijing, China, ⁴Beijing Institute for Brain Disorders, Beijing, China

Patients with focal drug-resistant epilepsy are potential candidates for surgery. Stereo-electroencephalograph (SEEG) is often considered as the "gold standard" to identify the epileptogenic zone (EZ) that accounts for the onset and propagation of epileptiform discharges. However, visual analysis of SEEG still prevails in clinical practice. In addition, epilepsy is increasingly understood to be the result of network disorder, but the specific organization of the epileptic network is still unclear. Therefore, it is necessary to quantitatively localize the EZ and investigate the nature of epileptogenic networks. In this study, intracranial recordings from 10 patients were analyzed through adaptive directed transfer function, and the out-degree of effective network was selected as the principal indicator to localize the epileptogenic area. Furthermore, a coupled neuronal population model was used to qualitatively simulate electrical activity in the brain. By removing individual populations, virtual surgery adjusting the network organization could be performed. Results suggested that the accuracy and detection rate of the EZ localization were 82.86 and 85.29%, respectively. In addition, the same stage shared a relatively stable connectivity pattern, while the patterns changed with transition to different processes. Meanwhile, eight cases of simulations indicated that networks in the ictal stage were more likely to generate rhythmic spikes. This indicated the existence of epileptogenic networks, which could enhance local excitability and facilitate synchronization. The removal of the EZ could correct these pathological networks and reduce the amount of spikes by at least 75%. This might be one reason why accurate resection could reduce or even suppress seizures. This study provides novel insights into epilepsy and surgical treatments from the network perspective.

Keywords: refractory focal epilepsy, stereo-electroencephalograph, epileptogenic zone localization, coupled neuronal population model, epileptogenic networks

INTRODUCTION

Focal epilepsy is characterized by paroxysmal abnormal discharge within local brain tissue, approximately 30% of which are medically refractory (1, 2). Drug-resistant epilepsy is defined as recurrent seizures despite adequate trials of at least two antiepileptic drugs (3, 4). For these patients, the surgical resection can be an effective way by which to achieve seizure freedom. However, the rather high rate of failure in epilepsy surgery underlines that the accurate localization of the epileptogenic area is still intractable. It is generally believed that the epileptogenic zone (EZ) acts as the driving hub of abnormal activity and sustains the seizures. Therefore, one of the purposes of research is to verify that causal analysis can assist in the localization. Furthermore, epilepsy is increasingly understood as the...
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algorithm based on the state space model is entirely different and can estimate the coefficient matrix at any time. The details of algorithm implementation were described in Arnold et al. (11). An example is presented in Figure 1, wherein the prediction error and the fluctuation of a coefficient are depicted.

At time $n$, the $p$ matrices jointly determine the causality relationships between different channels, and ADTF is then developed to integrate these into the frequency domain (25):

$$A(f, n) = I - \sum_{i=2}^{p} A_i(n) e^{-2\pi i n f},$$

$$ADTF_i(f, n) = |H_i(f, n)|^2,$$

where $u$ is the imaginary unit, $A(f, n)$ is the Fourier transformation of the coefficient matrix, and $H_i(f, n)$ is the element of $H(f, n)$, the inverse matrix of $A(f, n)$.

Since both the frequency and time can determine the value of $H_i(f, n)$, it may be more convenient to investigate the evolution with time if the information is integrated in a specific frequency band. Meanwhile, the ADTF is generally normalized with respect to the incoming flow, which leads to the spectrum-weighted ADTF (swADTF) (26):

$$\text{swADTF}_i(n) = \sum_{j \neq i}^{h} \sum_{k=1}^{K} \frac{|H_i(f, n)|^2}{|H_i(f', n)|^2} \sum_{i=1}^{K} \frac{|H_k(f, n)|^2}{|H_k(f', n)|^2}.$$

The swADTF is calculated in the frequency band (3, 45 Hz) to exclude the interference of background electrical activity and include the effect of low gamma rhythm simultaneously. Notably, the values of swADTF have a slight fluctuation even for stationary signals. Therefore, it is necessary and reasonable to take the average value every 0.5 s.

**Dynamic Effective Connectivity**

Effective connectivity is defined as the influence that one node exerts over another. In contrast to functional connectivity, it can describe the directionality of interactions and the path of information flow. The abnormal activity of the epileptogenic area can stimulate the connected neurons and cause sporadic or periodic spikes. The EZ often maintains robust causality relationships with neuronal populations. Therefore, the top $K$ non-diagonal elements of the causality matrix are selected to determine the dynamic threshold $\theta(n)$. The connection matrix $L$ is defined as follows:

$$\text{swADTF}_i(n) \geq \theta(n) \text{ and } i \neq j \to L_{ij}(n) = 1,$$

$$\text{swADTF}_i(n) \leq \theta(n) \text{ or } i = j \to L_{ii} = 0,$$

where $L_{ii}(n) = 1$ indicates that there exists an edge from node $j$ pointing to node $i$. The process of constructing dynamic effective connectivity is described in the left branch of Figure 2.

The out-degree is the number of outgoing edges from one node. A large out-degree means that this node dominated the network at that time. Its role is extremely similar to the impact of the EZ on other neural ensembles in the production of synchronous discharges. Therefore, the out-degree is selected as the principal indicator of the epileptic area.

**Coupled Neuronal Population Model**

In 2002, Wendling et al. proposed an exquisite single neuronal population model on the basis of previous research results (18, 27, 28). In this model, physiological insights have been considered as follows: (i) the non-uniform alteration of GABAergic inhibition in experimental epilepsy (29); (ii) the possible depression of GABA$_{ion}$ circuit activity by GABA$_{slow}$ inhibitory

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**TABLE 1 | Clinical patient characteristics.**

| Patient | Age (years) | Duration (years) | Side | Electrodes/contacts | Recorded seizures | Pathology |
|---------|-------------|------------------|------|---------------------|-------------------|------------|
| 1       | 16–20       | 12               | R    | 15/124              | 4                 | FCD Ia     |
| 2       | 6–10        | 5                | L    | 11/116              | 6                 | FCD Ib     |
| 3       | ≤5          | 7/12             | R    | 13/122              | 9                 | FCD Ila    |
| 4       | 6–10        | 1                | L    | 10/120              | 101               | FCD Ib + FCD Ic |
| 5       | ≤5          | 3                | R    | 10/108              | 2                 | FCD Ib     |
| 6       | 18–20       | 3                | L and R | 15/119          | 4                 | FCD Ib + FCD Ia |
| 7       | 30–35       | 22               | L and R | 9/126            | 9                 | HS         |
| 8       | 6–10        | 5                | L    | 13/116              | 1                 | FCD Ib     |
| 9       | 26–30       | 12               | L    | 8/108               | 5                 | HS         |
| 10      | 10–15       | 9                | R    | 8/117               | 17                | FCD Ib + GMH |

FCD, focal cortical dysplasia; HS, hippocampus sclerosis; GMH, gray matter heterotopia.
postsynaptic currents (30). This model includes three subsets of neurons, namely the main cells (e.g., pyramidal cells), the dendritic-projecting inhibitory interneurons (GABA A,slow receptors) and the somatic-projecting inhibitory interneurons (GABA A,fast receptors). Excitatory and inhibitory connections between subsets are depicted in Figure 3A, and their intrinsic mechanism is described in the form of block diagram (Figure 3B).

Neuronal populations are connected and cooperate to achieve specific functions. To account for this organization, a
coupled neuronal population model is established (Figure 4), and the connection strength is represented by the following equation:

\[
\begin{align*}
W_i(n) = \gamma \text{swADTF}_i(n) &; \ i \neq j \\
W_i = 0 &; \ i = j
\end{align*}
\]  

where \( \gamma \) is the amplification factor that neutralizes the impact of normalization. Its value depends on the size of the network, which generally takes 320–360.

According to the block diagram, the following set of 12 × K differential equations that describe the state of every population can be established:

\[
\begin{align*}
\dot{y}_i &= y_i \\
\dot{y}_i &= \text{EXC} \cdot aS(y_i - y_i - y_i) - 2ay_i - a^2y_i \\
\dot{y}_i &= \text{EXC} \cdot a \left[ p(t) + C_3S(C_3y_i) + \sum_{j=1, j \neq i}^K W_i y_j \right] - 2ay_i - a^2y_i \\
\dot{y}_i &= y_i \beta \\
\dot{y}_i &= \text{SDI} \cdot bC_4S(C_4y_i) - 2by_i - b^2y_i \\
\dot{y}_i &= y_i \\
\dot{y}_i &= \text{FSI} \cdot gC_5S(C_5y_i - C_5y_i) - 2gy_i - g^2y_i \\
\dot{y}_i &= y_i \\
\dot{y}_i &= \text{SDI} \cdot bS(C_6y_i) - 2by_i - b^2y_i \\
\dot{y}_i &= y_i \\
\dot{y}_i &= \text{EXC} \cdot aS(y_i - y_i - y_i) - 2ay_i - a^2y_i \\
\dot{y}_i &= y_i \\
\end{align*}
\]  

where \( p(t) \) is an impulse response function similar to \( h_{\text{EXC}}, h_{\text{SDI}}, \) and \( h_{\text{FSI}}. \)

\( | \text{Model parameters, interpretation, and standard values.} |

| Parameter | Interpretation | Standard value |
|-----------|---------------|----------------|
| EXC       | Excitatory synaptic gain | 3.25 mV |
| SDI       | Slow inhibitory synaptic gain | 22 mV |
| FSI       | Fast inhibitory synaptic gain | 20 mV |
| 1/a       | Time constant in the feedback excitatory loop | \( a = 100 \text{ s}^{-1} \) |
| 1/b       | Time constant in the slow feedback inhibitory loop | \( b = 50 \text{ s}^{-1} \) |
| 1/g       | Time constant in the fast feedback inhibitory loop | \( g = 500 \text{ s}^{-1} \) |
| 1/\( ab \) | Time constant associated to connections | \( ab = 30 \text{ s}^{-1} \) |
| \( C_1, C_2 \) | Average number of synaptic contacts in the slow feedback inhibitory loop | \( C_1 = C_2 \) |
| \( C_3, C_4 \) | Average number of synaptic contacts in the slow feedback inhibitory loop | \( C_3 = 0.25C \) |
| \( C_5, C_6 \) | Average number of synaptic contacts in the fast feedback inhibitory loop | \( C_5 = 0.3C \) |
| \( C_7 \) | Average number of synaptic contacts between slow and fast inhibitory interneurons | \( C_7 = 0.8C \) |

The parameters’ interpretation and standard values are listed in Table 2 (18, 28). In the process of simulation, it is assumed that all populations have the same parameters values so that only the relationship between seizures and networks is investigated.

It is also possible to perform virtual surgery and observe outcomes by removing the specified population from the model. For instance, if population \( u \) is selected for removal, the coupling matrix \( W(n) \) can be changed into \( W'(n) \)

\[
\begin{align*}
W_i(n) = W_i(n); i \neq u \quad \text{and} \quad j \neq u \\
W_i(n) = 0; i = u \quad \text{or} \quad j = u
\end{align*}
\]
Thus, it is easy to investigate the outcomes of different operation schemes, including the removal of the EZ based on theoretical calculation.

RESULTS

Localizing the EZ

The comparison between the out-degree calculation results and clinical conclusions is shown in Table 3. The electrode contacts with the maximum out-degree in various stages are regarded as the theoretical epileptogenic focus. SEEG reports accomplished by several experienced electrophysiologists indicate which channels are involved in the onset of epileptiform discharges. All of the epileptogenic channels in the SEEG reports are located in the resection region. Fine surgical outcomes can ensure that the clinical conclusions are credible enough to evaluate the theoretical results. In addition, the pre-ictal stage is combined with the ictal process since it has long been recognized as the sign of seizure onset (22).

The accuracy is the proportion of true positives to all calculated results. The detection rate measures corresponding percentage in clinical conclusions. They describe the influence of false positives and false negatives, respectively. According to Table 3, SEEG reports list 34 epileptogenic channels, while the calculated results show 35, 29 of which are actual positives, meaning that the accuracy is 82.86%, and the detection rate is 85.29%. In addition, the results from eight patients demonstrate that the driving hubs (the channels with the maximum out-degree) of the interictal activity overlap with those in the ictal stage. This implies that the cause of seizures may lurk in the interictal activity.

Particularly, the out-degree of each channel with time is depicted for patient No. 2 in Figure 5A. The stages are divided according to the features of the SEEG data and clinical symptoms. Although the salient characteristics of the brain networks are relatively stable within a certain stage, they differ from one stage to another. To clearly reveal the transient process, a compromise is struck. Only the dots whose out-degree is above 30 are marked.

Meanwhile, the networks in the same stage are summed and then averaged. The distribution of the mean out-degree is shown to interpret the difference of connectivity patterns (Figure 5B). In addition, the results are in good agreement during the two episodes.

It is shown that E04, the fourth contact of electrode E, governs the whole networks in both the interictal stage and the pre-IP. Similarly, J14 and M08 dominate activities in the ictal stage. The SEEG report verifies that intermittent spikes can be observed in E04 during the interictal period (IIP). Electrophysiologists deem that the abnormal discharges spread from E04 to electrode J and M. The calculated results are included within the resection area (Figures 5C,D). Fine surgical outcome also proves that this method is effective and accurate for localizing the EZ.

The Correction of Pathological Networks

Spikes are one of the most common biomarkers for indicating abnormal brain activity. In the simulation process, sporadic spikes are seen in the interictal stage and rhythmic spikes are observed during the IP (Figure 6B), which are consistent with clinical findings (Figure 6A). Virtual surgical resections are then carried out, and the outcomes of different operations are observed (Figures 6C,D). These simulations in three conditions (no resection, random resection, and removing the epileptogenic channel) are repeated 30 times, respectively. Compared with random resection, the removal of the EZ is more effective to reduce the amount of spikes (at least 75%) (Figure 6E). It is worth mentioning that the clinical resection range is wider than single channel, so sporadic spikes can be further eliminated.

However, two cases of simulations are not in accordance with the real situation. In Figure 6F, periodic spikes occupy almost the entire IIP, while there are only partial spikes during the ictal process. Nevertheless, the channels with the maximum out-degree in the interictal stage were also the targets for surgical resection. These spikes can be eliminated after removing the calculated source. Such interictal networks are also pathological and may well be the cause of seizures.

| TABLE 3 | Calculated results compared with clinical conclusions. |

| No. | Calculated result | Stereo-electroencephalograph report | Surgery region |
|---|---|---|---|
| | IIP | IP | IIP | IP | A.B. | P.B. | S.B. | I.B. |
| 1 | D02 | D11, D12 | D11 | D12 | MTG | RF. of CG. on L.Sur. | SMG. to IPS | BT |
| 2 | E04 | E04, J14, M08 | E04 | E04, J13, M05 | CS. | PreCS. | L.Sur. | CC. |
| 3 | P08, E07 | E07, M04 | P08, E07 | E07, G06 | 6 cm before PreCS. | PreCS. | L.Sur. | TF. |
| 4 | L11, H13, J11 | L09, H13, H13 | L11, H13 | L09, H13 | 2 gyri before PreCS. | PreCS. | L.Sur. | SF. |
| 5 | P08, D03 | D03 | P08, D03 | D03 | FP. | CSut. | L.Sur. | SF. |
| 6 | L04 | D04 | L04, H04 | D04 | FP. | CSut. | SFS. | SF. |
| 7 | E10 | C02, D01 | E10 | C02, D01 | Residual tissue | 5.5 cm to TP | Hippo. | R.G. |
| 8 | G10 | G10 | G10 | G10, H10 | 3 cm to TP | 6 cm to TP | HF. | SF. |
| 9 | A01, B01 | A01, B01, E03 | A01, B01 | A01, B01 | SAH. (amygdala + hippocampus head and 3 cm tail) | FP. | PreCS. | SF. |
| 10 | K09, G11 | K09 | K09, G11 | K09, G11 | | | | |

IIP: interictal period; IP: ictal period; A.B.: anterior boundary; P.B.: posterior boundary; S.B.: superior boundary; I.B.: inferior boundary; BT: basal surface of temporal lobe; CC.: corpus callosum; CG.: cuneus gyrus; CS.: central sulcus; CSut.: coronal suture; FP.: frontal pole; Hippo.: hippocampus; IPS.: inferior frontal sulcus; SF.: superior frontal sulcus; SMG.: superior marginal gyrus; STS.: superior temporal sulcus; TP.: temporal pole; TF.: tail of sylvian fissure.
DISCUSSION

This study developed a framework to analyze long-term SEEG data and localize the epileptogenic area. The electrode contacts with the highest total out-degree corresponded well with epileptogenic region defined by electrophysiologists. The same stage shared a relatively stable connectivity pattern, while the patterns varied from stages. Coupled neuronal population model was
established to investigate the presence of epileptogenic patterns. Simulations indicated that most networks during the IP were epileptogenic, and the nature of epileptogenic network was to enhance local excitability and facilitate synchronization. The fact that removing the EZ could significantly reduce the amount of spikes suggested that the EZ should be responsible for the formation of these pathological networks.

The Value of the Interictal Process
Epilepsy is regarded as a network-level phenomenon (32). This type of network disorder may have already existed before clinical seizures. Intermittent spikes during this period can be regarded as implications. Previous studies have demonstrated that interictal epileptic events were helpful for confirming the range of abnormal areas (33, 34). Wilke et al. concluded that the sources of interictal spike activity might be the generators underlying the clinical seizures (13). In Table 3, eight patients demonstrated that the sources of the interictal activity overlapped with the locations that the ictal activity originated from. In addition, some networks in the interictal stages exhibited epileptogenic behaviors and rhythmic spikes were observed in the simulations. This organization that is prone to producing spikes may push brain function to the edge of disorder. In other words, eliminating the interictal spikes or correcting these pathological networks should be part of epilepsy treatment.

Epileptogenic Networks
It is increasingly understood that seizures arise from epileptogenic networks, through which epileptiform discharges are produced and propagated (35). However, it is still unclear that which type of network can be considered as epileptogenic connectivity. Epileptogenic networks are not equal to the networks in the ictal stage. Some networks in the interictal stage were also epileptogenic. Varotto et al. reached a similar conclusion on the basis of SEEG recordings from patients with type II focal cortical dysplasia (36). Brain functional connectivity identified by the fMRI also showed an abnormal pattern in the absence of epileptiform discharges (37).

We have known that the channel with the maximum out-degree varies from stages. However, the occurrence of seizures is not only due to the transfer of important nodes. Qualitative change of connectivity patterns must follow. Coupled neuronal population model was used to simulate the time course in different patterns. An increased number of spikes indicate that this connectivity pattern is more likely to generate ictal activity. The nature of
epileptogenic network should be a specific organization which likely enhances local excitability and facilitates synchronization. Because the epileptogenic region affects other neural ensembles and enables them to synchronize, it should be responsible for the formation of these pathological networks. Virtual surgery also suggested that the removal of the EZ could significantly reduce the amount of spikes. Sinha et al. also concluded that the resection of these regions could reduce the overall likelihood of seizures after conducting simulations with a bi-stable model (21). Hence, it is helpful to explain why seizures are reduced or suppressed after accurate resection.

Model Simulation and Virtual Surgery
The distribution of the out-degree could aid identification of the EZ and deepen our understanding of the difference between connectivity patterns. However, it failed to describe the qualitative changes of network organization. Coupled neuronal population model was used to demonstrate the network behavior and judge whether the network was epileptogenic or not. It was also convenient to observe the outcomes of different operation schemes. There are two reasons to carry out virtual surgery. First, contradiction may appear when the EZ overlaps with the brain function area. Hypothetical surgery provides new options for controlling seizures with regard to preserving major function. Second, virtual surgery is beneficial in terms of balancing the resection range and therapeutic effect (38, 39). It is well known that a smaller resection region means lower physiological cost. By removing one or more channels, it is possible to assess the corresponding surgical effect, and the optimal scope can be determined. In a word, the treatment for refractory focal epilepsy should be a comprehensive and systematic project instead of just achieving seizure freedom.

The Relationships Between the Two Branches
This study consists of the EZ localization and coupled neuronal population model. The former aims at identifying the electrode contacts triggering the abnormal discharge. However, seizures are often associated with abnormal changes in brain synchronization mechanisms. This model is used to find epileptogenic connectivity patterns and evaluate the role of the EZ on pathological network. The results suggested that the presence of the EZ made the network more easily synchronized. Accordingly, if the EZ is removed, pathological network can be corrected and seizures are reduced or suppressed. It is helpful to understand the relationship between the EZ and epileptogenic networks.

CONCLUSION
It was shown that channels with the maximum out-degree corresponded to the epileptogenic area after applying the ADTF technique to SEEG recordings from 10 patients. All of these recordings from the interictal, pre-ictal and ictal stages were valuable for the EZ localization. Interestingly, we found that the same stage shared a relatively stable connectivity pattern, while the pattern evolved at different stages. In addition, some of these patterns were verified to be epileptogenic. Model simulations confirmed this and suggested that these pathological networks could be corrected if the EZ was removed. It indicated that the nature of the epileptogenic network should be a specific organization enhancing local excitability and facilitating synchronization.

ETHICS STATEMENT
This study was carried out in accordance with the recommendations of the Ethics Committee of the Sanbo Brain Hospital of Capital Medical University with written informed consent from all subjects. All subjects gave written informed consent in accordance with the Declaration of Helsinki. The protocol was approved by the Ethics Committee of the Sanbo Hospital of Capital Medical University.

AUTHOR CONTRIBUTIONS
CY, GL, QW, ZL, FZ, and QYW collected and analyzed these clinical recordings to localize the epileptogenic zone and wrote the paper. CY and QYW built mathematical model and carried out a series of simulations.

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REFERENCES
1. Souriut Z, Sghir A, Belkh R, Messouak O. Focal drug-resistant epilepsy: progress in care and barriers, a Moroccan perspective. J Clin Neurosci (2016) 34:276–80. doi:10.1016/j.jocn.2016.08.002
2. Pedersen M, Omidvarnia A, Curwood EK, Walz JM, Rayner G, Jackson GD. The dynamics of functional connectivity in neocortical focal epilepsy. Neuroimage Clin (2017) 15(C):209. doi:10.1016/j.nicl.2017.04.005
3. Kwan P, Arzimanoglu A, Berg AT, Brodie MJ, Allen HW, Mathern G, et al. Definition of drug resistant epilepsy: consensus proposal by the ad hoc task force of the ILAE commission on therapeutic strategies. Epilepsia (2010) 51(6):1069–77. doi:10.1111/j.1528-1167.2010.02744.x
4. Poole LS, Castagna CE, Williams S, Miller AB, Story TJ. Association between antiepileptic drug dose and long-term response in patients with refractory epilepsy. Epilepsy Behav (2017) 69:59–68. doi:10.1016/j.yebeh.2016.10.010
5. Hupalo M, Wojcik R, Jaskolski DJ. Intracranial video-EEG monitoring in presurgical evaluation of patients with refractory epilepsy. Neurol Neurochir Pol (2017) 51(3):201–7. doi:10.1016/j.pjnn.2017.02.002
6. Cosset M, Cardinale F, Colombo N, Mai R, Nobili L, Sartori I, et al. Stereoelectroencephalography in the presurgical evaluation of children with drug-resistant focal epilepsy. J Neurosurg (2005) 103(4 Suppl):333–43. doi:10.3171/ped.2005.103.4.0333
7. Harvey AS, Cross JH, Shinnar S, Mathern GW; ILAE Pediatric Epilepsy Surgery Survey Taskforce. Defining the spectrum of international practice in
assessment of significance. *Biol Cybern* (2001) 85(2):145–57. doi:10.1007/s004220000235

26. Mierlo PV, Carrette E, Halletz H, Raedt R, Meurs A, Vandenberghe S, et al. Lctal-onset localization through connectivity analysis of intracranial EEG signals in patients with refractory epilepsy. *Epilepsia* (2013) 54(8):1409–18. doi:10.1111/epi.12206

27. Jansen BH, Zouridakis G, Brandt ME. A neurophysiologically-based mathematical model of flash visual evoked potentials. *Biol Cybern* (1993) 68(3):275–83. doi:10.1007/BF00224863

28. Jansen BH, Rit VG. Electroencephalogram and visual evoked potential generation in a mathematical model of coupled cortical columns. *Biol Cybern* (1995) 73(4):357–66. doi:10.1007/BF00198091

29. Cossart R, Dutoitcourt C, Hirsch JC, Merchán-Pérez A, De Felipe J, Ben-Ari Y, et al. Dendritic but not somatic GABAeric inhibition is decreased in experimental epilepsy. *Nat Neurosci* (2001) 4(1):52–62. doi:10.1038/82900

30. White JA, Banks MI, Pearce RA, Kopell NJ. Networks of interneurons with fast and slow gamma-aminobutyric acid type A (GABAA) kinetics provide substrate for mixed gamma-theta rhythm. *Proc Natl Acad Sci U S A* (2000) 97(14):8128–33. doi:10.1073/pnas.100124097

31. Wendling F, Chauvel P. Transition to ictal activity in temporal lobe epilepsy: insights from macroscopic models. In: Soltesz I, Staley K, editor. *Computational Neuroscience in Epilepsy*. London: Academic Press, Elsevier (2008). p. 356–86.

32. Cleenert E, Premereur E, Casteels G, Goffin K, Janssens P, Paesschen WV. The effective connectivity of the seizure onset zone and ictal perfusion changes in amygdala kindled thesus monkeys. *Neuroimage* (2016) 12(12):252–61. doi:10.1016/j.neuroimage.2016.05.020

33. Alarcon G, Garcia Seoane JJ, Binnie CD, Martin Miguel MC, Juler J, Polkey CE, et al. Origin and propagation of interictal discharges in the acute electrocorticogram. Implications for pathophysiology and surgical treatment of temporal lobe epilepsy. *Brain* (1997) 120(12):2359–82. doi:10.1093/brain/120.12.2359

34. Hufnagel A, Dümppelmann M, Zentner J, Schijns O, Elger CE. Clinical relevance of quantified intracranial interictal spike activity in presurgical evaluation of epilepsy. *Epilepsia* (2000) 41(4):467–78. doi:10.1111/j.1528-1157.2000.tb00191.x

35. Korzeniewska A, Cervenka MC, Jouny CC, Perilla JR, Harezlak J, Bergey GK, et al. Ictal propagation of high frequency activity is recapitulated in interictal recordings: effective connectivity of epileptic networks recorded with intracranial EEG. *Neuroimage* (2014) 101:96–113. doi:10.1016/j.neuroimage.2014.06.078

36. Varotto G, Tassi L, Franceschetti S, Spreafico R, Panzica F. Epileptic networks of type II focal cortical dysplasia: a stereo-EEG study. *Neuroimage* (2012) 61(3):591–98. doi:10.1016/j.neuroimage.2012.03.090

37. Pittau F. Epileptogenic networks as identified by the fMRI. *Clin Neurophysiol* (2016) 127(3):e47. doi:10.1016/j.clinph.2015.11.152

38. Elskarkawy AE, Alabbi AA, Pannek H, Schulz R, Hoppe M, Palu G, et al. Outcome of frontal lobe epilepsy surgery in adults. *Epilepsie Res* (2008) 81(2–3):97–106. doi:10.1016/j.eplepsres.2008.04.017

39. Spencer S, Huh L. Outcomes of epilepsy surgery in adults and children. *Lancet Neurol* (2008) 7(6):525–37. doi:10.1016/S1474-4422(08)70109-1

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