The therapeutic mechanism of epilepsy seizures in different target areas: Research on a theoretical model

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Abstract.
BACKGROUND: The selection of optimal target areas in the surgical treatment of epilepsy is always a difficult problem in medicine.
OBJECTIVE: We employed a theoretical calculation model to explore the control mechanism of seizures by an external voltage stimulus acting in different nerve nuclei.
METHODS: Theoretical analysis and numerical simulation were combined.
RESULTS: The globus pallidus, excitatory pyramidal neurons, striatal D1 neurons, thalamic reticular nucleus and specific relay nuclei were selected, we analyzed that the electrical stimulation has different effects in these target areas.
CONCLUSIONS: The data selected were reasonable in study, the results may give a theoretical support for similar studies in clinical.

Keywords: Epilepsy, a computing network, voltage stimulation, nerve nuclei

1. Introduction

Deep brain stimulation technology has been proved to be effective in relieving seizures for intractable epilepsy patients [1,2]. Cortex and thalamus are the main seizure areas of epilepsy [3], and the subthalamic nucleus (STN) is a common stimulus target for treating seizures [4]. Basal ganglia (BG) as an important unit in the brain has closely input and output relationship with cortical and thalamic circuit [5–7]. Thus, in addition to the STN, other organizations in the BG may also be effective DBS targets in controlling seizures. Vuong and Devergnas reviewed the possible roles of the BG in regulating absence, neocortical seizures and temporal lobe [8], di Giacopo et al. observed that substantia nigra DBS can inhibit myoclonus for myoclonic seizures [9]. The electrophysiological and lesion results implied that substantia nigra pars reticulata (SNr) anterior can inhibit convulsant in seizures [10]. Guo et al. showed that SNr-DBS can effectively control seizure activities in rats [11]. SNr-DBS can also relieve amygdala-kindled seizures for a long time [12]. Chua et al. found that the quality of life can be improved with DBS of the ventral pallidum (VP) in rat models [13]. Yu et al. found that VP-DBS can reduce seizure activity and behavior in pilocarpine-treated rats [14]. Cheng et al. observed that low-frequency stimulation at the external globus

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palladium (GPe) can inhibit tonic-colonic generalized seizures and temporal lobe epilepsy [15]. However, the therapeutic mechanism and effect of these targets are different, the choice of the best stimulation target area is an important direction of clinical medicine research in the future, which need the support of theoretical research results. Also, many studies implied that striatum may involve in controlling epileptic seizures [16], however, if the striatum can be taken as an effective target in the DBS, it still needs a lot of theoretical evidences to support.

Abnormal interactions of cortex and thalamus circuits have been considered to be a key factor in inducing epilepsy seizures, recently, some studies have shown that the inhibition effect is also obvious as the external stimulation was exerted in the thalamocortical system directly [17,18]. Molnar et al. found that thalamic DBS may increase short-interval intracortical inhibition, which is similar to the treatment effect of antiepileptic drugs [19]. Lee et al. shown that using the DBS at the anterior thalamic nuclei can significant relief epileptic symptoms in the patient [20]. Son et al. observed that centromedian thalamic nucleus DBS is effective and safe in controlling refractory seizures [21]. Recently, Elhadd et al. found that generalised epilepsy can be successfully managed by thalamic DBS [22]. Grewal et al. shown that chronic subthreshold cortical stimulation may be effective for focal epilepsy [23]. Young et al. pointed out that the piriform cortex may relieve seizures in focal and generalized epilepsy [24]. Lundström et al. found that chronic subthreshold cortical stimulation can reduce the probability of seizures [25,26]. However, the control mechanism of seizures is complex, which should be intensively studied in the dynamic model.

Therefore, the mechanism of DBS in epilepsy is still unclear, the choice of the best treatment plan and region is a difficult problem to be solved. In this paper, we explored the treatment principle of absence epilepsy through exerting voltage stimulation in different neural nuclei of brain network. Different mechanisms were derived, which may be the theoretical basis for further research.

2. The calculation model and method

The coupling relation of different populations is presented in Fig. 1. The specific meaning and symbols of this model can be referred from [27–30]. The red lines and black lines represent excitatory and inhibitory inputs, respectively. The electrical stimulation was exerted in different targets, which were denoted in red.

The dynamic equations that described the network structure of the model were derived from previous studies [27–32]:

\[
\frac{d\phi_e(t)}{dt} = \dot{\phi}_e(t)
\]

\[
\frac{d\phi_c(t)}{dt} = \gamma_c^2 [-\phi_c(t) + F(V_e(t))] - 2\gamma_c \dot{\phi}_c(t)
\]

\[
\frac{dX(t)}{dt} = \dot{X}(t)
\]

\[
X(t) = [V_c(t), V_{d_1}(t), V_{d_2}(t), V_{p_1}(t), V_{p_2}(t), V_s(t), V_r(t), V_s(t)]^T
\]

\[
\frac{dV_e(t)}{dt} = \alpha \beta (\nu_{ee} \phi_c(t) + \nu_{ei} F(V_e(t)) + \nu_{es} F(V_s(t)) - V_e(t)) - (\alpha + \beta) \dot{V}_e(t)
\]

\[
\frac{dV_{d_1}(t)}{dt} = \alpha \beta (\nu_{d_1 e} \phi_c(t) + \nu_{d_1 d_1} F(V_{d_1}(t)) + \nu_{d_1 s} F(V_s(t)) - V_{d_1}(t)) - (\alpha + \beta) \dot{V}_{d_1}(t)
\]
Fig. 1. Schematic diagram of the calculation model used in this paper, the symbol and biological significance of which can be referred from [27–30].

\[
\begin{align*}
\frac{dV_{d_2}(t)}{dt} & = \alpha \beta (\nu_{d_2} \phi_e + \nu_{d_2} d_2 F (V_{d_2} (t)) + \nu_{d_2} s F (V_s (t)) - V_{d_2} (t)) - (\alpha + \beta) \dot{V}_{d_2} (t) \\
\frac{dV_{p_1}(t)}{dt} & = \alpha \beta (\nu_{p_1} d_1 F (V_{d_1} (t)) + \nu_{p_1} p_2 F (V_{p_2} (t)) + \nu_{p_1} \zeta F (V_{\zeta} (t)) - V_{p_1} (t)) - (\alpha + \beta) \dot{V}_{p_1} (t) \\
\frac{dV_{p_2}(t)}{dt} & = \alpha \beta (\nu_{p_2} d_2 F (V_{d_2} (t)) + \nu_{p_2} p_2 F (V_{p_2} (t)) + \nu_{p_2} \zeta F (V_{\zeta} (t)) - V_{p_2} (t)) - (\alpha + \beta) \dot{V}_{p_2} (t) \\
\frac{dV_{\zeta}(t)}{dt} & = \alpha \beta (\nu_{\zeta} \phi_e + \nu_{\zeta} p_2 F (V_{p_2} (t)) - V_{\zeta} (t) + \zeta) - (\alpha + \beta) \dot{V}_{\zeta} (t) \\
\frac{dV_{r}(t)}{dt} & = \alpha \beta (\nu_{r} e \phi_e + \nu_{r} p_1 F (V_{p_1} (t)) + \nu_{r} s F (V_s (t)) - V_{r} (t)) - (\alpha + \beta) \dot{V}_{r} (t) \\
\frac{dV_{s}(t)}{dt} & = \alpha \beta (\nu_{s} \phi_e + \nu_{s} p_1 F (V_{p_1} (t)) + \nu_{s} A F (V_s) + \nu_{s} B F (V_{r} (T - \tau)) - V_{s} (t) + \phi_s) \\
& \quad - (\alpha + \beta) \dot{V}_{s} (t)
\end{align*}
\]

The meaning of parameters in the above equations were listed in Table 1 [27–29]. Through the numerical calculation of the above equations [27,28], we got the main results of this paper. The default simulation data were derived from previous literature [27–32], which are in a reasonable range: \(\beta = 200 \text{ s}^{-1}, \alpha = 50 \text{ s}^{-1}, \tau = 50 \text{ ms}\); The unit in mV: \(\sigma = 6, \theta_{p_2} = 9, \theta_{d_1} = 19, \theta_{p_1} = \theta_e = 10, \theta_i = \theta_s = 15\); The unit in mV s: \(\nu_{ei} = \nu_{es} = 1.8, \nu_{p_2} = 0.45, \nu_{d_2} = 0.7, \nu_{p_1} = 0.3, \nu_{d_1} = 0.2, \phi_i = 2, \nu_{d_1} = 1, \nu_{p_1} = 0.03, \nu_{p_2} = 0.03, \nu_{p_1} = 0.035, \nu_{p_2} = 0.04, \nu_{r} = 0.5, \nu_{p_2} = 0.075, \nu_{d_2} = 0.05, \nu_{s} = 2.2\); The unit in Hz: \(Q_{p_2}^{\text{max}} = 300, Q_{d_2}^{\text{max}} = 45, Q_{s}^{\text{max}} = 500, Q_{p_1}^{\text{max}} = Q_{r}^{\text{max}} = Q_{e}^{\text{max}}, Q_{i}^{\text{max}} = 250\).
Table 1
Description of the parameters in the calculation equations [27–29]

| Parameter | Meaning | Parameter | Meaning |
|-----------|---------|-----------|---------|
| $Q_{e}^{\text{max}}$ | Maximum cortical discharge rate | $\theta_{e}, \theta_{i}$ | Average discharge threshold (ADT) of the cortical nuclei |
| $Q_{d1}^{\text{max}}, Q_{d2}^{\text{max}}$ | Maximum striatal discharge rate | $\theta_{d1}, \theta_{d2}$ | The ADT of the striatum |
| $Q_{\text{SN}}^{\text{max}}$ | Maximum STN discharge rate | $\theta_{\text{SN}}$ | The ADT of the SNR |
| $Q_{\text{GPe}}^{\text{max}}$ | Maximum GPe discharge rate | $\theta_{\text{GPe}}$ | The ADT of the TRN |
| $Q_{\text{STN}}^{\text{max}}$ | Maximum TRN discharge rate | $\theta_{\text{STN}}$ | The ADT of the STN |
| $Q_{\text{SRN}}^{\text{max}}$ | Maximum SRN discharge rate | $\theta_{\text{SRN}}$ | The ADT of the GPe |
| $Q_{\text{SNr}}^{\text{max}}$ | Maximum SNr discharge rate | $\theta_{\text{SNr}}$ | The ADT of the SNr |
| $Q_{\text{GPe}}^{\text{max}}$ | Maximum GPe discharge rate | $\theta_{\text{GPe}}$ | The ADT of the GPe |
| $\tau$ | The delay induced by the receptors $\text{GABA}_{B}$ | $\alpha$ | The decay time constant of synaptodendritic |
| $\beta$ | The rising time constant of synaptodendritic | $\gamma_{e}$ | The damping rate in cortex |
| $\sigma$ | Change in discharge rate threshold | $\phi_{n}$ | The nonspecific subthalamic input |
| $\nu_{ab}$ | The connection weight from the population “$b$” to the population “$a$” |

Fig. 2. The stimulation $V$ was acted in GPi, EPN, GPe, SRN, TRN and striatum D1 neurons, respectively. The AE may be well relieved by tuning $V$. Here, we set $\nu_{GPe} = 0.05$ and $-\nu_{SRN} = 1$.

3. Main results

In this section, we explored the control mechanism of seizures by a small voltage $V$ acting in different nuclei. The simulation results were presented in Fig. 2, which is a bifurcation diagram of different states in the EPN. The symbol A represents the low firing state, the B is the simple periodic discharge state, the C is the seizure state and D represents the saturation firing state. The specific meaning and characteristics of these four states can be referred from [27–29]. $KK$ is a proportional coefficient between “GPi $\rightarrow$ SRN” and “GPi $\rightarrow$ TRN”, i.e., $\nu_{GPi} = kK \nu_{SP}$ [27]. Figure 2a was obtained by exerting the V in the GPi. In this
model, we inferred that the “GPi → SRN” affected the EPN by the projection “GPi → SRN → EPN”, so the $\nu_{sp}$ exerted an inhibitory role in EPN. The “GPi → TRN” affected the EPN by the projection “GPi → TRN → SRN → EPN”, so the $\nu_{rp}$, exerted an excitatory role in EPN. As defined above, the effect of $\nu_{rp}$ strengthened with increasing in $KK$. Therefore, an increase in $KK$ promoted the discharge of EPN, as indicated in Fig. 2a. Therefore, when $KK$ was small, the effect of $\nu_{sp}$ was major, and the state C was inhibited by transferring into the A. As $KK$ increased to large enough, the role of $\nu_{rp}$ strengthened, and the C was pushed into the D. Figure 2b describes the regulation effect via the V stimulating in the EPN directly. We noticed that the V is positive, which improved the firing ability of EPN. So, when the stimulus intensity increased to a certain value, the C disappeared and was pushed into the B or the D. Figure 2c was simulated by putting the V in GPe. Here, we taken $KK = 0.01$, which was small. From the above analysis, we known that the GPe mainly affected the EPN by the projection “GPe → GPi → SRN → EPN”, which was generally excited. Therefore, the C can be pushed into the B by reducing the V, as implied by the arrow. Figure 2d is a state transition process of the B and the C as the V acting in SRN. Because SRN exerted a direct excitatory input to the EPN, seizures were changed to the B when V was small. Therefore, we inferred that the inhibitory stimulation in SRN may be more conducive to the control of seizures. In the contrary, when the V was putted in TRN, it gave an inhibitory effect to EPN through the projection “TRN → SRN → EPN”. Thus, the C was inhibited with an increase in V, as clearly presented in Fig. 2e. Striatum D1 neurons exerted inhibitory projections to GPi. The V in GPe and striatum D1 neurons have similar mechanisms, which can be inferred from Fig. 1. Therefore, transfer processes in Fig. 2e and f are similar, both were simulated as $KK$ is small.

4. Conclusion

Although there are many reports on the experiment about various potential target areas for the treatment of epilepsy, the theoretical mechanism research based on the model is limited. In this paper, we exerted a stimulation voltage V in the thalamus, GPi, GPe, striatum and EPN respectively to explore the adjustment mechanism of seizures in a dynamic calculation model. We found that positive voltages were suitable for different targets and the stimulus intensity needed to inhibit the seizure was also different. For example, we observed that the intensities needed to inhibit the seizures in SRN and TRN were smaller than that in GPi and EPN. The decrease in V can lead the seizures in Fig. 2c and f to disappear. The parameter values in the model have some influences on the control effect. For example, as shown in Fig. 2a, the excitatory stimulation protocol was effective, but the seizure was transferred to the state A when $KK$ was small, and the seizure was pushed into the state D when $KK$ increased to large enough. As indicated in Fig. 2c and f, the stimulus protocol may be suitable in GPe and striatum D1 when $KK$ was small. The minimum voltage intensity needed to inhibit seizures in SRN and TRN may be relative small as implied in Fig. 2d and e. These different dynamic mechanisms may give theoretical basis for the choice of the optimal scheme and precise treatment in the experiment.

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Conflict of interest

None to report.

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