Review Article

Role of Thalamus in Recovery of Traumatic Brain Injury

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ABSTRACT

Degree of recovery after traumatic brain injury is highly variable that lasts for many weeks to months. The evidence of brain structures involved in recovery mechanisms is limited. This review highlights evidence of the brain structure particularly thalamus in neuroplasticity mechanism. Thalamus with its complex global networking has potential role in refining the cortical and other brain structures. Thalamic nuclei activation both naturally or by neurorehabilitation in injured brain can enhance and facilitate the improvement of posttraumatic symptoms. This review provides evidence from literature that thalamus plays a key role in recovery mechanism after injury. The study also emphasize that thalamus should be specifically targeted in neurorehabilitation following brain injury.

KEYWORDS: Anisotropy, neuroplasticity, posttraumatic symptoms, recovery, thalamus, traumatic brain injury

INTRODUCTION

Neuroplasticity following traumatic brain injury

T

raumatic brain injury (TBI) is a critical public and medical health problem, which causes morbidity and mortality, especially among productive age group. Although TBI is commonly used all over, no two brain injuries are same in any aspect. TBI is heterogeneous in manifestation that may be due to mechanism of injury, force by which patient’s brain was hit, environmental factors, and patient factors (age, genetic, etc). Posttraumatic symptoms can be compared to spectrum of symptoms that affect daily activities of life. As such, the degree of recovery is highly variable, with some individuals regaining complete or near complete functions while others are disabled. There is uncertainty in recovery mechanism following brain injury.1-4 In any severity of brain injury, there is substantial spontaneous recovery that occurs over weeks to months after injury. The human brain is capable of widespread structural plasticity leading to restoration of function. There is experimentation that has demonstrated many anatomical and physiological examples of neuroplasticity.5-6 During this neuroplasticity phase, few brain parts play a potential role in enhancing and facilitating recovery process.3 Among them, thalamus, a subcortical brain structure with its global network connections plays a key role.7,8

METHODS AND DISCUSSION

Thalamic connections and its functions in neuroplasticity following traumatic brain injury

The thalamus is a deep-seated subcortical structure, that is, situated in the forebrain above the midbrain. The thalamic nuclei play an important role in coordinating and maintaining consciousness in human beings.9-11 A positron emission tomographic study on vegetative state, TBI patients showed that there was a significant difference in modulation between thalamic nuclei, prefrontal, and anterior cingulate cortices as compared with healthy controls. After recovery, the thalamocortical modulation was same as controls. The study supports that thalamocortical connections are associated in the maintenance of consciousness.9 Thalamic nuclei receive ascending fibers from brain stem/basal forebrain, that is, arousal system that regulates cortical activities. The same thalamic neurons are innervated from descending fibers from cortices. Collectively, the nuclei of thalamus appear to modulate the level of arousal and alertness that is important to carry out cognitive task and maintain wakeful state.11-13

The thalamus with its complex global network connections has anatomical and physiological specialization that supports its key role in consciousness,9 arousal,11 cognition,14 behavior,15 working memory,16 executive function,11 motor control,17 sustained, and vigilent attention.18 The recent microcircuit proposes that central thalamic neurons inhibit GABAergic pallidal neurons which in turn have tonic inhibition on central thalamic neurons, therefore, creating positive feedback loop. The thalamocortical and thalamostriatal show progressive deafferentation in proportion to severity of structural brain injuries, causing disinhibition of GABAergic pallidal neurons which in turn cause persistent tonic inhibition of central thalamic nuclei as a whole the thalamic network connection decreases.10,19,20 The concept of central thalamus

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deep brain stimulation (CT-DBS) is to activate thalamostriatial and thalamocortical network connections to regulate and facilitate thalamic function. Thalamic nuclei with its ascending and descending fibers regulate and facilitate many physiological functions. Few studies have evidenced that thalamus by natural means,[7,8,21] rehabilitation means,[22‑24] and by invasive stimulation methods[8] have crucial role in recovery mechanism. However, the exact evidence and mechanism for thalamic global dynamic network are not understood in detail.

**Preclinical and histopathological studies evidence of thalamus with its network connection in recovery of traumatic brain injuries**

In rodents, electrical stimulation of central thalamus is linked with activation of cortical, striatal, hippocampal neural circuits. This network activation is associated with improvements in learning, working memory, arousal system, and waking of rat from asleep.[10,23,26] Recovery of somatosensory function among rats was associated with intact myelinated fibers connecting thalamic nucleus and somatosensory cortex as was evidenced by blood-oxygen-level-dependent response of magnetic resonance imaging (MRI) scan.[27] TBI rats treated with allopregnanolone had shown better performance in spatial performance task; allopregnanolone, a metabolite of progesterone, has shown to enhance gamma-aminobutyric acid neurotransmission in mediodorsal nucleus of thalamus.[28]

Electrophysiological studies of alert monkeys have shown that central thalamus is directly involved in regulating firing pattern for performing a task and increases in firing pattern while performing task.[20] In a histopathological study, one patient with diffuse axonal injuries (DAI) and transneuronal degeneration with no structural damage to thalamus had survived for more months as compared to other DAI patients.[30] Reason for the same is not known.

**Imaging evidence: Thalamic role in recovery of cognitive domains, postconcussion symptoms, pain, and behavioral functions following brain injuries**

There are very few studies that have provided evidence of brain structures and connections that are involved in recovery following brain injuries [Table 1].

**Direct evidence**

Thirteen mild TBI patients at 4 months showed that thalamic functional connectivity (FC) was normalizing to baseline as compared to impaired 6 weeks thalamic FC. The decrease in patient’s pain severity and postconcussion symptoms was significantly associated with normalization of thalamic dorsal attention network FC at 4 months.[31] A three-time-point imaging study on mild TBI patients hypothesized that increase in mean thalamic volume from baseline and appearance of fractional anisotropy of thalamocortical connection after 3–4 months of injury was significantly associated with improvements in cognitive domains.[31] Two moderate TBI pediatric patients after 9 months of injury had poor cognitive performance (<5 percentile), moderately disabled with 7–8 concussive symptoms. The patients were subjected to 20 sessions of electroencephalogram (EEG) neurofeedback therapy during which pre- and post-MRI scan and clinical parameters were documented. The study reported that emergence of fractional anisotropy value of thalamocortical connections had significant association with clinical improvements.[18]

A study on single severe TBI patient reported progress from coma (day 8) to minimally conscious state (day 44) to posttraumatic confusional state (day 198) to fully community integration (day 366) was significantly associated with an increase in apparent diffusion coefficient and decrease in anisotropy value of thalamus.[22] In 2007, a severe TBI patient with minimal conscious state was subjected to CT-DBS, 6.5 years after injury. The patient showed improvements in behavioral, arousal, and motor function, that is, exclusively due to broad recruitment of frontal-striatal resource with thalamic stimulation.[15]

**Indirect evidence**

A single severe TBI patient with persistent right-sided sensory motor deficit and difficulty in maintaining both static/dynamic balance, 43 months following injury was given intermittent theta burst stimulation to cerebellum and neuromotor training for 3 weeks. The patient showed a significant improvement in right-sided limb sensation and balance while walking. The improvement was due to enforcement of cerebello-thalamo-cerebral circuit.[24]

In a study, the diffusion-tensor tractography revealed that the ascending reticular activating systems (connects pons and thalamus) tract volume and fractional anisotropy value increased significantly at 40 months as compared to 4 months was associated with patient Glasgow coma scale score improvement from 3 to 15.[23] In mild TBIs, without any intervention, majority of patients recovers where thalamus plays a major role by means of adaptive neuroplasticity.[9,21,31]

In moderate to severe TBIs, specific neurorehabilitation helps in recovery,[15,22,24] where thalamus has a significant role to play. The thalamus with its broad connections has a potential part in recovery of insult brain.[6,9,15,21‑24,31] Therefore, thalamus has a key role in posttraumatic neuroplasticity.

**Future directions**

One of the challenges in TBI neuroplasticity is lack of quantitative evidence. If there was an objective metric that could be tracked over time in any severity of injury, TBI patients would be benefited. The longitudinal studies in all severities of brain injuries should focus and evaluate thalamus and its network’s specific role in postinjury phase. These studies may give hint to by what threshold of preserved thalamic connections will enhance and facilitate recovery process and what specific rehabilitation and what time to be given. TBI neurorehabilitation should mainly focus on thalamic nuclei activation. For example, in mild to moderate, the noninvasive neurorehabilitation can be initiated like EEG neurofeedback therapy that specifically activates thalamic rhythm. Among severe TBI with minimal conscious state, CT-DBS mode may be beneficial.
Conclusion
Not much is known about the basics of neuroplasticity of TBI and brain structures involved in it. Our review provides evidence that thalamus is naturally involved in recovery process as in mild TBIs. Few studies from available knowledge on pathophysiological process and brain structures involved in brain injury recovery mechanisms are targeting thalamus by means of activating thalamic nuclei by DBS, a specific neurorehabilitation that targets in activating rhythm of thalamus and by metabolites that targets and activates thalamus. By activating thalamus, it refines global network connections as a result there are significant improvements in posttraumatic symptoms. Postinjury neuroplasticity may be coordinated by many brain structures; however, among them, we emphasize that thalamus plays a major key role.

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Conflicts of interest
There are no conflicts of interest.

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Table 1: Evidence of role of thalamus in improvement of posttraumatic symptoms

| Authors and year | Severities of TBI (n) | Intervention | Summary of improvements |
|------------------|----------------------|--------------|-------------------------|
| Direct           |                      |              |                         |
| Banks et al., 2016[21] | Mild (13) | No | Thalamic FC ↓ |
| Munivenkatappa et al., 2015[31] | Mild (21) | No | Thalamic DAN- FC ↓ was associated with ↓ in pain and postconcussive symptoms scores |
| Munivenkatappa et al., 2014[8] | Moderate (2) | EEG-neurofeedback therapy | Thalamic volume ↑ |
| Munivenkatappa et al., 2014[8] | Moderate (2) | EEG-neurofeedback therapy | Thalamocortical FA ↑ |
| Munivenkatappa et al., 2014[8] | Moderate (2) | EEG-neurofeedback therapy | Thalamus diffusivity values ↓ was associated with ↑ in memory scores |
| Munivenkatappa et al., 2014[8] | Moderate (2) | EEG-neurofeedback therapy | Thalamic diffusivity values ↓ was associated with ↑ attention scores |
| Edlow et al., 2013[22] | Severe (1) | Comprehensive neurorehabilitation | Cortical gray matter volume ↑ |
| Edlow et al., 2013[22] | Severe (1) | Comprehensive neurorehabilitation | FA ↑ of white matter tracts |
| Schiff et al., 2007[15] | Severe (1) | CT-DBS | Thalamus ADC ↑ and FA ↓ |
| Indirect          |                      |              |                         |
| Martino Cinnera et al., 2016[24] | Severe (1) | rTMS+neurorehabilitation | The improvement in interactive behavior, arousal level, and motor control |
| Jang et al., 2015[23] | Severe (1) | No | Emergence of new gait patterns was due to activation of cerebello-thalamo-cerebral circuit |

FC: Functional connectivity, DAN: Dorsal attention network, FA: Fractional anisotropy, ARAS: Ascending reticular activating system, GCS: Glasgow coma scale, EEG: Electroencephalogram, CT-DBS: Central thalamus deep brain stimulation, rTMS: Repetitive transcranial magnetic stimulation, TBI: Traumatic brain injury, ↓: Decrease, ↑: Increase
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