Case Report: Thalamic Stroke Patient with Neuropsychiatric Symptoms

Ishwari Jadhav*, Kunal Doshi

Maharashtra Medical Council

Abstract

Vascular dementia presents a complex clinical picture when compounded by rare vascular events of such as thalamic stroke. This case report highlights the pivotal role of early psychiatric consultation and neuropsychological testing in the management of vascular dementia, particularly in the context of a thalamic stroke. A 81-year-old male presented with cognitive decline, depressive symptoms, and functional impairment following a vascular stroke. Prompt psychiatric evaluation and comprehensive neuropsychological assessment revealed concurrent vascular dementia and geriatric depression along with localization of vascular stroke. Early intervention with tailored pharmacotherapy targeting both cognitive and mood symptoms yielded significant improvement in cognitive function, mood, management of delirium and overall functional status. Moreover, the incorporation of psychiatric rehabilitation strategies facilitated enhanced recovery and long-term prognosis. This case underscores the critical importance of multidisciplinary collaboration involving psychiatrists, neurologists, and rehabilitation specialists in the holistic management of vascular dementia. Early identification of psychiatric symptoms through thorough evaluation and timely intervention not only improves short-term outcomes but also holds promise for mitigating long-term disability and optimizing the quality of life in patients with vascular dementia and associated thalamic stroke. Further research is warranted to delineate optimal treatment approaches and rehabilitation strategies in this vulnerable population.

ARTICLE INFO

*Correspondence:

Ishwari Jadhav ishwari.jadhav.ij@gmail. com Maharashtra Medical Council

Dates:

Received: 14-02-2024 Accepted: 28-04-2024 Published: 05-06-2024

Keywords:

Cognitive impairment, Stroke, Geriatric, Delerium, Psychiatrist referral thalamic.

How to Cite:

Jadhav I, Doshi K.
Case Report: Thalamic
Stroke Patient with
Neuropsychiatric
Symptoms. Indian
Journal of Clinical
Psychiatry.2024;4(1):108-111.
doi:10.54169/ijocp.v4i01.117

INTRODUCTION

The thalamus is a complex structure acting as a relay for various tracts to convey Information and mediate action. The reticular activating system also forms a part of the thalamus and, affects arousal attention and mediates voluntary movement. Thalamic strokes, although rare, constitute 3% of all CNS strokes with a telling symptom of excessive daytime somnolence and impaired arousal.¹ It can present with vegetative symptoms and can present as geriatric depression and sometimes even with delerium.² This case report highlights the clinical presentation of thalamic stroke in the form of avolition and neurological negative emotionalism with a peculiar quality of depression with themes of identity loss that were not typical of a simple mood episode with negative symptoms. We recorded a case of thalamic stroke, which presented with

© IJOCP, 2024. Open Access This article is licensed under a Creative Commons Attribution-NonCommercial-ShareAlike 4.0 International (CC BY-NC-SA 4.0) License, which allows users to download and share the article for non-commercial purposes, so long as the article is reproduced in the whole without changes, and the original authorship is acknowledged. If you remix, transform, or build upon the material, you must distribute your contributions under the same license as the original. If your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit https://creativecommons.org/licenses/by-nc-sa/4.0/

severe depression and delirium and was detected and treated early due to an early psychiatric referral.

Case Report

Mr. J. D., an 81 year old retired male, previously working as a medical practitioner, presented with the following complaints of cognitive and emotional changes that were acute in onset and with progressive decline in functioning, including prominent forgetfulness and naming difficulty. The patient and relative complained of excess daytime drowsiness with a total sleep duration amounting to 21 hours. The patient reported of severe lack of energy and sadness of mood experienced for most of the day for most days in the last three months. Relatives reported poor self-care, with the patient needing assistance for all activities of daily living. The patient voiced repeated passive death wishes with a constant theme of loss of sense of meaningful existence. On tracking the progress of the illness, we uncovered a previous similar episode with the deterioration that could be graphed as a progressive step ladder pattern of deteriorating illness over the last four years. The patient was a known case of chronic dermatological illness of psoriasis. The patient lived with his wife and had no history of substance use. Patient had a past psychiatric history of moderate depressive episodes without psychotic symptoms, which occurred six years ago and lasted for nine months but a treatment history was not available.

Mr. J.D. presented to the neurology clinic with complaints of cognitive decline and emotional disturbances following a recent stroke. He reported the sudden onset of symptoms approximately six weeks ago, characterized by weakness on his left side and difficulty speaking. The patient needed assistance with all activities of daily living. On clinical assessment, there was evidence of a right trigeminal nerve dermatome with herpes lesions. Magnetic resonance imaging (MRI) of the brain revealed an acute infarct in the right medial and inferior thalamus region and lacunar infarcts in the periventricular region. The patient's neurological examination revealed Mr. J.D. presented with left-sided hemiparesis and mild dysarthria. Cranial nerve examination was within normal limits except for the right facial nerve, with Ramsey Hunt syndrome evident. Sensory examination showed decreased sensation to pinprick and light touch on the left side. Reflexes were brisk on the right side, with an extensor plantar response. The cognitive assessment revealed deficits in attention, executive function, and memory, and hemispatial neglect on the clock drawing test. Mood assessment demonstrated symptoms of depression, including low mood, anhedonia, and feelings of worthlessness, along with severe apathy and avolition.

Diagnostic evaluation was done as follows with an MRI brain showing an acute infarct involving the right thalamus, consistent with a lacunar stroke. Laboratory tests done were routine blood tests including complete blood count, electrolytes, renal function, and thyroid function tests, which were within normal limits.

Neuropsychological Evaluation

Formal neuropsychological testing revealed deficits in attention, executive function, and verbal memory consistent with thalamic involvement. Clock drawing test showed visuospatial impairment and hemineglect on left side with crowding of numbers on right side.

Diagnosis

As per ICD 11code 6D81, dementia due to cerebrovascular disease and 6E62 Secondary mood syndrome- boundary with dementia. [Mr. J.D. was diagnosed with a thalamic stroke resulting in cognitive impairment and neuropsychiatric symptoms, including depression (vascular dementia, poststroke depression).

Management was as follows, with medical management for Mr. J.D., he was started on antiplatelet therapy (aspirin) and statin therapy to reduce the risk of further cardiovascular events. His blood pressure and lipid profile were optimized. Dermatology referral and management of herpes. For physical therapy, he was referred to a rehabilitation program for physical and occupational therapy to improve motor function and activities of daily living. For neuropsychiatric treatment, a multidisciplinary approach involving neurology and psychiatry was employed. Mr. J.D. was started on an antidepressant (selective serotonin reuptake inhibitor) for the treatment of depression. Transient delirium was managed with low-dose antipsychotics along with non-pharmacological

management with daily schedule and activity planning. Multivitamin injections with age-appropriate nutrition management were done. Psychoeducation and supportive therapy were provided to address emotional adjustment to stroke-related disability. For cognitive rehabilitation, strategies focusing on attention, memory, and executive function were implemented to help mitigate cognitive deficits. While admitted for the duration of one week, Mr. J.D. was scheduled for regular follow-up appointments with neurology, psychiatry, and rehabilitation services. He demonstrated gradual improvement in motor function and cognitive symptoms over the subsequent week. His mood symptoms also showed significant improvement with pharmacotherapy and supportive therapy.

DISCUSSION

Thalamic strokes can result in a wide range of cognitive and neuropsychiatric symptoms due to the thalamus's role in relaying sensory and motor signals and its involvement in cognitive and emotional processing. ^{1,2} Cognitive deficits commonly include attention, memory, and executive dysfunction, while neuropsychiatric symptoms such as depression can significantly impact functional outcomes. Multidisciplinary management involving neurology, psychiatry, rehabilitation, and supportive services is essential for optimizing outcomes in patients with thalamic stroke. The patient showed 50% improvement at the time of discharge.

The thalamus consists mostly of grey matter but also has white matter structures composed of external and internal laminate, which cover the lateral surface of the thalamus while dividing the thalamic nuclear into many divisions.3 Functional division of the thalamus includes reticular and intralaminar nuclei (arousal and pain), sensory nuclei, effector nuclei, associative nuclei (cognition), limbic nuclei (mood and motivation). The circuit of the reticular activating system is responsible for arousal, attention, sleep-wake transitions, and circadian rhythm. Within the RAS:The cortical-hippocampal-cortical circuit is responsible for sleepy-dependant memory consolidation while basal forebrain circuit is responsible for REM sleep, cortical activation, and attention. The prefrontal-amygdala circuit for sleep-related

emotional reactivity, and attention. It was clinically evident in our patient that the excessive daytime sleepiness amounting to over 20 hours of sleep and with a score of 22 on the Epworth sleepiness scale, with the above symptoms that we were looking at geriatric depression, which was actually a result of CVA, probably affecting thalamus and was masking the vascular event. During wakefulness, the RAS originates in the brainstem and activates the thalamus and cortex *via* well defined "Bottom -up "pathway; however, another "Top-down" pathway includes projections for the sapience network of the neocortex; amygdala, and hypothalamus.⁴ Hypersomnolence is a typical feature seen in 80% of anterior strokes and 20% of posterior thalamic strokes.

Sleep and Neuroplasticity

Deep sleep characterized by sleep spindles, are important for neuroplasticity. Changes in the sleep micro architecture following Thalamic stroke could underline cognitive deficits like thalamic amnesia and dementia. Thalamic dementia typically caused by anterior and paramedian thalamic strokes. Characterised by temporal and spatial disorientation, behavioural changes: Apathy or agitation, impaired executive functions: Working memory, cognitive flexibility, abstract thinking, attention deficits. Association of depression and anxiety is higher in anterior thalamic lesions; and behavioral disturbances and delirium is associated with paramedian stroke of the thalamus.

Thalamic stroke has shown to have not complete recovery along with reduced neuroplasticity related to changed sleep. Patients with incomplete recovery need high amount of sleep immediately and even after one year of the thalamic stroke.8 This is most pronounced in anterior strokes. We learn from this that along with the psychiatric assessment and neurological assessment that guided the treatment, it is important to pay as much importance to the descriptive quality of the emotion that the patient experiences. Assessment of states of arousal was thoroughly done which led us towards the clinical diagnosis of a thalamic stroke and prompted apt treatment, however, the need for better rehabilitation services was felt with an urgency in the tribal region of Palghar. Although patient recovered symptomatically after a week of indoor admission, the stigma about the depressive illness led to the discontinuation of treatment one week after discharge. In

CONCLUSION

This case highlights the complex interplay between neurological and psychiatric symptoms following a thalamic stroke. It cannot be stressed enough that early psychiatric referral and treatment initiation can change the treatment outcome and improve the quality of life of the patient significantly. A comprehensive approach addressing medical, rehabilitative, and neuropsychiatric needs is crucial for maximizing functional recovery and quality of life in these patients. Long-term monitoring and support are essential to address ongoing cognitive and emotional challenges.

DECLARATION OF FUNDING

Nil

CONFLICT OF INTEREST

Nil

REFERENCES

 Kumral E, Evyapan D, Kutluhan S. Pure thalamic infarctions: clinical findings. Journal of Stroke and Cerebro-

- vascular Diseases. 2000 Nov 1;9(6):287-97. https://doi.org/10.1053/jscd.2000.18741.
- 2. Amici S. Thalamic infarcts and hemorrhages. (2012) Frontiers of neurology and neuroscience. 30: 132-6. Doi:10.1159/000333611
- Savage LM, Sweet AJ, Castillo R, Langlais PJ. The effects of lesions to thalamic lateral internal medullary lamina and posterior nuclei on learning, memory and habituation in the rat. Behav Brain Res. 1997 Jan;82(2):133-47.
- Filchenko I, Bassetti CLA, Gutierrez Herrera C. Thalamic Stroke: An Opportunity to Study the Brain Mechanisms of Sleep–Wake and Cognition. Clinical and Translational Neuroscience. 2023; 7(4):36. https://doi.org/10.3390/ ctn7040036
- Warburton, E.C.; Aggleton, J.P. Differential deficits in the Morris water maze following cytotoxic lesions of the anterior thalamus and fornix transection. Behav. Brain Res. 1999, 98, 27–38. [Google Scholar] [CrossRef] [PubMed]
- 6. Frohardt, R.J.; Bassett, J.P.; Taube, J.S. Path integration and lesions within the head direction cell circuit: Comparison between the roles of the anterodorsal thalamus and dorsal tegmental nucleus. Behav. Neurosci. 2006, 120, 135–149.
- Bogousslavsky, J et al. "Manic delirium and frontal-like syndrome with paramedian infarction of the right thalamus." Journal of neurology, neurosurgery, and psychiatry vol. 51,1 (1988): 116-9. Doi:10.1136/jnnp.51.1.116
- 8. Scharf, A.C.; Gronewold, J.; Todica, O.; Moenninghoff, C.; Doeppner, T.R.; de Haan, B.; Bassetti, C.L.; Hermann, D.M. Evolution of Neuropsychological Deficits in First-Ever Isolated Ischemic Thalamic Stroke and Their Association with Stroke Topography: A Case-Control Study. Stroke 2022, 53, 1904–1914.