VASCULAR DEMENTIA BY THALAMIC STRATEGIC INFARCT

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Strokes in privileged sites of the cognitive and behavioral circuits constitute strategic infarct vascular dementia (VaD) subtypes (SIVaD)1-3. Among such eloquent regions the thalamic nuclei are important targets4-6. Such lesions are not common, and those involving thalamus and basal ganglia represent 4.6% of VaD cases4-6. These strategic thalamic nuclei play an important integrative role and interconnects with several cortical areas (hippocampal, prefrontal). Amnesia, executive dysfunction and dysphasia are common clinical presentations, besides behavioral and psychological symptoms4-6. The thalamic nuclei may be classified in five functional classes and include associative and limbic ones. Branches from the posterior cerebral circulation supply different nuclear groups and intrathalamic tract fibers. The syndromes may be classified considering functional characteristics of the damaged nuclei and the affected vascular territories7. Such reports are rare in Brazil, and only two papers were published focusing subcortical structures lesions, including the thalamus.

In the present report a typical case of thalamic SIVaD is described, highlighting the anatomical substrates related to the clinical manifestations10-11. The study on VaD was approved by the Ethics Committee of IPUB/UFRJ and signed consent was obtained from the patient and the responsible relative.

CASE

The patient is a 54-years-old female, right handed, with 9 years of education. She was on treatment for chronic arterial hypertension, dyslipidemia and depression. The caregiver reported that in January 2006 she woke up at night looking for her son, who was sleeping in the next room, without finding him. In the morning she presented amnesia for this episode associated to dysphasia, incoherent discourse and memory impairment. This episode was diagnosed as a stroke by the patient's general practitioner and treatment was instituted. The patient was assessed for the first time ten months after this stroke in the outpatient ambulatory of the CDA-UFRJ. She reported memory decline and apathy in the last three months. There were no motor or sensorial impairment, and mild expression dysphasia was found on the neurological examination. The neuropsychological assessment showed Mini-Mental State Examination (MMSE)3=20/30 (with impairment in orientation for time, attention and recall); Cambridge Cognitive Examination (CAMCOG)3=66/107 (performance was worse in short and long term memory [10/27], abstraction [4/8], and attention [3/7], with mild to moderate impairment in other domains, as orientation [7/10], language [25/30], praxis [8/12], perception [7/11]. An abbreviated Boston naming test=2 errors. The frontal functions were assessed with a modified Porteus mazes test=normal performance; Trail Making Test (TMT) part A (1min47seg) (percentile <10) and part B (10min47seg) (percentile <10), both very slowed, mainly part B; semantic (animals) verbal fluency test (VF)=9 items/min. The BPSD assessment was performed with the Neuropsychiatric Inventory (NPI)=17 points (mainly depression, apathy and mild anxiety) and the Cornell Depression Scale=8 points (mainly anhedonia, loss of energy and pessimism). Pfeffer's Functional Activities Questionnaire=20/30. Ischaemic score of Hachinski=12 points. Clinical Dementia Rating scale – CDR=2.

The neuroimaging study with brain magnetic resonance (MR) showed a thalamic infarct in the left anteromedial region involving parts of the anteromedial and medial dorsal nuclei, and possibly the mamilothalamic tract and the ventral amygdalofugal path, vascular territory of the tuberohalamic and/or paramedian artery, and a small infarct in the right paramedian thalamus. The hippocampi showed bilateral preserved volume and signal (Fig 1). The hippocampal and posterior cingulum MR proton spectroscopy did not show significant changes. Cerebral perfusion with SPECT revealed poor fixation of the radiopharmaceutical on frontal lobes bilaterally, larger on the left side, and on temporal (anterior pole), posterior parietal and basal ganglia on left side (Fig 2). EEG was normal.

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DISCUSSION

Thalamic SILVaD may include several cognitive manifestations affecting memory and other functional domains depending on the involved nuclei and its connections. The main nuclei related to the mnemonic deficits are the anterior group (medial, lateral) and the medial dorsal nucleus that maintain close connections with the hippocampus, as well as with prefrontal areas, and receive their vascular supply through the tuberoinfundibular and/or paraventricular arteries. Other case reports emphasize hippocampal related symptoms with the occurrence of lesions of the anterior thalamic nuclei, considering their connections through the mammillothalamic tract.

The present case showed learning deficiency and short-term memory impairment (episodic and fixation, verbal recall, the visual recall being mildly affected) associated to temporal disorientation (revealed by MMSE and CAMCOG), compatible with hippocampal pathways related symptoms, and mild impairment of long-term memory. Disconnection of part of these pathways that cross the internal medullary lamina causes in addition an interruption of efferent amygdala fibers to the medial dorsal nucleus, intensifying the amnesic symptoms due to impairment of semantic and autobiographic (long-term) memory. These symptoms may also occur in cases where anterior nuclei are affected, being more severe in cases of paramedian thalamic SILVaD.

The interruption of thalamic-frontal connections (pertaining to the basal ganglia-thalamus-prefrontal circuits) is related to executive dysfunction symptoms as revealed by the TMT, particularly part B, with very slow performance, impulsivity, as well as sequencing and inattention errors. Attention impairment was also expressive (shown by MMSE and CAMCOG). There was also poor spontaneous speech and low VF performance related to executive dysfunction.

A low performance in items related to abstract thinking was also found. There was additionally mild motor transcortical aphasia, with preserved repetition, consistent with the reports found in the literature on speech disorders in connection with left anterior nucleus infarct.
Mood disorder with depression and severe apathy, as revealed by NPI and Cornell are frequent symptoms of limbic circuit lesions as seen in the present case. Pfeffer’s functional scale indicated moderate dependency. CDR characterized moderate dementia, stable since the stroke occurrence. MRI clearly showed the lesion, particularly the T2 weighted images, a peculiarity observed in thalamic infarcts. SPECT showed hypoperfusion in the frontal region, larger on the left side which can be considered as a probable consequence of thalamic-frontal disconnection.

The correlation of clinical, neuroimage, and chronological findings allowed for the diagnosis of a probable thalamic SIVaD, according to NINDS-AIREN criteria. Strategic infarcts show a peculiar characteristic of causing dementia in an acute way, and of remaining permanently once established at the time of the ischemic stroke. Well restricted thalamic lesions show this aspect in a clear fashion, as the sole cause for the appearance of the dementia picture, as illustrated by this case.

In conclusion, the present case represents a typical SIVaD, with memory impairment, attention deficit and executive dysfunction as the main symptoms. These manifestations are clearly related to an anteromedian region thalamic lesion, as shown by MRI, and of thalamic-frontal disconnection, as shown by SPECT images.

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