Frontal Lobe Dysfunction Following Infarction of the Left-Sided Medial Thalamus

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- We treated a 62-year-old woman who developed a dramatic change in personality and behavior following a discrete left-sided medial thalamic infarction involving the dorsomedial nucleus. Neuropsychological testing demonstrated severe impairment of complex executive behaviors that are usually associated with frontal lobe function. Electroencephalography and single-photon emission computed tomography strongly implicated dysfunction of the ipsilateral frontal lobe. This case further supports a functional and physiologic thalamofrontal linkage as part of a broader cerebral network modulating complex human behavior.

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Lesions in the cerebral cortex and thalamus can result in similar cognitive deficits. The existing evidence, obtained primarily from experimental animals, suggests that lesions in a given thalamic nucleus may produce cognitive deficits similar to those seen following injury to the primary area of cortical connectivity. Associations have been established between injury to specific thalamic nuclei and the development of amnesia, aphasia, and hemispatial neglect. Much less is known about the relationship between the thalamus and complex executive behaviors associated with frontal lobe function. Because there are extensive connections between the dorsomedial (DM) nucleus and prefrontal cortex, one would predict that injury to the DM nucleus could result in behavioral and personality changes similar to those seen with prefrontal lesions, and perhaps even electrophysiologic and metabolic changes indicative of ipsilateral frontal lobe dysfunction. Surprisingly few such cases have been described in the literature.

REPORT OF A CASE

A 62-year-old right-handed female executive administrative assistant was noted by her family to have a sudden, dramatic change in personality and behavior. She was previously described as gregarious, talkative, and extremely interested in people and current events. Initially, she became lethargic and slept excessively. The next day, she was noted to be apathetic, quiet, forgetful, and unconcerned, and showed a striking lack of spontaneity and initiative. She was described by her son as behaving "like she had a lobotomy." Neurologic consultation was obtained on the fourth day. The patient had been recently diagnosed as hypertensive and diabetic.

She was alert and oriented to person, place, and time. She lacked any insight into her condition and could not understand why she was being evaluated. She did not initiate any conversation and was very slow to respond to questions. Digit span was six digits forward and five digits backward. In a test of verbal fluency,1 she could generate only two words beginning with the letter "F" and three words beginning with the letter "A" in 1 minute. She perseverated on a written test of alternating sequences and was unable to complete the Trail Making Test, Part B.2 She made 10 errors of commission on the 50-item Stroop Color Interference Test3 and took approximately 2 minutes to finish. She had great difficulty with Luria motor sequencing.4 Interpretation of proverbs and similarities was extremely concrete. Memory for repeating a cube in three dimensions. Performed well on the three words of Three Shapes Test,5 she could not recall any of three written words after 5 minutes but was able to correctly identify all three words from a list. She was able to correctly reproduce all three drawn shapes after 5 minutes. Speech was sparse, fluent, and mildly hypophonic. She named only 18 of 30 items correctly when given every other item from the Boston Naming Test.6 Comprehension, repetition, reading, and writing were intact. Her clock drawing demonstrated poor planning, and she had difficulty constructing a cube in three dimensions. Performance of simple calculations and left-right discrimination were normal. There was no evidence of neglect on a letter cancellation test.7

An elementary neurologic examination yielded normal findings, except for a mild right-sided pronator drift and distal sensory loss. Laboratory data included a serum glucose level of 19.2 mmol/L and a fasting total cholesterol of 12.55 mmol/L.

Cranial computed tomography (CT) on the day of admission revealed a small area of slightly decreased attenuation in the left medial thalamic infarct. Repeated CT 4 days later showed the area of decreased attenuation to be better defined, consistent with a recent infarct. Magnetic resonance imaging (MRI) with 1.5-mm sections also demonstrated the left medial thalamic infarct (Fig 1) as well as several tiny foci of increased T1 involving the white matter of both hemispheres. With use of a stereotactic method described in detail by Krichevsky et al,8 the lesion was plotted on sagittal plates from the Schaltenbrand and Wahren9 atlas. The lesion primarily involved the DM nucleus, as well as adjacent internal medullary lamina, mammillothalamic tract, and ventral lateral nucleus. For illustrative purposes, a magnified view of the coronal MRI scan and corresponding coronal model drawing from the Andrew and Watkins10 atlas are shown in Fig 2. Electroencephalography (EEG) demonstrated frequent periods of focal polymorphic delta activity in the left frontal region. A single-photon emission CT (SPECT) perfusion study with technetium Te 99m-labeled hexamethylpropyleneamine oxime, with a resolution of 8.2 mm,11 showed markedly decreased uptake in the left thalamus and a 30% decrease in the left dorsolateral and anteromedial frontal lobe compared with the right (Fig 3).

During the 8 months following discharge, the patient's family noted that she showed increased initiative and independence in performing household chores but was still unable to write. When asked to write, she could not describe her family or write a letter. She had renewed interest in current events and reading. Spontaneous speech output was more productive and no longer hypophonic. She scored in the 92nd percentile on the Raven's Standard Progressive Matrices,12 a test of analogic reasoning skills. There was moderate improvement in verbal fluency, Trail Making Test, Part B, Stroop Color Interference Test, Luria motor sequencing, Boston Naming Test, and constructional tasks (Table).

By 8 months following discharge, her family felt that she was closer to her baseline personality. She had returned to work full-time, was driving without difficulty, and was performing all of her prior household and family responsibilities. However, they noted that the patient still lacked her premorbid "feisty spirit" and was slow to react to her surroundings. She had more difficulty comprehending complex ideas than before the stroke and was less interested in her grandchild. She was much more interactive and engaging during the examination. She continued to exhibit modest improvement on cognitive testing, but deficits persisted (Table).
Herein, we describe a 62-year-old woman who developed a sudden change in personality and behavior. Neuropsychologic testing demonstrated three types of deficits: (1) an impairment of complex executive behaviors that are most often associated with frontal lobe function, (2) a naming impairment, and (3) a retrieval deficit for verbal material. Imaging studies demonstrated a discrete thalamic infarction predominantly involving the DM nucleus but probably also including adjacent mamillothalamic tract, internal medullary lamina, intralaminar nuclei, and ventrolateral nucleus. The lesion appeared to be primarily in the distribution of the tuberoinfundibular artery, also called the polar artery.15,16 The language and memory disturbances in this patient are consistent with established anatomic-behavioral correlates associated with thalamic lesions. Aphasia has been associated with involvement of the ventrolateral nucleus,17,18 while verbal memory dysfunction has been described following injury to either the mamillothalamic tract or the left DM nucleus.17,18,63 Much less is known about the specific anatomic-behavioral correlates associated with a disruption in executive functions following thalamic injury. The most salient and disabling features in this case were decreased spontaneity, initiative, insight, judgment, abstraction, perseverence, and response inhibition— a constellation of behavioral disturbances that are commonly seen following prefrontal damage.21 We believe that this impairment in complex behavioral functions resulted from injury to the DM nucleus, thereby disrupting an important subcortical component of the frontal networks.21,22 This is supported by the frontal lobe dysfunction ipsilateral to the thalamic infarct documented by EEG and SPECT.

The several small foci of increased T2 signal seen on this patient's MRI scan are unlikely to have made a major contribution to her clinical presentation. Unlike other cases in which multifocal white-matter lesions were associated with frontal systems dysfunction,21 no such lesions were noted on our patient's CT scan or T1-weighted MRI image corresponding to the foci of increased T2 signal. One or more such areas of increased T2 signal are identified in 92% of patients older than 60 years,21 and no consistent correlation has been found...
between neuropsychological dysfunction and the presence of white-matter hyperintensities in the elderly. Furthermore, the patient developed a sudden change in behavior that was clinically attributable to the new thalamic lesion. While the foci of increased $T_2$ signal were small and bilateral, the EEG and SPECT studies revealed frontal lobe dysfunction ipsilateral to the new thalamic lesion.

Studies involving humans and other mammals have shown that the DM nucleus has reciprocal connections with prefrontal cortex and is part of several parallel thalamocortical circuits involving different regions of prefrontal cortex. These pathways are organized such that the parvocellularis is reciprocally interconnected with the dorsolateral prefrontal cortex, while the pars magnocellularis is reciprocally interconnected with the orbitofrontal cortex. The pars magnocellularis receives additional afferents from limbic and paralimbic structures, including the amygdala, piriform cortex, hypothalamic nuclei, and the inferotemporal cortex. Lesions of the DM nucleus in rats and monkeys may result in behavioral sequelae similar to those associated with prefrontal lesion.

The remote effect of thalamic injury on prefrontal cortex has been demonstrated by several functional neuroimaging studies. Baron et al. and Szekely et al. performed positron emission tomography on patients with thalamic strokes localized by CT. Those patients with medial thalamic lesions demonstrated hypometabolism predominantly affecting the ipsilateral frontal lobes. Posterior thalamic lesions resulted in metabolic depression affecting the temporoparietal regions most severely. Neither study employed MRI or a stereotactic method to determine which thalamic nuclei or other structures were involved. Hemmerici et al. described four patients with abulia, memory disturbances, a "premotor syndrome," and mild pure motor hemiparesis attributed to lesions in the territory of the anterior thalamoperforating arteries. The positron emission tomographic scans showed hypometabolism in the frontal and anterior regions ipsilateral to the thalamic lesions. Similarly, Bogossiansky et al. described a patient with manic delirium and frontal system dysfunction following a right thalamic infarction. The SPECT scan of this patient revealed hypoperfusion of the overlying cortex, predominantly in the frontal region.

Neuropsychological studies have also supported the predicted thalamocortical association. Stereotactically induced
thalamic lesions in cats have been shown to result in focal delta slowing overlying the cortical projection area of the destroyed thalamic nucleus. In humans, Schaul et al.14 found focal or lateralized EEG abnormalities in 59% of patients with thalamic lesions. However, the exact location and types of lesions as well as the distribution of the focal EEG abnormalities were not discussed. Most case reports of thalamic infarction have noted ipsilateral or diffuse slowing on EEG.15-16 However, there has been at least one prior report of focal delta activity corresponding to the projection area of a discrete medial thalamic lesion,17 as was seen in our case.

During the 8 months following the thalamic infarction, our patient progressively improved to a point at which her family believed she had almost returned to her baseline personality and behavioral status. Serial evaluations revealed improved spontaneity, sustained behavioral output, shifting of mental set, and inhibition of inappropriate responses. However, modest deficits persisted. This improvement is in keeping with outcomes observed following unilateral damage to prefrontal cortex. Currently, there are insufficient data to determine whether injury to the subcortical components of the frontal network results in a more favorable prognosis.

The case described herein is noteworthy because salient changes in personality and behavior consistent with a frontal lobe syndrome developed following a discrete medial thalamic infarction. As might be predicted from established anatomical components and connecting pathways of the frontal network, similar disturbances in complex executive behavior have been described not only in patients with frontal lobe injury but also in association with lesions in the caudate nucleus and with multiple subcortical lacunes.18-20 Our case adds to the literature supporting the possibility of such a behavioral outcome following injury to the medial thalamus. Moreover, it suggests that disruption of the distributed frontal network at different points can result in similar behavioral, electrophysiological, and metabolic dysfunction.

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References