Electrophysiologic recordings in a patient with a discrete unilateral thalamic infarction

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SUMMARY The electroencephalogram, and somatosensory and auditory evoked potentials were recorded from a patient, who, at necropsy, showed a restricted unilateral thalamic infarct involving predominantly the anterior and lateral thalamus. The electroencephalogram showed distinct monomorphomorphic delta activity and a suppression of the alpha rhythm over the side of the lesion. Short latency somatosensory evoked potentials were absent ipsilaterally; mid-latency somatosensory evoked potentials were absent bilaterally. Both mid-latency and long latency auditory evoked potentials were normal.

Methods

Case report The patient was a 66-year-old male with a long history of hypertension, who, while working in the sun, became weak and collapsed. At the time of hospital admission examination showed confusion, a mixed transcortical aphasia, right hemiparesis and right hemisensory deficit. Ten days after admission neurological examination revealed a mild right facial weakness, a right hemiparesis with arm weaker than leg and distal muscles weaker than proximal muscles, and a right hemisensory deficit for graphesthesia and stereognosis, vibration, pin and touch sensation. Grasp, suck, and root responses were present. Reflexes were enhanced and the Babinski sign was present bilaterally. Cerebellar examination was within normal limits. CT revealed a radiolucent area confined to the left thalamus. Three days after electrophysiologic recordings were made the patient suffered cardiorespiratory arrest from which he could not be resuscitated.

Postmortem examination of the brain revealed atherosclerosis involving the major cerebral arteries with marked involvement of the left posterior cerebral artery just distal to the junction of the left posterior communicating artery. A thrombus was found in this region which extended for approximately 1.0 cm. This thrombus apparently occluded perforating vessels which serve portions of the thalamus. Horizontal sections of the brain demonstrated an area of infarction involving the lateral portions of the left thalamus (fig 1). This infarct involved all of the ventral anterior, ventral lateral and lateral posterior nuclei. It also involved the anterior portion of the pulvinar and the lateral portion of the dorsomedial nucleus of the thalamus. The anterior nucleus of the thalamus was spared as well as the majority of the dorsomedial nucleus. The lesion extended rostrally to involve a small region of the prerubral radiations. Involvement of the internal capsule on the left was very minimal.
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Results

The electroencephalogram showed distinct monomorphic delta waves localised in the left fronto-central region. A clear alpha rhythm which was present on the right, was absent on the left (fig 2).

Recordings from the vertex disclosed three positive far-field median nerve-evoked potential components with peak latencies of 9-0, 12-5, and 15-0 ms (fig 3A). These components were recorded in response to both left and right median nerve stimulation. Left and right median nerve stimulation also evoked a negative component recorded over the neck peaking at about 14 ms (fig 3A, bottom traces). In response to left median nerve stimulation a clear N22–P28 potential was recorded over the right somatosensory cortex; a frontal N33 component was also recorded on the right (fig 3B). In contrast, stimulation of the right median nerve evoked no potentials over the left somatosensory or motor cortex, even if the stimulus raised to twice the motor threshold (fig 3B).

With auditory stimulation no hemispheric asymmetries were noted in the mid-latency ("N1–P2") or long latency ("N1–P2") components*(fig 3C).

Discussion

At the time of the recording sessions neuroradiological findings had suggested a discrete thalamic lesion and although a major object of this study was a delineation of the sources of the short and mid-latency somatosensory evoked potentials (SLSEPs and MLSEPs), some pathophysiological correlations could also be made for the EEG. The EEG of our patient was characterised by a prominent left fronto-central delta rhythm and absence of the alpha rhythm on the left. The pathophysiology of delta activity in the human EEG is poorly understood.\(^5,6,11\) Gloor and colleagues have recently showed in cats,\(^12\) that discrete thalamic lesions caused unilateral focal delta waves overlying the lesion. The frontal delta rhythm in our patient, therefore, was likely to have been caused by destruction of anterior, lateral, and dorsomedial nuclei and interruption of their projection fibres. The unilateral suppression of the alpha rhythm is similar to recordings in patients with anterior and mesial thalamic tumours presented by Jasper and Van Buren,\(^13\) who felt that the most critical projections necessary for maintenance of the alpha rhythm arose from these areas.

Although evoked potentials are widely used to study and assess the human somatosensory pathway, the actual neural sources of many components are not firmly established. With regard to the early
Fig 2  Electroencephalogram during waking state. Prominent delta waves and suppression of the alpha rhythm on the left were recorded throughout the record. Calibrations: 1 s and 10 μV.

Fig 3  (A) Short latency somatosensory evoked potentials (median n stimulation), recorded from scalp location Fz and from over the seventh cervical vertebra; the reference electrode was on the unstimulated arm. Arrows indicate far field scalp positive waves and nuchal negativity. (B) Middle latency somatosensory evoked potentials (median n stimulation) recorded from scalp locations F3, Fz, F4 (referred to Fz). Note absence of potentials over left hemisphere. (C) Middle latency and long latency auditory evoked potentials. (Binaural stimulation, 60 dBSL) recorded from C3 and C4 referred to seventh cervical vertebra.

components, Mauguier and Courjon have recently described two patients, one with a capsulo-thalamic haemorrhage and the other with a circatricial ischaemic lesion of the posterior thalamus (CT data); in both of these patients the SLSEPs could be recorded but the later MLSEPs were absent over the cortex ipsilateral to the lesion. Our patient who had an infarction involving the anterior and lateral thalamic nuclei also presented with intact SLSEPs but absent MLSEPs over the affected hemisphere indicating that the SLSEPs occurring within 15 ms post-stimulus are generated at a site caudal to the thalamus.

The exact neural source of the N20 of the MLSEPs is unknown; its generator is probably from within the primary somatosensory cortex or from
the presynaptic thalamocortical fibres. Chiappa et al., however, have presented CT data from a patient with a unilateral centrum semiovale leucoencephalopathy disconnecting the thalamus from the cortex, and have found that the N20 to be intact, suggesting that this potential is generated from within the thalamus. The lesion in our patient was restricted to the thalamus and all parietal and frontal MLSEPs were absent ipsilateral to the lesion, indicating that the thalamus is necessary for generation of N20 and succeeding components, but not required for components with a latency of less than 20 ms. The intact middle and long latency auditory potentials in the patient reflect lack of damage to the medial geniculate and medial parts of the thalamus which are thought to be necessary for their generation.

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