Dear Sir,

Astasia designates motor incoordination with inability to stand and is occasionally observed in patients with conversion hysteria [1]. Organic causes of astasia have been rarely reported. Unilateral asterixis may be caused by a focal structural brain lesion including stroke involving the thalamus, parietal lobe, frontal lobe, midbrain, basal ganglia and internal capsule, although asterixis was classically associated with metabolic derangement, particularly hepatic encephalopathy, which occurs usually bilaterally [2–5]. In addition, concurrent unilateral manifestation of astasia and asterixis due to focal cerebral lesions is extremely rare [2, 4]. To our knowledge, sudden concurrent development of asterixis and astasia as a consequence of a rostral midbrain infarction has not yet been reported. We report a patient with concurrent unilateral astasia and asterixis developed due to acute cerebral infarctions involving the rostral level of the midbrain without significant metabolic disorder.

Case Report

A 70-year-old man was admitted due to sudden onset of postural instability and dysarthria. His medical record revealed a cerebral infarction in the right centrum semiovale 8 years previously but showed no neurological condition such as essential tremor or extrapyramidal disease before the present event. He also had an 8-year history of hypertension and a 5-year history of diabetes mellitus, which were well controlled. However, there were no histories of hepatitis, smoking and heavy alcohol consumption. He has taken aspirin 100 mg, clindipine 10 mg, imidapril 10 mg and acarbose 50 mg daily. At the emergency room, the blood pressure was 160/90 mm Hg, the pulse rate 78/min, the temperature 36.8 °C. On neurologic examination, he was alert and oriented with fluent responses were flexor. When both his hands were outstretched, postural lapses with arrhythmic losses of extensor muscle tone that developed upon instruction to maintain the wrist in an extended position, so-called asterixis, or flapping tremor, was observed in the right hand. Finger-to-nose test, rapid alternation movement and heel-to-shin test were all normal. On attempt to stand, however, he showed postural instability and gait disturbance unassisted due to a tendency to tilt backward or toward right, while his ability to sit was relatively preserved. Laboratory studies, including a complete blood cell and platelet count, the erythrocyte sedimentation rate, blood electrolytes, creatinine, liver enzymes, cholesterol, triglycerides and the prothrombin and partial thromboplastin time were all normal. Magnetic resonance images of the brain on the admission day revealed high signal intensities on an apparent diffusion coefficient map in the left rostral paramedian mesencephalon region (fig. 1). One day later, we performed surface electromyographic (EMG) recording to verify unilateral postural lapses of the right upper extremity. Surface EMG recording from right wrist extensor muscle showed an irregular brief pause of muscle activities with a duration of approximately 30–50 msec, which indicated asterixis (fig. 2). Waking and sleep electroencephalography performed to exclude partial seizure showed neither epileptic discharges nor other abnormal slow waves. On the 4th hospital day, asterixis disappeared, while the tendency to fall to the right posterolateral side was resolved on the 5th hospital day.

Discussion

The term astasia is defined as motor incoordination with inability to stand despite good motor strength. The disturbance differs from cerebellar ataxia in that gait is not broad-based or lurching. It resembles the marked balance impairment of patients with vestibulocerebellar disease [4]. The responsible anatomical region for astasia or astasia-abasia has been known to be related to the thalamic lesion affecting mainly the medial ventrolateral nucleus of the thalamus because the fastigial fibers of...
Concurrent Astasia and Asterixis due to Mesencephalic Infarction

**Fig. 1.** Magnetic resonance images of the brain on admission day. **A** Diffusion weighted imaging of brain shows small and discrete high signal intensities only in the left rostral paramedian mesencephalon without lesion of other areas including thalamus. **B** Apparent diffuse coefficient map shows low signal intensities at comparable site.

**Fig. 2.** Surface electromyographic recording from right wrist extensor shows irregular brief pause of muscle activities.
the vestibulocerebellar pathway project to the medial ventrolateral nucleus, and disruption of this pathway may cause thalamic astasia [6]. Inability to walk despite good motor strength was also reported in a patient with a red nucleus infarction involving the ascending fibers of the crossed dentatorubrothalamic tract, in a patient with a small medial capsular hemorrhage extending to the lateral portion of the ventrolateral nucleus of the thalamus and in a patient with a hemorrhage at the pontomesencephalic junction (midbrain tegmental lesion) involving the pedunculopontine area [7–9]. According to their previous report, Masdeu and Gorelick [1] also asserted that the propensity of a thalamic lesion to cause impairment of balance and gait might be because of a disconnection from vestibulocerebellar information to the vestibular cortex. However, the exact anatomical origin of astasia is not yet known.

Asterixis, one type of negative myoclonus, results from a sudden cessation of electrical activity in the extension of the limb [5]. Asterixis may occur after focal cerebral lesions involving the midbrain, thalamus, parietal lobe or the frontal cortex [2, 3, 10, 11]. The pathogenic mechanism for asterixis remains elusive. Previous authors suggested that asterixis is a negative myoclonus that arose as a result of damage to a neuronal circuit sustaining postural muscle contraction [10]. This postural stability or tonic control of the extremities is related to multiple brainstem-spinal pathways such as the vestibulospinal, reticulospinal or rubrospinal tracts.

These systems are in turn regulated by supratentorial structures; the ventrolateral nucleus of the thalamus is the area in which cerebellar-rubral or vestibulocerebellar fibers converge and is also heavily connected with the prefrontal area [3]. Since astasia and asterixis are considered to be manifestations of disordered postural or tonic balance regulated by the fastigial-thalamocortical pathway, we could explain that the lesions involving this circuit may produce these symptoms at the same time.

Our patient showed transient co-occurrence of astasia and contralateral asterixis due to acute left rostral paramedian mesencephalic infarction. The paucity or absence of motor weakness and sensory dysfunction illustrated that the corticospinal and spinothalamic tracts were generally spared. Therefore, concurrent manifestation of astasia and unilateral asterixis in our patient could be explained by disruption of cerebellar-rubral or vestibulocerebellar fiber, which projected from the fastigial fiber to the ventrolateral nucleus of the thalamus, at the rostral midbrain or midbrain-diencephalon junction level. In conclusion, we suggest that clinicians must remember to develop focal asterixis and astasia by disconnection of projection from the fastigial fiber to the ventrolateral nucleus of thalamus.

References
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