Contralateral Body Half Hypalgesia in a Patient with Lateral Medullary Infarction: Atypical Wallenberg Syndrome

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On admission she was normotensive, and her general examination was normal. Neurological examination showed left Horner’s syndrome, with the left eye mildly ptotic (fig. 1a). There was no nystagmus in both eyes. The left pupil was about 3 mm and the right one 4 mm in diameter; both pupils were alert to light. The left palate was paraparetic, and the left throat reflex was decreased. The tongue moved well and there was no facial weakness. Sensation to pain and heat was decreased substantially in the right trunk and limbs, but decreased mildly in the right face, and normal in the left face and the left body. Bilateral vibratory and postural sensation was preserved. The left finger-nose test and the heel-knee-shin test were minimally incoordinated, compared with those of the right. When standing upright and closing the eyes, she felt mild lateropulsion to the left. There was no weakness of the limbs.

Cerebral MRI detected a small band-like ischemic lesion in the left retro-olivary midlateral tegmentum, with preservation of the far dorsal vestibular nucleus and inferior cerebellar peduncle (fig. 1c). Angiographic sequences disclosed congenital dysplasia of the left vertebral artery and segmental stenosis at the origin of the posterior inferior cerebellar artery (fig. 1b).

Discussion

Symptoms and signs found in patients with lateral medullary infarction depend on the dorsal-ventral, medial-lateral and retrocaudal locations of the lesion. Vestibulocerebellar symptoms and signs are nearly always present in these patients [3]. The present patient presented with left Horner’s syndrome, dysphagia, mild hoarseness and mild lateropulsion to the left. These clinical findings, together with the radiological findings of the left medulla, confirmed the diagnosis of lateral medullary infarction without any doubt. What was extraordinary in this patient was that she merely felt dizziness, which gradually disappeared in half an hour after onset. In addition, she did not feel obvious vertigo, and the left finger-nose test and the heel-knee-shin test were only minimally incoordinated, indicating that the infarction did not involve the vestibular nucleus and the left inferior cerebellar peduncle. This condition was later explained by the MRI findings, which disclosed normal signals at the dorsal edge of the medulla. It is common knowledge that the vestibular nucleus is located far dorsally to the bottom of the fourth ventricle. This type of infarction without involvement of the vestibular nucleus and inferior cerebellar peduncle is a rare condition.
Moreover, the characteristic crossed sensory presentations including hypalgesia in the ipsilateral face and the contralateral trunk and limbs were not seen. This classic pattern of sensory symptoms (type I) is due to the involvement of the crossed lateral spinothalamic tract and the ipsilateral descending spinal trigeminal tract and its nucleus [4, 7]. This crossed type corresponds to far dorsal lateral medullary lesions (fig. 1e). If the lesion extends more ventrally and medially, the crossed trigeminotectal tract (which carries pain and heat sensation from the contralateral side of the face) may be involved, and the patient may...
present with hypalgesia in the whole contralateral body, including the face, trunk and limbs, in addition to the ipsilateral face (fig. 1f); this type of sensory defect is type II. In our patient, however, the ipsilateral descending spinal trigeminal tract and its nucleus (which lies more dorsally to the spinothalamic tract) had obviously been spared, and the ventral crossed trigeminothalamic tract had been involved (fig. 1g). This is why our patient did not feel numbness or hypesthesia in the left face and why the left corneal reflex was sensitive. This type of sensory defect that only involves the contralateral body is type III.

Still, there are other less common sensory defect patterns in lateral medullary infarction, which have been clearly described by Matsumoto et al. [6]. One pattern presents with hypalgesia in the ipsilateral face and contralateral lower trunk and leg (type IV). This type of sensory defect is caused by the infarction of far lateral location, which only involves the lateral descending spinal trigeminal tract and its nucleus, and the ventral lateral part of the spinal thalamic tract (fig. 1h). An-
other pattern of sensory defect manifests hypalgesia only in the contralateral face, arm and upper trunk, without involvement of the ipsilateral face. This rare type of sensory defect (type V) is caused by restricted infarction of the mediolateral medulla, which only involves the crossed trigeminal thalamic tract and the medial part of the spinal thalamic tract (fig. 1i) [5, 6]. These partial sensory defects on the contralateral side of the body are due to somatotopical organization of the spinothalamic tract (fig. 1d). The sacral afferent fibers are located in the lateral medullary part, and the cervical afferent fibers ascend more medially. The crossed ventral trigeminothalamic tract appends on the medial part of the spinothalamic tract and carries pain and heat sensation from the contralateral side of the face [8].

In conclusion, we report a rare type of lateral medullary infarction, which presents without obvious vertigo, and the sensory defect is entirely in the contralateral body half.

References