Unusual neurological cause of intractable hiccups: A case report

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Abstract
Intractable hiccups is a poorly understood medical problem and the pathophysiology is highly debatable. Here we report an interesting case of a 40 year male patient who presented to us with complaints of difficulty in swallowing and hiccups for 3 weeks which was resistant to medication. MRI scans revealed Chiari I malformation and cervical syrinx. He underwent suboccipital craniectomy including a C1 laminectomy following which he was relieved of intractable hiccups.

Introduction
Intractable hiccups is a poorly understood medical problem and the pathophysiology is highly debatable. Hiccups consist of brief bursts of intense inspiratory activity involving the diaphragm and the inspiratory intercostal muscles, with reciprocal inhibition of expiratory intercostals muscles, the sound and discomfort resulting from glottis closure immediately after the onset of diaphragmatic contraction.1 Most episodes of hiccups are self-limited, but prolonged or intractable hiccups should prompt a search for a cause. Hiccups are defined as intractable when they persist for 24 hours. The afferent pathway of the hiccup reflex arc includes the sensory branches of the phrenic and vagus nerves and dorsal sympathetic fibers; the main efferent limb, causing spasm of the diaphragm, is composed of motor fibers of the phrenic nerve.2

Case report
A 40 year male patient presented to us with complaints of difficulty in swallowing and hiccups for 3 weeks. His hiccups were refractory to baclofen, metoclopramide and chlorpromazine. Blood urea was normal and evaluation by other departments could not reveal any cause. Neurological examination revealed hyperreflexia of all 4 limbs and suspended sensory loss in the upper extremities. Cranial nerve examination demonstrated no positive neurological findings, including bilateral gag reflexes, except for the persisting hiccups. Preoperative magnetic resonance imaging (MRI) scans revealed Chiari I malformation and cervical syrinx.

Figure 1. MRI showing Chiari I with syrinx
Suboccipital craniectomy including a C1 laminectomy was performed. Dura was opened in Y shaped fashion, subpial tonsillar resection and duraplasty was done. Post operative course was uneventful. Hiccups disappeared for 2 days post op but reappeared again with decreased frequency and finally after 2 weeks patient was relieved of the vexing problem.
Discussion

Although their pathophysiological cause remains unclear, hiccups or singultus are thought to be related to stimulation, probably by damage or irritation, of one or more points of the hiccup reflex arc. The hiccup reflex arc includes an afferent portion that involves the vagus nerve, the phrenic nerve, and the sympathetic chain arising from T6–T12. The hiccup centers are thought to be located either in the brainstem close to the inspiratory centers or in the cervical cord between C3 and C5. The efferent limb consists primarily of the phrenic nerve and its interactions with the glottis and accessory respiratory muscles, and the complex interactions between the brainstem and midbrain areas, including the dorsal portion of the nucleus tractus solitarius (inspiratory center), the ventral group of the nucleus ambiguus (expiratory center), phrenic nerve nuclei, medullary reticular formation, and hypothalamus.4

Intractable hiccups as a presenting symptom of cerebellar hemangioblastoma5 medulla oblongata cavernoma6, and rare cases of medullary compressive lesions presenting with intractable hiccups7 have been reported. Fisher et al.8 reported dorsal and lateral medullary infarctions presenting with persistent hiccup. Kumral and Acarer9 described an unusual case of primary medullary hemorrhage with intractable hiccup. There have been some reports describing cases of intractable hiccups associated with extensive syringomyelia and Chiari malformation.10,11 Because compression of the medulla oblongata by the herniated tonsil is a common finding in syringomyelia associated with Chiari malformations. It is likely that medullary compression played a role in pathogenesis in our patient. It is often difficult to treat intractable hiccups. Baclofen, haloperidol, carbamazepine, and chlorpromazine have been recommended for the treatment of intractable hiccups. Electrical stimulation of the phrenic or vagus nerves and microvascular decompression of the vagus nerve have been reported for the treatment of hiccups. In our case, intractable hiccups associated with syringomyelia was successfully treated with foramen magnum decompression. When persistent intractable hiccups occur, with or without slow neurological deterioration, intracranial lesion should be suspected. MRI scans obtained at the time of the initial diagnosis and at follow-up assessments are valuable. Hiccups caused by syringomyelia and Chiari malformations represent a surgically treatable disorder, although the incidence is low.

Conflict of Interest: None declared

References