Locked-in Syndrome due to Central Pontine Myelinolysis: Case Report

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Central pontine myelinolysis (CPM) classically occurs in alcoholics, malnourished individuals, chronic liver diseases, and rapid correction of hyponatremia. This report presents locked-in syndrome due to CPM following rapid correction of hyponatremia. A 44-year-old male came to the hospital due to a short period of loss of consciousness. He was alert and had no focal neurological abnormalities at admission. The serum sodium concentration was 118 mEq/L and was corrected to 134 mEq/L in the first 18 hours. One week later, progressive weakness in limbs developed and he progressed to a complete quadriplegic state and bulbar palsy, with only eye blinking preserved. Brain magnetic resonance imaging revealed a characteristic hyperintense signal abnormality in both pons, so he was diagnosed to locked-in syndrome caused by CPM. The patient gradually improved following continuous intensive rehabilitation for more than 2 years. He was able to move all joint muscles against gravity in generally and he could gait under supervision.

Keywords Central pontine myelinolysis, Quadriplegia, Recovery of function

INTRODUCTION

Central pontine myelinolysis (CPM), first described by Adams et al. [1], is caused by disappearance of neurons and myelin comprising the central pontine. The exact incidence of CPM is unknown. In a study of 3,000 brains examined postmortem 15 cases of asymptomatic CPM were evident [2]. Lampl and Yazdi [3] reported 39.4% of patients of CPM were diagnosed with alcohol poisoning, 21.5% had a history of correcting hyponatremia, and 17.4% received a liver transplantation.

Clinical presentation of CPM is highly variable from asymptomatic to coma depending on the location of the lesion. The severity of the clinical presentation does not correlate with the size of the lesion [4]. Patients typically experience a rapidly evolving paraparesis or quadriparesis, and pseudobulbar symptoms like dysarthria and dysphagia. They may present with ‘locked-in syndrome’ in which intellectual activity is preserved but cannot be expressed because of a total incapacity to produce voluntary responses.

Although earlier reports on CPM described very poor outcome with mortality rates as high as 50% in the first 2 weeks after presentation and 90% at 6 months, recent reports suggest the decreased mortality mainly due to early diagnosis and improved intensive care treatment [5]. When patients survive their illness, most have significant
neurological deficit. There were no reports of neurologic recovery after diagnosis of locked-in syndrome caused by CPM in Korea.

We report a patient diagnosed with locked-in syndrome due to CPM, whose ability to perform daily living and muscle power were continuously improved after comprehensive rehabilitation treatment for more than 2 years.

CASE REPORT

A 44-year-old Korean male was admitted to the hospital because of a dizziness-induced fall and 1-minute loss of consciousness. The patient had a history of drinking two or more bottles of alcohol every day and had not eaten for the previous month.

On initial physical examination, he was alert and no focal neurological abnormalities were noted. Laboratory data revealed hyponatremia (serum sodium 118 mEq/L), hypokalemia (potassium 2.5 mEq/L), hypoalbuminemia (albumin 3.2 g/dL), abnormal liver function (aspartate transaminase 175 IU/L and alanine transaminase 68 IU/
L). Serum sodium concentration was corrected to 134 mEq/L during the first 14 hours and 137 mEq/L within 18 hours following intravenous injection of 3% saline (Fig. 1).

One week after admission, progressive dysarthria, dysphagia, and weakness in limbs developed, so he was transferred to the intensive care unit. Feeding nasogastric and tracheostomy tubes were inserted and ventilator care was started. Brain magnetic resonance imaging (MRI) showed symmetrical areas of signal hyperintensity in the bilateral pons on T2-weighted images and diffusion-weighted imaging, consistent with a diagnosis of CPM (Fig. 2A, 2B). Although he remained conscious, complete facial, bulbar, and limb muscle palsy developed with only eye blinking and eyeball movement preserved. He was moved to the general ward 3 weeks later and transferred to the Department of Rehabilitation Medicine 2 months later. His transfer was delayed due to fever and anal abscess.

At transfer to rehabilitation the patient presented with severe flaccid bulbar and limb paralysis, and the Korean version of Modified Barthel Index (K-MBI) score was 0. Median and tibial somatosensory evoked potentials showed no response bilaterally. Comprehensive rehabilitation including general reconditioning and functional training to prevent deconditioning and contracture, and improve muscle power and cardiopulmonary function, electrical stimulation, and oromotor facilitation training were performed. Muscle power and physical function gradually improved. After 2 years, he could walk with supervision (Motricity index: upper extremity 76/76, lower extremity 75/75, total 76/76, Berg Balance Score 42; 10-m walking test 17 seconds; timed to up and go 22 seconds; 6-minute walking distance 90 m). He could use his hands in activity (Purdue pegboard score 6/6, grip power 51 lb/48 lb) (Table 1). He could write a letter and communicate by language. The tracheostomy tube was removed. Although dysphagia gradually improved, he still maintained percutaneous endoscopic gastrostomy.

Follow-up brain MRI repeated one year later showed encephalomalacia at the site of T2-high signal intensity lesion in pons (Fig. 2C, 2D). And follow-up median and tibial somatosensory evoked potentials showed prolonged latency of N19, P23, P37, N45 potentials bilaterally (Fig. 3).

**DISCUSSION**

CPM has traditionally been associated with rapid correction of hyponatremia, but the etiology has not been clearly established. Alcoholism, chronic malnutrition, and sodium imbalance are the primary conditions reported with CPM [1]. Diagnosis of CPM is based on clinical neurologic examination and confirmed by imaging studies. MRI is the primary method for diagnosis and is superior to computed tomography [6].

Rapid correction of hyponatremia (i.e., >12 mEq/day) poses a higher risk of developing CPM [7]. In this case, sodium level rapidly corrected to 137 mEq/L from 118 mEq/L within 18 hours. Neurological symptoms that included quadriplegia, dysphagia, and dysarthria developed after 1 week. Brain MRI showed symmetrical areas of signal hyperintensity in the pons on T2-weighted images, consistent with a diagnosis of CPM. Diffusion-weighted imaging confirmed the presence of hyperintensity at the same areas. The patient developed severe

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**Table 1. Clinical data after 2 years of comprehensive rehabilitation**

| Variable                        | Value          |
|---------------------------------|----------------|
| Medical Research Council        |                |
| Shoulder abductor               | 4 (5) / 4 (5)  |
| Elbow flexor                    | 4 (5) / 4 (5)  |
| Finger flexor                   | 4 (5) / 4 (5)  |
| Finger extensor                 | 4 (5) / 4 (5)  |
| Hip flexor                      | 4 (5) / 4 (5)  |
| Knee extensor                   | 4 (5) / 4 (5)  |
| Ankle dorsiflexor               | 4 (5) / 4 (5)  |
| Motricity index                 |                |
| Upper extremity                 | 76 (100) / 76 (100) |
| Lower extremity                 | 75 (100) / 75 (100) |
| Total                           | 76 (100) / 76 (100) |
| Berg Balance Score              | 42 (56)        |
| Purdue pegboard test            | 6 (14±2.3) / 6 (14±2.1) |
| Grip power (lb)                 | 51 (109.9) / 48 (100.8) |
| Box-block test                  | 31.25 (76.9) / 31.84 (75.8) |
| Nine-hole pegboard test (second)| 46 (18.35) / 48 (19.57) |

Values are presented as number (full score of scale or age-and sex-matched normal control data for the Purdue pegboard test, grip power, box-block test, and nine-hole pegboard).
Recovery of Fx in Locked-In Syndrome due to CPM

Flaccid quadriplegia associated with dysarthria and dysphasia. Eventually, the patient was diagnosed as locked-in syndrome caused by CPM.

Previous studies have reported on the heterogeneous prognosis of quadriplegia in patients with locked-in syndrome [7,8]. These studies suggested the possibility of several types of motor recovery mechanisms for locked-in syndrome. Kwon and Jang [6] reported on a quadriplegic patient with bilateral pontine infarcts, whose motor function appeared to be reorganized into the lateral areas of the pontine infarcts. Less severe hyponatremia (>115 mEq/L), absence of superadded hypokalemia, and higher Glasgow Coma Scale (≥11) at the time of admission and at discharge are good prognostic factors in osmotic demyelination syndrome [8]. Also, the neurologic recovery rate of locked-in syndrome caused by non-vascular lesion is faster than that of avascular causes and more complete recovery is evident in locked-in syndrome due to non-vascular cause [9]. In this case, the patient was in a complete quadriplegic state but gradually recovered to antigravity power and could walk unaided after 2 years of comprehensive rehabilitation. Although he had very severe locked-in syndrome, some favorable prognostic factors were present. These included non-vascular cause locked-in syndrome, initial sodium level of 118 mEq/L (≥115 mEq/L), good Glasgow Coma Scale score, and preserved lateral area of the pons on brain MRI. In addition, comprehensive rehabilitation worked as a good prognostic factor.

In conclusion, if blood test showed hyponatremia, care must be taken to avoid develop CPM following rapid correction of hyponatremia. Diagnosis as CPM with clinically confirmed locked-in syndrome necessitates further studies to evaluate prognostic factors, and early comprehensive rehabilitation are recommended for recovery from neurological deficits as soon as possible.

CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was reported.
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