Central pontine myelinolysis associated with hypokalaemia in anorexia nervosa

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A 31 year old man was admitted to hospital with anorexia, binge eating, and self induced vomiting. On admission, he showed a pronounced low weight and disturbance of the body image and was diagnosed as having anorexia nervosa. In addition, electrolyte abnormalities, mainly hypokalaemia, and increased serum renin and aldosterone concentrations were recorded, suggesting pseudo-Bartter syndrome as a complication. Under frequent monitoring of the serum potassium and sodium concentrations, serum electrolytes were gradually corrected, but brain magnetic resonance imaging revealed reversible central pontine myelinolysis (CPM). Although attention has been mainly paid to the association of CPM with rapid correction of hyponatraemia and abnormal osmolality, this case report strongly suggested the involvement of hypokalaemia in the pathogenesis of CPM.

Central pontine myelinolysis (CPM) is a demyelinating lesion in the central pons that was first reported in necropsy cases of alcoholism and malnutrition.1 CPM is characterised by disturbance of consciousness, quadriplegia, and mutism and has been considered to have a poor prognosis.2,3 However, with advances in diagnostic imaging techniques such as magnetic resonance imaging (MRI), some patients with asymptomatic CPM, which had been overlooked in the past, have been reported.4 Since the pathogenesis of CPM, abnormalities in serum electrolytes and serum osmolality have mainly attracted attention, and many patients with CPM caused by rapid correction of hyponatraemia have been reported.4 However, normonatraemic and hypernatraemic patients have also been described.5–9 Recently, hypokalaemia related CPM irrespective of the serum sodium concentration has been also reported.10–11 Thus, there is no consensus about what abnormalities in electrolytes or osmolality are important pathogenic factors of CPM.

Several patients with CPM as a complication of anorexia nervosa have been reported. In each patient, the mechanism of the development of CPM was discussed in association with serum electrolyte abnormalities attributable to malnutrition, excessive water intake, or laxative and/or diuretic misuse and their rapid correction.12–14 We report a patient with anorexia nervosa complicated by pseudo-Bartter syndrome who repeated self induced vomiting and showed a reversible CPM lesion on MRI images despite gradual correction of electrolyte abnormalities mainly hypokalaemia.

CASE REPORT

The patient was a 31 year old man without remarkable family history or past history. Though his school record was good, he developed pilonidal abscess in the gluteal region at the age of 14 years and could not concentrate on studying because of pain induced stress and repeated operations. He could not enter the high school he hoped and began to repeatedly exhibit anorexia, binge eating, and self induced vomiting. The body weight was 84 kg at the age of 14 years but decreased to 50 kg at the age of 25 years when he got a job. After he began to work on a day/night shift in April 2001, the frequency of binge eating and self induced vomiting increased. On 23 July, he visited a local hospital because of general malaise. At this time, the body weight was 34 kg (body mass index =12.7), and abnormalities of electrolytes mainly hypokalaemia were observed (serum sodium concentration, 140 mEq/l; serum potassium concentration, 2.8 mEq/l). He was referred to the medical department of our hospital and admitted on the same day. Immediately, intravenous fluid therapy via a peripheral vein was initiated for the normalisation of the hypokalaemia and nutritional supplementation. Based on increased serum renin and aldosterone values and hypokalaemic alkalosis (renin, 24.6 ng/ml/h; aldosterone, 572 ng/dl; pH, 7.51), a diagnosis of pseudo-Bartter syndrome caused by self induced vomiting was made. After admission, self induced vomiting was observed after almost each meal, and the body weight did not increase. On 20 August, intravenous hyperalimentation (IVH) was started, but behavioural problems such as going out without permission and binge eating appeared. On 6 September, he was transferred to the psychiatric ward. During this period, the serum potassium concentration markedly changed, often resulting in severe hypokalaemia (fig 1). However, the serum sodium concentration and osmolality changed almost in the normal range (fig 1).

At the time of the transfer to the psychiatric ward, marked low body weight (<60% of standard weight) was observed. He had a strong fear of obesity and misperception of the body image and was diagnosed as having anorexia nervosa according to the DSM-IV diagnostic criteria.14 On 10 September, brain MRI revealed hyperintensity in the central pons on T2 weighted images (fig 2A), indicating asymptomatic CPM (absence of neurological or psychiatric symptoms in the foreground). IVH was continued and changed to oral ingestion when the body weight reached 40 kg. To inhibit self induced vomiting, behavioural restrictions were imposed (stepwise according to body weight) by making him to stay at rest in his room for a certain time after each meal. However, he could not heed these behavioural restrictions and repeatedly vomited in secret, and the body weight decreased to 32 kg. Therefore, from 15 November, IVH was performed again, and behavioural restrictions were tightened. The frequency of vomiting decreased, and abnormalities of electrolytes, mainly hypokalaemia, were normalised. Subsequently, CPM findings on MRI images disappeared (fig 2B).

Abbreviations: CPM, central pontine myelinolysis; MRI, magnetic resonance imaging

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DISCUSSION

CPM produces neurological symptoms such as quadriplegia, pseudobulbar palsy, oculomotor paresis, and locked in syndrome and psychiatric symptoms such as changes in the level of consciousness and behaviour or cognitive dysfunction. In this patient, none of these clinical symptoms was observed in the foreground, and asymptomatic CPM was considered. With recent advances in diagnostic imaging techniques such as MRI, cases of asymptomatic CPM, which may have been overlooked previously, have been reported. There may be many patients with latent reversible CPM as in this patient.

The pathogenic factors of CPM first reported were alcoholism, malnutrition, and chronic liver disease. Subsequently, rapid correction of hyponatraemia has attracted attention as an important factor. In addition, abnormal osmotic pressure and electrolyte abnormalities such as hypokalaemia and acute changes in the blood glucose concentration have also been suggested. In this patient with anorexia nervosa, there was no history of alcohol misuse or liver dysfunction, and the serum sodium concentration and osmole changed almost in the normal range. The fasting blood glucose concentration was 70–100 mg/dl without marked changes. On the other hand, the serum potassium concentration considerably changed, often showing severe hypokalaemia. Patients who vomit lose hydrogen, chloride, and water and develop hypokalaemia not only because of shift of potassium ions into cells as a result of a decreased concentration of Na,K-ATPase but also because of increased potassium ion excretion in the urine as a result of reabsorption of hydrogen ions. Secondary aldosteronism attributable to the decrease of total body water also causes an increase of potassium ion excretion in the urine. As this patient also frequently repeated vomiting, pseudo-Bartter syndrome may have developed by such a mechanism, resulting in marked hypokalaemia. Our review of the literature showed many patients with CPM complicated by hypokalaemia but only three patients with CPM with hypokalaemia but normonatraemia as was observed in our patient. The underlying disease was chronic alcohol misuse in two of the three patients and Sjögren’s syndrome in the other; anorexia nervosa was not observed.

The developmental mechanism of CPM is unknown. Proposed hypotheses include osmotic injury to the endothelium resulting in release of myelinotoxic factors or vasogenic oedema and brain dehydration resulting in separation of the axon from its myelin sheath with resultant injury of oligodendrocytes, particularly at interfaces of grey and white matter. However, a recent study suggested that CPM tends to occur in hyponatraemia complicated by hypokalaemia because of a decreased concentration of Na,K-ATPase in endothelial cell membrane during hypokalaemia may predispose the cell susceptible to injury by osmotic stress associated with the rapid rise in the serum sodium concentration. This patient also developed hypokalaemia caused by frequent self induced vomiting. Therefore, there is a possibility that CPM was induced by a slight increase in osmotic pressure attributable to fluid infusion such as of electrolytes and glucose in the presence of severe hypokalaemia (even if the increase rate was so slow as not to induce injury in the normal state).

To our knowledge, there have been reports of only two patients with anorexia nervosa in whom CPM developed after rapid correction of marked serum electrolyte abnormalities attributable to water intoxication (excessive water intake) or...
laxative misuse. In our patient, however, neither excessive water intake nor laxative abuse was observed, and electrolyte abnormalities, mainly hypokalaemia, caused by frequent self-induced vomiting developed (the serum sodium concentration was nearly normal), and CPM occurred despite gradual electrolyte correction. In patients with anorexia nervosa, the complication rate of vomiting is 28% and that of hypokalaemia is 20%. This case report suggests that further caution is necessary when serum electrolytes are corrected in such patients.

REFERENCES