Wernicke’s Encephalopathy, A Curable Cause of Acute Bilateral Vestibulopathy

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Introduction

Carl Wernicke a German physician was the first to describe Wernicke’s encephalopathy (WE) in 1881 [1]. WE are defined as an acute neurologic disorder resulting from thiamine (vitamin B1) deficiency [2]. WE is known to be common in alcoholics, but it was reported to be associated with conditions which cause nutritional deficiencies such as gastrointestinal malignancies, Hyperemesis Gravidarum, chemotherapy, HIV, prolonged therapeutic fasting, prolonged parenteral nutrition and bariatric surgery, anorexia nervosa and secondary to malnutrition due to poverty [3-6]. The classic symptom triad of WE consists of mental status changes, ophthalmoplegia, and gait ataxia, but research showed that this triad only presents in 16-19% of cases [2,7,8]. This has been taken into account by published diagnostic guidelines such as the EFNS guidelines, requiring only two out of four signs (dietary deficiencies, eye signs, cerebellar dysfunction, and either altered mental state or mild memory impairment) [9]. Variety of neuro-ophtalmological signs such as horizontal, vertical, and gaze-evoked nystagmus, abducens palsy, ophthalmoplegia and internuclear-opthalmoplegia could be found [10]. Nystagmus is the most commonly described sign, recent reports showed that bilateral severe impairment of lateral canal vestibulo-ocular reflex is commonly associated with Wernicke’s encephalopathy documented by bedside head impulse test, caloric testing and rotary chair testing [11-13]. The availability of video Head Impulse Test systems in the last few years made it possible to test the functions of the six semicircular canals [14-16]. The fact that afferents from horizontal canals relay mainly in medial vestibular nuclei and afferents from vertical canals relay mainly in lateral vestibular nuclei, and the fact that the medial vestibular nucleus is vulnerable to thiamine deficiency explains the selective impairment of vestibulo-ocular reflex of horizontal canals in cases of Wernicke’s Encephalopathy [17,18]. Being a treatable cause of acute vestibulopathy, WE should be detected early using the vHIT and thiamine replacement should be initiated in a proper and rapid manner [19]. MRI brain can help in diagnosis of WE by showing the characteristic findings such as symmetrical alterations in the thalami, mammillary bodies, tectal plate, and periaqueductal area, but atypical alterations may also be seen [18,20]. Direct measurement direct of thiamine level in human blood is possible by a technique called HPLC. Adult normal range (60-220nM). The blood sample should be taken before administration of thiamine and should be protected from light. However, normal thiamine levels can be found in cases of WE due to rare condition of thiamine transporter gene mutation [9,21-23]. Regarding the treatment of WE, there is no consensus on the optimal dose, form and duration of of thiamine treatment. Treatment with 200 mg thiamine given intravenously three times daily until improvement of the symptoms and signs or until no further improvement could be obtained [9,24,25].

Case Report

A 77-year-old man, with recent history of post-traumatic stress disorder after he had witnessed the sudden death of his son, the patient suffered loss of appetite, anorexia and vomiting over the last two months. He had no history of drinking alcoholics and he was in a good health until that incident. He presented to the specialized Hearing & Balance clinic with unsteadiness, vertigo, blurring of vision during head movements, tingling in his four extremities, there was no concerns regards memory. On examination ; the patient was ataxic and unable to stand unsupported, there was horizontal gaze-evoked nystagmus, vHIT showed bilateral selective severe lateral semicircular canal hypo functions (gain below 0.2) with compensatory saccades with normal VOR gain of vertical canals on RALP and LARP planes.MRI brain done and revealed unremarkable. Neurological examination showed no motor deficits. Serum vitamin B1 was...
assessed and the result showed decreased serum level. Vitamin B12 and B6 levels were of normal levels. Vitamin B1 (Thiamine) was given to the patient in a dose of 200mg intramuscular injection twice daily for two weeks. Gaze-evoked nystagmus disappeared at the 10th day of therapy. Ataxia improved after two weeks of treatment. vHIT was repeated after two weeks of treatment and showed marked recovery of the lateral canal VOR gain in both sides with disappearance of compensatory saccades Figure 1.

Discussion

The patient was presented with acute vestibular syndrome in the form of rapid onset but persistent unsteadiness and vertigo. The findings on clinical examination was suggestive of cerebellar involvement in the form of ataxia and gaze evoked nystagmus, testing the patient with vHIT revealed the characteristic finding of severe selective lateral canal hypofunction, which directed our attention to WE as the most probable diagnosis. The clear history of nutritional deficiency for two months in this case explains the thiamin deficiency. The low serum level of Thiamin supported the diagnosis of WE. The absent memory and mental changes as well as the absence of classical findings in MRI brain could be explained by the early stage of WE. The dramatic response to timely parental thiamin treatment confirmed the diagnosis. The findings in this case are in agreement with what has been recently reported in the literature. The use of vHIT for assessment of the six SCCs can enable the clinicians to easily detect the characteristic pattern of bilateral severe selective lateral canal hypofunction in cases of WE, and hence the timely and prompt initiation of thiamin treatment. WE is a curable cause of bilateral vestibulopathy which should get more attention by clinicians Figure 2.

References


