Midbrain stroke syndrome: Weber’s syndrome in two male HIV-positive patients with cerebral toxoplasmosis

Jorge-Luis Wong-Armas* 1,2,a; César-Justo Obregón-Manzur 2,b; Bruce-Jorge Wong-Ramírez 2,b; Tian-Hanayka-Del Carmen Wong-Ramírez 2,b; Carlos-Alberto Ortiz-Castillo 1,2,c

ABSTRACT
This report describes the case of two male HIV-positive patients who were not receiving treatment and whose diagnosis was confirmed by serology testing. Both patients developed midbrain stroke syndrome and the structural injury, which was caused by a central nervous system infection due to cerebral toxoplasmosis, was verified by brain magnetic resonance imaging. This condition was confirmed by cerebrospinal fluid serology testing.

Weber’s syndrome is a very rare type of midbrain stroke syndrome with little information available on its symptoms and evolution. It was first described by Sir Herman David Weber in 1863 and is characterized by ipsilateral injury of the third cranial nerve fascicle with contralateral hemiparesis due to injury of the corticospinal and corticobulbar tracts of the cerebral peduncle. Its causes range from ischemic or hemorrhagic processes, which damage the third cranial nerve fascicle and cerebral peduncle, to brain tumors, aneurysms, extradural hematomas and infectious diseases that behave like spreading processes.

The assessed patients showed clinical signs and symptoms such as ptosis; vertical double vision; difficulty standing up; hemiparesis; hyporeflexia; decreased superficial and deep sensation; poor balance and coordination; third cranial nerve palsy; slurred speech; intermittent disorientation in time, place and person; and facial asymmetry. Oral trimethoprim-sulfamethoxazole, clindamycin and prednisone were administered as specific treatment for toxoplasmosis, which enabled the improvement of the clinical picture.

This case report presents the sequence of events, treatment and a brief review of the literature to be considered in the diagnosis and determine its importance in patients with HIV-toxoplasmosis of the central nervous system.

Keywords: Brain Stem Infarctions; HIV; Toxoplasmosis (Source: MeSH NLM).

Síndrome alterno mesencefálico: síndrome de Weber en dos pacientes varones VIH positivos con toxoplasmosis cerebral

RESUMEN
En este reporte se describen dos casos de pacientes varones VIH diagnosticados por serología, que no estaban recibiendo tratamiento. Ambos pacientes desarrollaron el síndrome alterno mesencefálico y la lesión estructural, causada por la infección del sistema nervioso central por toxoplasmosis cerebral, confirmada mediante una resonancia magnética cerebral. Esta condición se constató por serología en líquido cefalorraquídeo.

El síndrome de Weber es un tipo de síndrome alterno mesencefálico poco frecuente y existe poca información de su clínica y evolución. Fue descrito por sir Herman David Weber en 1863, y se caracteriza por la lesión ipsilateral del fascículo del III par craneal, con presencia de hemiparesia contralateral debida a la lesión de la vía corticoespinal y corticobulbar del pedúnculo cerebral. Las causas que lo originan incluyen tanto procesos isquémicos o hemorrágicos, que lesionan el fascículo del III par craneal y pedúnculo cerebral, como neoplasias intraencefálicas, aneurismas, hematomas extradurales y procesos infecciosos que se comportan como procesos expansivos.

Los pacientes evaluados presentaron clínica de ptosis palpebral, visión doble vertical, dificultad para bipedestación, hemiparesia, hiperreflexia, sensibilidad superficial y profunda disminuidas, equilibrio y coordinación alterados, III par parético, habla incoherente, desorientación en tiempo, espacio y persona de manera intermitente y asimetría facial. Para la toxoplasmosis se aplicó un tratamiento específico con trimetoprim-sulfametoxasol, clindamicina y
INTRODUCTION

Midbrain stroke syndrome, described by Sir Herman David Weber in 1863, is characterized by ipsilateral injury of the third cranial nerve fascicle with contralateral hemiparesis due to injury of the corticospinal and corticobulbar tracts of the cerebral peduncle (1,2). Its causes range from ischemic or hemorrhagic processes, which damage the third cranial nerve fascicle and cerebral peduncle, to brain tumors, aneurysms, extradural hematomas and infectious diseases that behave like spreading processes (3,4).

Toxoplasmic encephalitis is the most frequent opportunistic infection that affects the central nervous system (CNS) of 26% of patients with HIV/AIDS and is associated with multiple focal injuries in the brain (5).

The criteria to diagnose cerebral toxoplasmosis are based on histology testing and laboratory tests to determine the definitive diagnosis—along with the signs and symptoms shown by patients and the injury evidenced by neuroimaging. Nevertheless, the presumptive diagnosis is more frequently based only on the signs and symptoms of patients, compatible injuries found with neuroimaging and the radiological response within 10-14 days after the empirical start of the anti-toxoplasma treatment (6). In these two cases we will describe the clinical presentation of two male HIV-positive patients with CNS toxoplasmosis in the Department of Neuroinfectious Diseases at Instituto Nacional de Ciencias Neurológicas (INCN, National Institute of Neurological Sciences) in Lima, Peru.

CLINICAL CASE I

This is the case of a 28-year-old male patient recently diagnosed with HIV, without antiretroviral therapy. He was admitted to the hospital with the following symptoms: ptosis of the right eyelid, vertical double vision, dragging of the left foot, inability to keep standing, left hemiparesis (muscle strength: upper limb 2/5, lower limb 2/5), left hypotony, deep tendon reflexes (DTRs): left hyporeflexia (++/+), decreased left superficial and deep sensation, poor balance and coordination, third cranial nerve palsy (75%), 5 mm right pupil unreactive to light and absence of consensual pupillary reflex. A contrast-enhanced magnetic resonance imaging (MRI) scan showed an image compatible with cerebral toxoplasmosis (Figure 1). Moreover, cerebrospinal fluid (CSF) serology testing for cerebral toxoplasmosis using chemiluminescence (IMMULITE 2000 Immunoassay System) found IgM antibodies > 1.1 with “reactive” result, which revealed an active infection. Therefore, a protocol-based treatment was prescribed at the institution and recovery was observed; the patient made good progress and his symptoms improved.

Figure 1. Contrast-enhanced brain MRI. Contrast uptake showed 3-5 mm images on the midbrain (white arrow).
CLINICAL CASE II

This is the case of a 40-year-old male patient diagnosed with HIV about a year before, without antiretroviral therapy and without controls. He went to the hospital because he had presented ptosis of the right eyelid and slurred speech for approximately four days. The assessment determined intermittent disorientation in time, place and person; left facial asymmetry; left hemiparesis (4/5); left hyporeflexia (++/+++); decreased sensation in the left hemibody; absence of meningeal signs; third cranial nerve palsy; 3.5 mm right pupil hyporeactive to light; disflueney and slurry speech. A contrast-enhanced MRI scan showed an injury compatible with cerebral toxoplasmosis (Figure 2). Moreover, CSF serology testing for cerebral toxoplasmosis using chemiluminescence (IMMULITE 2000 Immunoassay System) found IgM antibodies > 1.1 with “reactive” result, which revealed an active infection. Therefore, a protocol-based treatment was prescribed at the institution and recovery was observed; the patient made good progress and his symptoms improved.

DISCUSSION

HIV-seropositive patients may suffer many opportunistic infections, and cerebral toxoplasmosis is one of the most frequent in the CNS with a rate of about 26 %, particularly in countries with a high burden of infection and limited access to antiretroviral therapy \(^{(5,7)}\). In most of the cases in which patients live with HIV, cerebral toxoplasmosis manifests clinically with an HIV viral load over 50 copies/ml and CD4+ cell count < 100/mm\(^3\) \(^{(8)}\). The brain is one of the tissues preferred by toxoplasma because of its low inflammatory reaction and, in most cases, cerebral toxoplasmosis occurs because of the reactivation of a latent infection \(^{(6,10)}\). Midbrain injuries that cause ipsilateral palsy of the third cranial nerve and hemiplegia or contralateral hemiparesis are characteristic of Weber’s syndrome \(^{(11,12)}\). The occurrence of stroke syndromes in the context of HIV and cerebral toxoplasmosis is quite rare since their onset manifests with very specific symptoms such as homolateral ophthamoplegia and contralateral palsy of the lower face, tongue, arm and leg. The palsy of the medial rectus muscle causes ptosis of the upper eyelid; if the Edinger-Westphal nucleus is damaged, the pupil will be dilated and fixed to light and accommodation because of the damage to the third cranial nerve \(^{(2-4)}\). The possible etiologies of Weber’s syndrome are compression injuries of the interpeduncular fossa such as meningeal diseases, midbrain infarctions, aneurysms, brain tumors or infections, as shown in the cases presented in the literature, in which patients had CNS infection caused by cerebral toxoplasmosis with damage to the midbrain and the third cranial nerve \(^{(6,13)}\).

The patients received alternative and standard treatment established by the Ministry of Health of Peru. The following scheme was used: trimethoprim (TMP) 320 mg-sulfamethoxazole 1,600 mg bid, and clindamycin 600 mg every six hours for at least six weeks (duration could be extended if the clinical and imaging responses were incomplete after six weeks). In the clinical cases presented, the patients showed significant improvement between the sixth and eighth week since the onset of their neurological signs and symptoms. Also, they started highly active antiretroviral therapy (HAART) and were followed up for six months with maintenance therapy for toxoplasmosis until their CD4+ cell count improved \(^{(14,15)}\). These cases show the importance of considering midbrain stroke syndrome as a probable diagnosis in the context of an HIV infection and CNS toxoplasmosis, and making a correct diagnosis for the prompt treatment to minimize the possible sequelae.
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