Viral encephalitis complicated by acute retinal necrosis syndrome: A case report

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Abstract. Acute retinal necrosis syndrome (ARN) is a viral infection characterized by focal retinal necrosis. Viral meningitis complicated by ARN is relatively rare. In the present case study, a 44-year-old male presented with fever, headache and mental disorder. After four days, the patient developed blurred vision. The patient was diagnosed with viral encephalitis complicated by bilateral ARN, based on the examination results. After treatment with antivirals and systemic glucocorticoids, the symptoms of the patient improved.

Viral encephalitis may be an important risk factor for ARN. For a patient with viral encephalitis who experiences decreased visual acuity or vitreous opacification, the possibility of ARN should be considered.

Introduction

Viral encephalitis is an inflammation of the central nervous system caused by a specific virus. The prognosis is often good if treated early. Acute retinal necrosis syndrome (ARN) is a viral infection characterized by retinal focal necrosis (1). A previous study reported an incidence of 1 case of ARN per 1.6–2.0 million population (2). ARN is a viral inflammatory condition that manifests with vitreitis, severe retinal vasculitis and progressive peripheral retinal necrosis (3). Previous studies have shown that ARN may exist concurrently with viral encephalitis (4,5). The present study describes the case of a patient with viral encephalitis complicated by bilateral ARN. Written informed consent was obtained from the patient. The case report and a review of the literature were subsequently utilized to investigate the possible pathogenesis of this disease.

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Case report

A 44-year-old male patient was admitted to the Department of Neurology in the Affiliated Yantai Yuhuangding Hospital of Qingdao University Medical College (Yantai, China) on June 2, 2012 with a fever and headache that had persisted for 12 days, and confusion and divagation that had lasted for 1 day. The patient had no remarkable clinical or family history and had no history of drug allergies. Physical examination revealed that the patient was restless, irritable, had neck stiffness and was positive for Kernig's sign. The patient's deep tendon reflexes were normal and no paresis was present. Blood chemistry and urinalysis tests were normal. The results from the cerebrospinal fluid (CSF) analysis were as follows: leucocytes, 805.3 mg/l; monocytes (90%), 130x10^6/l; and herpes simplex virus I (HSV-I) immunoglobulin M, positive. Polymerase chain reaction analysis revealed positive results for HSV in the CSF and blood, while tests for cytomegalovirus (CMV) allotype, human immunodeficiency virus and hepatitis C were negative. The culture of CSF produced no acid-fast bacilli, common bacteria or Cryptococci. Based on these results, the patient was diagnosed with viral encephalitis and administered antiviral therapy (500 mg acyclovir intravenously every 8 h, dropwise).

Following antiviral therapy, the general condition of the patient improved; however, on June 6, 2012 he presented with blurred vision. Ophthalmological examinations revealed mild opacification of the lenses and moderate opacification of the vitreous body. The fundus appeared hazy and had blurred disc margins. Ophthalmic ultrasound showed a dense hypochrome focal lesion (Fig. 1). The mean visual evoked potential amplitude was slightly decreased and the time-to-peak was substantially prolonged. Methylprednisolone pulse therapy (1.0 g/day) was administered for three days, followed by orally administered prednisone. A second ophthalmological examination (June 16, 2012) revealed that vitreous opacification was present in both eyes. The majority of the retinal artery was occluded, particularly in the left eye. Retinal necrotic lesions with mild bleeding were visible.

The examination results confirmed the patient had viral encephalitis with bilateral ARN, and retinal laser treatment was administered. Two days after retinal laser treatment, visual acuity had improved. The CSF tests (June 18, 2012) were found to be normal. Fluorescein fundus angiography revealed that...
the majority of the retinal artery, shown as a white line, was occluded, particularly in the left eye (Fig. 2). Retinal necrotic lesions, together with mild bleeding, were visible along the white line. Visual acuity was markedly improved following treatment.

**Discussion**

Viral meningitis complicated by ARN is relatively rare, and is most common in immunocompromised individuals. Bilateral eye involvement occurs only in approximately one-third of patients with ARN (6). The present case study involved an adult male with viral encephalitis complicated by bilateral ARN. The mechanism by which viruses induce ARN has yet to be elucidated; however, it has been suggested that the virus may be transmitted along the brain-optic nerve axon-retina pathway (7,8).

Viral encephalitis may be an important risk factor for ARN. ARN syndrome is known to occur occasionally alongside, or shortly after, herpetic encephalitis (9). A review of the available literature (10), as well as our own observations, indicates that there is no universal guideline on how long antiviral and anti-inflammatory treatment should be continued. Early retinal laser treatment may prevent retinal detachment in patients with ARN (11). At present, a full course of antiviral therapy is essential to prevent the development of ARN as a result of viral
encephalitis, as well as to prevent the progression of ARN from one eye to the other (12). Systemic glucocorticoids may alleviate retinal inflammation, and have also been shown to protect the retina and optic nerve. Prophylactic laser coagulation of retinal defects and margins may also prevent retinal detachment (13).

In conclusion, if a patient with viral encephalitis experiences decreased visual acuity or has vitreous opacification, the possibility of ARN should be considered. Fundus fluorescein angiography should be performed and early active treatment should be applied.

References