Do Not Overlook Zona Herpes Ophthalmicus in a Patient Presenting with Pupil-Sparing Oculomotor Nerve Palsy

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Herpes Zoster (HZ), relatively common syndrome, is the result of reactivation of Varicella-Zoster Virus (VZV) that lying in the ganglion after a prior bout of varicella. Reactivation is usually caused by a decline in the specific cell-mediated immunity to VZV with aging, immunosuppression, trauma, and psychological stress [1].

In the HZ, various neurologic complications are seen. Those are postherpetic neuralgia (chronic pain), VZV vasculopathy, meningoencephalitis, meningoencephalitis, cerebellitis, myelopathy, and involvement of cranial nerves [2]. When HZ is involved in the ophthalmic branch of the trigeminal nerve, it is termed as Herpes Zoster Ophthalmicus (HZO). HZO accounts for 10-15% of all herpes zoster cases. Half of the HZO cases involve the ocular component like keratitis, cataracts, glaucoma, uveitis [3].

Oculomotor Nerve Palsy (ONP) could result from lesions anywhere along its path between the oculomotor nucleus and extraocular muscles. Differential diagnosis of ONP can be made according to presence of pupil involvement. Pupil sparing oculomotor nerve palsy typically result from ischemic cranial neuropathy, often associated with vascular risk factors, which improves (and usually fully resolves) within 3 months.

We present herein a patient exhibiting oculomotor nerve palsy associated with herpes zoster ophthalmicus.

**Presentation**

A 70-year-old man presented to the neurology outpatient clinic with acute onset of painful ptosis of his right eye. Past medical history revealed hypertension, hyperlipidemia and cerebrovascular disease. Family, social and surgical history was not significant. His blood pressure, pulse, respiration and body temperature were within normal limits. At first physical examination of patient showed complete ptosis and absence of supraduction, infraduction and adduction movements of eye ball. The pupils was equally reactive to light and isocoric. There were some squamatous and hyperemic cutaneous lesion along ophthalmic division of right trigeminal nerve, which is consistent with healing herpes zoster lesion (Figure 1). Visual acurity of both eye and rest of the examination were normal.

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Figure 2: Brain Magnetic Resonance Imaging (MRI) shown right peri-insular old infarct (red arrow).

Figure 3: Digital subtraction angiography (DSA) of brain vessel shown that there is no intracranial aneurism.
Brain Magnetic Resonance Imaging (MRI) revealed right peri-insular old infarct (Figure 2). There was no pathologic signal that considered as oculomotor nerve palsy. Digital subtraction angiography of brain vessel that was done 3 months ago due to acute stroke treatment was re-evaluated and revealed no brain aneurism (Figure 3). Blood biochemistry, hemogram, sedimentation was normal.

Ten days ago, the patient presented to the dermatology clinic with sharp pain and sudden onset vesicular cutaneous eruption at right periorbital region and was diagnosed as herpes zoster ophthalmicus. Oral valacyclovir treatment initiated within 24 hours of symptoms and continued for 10 days. Ophthalmologic examination was within normal limits.

Third nerve palsy associated with HZO was considered and deflazacort 60 mg/day was given and tapered. At three weeks later, patient symptom improve and at the 2 months, there was no residual symptoms (Figure 4).

Discussion

HZ is a localized disease characterized by unilateral radicular pain with vesicular eruption and nearly 1 in 3 people will develop it during their lifetime [1]. After thoracic dermatomes, cranial nerves are commonly affected by herpes zoster. Beside the sensory nerves, motor neurons very rarely could be affected by HZ. HZO occur between 10%-25% of all cases of herpes zoster and half of these cases affect the eye [3,4]. Ocular signs include conjunctivitis, keratitis, episcleritis, scleritis, uveitis, secondary glaucoma, cataract, and retinal necrosis [4].

Ophthalmoplegia is seen in 5%-31% of patients with HZO. The oculomotor nerve is the most common site amongst them and the trochlear nerve is the least. Complete ophthalmoplegia is seen very rarely. The extra-ocular muscle palsies usually are seen 2-4 weeks after the vesicular eruption, but sometimes occurs simultaneously with the eruption or more than 1 month later [3]. In our patient, there was painful-pupil sparing complete oculomotor nerve palsy that appeared 10 days after the rush.

The pathogenesis of oculomotor nerve palsy is controversial and several mechanisms have been suggested. Some of them are following; direct cytopathic effect of virus, immune response of the central nervous system to the virus, occlusive vasculitis induced by the virus and another latent neuropathic virus activation within the brain [5]. Given that the patient has vascular risk factors and pace of symptom recovery, occlusive vasculitis and microinfarction of oculomotor nerve could be responsible in our patient.

Treatment of herpes zoster ophthalmicus require systemic antiviral drugs. It is most effective if it is started within 72 hours of symptom onset. Administration of antiviral medications reduce viral spreading and ocular complications. The use of systemic steroid along with antiviral treatment is recommended because it may be effective to prevent occlusive vasculitis and reduce duration of acute pain [1]. This patient received valacyclovir for 10 days before the our presentation. After oculomotor palsy related to HZO was confirmed, we had started oral steroid for 3 weeks. It was shown that duration of ptosis and diplopia related to third nerve was from 2 months to 23 months. In our patient, pupil sparing oculomotor nerve palsy has improved and show complete recovery within the 2 months.

It is important to consider zona zoster as a reversible cause of oculomotor nerve palsy. Therefore physicians should examine the periorbital skin of patients and look for any healing lesion.

References