Syphilitic Optic Neuritis Initially Misdiagnosed as Ischemic Optic Neuropathy

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Purpose: To report a syphilitic optic neuritis case, initially misdiagnosed as arteritic ischemic optic neuropathy (AION), in which prior transient posterior placoid chorioretinitis was an important clue to the correct diagnosis.

Case summary: A 50-year-old man presented with blurry vision in the right eye. Funduscopy revealed optic disc swelling. Due to an elevated erythrocyte sedimentation ratio and C-reactive protein, our initial impression was AION. However, the diagnosis was corrected after reviewing a previous fundus photo revealing a large, pale yellow placoid lesion in the macula of the right eye. Serological examinations revealed confirmed syphilis infection. After a 2-week treatment with penicillin G, visual symptoms and signs fully resolved.

Conclusions: Optic neuropathy with an elevated erythrocyte sedimentation ratio and C-reactive protein should prompt suspicion for syphilitic optic neuritis. Misdiagnosis as AION could lead to steroid therapy without antibiotics, which can worsen prognosis.

Keywords: Acute syphilitic posterior placoid chorioretinitis; Ischemic optic neuropathy; Ocular syphilis; Optic neuritis; Syphilitic optic neuritis

Introduction

Syphilis is a sexually transmitted disease caused by the spirochete Treponema pallidum. Ocular syphilis is an uncommon but diagnostically important manifestation of the disease. It may be silent or present as anterior uveitis, chorioiditis, interstitial keratitis, retinal vasculitis, retinitis, optic neuritis, dacryoadenitis, or scleritis in both human immunodeficiency virus (HIV) positive and HIV negative patients [1]. Uveitis is reportedly the most common ocular manifestation of ocular syphilis [1]. Optic nerve involvement in syphilis, the more infrequent type of ocular syphilis, may be unilateral or bilateral, and becomes apparent as perineuritis, anterior or retrobulbar optic neuritis, or papilledema [2].

Delayed anti-treponemal therapy can result in permanent visual loss, and steroid therapy without adjunctive antibiotic treatment may increase the treponemal load. Thus, it is important to rule out treponemal disease at an early stage. However, the characteristics distinguishing syphilitic optic neuritis from nonsyphilitic involvement of a similar distribution are unclear.

We present a case of syphilis in an immunocompetent pa-
Patient who was initially misdiagnosed with arteritic ischemic optic neuropathy (AION). We belatedly reviewed the initial fundus photo provided by a clinic that the patient had previously visited and corrected the diagnosis to acute syphilitic posterior placoid chorioretinitis (ASPPC).

**Case Report**

A 50-year-old male presented with a 7-day history of significant visual loss in the right eye. Examination revealed a best-corrected visual acuity of 20/100 in the right eye and 20/20 in the left eye. The patient had diabetes and was taking diabetes medication. A right relative afferent pupillary defect (RAPD) was elicited. The Ishihara test showed decreased color vision. Funduscopy revealed optic disc swelling with blurring of the disc margin in both eyes, which was prominent in the right eye and otherwise unremarkable (Fig. 1A-D). There were no signs of inflammation in the anterior segments or the vitreous. Humphrey perimetry (24-2 Swedish Interactive Threshold Algorithm Standard) testing revealed an extensive central scotoma in the right eye, leading to a lack of reliability in the left eye (Fig. 1E, F). Initial laboratory findings revealed an elevated erythrocyte sedimentation ratio (ESR) and C-reactive protein (CRP).

The patient had visited another clinic 4 days previously. His visual symptoms had partially improved without any treatment. However, the vision in the right eye worsened, prompting the visit to our clinic. Due to the elevated ESR and CRP in our initial blood test, AION was the provisional diagnosis. For the AION work-up, we consulted with a rheumatologist about this case and further blood screening including platelets was performed. We considered systemic steroids and temporal artery biopsy and the patient was asked to return the following day with the data from the previous clinic visit.

The following day, we reviewed the fundus photo from the previous clinic, which revealed a large, yellow-white, placoid lesion of the right macula (Fig. 2A, B). A fundus fluorescein angiogram conducted at the previous clinic demonstrated early and late phase staining of the optic disc, but no clear sign of leakage. We also observed early central hypofluorescence and progressive late hyperfluorescence in the area of the placoid lesion (Fig. 2C, D). There were no signs of any retinal edema or serous detachment on macular optical coherence tomography (Fig. 2E, F). Thus, we had a high level of clinical suspicion regarding ASPPC, and conducted a rapid plasma regain test as a screening for acute syphilis infection. After a positive rapid plasma regain test (RPR; 1:128), we performed a T. pallidum particle agglutination assay and a serum fluorescent treponemal antibody absorption test to confirm the diagnosis and discontinued the work-up for AION. Finally, these confirmative tests revealed positive results.

A cerebrospinal fluid (CSF) tap revealed CSF protein and glucose within the normal range. The CSF white cell count was slightly elevated and the CSF RPR was negative. The patient did not have any systemic symptoms of syphilis, such as malaise, headache, nausea, or constipation. There were no other systemic manifestations including chancres, condyloma lata, macular papular rash, or lymphadenopathy. His personal, family and social histories were unremarkable and...
his HIV titers were negative.

A diagnosis of syphilitic optic neuritis accompanied by syphilitic posterior placoid chorioretinitis in the acute phase was made. As optic neuritis and retinitis are generally considered neurosyphilis and should be managed accordingly, he was admitted for a 2-week course of 1.2 g intravenous benzylpenicillin every 4 hours. Visual acuity, color vision, and optic disc swelling fully recovered after 2 weeks of penicillin treatment (Fig. 3). Three months after treatment, the RPR titer was reduced from 1:128 to 1:32.

Discussion

In patients with optic neuropathy and elevated ESR and CRP, a high degree of suspicion should be present for AION. Systemic steroids should be given immediately once AION is suspected. However, our case showed that infectious dis-

ease, especially ocular syphilis, should always be on the list of differential diagnoses in optic neuritis with elevated ESR and CRP. Misdiagnosis of ocular syphilis as AION would lead to unnecessary temporal artery biopsy and steroid therapy without adjunctive antibiotics, which would worsen the prognosis by increasing the treponemal load.

Ocular syphilis has diverse possible clinical signs and symptoms. It should be ruled out in all patients with ocular inflammation including optic neuritis. The sooner the treatment is initiated, the better the outcome, partly because secondary irreversible injuries like optic nerve atrophy or ganglion cell degeneration can be prevented. However, due to the lack of distinguishing characteristics of syphilitic optic neuritis and nonsyphilitic involvement, prompt diagnosis is often challenging, especially when there are no systemic manifestations, as in the present case.

There is often cellular activity in the posterior vitreous in anterior syphilitic optic neuritis. However, there were no such vitreous signs in our patient. Therefore, the ASPPC macular lesion identified in the present case was invaluable for diagnosis. ASPPC is a distinctive manifestation of ocular syphilis characterized by the development of a placoid macular deposit in the outer retina.

However, problematically, ASPPC can be evident only transiently. In our patient, the ASPPC had resolved sponta-
neously without systemic penicillin treatment. By the time of the visit to our clinic, this clinical clue was no longer apparent on a fundus exam. Therefore, even though there may be no abnormal funduscopic finding other than optic disc swelling, a high degree of clinical suspicion about syphilitic optic neuritis should be retained. As in the present case, review of the initial fundus photo or charts from previous clinics that a patient visited can detect this important but transient clue.

If a clinic is equipped with spectral domain-optical computed tomography (SD-OCT), detection of an already resolved ASPPC lesion is possible. SD-OCT imaging can reveal prominent nodular elevation of the RPE that can persist for up to 1 month after treatment [3].

In patients with ASPPC, circulating T. pallidum enters the choroidal circulation and gains access to the outer retina where the choroidal vascular supply is greatest, especially in the macula [4]. This can explain the presently observed simultaneous development of ASPPC and optic neuritis, because the posterior ciliary artery derived from the choroidal circulation is not only the main source of blood supply to the macular lesion but also the optic head, except for the nerve fiber layer.

Even though the outer retina of the macula and optic nerve were involved simultaneously in our case, ASPPC resolves quickly without treatment, while optic neuritis does not. The retinal pigment epithelium (RPE) may contribute to this difference. The RPE shares several functions with macrophages, including expression of Toll-like receptors, complement, and bacterial phagocytosis [5]. Since spirochetes are effectively eradicated by the engulfing activity of macrophages, we can deduce that phagocytosis of the RPE is also effective in eliminating this microorganism. SD-OCT imaging of ASPPC reveals irregular hyperreflectivity of the RPE with prominent nodular elevation [3]. This also suggests the possibility that RPE layers are crucially involved in the mechanism of ASPPC.

Even though capillaries of the optic nerve head have tight junctions and nonfenestrated endothelium, which represent a nerve-blood barrier against microorganisms, they are just passive physical barriers and do not have an active function like the RPE. Clinicopathologic examination combined with molecular analysis is required to confirm this hypothesis.

In conclusion, even though optic neuropathy with elevated ESR and CRP is an important feature of AION, a high degree of clinical suspicion regarding syphilitic optic neuritis is also vital in these patients. Misdiagnosis of syphilitic optic neuritis as AION can lead to steroid therapy without adjunctive antibiotics, which will worsen the prognosis. Even when there is no abnormal funduscopic finding other than optic disc swelling, previous fundus photographs or chart data should be reviewed. These efforts can hasten diagnosis and result in prompt treatment with benzylpenicillin in cases of transient ASPPC, as in our patient.

**Conflicts of interest**
There are no conflicts of interest.

**References**

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