Clinical Diagnosis of Chorioretinitis Sclopetaria in a Patient Presenting without a History of Trauma

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ABSTRACT

Chorioretinitis sclopetaria is a rare and unique manifestation of ocular trauma caused by high-velocity objects that pass adjacent to, but do not penetrate the globe. In these patients, anamnestic information is very helpful for diagnosis and it is not expected that a patient could completely forget the trauma as these traumas are generally severe. Diagnosis can be difficult in the case of patients who do not state trauma in anamnesis as in our case. It is crucial to recognize the path-ognomonic fundus findings in this rarely seen condition because these findings are invaluable in the absence of history of trauma. In this case report, we de-
scribe a case of chorioretinitis sclopetaria who was injured with a ball-bearing (BB) gun in childhood and did not realize.

Key Words: Chorioretinitis, choroid injuries, sclopetaria, trauma.

INTRODUCTION

Ocular trauma is one of the leading causes of unilateral visual loss in children and young adults. Non-penetrating trauma or closed globe injury where the corneoscleral wall of the globe remains intact may also be associated with various vitreoretinal complications such as choroidal rupture, optic nerve injury, macular hole, and retinal detachment (1). Chorioretinitis sclopetaria is a rare clinical presentation resulting from trauma caused by a high-velocity projectile object passing adjacent to or through the orbit without penetrating the globe (2,3). The pathophysiolog-ical mechanism of the injury is caused by the energy release resulting from the high speed of the projectile when reaching the orbital cavity and passing close to the eyeball in its path (4). The descriptive term “chorioretinitis” initially re-
flected to white areas seen on fundus examination that were thought to be non-inflammatory, but it is clear that the chorioretinal rupture is a result of trauma and not an inflammatory process (3,4). Despite widespread injury to the reti-
ina, retinal pigment epithelium and choroid, the retina often remains attached even without any treatment, as a result of chorioretinal scarring and the sub-
sequent adhesion between the retina and underlying tissues (5).

In this case report, we describe a rare case of chorioretinitis sclopetaria who was injured with a ball-bearing (BB) gun in childhood and did not realize and discuss the clinical findings and treatment options.

CASE REPORT

A 22-year-old male was referred to our clinic because of low vision in the right eye that was established during a military health and welfare inspection. He did not notice the lack of vision and ophthalmologic examination was not previously performed. Upon examination, the uncorrected visual acuity (VA) in the right eye was finger count at 1.5 meters and 10/10 in the left eye with normal ocular motility. The relative afferent pupillary defect (RAPD) was posi-
tive in the right eye. The intraocular pressure (IOP) measurements with Gold-
mann applanation tonometry were 15 mmHg in the right eye and 14 mmHg in the left eye. Anterior segment examination of the right and left eyes was nor-
mal. Fundus examination of the right eye revealed hyperplasia of the retina pigment epithelium, epiretinal membrane and subretinal membrane, extend-
ing from the macula and optic nerve to the peripheral retina (Figure 1). In-
traretinal-periretinal hemorrhage, macular scar, macular hole, retinal detach-
ment/hole, and atrophy-hemiatrophy of the optic disc were not present. Fund-
us examination of the left eye was normal. Optical coherence tomography of the right macula revealed hyper-reflectivity from the retinal pigment epithe-
lium (RPE) hyperplasia and loss of outer retinal layers adjacent to the RPE hyperplasia (Figure 2).
The patient's past history was negative for any medical or surgical diseases. When asked about trauma, his parents recalled that the patient applied to the emergency service with conjunctival hemorrhage complaint nearly 15 years ago. Topical antibiotic therapy was started with 2 weeks duration and conjunctival hemorrhage recovered after this treatment. Based on the history and ophthalmological examination findings, a diagnosis of chorioretinitis scleropetaria was suspected and posteroanterior and right lateral radiographs of the orbit were performed to confirm the diagnosis. X-rays revealed an intraorbital foreign body in the right orbit, and this was confirmed on computerized tomography (CT) images were analyzed. No surgical intervention was undertaken. The patient was warned about the potential risks of the metal foreign body adjacent to the optic nerve in the orbital apex especially during magnetic resonance imaging (MRI).

Chorioretinitis scleropetaria may be defined as a full-thickness break of the choroid, Bruch’s membrane, and retina as a result of a high-velocity missile striking or passing adjacent to, but not penetrating, the globe (5). Two mechanisms have been considered here: damage adjacent to the pathway of the missile is responsible for the direct injury, and the shock waves transmitted to the globe cause the indirect injury. The extent of ocular damage is determined by the missile’s proximity to the globe, its velocity and its size (6).

There are various eye problems accompanying scleropetaria at presentation, such as lid laceration, canthus laceration, subconjunctival hemorrhage, conjunctival laceration, hyphaema, vitreous hemorrhage, subretinal and macular hemorrhage, hemorrhage on optic disc, retinal edema, choroidal and retinal ruptures, optic nerve transaction and transection of the rectus muscle (2.5-8).

In the acute phase of chorioretinitis scleropetaria, the disease is easily diagnosed due to clear findings, intraorbital foreign body and history of trauma, while in the advanced period in situations where the patient forgets trauma on amnesia, hides trauma due to family or patient legal reasons or due to memory loss linked to the effect of trauma posterior segment, findings carry great importance. Recognizing these pathognomonic fundus findings can ensure the clinician suspects chorioretinitis scleropetaria in spite of the lack of trauma in patient history. Histopathologic studies about scleropetaria show partial loss of the nerve fiber and ganglion cell layers in the macula, temporal peripapillary and macular loss of photoreceptors with hypertrophy and hyperplasia of the RPE, an epiretinal membrane, and defects in Bruch’s membrane, choroid, and retina.

With time, these structures are replaced by loose and dense fibrous connective tissue (9). Besides, chorioretinitis scleropetaria is accompanied by bare sclera, typically. Scleral attachments to the choroid are practically nonexistent between the ora serrata and the equator and the choroid is under constant tension and will retract, exposing the bare sclera when cut (4). In the late period, a sharply demarcated area in the peripheral retina of blotches of pigment proliferation along with fibrosis, bare sclera, pale optic disc, macular pigmentary changes and retinal and choroidal tears is suggestive of chorioretinitis scleropetaria (6). Also, macular scar, choroidal neovascularization and fibrovascular proliferation are late complications developed in response to the damage to Bruch’s membrane (10).

The risk of retinal detachment following chorioretinitis scleropetaria is low and based on the proliferation of fibrous tissue, which causes firm adherence of retina and choroid to the sclera, along with the typically intact posterior hyaloid frequently seen in young patients, which prevents fluid access to the subretinal space and lowers the risk of subretinal neovascular membrane formation and acute retinal detachment (11,12). Ahmadabadi et al. published one of the largest chorioretinitis scleropetaria study series, examining 13 affected eyes in 13 patients. Among these 13 cases of chorioretinitis scleropetaria, only one patient had developed acute retinal detachment; the retina remained attached in the other patients throughout the follow-up period (6).

In the acute phase of chorioretinitis scleropetaria, hemorrhages (vitreous, retinal and/or subretinal), and choroidal and retinal ruptures can be seen, besides retinal edema (11). For differential diagnosis, ruptured globe, intraocular foreign body, commotio retina, choroidal rupture, and optic nerve avulsion should be considered. Ruptured globe and intraocular foreign body are excluded by clinical exam, B-scan ultrasonography, and computed tomography of the eye and orbit (13). It is known that the choroid ruptures are associated with blunt trauma, either by direct or indirect mechanism. Choroid, Bruch’s membrane and RPE injury are described.
Neurosensory retina is not involved, usually white or yellow crescent-shaped subretinal streaks are seen usually concentric to the optic nerve. Subretinal neovascular membrane may develop. Injury to Bruch’s membrane increase the risk of subretinal neovascularization development (10). Retinal and macula whitening with a cherry aspect (Berlin edema) have also been reported after eye injuries, a condition that is known as commotio retinae. Both can appear in chorioretinitis sclopetaria, as it also has a blunt component in its pathophysiology. In optic nerve avulsions, due to the accompanying anterior segment pathologies and appearance of optic nerve and retina with vitreous hemorrhage, diagnosis is obstructed. In these types of cases, imaging methods like electrodiagnostic tests, B-scan ultrasonography, CT or MR may be beneficial (13).

The late complication of the chorioretinitis sclopetaria is the same as chorioidal rupture in other situations, i.e., the possibility of developing neovascularization from the choroid and fibrovascular proliferation. This occurs in 25 to 50% one year after the chorioid ruptures (10).

CONCLUSION

Chorioretinitis sclopetaria is a rare manifestation of ocular trauma and has no specific treatment. In the acute phase, a detailed examination and recognizing the pathognomonic findings allow immediate diagnosis. As in our case, if patients do not state trauma in the anamnesis, accurate diagnosis can be difficult after a few years. After intense findings in the acute period regress, the appearance of the fundus should still contain findings reminiscent of severe trauma. It is important and necessary to recognize this characteristic fundus appearance, because an accurate diagnosis can prevent unwarranted surgical intervention.

Conflict of interest

No conflict of interest was declared by the authors.

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