Retinal Angiomatous Proliferation in Adult Onset Coat’s Disease

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Abstract

A 31 year old female patient had decreased vision in her left eye for several months. Visual acuity was 20/20 in right eye and counting fingers from 2 m in left eye. Examination of left eye demonstrated massive macular exudation with grayish discoloration of central macula and, aneurysmal and telangiectatic vessels with surrounding exudation in temporal mid peripheral retina. Optical coherence tomography of left eye showed findings consistent with neovascular complex, pigment epithelial detachment, subretinal fluid and involvement of choroid. There was hyperfluorescent focus with perfusing retinal arteriole and draining venule at the fovea which was compatible with retinal angiomatic proliferation (RAP) in early phase fundus fluorescein angiography and leakage was observed in late phase. After 3 doses of monthly-injected intravitreal ranibizumab and argon laser photocoagulation applied on aneurysmal and telangiectatic vessels, we observed central macular fibrosis, minimal lipid exudation and no macular edema. So retinal angiomatic proliferation may be a feature of adult onset Coats’ disease.

Introduction

Adult onset Coats’ disease is a rare, idiopathic and exudative retinal vascular disease without an inciting factor and with slow progression. It frequently occurs in males and is typically unilateral. Localized lipid deposition and hemorrhage around macroaneurysms are often present both at peripheral and juxtamacular areas [1,2].

Retinal Angiomatosis Proliferation (RAP) (type 3 neovascularization) emphasizes the intraretinal location of the neovascularization and originates not only from deep retinal capillaries but also from the choroid. It is characterized by telangiectatic response associated with intraretinal proliferative activity, leading to retinal-retinal and retino-choroidal anastomosis [3].

Recently RAP has been reported in 24% (5 of 21 patients) of childhood Coats’ disease by Sigler et al. [4]. But there is no adult onset Coats’ disease associated with RAP in the literature. Herein we describe the first adult onset Coats’ disease associated with RAP.

Case Report

A 31 year old female patient had decreased vision in the left eye for several months. She did not have any systemic diseases. Best corrected visual acuity was 20/20 in the right eye and counting fingers from 2 m in the left eye. Anterior segment examination did not reveal any abnormal findings. Fundus examination of right eye was normal. There was dense macular exudation with grayish discoloration of central macula and aneurysmal and telangiectatic vessels with surrounding exudation at temporal mid peripheral retina (Figures 1A and 1B).

After complete ophthalmological examination, optical coherence tomography (OCT) (Retinascan Advanced RS-3000; NIDEK, Japan) imaging and fundus fluorescein angiography (FFA) (Canon Inc., Tokyo, Japan) were performed. OCT demonstrated subretinal fluid, subfoveal neovascular complex extending into the choroid with disruption of RPE/Bruch membrane complex and hyporeflective vessel tracks in neovascular complex (black arrows) (Figure 1C). Early phase FFA showed hyperfluorescent angiomia with perfusing retinal arteriole (arrow) and draining venule (arrowhead) (retinal-retinal anastomosis) at fovea and there was leakage due to neovascular complex and filling of surrounding cystoid spaces in late phase (Figures 1D and 1E). And FFA of temporal midperipheral retina showed aneurysmal and telangiectatic vessels (Figure 1F).

After 3 consecutive monthly-injected intravitreal ranibizumab and argon laser photocoagulation applied on temporal aneurysms and telangiectatic vessels, there was central macular fibrosis, no macular edema and reduced lipid exudation (Figure 2A). OCT also revealed sharply demarcated borders of the subretinal fibrotic nodule with retinal vessel inside (black arrow), but there was no subretinal and intraretinal edema (Figure 2B). Hyperfluoresans caused by staining of this subretinal fibrotic nodule with a third-order arteriole entering into it by FFA (Figures 2C and 2D). Best corrected visual acuity was counting fingers from 4 m in the left eye.
Discussion

Adult onset Coats’ disease has retinal features different from the childhood onset Coats’ disease. Although retinal exudation and telangiectasia are hallmark for diagnosis as in childhood Coats’ disease, less extensive retinal involvement, more benign natural course and a more favorable treatment outcome were reported in adult onset Coats’ disease. In their series of 48 eyes, Rishi et al. [1] reported that decreased vision was the most common (83%) presenting sign probably due to more prevalent macular involvement in adult onset Coats’ disease. Macular involvement due to macular fibrosis, macular edema, macular exudation, epiretinal membrane, foveal ischemia, sensory macular detachment have been reported in adult onset Coats’ disease [1,5].

RAP has rarely been reported in conditions other than age-related macular degeneration, such as radiation retinopathy, syphilitic retinopathy, retinitis pigmentosa and childhood Coats’ disease [4,6-8]. Sigler et al. [4] reported presence of RAP lesion with chorioretinal anastomosis in 5 of 21 patients with childhood Coats’ disease. And Rishi et al. [9] also showed chorioretinal anastomosis in a 4-year old patient with Coats’ disease. Macular fibrosis in Coats’ disease reveals subretinal white-gray scar which is usually associated with vascular component at subfoveal area. Association of macular fibrosis with retinal-retinal anastomosis, intraretinal, subretinal and choroidal neovascularization were reported in previous studies [4,9,10]. Although factors initiating neovascularization in Coats’ disease are not clear yet, we agree with the previous manuscripts that severe macular lipid exudation leading to inflammatory stimulus, elevated vascular endothelial growth factor levels from ischemic peripheral retina and abortive defect during angiogenesis of retinal mid-capillary plexus can be possible causes [9,10]. Also we agree with Sigler and Rishi [4,9] that aberrant vascular communicating channels which are seen in Coats’ disease could favor formation of retinal-retinal and retina-choroidal anastomosis which are poor prognostic sign for treatment of neovascular complex.

As far as we know, this is the only case in the current literature who has RAP (type 3 neovascularization) lesion complicating adult onset Coats’ disease. Although indocyanine green angiography is needed for the diagnosis of RAP, spectral domain OCT and FFA when evaluated together can help for the diagnosis, as the case we reported herein.

Figure 1: A and B. Left eye had massive macular exudation with grayish discoloration of central macula (left) and aneurysms and telangiectatic vessels with surrounding exudation in temporal midperipheral retina (right). C. Optical coherence tomography reveals subfoveal neovascular lesion, subretinal fluid and extension of the neovascular complex into the choroid with disruption of RPE/Bruch membrane. Dark arrows shows retina vessels within the neovascular complex. D. Early phase fundus fluorescein angiography shows hyperfluorescent angioma with retinal-retinal anastomosis (black arrow: perfusing retinal arteriole, arrowhead: draining retinal venule). E. And increased hyperfluorescence due to neovascular lesion and filling of surrounding cystoid spaces in late phase. F. Fundus fluorescein angiography of temporal mid peripheral retina showed aneurysms and telangiectatic vessels (RPE: retina pigment epithelium).

Figure 2: A. After 3 consecutive monthly-injected intravitreal ranibizumab and argon laser photocoagulation applied on temporal aneurysms and telangiectasia there was central macular fibrosis, no macular edema and reduced lipid exudation seen in color fundus photography B. Optical coherence tomography demonstrated sharply demarcated borders of the subretinal fibrotic nodule, but no subretinal and intraretinal edema. Black arrow shows retinal vessel inside fibrotic nodule C and D. Fundus florescein angiography showed hyperflorescans caused by staining of this subretinal fibrotic nodule with a third-order arteriole entering into it.
Further studies would be required to understand this phenomenon more.

References