Congenital Retinal Macrovessel with Normal Visual Acuity: A Case Report

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Abstract
Congenital retinal macrovessel is a large aberrant vessel discovered as an incidental finding. These aberrant vessels are mostly single, unilateral veins. CRMs do not affect vision unless there are associated foveolar cysts, foveal ectopia, pigmentary changes at fovea, macular haemorrhage, exudates, serous macular detachment or the macrovessel crosses the fovea. Thorough clinical evaluation and investigations are valuable in diagnosis and follow-up of such cases and aid in timely management of associated complications. In this report, we have discussed a case of CRM with unaffected visual acuity. To the best of our knowledge, such a case of CRM crossing fovea but sparing vision and findings substantiated by Spectral Domain Optical Coherence Tomography (OCT) has not been documented earlier.

Keywords
Congenital retinal macrovessels, Aberrant macrovessels, Visual acuity, Fluorescein angiography, Optical coherence tomography, Retinal thickness, Retinal pigment epithelium, Central macular thickness

Abbreviations

Introduction
Congenital retinal macrovessels are a rare finding. They are usually discovered as an incidental finding. The aberrant vessel can be an artery or a vein, the latter being more common [1]. The visual acuity remains unaffected in majority of the cases [1,2]. In this paper we have reported a case of congenital retinal macrovessel crossing the fovea but not affecting the visual acuity.

Case Report
A 16 years old female presented to our clinic for routine ophthalmic evaluation. Best corrected visual acuity (BCVA) in both eyes was 20/20. Anterior segment in both eyes revealed no abnormal findings. Posterior segment evaluation of right eye revealed an aberrant vessel originating at the optic nerve head and traversing temporally across the macula and involving the fovea. The calibre of the macrovessel was comparable to that of the veins and had numerous branches crossing the horizontal raphe. The rest of the fundus findings in both the eyes were normal with cup: disc ratio of 0.4:1 (Figure 1).

Fluorescein angiography revealed the filling of the macrovessel starting in the arteriovenous phase as laminar flow with complete filling in the venous phase. The filling occurred earlier and emptying later than the rest of the veins. The macrovessel had numerous tributaries spread over the macula but the foveal avascular zone was just spared. There were no arteriovenous anastomoses or abnormal leakage points (Figure 2).

RTVue spectral domain OCT (Optovue Inc., Fremont, California, USA) analysis of the right eye revealed normal foveal contour in the radial scan. There was attenuation of signals in deeper retinal layers inferior to the fovea due to the hyper-reflective aberrant vessel. The retinal thickness maps of both the eyes revealed thickness within normal range. There was no evidence of foveal cysts, submacular detachment (Figure 3). Over 4 months of follow-up no changes have been noticed.

Discussion
Congenital retinal macrovessels (CRM) were first described by Mauthner in 1869 [3]. A century later in1969, Ashton explained the formation of congenital retinal macrovessels [4]. These vessels are of mesenchymal origin and develop around 15-16 weeks of gestation when differentiation of arteries and veins occurs. If foveal hypoxia occurs, vascular proliferation may reach foveola. CRM was defined by Brown in 1982 as the large aberrant vessel that crossed the horizontal raphe without affecting the visual acuity [1].

CRMs are mostly unilateral, single veins [1,5]. These vessels may supply or drain blood from both inferior and superior retina. In the case in discussion, the macrovessel was of venous origin as seen clinically and suggested by the laminar flow and complete filling in the venous phase on fluorescein angiography. Fewer cases have been reported of arteriolar origin of CRM. There have been reports of anomalous retinal arteries with leaking macroaneurysms causing macular oedema, haemorrhage and exudates and hence decreased vision [6,7]. A case of congenital retinal arterial macrovessel associated with congenital simple hamartoma of retinal pigment epithelium has also been reported [8].

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associated with normal visual acuity [1,9,10]. Visual status tends to remain stable over long follow up periods [9-11]. Brown et al first reported 7 eyes with CRM all having normal vision [11]. de Crecchio et al have followed a case with impaired but stable visual acuity over 14 years [11]. The current case presented with normal visual acuity which has remained stable over 4 months of follow-up.

Figure 1: Right eye - Retinal macrovessel originating at optic nerve head and crossing the fovea just inferior to foveola; Left eye - Normal retinal findings.

Figure 2: Fluorescein angiographs of right eye showing early filling of CRM with laminar flow and delayed emptying; no leakage, no arteriovenous anastomoses. Left eye shows normal findings.
Visual acuity when affected can be attributed to the macrovessel crossing the foveola, foveolar cyst, foveal ectopia, pigmentary changes at fovea, macular haemorrhage, exudates or serous macular detachment [1-5,9,12,13]. In our patient, foveola was surrounded but not traversed by the macrovessel and its tributary and none of the above associations were noticed. Other proposed causes of decreased vision include macular thickening and distorted foveal architecture as seen on SD OCT [14,15]. Visual field analyses have revealed relative angioscotomas caused by decreased retinal sensitivity in the area crossed by the macrovessel [15,16]. In our subject, OCT revealed the shadow effect of the hyper-reflective vessel with normal foveal contour and no intraretinal or subretinal fluid. The macular thickness was within normal range in both the eyes.

Few authors have reported the occurrence of central serous chorioretinopathy in such cases [17-20]. The decompensation of an otherwise stable macrovessel has been associated with the alteration of pressure (Valsalva manoeuvre) or gravitational forces (roller-coaster rides or bungee jumping) [21,22]. Branch retinal artery occlusion occurring as a result of ischemic decompensation of CRM has also been discussed recently [23]. Amblyopia can occur in children by these vessels crossing fovea [24].

Fluorescein angiography reveals certain characteristic findings of CRMs which may include early filling and delayed emptying of the vessel, arteriovenous anastomoses, capillary capillary non perfusion, hyperfluorescence associated with RPE alterations and vascular leakage [1,2,5]. Except the early filling and delayed emptying, none of the other findings were seen in the case in discussion. CRMs can mimic certain vascular anomalies like capillary hemangiomas, racemose angiomas, retinal venous collaterals following venous occlusions and tumours like malignant melanoma and retinoblastoma [1,3,6]. Cavernous hemangiomas have also been found to coexist with such macrovessels [25]. Microvascular anomalies with CRMs have been reported in a case of neurofibromatosis-I [26]. In our case no anomalies were noted in the retinal vasculature. There also exists a possibility of brain vascular anomalies like Wyburn-Masson syndrome so neurological studies should be considered [1,3,5].

The limitation in our case was the visual field analysis, though performed by the young subject could not fetch us useful results due to highly unreliable indices. The follow-up period of 4 months is too short and it would be interesting to look for any new developments over a longer duration.

**Conclusion**

CRMs are a rare and incidental finding. Thorough clinical examination, fluorescein angiography and OCT are essential in the diagnosis and management of such cases. Although a rare occurrence, visual acuity may remain unaffected even with the macrovessel crossing the fovea. Regular follow-up is an important aspect to appropriately manage cases developing complications associated with CRMs.

**Acknowledgement**

I am grateful to Dr. Mohit Bhatia for providing assistance in writing and editing this manuscript.

**References**