Case Report

Orbital Lymphatic-Venous Malformation Accompanied by an Intraocular Vascular Malformation: A Rare Case Study

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Abstract
Lymphatic-venous malformations (LVMs) are development defects that result in abnormal connections between the lymphatic and venous systems. The authors describe a 7-weeks-old female infant who presented with a right orbital LVM extending to the ipsilateral cheek and subconjunctiva of the right eye, intracranial developmental venous anomalies in the right cerebellum, and a significant right eye intraocular retinal vascular malformation. Since orbital LVM is usually diagnosed in infancy or childhood, pediatric ophthalmologists should actively look for intraocular vascular malformations as such findings can poorly affect a patient’s vision.

Introduction

Lymphatic-venous malformations (LVMs) [1], also known as lymphangiomas, are a subgroup of slow-flow vascular malformations [2]. They are developmental defects caused by an improper connection between the lymphatic and venous system or abnormal development
of lymphatic vessels. LVMs are considered benign tumors, usually present at birth, that continue to grow at the normal rate of endothelial turnover throughout the subject’s life [3]. The most common (~60%) locations of LVMs are in the head and neck, but they can be found in any body part where lymphatic vessels are present [4–6].

Orbital LVMs account for about 1–3 percent of orbital tumors [7]. They typically present in childhood in the subconjunctival and periorcular tissues. However, orbital LVMs can extend to the frontotemporal region and the cheek and may be associated with intracranial vascular malformations, such as venous anomalies [8–10]. Lesions can be asymptomatic until they reach a large size, or intralesional hemorrhage occurs and causes swelling, proptosis, decreased motility of the eye, or pain [11]. Even though they are benign, these lesions can threaten vision by causing a compressive optic neuropathy [7]. We present a rare case in which an orbital LVM is accompanied by a significant retinal vascular disorder in the ipsilateral eye.

Case Report

A 7-weeks-old female was referred to our clinic because of an abnormal red reflex of the right eye. General development was appropriate to age. Pregnancy with the patient was achieved via IVF and sperm donation. The subject’s mother, Jewish of Iraqi-Ashkenazi origin, and sperm donor of Danish origin have no known ocular disorders (self-reported). Delivery was induced at 39 weeks of gestation due to preeclampsia and was traumatic (vacuum-assisted delivery, fractured clavicle, and meconium-stained amniotic fluid). The birth weight was 3,490 g.

On her exam, the subject blinked to light and had normal anterior segments in each eye. After dilation, a dense cloudy vitreous was seen in the right eye that obscured the posterior pole and did not enable visualization of the optic nerve or any other details of the posterior pole (shown in Fig. 1). The left eye had normal optic disc appearance, normal foveal reflex, and attached retina.

Examination under anesthesia (EUA) revealed a normal anterior segment in both eyes. After pupil dilation, RetCam images were taken, and an ultrasound showed an elevated, dome-shaped lesion (mass) in the macular area compatible with organized vitreous hemorrhage. Left fundus appeared normal.

Due to swelling of the right cheek (shown in Fig. 2), the patient underwent MRI, magnetic resonance angiography, and magnetic resonance venography of the brain and orbits. Imaging revealed a LVM in the right orbit extending anteriorly to the soft tissues of the right cheek and

Fig. 1. Right eye fundus photo demonstrating central opacity blocking visibility of the posterior pole.
posteriorly to the right cavernous sinus. Infiltration of the LVM behind the eyeball caused a slight flattening of the posterior aspect of the eyeball, along with slight forward displacement of the eyeball. In the brain, there was a venous anomaly in the right cerebellum (shown in Fig. 3).

The patient underwent a vitrectomy in her right eye to clear the old dense vitreal hemorrhage without any further interactions during surgery. Subsequent fluorescein angiography demonstrated a subretinal lesion in the posterior pole accompanied by capillary nonperfusion and microaneurysms at the retinal periphery 360° (shown in Fig. 4). Therefore, laser coagulation to the retinal periphery was performed.

A follow-up EUA also revealed a transparent nasal subconjunctival mass compatible with LVM (shown in Fig. 5), a central macular scar, and peripheral retinal laser scars in the right eye. The patient also underwent genetic evaluation. Whole-exome sequencing tests of the subject and her mother revealed a maternal inherited NM_001256071.1:c.13798-2A>C variant in the $RNF213$ gene (see online suppl. material 1; for all online suppl. material, see...
The mother had a normal physical exam but did not complete the recommended clinical workup, including a dilated eye exam. However, she has good visual acuity in both eyes.

The patient is now almost 2 years old and still continues regular follow-up visits in our clinic. She is continuously treated for amblyopia and seems to develop useful visual acuity in the right eye. She did not undergo any other surgical procedure to date.

**Discussion**

Presented is a unique case of co-occurrence of facial, orbital, and subconjunctival LVM, associated with a venous anomaly, and accompanied by an intraocular retinal vascular anomaly and vitreous hemorrhage in the ipsilateral eye of a baby girl with a genetic molecular change in her RNF213 gene. In the past, it was assumed there were no lymphatic vessels in the eye [12]. However, there have been significant advances in lymphatic research in recent years. Studies now support the presence of lymphatic components in the central nervous system, orbit, and inside the eye [13–15]. Yücel et al. [13] demonstrated the presence of lymphatic vessels in the ciliary body and confirmed drainage of radioactively labeled albumin from the anterior chamber to cervical, retropharyngeal, submandibular, and preauricular lymph nodes. Dickinson and Gausas [14] demonstrated the presence of lymphatic vessels in the dura surrounding the optic nerve.

Katz et al. [8] previously reported the association of orbital LVMs with noncontiguous intracranial venous anomalies in 28 percent of patients primarily diagnosed with orbital LVM. This association indicates a propensity for multifocal anomalous vascular formation. In their paper, the authors perceive LVM of the orbit as a spectrum of multicentric venous LVMs, probably depending on time of occurrence during embryonal development.

Two cases previously published [16] described orbital LVM accompanied by iris venous malformation, abnormal branching of retinal vessels, abnormal foveal reflex, and poor vision. The currently presented patient had extensive retinal vascular involvement that included the macula and the periphery, probably representing an extreme scenario of the continuum suggested. We suggest that the same embryonal sequence that induced the abnormal vascular development on the same side of the brain, orbit, and face also influenced the intraocular vasculature anomalous development, which caused very early vitreal hemorrhage and capillary nonperfusion.

Finally, our patient had a genetic variant in the RNF213 gene. Mutations in RNF213 are known to be associated with moyamoya disease, an idiopathic vascular disorder of intracranial

![Fig. 4. FA of the right eye demonstrating subretinal lesion in the posterior pole after vitrectomy and retinal periphery CNP (a); FA of the right eye post vitrectomy demonstrating macular scar and peripheral laser therapy scars (b); and fundus photo of the right eye post vitrectomy demonstrating macular scar and peripheral laser therapy scars (c). FA, fluorescein angiography; CNP, capillary nonperfusion.](image)
arteries. The inheritance may be autosomal recessive or autosomal dominant with incomplete penetrance. The c.13798-2A>C variant occurs in a highly conserved splice site area that has not been described previously in the literature. This variant has a very low frequency in healthy populations, 1: 0.0000041, as estimated by gnomAD. Altogether, the finding was concluded as a variant of unknown significance. Relevant clinical workup was recommended to the mother but has not yet been completed. Interestingly, RNF213 knockdown zebra fish showed severely abnormal sprouting vessels in the head region [17, 18]. However, no direct association has yet been found between this gene and LVMs in humans.

In conclusion, we suggest that in cases of diffuse orbital LVM, retinal vascular abnormalities should be prospectively sought. Since LVMs are most frequently diagnosed in childhood, EUAs, including fluorescein angiographies, should be undertaken. Further clinical investigation is needed to understand the frequency and severity of intraocular anomalous vascular component as part of diffuse orbital LVMs associated with venous anomalies, and the significance of the variant in the RNF213 gene.

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**Statement of Ethics**

Written informed consent was obtained from the patient’s parent, who is her legal guardian, for publication of this case report and any accompanying images.

**Conflict of Interest Statement**

The authors declare that they have no competing interests.
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Author Contributions

Ophthalmological material preparation and data collection were performed by K.S.B., I.Y., O.G., R.E., A.S., and M.E. Radiological material preparation and data collection were performed by L.K. Genetical material preparation and data collection were performed by M.R. The first draft of the manuscript was written by K.S.B. and M.E., and all the authors commented on previous versions of the manuscript. All the authors read and approved the final manuscript.

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