Hamartoma simple congenital of retina, with OCT and ultrasonorography evaluation

Case Report

An 11-year-old girl consulted with her parents and reported that the girl had low vision with her right eye. Her parents reported that the girl did not have trauma to that eye nor had a history of eye inflammation. Snellen’s visual acuity was 20/80 in her right eye and 20/20 in her left eye. In the biomicroscopy, there were no significant alterations, the ocular pressure remained within normal limits (14 mmHg in both eyes) and, in the fundoscopy of the right eye, revealed a circumscribed lesion in the paramacular area, with irregular borders, nodular type and dark brown pigmentation affecting the RPE, which presented slight hyperplasia, with an increase in retinal thickness in the lower paramacular area, with minimal invasion into the vitreous cavity. Optical coherence tomography (OCT) in this case showed increased margins of the tumor as well as the adjacent retina. All the information that was obtained in the evaluation of ocular tumors allows to rule out other vascular pathologies that could affect vision, such as the presence of iris neovascularization or PVR, and to obtain a differential diagnosis of these pathologies with the hamartoma, which is supposedly congenital, with stable presentation and located in the lower paramacular area, favoring this diagnosis to maintain an acceptable visual acuity and a good prognosis. The lesion is not the cause of low vision, it is located in the paramacular area away from the fovea, but the hyperplasia of the EPR permits unobstructed vision.

Discussion

The retinal Hamartoma was named by Dr. Donald M. Gassen in 1989 and described this pathology as raised, superficial and full thickness. The lesion presented in this case describes it as a tumor lesion with hyperplasia components of the RPE, without apparent vascularization. The hamartoma is supposedly congenital, with stable presentation and located in the lower paramacular area, favoring this diagnosis to maintain an acceptable visual acuity and a good prognosis. The lesion is not the cause of low vision, it is located in the paramacular area away from the fovea, but the hyperplasia of the EPR permits unobstructed vision.

Keywords: eye, tumor, retina, congenital simple hamartoma, retinal pigment epithelium, optical coherence tomography, diagnosis

References


Abstract

**Purpose:** To report in detail all the findings of optical coherence tomography (OCT) and ultrasound in a case of simple congenital hamartoma of the retinal pigment epithelium (RPE).

**Design:** Observational case report.

**Methods:** Fundus examination, retinography, ultrasonography and OCT were done in one case of simple congenital hamartoma of the retinal pigment epithelium.

**Results:** a considerable, paramacular, solitary and highly pigmented lesion was observed in the right eye of an 11-year-old girl. Visual acuity was 20/60 in the eye with paramacular lesion. The ultrasound showed a hyperreflective nodule with echoes of medium and high reflectivity. The OCT revealed a high reflectivity of the surface of retinal mass with full increase thickening, with abrupt and total shadow of the optical transmission. It presented a solid vitreomacular adhesion on the border of tumor lesion. Macular thickening did not affect the surrounding RPE and choroid.

**Conclusion:** OCT is an important diagnostic method, useful and non-invasive to evaluate this type of tumor lesion. This diagnostic study may show in some cases (and this in particular), a vitreoretinal adhesion with thickening of borders of the tumor and adjacent retina.

This type of tumor (simple congenital hamartoma of the retinal pigment epithelium (RPE) is a fairly rare lesion of low prevalence. Diagnosis is usually made casually in healthy and asymptomatic children, as well as in young adults, during a routine fundoscopy in general ophthalmological consultation. Optical Coherence Tomography (OCT) has recently become an important tool for the evaluation of retinal anatomy and retinal abnormalities. Few reported cases have been written about the role of OCT in the evaluation of ocular tumors. In this report, we describe the OCT findings of this highly pigmented retinal tumor.

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through the diagnostic methods together with the bibliography found confirms that it is a rare and infrequent tumor.

Figure 1(A) Retinography showing a heavily pigmented solitary macular lesion.

Figure 1(B) Ultrasonography showed a nodular high-echogenic mass at the macula.

Figure 1(C) OCT revealed prominent surface reflectivity of the full thickness retinal mass with abrupt and complete shadowing of optical transmission.

Figure 2 Retinography showing a pigmented solitary macular lesion.

Figure 3 Angiography showing a staining in the macular area.

Conclusion

OCT is an important diagnostic method, useful and non-invasive to evaluate this type of tumor lesion. This diagnostic study may show in some cases (and this in particular), a vitreoretinal adhesion with thickening of borders of the tumor and adjacent retina.

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Acknowledgment

None.

Conflicts of interest

Author declares there is no conflict of interest towards the manuscript.

References