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CASE REPORT

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Submillimeter retinoblastoma monitoring following transpupillary thermotherapy using hand-held optical coherence tomography

Mark McGarrey, Thamolwan Surakiatchanukul and Carol L. Shields*

Abstract

Background: Precise, submillimeter visualization of retinal microstructures is useful for treatment monitoring of retinoblastoma. Herein, we report the use of hand-held optical coherence tomography (HH-OCT) to document a nearly-invisible retinoblastoma and monitor tumor response to transpupillary thermotherapy (TTT).

Case presentation: A 3-week-old boy was diagnosed with unilateral familial retinoblastoma in the left eye, classified as group B, and treated with intravenous chemoreduction. At 13-month follow-up, the tumor in the left eye was regressed, and evaluation of the right eye revealed a microscopic, nearly invisible tumor measuring 372 μm in thickness and 1.51 mm in basal dimension. The tumor was confirmed on HH-OCT and the diagnosis was changed to bilateral familial retinoblastoma. The new tumor was subsequently treated with TTT and after 1 month, the tumor regressed to an optically dense scar 166 μm in thickness by HH-OCT. Five months after TTT, the scar was clinically flat and measured 73 μm .

Conclusions: HH-OCT has assumed a significant role in the monitoring of retinoblastoma in children. In this case, imaging with HH-OCT allowed precise localization of the tumor and submillimeter monitoring of treatment response following TTT.

Keywords: Eye, Retina, Retinoblastoma, Chemoreduction, Regression, Transpupillary thermotherapy, Optical coherence tomography

Background

Optical coherence tomography (OCT) is an imaging technology that utilizes the predictable properties of light refraction to produce precise cross sectional images of the human fundus [1]. Following the popularization of OCT since 1991, OCT technologies including spectral domain OCT (SD-OCT) have allowed for non-invasive, higher-resolution evaluation of retinal and choroidal microstructures. Subsequent technological advancements have given rise to hand-held OCT (HH-OCT), a portable SD-OCT unit, that can be

used intra-operatively to image the eyes of children and non-cooperative patients [2]. HH-OCT has proven particularly useful in detection and monitoring of small or “invisible” parafoveal retinoblastomas as well as estimating visual acuity potential in preverbal children [3–5].

Herein, we report a 3-week-old boy with familial retinoblastoma who was detected to have a minimally elevated submillimeter tumor, confirmed on HH-OCT. Following treatment, documentation of tumor regression by HH-OCT was important in confirming complete tumor response.

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Case presentation

A 3-week-old white male, with a family history of maternal retinoblastoma, presented for clinical evaluation after fetal ultrasound revealed a tumor in the left eye (OS). On examination, visual acuity was fix and follow in each eye and leukocoria OS was noted. Funduscopic evaluation right eye (OD) showed no tumor, whereas fundus evaluation OS (Fig. 1a) revealed a macular retinoblastoma measuring 10.0 mm in basal dimension and 5.0 mm in thickness, classified as group B familial retinoblastoma. The patient received six cycles of intravenous chemoreduction (CRD) using vincristine, etoposide, and carboplatin. Following CRD, the right eye remained normal and the left eye (Fig. 1b) showed type III regression, with partial calcification and regressed tumor measuring 6.0 mm in basal dimension and 2.1 mm in thickness.

At a 13-month follow-up, a new, minimally elevated parafoveal retinoblastoma measuring 1.51 mm in diameter located 6.0 mm temporal to the foveola OD (Fig. 2a) was detected by indirect ophthalmoscopy. The left eye remained stable with a partially calcified scar. By HH-OCT, the foveola OD was intact and the tiny parafoveal retinoblastoma measured 372 μm in thickness and was without subretinal fluid (Fig. 2b). Ultrasonography confirmed the tiny retinoblastoma OD of 1.5 mm thickness and the regressed calcified retinal scar OS of 2.1 mm thickness.

The diagnosis was changed to bilateral familial retinoblastoma, and the newly-diagnosed retinoblastoma OD was treated with transpupillary thermotherapy (TTT) using 300 mW power for 4 min with grey-white uptake. One month later, retinoblastoma regression was noted (Fig. 2c) and HH-OCT (Fig. 2d) revealed thickness reduction to 166 μm . Five months after TTT, the scar was clinically flat (Fig. 2e), measuring 73 μm on HH-OCT (Fig. 2f). There were no further tumors.

Discussion

Imaging technologies play a significant role in the detection, diagnosis, and selection of therapeutic options for retinoblastoma. Previously, cross-sectional imaging of retinoblastoma was only possible with ultrasonography,

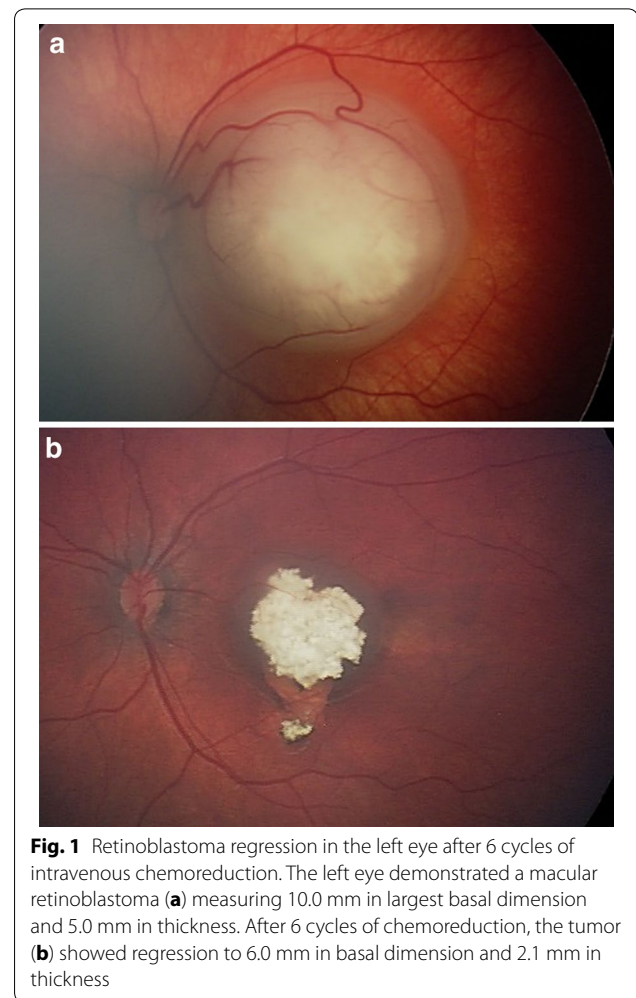


Fig. 1 Retinoblastoma regression in the left eye after 6 cycles of intravenous chemoreduction. The left eye demonstrated a macular retinoblastoma (a) measuring 10.0 mm in largest basal dimension and 5.0 mm in thickness. After 6 cycles of chemoreduction, the tumor (b) showed regression to 6.0 mm in basal dimension and 2.1 mm in thickness

computed tomography, or magnetic resonance imaging. These techniques were only able to detect medium to large retinoblastoma and provide information on tumor configuration, location, size, and extrascleral extension. However, high resolution HH-OCT has recently become available for detection of small retinoblastomas as depicted in Table 1 [3–8], and recently even those that are sub-millimeter or nearly clinically invisible, as in this case.

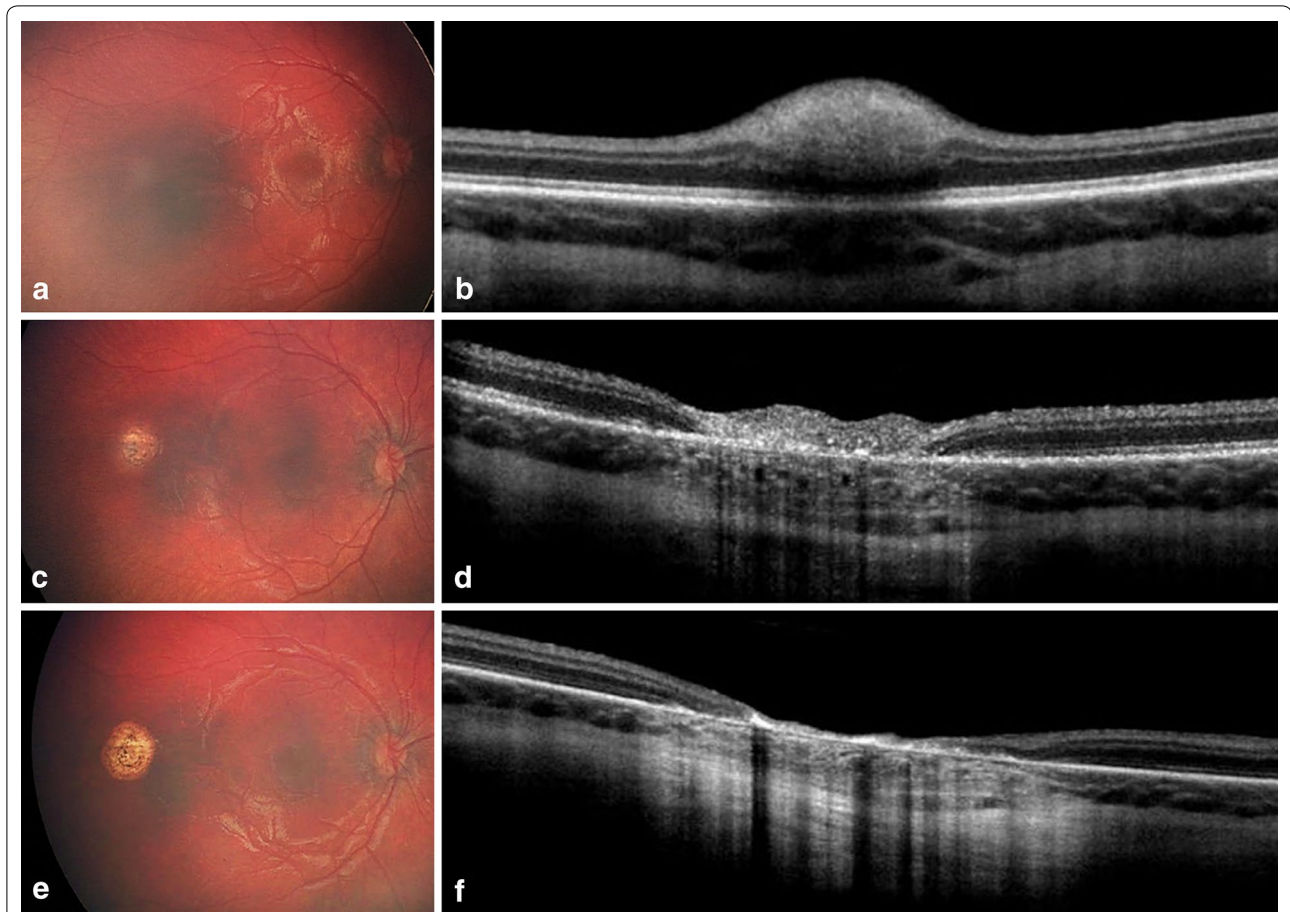


Fig. 2 Retinoblastoma regression documented on hand-held optical coherence tomography (HH-OCT) in the right eye. The initially unaffected right eye (**a**) demonstrated a small retinoblastoma temporal to the fovea of 1.51 mm in basal dimension and 372 μm in thickness (**b**). One month following transpupillary thermotherapy (TTT), the tumor (**c**) showed regression, confirmed as a (**d**) hyperreflective scar on OCT of 166 μm thickness. At 5 months after TTT, the flat scar (**e**) measured 73 μm thickness on HH-OCT (**f**)

HH-OCT is a portable SD-OCT unit, particularly useful for imaging young children and uncooperative patients. In 2004, Shields et al. [9] reported that time domain OCT was more sensitive than clinical examination in detection and monitoring of a variety of macular pathology in children such as cystoid macular edema and

subretinal fluid. Further studies have revealed that HH-OCT is important for monitoring tumor regression [10], detecting subclinical recurrence [6], identifying invisible retinoblastoma [4, 5], and elucidating foveal microanatomy in children following retinoblastoma treatment [11].

Table 1 Comparison of published cases of small retinoblastomas documenting tumor detection, regression, or recurrence by optical coherence tomography

Authors	Patient demographics	OCT findings					Descriptions	Machine
		Tumor diameter (mm)	Tumor thickness (μm)	Distance from foveola (mm)	Distance from optic nerve (mm)			
Cao et al. [3]	33-month-old male	8.0 (6.5) ^a	4000 (2200) ^a	0.0 (0.2) ^a	Unspecified (0.0) ^a	Exophytic mass draped by normal retina	iVue Optovue with iStand	
	9-month-old female	12.0 (8.0) ^a	6300 (3000) ^a	0.0 (0.5) ^a	Unspecified (3.5) ^a	Exophytic mass draped by normal retina		
	7-month-old male	11.0 (6.0) ^a	4000 (2600) ^a	0.0 (0.0) ^a	Unspecified (2.5) ^a	Exophytic mass draped by normal retina		
Saktanasate et al. [4]	2-month-old male	Unspecified	Unspecified	Unspecified	Unspecified	Ovoid; Outer nuclear layer	iVue Optovue	
Berry et al. [5]	7-week-old male	Unspecified (3 tumors)	Unspecified	Unspecified	Unspecified	Dome-shaped; Outer nuclear, outer plexiform, and inner nuclear layers	InVivoVue, Biop-tigen	
Park et al. [6]	3-week-old female	2.10	639	1.85	Unspecified	Dome-shaped; Outer nuclear layer ^b	iVue Optovue	
Soliman et al. [7]	Unspecified	Unspecified	Unspecified	Unspecified	Unspecified	Ovoid ^b ; Inner nuclear layer	Biop-tigen Envisu C2200/C2300	
Yarovaya et al. [8]	3-week-old female	2.75	792	0.62	Unspecified	Dome-shaped ^b ; Outer retinal layer ^b	iVue Optovue	
Current report	3-week-old male	1.51	372	6.0		9.0 Dome-shaped; Inner retinal layer	iVue Optovue	

Reports from peer-reviewed journals in 2014-17 were included

^a Dimensions were re-measured after therapy

^b Per our interpretation of the published images

Conclusions

Imaging with HH-OCT was helpful in this case as it confirmed tumor presence and allowed for precise, submillimeter monitoring of tumor thickness and regression, with preservation of foveal microanatomy. We encourage all clinicians who manage retinoblastoma to consider the use of HH-OCT for best monitoring of even the tiniest tumors.

Abbreviations

OD: Right eye; OS: Left eye; OCT: Optical coherence tomography; SD-OCT: Spectral-domain optical coherence tomography; HH-OCT: Hand-held optical coherence tomography; CRD: Chemoreduction; TTT: Transpupillary thermotherapy.

Authors' contributions

MM, TS, and CS participated in the design of the study. MM and TS collected data and drafted the manuscript. CS helped draft and finalize the manuscript. All authors read and approved the final manuscript.

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None.

Competing interests

The authors declare that they have no competing interests.

Availability of data and materials

All data generated or analyzed during this study are included in this published report.

Consent for publication

Consent for publication has been obtained from the parent or legal guardian of the study patient.

Ethics approval and consent to participate

The present study has been performed in accordance with the Declaration of Helsinki.

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