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Case report

### Projection-resolved optical coherence tomography angiography exhibiting early flow prior to clinically observed retinal angiomatous proliferation

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CASE REPORTS

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#### ABSTRACT

*Purpose:* The purpose of this study is to analyze early retinal angiomatous proliferation (RAP) utilizing a novel imaging modality, Projection-Resolved Optical Coherence Tomography Angiography (PR-OCTA). *Observations:* Five months prior to the diagnosis of a RAP lesion, cross-sectional PR-OCTA demonstrated flow in the outer retina contiguous with the deep retinal capillary plexus (DCP) and adjacent to a small pigment epithelial detachment. After development of a clinically visible RAP lesion, cross-sectional PR-OCTA demonstrated the RAP lesion connecting DCP and sub-retinal pigment epithelial neovascularization.

*Conclusions & importance:* This is the first report of PR-OCTA demonstrating abnormal flow in the outer retina prior to the development of a clinically detectable RAP lesion. PR-OCTA may be useful for surveillance and to help further characterize and stage RAP lesions.

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#### 1. Introduction

Retinal angiomatous proliferation (RAP) or type 3 neovascularization is a well-recognized variant of neovascular agerelated macular degeneration (AMD) characterized by intraretinal hemorrhage (IRH) and cystic retinal edema.<sup>1–6</sup> Structural optical coherence tomography (OCT) may additionally demonstrate the presence of accompanying pigment epithelial detachment (PED) or intraretinal pigment migration.<sup>3</sup>

Optical coherence tomography angiography (OCTA) is a novel functional extension of OCT that enables non-invasive visualization of separate retinal capillary plexuses as well as choroidal neo-vascularization.<sup>7–12</sup> One limitation of OCTA is that moving red blood cells in the inner retinal vessels project fluctuating shadow artifact onto the deeper layers of the retina creating artificial flow signals. On cross-sectional OCTA, projection artifact appears as "tails" below *in situ* flow most prominent on hyperreflective

vasculature visible in deeper retinal tissue. A recent image processing algorithm termed projection-

structural regions, and en face images contain artificial inner retinal

resolved OCTA (PR-OCTA) mitigates projection artifact by resolving the ambiguity between true flow signal and projection artifacts.<sup>10</sup> PR-OCTA detects voxels with *in situ* flow as those where intensity-normalized decorrelation values are higher than all shallower voxels in the same axial scan line, providing artifact resolution for both *en face* and cross sectional OCTA. In commercial OCT angiography, the flow projection artifact is suppressed with a slab-subtraction (SS) algorithm. A notable limitation of the SS algorithm, in contrast to PR-OCTA, is the failure to remove prominent tail artifacts on cross-sectional OCT angiograms. Adequate resolution of these artifacts is particularly necessary to capture axially directed flow which is characteristic of RAP. We herein present OCTA and PR-OCTA findings of a RAP lesion prior to diagnosis, at clinical diagnosis, and with subsequent therapy.

#### 2. Materials and methods

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Following Institutional Review Board approval, multimodal retinal imaging including structural OCT, color fundus photography,

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and fluorescein angiography was retrospectively reviewed. OCT angiograms were acquired using the commercially available scan protocols with the spectral domain OCT (RTVue-XR Avanti) based on the split-spectrum amplitude decorrelation angiography (SSADA) algorithm.<sup>8</sup> These images were subsequently exported to the Casey Eye Reading Center for application of PR-OCTA algorithm and semi-automated segmentation.<sup>10</sup> The deep capillary plexus (DCP) of the retina was defined as flow between the outer half of inner nuclear layer and the outer boundary of the outer plexiform layer (OPL). Outer retinal flow was localized between the outer boundary of the OPL and Bruch's membrane (BM). Purple segmentation lines were utilized to depict the inner limiting membrane, yellow lines the interface of OPL and outer nuclear layer, and green lines the interface of retinal pigment epithelium (RPE) and BM. Inner retinal flow was depicted as purple, outer retinal flow as vellow, and choroidal flow as red.

DCP and outer retinal en face angiograms consisted of maximal flow projection along axial (Z) dimension. Thick (100  $\mu$ m) cross-sectional OCT angiograms consisted of 10 axial frames. Cross-sectional structural OCT images were represented by the reflectance signals in the middle of the thick cross-sectional OCTA (Fig. 2A,D).

#### 2.1. Case report

A 79-year-old male with neovascular AMD in the right eye presented with vision loss in the left eye. Visual acuity measured

20/40, IRH was detected superior to the fovea by fundus exam and was associated with leakage on fluorescein angiography (Fig. 1A–B). Cross-sectional OCTA showed intra-retinal fluid (IRF) and abnormal flow in the outer retina, however projection artifact is present on retinal pigment epithelium (RPE) and artifact "tails" of *in situ* flow limit depth discrimination (Fig. 1C). With cross-sectional PR-OCTA, projection artifact is removed allowing easier discrimination of RAP lesion depth (Fig. 1D). Three-dimensional volume rendering of OCTA illustrates axially directed flow within the RAP lesion (Supplementary Video).

Supplementary video related to this article can be found at https://doi.org/10.1016/j.ajoc.2017.10.001.

Five months prior, the patient had received OCTA imaging of the left eye as part of routine evaluation. Scans were reviewed focusing on the region superior to the fovea: *en face* PR-OCTA of DCP showed a single vessel that was slightly dilated and brighter compared to surrounding DCP (Fig. 2A). Flow could not be confirmed by conventional OCTA due to tail artifact (Fig. 2B). Cross-sectional PR-OCTA detected flow in the outer retina associated with a very small PED and no IRF was present (Fig. 2D). Thick cross-sectional OCTA revealed outer retinal flow was contiguous with DCP without extension into the sub-retinal pigment epithelial (RPE) space (Fig. 3). The axially oriented RAP lesion appeared small with *en face* OCTA (Fig. 3).

At the time of RAP diagnosis, monthly treatment with aflibercept was initiated. After three injections, treatment interval was extended to every six weeks. Serial PR-OCTA demonstrates RAP

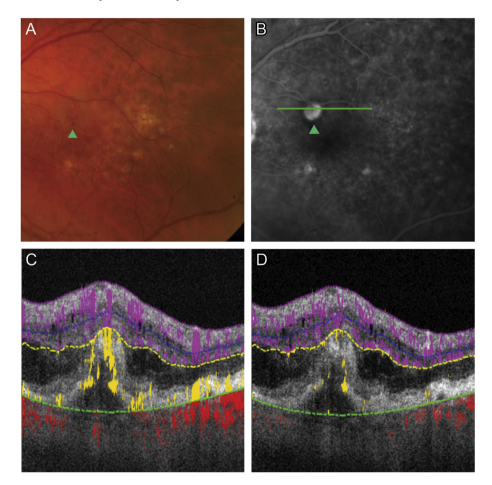


Fig. 1. Retinal angiomatous proliferation (RAP) at the time of diagnosis.

(A) Color photo and (B) fluorescein angiography of RAP lesion (green arrow). (C) Cross-sectional (corresponds to green line in B) optical coherence tomographic angiography (OCTA) revealed abnormal flow in the outer retina (yellow), however projection artifact is present. With cross-sectional PR-OCTA, projection artifact is removed allowing easier discrimination of RAP lesion depth (D). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

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