OPTICAL COHERENCE TOMOGRAPHY ANGIOGRAPHY TO DETECT AND MONITOR NONEXUDATIVE NEOVASCULARIZATION IN AGE-RELATED MACULAR DEGENERATION

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ABSTRACT

OCT angiography has improved the ability to identify silent or nonexudative neovessels in age-related macular degeneration. This promotes a rapid advancement in the diagnosis and in the necessity to expand our management decisions. This context is presented and the biomarkers are discussed.

Keywords: Optical coherence tomography; Optical coherence tomography angiography; age-related macular degeneration


INTRODUCTION

Age-related macular degeneration (AMD) has many new contexts. These are necessary given that prevalence is projected to rise in the next 20 years to as much as 75%, or 175 million patients. The current standard diagnostic tools routinely combine clinical retinal examination with dye-based angiographies and structural ocular coherence tomography (OCT). Using these resources we identified the site of the disease and monitored its activity. Another important aspect to consider is related to neovessel resilience in AMD even with prolonged anti-VEGF therapy, given that the MARINA Study found that these neovessels do not disappear.1,2

Further relevant information came from different clinical spectra. Nowadays the Gass-Freund classification is used and helps to provide a prognosis. Basically, there are 3 subtypes of neovascularization: (type 1) sub-retinal pigment epithelium (RPE) or occult; (type 2) subretinal or classic or well-defined; retinal angiomatosus proliferation or deep retinal neovascular complex (type 3).3,4

Recently the use of OCT angiography (OCTA) has revealed the presence of nonexudative neovessels undetected by any other resource and the necessity to establish the best form of case management. But this situation is not new at all. Almost 45 years ago histological studies showed sub-RPE neovessels in 11% of eyes that had previously been diagnosed as having intermediate AMD (iAMD). Probably the blockage of fluorescein caused by the RPE, the low flow inside these vessels, and the absence of exudation prevent the image from being captured by the resources used.5 With the use of indocyanine green angiography (ICGA) which has better access under the RPE this situation was detected in around 11% of the cases with iAMD and with negative intravenous fluorescein angiography (IVFA).
ICGA was able to show the different evolution (2.6 x more likely to exudate) or impaired visual acuity without exudation. Lesions tended to enlarge over time, causing visual distortions even in the absence of exudation.

The objective of this paper is to review the current information concerning the detection of neovessels in iAMD and the biomarkers that help in the evaluation of their activity.

**CLINICAL DETECTION AND BIOMARKERS**

During many years the risks for AMD cases are related to drusen size. After 10 years almost 15% of patients with medium drusen in both eyes will develop late AMD, and for fellow eye cases almost 60% or more will develop late AMD, depending on the number of risk factors.²

The introduction of OCT has pointed out indirect intraretinal spots, external limiting membrane discontinuity, ellipsoid breaks, and growth of pigment epithelium detachment are related to the presence of these neovessels. Some modifications in the structure of the retina are strong indicators, but the vessels are unseen.⁵,⁹

OCTA has given us new definitions. Nonexudative macular neovascularization in AMD (nneAMD) is defined as type 1 lesion detected with dye-based angiography or OCTA that is not associated with clinical or structural OCT evidence of exudation for at least 6 months. There is a 30% chance of exudation in that period.¹⁰⁻¹³ Another definition, “nonexudative detachment of the neurosensory retina” (NEDNR) characterizes the presence of subretinal fluid (SRF) in iAMD that was not associated with any detectable macular neovascularization using current state-of-the-art imaging, including SS-OCTA.¹⁴ (figure 1 and 2)

**Figure 1. NONEXUDATIVE NEOVASCULARIZATION IN AGE-RELATED MACULAR DEGENERATION - SS-OCT (DRI - OCT1 - TRITON Plus Topcon, Tokyo, Japan).**

Drusen are shown in 3D reconstruction (a) and color photograph (i). The OCT-B mode (b) and OCT-B flow analysis (g) show the presence of fluid. The OCT-A segmentation (c,d,e,f) identify a small neovascular tuft (arrow). The flow vessel density (h) localizes the tiny neovascularization.
Prevalence of nneAMD ranges from 5.5-34.0% (mean 21.1%) depending on the series. Studies have used different methodologies related to sample size, inclusion, exclusion, stage of AMD, equipment, manual or automatic segmentation. This heterogeneity explains this variation in the prevalence rates. But this average and the quality of observation is better than the routine combinations to achieve the same objective.\textsuperscript{11,14-17}

OCTA sensitivity is near to 80%, and probably a little bit lower if only type 1 is considered. This means that an important number of cases with nneAMD are not diagnosed with this technique. Specificity could be as high as 100%.\textsuperscript{18-21} Another relevant limitation is related to natural history. In nneAMD, exact evolution is poorly understood. The relative risk for exudation could be 15.2 times more compared with AMD without CMNV. In a year, 21.1-30% of these cases (versus 3.6% of normal cases) will develop active neovascularization.\textsuperscript{16}

A further consideration is that blood flow persists in all cases with nneAMD even after prolonged therapy. This is strong evidence that the treatment is not always necessary. Moreover, remission is not free from recurrence. Remission lasts longer in the case of eyes with low neovessels density and these cases must be closely monitored. Remission is caused by the RPE surrounding the tufts or regression of immature vessels. Lower vessel density is associated with subretinal fibrosis and low VA.\textsuperscript{18-23} One of the best explanations about this is called the “vascular abnormalization theory”.\textsuperscript{24}

The artifacts produced during OCTA image acquisition (segmentation errors, projections, shadowing) are great challenges to be overcome and can interfere with precise evaluation. It is also important to highlight that most of the studies of this subject excluded patients with AMD associated with high myopia, cataracts, and diabetic retinopathy. These exclusions limit the conclusions that can be drawn.\textsuperscript{13,25,26}
OCTA has made some considerable contributions to understand the pathogenesis. Standing out among them is the demonstration of dark areas and the flow void points also around the neovascular lesions. These halos should be correlated with low flow signs in structural OCTA. The situation reflects zones of bad perfusion (real ones or tiny vessels with sluggish flow) and is assumed to be related to primary AMD pathogenesis. Drusen can also co-localize choriocapillary impairment.\(^{13,27-29}\) Another important contribution derives from the analysis of the retinal and optic disc vessels in these cases. Both superficial and deep retinal and optic disc plexus are altered among patients with not-late AMD. The involvement of optic disc (deep and radial plexus) vessel density was also found to be reduced in the fellow eyes, and this could be a predictive sign. So choroidal thickness, vessel density and/or tortuosity could express vascular perfusion integrity, whereas vessel dispersion and rarefaction could measure damage to the retinal vessels.\(^{29-31}\) Observation of these modifications may improve sensitivity in detecting subclinical alterations occurring in these patients and lead to earlier therapeutic interventions.

Detection of biomarkers is of paramount importance for all patients with nneAMD. Both OCT-B and OCT-A have become the gold standard methods for detecting nnAMD. The combination of three of the situations can identify signs related to a worse prognosis: (1) a well-defined vessel with a lacy-wheel or sea-fan pattern (rather than long filamentous linear vessels); (2) great number and branching of small capillaries (rather than a few, large vessels); (3) detection of anastomoses and loops; (4) presence of peripheral arcade at the vessel terminus (not a “dead tree” appearance); (5) presence of a perilesional hypointense halo at the level of the choriocapillaris; (6) hyperreflective drusenoid lesions with flow evidence.\(^{8,13,17, 27,32-34}\)

In conclusion, nneAMD is easier to detect with OCTA than with other resources, although not in all cases, neovessels uses not disappear completely. The main limitations are related to artifacts, and biomarkers could help to decide the moment to begin treatment.

REFERENCES

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