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The evolving role of OCT in pathologic myopia

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Introduction
The global burden of myopia represents a significant public health concern that is expected to continue to increase in the near future. It is estimated that 50% of the world’s population will be affected by myopia by 2050, with a disproportionately high prevalence in Asia. High myopia, where the spherical equivalent refractive error is equal to or higher than 6.00 diopters, is expected to increase in prevalence from 2.7% to 10% during this period. The severity of myopia is of paramount concern to clinicians as higher levels are associated with pathologic myopia (PM) and increased risk of vision loss. Pathologic myopia, as recently defined by the International Myopia Institute, is an excessive axial elongation associated with myopia that leads to structural changes in the posterior segment of the eye that can lead to loss of best-corrected visual acuity. These structural changes and their complications include posterior staphyloma, myopic choroidal neovascularization, myopic maculopathy, myopic traction maculopathy, dome shaped maculopathy, optic disc changes and glaucoma associated with myopia, and retinal detachments.

The advent of optical coherence tomography (OCT) has facilitated the characterization, diagnosis, and management of several of these complications associated with PM and will be the focus of this article. Imaging the highly myopic eye represents a crucial step in the identification of these complications and poses its own unique challenges. Researchers have demonstrated the advantage of 3D cube scans in the detection of pathology compared to 1- and 5-line rasters. Using vertical scanning patterns aligning where the radius of curvature is larger relative to the horizontal plane of the myopic eye can minimize associated artifacts. Wide scans, facilitated by emerging technologies such as swept-source OCT and ultra wide-field OCT, are useful in cases of PM where the pathology can initiate peripherally.

Myopic Choroidal Neovascularization
It is estimated that 5-11% of patients with PM will develop myopic choroidal neovascularization (MCNV). Effective treatment, primarily with anti-VEGF agents, can alter the natural history of MCNV that would otherwise result in significant vision loss.

Diagnosis: MCNV often appears as a type 2 CNV on OCT with a hyperreflective mound above the retinal pigment epithelium (RPE) (Figure 1). Additional OCT features of MCNV include a loss of the ellipsoid layer and Bruch’s membranes, absence of the external limiting membrane, and retinal thickening. The degree of intra- and subretinal fluid by OCT is less in MCNV compared to other etiologies and thus may be a less sensitive marker for its detection if used in isolation. At the same time, features such as the absence of subretinal fluid, may aid in the differentiation of MCNV from hemorrhage related to lacquer cracks. However, to date, FA remains the gold standard for diagnosis of MCNV.

Figure 1. A) Pigment epithelial detachment (red) with overlying disruption of the ellipsoid and external limiting membrane (yellow arrow), subretinal hyper-reflective material (green arrow), and small amount of subretinal fluid (blue circle). B) Follow up OCT after 1 year of “treat and extend” anti-VEGF. Note the consolidation of the borders of the PED, and absence of subretinal fluid and sub-retinal hyperreflective material.
represents inner-outer macular schisis, stage 2 represents predominantly outer macular schisis, stage 3 represents macular schisis detachment, and stage 4 represents macular detachment. Stage a represents a normal foveal profile, stage b represents lamellar hole changes, and stage c represents macular hole changes. Epiretinal abnormalities are designated with a “+” and can occur at any stage.

**Disease Monitoring:** OCT provides a rapid, non-invasive means of monitoring disease activity and response to treatment. A characteristic “fuzzy area” of the hyperreflective lesion often corresponds with active FA leakage\(^\text{13}\). Subretinal hyperreflective exudation is similarly predictive for MCNV activity\(^\text{14}\). Several studies have sought to identify OCT characteristics capable of prognosticating development of MCNV, response to treatment, and recurrence. Eyes with MCNV have been found to have reduced choroidal thickness compared to fellow eyes\(^\text{15}\). Thinner choroids have also been associated with recurrence of MCNV at the 1-year follow up and with a lower resolution rate after single injections of anti-VEGF therapy\(^\text{16}\).

**Myopic Traction Maculopathy**
Myopic traction maculopathy (MTM) refers to a collection of conditions including vitreomacular traction, epiretinal membrane, lamellar hole, macular hole, myopic foveoschisis/macular schisis that are unified by their underlying etiology of tractional forces acting on the retina in highly myopic eyes\(^\text{2}\). MTM may affect up to 30% of patients with PM\(^\text{17}\). OCT findings in MTM have recently been organized into a staging system\(^\text{18}\) called the MTM Staging System (MSS) (Figure 2). The MSS organizes the evolution of changes from forces perpendicular to the retina (stage 1-4) with forces tangential to the retina (a-c), to arrive at a number and letter staging (e.g. 2B). Stage 1 represents inner-outer macular schisis, stage 2 represents predominantly outer macular schisis, stage 3 represents macular schisis detachment, and stage 4 represents macular detachment. Stage a represents a normal foveal profile, stage b represents lamellar hole changes, and stage c represents macular hole changes. Epiretinal abnormalities are designated with a “+” and can occur at any stage.

**Myopic Macular Schisis:** Myopic Macular Schisis (MMS), or foveoschisis, is evident on OCT as retinoschisis in multiple retinal layers, bridged vertically by presumed Müller cells. Separation of the internal limiting membrane (ILM) from the remaining retinal layers can occur. In early stages (stage 1-2 MSS), the ellipsoid zone is usually well-preserved and visual function is minimally affected. A large natural history study of MTM\(^\text{19}\) found that over 36.2 ± 6.2 months of follow up, 11.6% (n= 24/207) of eyes experienced progression of MTM (Figure 3), with higher rates of progression (42.9%) in eyes with more extensive MMS.

**Myopic Macular Hole:** Macular holes are reported to occur in approximately 8.4% of eyes with PM\(^\text{20}\). Data has shown that there is a differentiation between two types of macular holes in myopic eyes. The first is a “flat type”, characterized by cystic changes at the edge of the hole and similar success rates of surgical closure similar to macular holes in non-myopic eyes\(^\text{21}\). The second is a “schisis type”
characterized by accompanying retinoschisis changes which are evident at the borders of the hole. These latter holes have a lower success rate of closure and are at high risk of progression to macular hole retinal detachments.

Treatment: Intervention at the MMS stage is generally guided by progressively worsening visual symptoms, metamorphopsia, and reduction in visual acuity. Treatment consists largely of surgical intervention across the spectrum of MTM and comprises vitrectomy and/or macular buckling. In a prospective, non-randomized study of 62 consecutive eyes, rates of visual improvement with surgery were shown to be greater when foveal detachment or disruption was evident. Several variations of vitrectomy have been proposed, vitrectomy alone, with membrane peeling, with ILM peeling, ILM flaps, subretinal expansion, retinal incisions, and grafting procedures of amniotic membrane or retinal tissue. There has been renewed interest in macular buckling for the management of MTM with the hypothesis that anatomical and functional outcomes may be superior to vitrectomy alone (Figure 4). Several modifications to the standard macular buckling technique have been proposed to facilitate the surgery including modified buckle shapes and internal chandelier placement. Intraoperative OCT has been used to augment surgery in MTM eyes through the detection of residual epiretinal membrane (ERM), ILM, and cortical vitreous and has revealed undetected holes post peeling. Clinicians have proposed a treatment algorithm based on their MSS using permutations of vitrectomy and macular buckling depending on the stage of disease, with vitrectomy better suited to address tangential traction and macular buckling better suited to address perpendicular traction. Such studies underscore the utility of OCT in cases of MTM in potentially identifying optimal treatment strategies corresponding to pathophysiologic principles.

Myopic Maculopathy
Myopic maculopathy (MM), or myopic macular degeneration, are terms often used interchangeably to describe several of the degenerative features in PM that contribute loss of best corrected visual acuity in myopia. The most widely used classification system is the meta-analysis of pathologic myopia (META-PM) which divides PM into five distinct categories as follows: category ‘0’: no myopic retinal lesions, category ‘1’: tessellated fundus, category ‘2’: diffuse chorioretinal atrophy, category ‘3’: patchy chorioretinal atrophy, and category ‘4’: macular atrophy. Additional features of “plus lesions” can be assigned for lacquer cracks, choroidal neovascularization, and Fuchs’ spots. MM tends to progress. Over a mean follow up of 18 years in a retrospective observational case series of 810 eyes, the authors reported progression of MM in 58.6% of all eyes, and 74.3% in eyes with pre-existing PM. Notably, META-PM is based on fundus photographs

Figure 3. Progression of MTM from baseline (A) to predominantly outer retinal schisis/stage 2a at 3 years (B) followed by progressive outer retinal schisis and increasing lamellar changes/stage 2b at 4 years (C) and 5 years (D). 9 months after (D) the appearance of a macular schisis detachment/stage 3c was noted (E) with extension of the retinal detachment to outside the macula after another month (F). The patient was treatment with vitrectomy, membrane and ILM peel, and silicone oil with continued anatomical success and 20/200 vision at 2 years follow up but significant retinal atrophy (G)
and does not incorporate OCT findings. Researchers have recently attempted to supplement the META-PM classification with OCT features and demonstrated that choroidal thinning is associated with the progression of myopic maculopathy. They established cut-off values of choroidal thickness of \(<56.5\) μm located 3000 μm nasally from the fovea to define peripapillary choroidal thinning, and \(<62\) μm subfoveally to define macular choroidal thinning, two terms proposed as the OCT equivalent to the subtypes of category 2 diffuse chorioretinal atrophy in META-PM. Unfortunately, to date, no established treatment for MM exists.

**Conclusion**

OCT has emerged as a foundational technology in the management of PM. The relative hypopigmentation of myopic fundi and microstructural changes associated with pathology means that biomicroscopy alone is insufficient in the assessment of PM eyes. Advances in OCT imaging continue to enhance our understanding of PM and it is imperative for the eyecare specialist to leverage these advantages in the care of our patients.

**References**


Figure 4. Pre-operative OCT of stage 4b MTM in the right (A) and left eye (B) of a -12 myope with symptomatic visual decline to 20/100 and 20/70 respectively. Post-operative OCT after successful macular buckling surgery in the right (C) and left eye (D) with reduction in retinoschisis and resolution of macular detachment. Visual acuity improved to 20/30 OU with a concurrent reduction of 5 diopters of myopia.


