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REVIEW

Choroidal Cavitary Disorders

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Abstract: The structure and functions of the choroid have been long acknowledged but the pathophysiology behind various anomalies has been difficult to understand until the advent of optical coherence tomography (OCT). With OCT imaging, choroidal cavitations appear as optically empty spaces between the outer retinal and choroidal layers with attenuation or loss of outer retinal layers. Choroidal cavitations are found in the posterior pole and seen in conditions such as pathologic myopia, north carolina macular dystrophy (NCMD), focal choroidal excavation (FCE), and torpedo maculopathy (TM). To date, these disorders have not been linked. A commonality they all share is malformation of the RPE-photoreceptorchoroid complex. The following report describes the differences and similarities of choroidal cavitation amongst the different retinal disorders and emphasizes the importance of multimodal imaging in the detection and management of potential complications.

Keywords: peripapillary intrachoroidal cavitation, macular intrachoroidal cavitation, focal choroidal excavation, torpedo maculopathy, choroidal neovascularization

Key Points

Intrachoroidal cavitation secondary to pathologic myopia is associated with axial elongation and structural weakening of the involved sclera, RPE, and choroid, notably in the macula and peripapillary area near the myopic conus.

Macular intrachoroidal cavitation has also been reported in cases of north carolina macular dystrophy - in contrast, underlying mechanisms for ICC formation are thought to arise in utero.

Focal choroidal excavation, often an incidental finding on OCT, has gained attention due to its elusive pathophysiology and associated pachychoroid disorders.

Torpedo maculopathy, a congenital lesion that exhibits choroidal cavitation and thereby compromised RPE, can also exhibit vulnerability to the vascular effects of aging.

The relevance of these clinical findings becomes apparent when considering underlying pathophysiology, potential complications that may be vision threatening, and early detection/treatment of said complications.

Introduction

In primates, the choroid receives 85–90% of blood flow to the eye and delivers many times more blood to the eye than the retinal vasculature.^{1,2} The choroid functions to nourish the retina, influences scleral growth, and maintain the retina's optical position by virtue of thickness changes.³ The uveal tract is the vascular tunic of the eye, positioned between the sclera and the neuroepithelium. This tract is made of the iris, ciliary body and the choroid. The choroid extends anteriorly from the ora serrata

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posteriorly to the optic nerve. Spanning from the posterior of the retina to the sclera, the choroid is made of five layers. The basement membrane of the retinal pigment epithelium is Bruch's membrane. Posterior to this layer are the remaining choroidal layers, in order from nearest to the retina to the furthest: the choriocapillaris, Haller's layer, Sattler's layer, and the suprachoroid (Figure 1).³

The choroid receives its blood supply from the 2–3 ciliary branches of the ophthalmic artery which then further supply the 10–20 short posterior ciliary arteries.^{4,5} It is drained by the vortex veins. Two thirds of the oxygen consumed by the retina in the light-adapted state is delivered by the choroid.² A high rate of choroidal blood flow is not only necessary for maintaining high oxygen tension but also maintaining homeostatic temperature in a highly metabolic tissue.⁶

The choroidal vasculature are innervated by sympathetic fibers emanating from the superior cervical ganglion.⁷ Parasympathetic innervation is achieved by the facial nerve and pterygopalatine ganglion.⁸ The choroid is typically thickest subfoveally with an average thickness of 272 microns in a healthy middle-aged adult and decreases gradually with age, reaching around 80um by age 90.⁹ The choroid plays a role in retinal homeostasis and secretes signal molecules and growth factors including vascular endothelial growth factor (VEGF), basic fibroblast growth factor and hepatocyte growth factor.^{3,10}

Innovations in ophthalmic imaging including optical coherence tomography have vastly improved the visualization of pathology of the choroid. These innovations have allowed the further elucidation of known entities and led to the discovery of new choroidal conditions. In the text that follows, we review cavitary disorders of the choroid and the pathophysiologic mechanisms that underpin them. The connections made between the cavitary disorders in this review serve as one of the many of the initial building blocks in understanding common complications and treatment modalities.

Intrachoroidal Cavitation (ICC) Peripapillary Intrachoroidal Cavitation

In 2003, Freund et al first published the report of a localized peripapillary retinal pigment epithelial detachment (PED) in patients with high myopia.¹¹ Subsequent studies and more resolute time-domain OCT imaging by Toranzo et al¹² revealed such lesions to be comprised of choroidal tissue cavitation and a flat retinal pigment epithelium (RPE) profile, not consistent with a dome-shaped elevations seen in a PED. The entity was accordingly renamed peripapillary intrachoroidal cavitation (pICC).¹¹

Further advancements in spectral-domain OCT (SD-OCT) imaging in 2008 demonstrated there to be an associated choroidal thickening, with or without cavitation, in these peripapillary lesions.¹³ Hence the name "peripapillary choroidal thickening and cavitation" was proposed by Freund et al in 2011 to be a more accurate description of the lesion.¹⁴ In 2012, Spaide et al, using SS-OCT, significantly improved visualization of the scleral-choroidal junction which demonstrated a characteristic bowing of the sclera beneath the choroidal cavitation. This implied that the cavitation is an effect of posterior excursion of the sclera rather than anterior displacement of the retina and RPE.¹⁵

With fundus examination, the characteristic finding of pICC is a well-circumscribed, peripapillary lobular,



Figure I This is a SS-OCT of a normal left eye of a 45-year-old female. The blue arrow on the color fundus photograph indicates the position of the OCT B-scan. The yellow arrow indicates a Haller vessel, the blue arrow a Sattler vessel, and the red arrow the choriocapillaris, a 10–30 micron thick layer immediately subjacent to the Bruch's membrane. The choroidal thickness varies greatly from person to person, by location relative to the optic nerve and fovea, over time with aging, and in diseases such a central serous retinopathy (thicker than normal) and high myopia (thinner than normal). In the case shown here, the choroid is thicker between the fovea and optic nerve (white double headed arrow) than at the fovea (purple double headed arrow), and thinner nearer the optic nerve (double headed green arrow).

yellow-orange, elevation or thickening.^{11,15-17} These lesions are typically located inferior at the border of myopic conus but can occur less frequently temporal.^{14,18} In some cases pICC may be indistinguishable or resemble other retinal findings during fundus exam; therefore, OCT raster scans of the peripapillary region in highly myopic patients are advised so as to detect the presence of pICC in suspected cases.^{16,17}

The appearance is similar to a PED and they were initially described as such.^{7,19} However, the lesion is not typically elevated as in a PED and does not display the fluorescein angiographic (FA) findings of early localized hyperfluorescence with progressive dye pooling.¹² Fluorescein angiography of ICC lesions demonstrates early hypofluorescence with progressive staining of the peripapillary lesion without dye pooling. Indocyanine green angiography (ICGA) shows the area of peripapillary cavitation is hypofluorescent throughout the entire sequence.^{12,21} Literature on OCT-angiography (OCT-A) use in pICC is limited. A case report by Mazzaferro et al on a 67-year-old female with pathologic myopia and pICC demonstrated a hyporeflective area corresponding to the cavitation on OCTA which may indicate a sluggish choriocapillaris vasculature or a complete absence of choroid.²¹

Peripapillary ICC is predominantly found in pathologic or high myopia but may occur in lower degrees of myopia and rarely in hyperopia.^{11,16,17,19} Pathologic myopia is defined as a refractive error \geq -6.00 diopters and an axial length >26 mm.²² Complications arise secondary to excessive axial elongation of the globe and thinning of the sclera, choroid and RPE.^{23–25} These changes, along with advancing age, cause a weakening of the peripapillary tissue in the area of the myopic conus where choroid loss and cavitation occur.^{12,17,26}

Peripapillary ICC is rare in younger patients. Shimada et al reported only 1 of 31 cases of pICC were younger than 30 years of age.²⁶ Yeh et al found 79 of the 83 eyes with pICC were over thirty years of age.¹⁷ In addition, those patients with low refractive errors and pICC were significantly older than those with high myopia, underscoring the combined effects of time and structural weakness.¹⁷ Although pICC has been reported in patients with mild refractive error, these patients commonly exhibit findings typical of high myopia such as peripapillary atrophy, tilted disc and choroidal thinning.^{27,28}

The mechanism of the actual formation of the cavity following structural weakening has been debated. Forte et al have suggested the posterior scleral bowing and deep excavation of the myopic conus create "the consequent impossibility for the retina-RPE to follow this steep fall," thus detaching from the choroid and returning to a more natural anatomic level and in the process creates the area of cavitation.²⁹ Toranzo hypothesized that staphyloma progression breaks the collagenous limiting tissue of Elschnig (and/or Jacoby¹⁵), which serves as the connection between the choroid and the optic nerve.³⁰ The resultant retraction of the choroid from the nerve margin accounts for the gradual enlargement of the yellow-orange area of cavitation over time.¹²

Some studies have suggested that a cleft forms that connects the pICC and the vitreous cavity^{15,16,31,32} During peripapillary staphyloma progression and stretching of the retina, a cleft opens but the adherence of the retina and RPE at the conal margin prevents fluid from entering the subretinal space; instead, the fluid forms a cavity within the choroid.¹⁶ This is supported by observations of Spaide et al, who used SS-OCT to evaluate pICC.¹⁵ They conjectured that the sclera, thinned and weakened in the area of conus, bows over time in response to intraocular pressure creating space for cavitation to occur.¹⁵ Age is an associated risk factor for pICC and it has been suggested that poor fluid reabsorption associated with an aged retina, combined with gravity, produces the characteristic cavity at the inferior margin of the conus in pICC.¹⁷

Peripapillary ICC lesions can be stable and relatively benign without visual compromise.^{11,16} Complications that can arise include glaucomatous-like defects, macular detachment and retinoschisis^{17,19,20,26-29,32} Shimada et al found that 71% (22/31 eyes) of patients with pICC had glaucomatous visual field defects on Goldmann visual field perimetry.¹⁹ These defects, such as nasal step and arcuate defects, have been described and attributed in part to the RNFL damage occurring at the edge of conus excavation.²⁹

Retinoschisis and macular detachment occur in patients with pICC and is thought to result when the choroidal cavity extends and connects to the vitreous. Cavitary fluid then gains access to the subretinal or intraretinal space.^{28,31,32} Vitrectomy has shown benefit when used in the treatment of myopic macular retinoschisis, with and without detachment.³³ Employing pars plana vitrectomy in pICC has been shown to eliminate the choroidal-vitreal cavity communication and resolve the macular detachment and schisis.^{28,32}

Macular Intrachoroidal Cavitation

Choroidal cavitation that occurs in the macula is typically associated with posterior staphyloma and pathologic myopia. Pathologic myopia is associated with many structural causes of macular visual impairment, most notably: myopic choroidal neovascularization (CNV), myopic macular retinoschisis, and myopic chorioretinal atrophy. The myopic chorioretinal atrophies can be subdivided into diffuse chorioretinal atrophy (DCA) and patchy chorioretinal atrophy (PCA).^{34–37}

Diffuse chorioretinal atrophy is the milder form with a yellowish appearance and ill-defined borders. A thin choroid is interposed between the sclera and retina. DCA is distinguished from the more severe PCA, which shows sharply defined borders and white atrophy. Involved zones in PCA lack any choroid in front of the sclera accounting for the scleral coloration. Vision is severely impacted in PCA due to the absence of underlying choroid and, consequently, the degeneration of overlying RPE and photoreceptors.³⁶ The progression and expansion of these lesions is the leading cause of central vision loss in pathologic myopia.³⁷ Macular ICC has been shown to occur in 55.4% of patients with PCA. It occurs as an excavation zone of relatively thicker choroid immediately surrounding the PCA.³⁴

The pathophysiology appears to be similar to pICC.³⁶ Ohno-Matsui et al, in 2012, examined 56 eyes of 44 patients with PCA in pathologic myopia. Using SS-OCT, 31 of 56 eyes (55.4%) with PCA showed the characteristic posterior bowing of the sclera similar to pICC. This was not apparent in any of the 113 highly myopic eyes without PCA.³⁴ The cavitary lesions tend to occur on the slope of the staphyloma. These focal areas, devoid of choriocapillaris and RPE (along with the loss of overlying photoreceptors), are thought to weaken the structural integrity of the eye wall.³⁶ Intraocular pressure applied constantly and uniformly over time causes further weakening and thinning resulting cavitation. In addition, a direct communication between the mICC lesion and vitreal cavity was noted in 3 of 31 (9.7%) eyes with mICC.³⁴

The characteristic posterior scleral bowing distinguishes PCA from focal choroidal cavitation in which scleral bowing is absent.³⁶ We have also observed cases in which outward scleral bowing occurs in the absence of choroid (ie, within the patch of PCA) causing a cavitation separating the overlying retina from the underlying sclera (Figure 2). Not all outward bowing of the sclera in pathologic myopia is associated with mICC. For example, Figure 3 exhibits a case in which outward bowing of the sclera concomitant with scleral thinning, but instead of mICC, there is mild outer nuclear layer retinoschisis and localized choroidal thickening without mICC. North Carolina macular dystrophy (NCMD) is an autosomal dominant macular dystrophy originally described in a multigenerational Irish family located in the mountains of North Carolina.^{35,38,39,48} Schoenberger and Agarwal published 2 cases of NCMD with mICC.³⁵ Both patients had NCMD associated with posterior staphyloma and visual acuity in the 20/40–20/60 range. Imaging with EDI-OCT shows the characteristic bowing of the sclera that is associated with intrachoroidal cavitation. The cavitations appear nonreflective and, similar to mICC, tend to be along the slope of the focal posterior staphyloma in NCMD. The authors also describe a grayish-yellow color of NCMD-related mICC versus the yellow-orange color seen in myopic-related lesions.³⁵

Mechanisms for ICC formation in NCMD are thought to arise in utero. NCMD lesions have been seen in patients as young as 2 months of age.^{40,47} In contrast, mICC development in pathologic myopia involves a progression in axial length and a weakening of the sclera over time.

Focal Choroidal Excavation

Focal choroidal excavation (FCE) is an area of focal choroidal thinning and concavity with near-normal overlying retina, in the absence of scleral ectasia, posterior staphyloma, and choroidal or PED.42-45 Wakabavashi et al described 2 variations of FCE, which were later termed the "conforming" type and the "nonconforming" type by Margolis et al.⁴¹ The determining factor was whether the outer retinal layers in FCE conformed to the choroidal depression without disruption. If the layers followed the depression without separation it was designated the conforming type (Figure 4). The nonconforming type demonstrated a localized neurosensory detachment or separation between the photoreceptor tips and underlying RPE with subretinal fluid (SRF) presumably occupying the space (Figure 5).⁴³ While generally stable, FCE can change from conforming to nonconforming and vice versa over time.^{42,46} There are published reports of the transition of FCE in cases with⁴⁷ and without⁴⁸ central serous chorioretinopathy (CSC). Focal choroidal excavation lesions associated with serous retinal detachment in CSC or CNV have been reported to convert to conforming following resolution of SRF.^{42,49-51} Additionally, a nonconforming to conforming conversion occurred in FCE lesions with CNV treated with anti-VEGF therapy.⁵²

Despite typically occurring in the fovea or perifovea, FCE is associated with good visual acuity and a stable clinical course. The finding may be incidental on routine



Figure 2 Red free photograph (top) and spectral domain OCT image (bottom) from the left eye of a 68-year-old man with pathologic myopia and a previously undescribed variant of macular intrachoroidal cavitation. The lesion occurs in an area of RPE and choroidal atrophy in an area of excavation of the thickened subjacent sclera producing a pseudofovea of a sort different from that described in macular intrachoroidal cavitation.



Figure 3 Near infrared reflectance image (left panel) and B-scan OCT image (right panel) of a 68-year-old woman with pathologic myopia. The scleral thickness varies widely (compare the red double-head arrows in nasal and temporal macula, respectively). Where the sclera thins and bows outward, the choroid thickness (space between the two yellow arrows) compared to the choroidal thickness overlying thick sclera (space between two green arrows). However, instead of a choroidal cavitation, the associated response seen is outer nuclear layer retinoschisis (purple arrow). The blue arrows indicate a large intrascleral vein.

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Figure 4 SD-OCT of conforming FCE in the macula shows the outer retinal layers in apposition to the area of focal choroidal concavity.



Figure 5 SD-OCT of non-conforming FCE in the macula. In contrast to the conforming type FCE in Figure 4, non-conforming shows a separation of photo-receptor tips from RPE.

examination.^{46,53,54} Subjective disturbances such as metamorphopsia have been reported even in eyes with normal vision.⁴³ On examination of the fundus, small FCE lesions may not be apparent. Larger lesions show non-specific pigmentary disruption and with choroidal vasculature intact the lesion may appear reddish, yellowish, or orange.⁵⁵

Focal choroidal excavation is easily recognized on OCT imaging with a characteristic focal intrachoroidal concavity or downward deflection of the Bruch's membrane-RPE-choriocapillaris complex line.⁴⁴ The inner retinal layers appear relatively intact despite deformation.^{42,43,54} There

is a thickening of the outer nuclear layer which is thought to be from the traction between the excavation dip and vascular system of the inner retina; creating strain on the RPEphotoreceptor interface.⁵⁴

Subretinal fluid is thought to occupy the subretinal space in nonconforming FCE but vitelliform material occasionally accumulates.⁵⁶ The presence of an unusually high reflective choroidal material beneath the FCE is thought to represent scar formation and this scarring subsequently contracts leading to traction on RPE and FCE formation.^{49,51} Reduced vision and symptoms of metamorphopsia are more commonly associated with nonconforming FCE.^{49,57,58} The SRF in nonconforming FCE has been hypothesized to arise from traction on the photoreceptors, but OCT evidence of traction is found in a minority of cases.⁵³

The choroid is thin in the area of FCE, but the scleral contour and thickness are intact and normal.⁴⁹ A staphylomatous excavation of the sclerochoroidal junction needs to be excluded via OCT to diagnose FCE and avoid confusion with other choroidal cavitary disorders.^{45,61} Immediately beneath FCE, the choroid may be hyperreflective on OCT, with a paucity of vessels compared to the adjacent choroid.⁵⁹

Although initially described as a finding in otherwise healthy eyes, FCE has been shown to accompany a host of choroidal pathologies including CSC, 51,60,61 CNV, 52,62 polypoidal choroidal vasculopathy,63-66 choroidal osteoma,67 multiple evanescent white dot syndrome (MEWDS),68-70 multifocal choroiditis,⁷¹ Epstein Barr virus infection,⁶⁹ and Vogt-Koyanagi-Harada disease.⁷² Retinal disorders have also been associated with FCE such as age-related macular degeneration (AMD),^{63,73} Stargardt Disease,⁷⁴ Best pattern dystrophy,⁷⁴ cone dystrophy,⁷⁷ Disease,^{75,76} foveoschisis,⁷⁸ retinal hamartoma,^{79,80} vitreomacular traction and retinal hole,^{81,82} and torpedo maculopathy.⁴² Speculation remains on the exact relationship of FCE to the above findings but associated weakening or mechanical changes in the RPE-Bruch's membrane complex and consequences of disrupted choroidal vasculature are thought to be predisposing factors (Figure 6).⁵³

A large retrospective study examined 4436 SD-OCT patient scans in an 18-month period and found 16 patients with FCE ranging from 31-78 years.⁵³ There was no gender predilection and the refraction of both the involved eye and fellow eye was approximately -3 diopters (D). Macular pathology was present in four cases (25.0%) of eyes with FCE: one patient with macular pucker, one with parafoveal scarring, one associated with PCV and one with macular hemorrhage from lacquer crack formation in high myopia. The remaining 12 cases (75%) were not associated with concurrent macular pathology in the FCE eye. Among those 12 eyes with FCE but without macular pathology, fundus examination showed half of them had visible RPE changes while the rest were not discernible clinically and only seen by OCT imaging.⁵³

The size of the lesion is useful clinically as it has been found to have bearing on the risk of concurrent macular pathology. A greater length, area and volume of an FCE lesion impart greater risk of associated macular pathology.^{53,54} Chen et al found that in multimodal imaging of FCE, the most sensitive modality was near-infrared autofluorescence (NIA), with the ability to better image subretinal and choroidal melanin loss in the form of hypoautofluorescence. A greater area of FCE on autofluorescent imaging was shown to be associated with maculopathy such as CNV and CSC. Greater volume (NIA area) x (Depth measured by SD-OCT) of FCE was associated with more subjective distortion.⁵⁴

Fluorescein angiographic findings include varying degrees of hyper and hypo fluorescence associated with RPE window defects.⁵⁵ Late phase hypofluorescence and abnormal staining on ICGA has been described, as well as, choroidal vascular hyperpermeability.^{51,55} These findings suggest a correlation of FCE and pachychoroid spectrum disease.⁸³ Pachychoroid disease and its encompassing maladies of PCV, CSC and CNV have been related to FCE.^{59,63} The prevalence of FCE in eyes with CSC has been documented to be between 4.8% and 7.8%.^{47,51,60} However, the prevalence of CSC in eyes with FCE has been reported up to 24.4%, much higher than that of normal eyes.⁴⁷

The pathologic mechanism in FCE remains unclear. Inflammatory,^{68,71} infectious,⁸⁴ vascular,^{51,60} developmental⁷⁹ and congenital^{42,57,85} processes have all been suggested. Age varies significantly in patients that have FCE lesions with mean ages falling in the 45–55 range.^{42,46,49} Park et al found a relatively low prevalence



Figure 6 SD-OCT image of a patient with nonexudative AMD in which a PED developed with focal atrophy of the RPE at the crest of the PED (red arrow), but in which the Bruch's membrane remained intact (yellow arrow). The subjacent choroid (blue oval) suffered no choroidal excavation. Destruction of the Bruch's membrane is hypothesized to be a necessary precondition for FCE development in acquired cases.

in FCE among children and adolescents with only 3 of 1697 eyes with FCE.⁸⁶ Many studies have shown an association of myopia with FCE.^{50,51,63} It has been suggested that eyes with longer axial lengths have thinning of the choroid and weakening of RPE-Bruch's complex resulting in FCE formation.⁶³ Congenital or embryological developmental disorders have been proposed in light of the relatively intact outer retinal layers and choriocapillaris which are absent in chorioretinal scars.^{42,85} Kumano et al speculated an outward traction created by choroidal vascular abnormalities from embryonic developmental failure of the choroid.⁵⁷ Other clinicians have theorized that choroidal abnormalities may lead to ischemic changes causing RPE atrophy.⁶¹

Viral-related inflammation such as that seen in MEWDS has been associated with the development of FCE.^{68,69} Hashimoto et al suggested that the pathophysiology is related to inflammation at the level of the outer retina causing disruption of the RPE-Bruch membrane complex. Following this disruption, there is an adherence of the outer retinal and choroidal tissues to each other through the ruptured RPE-Bruch membrane complex. Intraocular pressure then causes the adherent complex to protrude outward through the Bruch membrane gap creating a choroidal excavation.⁶⁸ Similarly, the associated RPE and inner choroidal inflammation in multifocal choroiditis and panuveitis/punctate inner choroidopathy are associated with FCE formation.⁷¹

Vision is typically unaffected in FCE unless it is located near the fovea. Management involves excluding associated CNV, PCV and CSC. Choroidal neovascularization in FCE has shown to be amenable to anti-VEGF treatment.^{52,62,73,74} Both PCV and CSC with associated FCE have shown to respond to anti-VEGF injections and photodynamic therapy (PDT).^{58,64} Restoration of vision varies depending on residual retinal damage.⁵⁸

Torpedo Maculopathy

Torpedo maculopathy (TM) is a congenital, solitary, spindle-shaped chorioretinal lesion in the temporal macula.⁸⁷ It was originally classified as congenital hypertrophy of the RPE (CHRPE) by Gass in 1989⁸⁸ and revised to a "solitary hypopigmented nevus of the RPE" by Roseman and Gass in 1992.⁸⁹ It was later given the name Torpedo Maculopathy by Daily in 1993 due to its torpedo-like appearance.⁹⁰

Fundus exam shows a hypopigmented, horizontally oriented, oval-shaped lesion in the temporal macula

(Figure 7).^{91,92} Although typically unilateral, bilateral cases have been reported.^{93,94} TM is non-progressive and spares the fovea.⁹¹ There have been reports of growth and change in shape.^{95,96} While usually isolated, occasionally small, round satellite lesions temporal to the main lesion have been noted.^{88,92,97,98} Choroidal cavitation in TM differs from FCE in which lesions are not usually oval in shape, typically involve the fovea, and are associated with less outer retinal thinning and degeneration.⁴²

Since TM lesions are rarely located in the fovea, the patient is asymptomatic and the condition is found incidentally on exam.^{87,99-101} Visual field defects are present and correlate with the size, shape and location of the lesion.^{87,92,102} Shirley et al estimated the prevalence of TM as 2 per 100,000 population, but suggested that this is an underestimate due to lack of symptoms.⁹² There are no known associated systemic conditions with TM.^{87,92}

The TM lesion may have varying degrees of hyperpigmentation most commonly located at the temporal edge of the lesion.^{94,101} Geographic regions of hyperpigmentation within the central hypopigmented area are also possible.¹⁰³ The orientation of the lesion is horizontal with a comparatively sharper pointed "head" directed toward the center of the macula.^{87,90} Two different variations have been described for the tail: rounded or frayed.¹⁰¹ The tails with the "rounded" configuration tend to be smooth while the "frayed" tail may be associated with hyper or hypopigmentation. Typical TM lesions measure around 2 disc diameters horizontally and 1 disc diameter vertically.¹⁰¹

Spectral domain OCT demonstrates a thin RPE layer with increased signal transmission to the choroid, occasional loss of the ellipsoid zone line, and no overlying neuroretinal changes.^{80,85,104,105} Wong et al classified two types of TM lesions based on the presence of outer retinal and/or choroidal cavitation. Type 1 was defined by having attenuation of the outer retinal structures without retinal or inner choroidal cavitation; Type 2 featured attenuation of the outer retinal structures with subretinal cavitation.⁹⁷ Torpedo lesions that contain choroidal excavation were recently classified as type 3 TM by Tripathy et al and studied by Venkatesh et al.^{105–107}

Cavitation appears as an optically empty space on OCT as a result of thinning or loss of the outer retinal layers or choroidal layers. The distinction between cavitation and a neurosensory detachment that occurs in some cases can be made on the basis of preservation of retinal layer thickness in a detachment and attenuation or loss in cavitation.⁹⁷ It is speculated that the outer retinal



Figure 7 Color fundus photograph (A) near infrared reflectance image (B), and SD-OCT image (C) of an 8-year-old girl with torpedo maculopathy of the right eye. Her visual acuity was 20/20 bilaterally. The fovea is spared. The point of the torpedo lesion is closest to the fovea. The lesion would be classified as type 2 because of the cavitation (green arrow) of the outer retina. The outer nuclear layer over the cavitation (pink arrow) is attenuated. The choroid is attenuated, but the scleral-choroidal interface is not affected (yellow arrows). The retinal pigment epithelium at the base of the outer retinal cavity is hypopigmented in the color fundus photograph and hyporeflective in the OCT image.

excavation in the type 2 variant is created by loss of photoreceptors and/or RPE.^{87,103,105} However, thinning and disruption of the retinal layers is variable and may be due to the chronicity of the lesion or phenotypic differences.¹⁰⁴ The cavitation can be distinguished from the SRF found in other causes such as CSC by the absence of ragged photoreceptors.⁹⁶ Similarly, it is unclear if the choroidal excavation seen in type 3 TM is acquired or part of the underlying mechanism pathophysiology.¹⁰⁷

While some authors have reported these lesions to be nonprogressive,⁸⁷ others have shown mild progression over time.⁹⁷ Wong et al hypothesized that the Type 1 subtype progressed to type 2 over time but reports of type 2 TM in patients as young as 5 years old cast doubt on that hypothesis.⁹²

Fundus autofluorescence can vary in appearance but generally shows the large central area of the TM lesion to be hypoautofluorescent with surrounding edges demonstrating hyperautofluorescence.^{87,97,108,109} While some type 1 lesions demonstrate hypoautofluorescence, suggesting hypofunctioning RPE, all type 2 lesions are hypoautofluorescent. Wong et al have suggested that the hypoautofluorescence points to reduced interaction between RPE and outer segments that lead to a progressive loss of photoreceptor cells and outer retinal cavitation.⁹⁷ Adaptive optics imaging has been used to evaluate the photoreceptor density in TM. Venkatesh et al studied a type 2 lesion that had decreased cone density. However, because the existing photoreceptors and external limiting membrane were intact, visual acuity was not compromised.¹¹⁰

Scant information exists on perimetric function over TM. Available literature suggests a present, yet inconsistent, correlation of perimetry with lesion type.⁹⁷ Five of eight patients with a type 1 lesion were tested on a Humphrey field analyzer or microperimeter with only 1 patient demonstrating a - scotoma.^{100–112} All patients with type 2 lesions had documented scotomas.^{96,101,113} The variability between visual field defect and lesion may be due to the size of the TM lesions falling beneath the detection sensitivity of Humphrey field testing.⁹⁷ Although photoreceptor degeneration does not correlate with resulting acuity, it does seem to correlate with visual field defect.¹¹⁰

Wong et al proposed a hypothesis for TM formation in relation to the two types.⁹⁷ Type 1 forms during a lack of melanin production within the torpedo lesion leading to an altered interaction between the RPE and photoreceptor outer segments. This disruption leads to attenuation of the ellipsoid zone and interdigitation zone. Because the RPE continues to metabolize the outer segments, retinal sensitivity is preserved as demonstrated on perimetry testing. However, photoreceptor loss over time leads to loss of the outer nuclear layer thickness and cavitation. They noted these cavitations can appear similar to outer retinal cavitations in macular telangiectasia type2¹¹¹ and cone dystrophy or achromatopsia.⁹⁷ Choroidal cavitation occurring in TM, Wong surmised, was a result of reduced VEGF production from compromised RPE leading to a thinning and loss of inner choroidal vessels identified in type 2 lesions.⁹⁷

Shirley et al hypothesize that satellite lesions, which appear to be an extension of the main lesion, support the theory of a congenital defect of the RPE during fetal development. The phenotypic variation seen in TM, uniformity of location, orientation and general stability, adds weight to a congenital etiology.^{92,114,115} Shields et al also endorse an abnormality in fetal development (particularly the RPE) that occurs at a precise time.¹⁰¹ The fetal histology studies of Streeten¹¹⁶ showed that during early development there is a cone-shaped bulge temporal to the fovea in the vicinity of TM lesions. This bulge that occurs at 4–6 months of gestation lessens as development occurs between 6 and 8.5 months. A developmental disruption is speculated to occur at some point during the last 3 months and the variation in the time of occurrence accounts for different clinical TM phenotypes.^{92,101}

Alternatively, disruption of the choroidal vasculature in fetal development has been put forth as a possible etiology. In 1997, Teitelbaum et al suggested an abnormal event in the embryologic development of the choroidal vasculature underlying the macula could cause a disruption of the RPE and give rise to the torpedo-like lesion.¹¹⁵ With the advent of OCT-A, the hypothesis focuses on a malformation of the RPE-choriocapillaris complex.^{91,108,114,117,118} Using OCT-A, Giannakaki-Zimmermann et al evaluated 4 patients with TM lesions. In this study, all lesions demonstrated normal superficial and deep vascular capillary beds within the TM lesion, and choriocapillaris with on average 26% less signal when compared with healthy choriocapillaris in the same region.¹¹⁴ Additional studies have revealed a reduced capillary density within the choriocapillaris using OCT-A.^{101,105} Some reports have shown normal^{91,118} large caliber vessels of Sattler's and Haller's layer while others show them to be mildly attenuated.¹¹⁷ Giannakaki-Zimmermann et al suggested since the TM lesion extends into the choriocapillaris, congenital choroidal malformation is involved. However, it remains speculative whether the choroid is the primary site of malformation or related to a defect in the development of the RPE in view of the fact that the choroid needs an intact RPE to develop.^{105,114}

Further ideas on TM development have been proposed. Pian et al suggested that the lesion is a paramacular coloboma with the defect at the horizontal raphe of the nerve fiber layer.99 Trevino et al also endorse the idea of a colobomatous etiology. He cites the diagnostic criteria of Mann for macular coloboma and TM intersect when there is associated choroidal excavation in TM.¹⁰² Golchet et al suggest a malformation of the emissary canal of the long posterior ciliary artery and nerve given the anatomical proximity with TM lesions.⁸⁷ The temporal long posterior ciliary artery joins the long posterior ciliary nerve, which together penetrates through the sclera and runs temporarily towards the anterior of the eye through the suprachoroidal space and choroid. They hypothesize that the modification of the sclera during the embryological process allowing the nerve and artery to pass may have an effect on the developing choroid and RPE; thus leading to TM formation.⁸⁷

	Pathophysiology	Subtypes	Association	Appearance of Cavitation on OCT	Potential Complications
Intrachoroidal Cavitation (ICC)	Acquired following structural weakening of peripapillary tissue ^{12,17,26} Some forms of macular ICC are thought to arise in utero ^{40,47}	Peripapillary ICC (pICC) Macular ICC (mICC)	pICC - Pathologic myopia mICC- Pathologic myopia (PCA), posterior staphyloma (NCMD)	pICC- Posterior scleral bowing with choroidal tissue cavitation and a flat RPE ^{12,15} mICC lesions tend to occur on the slope of the staphyloma ^{35,36}	Retinoschisis and macular detachment ^{28–33}
Focal Choroidal Excavation (FCE)	Unclear - inflammatory, ^{68,71} infectious, ⁸⁴ vascular, ^{51,60} developmental, ⁷⁹ and congenital ^{42,57,85} etiology have all been suggested.	Conforming Nonconforming	Associated with a host of choroidal and retinal pathology - see text	Conforming- Area of focal choroidal thinning and concavity with normal overlying retina. Without scleral ectasia, posterior staphyloma, or PED ^{42–45} Noncomforming- Separation of photoreceptor tips and underlying RPE ⁴³	PCV, CSC, CNV ^{59,63}
Torpedo Maculopathy (TM)	Congenital abnormality of the RPE and/or choroid ^{105,114} Unclear if RPE or choroid is primary site of malformation in fetal development ¹⁰¹	Туре I Туре 2 Туре 3	No known associated conditions	Type I- Thinning of RPE, outer nuclear layer, interdigitation zone, ellipsoid zone, and inner choroid ^{80,85,104,105} Type 2-Type I features + attenuation of outer retinal structures and subretinal cavitation ⁹⁷ Type 3-Type I features + choroidal excavation ¹⁰⁵⁻¹⁰⁷	CNV ^{92,105}

Table I	Choroidal	Cavitary	Disorders	Summary
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Choroidal neovascularization has been shown to occur in TM. Shirley et al documented a case of a patient with TM-related CNV which responded well to a single injection of anti-VEGF (Ranibizumab).⁹² However, the authors stated that given the structural changes present, CNV remains an ongoing concern and should be monitored accordingly.

The choroidal cavitary disorders discussed in this review are summarized in Table 1.

Conclusion

A true visualization and a better understanding of choroidal cavitary disorders have eluded the medical community until recently with the introduction of optical coherence tomography. Over time as these conditions are encountered, tracked, and imaged with more sophisticated techniques we will gain much more knowledge about the potential impact on the patient and strategies to mitigate associated pathology. The connection between these choroidal cavitary disorders is difficult to establish as most hypotheses to explain pathogenesis cite poorly documented embryological mechanisms and will be difficult to test in their present level of articulation. Although these disorders can be asymptomatic, and at times even non-progressive, the possible complications are sight threatening and patients benefit from prompt diagnosis and treatment. To conclude, we have highlighted the known similarities and differences in an attempt to establish a preliminary connection to help aid in the future strategies to treat, and perhaps prevent, sight threatening complications from seemingly benign findings.

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