



CHAIR-SIDE REFERENCE: VITREOMACULAR INTERFACE ANOMALIES

Posterior vitreous detachment (PVD) is a gradual, typically age-related separation of the posterior vitreous from the inner retinal surface. As the vitreous contracts and liquefies, it detaches from the mid-peripheral retina, progressing to then detach from the perifoveal, foveal and peripapillary regions sequentially. When vitreous liquefaction occurs before vitreoretinal separation, localised traction may occur, leading to an anomalous PVD with potential complications at the macula. OCT (radial and raster B-scans) has revolutionised the understanding, diagnosis and management of anomalous PVDs and associated complications.

Retinal photo	Optical coherence tomography	Definition and aetiology	Clinical features and management
Vitreomacular Adhesion (VMA)			
		<ul style="list-style-type: none"> Vitreous adhesion to central macula with no evidence of retinal morphologic changes. Normal finding in natural course of PVD Classification based on size of adhesion <ul style="list-style-type: none"> Focal adhesion $\leq 1500\mu\text{m}$ Broad adhesion $>1500\mu\text{m}$ 	<ul style="list-style-type: none"> Unremarkable fundus appearance Attachment of the vitreous cortex within a 3mm radius of fovea centre with perifoveal vitreous detachment but no detectable distortion in foveal architecture <p><i>Review routinely</i></p>
Vitreomacular Traction (VMT)			
		<ul style="list-style-type: none"> Vitreous traction on the central macula causing anatomical distortion of foveal surface and/or intra-retinal abnormalities. Part of anomalous PVD progression Classification based on size of adhesion <ul style="list-style-type: none"> Focal adhesion $\leq 1500\mu\text{m}$ Broad adhesion $>1500\mu\text{m}$ 	<ul style="list-style-type: none"> May have a yellow discoloration of the central macula Attachment of the vitreous cortex within a 3mm radius of fovea centre with perifoveal vitreous detachment & associated foveal abnormalities Can involve intra-retinal/structural changes: foveal distortion, elevation, schisis, pseudocyst formation, elevation of the retina from the RPE or outer foveal microdefect (see below) <p><i>Consider referral if associated with symptoms/vision reduction or history of full thickness macula hole in the fellow eye (increased risk)</i></p>
Lamellar Macular Hole (LMH)			
		<ul style="list-style-type: none"> Partial thickness foveal defect Caused by anomalous PVD with partial avulsion of foveal tissue. Possible degenerative mechanism has also been speculated. Other non-PVD causes may also present with a LMH such as MacTel, partial closure of full-thickness macula hole, cystoid macula oedema (non-primary) 	<ul style="list-style-type: none"> Darker, reddish round lesion at the fovea Irregular foveal contour, focal cavity with undermined edges (assists in differentiating from ERM foveoschisis - see below) Signs suggestive of retinal tissue loss (pseudo-operculum, thinning at/around the fovea) May also show epiretinal proliferation (thick, iso-reflective, non-contractile material), foveal bump or EZ disruption <p><i>Consider referral if associated with symptoms/vision reduction</i></p>
Full-Thickness Macular Hole (FTMH)			
		<ul style="list-style-type: none"> Full-thickness foveal break Can be idiopathic or due to anomalous PVD (primary FTMH) Other aetiologies include trauma, myopia, iatrogenic, neuro-degenerative (e.g. MacTel) Classification based on size (narrowest aperture) and presence or absence of vitreomacular traction: Small: $\leq 250\mu\text{m}$, Medium: $250\mu\text{m}$-$400\mu\text{m}$, Large: $>400\mu\text{m}$ 	<ul style="list-style-type: none"> Appears as a round red lesion at central macula Interruption of all neural retinal layers from internal limiting membrane (ILM) up to (not including) the RPE Edges of the macular hole are typically rounded and pulled anteriorly, often containing pseudocysts <p><i>Typically requires prompt referral: Surgical outcome is better with shorter duration of symptoms, smaller hole size and pre-operative VA.</i></p>



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Epiretinal Membrane (ERM)			
		<ul style="list-style-type: none"> • A fibrocellular contractile proliferation over the internal limiting membrane (ILM) • Exact aetiology is unclear • Can occur at any stage of vitreous separation, referred to as idiopathic ERM • Can also be secondary to trauma, past surgery, chronic ocular disease 	<ul style="list-style-type: none"> • Thin distinct hyper reflective layer above ILM • Can cause tractional stress on underlying retina • May be associated with wrinkling of retinal surface, loss of foveal pit, retinal thickening and pseudocystic spaces <p><i>Consider referral if associated with symptoms/vision reduction</i></p>
Epiretinal Membrane (ERM) Foveoschisis			
		<ul style="list-style-type: none"> • Schisis at the fovea due to contractile epiretinal membrane (previously known as tractional lamellar hole) • Caused by mechanical displacement of retinal layers secondary to tractional force from the epiretinal membrane. 	<ul style="list-style-type: none"> • Roundish, slightly darker central lesion in the fovea • Foveoschisis at the level of Henle fibre layer (splitting in outer nuclear and plexiform layers) • Presence of contractile epiretinal membrane (differentiating from lamellar macula hole, see above) • May be associated with microcystoid spaces in inner nuclear layer, retinal thickening and retinal wrinkling <p><i>Consider referral if associated with symptoms/vision reduction</i></p>
Macular Pseudohole			
		<ul style="list-style-type: none"> • A lesion with a similar fundoscopic appearance to full thickness hole but without full thickness defect or signs of retinal tissue loss • Caused by mechanical displacement of retina towards the foveal centre via centripetal tangential traction 	<ul style="list-style-type: none"> • Discrete, red, round or oval lesion in the fovea • Presence of ERM with central opening sparing the fovea and associated surrounding retinal thickening • Verticalised/steepened foveal profile • May be associated with microcystoid spaces and near normal foveal thickness. • No loss of foveal tissue <p><i>Consider referral if associated with symptoms/vision reduction</i></p>
Outer foveal microdefects (OFMD)			
		<ul style="list-style-type: none"> • A small reddish foveal lesion with a small focal discontinuity within the outer retina • Current naming controversy: also includes macular microhole and foveal red spot syndrome • Common aetiologies include VMT, previous VMT or adhesion. • Other aetiologies include phototoxicity, blunt trauma, chronic macula oedema, spontaneous closure of full thickness macular hole, MacTel2 	<ul style="list-style-type: none"> • Appears as small reddish lesion at or adjacent to fovea • Focal discontinuity of RPE, photoreceptors and/or external limiting membrane • May present with irregular foveal pit, symptoms of PVD, overlying vitreoretinal traction/adhesion <p><i>Review routinely</i></p>

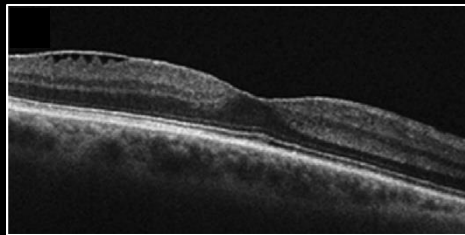


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In recent years, there have been several important developments in the use of OCT to detect prognostic markers or structural signs associated with ERM that can be used to predict visual outcomes. The key structural signs associated with lower visual acuity and poorer visual outcomes from surgery are the presence of ectopic inner foveal layers (EIFL) and disorganisation of the inner retinal layers. Although there is currently no widely accepted OCT staging system, the successive ERM stages identified by Govetto et al (2017) are associated with progressively worse visual acuity. Studies suggest referral for surgery prior to changes seen in stage 3 and 4 may be beneficial.

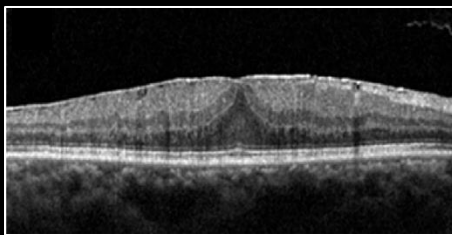
Epiretinal Membrane Progression

Stage 1 ERM



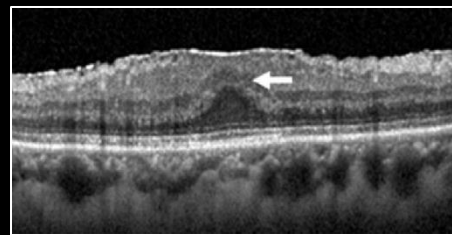
The foveal pit is preserved, and individual retinal layers are distinguishable.

Stage 2 ERM



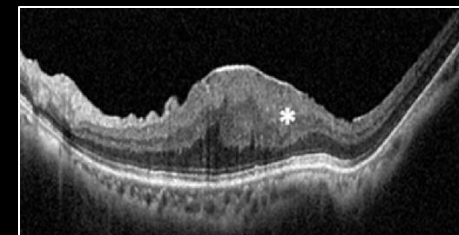
There is macular thickening and loss of the foveal pit, but retinal layers are still distinct

Stage 3 ERM



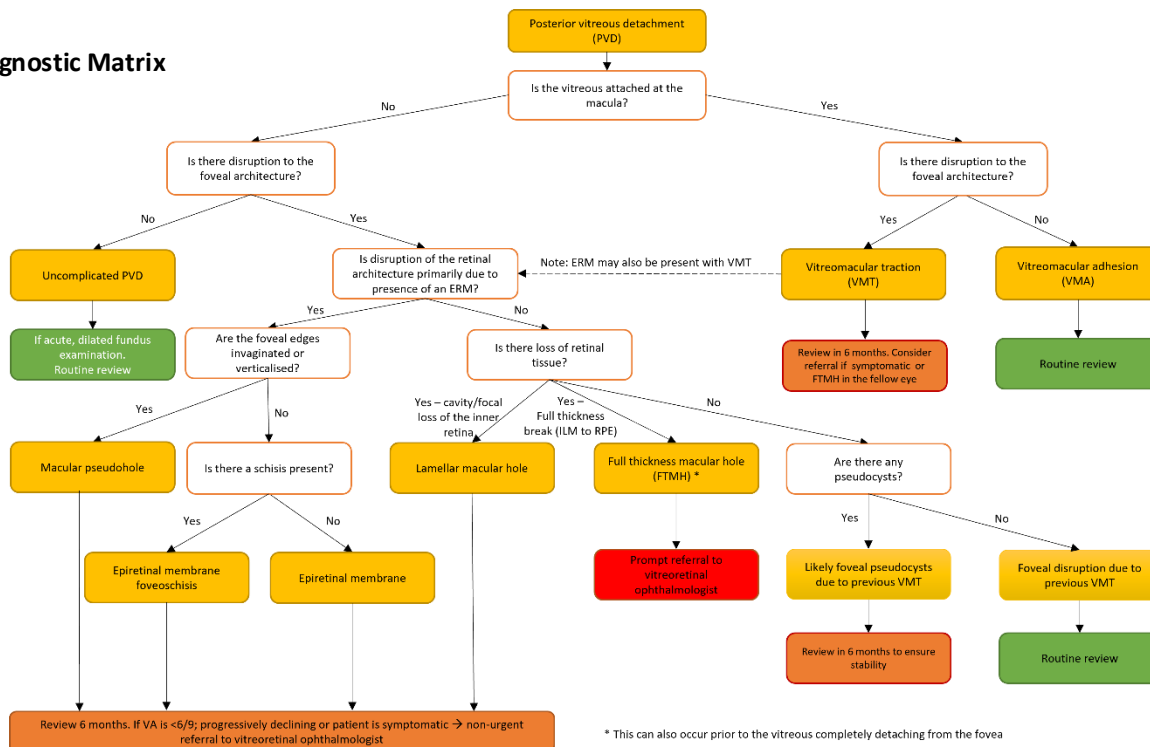
Presence of ectopic inner foveal layers (EIFL) seen as a continuous hyper or hyporeflective band extending from the INL and IPL (white arrow).

Stage 4 ERM



EIFL with disorganisation of the inner retinal layers (white asterisks)

VMI Diagnostic Matrix



Please note this chair-side reference was designed to assist optometrists in distinguishing between the different types of vitreomacular interface disorders. It provides general information only and may not be applicable to atypical cases. For more specific advice, our telehealth service is available to all optometrists in Australia.



- One-to-one consultation with a senior CFEH optometrist for non-urgent diagnosis and management support.
- Interactive, therapeutic CPD points.
- Consultations are at no charge, thanks to the generous support of Guide Dogs NSW/ACT.

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