

Putting the "Oh" in OCT

Danica J. Marrelli, OD, FAAO, AAO Diplomate
 Clinical Professor
 Assistant Dean of Clinical Education
 University of Houston College of Optometry

DMarrelli@uh.edu

Mark T. Dunbar, O.D., F.A.A.O.
 Bascom Palmer Eye Institute
 University of Miami Health System
 Miami, FL

mdunbar@miami.edu

1

Financial Disclosure - Marrelli

- Allergan
- Bausch & Lomb
- Carl Zeiss Meditec
- Glaukos
- M&S Technologies
- Thea

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Mark Dunbar: Disclosure

- Optometry Consultant/Advisory Board
 - Carl Zeiss
 - Orasis
 - Allergan
 - Visus
 - Regeneron
 - Tarsus
 - Astella
 - Topcon
 - Apellis
 - Avellino
 - B&L
 - Thea

Mark Dunbar does not own stock in any of the above companies

3

Reports

Optical Coherence Tomography 1991

DAVID HUANG, ERIC A. SWANSON, CHARLES P. LIN, JOEL S. SCHERMAN, WILLIAM G. STENSON, WARREN CHANG, MICHAEL R. HEH, THOMAS FLOTTE, KENTON GREGORY, CARMEN A. PULIAFITO, JAMES G. FUJIMOTO*

A technique called optical coherence tomography (OCT) has been developed for noninvasive cross-sectional imaging in biological systems. OCT uses low-coherence interferometry to produce a two-dimensional image of optical scattering from internal tissue microstructures in a way that is analogous to ultrasonic pulse-echo imaging. OCT has longitudinal and lateral spatial resolutions of a few micrometers and can detect reflected signals as small as $\sim 10^{-16}$ of the incident optical power. Tomographic imaging is demonstrated in vitro in the peripapillary area of the retina and in the coronary artery, two clinically relevant examples that are representative of transparent and turbid media, respectively.

4

The Evolution of OCT Imaging

- OCT has changed how clinicians look at the retina
- OCT has changed how we manage glaucoma
- The assessment of retinal abnormalities and glaucoma based on OCT imaging has advanced eye care
- OCT in Optometry practices ~ 70-85%
- As the technology has evolved -> prices continue to come down

5

Advances in SD-OCT

- Improving software
- **Faster – virtual angiography**
- Noise reduction/over sampling technology
- Wider and deeper scans
- Greater density in the scans
- Improvements in 3D imaging
- Enhanced depth imaging – imaging choroid
- Progression analysis software

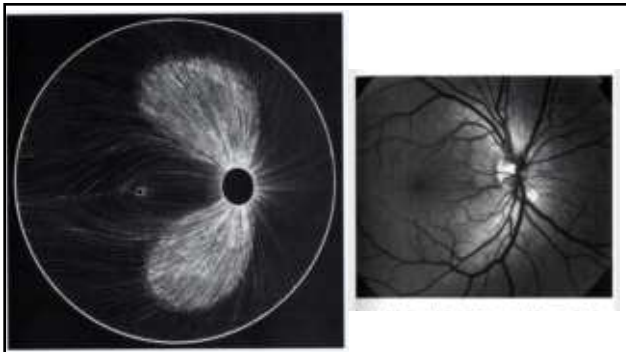
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The OCT in Glaucoma

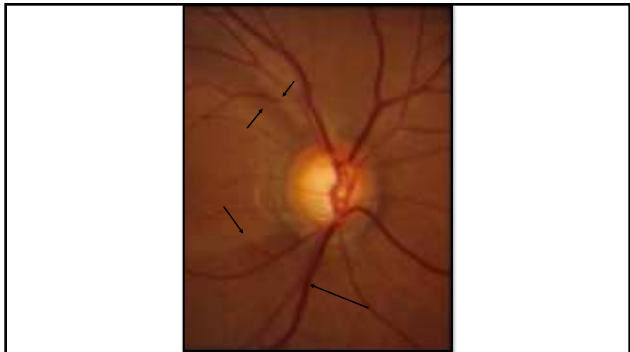
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- ## The OCT in Glaucoma
- When is it glaucoma?
 - Differentiating glaucoma nerves from physiologic nerves
 - Sometime it's very easy but not always
 - Following glaucoma suspects
 - Recognizing early change -> **green disease**
 - Recognizing when it's NOT glaucoma - **red disease**
 - Determining progression
 - When is the OCT not as helpful?

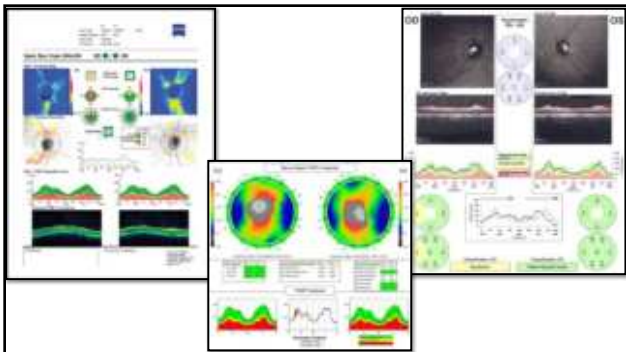
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11

- ## Quantifiable/Objective Imaging
- "Diagnostic capability":
 - Good for early glaucoma
 - Excellent for moderate to severe glaucoma
 - Improved when more than one parameter is evaluated
 - Many of us rely on imaging devices to identify glaucoma (and progression)
 - Has imaging become "the answer"?



12

OCT RNFL Basics:
How Do I Make Sense of This?

13

Systematic Strategy

- Quality
 - Signal Strength
 - Circle Placement
 - Movement?

14

Systematic Strategy

- Thickness Map
- Deviation Map

IMPORTANCE OF BLOOD VESSELS!!!!

15

Systematic Strategy

- Thickness Profiles
 - Compared to normative data
- KEYS:
 - SYMMETRY
 - MODULATION

Good at picking up notches in NRR

16

Systematic Strategy

- Quadrant and Clock Hour RNFL analysis

17

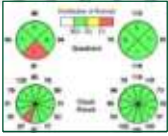
Do you only do 1 RNFL scan?

- How do you know how accurate/reliable that scan is?
- Instead do at least 3RNFL scans at a time
 - at a minimum do 2 scans
- Ensures consistency/reliability

18

What is the Reproducibility of RNFL OCT Clock Hour Measurements

- A. 0-3 microns
- B. About 4-5 microns**
- C. About 10 microns
- D. > 10 microns



19

Where Can We Go Wrong?

- Artifacts
- Interpretation Errors

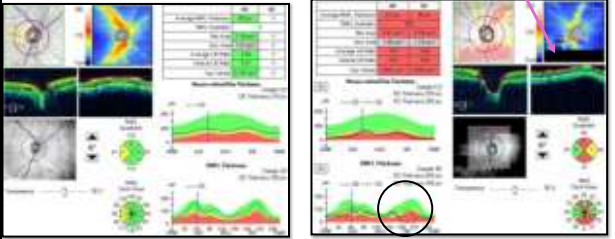
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Artifacts in Spectral Domain OCT (SD-OCT)

- **Artifacts are common in SD-OCT:**
 - 6-44% of scans will have some type of artifact
 - Artifacts increase in the presence of other pathology
 - Not all artifacts are visible on standard printout
- **Causes of Artifacts:**
 - Technician/Acquisition Errors
 - Software Errors
 - Patient Dependent:
 - Pupil Size
 - Media
 - Concurrent pathology (PVD, ERM)
 - Myopia-related changes

21

Acquisition Errors



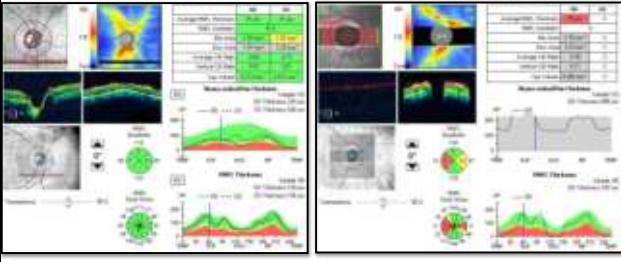
Circle Displacement

Image truncation

Images courtesy of Marcus Gonzales, OD, FAOD

22

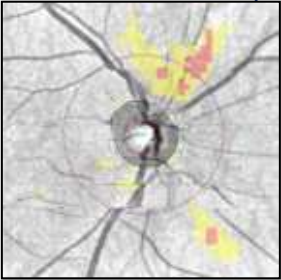
Acquisition Errors – Don't Blink!



Images courtesy of Marcus Gonzales, OD, FAOD

23

Acquisition Errors – Motion/Rotation



24

Software Errors

- Improper boundaries of optic nerve
- Improper segmentation of layers (RNFL, GC-IPL)
 - Pure segmentation error
 - Poor signal strength
 - Often related to coexisting disease

25

Segmentation Errors

Image courtesy of Andrew Rixon, OD

26

Clinical Interpretation Errors

- Reliance on reference database color indicators
 - Red Disease
 - Green Disease
- Failure to recognize non-glaucomatous causes of RNFL thinning

27

Reliance on Reference Database Color System

- Reference Database (aka “normative” database):
Reference population without disease in question, to which an individual patient’s data will be compared

Question: Does abnormal DATA mean you have DISEASE?

28

29

“Red Disease”

False Positive

30

REVIEW

Green disease in optical coherence tomography diagnosis of glaucoma

Abdennour B. Saad¹, Michael Manges^{2,3*} and Richard H. Lee⁴

Abstract of review:
Optical coherence tomography (OCT) has become an integral component of modern glaucoma practice. Utilizing color-coded, cross-sectional images, glaucoma diagnosis and disease severity can be determined by the presence of structural changes. However, green disease (GD) has become a growing concern as it can be mistaken for glaucoma, leading to incorrect diagnosis of glaucoma and to glaucoma management.

Keywords:
Glaucoma, green disease, optical coherence tomography.

31

“Green Disease”

False Negative

32

“Green Disease”

33

Non-glaucomatous Causes of RNFL thinning

Masqueraders

- Retinal Disease*
- Ischemic Optic Neuropathy*
- Other optic neuropathies
- Neuro-degenerative disease
- Myopia***

34

Kevin, 52yo WM

35

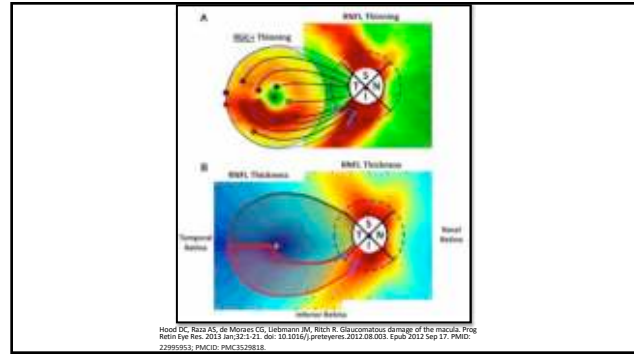


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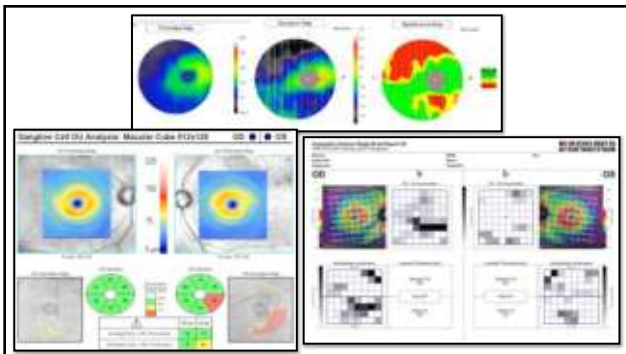
Newest Addition to Glaucoma Diagnosis Arsenal: Macular Imaging

- 1998: Zeimer et al reported on macular thickness loss in patients with known glaucomatous damage
- 2003: Greenfield reported correlation between total macular thickness and MD on VF in glaucoma patients (time domain OCT)
- 2013: Hood et al – extensive investigation of segmented “RGC+” (RGC + IPL) layer and description of the “Macular Vulnerability Zone” (MVZ)

37



38



39

Advantages of Macular Analysis

- Macula contains ~50% of retinal ganglion cells
 - Glaucoma is a disease of these cells
 - Macular thinning/irregularity cannot be detected during clinical exam
- More reproducible measure (if not using retinal nerve fiber layer) than peripapillary RNFL
 - Fewer blood vessels than other cell components
 - Less anatomic variation compared to optic disc/peripapillary region
- Better superior/inferior symmetry and symmetry between eyes than peripapillary RNFL

40

Disadvantages of Macular Imaging

- Macular imaging is not helpful in glaucoma cases in which patients have concurrent macular disease
 - AMD
 - ERM
 - CME
 - DME
 - Macular hole

41

Glaucoma and Myopia

A Diagnostic Dilemma

- **Myopia and Glaucoma**
 - Myopic epidemic: 5 Billion myopes by 2050
 - Myopia is a risk factor for glaucoma development
- **Myopic Discs can be difficult to evaluate**
 - Tilt
 - Peripapillary changes
 - Flattening of cup
- **Challenges with OCT in myopic eyes**
 - Difficult to acquire image
 - Higher incidence of segmentation errors
 - RNFL database not typically adjusted for RE or AL
 - RNFL and macular thickness may be affected by increased AL

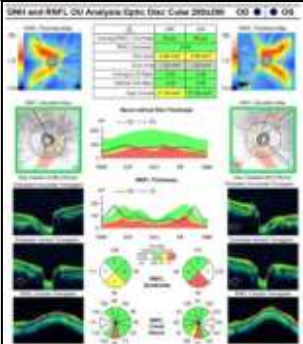
42

Glaucoma and Myopia *A Diagnostic Dilemma*

- **RNFL**
 - Decreased RNFL with increasing AL in S, I, N sectors
 - RNFL more temporally located (S/I peaks shifted)
 - Increasing axial length associated with "false positive" (red disease) RNFL OCT
 - Poor agreement with VF
- **Macular Ganglion Cell**
 - Average thickness reduced in high myopia
 - Tend to have diffuse circular thinning with irregular inner margin
 - "GCIPL Hemifield Test" shown to have high sensitivity and specificity in high myopia (Kim YC, et al. IOVS 2016;57:5856-63)

43

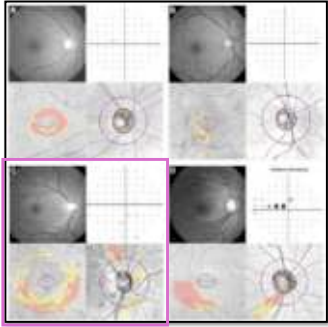
Temporally displaced RNFL peaks



Tan, et al. Br J Ophthalmol 2019;103:1347-1355

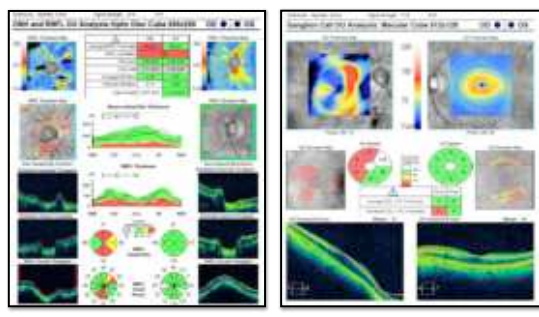
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GCIPL False+



Kim, et al. Ophthalmology 2015;122(3):502-510

45



46

Case for Discussion

CM, 51yo HM

- Referred for glaucoma suspicion due to ONH appearance
- **POH:**
 - LASIK OU (2000), PRK OS (2014)
- **FOH:**
 - (+) glaucoma (maternal gm)
- **PMH:**
 - unremarkable

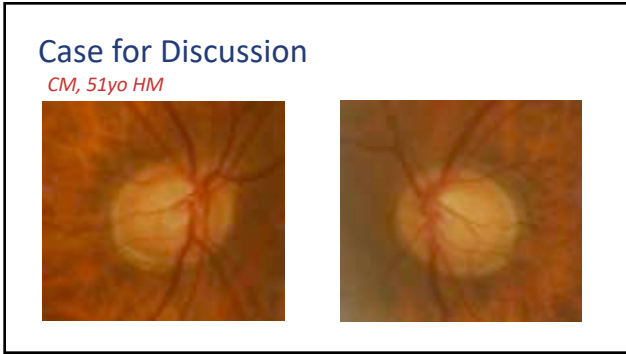
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Case for Discussion

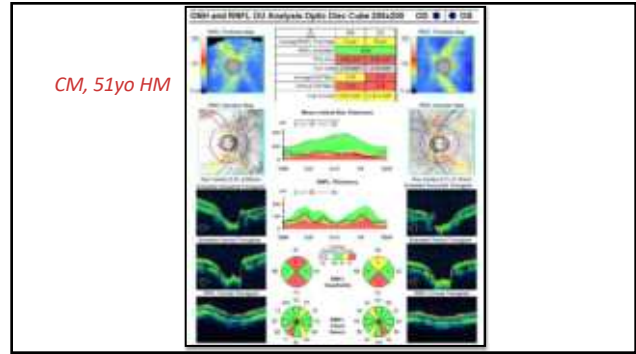
CM, 51yo HM

- BCVA: 20/20 OD, OS
- Normal pupils, motility, CVF
- SLE: LASIK flaps visible, otherwise normal
- Gonioscopy: open to CB 360° OD, OS
- Tmax: 18mmHg OU
- CCT: 523 OD 489 OS

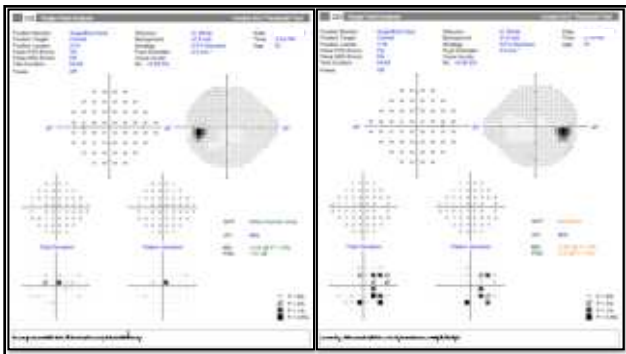
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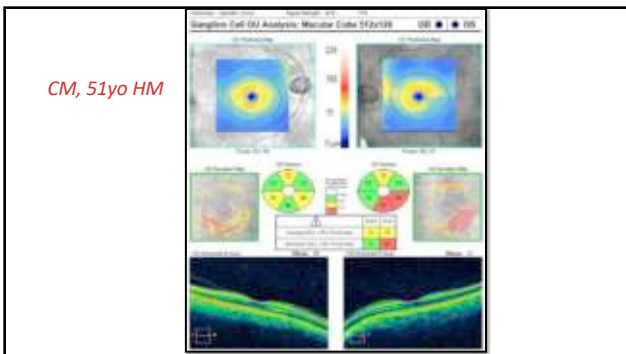


51

Case for Discussion
CM, 51yo HM

- Do you think this is glaucoma?
- What else should we do?

52

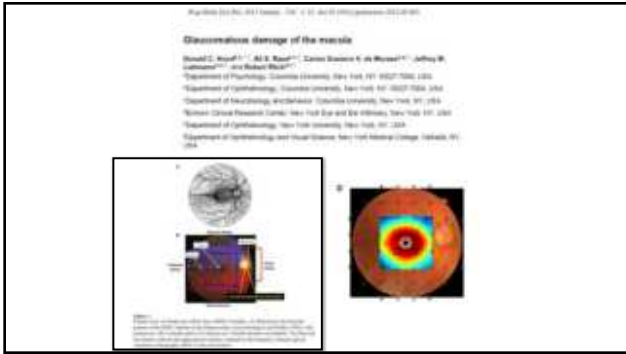


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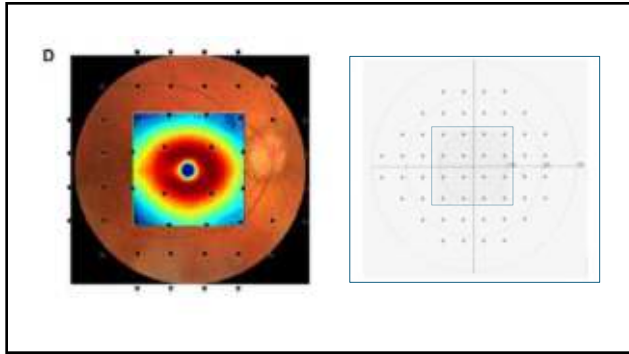
What about the 10-2 VF?

- Central 8 degrees from the center of the foveal contains more than 30% of retinal ganglion cells
- 24-2 and 30-2 test strategies use a 6 degree test grid pattern; these points fall outside of the densist region of ganglion cells
- 10-2 test strategy uses a 2 degree test grid
- Recent research has shown that in some patients with small regions of macular ganglion cell loss, 10-2 testing may be better able to detect VF loss

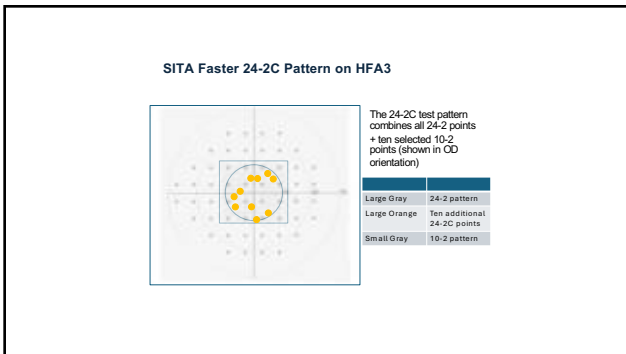
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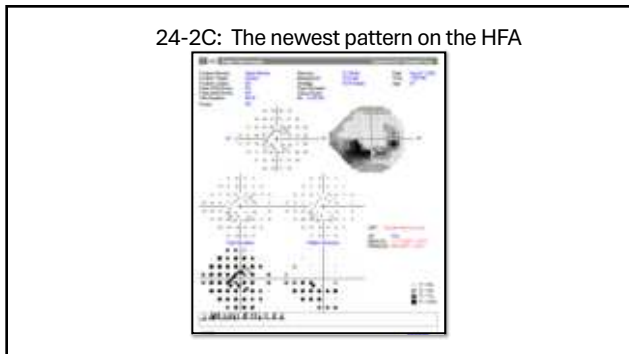
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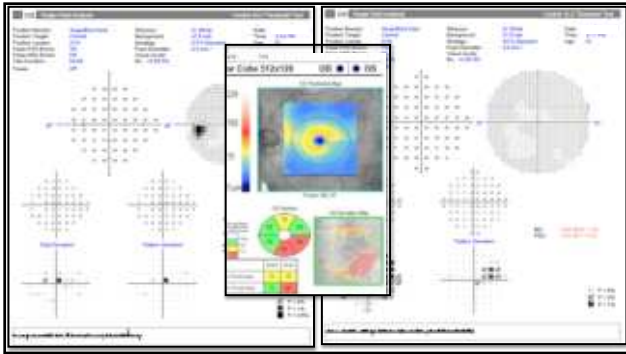
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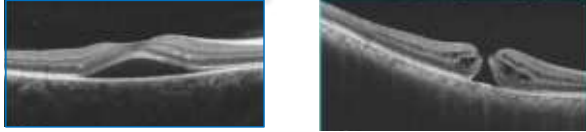
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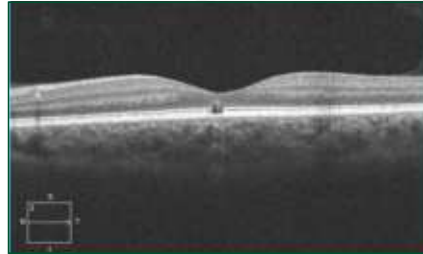
Don't Make It More Complicated Than it Needs to be

- Many macular disease conditions have a "signature" OCT feature
- Learn what those are and the diagnosis and interpretation becomes easier



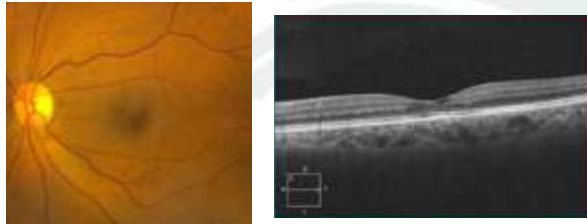
61

What does This patient have?



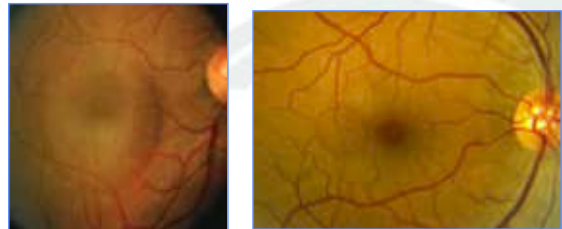
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What does this "signature" OCT represent?



63

Where is the Fluid?



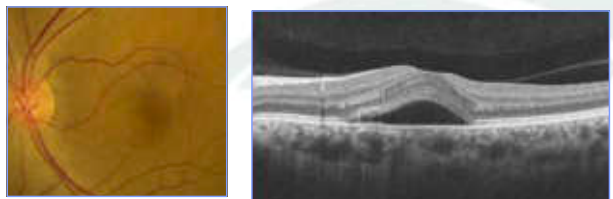
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Where is the Fluid?



65

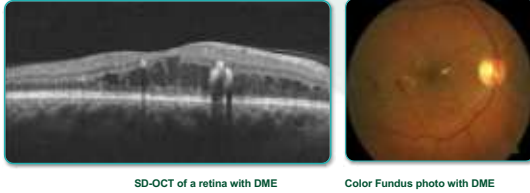
Central Serous Chorioretinopathy (CSR)



44 y/o Female: Notes blur in the LE X 1 mo BCVA: 20/25

66

Diabetic Macular Edema (DME)



SD-OCT of a retina with DME

Color Fundus photo with DME

67

The Macula in Diabetes



68

The Macula in Diabetes

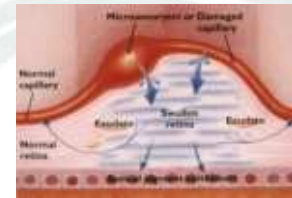
- Is there retinopathy?
- Is there retinal thickening?
- Is there fluid?
- How close is it to the macula?



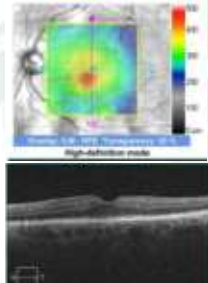
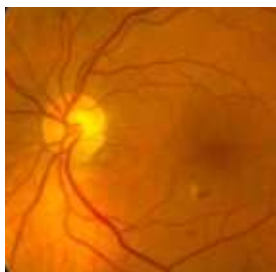
69

Macular Edema

- Thickening of the retina
- Secondary to leaky microaneurysms
- **90% of visual loss in diabetes**



70



71

CSME

- Retinal thickening within 500 microns from the center of the FAZ
- Hard exudates associated with retinal thickening 500 microns from center of FAZ
- Zones of retinal thickening > 1 DD in area, any part of which is 1 DD from the center of the fovea

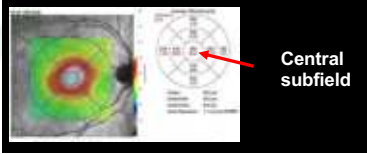


72

2017 DME Classification:

Center Involved or Not?

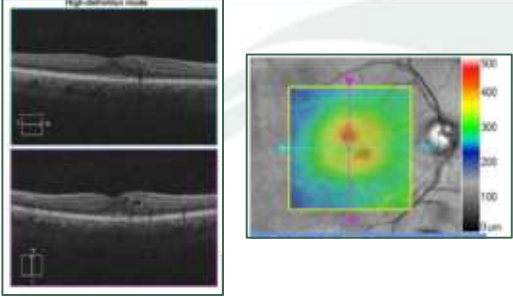
- ETDRS definition of "clinically significant macular edema" modified in era of OCT
- Randomized clinical trials of anti-VEGF agents used presence of DME in **OCT central subfield**



Central subfield


Figure 1. Guan Dong, et al. "Risk factors for diabetic macular edema: results from 2 phase II randomized trials: RISE and RISE2." Ophthalmology 119:4 (2012): 808-815.
Figure 2. Chakrabarti, et al. "Intravitreal aflibercept for diabetic macular edema: 100-week results from the VISTA and VIVID studies." Ophthalmology 122:10 (2015): 2044-2052.

73



74

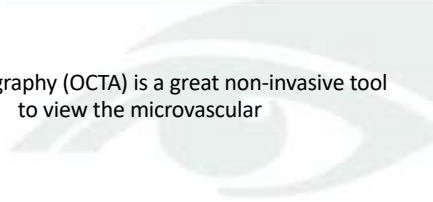
Diabetic Macular Edema (DME)



SD-OCT of a retina with DME **Color Fundus photo with DME**

75

OCT Angiography (OCTA) is a great non-invasive tool to view the microvascular

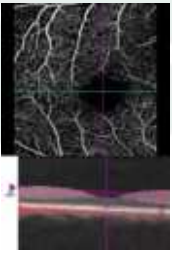


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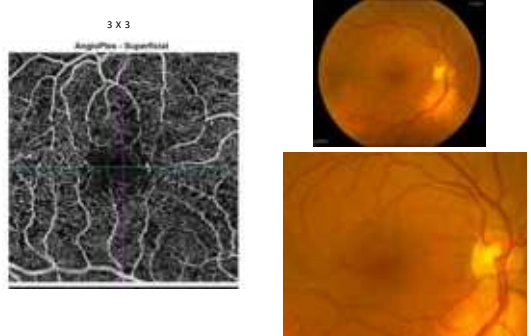
OCT Angiography (OCTA)

The Basic Idea of How it Works:

- Capturing motion** in the retina
- Scans at 68,000 A-scans per second
 - Traditional SD OCT scan at 28,000 to 40,000 A-scans per second
- Compares **repeat scans** acquired at the **same position** in the retina to look for changes - motion

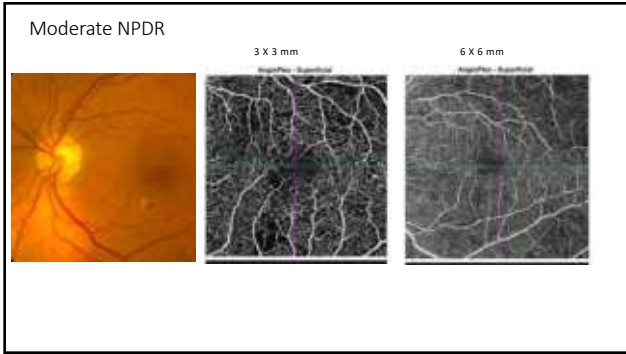


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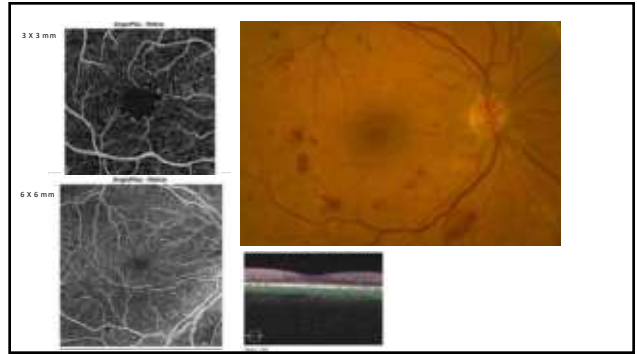


3 x 3
AngioPlex - Superficial

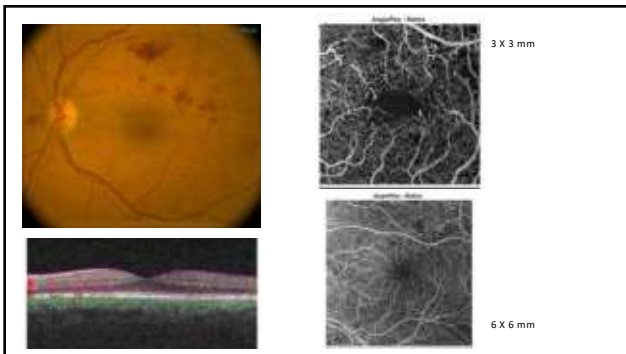
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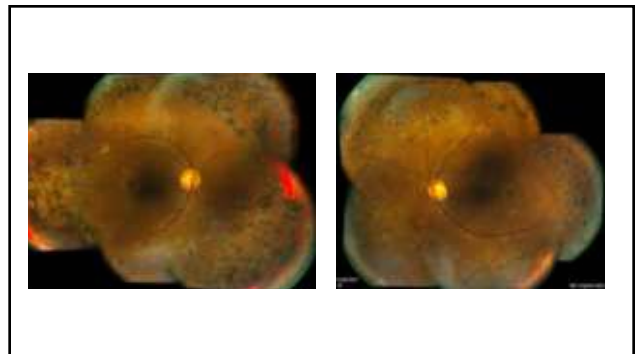
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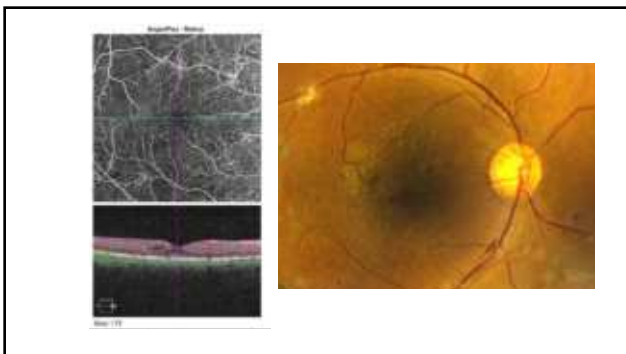
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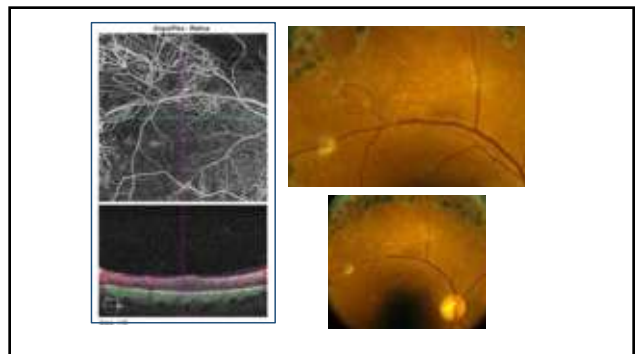
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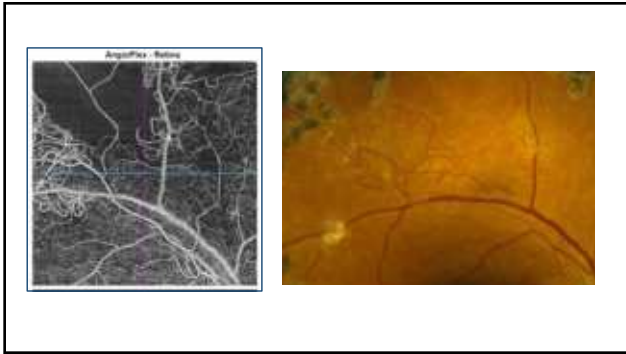
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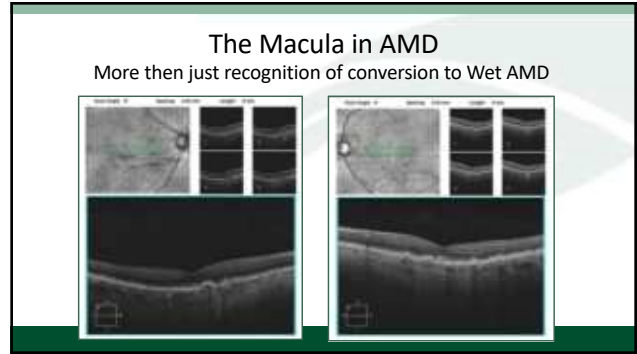
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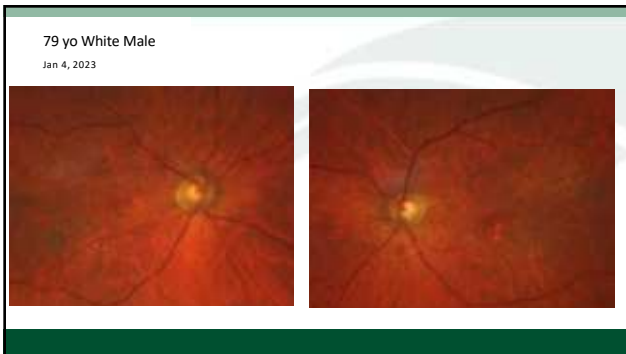
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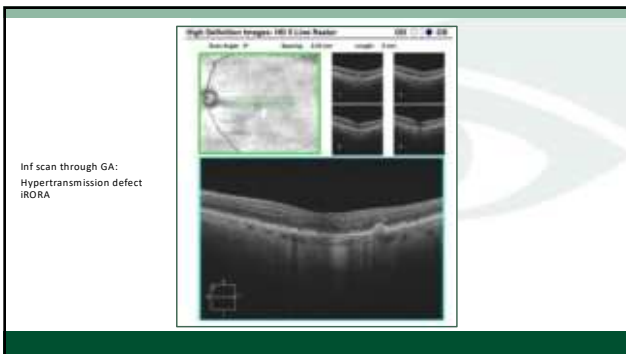
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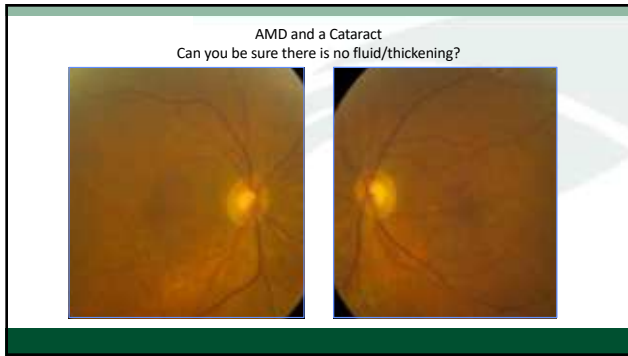
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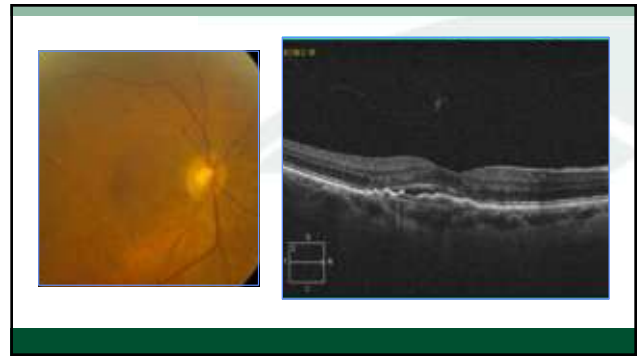
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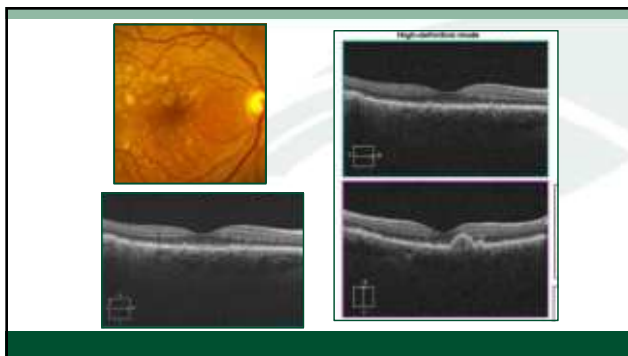
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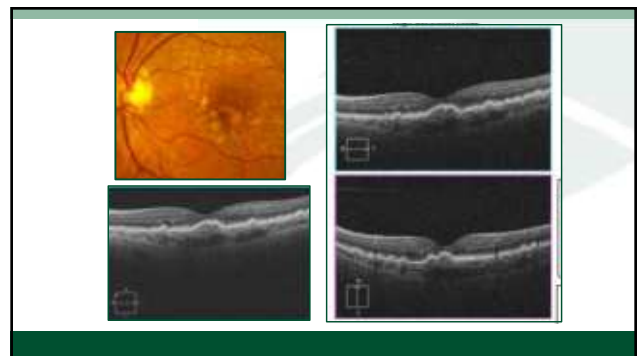
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92



93



94

Risk Factors for Progression to Wet AMD

- Traditionally based on clinical appearance
- Intermediate AMD
 - Large drusen > 125 microns
 - RPE mottling/pigmentary abnormalities
- Risk of conversion to wet AMD over 5 years > 50%

95

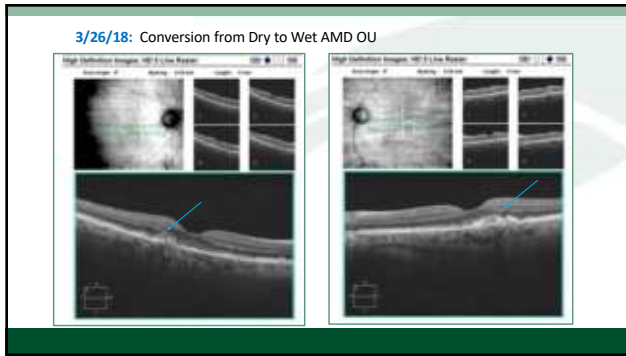
AMD Is the Leading Cause of Blindness for Caucasians in the US¹

10-year risk of progression for the highest risk category (AREDS simple scale)²

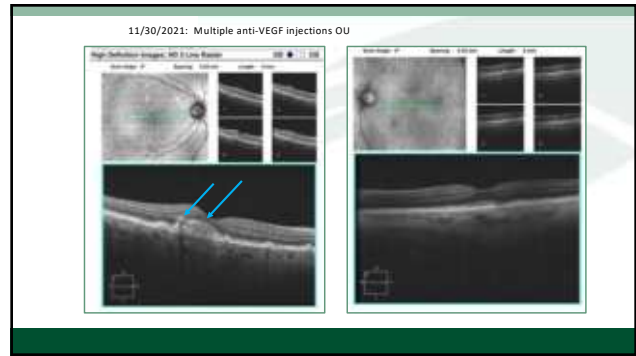
- Early AMD
- Intermediate AMD
- Central GA: 53.9%
- Advanced AMD
- Neovascular AMD (nvAMD): 47.6%

AMD, age-related macular degeneration; AREDS, Age-Related Eye Disease Study; GA, geographic atrophy; nvAMD, neovascular AMD. 1. Eye Diseases Prevalence Research Group. Arch Ophthalmol. 2006;124(4):477-483. 2. Fain GL, et al. Ophthalmology. 2013;120(6):1844-851. 3. Chew DF, et al. JAMA Ophthalmol. 2014;132(3):272-277. 4. Age-Related Study Disease Study Research Group. Arch Ophthalmol. 2002;120(11):1570-1574.

96



103



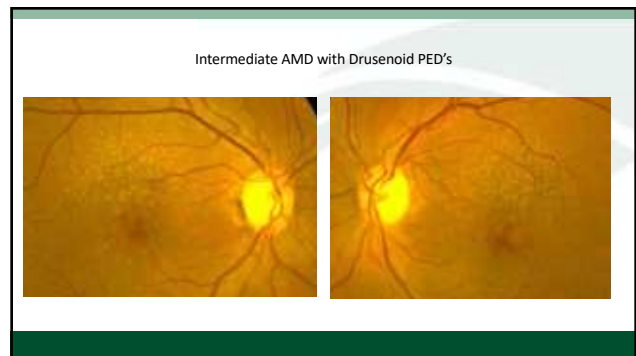
104

Hyperreflective Retinal Foci (HRF)

- Secondary to:
 - DR/DME, RVO, AMD, CSR, Uveitis, MacTel; IRD
- Most likely activated inflammatory microglia cells
- **Biomarker for disease progression**

Leung K, et al. Hyperreflective Retinal Foci (HRF) in the Macula. Invest Ophthalmol Vis Sci. 2015;56(10):3073-3077.
 Chakraborty S, et al. Hyperreflective Retinal Foci (HRF) in the Macula. Invest Ophthalmol Vis Sci. 2015;56(10):3078-3082.
 Fingert J, et al. Hyperreflective Retinal Foci (HRF) in the Macula. Invest Ophthalmol Vis Sci. 2015;56(10):3083-3087.
 Fingert J, et al. Hyperreflective Retinal Foci (HRF) in the Macula. Invest Ophthalmol Vis Sci. 2015;56(10):3088-3092.

105



106

Reticular Pseudo Drusen

- Subretinal collections of granular, interlacing, hyper-reflective material located above RPE
- Commonly found in the superior macula or close to superotemporal arcade
- Undergo a characteristic lifecycle of growth, invasion into the ellipsoid zone, and finally regression
- Reticular pseudodrusen is associated **with an additional 2-6-fold increased risk of progression to nAMD or central GA**,
 - Risk of progression higher for reticular pseudodrusen located outside the macula

107



108



109

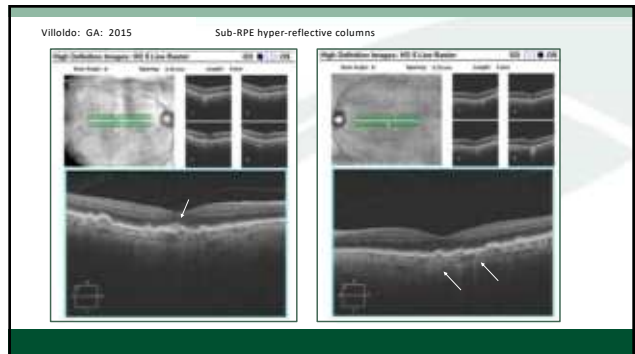


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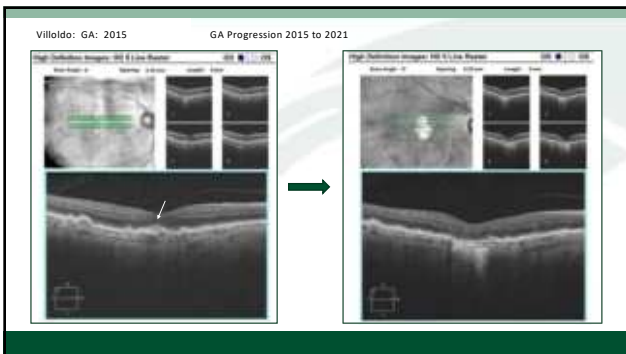
Hyper-reflective Columns

- Narrow strips of light transmission
- Overlying RPE appears intact
 - May represent micro-cracks
- Increased risk of progression to GA
 - Present in 27% of eyes that progressed to GA nAMD

111



112



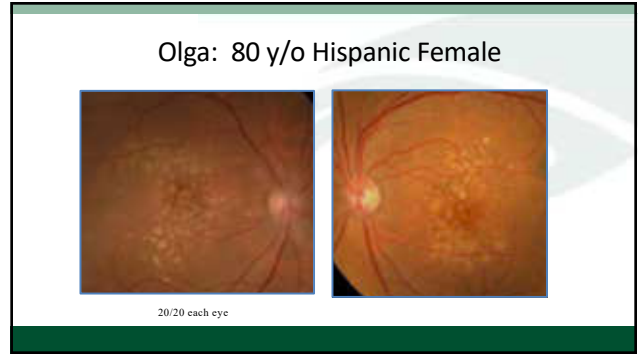
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114



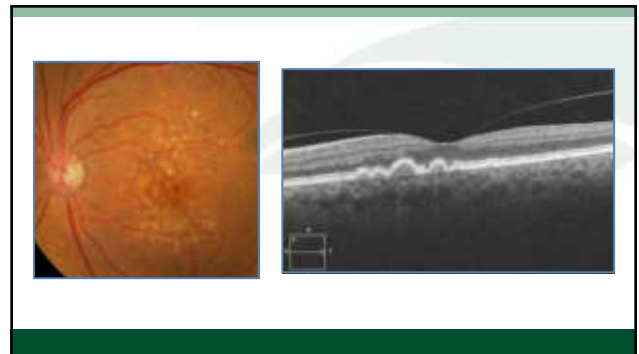
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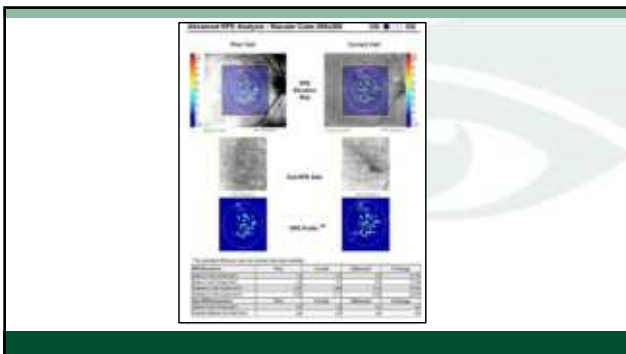
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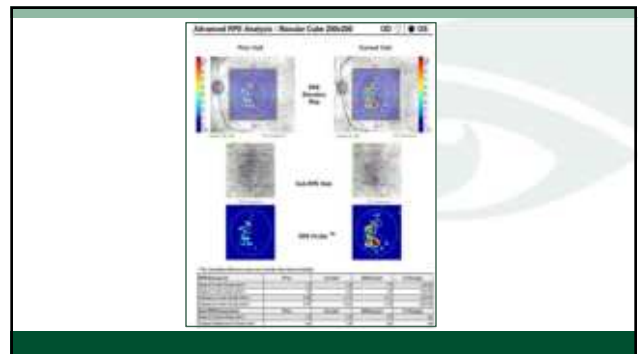
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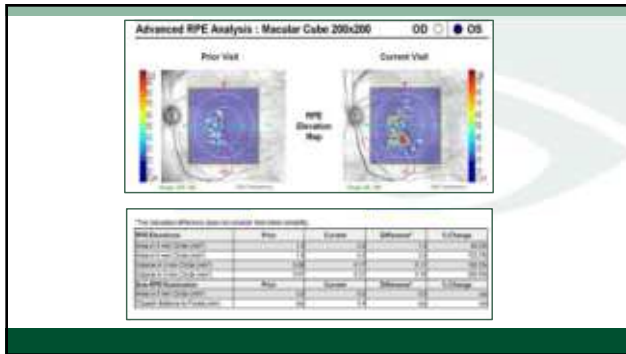
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119



120



121

Table 3. Progressive Macular Changes in AMD

Observation	Imaging Findings	Mechanism	Prevalence in AMD	Expected Progression (OR)
Drusen volume	Drusen volume increase	Displacement or deterioration of photoreceptor layer	88% ¹	1.21 risk of progression to advanced AMD for each 0.1 mm ³ of drusen volume increase [1,2]
RPE thickness complex (RC) Advanced analysis	RC ²	RPE swelling and drusen expansion	88% ¹	1.02 risk of developing central LA for each 0.5% increase in RC thickness [1,3]
AMD	Progressive hyperreflective lesions	Advanced migration of fully pigmented RPE cells, differentiation into elongated cell sub-population	80% in AMD	8 risk of 2 year progression to LA [1,2]
AMD	Small yellow deposits outside of the macula or in peripapillary region	Photoreceptor or choroidal macrophage (foveal) phagocytosis or choroidal lesions [1,2]	70% in 70% in AMD patients	2.26-3.8 risk of progression to advanced AMD [1,2]
AMD	Thickness of the GCL ⁴ and IPL ⁵ with a normal drusen average	Thinning of outer (outer depth)	7% in nonexudative AMD [1,2]	1.12 risk of progression to central LA [1,2]
Hyporeflective lesions	Collection of lipid or hyperreflective RPE layer	Indicates early RPE layer	37% in AMD patients [4]	AMD
AMD	Internal heterogeneity	Molecular instability	8% in soft drusen	1.9 risk of progression to non-dry AMD [1,2]
Non-exudative bilateral macular atrophy	Non-exudative lesion with no drusen	Photoreceptor non-attachment against retinal	4.2% in 17% in the follow-up of macular AMD [1,2]	1.22 risk of progression to advanced AMD at 2 year [1,2]

¹ Chikrii et al; ² not measured; ³ RPE thickness; ⁴ Outer Photoreceptor Layer; ⁵ Inner-outer layer

122

Double Layer Sign

- Shallow, irregular retinal pigment epithelium (RPE) elevation, or **SIRE**
 - Stemmed from the double-layer sign initially described in polypoidal choroidal vasculopathy.
- When the RPE is elevated due to (macular neovascularization), the underlying hyperreflective Bruch's membrane becomes visible
 - Creates 2 hyperreflective lines or a double-layer sign
- In patients with nonexudative AMD - SIRE may predict that 1 in 4 of these patients have an underlying MNV that is not yet exudative
- Features of SIRE include:
 - length of more than 1000 μm
 - RPE elevation < less than 100 μm (resulting in shallow morphologic features),
 - Irregular overlying RPE layer,
 - Nonhomogenous reflectivity

123

124

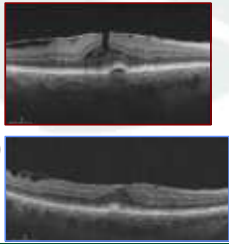
125

Pay Attention to the Vitreoretinal Interface

126

5 Diseases Arising from the Vitreomacular Interface (VMI):

- Vitreomacular traction (VMT)
- Full Thickness macular hole (FTMH)
- Lamellar macular hole
- Epiretinal membrane (ERM)
- Myopic tractional maculopathy (MTM)
 - Myopic macular schisis



127

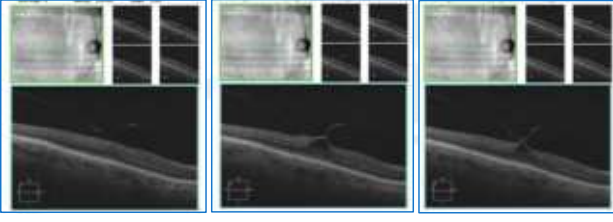
10/30/2018 20/40



Harrison

128

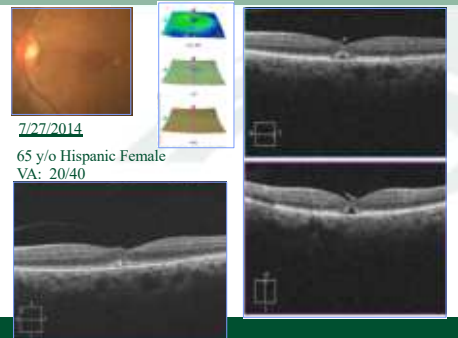
12/21/21 PCO -> Yag -> 20/30+



(Harrison 0341614)

129

7/27/2014
65 y/o Hispanic Female
VA: 20/40



130

9 weeks Later
Worse!
20/50

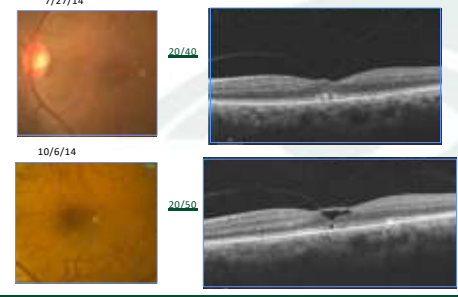


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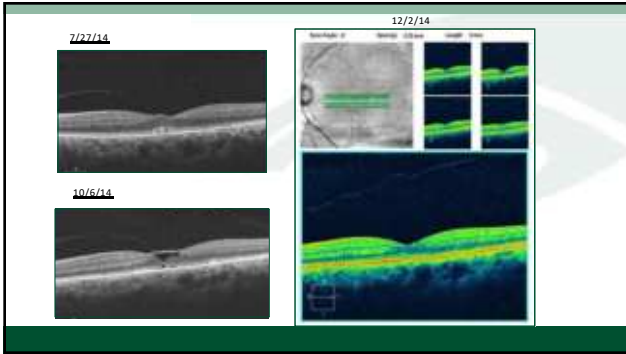
VMT evolving into a Macular Hole

7/27/14 20/40

10/6/14 20/50



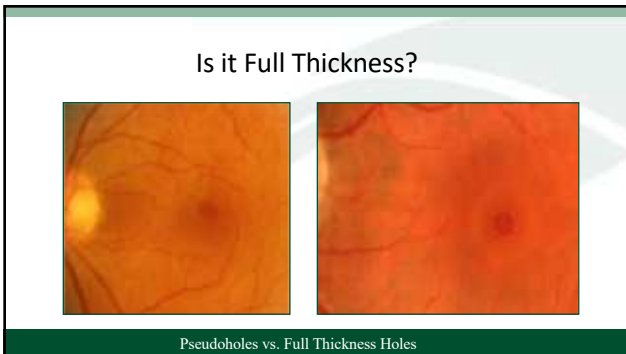
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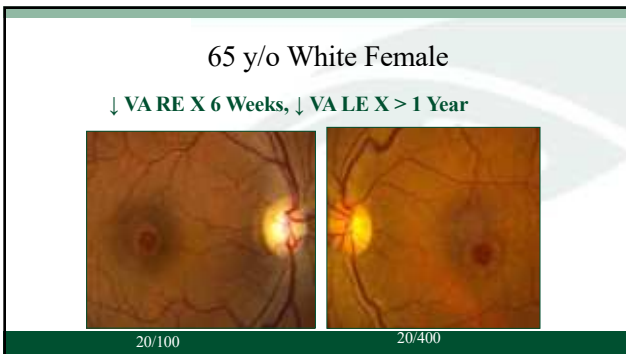
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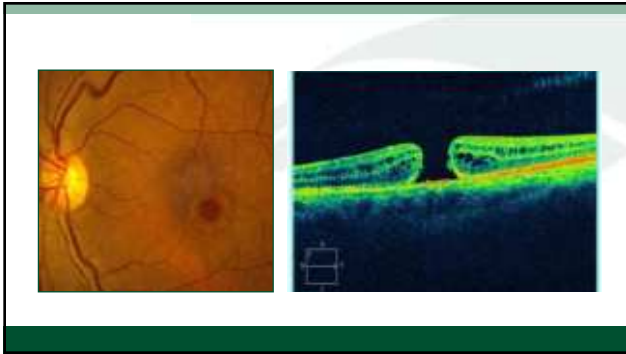
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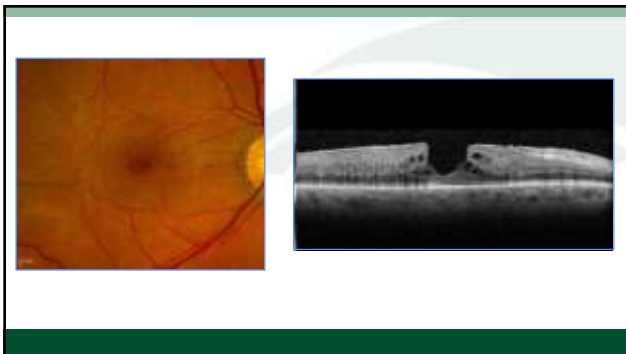
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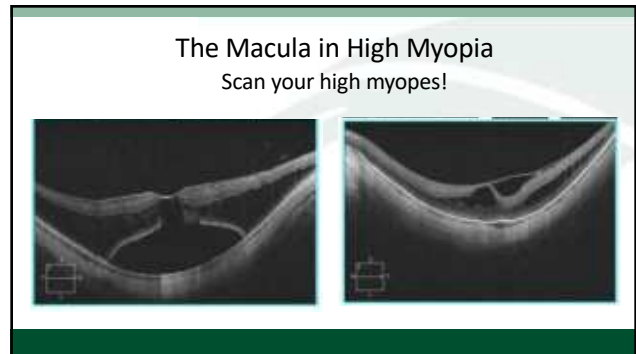
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140



141



142

Myopic Macular Retinoschisis
Myopic Tractional Maculopathy

- ◆ Seen in 9% of highly myopic eyes with posterior staphyloma
- ◆ 50% progress to macular hole formation or macular detachment within 2 years
- ◆ **Caused by rigidity of ILM that induces traction**

143

Jeff: mid-50's Attorney, High Myopia
Hx of RD Repair in both eyes: RE: 1985 LE 1989

- Never recovers vision in the RE
- He is followed through the 90's with a progressive NS and declining Va ~ 20/70
 - 1 eyed patient and reluctant to have CE
- Eventually has CE/IOL 90's-early 2000's and does well
 - VA 20/25 low refractive error

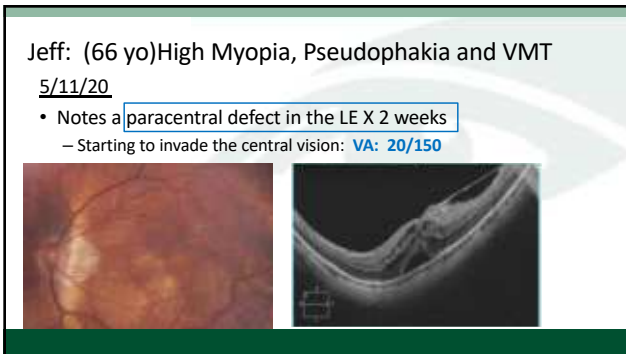
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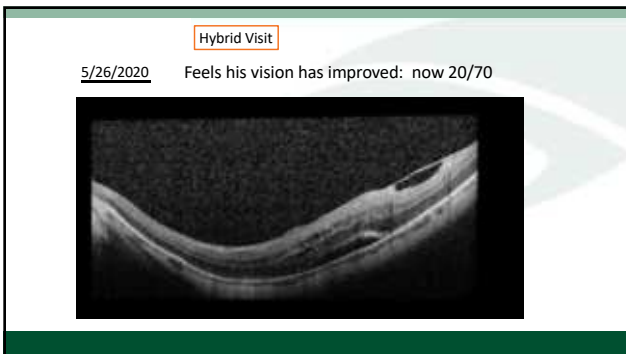
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148



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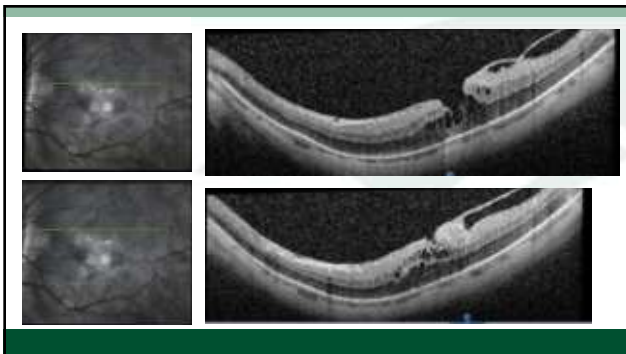
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151



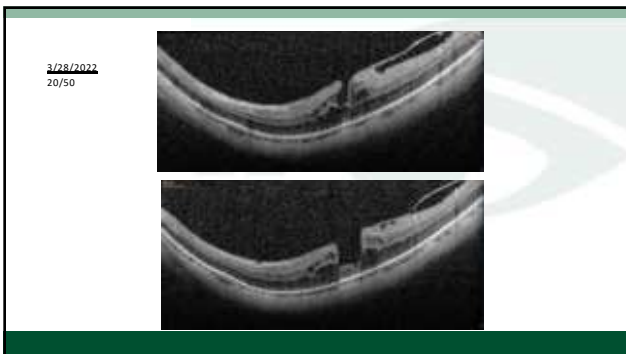
152



153



154



155



156

What is going to happen?
 Will he progress to macular hole
 Would he benefit from a vitrectomy?

157

Summary: OCT in Retina

- SD OCT has emerged as a critical tool in the diagnosis and treatment of retinal disease
- It has changed how we evaluate the macula
- Helps establish a diagnosis that is difficult to determine with only standard ophthalmoscopy
- Advancing software has provided expanded uses OCT
- OCT Angiography has taken OCT to the next level

158

Summary OCT in Glaucoma

- OCT provides another piece information for the “glaucoma puzzle”
 - Along with IOP, visual fields and clinical appearance of the nerve
- It provides an objective means of comparing “glaucomatous” nerves from normal or physiologic optic nerve
- It provides an objectives means of determining progression

159