



**Marshall B.
KETCHUM UNIVERSITY**
Southern California College of Optometry

Joint SCCO | USC | VA Symposium

Live Interactive CE Webinar | AM Session

Sunday | September 19, 2021 | 8:00 a.m. - 11:50 a.m.



Joint SCCO | VA | USC Symposium



**Marshall B.
KETCHUM UNIVERSITY**
Southern California College of Optometry
Department of Continuing Education

Sunday, September 19

Pacific Time Zone | Live Webinar | Pending COPE-Approval

Morning Session

8:00 a.m. - 8:55 a.m.

Keratoconus in Youth: An Urgent Issue?

Gloria Chiu, OD

8:55 a.m. - 9:50 a.m.

Updates in Clinical Glaucoma Management

Brian Song, MD

10:00 a.m. - 11:50 a.m.

Refer or Relax? Macula

Steven Ferrucci, OD

11:50 a.m. - 12:10 p.m.

Break

Afternoon Session

12:10 p.m. - 1:05 p.m.

Optic Disc Edema

Jessica Chang, MD

1:05 p.m. - 2:00 p.m.

Herpetic Eye Disease

Brian Toy, MD

2:10 p.m. - 3:05 p.m.

The Calm in the Eye of the Storm: Re-Purposed Medications for COVID-19

Judy Tong, OD

3:05 p.m. - 4:00 p.m.

IPC: A Case for Collaboration

John Nishimoto, OD and Julie Tyler, OD

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Instructor Biographies



**Marshall B.
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Southern California College of Optometry
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Gloria Chiu, OD

Associate Professor of Clinical Ophthalmology, USC Roski Eye Institute

Dr. Gloria Chiu completed her Bachelor of Arts from the University of California, Berkeley. She remained at the University of California, Berkeley, where she obtained her Doctor of Optometry degree in 2008. Following completion of her residency in Cornea and Contact Lenses at Southern California College of Optometry, Dr. Chiu pursued further fellowship training in Prosthetic Replacement of the Ocular Surface Ecosystem (PROSE) treatment at the Boston Foundation for Sight. Dr. Chiu developed and supervises the USC PROSE service and is actively conducting research in the areas of irregular corneas and ocular surface disease.

Brian Song, MD

Assistant Professor Of Clinical Ophthalmology & Director of Education, USC Department of Ophthalmology

Dr. Brian J. Song is Assistant Professor of Clinical Ophthalmology and Director of Education in the Department of Ophthalmology at the USC Keck School of Medicine. He received his undergraduate degree from Johns Hopkins University and his medical degree from the University of Texas Medical Branch. He then completed his ophthalmology residency at the Harkness Eye Institute of Columbia University Medical Center – New York Presbyterian Hospital followed by a glaucoma fellowship at the UCLA Stein Eye Institute. His current research interests include ophthalmic ultrasound and imaging methods to evaluate optic nerve biomechanics and blood flow abnormalities in glaucoma.

Steven Ferrucci, OD

Chief, Optometry Section at Sepulveda VA Ambulatory Care Center
Professor, MBKU | SCCO

Dr. Steven Ferrucci, a 1994 graduate of the New England College of Optometry, completed his Residency in Primary Care/Hospital Based/Geriatric Optometry at the Sepulveda VA Hospital in Sepulveda CA. He is currently Chief of Optometry at the Sepulveda VA Ambulatory Care Center and Nursing Home. He is also the Residency Director at his sight, and a Professor at the Southern California College of Optometry at Marshall B. Ketchum University. Dr. Ferrucci has lectured extensively, with a special interest in Diabetes, Diabetic Eye Disease, Age-Related Macular Degeneration, and Fluorescein Angiography. He has also published several articles in optometric journals, including The New England Journal of Optometry, Optometry and Vision Science, Optometry: Journal of The AOA and Review of Optometry. Currently, he serves on the Editorial Board for both Review of Optometry and Optometry Times. He is an active member in the American Optometric Association and the California Optometric Association, as well as a fellow in both the American Academy of Optometry and the Optometric Retinal Society.

Jessica Chang, MD

Clinical Assistant Professor of Ophthalmology, USC Roski Eye Institute

After undergraduate studies at Yale University, Dr. Jessica Chang spent a year doing volunteer work in China and then attended Duke University School of Medicine. As a medical student, she was selected as a Howard Hughes Medical Institute NIH Research Scholar and spent two years at the National Eye Institute doing basic and clinical research in ophthalmology. She then completed ophthalmology residency at the Wilmer Eye Institute, followed by American Society of Ophthalmic Plastic and Reconstructive Surgery fellowship training in Oculoplastics and Neuro-ophthalmology.

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Instructor Biographies



**Marshall B.
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Brian Toy, MD

Assistant Professor of Clinical Ophthalmology, Director of Clinical Informatics and Information Technology
Service Chief of the Uveitis and Ocular Inflammation Service, USC Roski Eye Institute

Dr. Brian Toy is a vitreoretinal fellow and clinical instructor at the USC Roski Eye Institute. He graduated magna cum laude from the University of California, Berkeley, with a degree in bioengineering, received an MD with distinction from the University of California, San Francisco, and completed a clinical research fellowship at the National Institutes of Health. He completed an internship at Santa Clara Valley Medical Center, a major county healthcare system in the Bay Area, and then completed an ophthalmology residency at Stanford University. Toy's professional interests include retinal imaging, telemedicine and safety net care, particularly as applied to diabetic retinopathy. Outside of work, he enjoys hiking, skiing, and spending time with friends and family.

Judy Tong, OD

Associate Professor & Assistant Dean of Residencies | MBKU | SCCO

Dr. Judy Tong is an Associate Professor of Optometry and Assistant Dean of Residencies at the Southern California College of Optometry of the Marshall B. Ketchum University. She received her BS degree in Genetics from the University of California, Berkeley and her OD degree from the Southern California College of Optometry. She completed a one-year residency in Primary Care Optometry at the Eye Institute of the Pennsylvania College of Optometry (Salus University). As the Assistant Dean of Residencies since November 2003, Dr. Tong serves to provide global administrative and educational direction to 24 residency programs across 6 different states. Her main academic responsibility is teaching anterior segment diseases, basic and advanced procedures including lasers, injection, and suturing. Dr. Tong is one of the core instructors of the glaucoma certification courses and grand rounds program in California. Dr. Tong's research activities include being the Principal Investigator and Co-Investigator on two phase III antibiotic drug trials and major allergy study.

John Nishimoto, OD, MBA

Professor & Senior Associate Dean for Professional Affairs, MBKU | SCCO

Dr. John Nishimoto received a Doctor of Optometry degree from the Southern California College of Optometry in 1987. In 1988, he completed a one-year residency in Hospital-Based Geriatric Optometry at the West Los Angeles VA Medical Center. Dr. Nishimoto is currently a Professor and the Senior Associate Dean for Professional Affairs. In 1997, he received a Health Care Executive Masters in Business Administration from the University of California, Irvine. Dr. Nishimoto has been a frequent contributor of articles and lectured on topics especially related to primary care and ocular disease. He is the co-author of the text "Differential Diagnosis in Primary Eye Care." Dr. Nishimoto is also currently a clinical faculty member in primary care and ocular disease at the University Eye Center at Ketchum Health. Dr. Nishimoto is a fellow of the American Academy of Optometry and served as Chair of the Section on Ocular Disease.

Julie Tyler, OD

Primary Care Department Chair & Associate Professor, MBKU | SCCO

Dr. Julie A. Tyler received her B.A. from Creighton University and her Doctor of Optometry Degree from Indiana University School of Optometry (IU). Following graduation, Dr. Tyler completed a Residency at Nova Southeastern University (NSU) and served as Chief Resident prior to joining the faculty at NSU full-time. More recently, Dr. Tyler joined the faculty at Southern California College of Optometry (SCCO) at MBKU. Dr. Tyler has served in a variety of clinical and academic roles including chief of service and instructor of record for various clinical and didactic coursework. She has been promoted to Associate Professor and received numerous teaching awards, as well as recognition as a faculty member of Gold Key Honor Society and in 2019 was inducted into Phi Kappa Phi honor society that recognizes individuals in all academic disciplines. Dr. Tyler has authored posters and published journal articles on a variety of topics in the areas of primary care and ocular disease and is a Fellow of the American Academy of Optometry and is also a member of the COA and AOA.

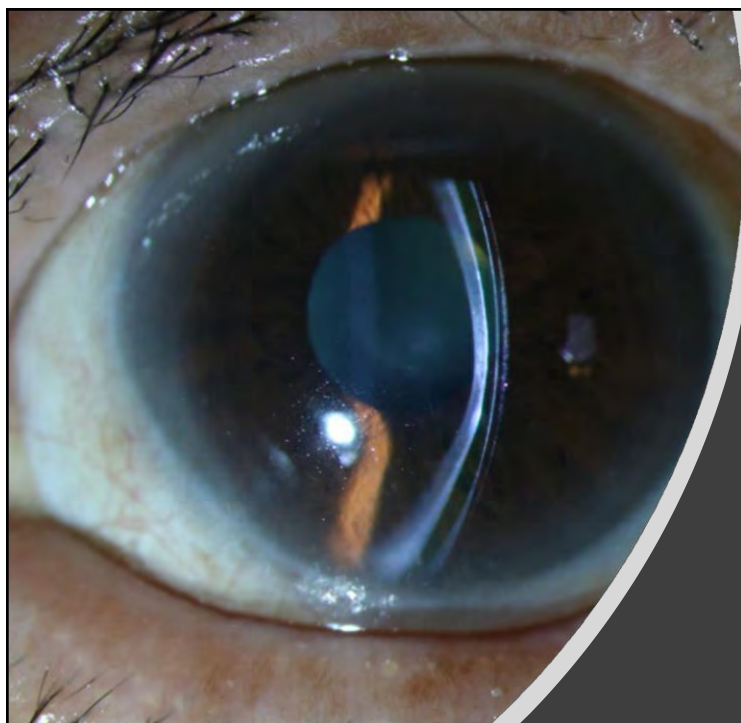


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Keratoconus in Youth: An Urgent Issue?

Gloria Chiu, OD





Keratoconus in Youth: An Urgent Issue?

Southern California College of Optometry at MBKU
USC Joint Symposium
September 19, 2021

Gloria B Chiu, OD, FAAO, FSLs
Associate Professor of Clinical Ophthalmology
USC Roski Eye Institute, Dept of Ophthalmology
Keck Medicine of USC
Adjunct Faculty at Southern California College of Optometry

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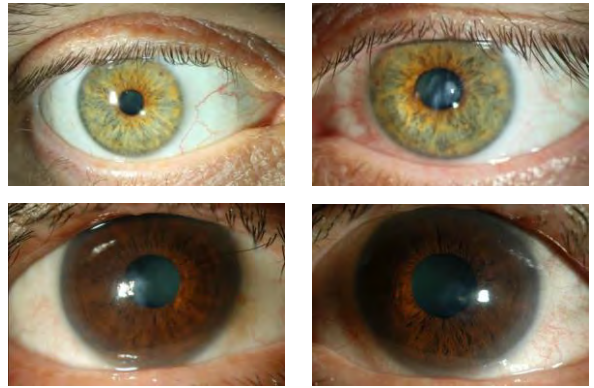
Financial Disclosures

- Avedro/Glaukos – Consultant
- Evolve Medical Education – Speaker
- Acculens – Received Honorarium

2

Keratoconus Overview

- Corneal Disorder with central thinning and steepening
 - Greek words: Kerato (cornea) and Conos (cone)
 - Bilateral
 - Asymmetric
 - Progressive
 - Non-inflammatory
- Onset
 - Teens/puberty
- Prevalence and Incidence
 - Varies with geography, ethnicity, study
 - Diagnosis often from ODs

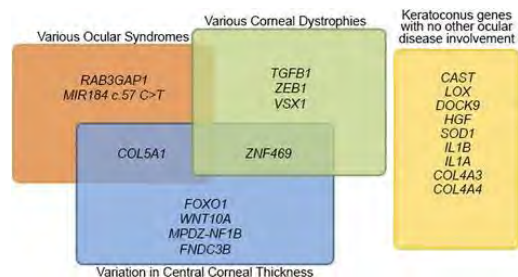


The Open Ophthalmology Journal: Epidemiology of Keratoconus Worldwide,
<https://openophthalmologyjournal.com/VOLUME/12/PAGE/289/FULLTEXT/>

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Etiology

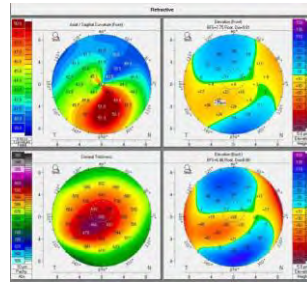
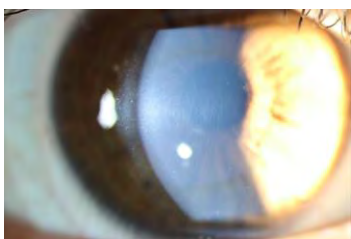
- Genetic
 - Familial Inheritance
 - Autosomal Dominant
 - Autosomal Recessive
- Environment
 - UV exposure
 - Atopic conditions
 - Contact Lens wear
- Risk Factors
 - Family History
 - Eye rubbing
 - Age
- Genetic Testing
 - Universal Genetic Test
 - AvaGen test



[Y Bykhovskaya](#), [B Margines](#), [Y Rabinowitz](#) Genetics in Keratoconus: where are we? [Eye Vis \(Lond\)](#). 2016; 3: 16.

4

Keratoconus Diagnosis Made

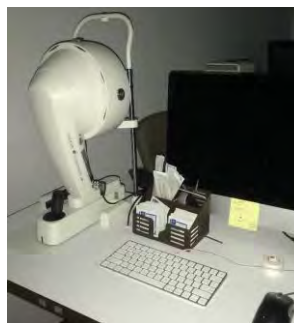


Al-Mahrouqi H, Oraba SB, Al-Habsi S, Mundemkattil N, Babu J, Panchatcharam SM, Al-Saidi R, Al-Raisi A. **Retinoscopy** as a Screening Tool for **Keratoconus**. *Cornea*. 2019 Apr;38(4):442-445.

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Helpful Equipment

- Corneal Topographer
 - 2-D Imaging
 - Placido disc based
 - Maps surface curvature
- Scheimpflug Tomographer
 - 3-D Imaging
 - Maps front and back corneal surface, thickness
- Anterior Segment Optical Coherence Tomography (AS-OCT)
 - Described in 1994 (Izatt JA)
 - initially for retina
 - High resolution cross-sectional imaging
 - Many applications



Izatt JA, Hee MR, Swanson EA, et al. Micrometer-scale resolution imaging of the anterior eye in vivo with optical coherence tomography. *Arch Ophthalmol*. 1994;112:1584-1589

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Questions for thought

- Do you screen for keratoconus in every patient?
- Should we screen for KC in every patient?
- At what age should we start screening for KC?

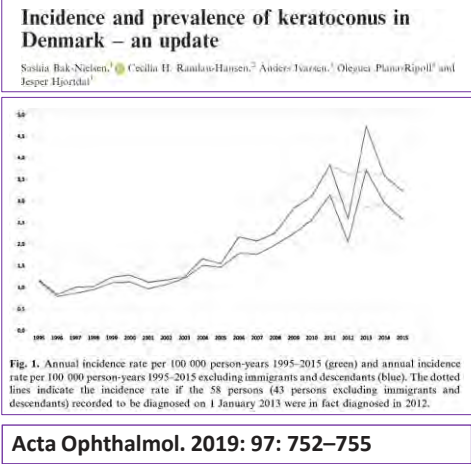


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Reported Incidence & Prevalence of keratoconus

Reference	Prevalence	Geography
Kennedy et al. 1986	0.05% or 1:2000	US
Jonas et al. 2009	2.3%	India
Millodot et al. 2011	2.3%	Israel
Xu et al. 2012	0.9%	China
Hashemi et al. 2014	2.5%	Iran
Godefrooij et al. 2017	0.26% or 1:375	Netherlands
Torres Netto et al. 2018	4.79%	Saudi Arabia
Chan et al. 2020	1.2% or 1:84	Australia
Hashemi et al. 2020*	0.14% or 1:700	Global Meta-Analysis

*Hashemi H, Heydarian S, Hooshmand E, et al. The Prevalence and Risk Factors for Keratoconus: A Systematic Review and Meta-Analysis. *Cornea*. 2020;39(2):263-270



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Keratoconus Progression

- Study: Ferdi A, Nguyen V, Kandel H, Tan JCK, Arnalich-Montiel F, Abbondanza M, Watson S. **Predictors of progression in untreated keratoconus: a Save Sight Keratoconus Registry study.** Br J Ophthalmol. 2021 Mar 30
 - Aim to identify risk factors for progression in untreated KC pts
 - 34 centers, across Australia, New Zealand, Spain and Italy
- Steeper Max-K and younger age were the most clinically useful baseline predictors of progression
 - Every 1D steeper Max-K was associated with a 7% and 3% greater risk of worsening VA and thinning TCT, respectively
 - Each 1 year younger was associated with a 4% and 2% greater risk of steepening Max-K and thinning TCT, respectively.

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Study: A Systematic Review and Meta-analysis of 11 529 Eyes Study

- Ferdi AC, Nguyen V, Gore DM, Allan BD, Rozema JJ, Watson SL. **Keratoconus Natural Progression: A Systematic Review and Meta-analysis of 11 529 Eyes.** Ophthalmology. 2019 Jul;126(7):935-945. Epub 2019 Mar 8.
 - 41 publications in systematic review
 - 23 in meta-analysis
- Younger patients and those with K_{max} steeper than 55 D at presentation have a greater risk of progression
- **Closer follow-up and a lower threshold for cross-linking should be adopted in patients younger than 17 years and steeper than 55 D Kmax.**

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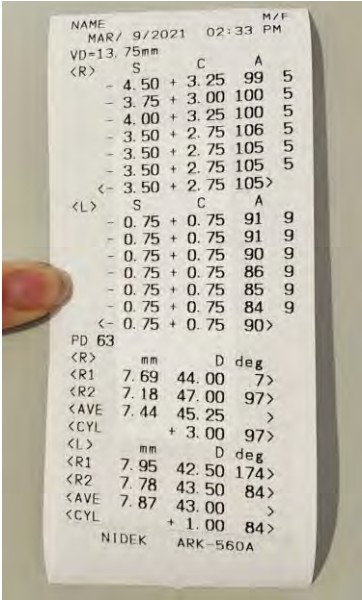
Keratoconus Progression

- Study: Ferdi A, Nguyen V, Kandel H, Tan JCK, Arnalich-Montiel F, Abbondanza M, Watson S. **Keratoconus after 40 years of age: a longitudinal comparative population-based study.** Int Ophthalmol. 2020 Mar;40(3):583-589.
 - Determine 5-yr changes in KC indices in 40-64 y/o compared to normal subjects
 - Posterior corneal steepening and thinning in keratoconus patients continue after the age of 40 years, but clinically negligible
 - Changes are independent of normal age-related changes and appear to be slower in cases with steeper and thinner corneas

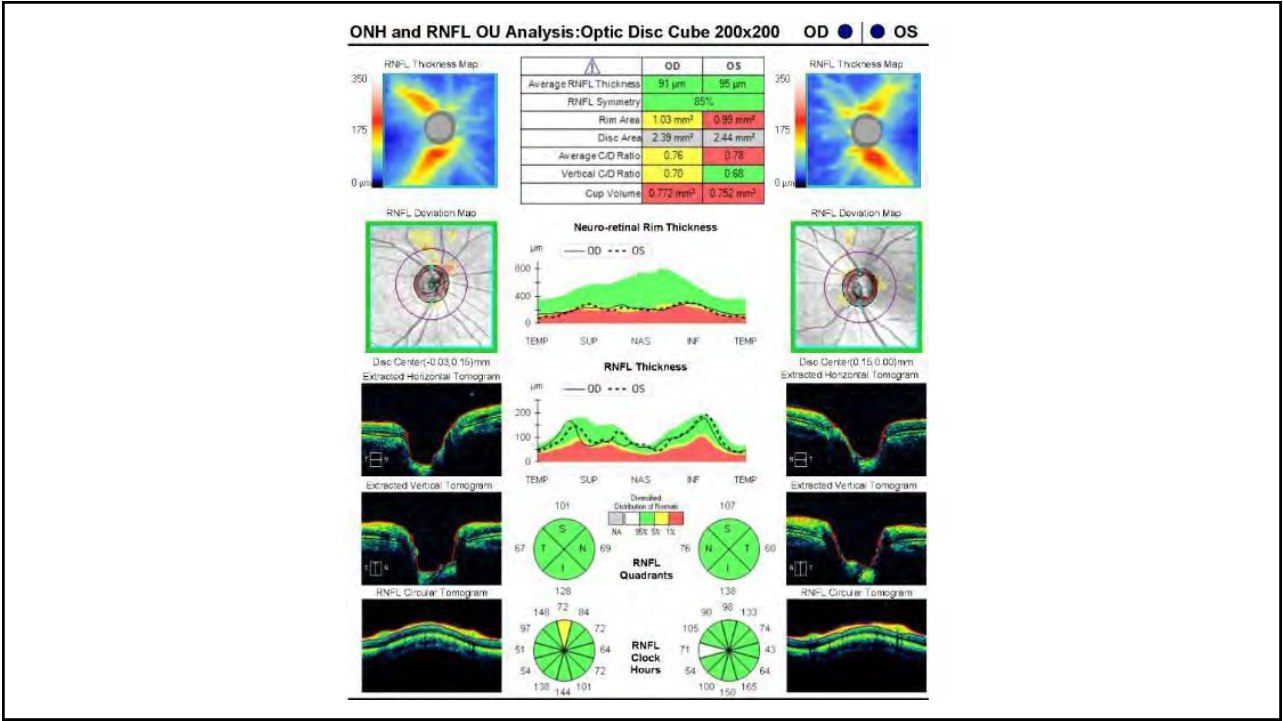
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Case Example

- 31 y/o male
- Referred for CLs
- sc VA 20/30+ OD and 20/20 OS
- Final SRx:
 - -0.50 +1.50 x 150 20/20 OD
 - Plano 20/20 OS

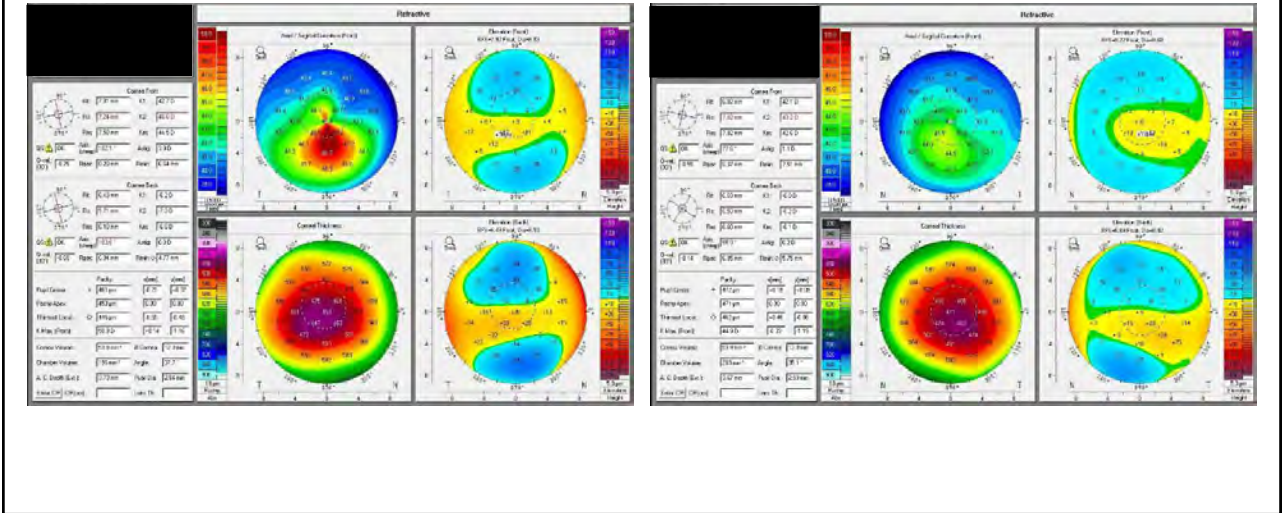


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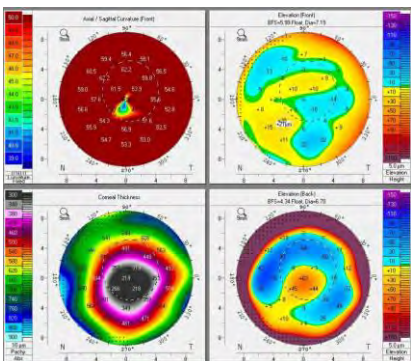
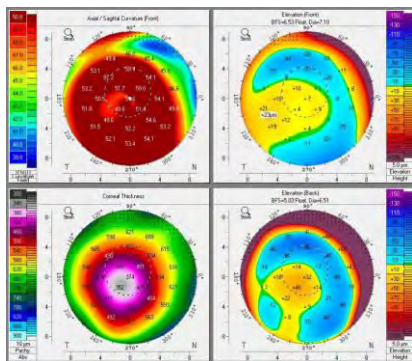
Tomography Images – Mild Keratoconus



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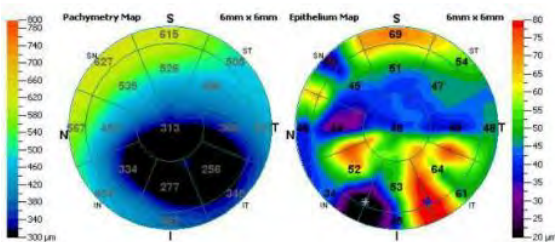
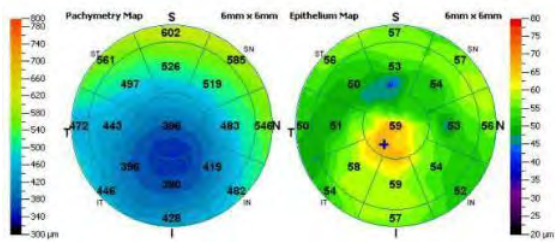
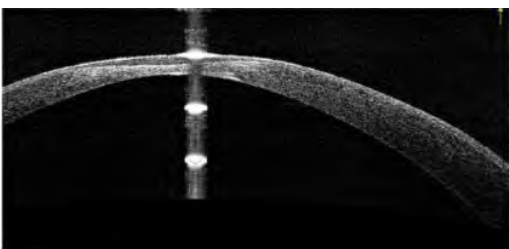
Case Example: KC Complications

- 26 y/o F
 - FHx: KC in sister
 - Hx Hybrid CL wear, switched to scleral lenses
 - s/p CXL off-label epi-on 2014 with outside MD
- 3-2019: OD: 54.6/52.1 D, 349 CCT OS: 55.3/52.8 D, **193 CCT**
- 4-2020: Urgent call, White scar upon waking OS: Hydrops



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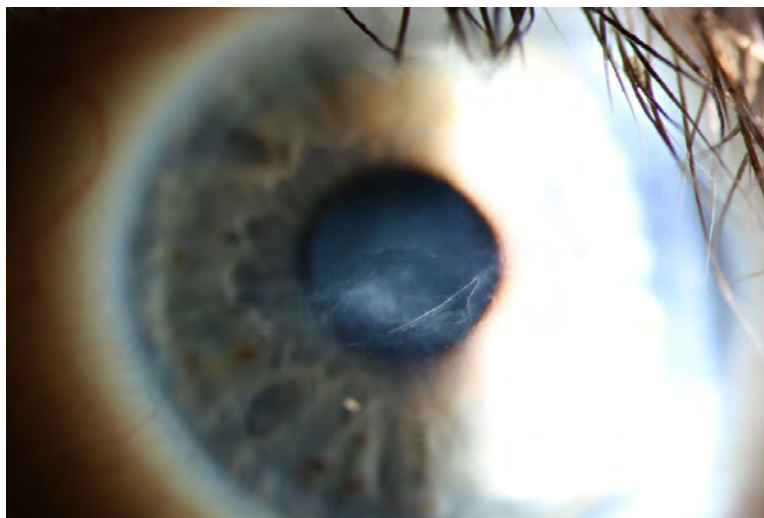
AS-OCT Images



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After Corneal Hydrops

BCVA 20/50-



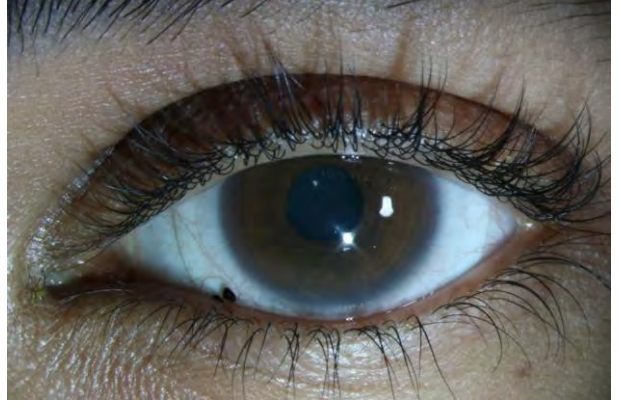
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Case Example: Advanced KC, complications

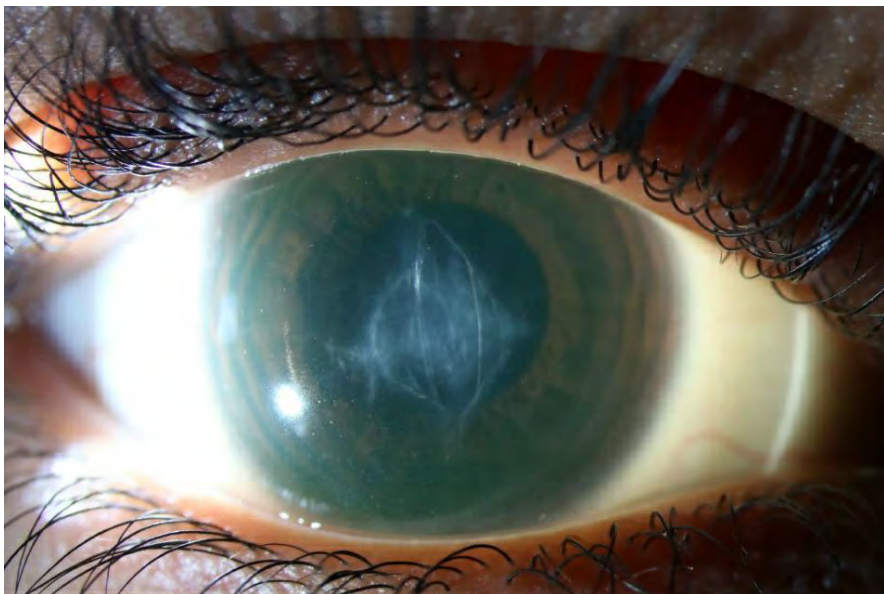
- 25 y/o F, referred for scleral lenses OU
 - Ocular itching and admits to eye rubbing
- KC OD>OS
- CXL OS 2017
- Hydrops OD 2018
 - BCVA 20/50-60
- Does not want to have surgery

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After Corneal Hydrops



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Case Example

- 15 y/o Caucasian male
- Incoming sc VA
 - 20/300 OD
 - 20/100 OS
- Tomography Ks
 - 60.4/53.3 OD
 - 59.0/51.4 OS
- SRx BCVA
 - -1.25 +0.50 x 080 20/125 OD
 - -1.25 +1.00 x 135 20/60 OS
- Central Striae and mild scars OU
- Send immediately for CXL

NAME
APR/21/2021 07:27 AM M/F
VD=13.75mm

<L>	S	C	A	
	-26.00	+13.50	85	5
	-25.25	+13.00	84	5
	-25.25	+13.75	83	5
	-25.25	+13.75	81	5
	-25.50	+13.75	81	5
	-25.50	+13.75	86	5
	<-25.50	+13.75	84>	

PD 54

<R>	mm	D	deg
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<R2	5.19	65.00	103>
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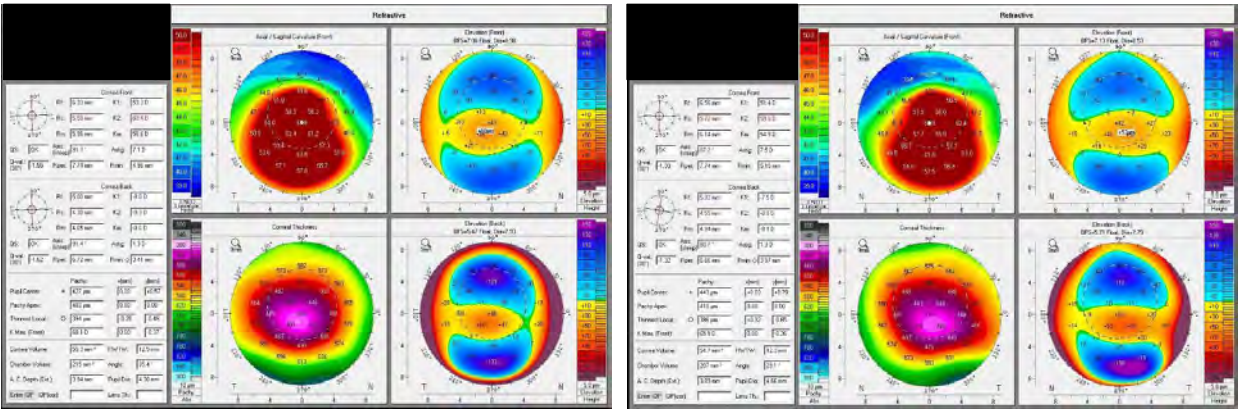
NIDEK ARK-560A

1.Ferdi AC, Nguyen V, Gore DM, Allan BD, Rozema JJ, Watson SL.Ophthalmology. **Keratoconus Natural Progression: A Systematic Review and Meta-analysis of 11 529 Eyes.** 2019 Jul;126(7):935-945. doi: 10.1016/j.ophtha.2019.02.029. Epub 2019 Mar 8.

2.Shah H, Pagano L, Vakharia A, Coco G, Gadhvi KA, Kaye SB, Romano V. **Impact of COVID-19 on keratoconus patients waiting for corneal cross linking.** Eur J Ophthalmol. 2021 Mar 15:112067212111001315.

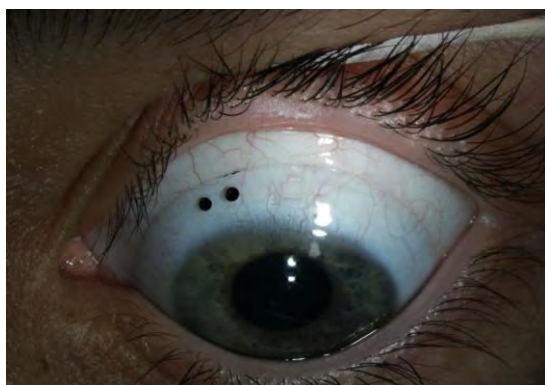
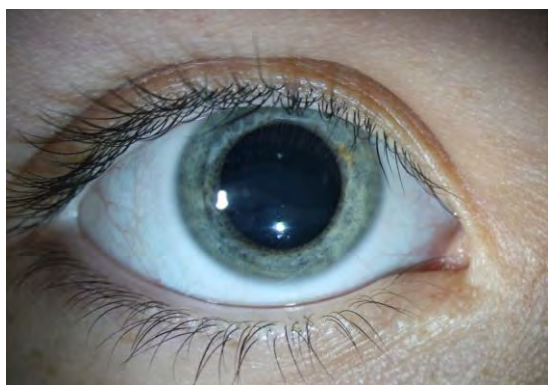
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Tomography Images



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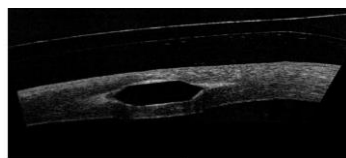
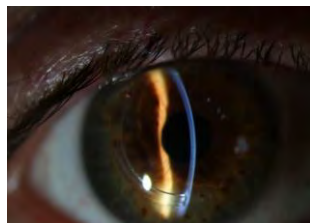
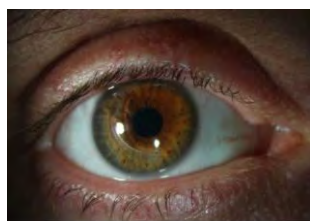
Fit with Scleral Lenses, 20/20 OD and OS



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Keratoconus Procedures to Help Improve Vision

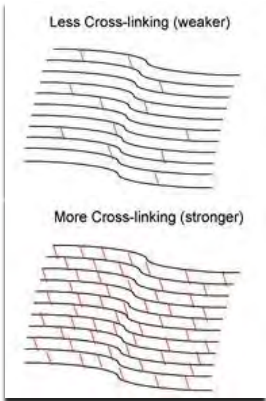
- Corneal Intacs
 - FDA approved for KC in 2004
 - “Flattens” cornea
- Topography Guided PRK (TG-PRK)
 - “Smooths” cornea, touch-up
- Corneal Transplantation
 - Replaces cornea



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Corneal Cross-linking (CXL) to stabilize and slow progression

- FDA approved in April 2016
 - KXL UV System, with Photrex and Photrex Viscous (riboflavin)
- Indications:
 - Progressive Keratoconus
 - Post-LASIK Ectasia
- Procedure involves:
 - Epithelium removal (Epi-off)
 - 30 min riboflavin application
 - Intraoperative Corneal thickness minimum: **400 microns**
 - 30 min exposure 365 nm UV-A light, 3.0 mW/cm2
- Activated riboflavin and reactive oxygen species interact in cornea to form crosslinks: stiffens cornea ¹
 - 328.9% increase in biomechanical rigidity ²

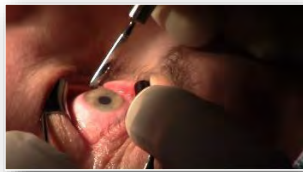


1. Beshtawi IM, O'Donnell C, Radhakrishnan H. Biomechanical properties of corneal tissue after ultraviolet-A-riboflavin crosslinking. J Cataract Refract Surg. 2013;39(3):451-62

2. Wollensak G. Crosslinking treatment of progressive kera-toconus: New hope. Curr Opin Ophthalmol. 2006;17:356-60.

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Cross-Linking Procedure Summary*



1. Remove epithelium.



2. Soak cornea with Photrex® Viscous (riboflavin 5'-phosphate in 20% dextran ophthalmic solution).

30 minutes

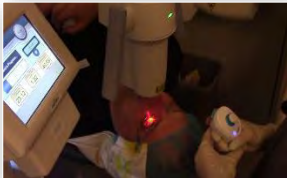


3. Check for flare.



4. Once flare is observed, measure corneal thickness.

If corneal thickness is less than 400 µm, instill 2 drops of Photrex (riboflavin 5'-phosphate in ophthalmic solution) until the corneal thickness increases to at least 400 µm.



5. Irradiate for 30 minutes.

Continue applying Photrex Viscous (riboflavin 5'-phosphate in 20% dextran ophthalmic solution) during irradiation.

Procedural images courtesy of Dr. Raj Rajpal

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Corneal Cross-linking (CXL)

- KXL System
 - Photrexa Viscous (riboflavin 5'-phosphate in 20% dextran solution)
 - Photrexa (riboflavin 5'-phosphate ophthalmic solution, hypotonic; swelling effect)
 - KXL UV light delivery system
- No other riboflavin solutions or UV devices can be used in the U.S. outside of a formal IDE (device) or IND (drug) study
- No specific age range limitations
 - Patients ages 14-65, included in FDA studies
- Not advised during pregnancy

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CXL Procedure Room



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CXL Expectations

- We need to **educate** our patients
 - Aim to slow or stop progression
 - NOT refractive surgery
 - Will not remove scarring
 - Treat DES to enhance healing
 - May still need visual correction
- Recovery Period
 - Bandage Contact Lens
 - May be discomfort
 - Do not rub eyes
 - Call if sudden pain or VA decline



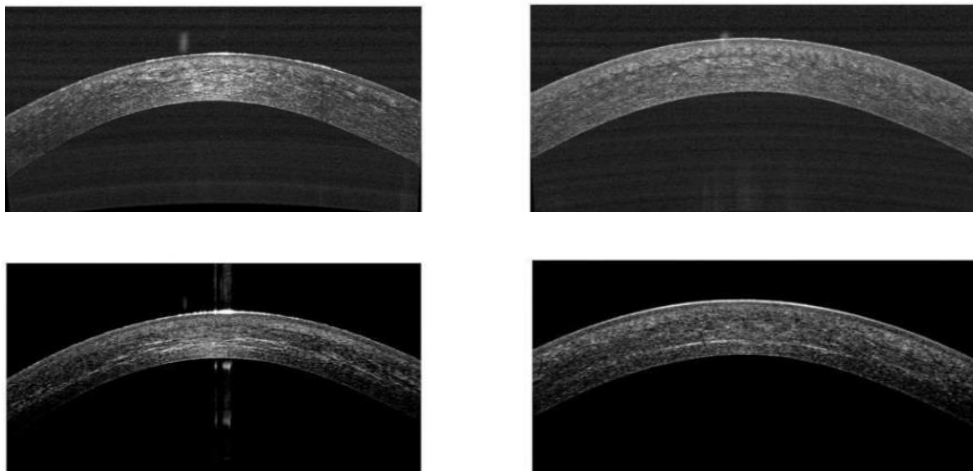
29

CXL Post-operative Considerations

- After procedure
 - Topical Antibiotic, Steroid (NSAID)
 - Lubrication
 - Placement of bandage SCL – No eye rubbing!
- Week 1:
 - Topical meds, lubrication
 - Remove bandage SCL once epithelium healed
- Month 1:
 - Assess vision
 - Corneal Imaging – stromal remodeling
 - Consider CL fitting
- Months 3, 6, 12:
 - Assess vision – MR and BCVA often change
 - Corneal Imaging
- **Zero Global Period: Visits can be billed to insurance**

30

AS-OCT before CXL and 2 mo after: Corneal Remodeling – Demarcation Line



[Spadea L, Tonti E, Vingolo EM. Corneal stromal demarcation line after collagen cross-linking in corneal ectatic diseases: a review of the literature. Clin Ophthalmol. 2016 Sep 19;10:1803-1810](#)

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Efficacy of CXL Supported by Literature

- [Riboflavin/ultraviolet-a-induced collagen crosslinking for the treatment of keratoconus.](#) Wollensak G, Spoerl E, Seiler T. Am J Ophthalmol. **2003** May;135(5):620-7. doi: 10.1016/s0002-9394(02)02220-1.
 - **Conclusions:** Collagen crosslinking may be a **new way** for stopping the progression of keratectasia in patients with keratoconus. The need for penetrating keratoplasty might then be significantly reduced in keratoconus. Long-term results are necessary to evaluate the duration of the stiffening effect and to exclude long term side-effects.
- [Corneal Collagen Cross-Linking for Keratoconus: Systematic Review.](#) Kobashi H, Rong SS. Biomed Res Int. 2017;2017:8145651. doi: 10.1155/2017/8145651. Epub 2017 Jun 11.
- [Corneal collagen crosslinking with riboflavin and ultraviolet-A light in progressive keratoconus: ten-year results.](#) Raiskup F, Theuring A, Pillunat LE, Spoerl E. J Cataract Refract Surg. 2015 Jan;41(1):41-6. doi: 10.1016/j.jcrs.2014.09.033.
- [Long-term results of cornea collagen cross-linking with riboflavin for keratoconus.](#) Agrawal V. Indian J Ophthalmol. 2013 Aug;61(8):433-4. doi: 10.4103/0301-4738.116072.

32

Epi-On vs Epi-Off

- Transepithelial = Epi-ON
 - All Epi-On procedures are off-label and investigational
 - Many studies have evaluated and are **continuing to evaluate** for best protocol
 - Studies change variables in procedure (UV light source, time, riboflavin formula)
- [Efficacy and safety of transepithelial corneal collagen crosslinking surgery versus standard corneal collagen crosslinking surgery for keratoconus: a meta-analysis of randomized controlled trials.](#) Li W, Wang B. BMC Ophthalmol. 2017 Dec 28;17(1):262. doi: 10.1186/s12886-017-0657-2.
 - [Prospective 2-year study of accelerated pulsed transepithelial corneal crosslinking outcomes for Keratoconus.](#) Ziaei M, Vellara H, Gokul A, Patel D, McGhee CNJ. Eye (Lond). 2019 Dec;33(12):1897-1903. doi: 10.1038/s41433-019-0502-3. Epub 2019 Jul 4. PMID: 31273313
 - [Comparison of transepithelial corneal collagen crosslinking with epithelium-off crosslinking in progressive keratoconus.](#) Kocak I, Aydin A, Kaya F, Koc H. J Fr Ophtalmol. 2014 May;37(5):371-6. doi: 10.1016/j.jfo.2013.11.012. Epub 2014 Mar 26.
 - [Transepithelial corneal crosslinking for keratoconus.](#) Hersh PS, Lai MJ, Gelles JD, Lesniak SP. J Cataract Refract Surg. 2018 Mar;44(3):313-322. doi: 10.1016/j.jcrs.2017.12.022.

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CXL Potential Complications

- Infections
- Non-healing epithelium
- Corneal Haze
- Corneal scarring
- Endothelial cell damage
- Continued progression

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Case Example

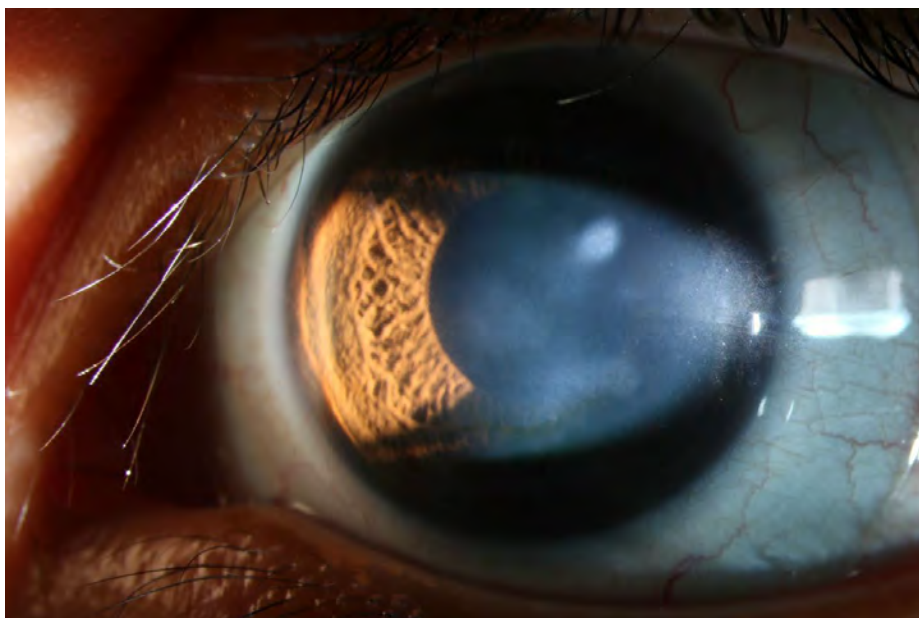
- 16 y/o Hispanic M
- KC Dx 2018 (age 13)
- CXL OS 2019
 - Resulted in diffuse persistent K haze/scarring
- BCVA with scleral lenses
 - 20/20 OD and 20/25- OS

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Corneal Haze s/p CXL



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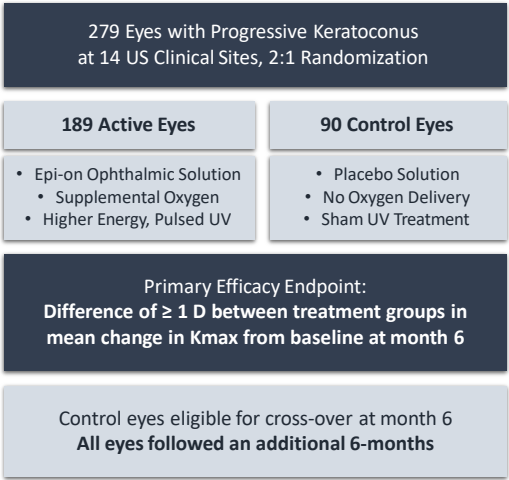
New protocols for CXL?

- LOTS to look forward to...
- May 2019 – Enrollment completed in Phase 3 Epi-On CXL Clinical trial for progressive KC
 - Multicenter (14 centers), randomized sham-controlled study
 - Latest-generation UV light source
 - Supplemental oxygen
 - New drug to penetrate K epithelium
 - Reduce treatment time

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US Phase III Pivotal Trial of Epi-on Cross-linking Therapy

Epi-on Treatment Demonstrated the Ability to Halt or Reduce the Progression of Keratoconus versus Observed Disease Progression in a Placebo-Control Arm



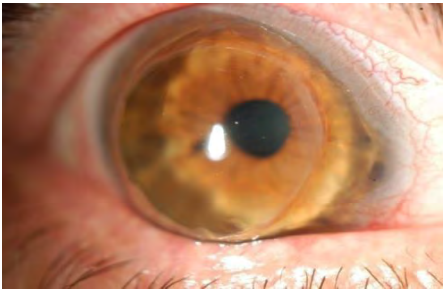
- Achieved prospectively defined primary efficacy outcome, demonstrating Kmax treatment effect of -1.0D ($p = 0.0004$)
- 98% of placebo randomized patients elected to cross-over to epi-on treatment
- Well-tolerated, majority of adverse events reported were mild and transient in nature, no change in corneal endothelial cell count over the course of the trial
- Forms the basis for planned regulatory submission (U.S. NDA) by Glaukos in 2022

Epithelium-on cross-linking is not approved by the US FDA

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Importance of Early KC Intervention

- Preserve vision
 - Maintain ability to wear contact lenses
 - Maintain quality of life
- Prevent progression
 - Corneal striae and scars reduce BCVA
- Protect against need for surgical intervention
 - Potential complications
 - Graft rejection
 - Cost



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How To Refer?

- <https://www.livingwithkeratoconus.com/>
- www.livingwithkc.com

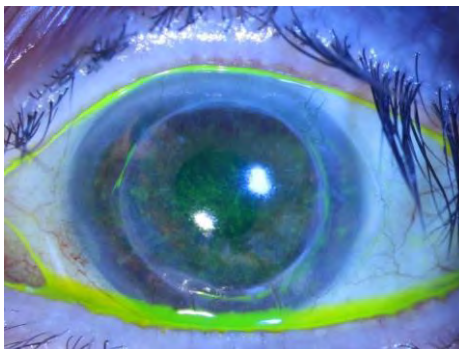


An Interactive Guide to Understanding Your Options

Does your insurance cover cross-linking? What are your options if cross-linking is not covered? Does your doctor perform FDA approved cross-linking? Finding the answers to all of these questions can feel very overwhelming! Follow this interactive guide to understand your options based on your unique situation.

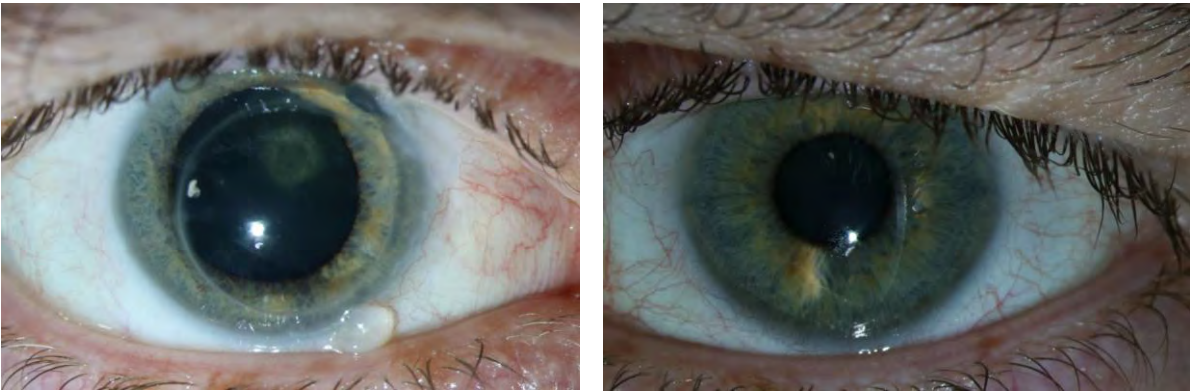
41

Better to Refer for 2nd Opinion or CXL
....than to Refer for a Cornea Transplant



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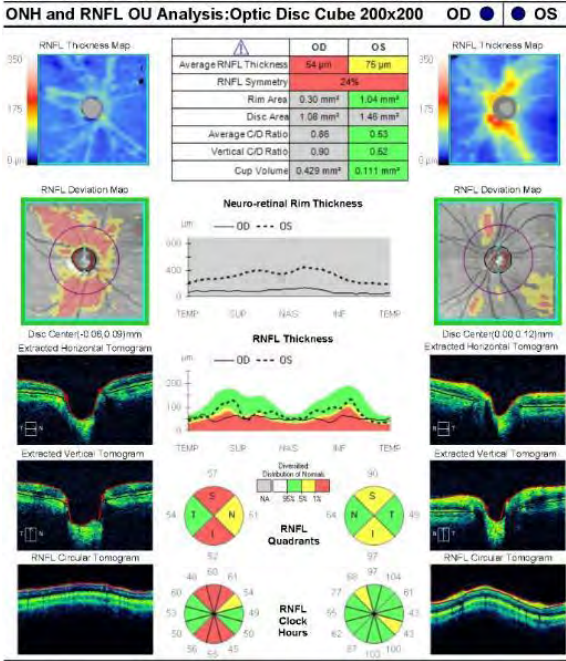
Case: Complications after Cornea Transplant



Study Conclusion: Physicians should maximize use of scleral or RGP CL because patients who successfully use CL have almost **one-fifth the risk of undergoing keratoplasty**.

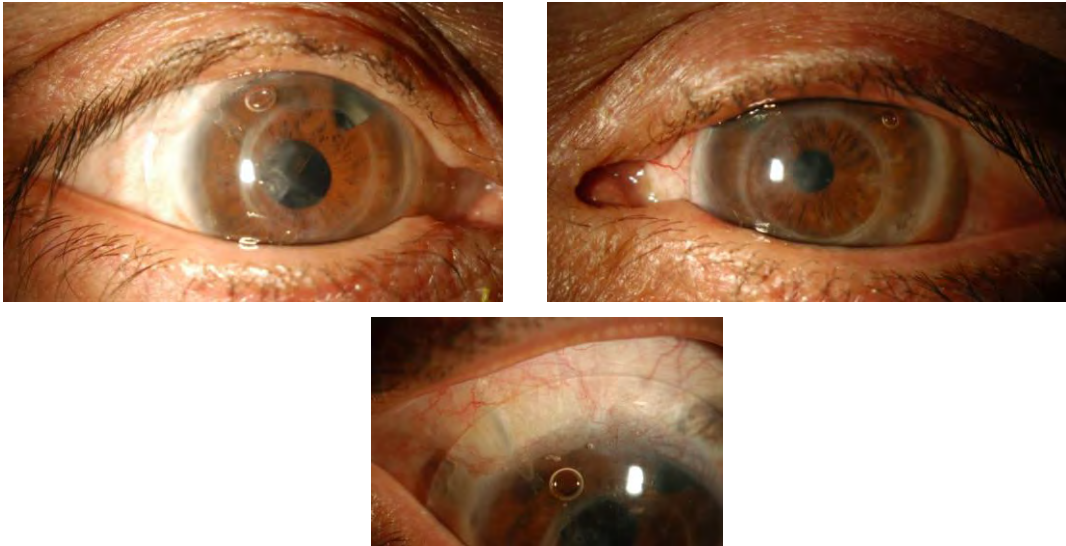
Ling JJ, Mian SI, Stein JD, Rahman M, Poliskey J, Woodward MA. **Impact of Scleral Contact Lens Use on the Rate of Corneal Transplantation for Keratoconus.** Cornea. 2021 Jan;40(1):39-42.

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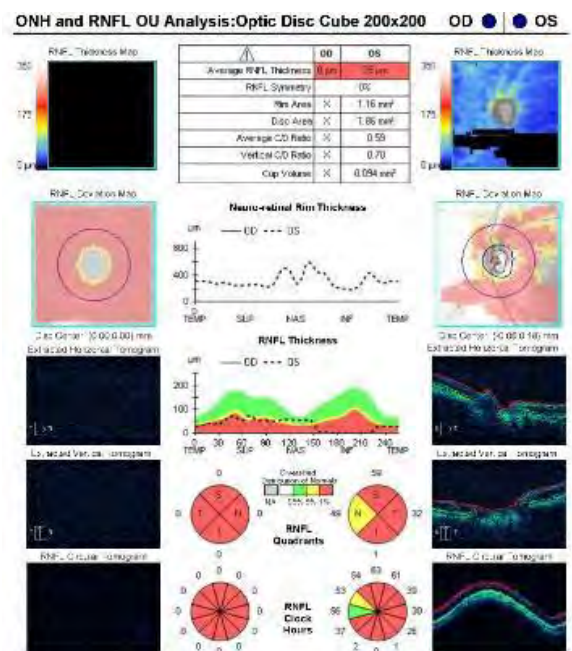
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Case: Complications after Corneal Transplant



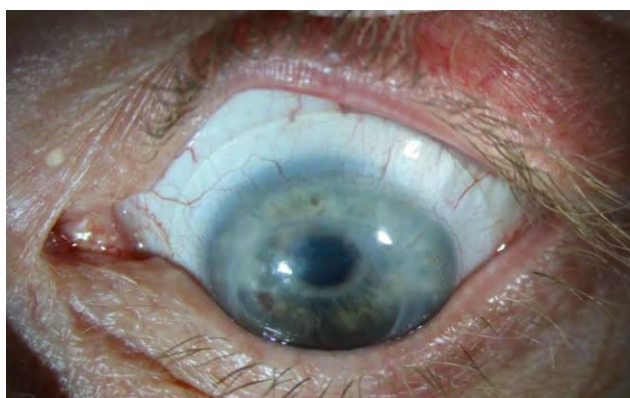
45

- 51 y/o Hispanic M
- PK OU 2 KC, in 30s
- Developed secondary GLC OU
- Tube-shunts OU
- Developed ET, had EOM Sx
 - Endophthalmitis OD s/p strab Sx
 - Monocular: HM VA OD
- Wears RGP OS now
 - BCVA 20/70



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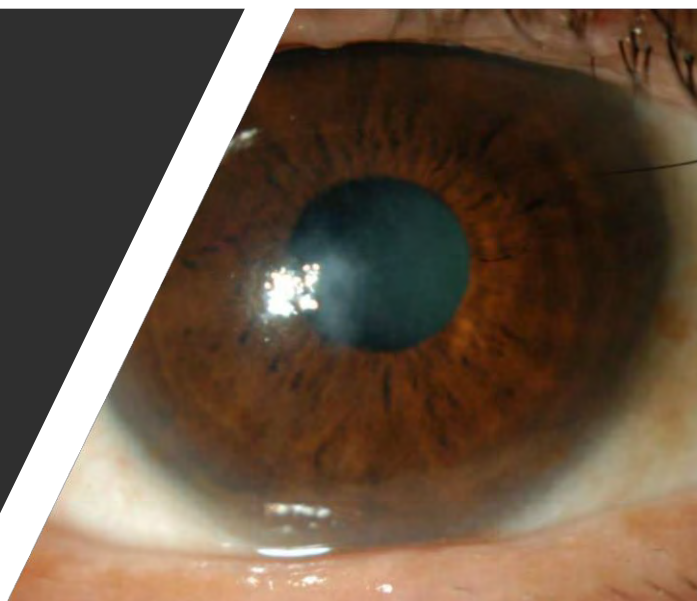
Case: Sclerals after Corneal Transplant



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Questions for thought

- Do you screen for keratoconus in every patient?
 - 1. Look for asymmetry on K
 - 2. Asymmetry between eyes
 - 3. Abnormal/high K values (>47 D)
- Should we screen for KC in every patient?
 - Yes. Unlike before, we now have CXL to slow/halt progression
- At what age should we start screening for KC?
 - Consider baseline imaging/topography at age 10-13
- Can we screen for KC as we do for glaucoma?



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Summary

- Keratoconus tends to present in early teens
 - Earlier presentation tends to be more aggressive
 - **Closer follow-up and lower threshold for CXL should be adopted in patients < 17 years and steeper than 55 D Kmax**
- Rather than monitoring the progression, we should detect and ACT to halt/slow the progression
- Visual and physiological changes from KC can greatly affect QOV/QOL
 - Aim to prevent changes associated with advanced KC and K transplantation
- Optometrists have vital role in detection and treatment of KC

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Thank You!

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**Marshall B.
KETCHUM UNIVERSITY**
Southern California College of Optometry

Updates in Clinical Glaucoma Management

Brian Song, MD



Updates in Clinical Glaucoma Management: Cases and Pearls

Brian J. Song, MD, MPH, FACS
Assistant Professor of Clinical Ophthalmology
Keck School of Medicine of USC
September 19, 2021

Disclosures

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- Previously supported by NEI/NIH K12 2K12 EY016335-11, Joanne Angle Investigator Award (Prevent Blindness), and Mentoring for the Advancement of Physician Scientists and Young Clinician Scientist Award (American Glaucoma Society).
- No proprietary interest in any of the devices or services mentioned in this presentation.

Why This Will Be One of the Most Important Hours of this Course:

- Glaucoma is the 2nd leading cause of blindness worldwide
- We will ALL come across glaucoma cases in practice at some point
- If you are not a glaucoma specialist, the key is to recognize:
 - Which cases are urgent and require immediate attention
 - How to treat or temporize the situation until you can get more help
 - Identify the source!

Goals of Today's Talk:

- To understand that glaucoma is an “umbrella” term that encompasses multiple diseases with common features
- To appreciate the multi-factorial nature of glaucoma
- To identify when surgical intervention is needed, or even preferred
- To use real-life case examples to illustrate the above

Case 1

Patient History

- 22 year old white male referred by his PCP to evaluate for “cataracts and elevated eye pressure” due to long-term steroid use
- No subjective complaints
- **PMH**
 - CNS (central nervous system) vasculitis
- **POH**
 - None

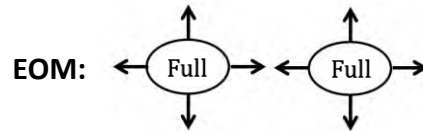
Ophthalmic Vitals

BCVA < $\begin{matrix} 20/20 \\ 20/20 -1 \end{matrix}$

IOP < $\begin{matrix} 25 \\ 24 \end{matrix}$

CVF: Full OU

Pupils < $\begin{matrix} 4 \rightarrow 3, \text{ brisk, no rAPD} \\ 4 \rightarrow 3, \text{ brisk, no rAPD} \end{matrix}$

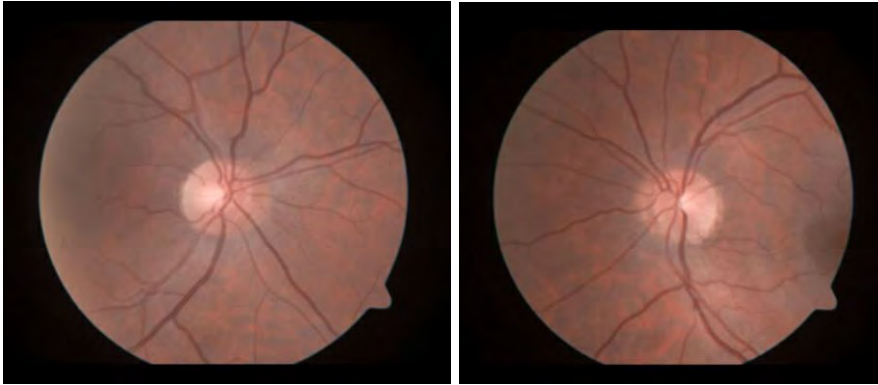


Gonioscopy: C40r1+ OU

Slit Lamp Exam

	OD	OS
Orbits/Adnexa	Normal	Normal
Lids/Lashes	Blepharitis	Blepharitis
Conjunctiva/Sclera	White and quiet	White and quiet
Cornea	1+ SPK	1+ SPK
Anterior Chamber	Deep and quiet	Deep and quiet
Iris	Flat, round	Flat, round
Lens	Clear	Clear
Anterior Vitreous	Clear	Clear

Fundus Exam

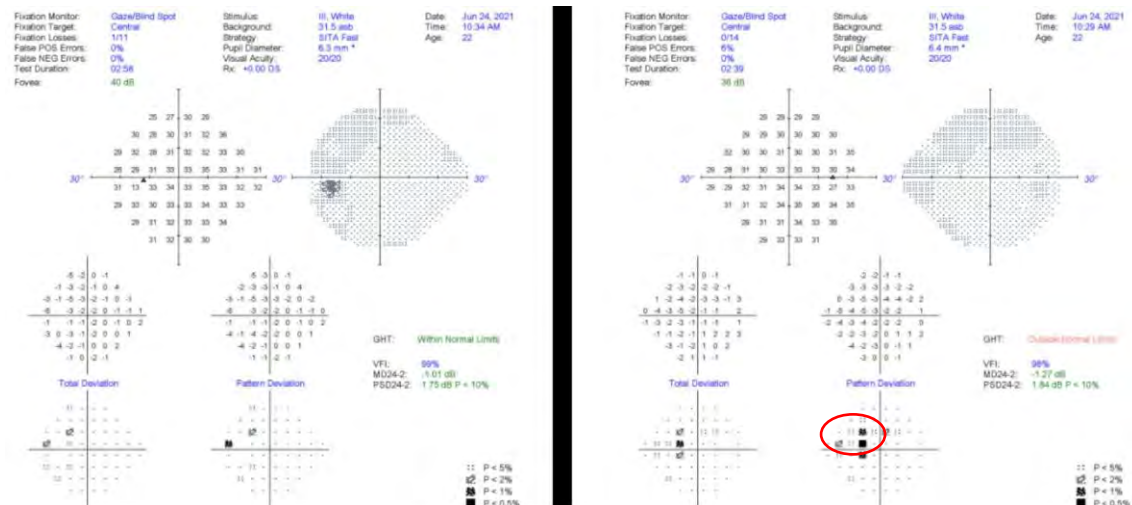


- Looks pretty normal to me?
- These small discs could not possibly be glaucoma?

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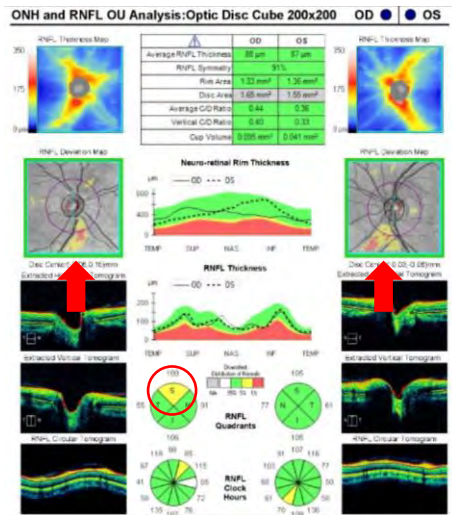
Humphrey Visual Field



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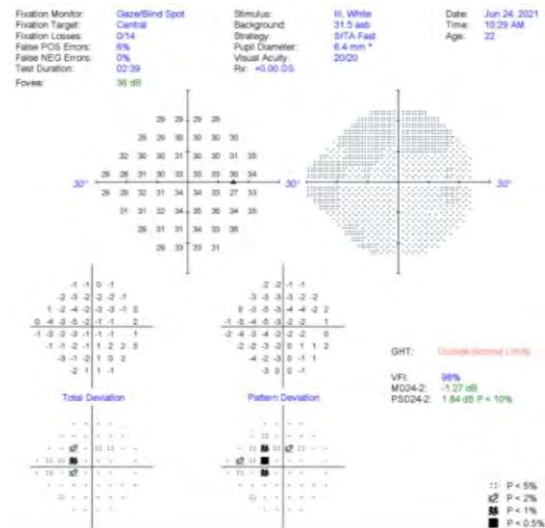
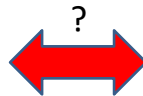
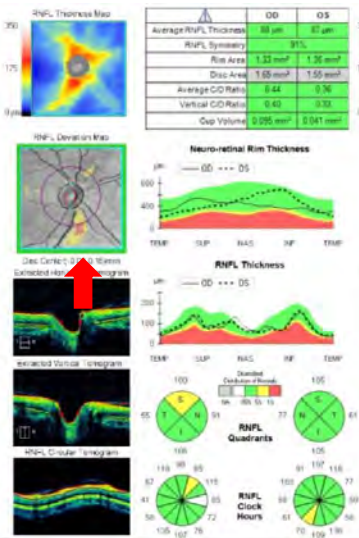
10

Optical Coherence Tomography (OCT)



- Quadrant scan:
 - Early superior RNFL thinning
- Does this correlate?
- What is going on?

Structure-Function Dissociation?



Assessment and Plan

- **Mixed mechanism open angle glaucoma OD > OS**
 - Primary Open Angle + Steroid Response
- What would you do next?
 - Selective laser trabeculoplasty
 - Angle surgery
 - Prostaglandin analogue
 - Rho-kinase inhibitor
 - Aqueous suppressant (beta-blocker or alpha-agonist)

COMBINED Mechanism vs MIXED Mechanism

- NOT the same thing
- ICD-9/10 and many ophthalmologists get this wrong
- COMBINED mechanism glaucoma
 - Open-angle with components of angle closure
 - e.g. Angle closure s/p LPI or cataract surgery, now with open angles
- MIXED mechanism glaucoma
 - Two or more etiologies contributing to glaucoma
 - e.g. underlying POAG with a steroid response

Pearls for Case 1

- Size matters!
 - Small discs can have small cups and still have glaucomatous optic neuropathy
- Risk factors deserve a work-up!
- Establish baselines!
- Look for structure-function correlation!
- Always interpret your own images. Do not rely on algorithms to make a diagnosis for you!



Case 2

Patient History

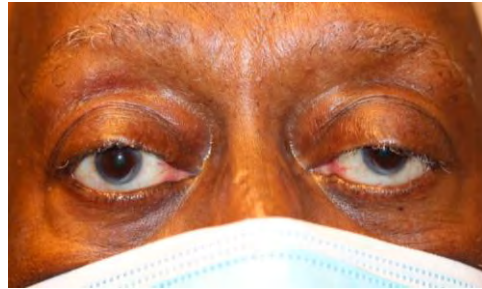
- 67 year old black male with thyroid eye disease (TED) OU with severe exophthalmos
- Presents May 2021 to re-establish care after being lost to follow-up since 2019

- **PMH**

- HTN
- Grave's disease

- **POH**

- TED OU
- Mixed mechanism glaucoma OU
- Iritis OD
- Hypertensive retinopathy OU



Past Ocular History

- Thyroid Eye Disease due to Graves Disease (dx ~ 2000)
 - s/p bilateral medial and lateral tarsorrhaphy OU 2015
 - Complicated by lagophthalmos, severe exophthalmos, and lid retraction causing exposure keratopathy and persistent retrobulbar pain
 - Thyroid disease currently inactive (CAS 0), not taking any thyroid meds. Follows closely with PCP
- Mixed Mechanism Glaucoma OS >> OD
- HTN retinopathy OU
- Iritis OD – quiescent
- Cataracts OU - visually significant

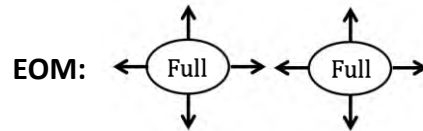
Ophthalmic Vitals

BCVA < $\frac{20}{25-2}$
20/50+2

IOP < $\frac{24}{27}$

CVF: Full OU

Pupils < $\frac{4 \rightarrow 3, \text{ brisk, no RAPD}}{4 \rightarrow 3, \text{ brisk, no RAPD}}$



Exophthalmometry:
Hertel 25.5 | 26 (Base: 102)

Slit Lamp Exam

	OD	OS
Orbits/Adnexa	Proptosis	Proptosis
Lids/Lashes	Collarettes	Collarettes
Conjunctiva/Sclera	White and quiet	White and quiet
Cornea	Arcus, 3+ SPK inferiorly	2+ SPK inferiorly
Anterior Chamber	Deep and quiet	Deep and quiet
Iris	Flat, round	Flat, round
Lens	2+ NSC	2+ NSC
Anterior Vitreous	Clear	Clear

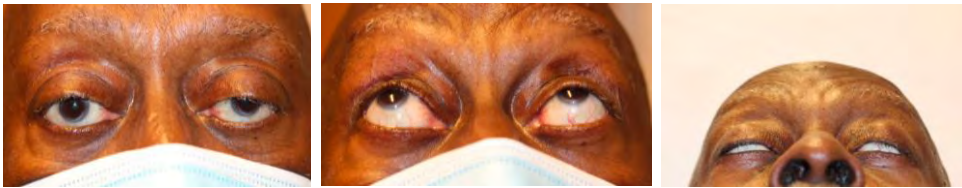
Fundus Exam

	OD	OS
CDR	0.65	0.9

December 2014:



June 2021:



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Assessment & Plan

- Thyroid Eye Disease, euthyroid levels
 - Recommend orbital decompression and ptosis surgery
 - Patient defers surgery again
 - Artificial tears and lubricating ointment QHS OU PRN
 - Follow-up 4 months for reassessment for surgery

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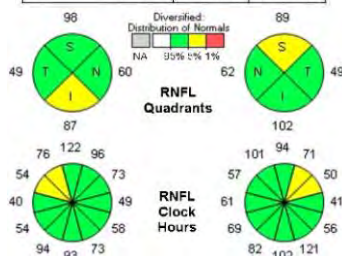
But....



OCT-RNFL

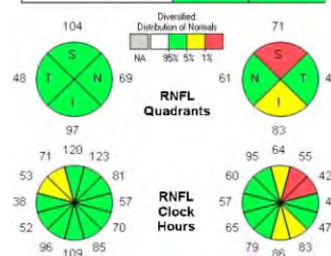
2015

	OD	OS
Average RNFL Thickness	74 μm	75 μm
RNFL Symmetry	81%	
Rim Area	1.13 mm^2	0.94 mm^2
Disc Area	2.07 mm^2	2.21 mm^2
Average C/D Ratio	0.66	0.75
Vertical C/D Ratio	0.69	0.76
Cup Volume	0.279 mm^3	0.405 mm^3



2021

	OD	OS
Average RNFL Thickness	80 μm	65 μm
RNFL Symmetry	77%	
Rim Area	1.04 mm^2	0.80 mm^2
Disc Area	2.07 mm^2	2.28 mm^2
Average C/D Ratio	0.70	0.81
Vertical C/D Ratio	0.73	0.81
Cup Volume	0.321 mm^3	0.519 mm^3



Fundus Exam



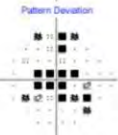
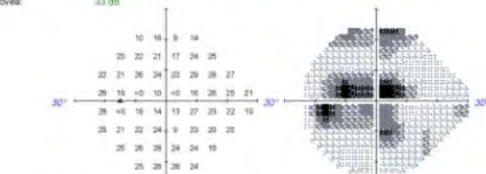
- Significant cupping OS > OD
- Is it enough to cause visual field loss?

Humphrey Visual Field

Fixation Monitor: Gaze/Blind Spot Central
 Fixation Target: 3/17
 Fixation Losses: 1%
 False POB Errors: 7%
 False NEG Errors: 06:18
 Test Duration: 33 dB
 Fovea: 33 dB

Stimulus: III, White
 Background: 31.5 asb
 Strategy: SITA Standard
 Pupil Diameter: <1 mm
 Visual Acuity: 20/60
 Rx: +3.25 DS

Date: May 19, 2021
 Time: 8:24 AM
 Age: 67



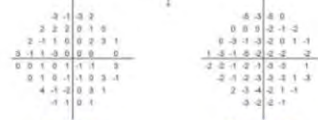
GHT: Glaucoma Suspect (1 error)
 VFI: 67%
 MD24-2: -5.23 dB P < 0.01
 PSD24-2: 9.57 dB P < 0.01

■ P < 5%
 ■ P < 2%
 ■ P < 1%
 ■ P < 0.5%

Fixation Monitor: Gaze/Blind Spot Central
 Fixation Target: 1/13
 Fixation Losses: 2%
 False POB Errors: 0%
 False NEG Errors: 05:04
 Test Duration: 34 dB
 Fovea: 34 dB

Stimulus: III, White
 Background: 31.5 asb
 Strategy: SITA Standard
 Pupil Diameter: <1 mm
 Visual Acuity: 20/60
 Rx: +3.25 DS

Date: May 19, 2021
 Time: 8:16 AM
 Age: 67



GHT: Within Normal Limits
 VFI: 100%
 MD24-2: 0.30 dB
 PSD24-2: 1.53 dB

■ P < 5%
 ■ P < 2%
 ■ P < 1%
 ■ P < 0.5%

Assessment & Plan (Continued)

- Mixed Mechanism Glaucoma OS >> OD
 - Uveitic glaucoma (hx of iritis)
 - Steroid response
 - Increased episcleral venous pressure due to TED
- Intolerant to multiple topical glaucoma medications
- After counseling, patient amenable to surgical decompression
- Recommend orbital decompression to reduce episcleral venous pressure and IOP ASAP!

Surgical Course

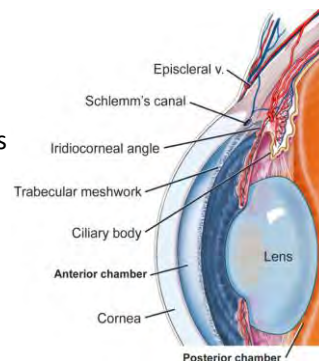
- **June 08, 2021** – Orbital Decompression OS
 - Pre-Op: IOP 24 mmHg
 - POD#3: IOP 15 mmHg
 - POW#1: IOP 14 mmHg; significant nasal chemosis, s/p conjunctival incision and close monitoring
 - POW#3: IOP 17 mmHg
 - POM#1: IOP 20 mmHg; IOP was improving but now likely steroid response
 - POM#1.5: 15 mmHg; IOP much improved but not at target

Surgical Course

- **June 08, 2021 – Orbital Decompression OS**
 - Pre-Op: IOP 24 mmHg
 - POD#3: IOP 15 mmHg
 - POW#1: IOP 14 mmHg; significant nasal chemosis, s/p conjunctival incision and close monitoring
 - POW#3: IOP 17 mmHg
 - POM#1: IOP 20 mmHg; IOP was improving but now likely steroid response
 - POM#1.5: 15 mmHg; IOP much improved but not at target
- **June 22, 2021 – Orbital Decompression OD**
 - Pre-Op: IOP 27 mmHg
 - POD#1: IOP 15 mmHg
 - POD#6: nasal chemosis– drained in clinic
 - POW#2: IOP 21 mmHg; IOP was improving but now likely steroid response
 - POM#1: IOP 17 mmHg

Glaucoma – An Extension of Thyroid Disease?

- **Mechanisms for IOP elevation**
 - Congestive orbitopathy with secondary elevation of **episcleral venous pressure**
 - Conventional aqueous humor drainage → aqueous and episcleral veins
 - Increased retrobulbar pressure
 - Contraction of extra-ocular muscles with increased pressure on the globe
 - Increased mucopolysaccharide deposition within the trabecular meshwork
 - Genetic predisposition
- Ocular hypertension in TED estimated to be ~3.1%–24%
 - Strong risk factor for open-angle glaucoma (OAG)
- Progression to glaucoma in TED estimated to be ~0.8–13%
- Association between OAG and TED (?) is controversial



Treatment of Thyroid Eye Disease

- Reduce IOP
- Corneal protection
- Comfort
- Cosmesis
- Prevention of irreversible damage to cornea or optic nerve
 - i.e. Exposure keratopathy or optic neuropathy

Medical Treatment

- **Corticosteroids** traditionally used in *active disease* → reduce inflammation and lower IOP
- **IOP lowering medications**
 - Aqueous suppression
 - Alpha-2 agonists (i.e. Brimonidine); β blockers (i.e. Timolol); Carbonic anhydrase inhibitors (i.e. Dorzolamide, Brinzolamide, Acetazolamide, Methazolamide)
 - Prostaglandin analogues
 - Periorbital fat atrophy / atrophy of adipocytes may be helpful?
 - Eftekhari et al. demonstrated decreased adipocyte density with orbital injection of bimatoprost compared to control orbits injected with saline (2018)
 - Rho kinase inhibitors
 - Increased trabecular outflow; decreased aqueous production and episcleral venous pressure

Surgical Treatment

- Orbital Decompression
 - Lowers IOP by decreasing episcleral venous pressure
 - Increase superior ophthalmic venous flow
 - Increase in choroidal perfusion
- Trabeculectomy / Tube Shunt
 - Beware of lid positioning!
- Minimally Invasive Glaucoma Surgery?



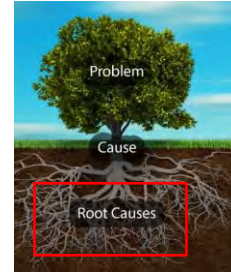
Surgical Treatment

- Angle Surgery?
 - Lowers IOP by overcoming resistance at the level of the trabecular meshwork
 - What happens if episcleral venous pressure is high though?
 - Proceed with caution!



Pearls for Case 2

- Glaucoma is oftentimes a “systemic” disease
 - Obtain a good history and be aware of the patient’s medical history
- When your roof is leaking, the answer is NOT to “buy more buckets”
 - Get to the ROOT of the problem
 - The best treatment is not always a glaucoma medication or surgery



Case 3

Patient History

- 82 year old Asian female referred by her general ophthalmologist for ocular surface disease and glaucoma progression despite maximum medical therapy
- Complains of red, irritated eyes and “tearing” OU
- **PMH**
 - Hypertension
 - History of pulmonary embolus – on Rivaroxaban
- **POH**
 - Combined mechanism glaucoma OU
 - Status post cataract surgery OU (2015)
 - Dry eye syndrome / ocular surface disease

Past Ocular History

- Combined mechanism glaucoma OU
 - Status post selective laser trabeculoplasty (SLT) OU (2018)
 - Status post trabeculectomy OU (2019)
 - Netarsudil/Latanoprost QHS OU; Brimonidine BID OU; Dorzolamide/Timolol BID OU
- Pseudophakia OU
 - Status post cataract surgery OU in 2018
- Dry Eye / Ocular Surface Disease OU
 - Artificial tears QID OU
 - Cyclosporine BID OU

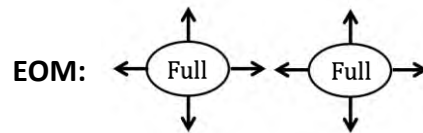
Ophthalmic Vitals

BCVA < $\begin{matrix} 20/25 \\ 20/40 \end{matrix}$

IOP < $\begin{matrix} 14 \\ 11 \end{matrix}$

CVF: Full OU

Pupils < $\begin{matrix} 4 \rightarrow 2, \text{ brisk, no RAPD} \\ 4 \rightarrow 2, \text{ brisk, no RAPD} \end{matrix}$

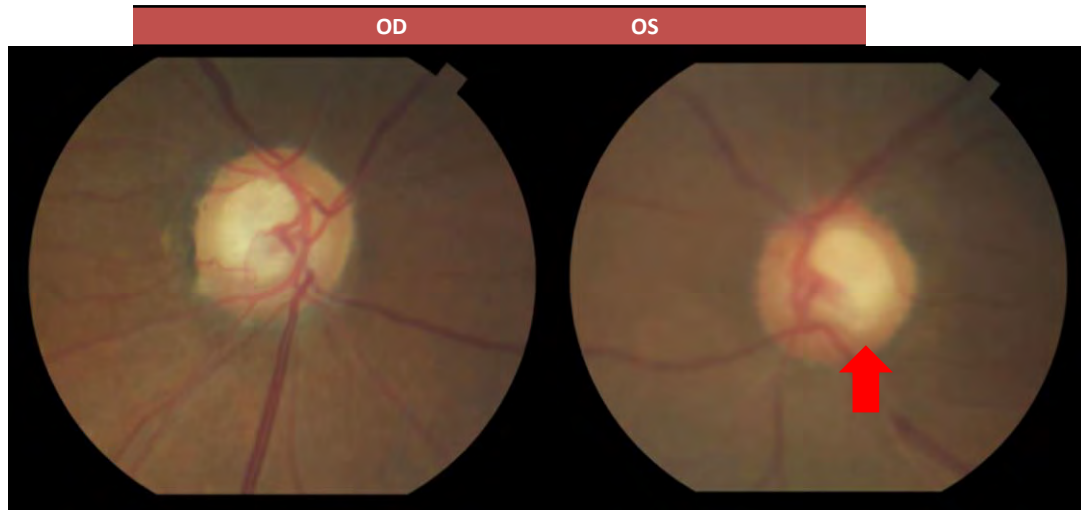


Gonioscopy: A-C35r3+ with scattered PAS OU

Slit Lamp Exam

	OD	OS
Orbits/Adnexa	Normal	Normal
Lids/Lashes	Blepharitis, periorbitopathy	Blepharitis, periorbitopathy
Conjunctiva/Sclera	1+ injection; flat superior bleb	1+ injection; flat superior bleb
Cornea	3+ PEE	3+ PEE
Anterior Chamber	Deep and quiet	Deep and quiet
Iris	Round, superior iridectomy	Round, superior iridectomy
Lens	PCIOL	PCIOL
Anterior Vitreous	Clear	Clear

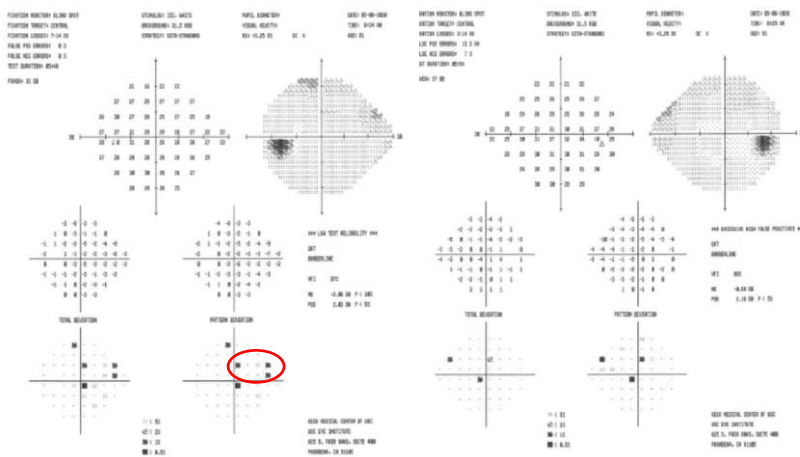
Fundus Exam



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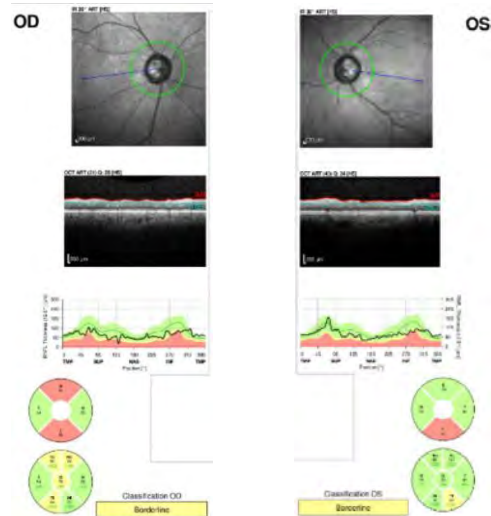
Humphrey Visual Field



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OCT-RNFL



Assessment and Plan

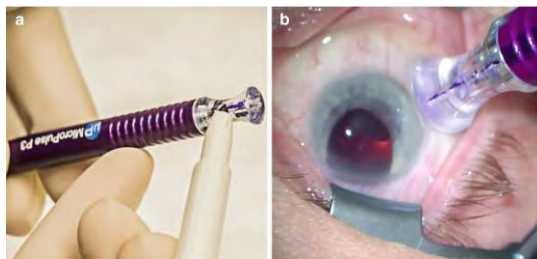
- **Combined mechanism open angle glaucoma OS > OD**
- What would you do next?
 - Continue present management
 - Change Dorzolamide/Timolol to preservative-free
 - Discontinue Brimonidine
 - Discontinue Netarsudil/Latanoprost combination
 - Selective laser trabeculoplasty

Clinical Course

• August 14, 2020

- Follow-up IOP check and reassessment of symptoms
 - IOP: 15 mmHg OD || 13 mmHg OS
 - Target IOP = low teens OU
- Minimal change in symptoms since stopping Brimonidine and switching to Dorzolamide/Timolol preservative-free
- Recommend micropulse cyclophotocoagulation OD first

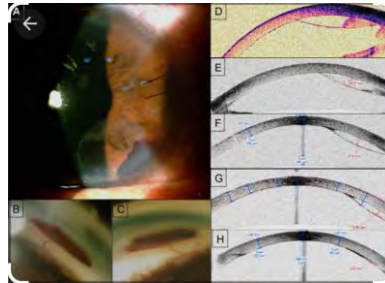
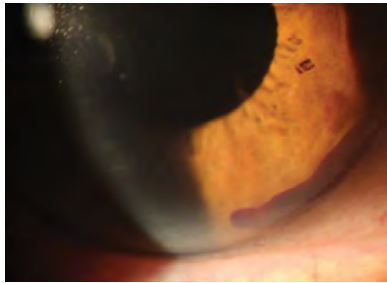
Micropulse Cyclophotocoagulation



• Why?

- Patient has relatively mild glaucoma
- Significant bleeding risk from trans-scleral filtration surgery due to anti-coagulation use

Micropulse Cyclophotocoagulation



• How about MIGS?

- Significant hyphema risk due to anti-coagulation use
- Risk of Descemet's detachment after prior trabeculectomy

Transcleral Cyclophotocoagulation

DEVICE COMPARISON	G-PROBE	G-PROBE illuminate	MP3 PROBE
Glaucoma Treatment Stage	Refractory	Refractory	Primary Open Angle
Repeatable Procedure	Yes	Yes	Yes
MicroPulse Technology	No	No	Yes
Transillumination	No	Yes	No
Destructive	Yes	Yes	No ²²
Therapy Location	Office & OR	Office & OR	Office & OR
CPT Code	66710	66710	66710
SmartProbe Technology	Laser Parameter Memory Enabled	Laser Parameter Memory Enabled	Laser Parameter Memory Enabled
Patented Probe Design	Wedge	Wedge	Curve



- Micropulse is not believed to be cyclo-destructive
 - Delivers short pulses of energy with resting periods in between
 - Results in “cooling period” between bursts and less collateral tissue damage

Clinical Course

- **July 23, 2021**
 - Status post MP-CPC OD 11/12/2020 and OS 4/6/2021
 - IOP: 12 mmHg OD || 9 mmHg OS
 - IOP at goal on only Dorzolamide/Timolol preservative-free BID OU
 - Conjunctival hyperemia improved/resolved
 - OCT-RNFL and HVF stable from 2020

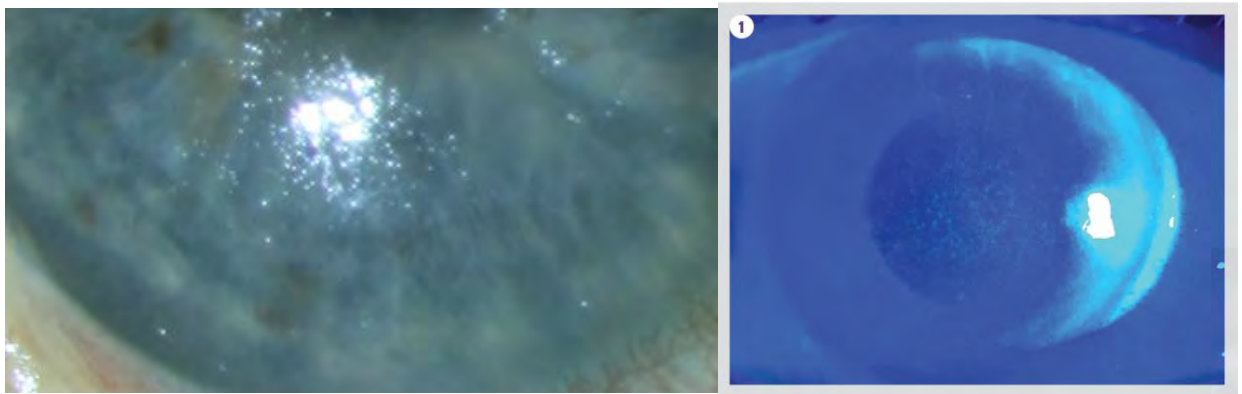
Ocular Surface Disease in Glaucoma Patients

- Glaucoma treatment may cause chronic inflammation or aggravate a concomitant ocular surface disease
- Topical glaucoma medications can cause burning, irritation, itching, tearing, and decreases in visual acuity within three months of medication initiation
- Glaucoma therapy-related ocular surface disease:
 - “Imbalance of the ocular surface homeostasis caused by the toxic effect of chronic topical medication, which leads to tear film instability, epithelial damage, and inflammation.”

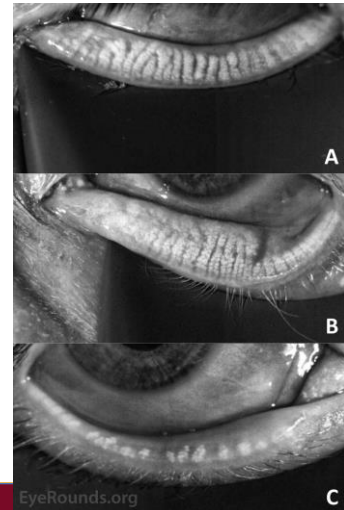
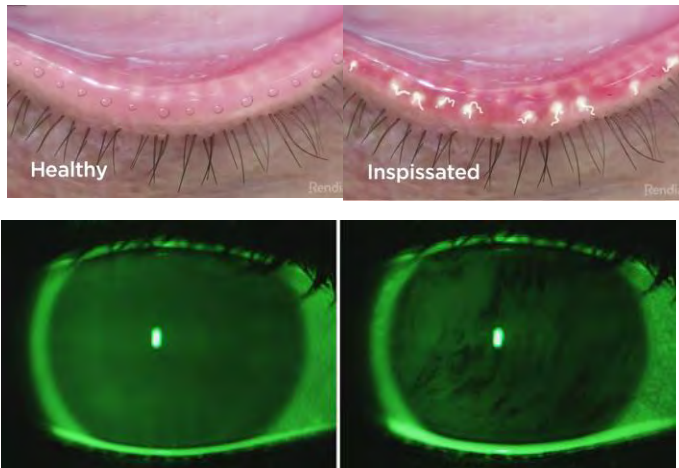
Clinical Manifestations of Surface Issues in Glaucoma Patients

- Punctate epitheliopathy
- Dry eye disease
- Meibomian gland dysfunction and tear film instability
- Allergy
- Pseudopemphigoid

Punctate epitheliopathy/Toxic keratitis



Meibomian Gland Dysfunction

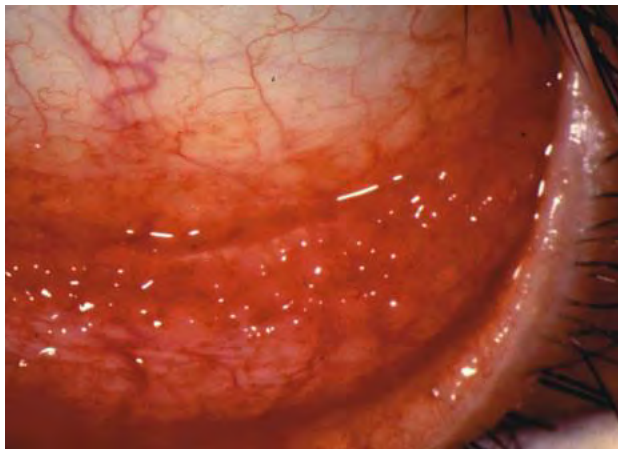


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<https://www.eyerounds.org/tutorials/Meibomian-gland-dysfunction-related-evap-dry-eye-syndrome/Fig4-LRG.jpg>
<https://ilux.myalcon.com/>

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Allergy

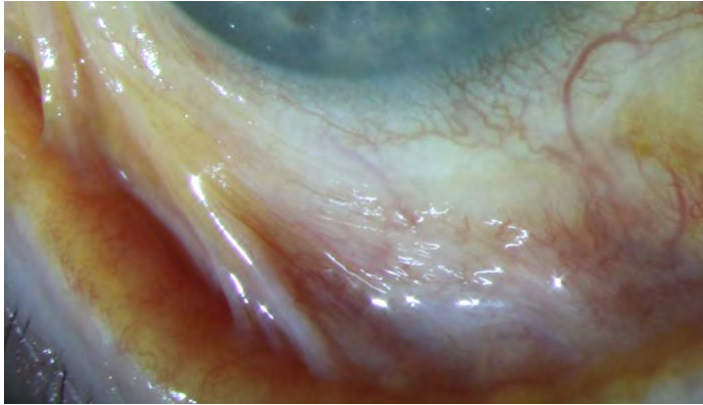


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<https://lh3.googleusercontent.com/proxy/633psEd>

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Pseudopemphigoid



Pathology of Drop Toxicity

- Preservatives
 - Benzalkonium chloride (BAK)
 - Bacteriostatic and bactericidal by destroying cell membranes
 - Pro-inflammatory and toxic
- Prostaglandin analogues (PGAs)
 - MGD and atrophy, reactivation of herpetic keratitis
- Alpha-adrenergic agonists
 - High ocular allergy rate
- Carbonic anhydrase inhibitors
 - Increased corneal thickness
- Beta-blockers
 - Reduce basal tear and alter mucus production

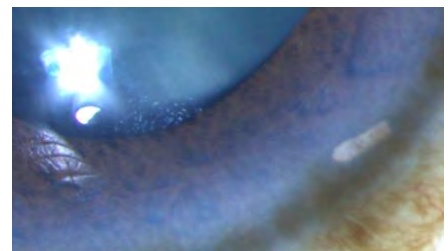
Medical Treatment

- Avoid BAK preservative
- **BETTER:**
 - Alternative preservatives
 - Purite
 - SofZia
- **BEST:**
 - Preservative-free hypotensive medications



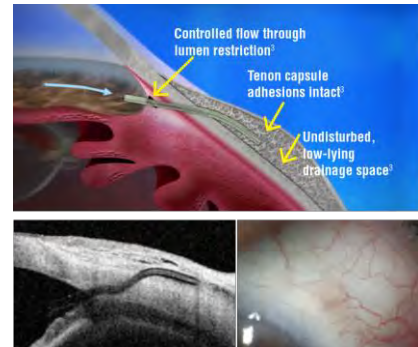
Medical Treatment

- Discontinue ineffective medications
- Lubrication without added preservatives
- Topical cyclosporine or lifitegrast
- Warm compresses/lid hygiene, IPL, thermal pulsation (if MGD present)
- New forms of drug delivery
 - Drug-eluting punctal plugs and contact lenses
 - Implants: Bimatoprost SR (Durysta)



Surgical Treatment

- Selective Laser Trabeculoplasty (SLT)
- Minimally Invasive Glaucoma Surgery (MIGS)
- Filtration Surgery
 - Trabeculectomy
 - Glaucoma tube shunt
 - XEN gel stent
- Cyclophotocoagulation?



First-line option? Treatment Paradigm shift?

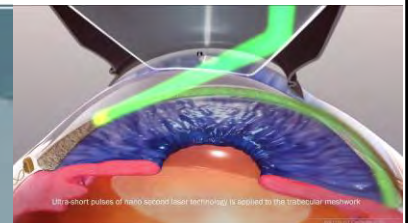
THE LANCET

ARTICLES | VOLUME 393, ISSUE 10180, P1505-1516, APRIL 13, 2019

Selective laser trabeculoplasty versus eye drops for first-line treatment of ocular hypertension and glaucoma (LiGHT): a multicentre randomised controlled trial

Gus Gazzard, FRCOphth • Evgenia Konstantakopoulou, PhD • Prof David Garway-Heath, MD • Anurag Garg, FRCOphth
Victoria Vickerstaff, MSc • Rachael Hunter, MSc • et al. Show all authors • Show footnotes

Open Access • Published: March 09, 2019 • DOI: [https://doi.org/10.1016/S0140-6736\(18\)32213-X](https://doi.org/10.1016/S0140-6736(18)32213-X) • Check for updates



Pearls for Case 3

- Ocular surface disease is often exacerbated in glaucoma patients, especially with the use of BAK-preserved topical anti-hypotensive.
- In glaucoma patients with dry eyes, care should be taken to reduce the burden of topical glaucoma medications in order to improve quality of life and treatment compliance and effect.
 - Modern medicine affords us good options to achieve this including:
 - Selective laser trabeculoplasty
 - Drug-eluting implants
 - Minimally invasive glaucoma surgery
 - Filtration surgery
- Cyclophotocoagulation, especially micro-pulse, is a reasonable option even for well-seeing eyes in the right circumstances

Summary and Take Home Points from Today's Cases

- Glaucoma is an “umbrella” term
- Not all glaucomas are the same
 - Just like not all cancers are the same
- A *group* of diseases characterized by:
 - *Progressive* optic neuropathy
 - *Characteristic* visual field loss
 - In total, there are > 20 types of glaucoma
- Intraocular pressure (IOP) is NOT used to define glaucoma
 - Just like we do not define lung cancer by smoking
 - IOP is the primary risk factor for glaucoma, but it is not a part of its definition
 - Treat the *disease* NOT the pressure



Summary

- Glaucoma is a *central nervous system* disease with *systemic relationships*
 - If you take the time to understand the **root** cause, you and your patients will be better off



Thank You



(Virtual) Questions?





**Marshall B.
KETCHUM UNIVERSITY**
Southern California College of Optometry

Refer or Relax? Macula

Steven Ferrucci, OD



REFER OR RELAX: macula

Steven Ferrucci, OD, FAAO
Chief, Optometry, Sepulveda VA
Professor, MBKU/SCCO

1

Disclosures

- Alcon
- Centervue
- Genentech
- Maculogix
- Optovue
- Regeneron
- Science Based Health
- Visible Genomics

2

INTRODUCTION

- Various macula cases will be presented
- Question is should the case be referred to a retina specialist OR can you monitor it yourself
- There are no right or wrong answers, just differences of opinion
- **JUST KIDDING, THERE ARE WRONG ANSWERS!**

3

Dry AMD

- **Currently mainstay treatment for Dry AMD revolves around prevention of progression through vitamins, nutrition and lifestyle changes**
 - Rheophoresis, Laser, Anecortave Acetate did not prove effective
 - Smoking #1 modifiable risk factor for getting AMD as well as its progression!
 - One study showed 90% of pts with AMD were not advised to quit smoking
- **Early detection of conversion from dry to wet may result in better treatment for patients**

4

AREDS

- First large scale study looking at nutrition and ocular health
- 3640 pts followed on average for 6.3 years
 - Results released October 2001
- Results showed that 25% risk reduction to developing advanced AMD in pts with intermediate (stage 3) AMD or worse
 - 500 mg vitamin C
 - 400 IU vitamin E
 - 15 mg vitamin A (25,000 IU beta carotene)
 - 80 mg zinc
 - 2 mg copper

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AREDS 2

- AREDS 2: Enrollment ended June 2008 with \approx 4200 patients followed for six years
 - Effect of lutein, zeaxanthin and omega 3 on AMD
 - Effect of eliminating beta carotene on AMD
 - Effect of reducing zinc on AMD
 - Effect of supplements on cataracts
 - Validate the AMD scale from original AREDS
- Results released May 5, 2013

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AREDS 2

■ Major Conclusions:

- The addition of lutein and zeaxanthin, DHA and EPA or both to the AREDS formulation did not further reduce the risk of progression to advanced AMD
- Substituting L/Z (10 mg/2 mg) for beta carotene is an appropriate substitution, because of potential increased incidence of lung cancer in former smokers

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Additional findings

■ Lutein and zeaxanthin did provide an additional 10% reduced risk over current supplements

- In patients with lowest dietary intake of l/z, additional 26% reduced risk

■ Decreasing zinc from 80 mg to 25 mg had no significant effect

- No change recommended (?)
- Deserves further study

■ Competitive absorption of carotenoids

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AREDS2 Formulation

Vitamin C (500 mg)

Vitamin E (400 IU)

~~Beta Carotene (15 mg)~~

Lutein (10 mg)/Zeaxanthin (2 mg)

Zinc (80 mg zinc oxide)

Copper (2 mg cupric oxide)

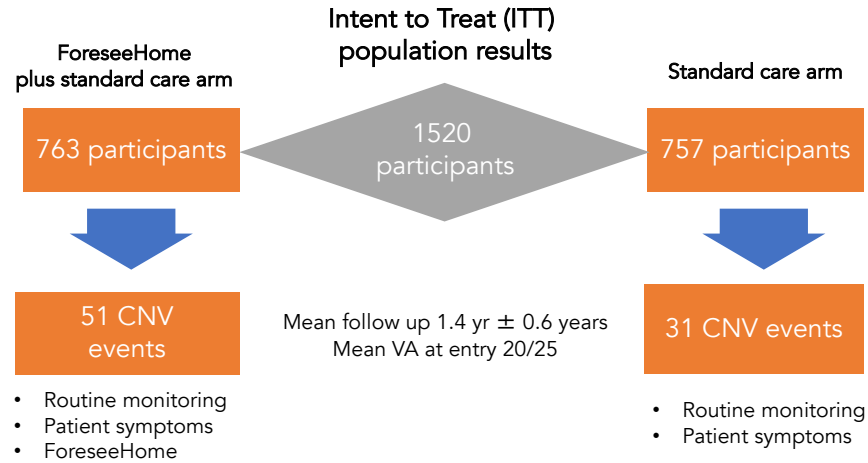
9

Dark adaptometer

- Dark adaptation is a sensitive marker for early AMD
- The AdaptDx measures dark adaptation
- A rapid test of dark adaptation using the AdaptDx has been found to have a 90% sensitivity for detecting dark adaptation impairment associated with AMD
- Decreased dark adaptation may precede clinical findings of AMD by as much as 3 years
- Dark adaptation is more sensitive than other tests such as Snellen acuity, contrast sensitivity, or visual fields which are about 25% sensitive.

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AREDS2-HOME Study



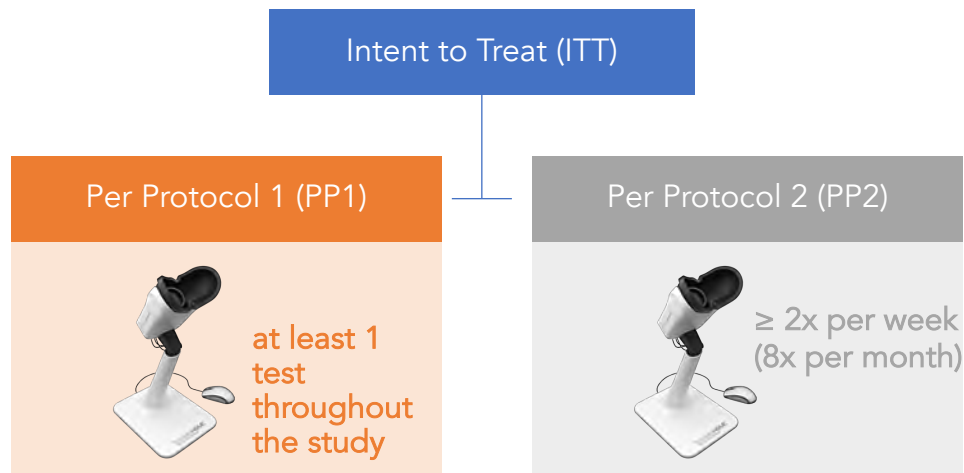
**Primary outcome: Change in BCVA from baseline to CNV detection*

Reference: AREDS2-HOME Study Research Group. Ophthalmology. 2014;121(2):535-544.

11

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ForeseeHome Arm

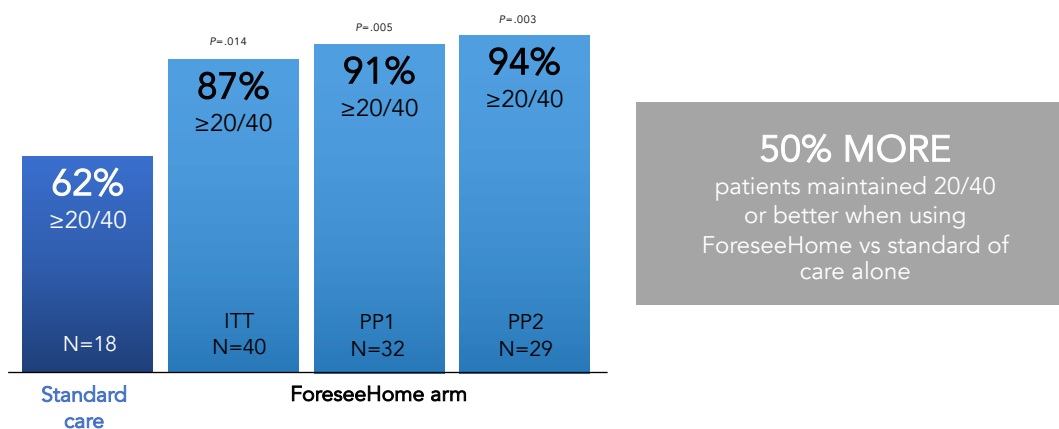


Reference: AREDS2-HOME Study Research Group. Ophthalmology. 2014;121(2):535-544.

12

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More patients who used ForeseeHome maintained $\geq 20/40$ VA



*94% of patients maintained 20/40 at time of wet AMD diagnosis;
Absolute visual acuity at time of wet AMD diagnosis is critical to visual acuity outcomes at year 1*

Reference: AREDS2-HOME Study Research Group. Ophthalmology. 2014;121(2):535-544.

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Home OCT

- Notal OCT Analyzer (NOA)
- “Uses computer image analysis algorithm to provide automated detection of pathological fluid in exudative retinal disease, including wet AMD, macular edema and retinal vein occlusion”
- Performance validated in study comparing sensitivity , specificity and accuracy with 3 retinal specialist

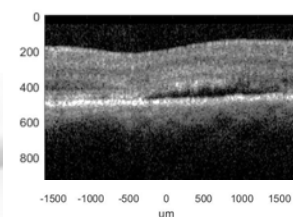
14

Patient Self-operated Home OCT provides high quality images

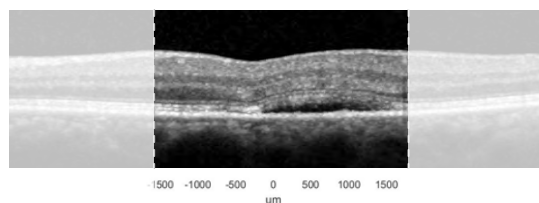
- Patient self-installed and self-operated OCT device
- Monitoring of intra- and subretinal fluid in between office visits
- Provides cross sectional images of the central 10 deg. (3 mm x 3 mm) of the macula in patients with exudative AMD
- 88 B-scans with dense 34 μm spacing ensure high sensitivity of fluid detection
- Test takes approximately 10 sec. per eye
- Device uploads OCT data to cloud



Home OCT



Heidelberg Spectralis (in-office device)



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Home OCT Performance and Roadmap

- US clinical trial demonstrated 90% of 196 elderly wet AMD patients with VA > 20/400 could self-operate and self-capture readable images following a 2-minute video tutorial (presented at ASRS 2019)
- Human graders identified fluid with SENSITIVITY = 91.5% and SPECIFICITY = 97.0% for Notal Home OCT V2.5 when compared to commercial OCT devices (presented at ASRS 2019)
- Notal Vision's patient-operated, AI-enabled Home OCT system was granted FDA Breakthrough Device Designation Status, and was selected to participate in FDA's OCT Innovation Pilot Program
- Notal Vision plans to bring first devices to patients' homes in 2020 as part of clinical trials with a commercial launch in 2021

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Is AMD in our DNA?

- AMD is a genetic disease with known markers accounting for at least 70% of the population attributable risk
- Other 30% is environmental/lifestyle
- Risk factors
 - Non-modifiable: age, race, gender
 - Modifiable: Smoking, increased BMI, poor diet/nutrition, UV exposure

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AMD is a Genetic Disease

Population Attributable Risk	
Condition	Genetics (%)
Colorectal Cancer	35
Diabetes II	26
Coronary Artery Disease	40
AMD	70

Those with stronger genetic risk develop more advanced disease earlier in life.

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Major genetic factors

- CFH
 - Single most important genetic component
 - CFH Y402H
- ARMS2/HTRA1
 - Second most important gene in AMD
- C3
 - Another component of the complement system
- Others
 - Less important

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Genetic Factors and Risk: More than additive!

- Former Smokers: 1.29x
- Current Smokers: 2.4X
- Non-Smoker and CFH,Y402H: 7.6X
- Current smoker and CFH,Y420H: 34X

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AMD Genetic Testing: Arctic DX

Macula Risk NXG

Looks at 15 SNPs as well as smoking, BMI, age and AMD status to determine AMD patients who may progress to advanced AMD and vision loss in

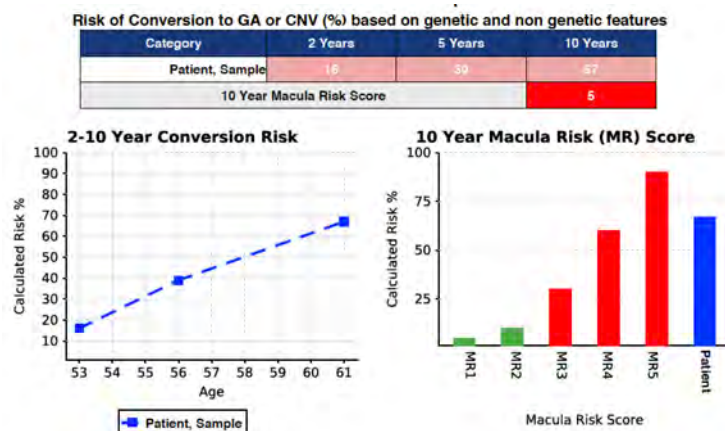
- 2 years
- 5 years
- 10 years



Cheek Swab

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Patient Report



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“Wet” AMD

- Neovascular “wet” AMD
 - Mainstay of treatment consists of serial intravitreal injection of anti-VEGF agents

Anti-VEGF Agents	Pegaptanib (Macugen®)	Ranibizumab (Lucentis®)	Aflibercept (Eylea®)	Brolucizumab (Beovu®)	Bevacizumab (Avastin®)
FDA approval	2004	2006	2011	2019	Not approved
Pivotal studies	VISION	ANCHOR MARINA IVAN	VIEW 1 and 2	HAWK HARRIER	CATT

- VEGF inhibitors have demonstrated *improved visual and anatomic outcomes* compared with other therapies

VEGF = vascular endothelial growth factor.

AAO. AMD preferred practice guidelines, 2019 (www.aao.org/preferred-practice-pattern/age-related-macular-degeneration-ppp). Kulkarni K, Prenner JL. *Rev Ophthalmol.* 1/13/2006. (www.reviewofophthalmology.com/article/an-update-on-macugen-trials). URLs accessed 5/30/2020.

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Anti-VEGF Agents

- VEGF is a primary driver of blood vessel growth and leakage in AMD
- Anti-VEGF agents block and neutralize VEGF
 - Results in decreased intra- and sub-retinal fluid
 - May also decrease risk of scar tissue formation
- Serious adverse effects (endophthalmitis) rare
- Less serious events (subconjunctival hemorrhage, vitreous hemorrhage, floaters) are also uncommon

Pongsachareonnon P, et al. *Clin Ophthalmol.* 2018;12:1877-1885. Yeo NJY, et al. *Front Pharmacol.* 2019;10:1363. Holz FG, et al. *Br J Ophthalmol.* 2016;100:1623-1628.

American Society of Retina Specialists (ASRS). Intravitreal injections. (www.asrs.org/content/documents/fact-sheet-30-intravitreal-injections.pdf). Sukgen EA, et al. *Int Ophthalmol.* 2017;37:215-219. Living well with low vision. (<https://lowvision.preventblindness.org/2013/06/25/betadine-and-eye-pain/>). URLs accessed 5/30/2020

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Anti-VEGF Agents: Delivery and Dosage

- Delivered intravitreally
- Dosing schedule and agent used varies
- In general
 - Loading dose with 1 injection per month for 3 months, then inject based on FA, OCT, or other clinical findings
 - Reduces patient burden while still delivering good results

Holz FG, et al. *Br J Ophthalmol*. 2016;100:1623-1628. Kress B. *Review of Optometry*. 2019 (<https://www.reviewofoptometry.com/article/antivegf-where-are-we-now>). Accessed June 10, 2020. Treating Wet AMD with Anti-VEGF drugs. 2016 (<https://www.reviewofophthalmology.com/article/treating-wet-amd-with-antivegf-drugs>). Accessed June 10, 2020

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Anti-VEGF Agents: Outcomes

Lucentis¹

- **94% stable vision at 2 years**
- **34–41% gained 15 letters or more**
- **Average gain of 11.3 letters at 1 year and 10.7 letters at 2 years**

Eylea^{2,3}

- 95% of patients treated maintained acuity
- 7.9–10.9 letters mean improvement of vision

Beovu⁴

- ~30% gained at least 15 letters by year 1
- Less fluid and greater reduction in CST vs aflibercept
- At 1 year, half of subjects on 3-month dosing

1. Brown DM, et al. *Ophthalmology*. 2009;116:57-65.e5. 2. Nguyen QD, et al. *Invest Ophthalmol Vis Sci*. 2011;52: abstract 3073. 3. Schmidt-Erfurth U, et al. *Invest Ophthalmol Vis Sci*. 2011;52:E-Abstract 1650. 4. Dugel PU, et al. *Ophthalmology*. 2020;127:72-84.

26

Other Treatment Options

Possible

- Photodynamic therapy (PDT) with verteporfin
- Intravitreal steroids in combination with PDT or anti-VEGF agents
- Laser photocoagulation
- Observation/suspension of treatment

Not advised

- Intravitreal steroids as monotherapy
- Radiation therapy
- Electrical stimulation
- Macular translocation therapy

AAO. AMD preferred practice guidelines, 2019 (www.aao.org/preferred-practice-pattern/age-related-macular-degeneration-ppp). Accessed 5/30/2020. Wong DT, et al. *Retina*. 2020;40:1010-1020. Giampoli E, et al. *J Ophthalmol*. 2018;56:12342. Veritti D, et al. *Expert Rev Ophthalmol*. 2010;5(5):681-688.

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Beovu (brolucizumab)

- Novartis
- FDA approved Oct 9, 2019
- Greater fluid resolution than previous agents with similar vision gains on 3 mos dosing
- Based on Hawk and Harrier Phase 3 trials

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Beovu (brolucizumab)

- Hawk and Harrier Study: compared to Eylea
 - 30% of pts gained at least 15 letters by year 1
 - Greater reduction in central retinal thickness at week 16 and 1 year than Eylea
 - Fewer pts with subretinal fluid than Eylea
 - Real key is extended dosing
 - After 3 monthly loading doses
 - By year 1, > ½ pts on 3 mos dosing
 - Rest were 2 mos dosing
 - Safety profile similar to Eylea

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Beovu update

- In Feb, 2020, American Society of Retinal Specialists (ASRS) issued a warning reporting 14 cases of retinal vasculitis following injection of Beovu
 - 11/14 were occlusive and resulted in vision loss
- In March, Novartis concluded that retinal vasculitis, retinal artery occlusion, or severe vision loss occurred in 8.75-10.08 out of 10,000 injection
- Added to warning label
 - Intraocular inflammation in 4% of pts
 - Artery occlusion in 1%
- Advised to avoid if pts had h/o inflammation to any other anti-Vegf agent

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Viagra and CSR

- Retina 2008: Fraunfelder and Fraunfelder
- 11 reported cases of CSR in men taking Viagra
 - In 8/11, pts stopped taking Viagra
 - In 6/8, vision improved with cessation
 - In 3 cases, CSR returned when started med again
 - 2 pts continued to have CSR after cessation
- Might consider recommending cessation of Viagra if active CSR, but relationship is unknown at this time

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Central Serous Retinopathy

- Common disorder of unknown etiology which typically affects men between age 20 and 45
 - Males to females 10:1
- Serous detachment of neurosensory retina due to leakage from small defect in RPE

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Central Serous Retinopathy

- Pt typically presents with fairly recent onset of blurred VA in one eye with a scotoma, micropsia, or metamorphopsia
 - VA typically 20/30-20/70
 - Often correctable with low hyperopic RX
 - Unilateral in 70% of cases

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Central Serous Retinopathy

- Appears as a shallow round or oval elevation of the sensory retina often outlined by a glistening reflex
- FA is helpful in providing definitive diagnosis
 - Classic Smoke stack appearance (occasionally)
 - Ink-blot appearance
- OCT shows marked elevation

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CSR: Risk Factors

TRADITIONAL

- Male > Female 10:1
- Age: Peak 20-45
- Type A personality
- Stress
- Pregnancy

OTHERS

- Steroid use
 - Oral
 - Topical?
 - Inhaled?
 - Injection?
- Choroidal Thickness
- Sleep apnea?
- Genes?
- Viagra?

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Central Serous Retinopathy

- 80-90% of pts will undergo spontaneous resolution and return to normal (or near normal) VA within 1-6 mos.
 - >60% resolve back to 20/20
 - Rare to have vision remain < 20/40
- Approx 40% will get recurrence
- CNVM is VERY rare occurrence, but possible

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CSR

- **When to worry/refer**
 - If VA worse than 20/70
 - If pt demographics do not support
 - If does not resolve in 6 mos
 - If gets worse rather than better
 - FA/ OCT does not support diagnosis
 - “Just doesn’t feel right”
 - Pt is unable to accept vision/prognosis

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Treatment

- | | |
|-----------------------------|--------------------------|
| • Observation | • Acetazolamide |
| • PDT | • Aspirin |
| • Anti-VEGF | • Metoprolol |
| • Anti-corticosteroids | • H.pylori treatment |
| • Rifampin | • Methotrexate |
| • Mifepristone | • Behavior Modification! |
| • Ketoconazole | |
| • Spironolactone/eplerenone | |
| • Finasteride | |

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LMH

- Lamellar Macula Hole OS
 - Also called partial thickness macular hole
- Pt ed.
- Monitor in 3 mos.
- Repeat OCT
- Consider retina referral if worsens

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LMH

- Symptoms
 - mild metamorphopsia,
 - limited acuity loss
 - stable vision
- Surgery is controversial
 - 25% to 75% improved visual acuity
- Therefore, monitoring seems reasonable

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FTMH

- Definition: Full thickness macular hole that affects all macular layers from ILM to RPE
- Size
 - Small: ≤ 250 μm
 - Medium: 250 μm to 400 μm
 - Large ≥ 400 μm
- Presence or absence of VMT
- By cause
 - Primary: Initiated by VMT (formerly idiopathic)
 - Secondary: from associated disease or trauma

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FTMH

- Small holes
 - Small rate of spontaneous closure
 - Very high surgical closure rate (almost 100%)
 - Best response to pharmacologic vitreolysis
- Medium holes
 - High surgical closure rate ($>90\%$)
 - Decent response to pharmacologic vitreolysis
- Large holes
 - High surgical closure rate (75-90%)
 - No response to pharmacologic vitreolysis
 - $\frac{1}{2}$ of all holes are large at time of diagnosis

42

antiVEGF

- Lucentis, Avastin, Eylea
- Shown in multiple studies to be beneficial for DME
 - RISE
 - 18.1% of pts in sham gained ≥ 15 letters vs. 44.8% (0.3 mg) or 39.2% (0.5 mg)
 - 2.6 letters gained in sham vs. 12.5 (0.3mg) or 11.9 (0.5mg)
 - RIDE
 - READ
 - VISTA
 - VIVID

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Protocol V

- 702 pts with CI-DME with VA 20/25 or better
- 3 treatment groups
 - Eylea
 - FML
 - Observation
- At end of 2 years, rate of loss of 5 letters or more similar in all 3 groups
- Avg acuity in all 3 groups was 20/20
- Bottom line: pts with CI-DME and good VA can be observed

44

DM/DME

- Refer if center involved DME/CSME evident on OCT in 1-2 weeks
- If not center involved, follow closely in 3- 6 mos
- Pt ed re role of BS/BP control
- Treatment: FML vs. serial anti-VEGF

45

ERM

AGE	INCIDENCE
< 60	1.7%
60-69	7.2%
70-79	11.6%
80+	9.3%

BLUE MOUNTAIN EYE STUDY, AUSTRALIA

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ERM

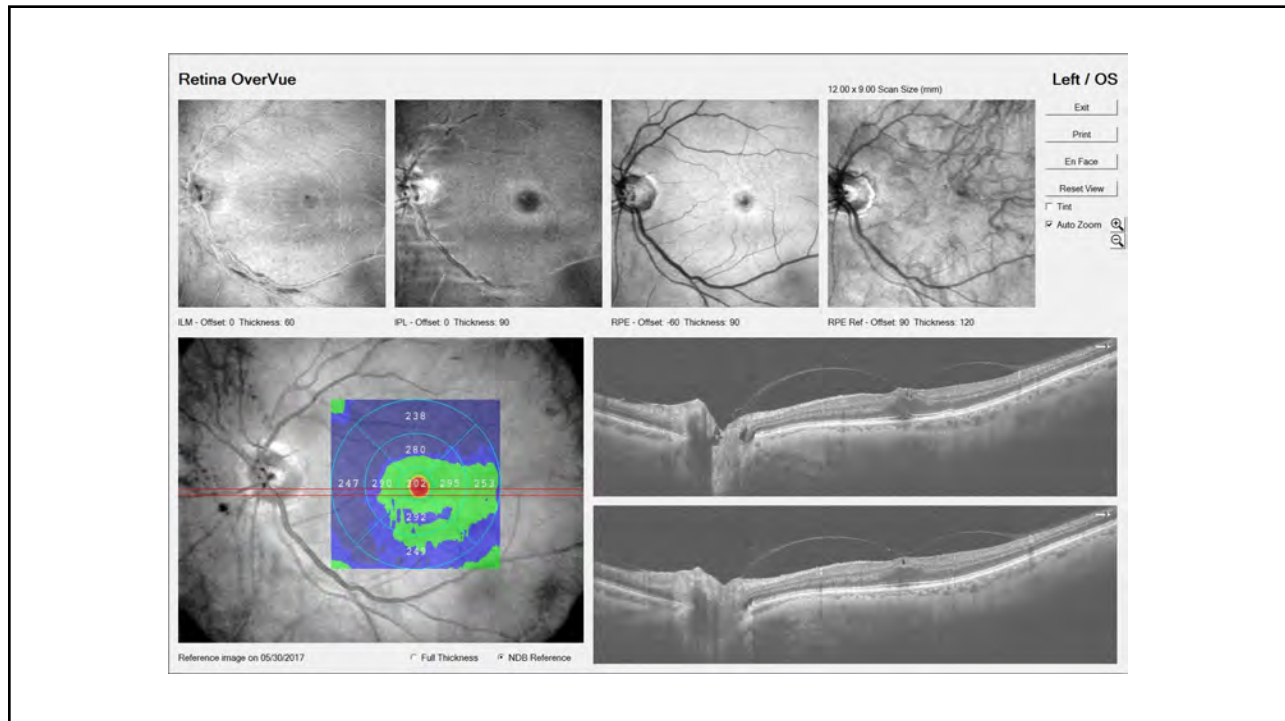
- Consider surgery if:
 - VA 20/40-ish or worse
 - Symptomatic
 - Visual need of patient
- Make sure you have an experienced surgeon!!

47

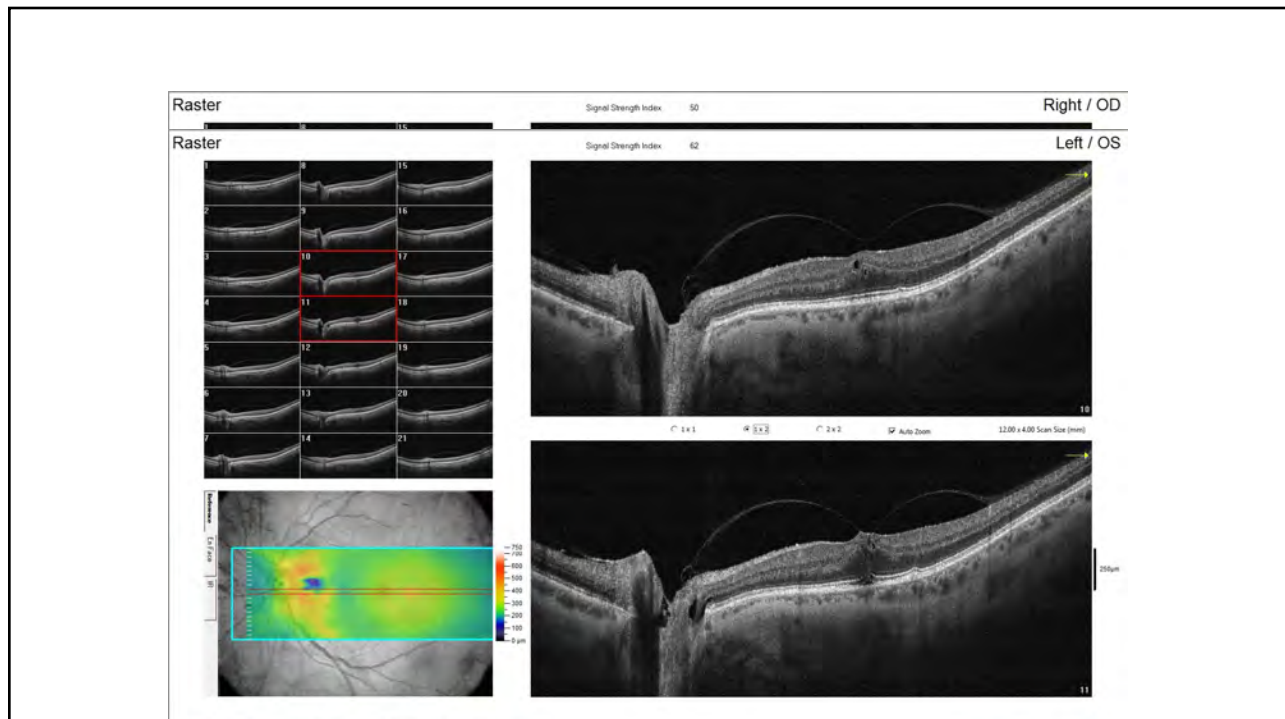
Case 7

- 70 year old male in for routine exam
- Notes mild change in distance vision, both eyes, since last exam 1 yr ago
- Thinks he needs new glasses
- 20/20 OD, 20/50 OS
 - Pt surprised that VA OS was decreased. Did not notice until exam today

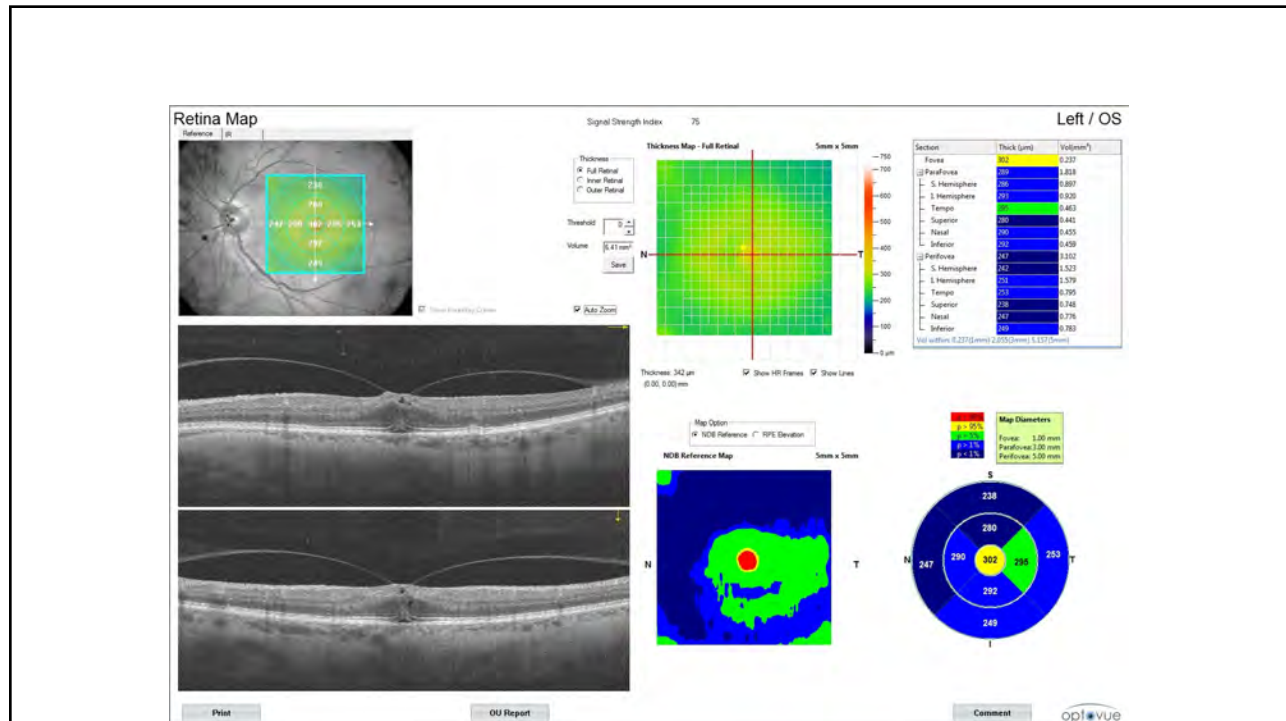
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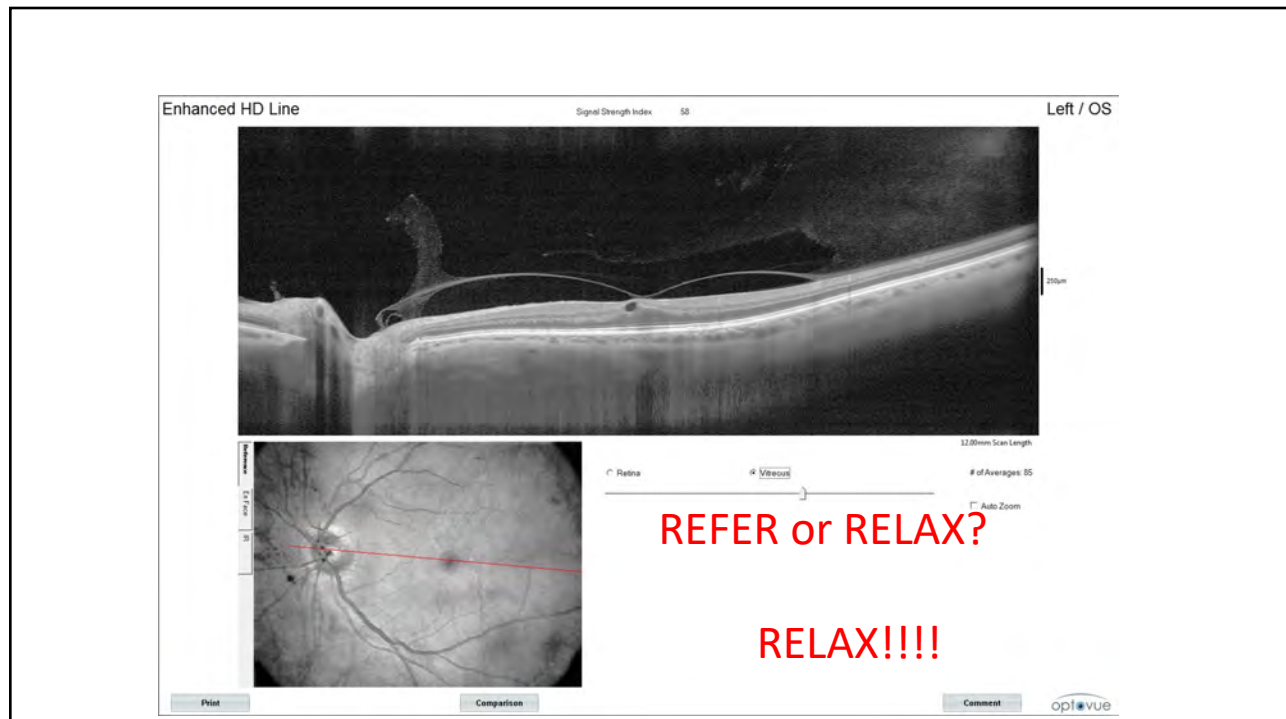
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50



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BRVO/CRVO

- Management includes diagnosis and management of underlying etiology
- Most often associated with DM and HTN
- However many other possible etiologies
 - Carotid artery disease
 - Hyperlipidemia/hypercholesterolemia
 - Altered platelet function
 - Coats disease
 - Von-Hippel Lindau
 - Eales' disease
 - Trauma

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BRVO/CRVO

- | | |
|---|--|
| <ul style="list-style-type: none"> • At minimum, should have <ul style="list-style-type: none"> • BP evaluated • Fasting Blood sugars (FBS)/A1c • CBC • Lipid profile | <ul style="list-style-type: none"> • Additional tests might include <ul style="list-style-type: none"> • Carotid artery evaluation • Cardiac evaluation • Additional blood tests <ul style="list-style-type: none"> • ANA • RF • FTA/ABS • ESR |
|---|--|

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Anti-VEGF:Lucentis

- CRUISE (CRVO) Study:
 - Vision improved > 15 letters in almost 50% of patients vs. 17% with sham at 6 mos
 - mean VA gain of almost 15 letters
- BRAVO (BRVO) Study:
 - Vision improved > 15 letters in over 60% of patients vs. 28% with sham
 - Mean VA gain of approx 18 letters
- Few side effects in either group

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Anti-VEGF: Eylea® (afilbercept)

- FDA approved Sept, 2012 for treatment of macula edema secondary to CRVO
- COPERNICUS and GALILEO studies:
 - % of pts gaining 15 letters or more of BCVA
 - Injection q 2 mos for 24 weeks
- COPERNICUS:
 - 56% vs. 12% with sham
 - 17.3 letters gained vs. 4.0 lost with sham
- GALILEO:
 - 60% vs. 22% with sham
 - 18.0 letters gained vs. 3.3 lost with sham

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CRVO/BRVO

- Refer if macula edema within 1 week
 - Laser vs. injection in BRVO
 - Injection CRVO
 - Steroids?
- Systemic workup recommended
 - DM
 - HTN
 - Cholesterol panel
 - Carotid Doppler
- Look for NV/NVI/NVA/NVG esp. in CRVO, esp. if ischemic



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KETCHUM UNIVERSITY**
Southern California College of Optometry

Optic Disc Edema

Jessica Chang, MD



Optic Disc Edema

Jessica R. Chang, MD
Clinical Assistant Professor of Ophthalmology

Disclosures

- Horizon Therapeutics Advisory Board
- Please do not share or copy patient images

Outline

- Papilledema
- Pseudopapilledema
- AION—arteritic and non-arteritic
- Diabetic papillopathy
- Atypical Optic neuritis, Optic Peri-neuritis
- Neuroretinitis
- Other causes
- Cases
- Summary



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Papilledema

- Optic disc edema from raised intracranial pressure (ICP)
- Edema tends to start nasally, wide range from mild to severe with hemorrhages and cotton wool spots
- May be asymmetric between the two eyes
- Visual symptoms:
 - Sometimes no visual symptoms, sometimes severe vision loss
 - Transient visual obscurations,
 - Enlarged blindspot on HVF
- Nonvisual symptoms:
 - whooshing pulsatile tinnitus, positional headaches



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Papilledema

- Workup:
 - Emergent neuroimaging to rule out mass lesion
 - LP to measure ICP
 - Medication and/or surgery to reduce ICP (and address any mass lesion if present)



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Pseudo-Papilledema

- With or without optic disc drusen
- May be asymptomatic or may cause VF deficits
- Drusen have been hypothesized to increase risk of NAION
- Buried drusen may become more apparent over lifetime
- Drusen diagnosed with Autofluorescence, FA, B-scan, EDI-OCT (and CT if calcified)



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Hickam's Dictum vs. Occam's Razor

"A patient can have as many diseases as he d--n well pleases"

- Keep an open mind for multiple diagnoses
- Remember to rethink an initial diagnosis if something doesn't fit
- One can have disc drusen AND raised ICP, or disc drusen AND glaucoma, for example

Anterior Ischemic Optic Neuropathy (AION)

- Arteritic (AAION): Giant cell arteritis, patient age >50yrs, F>M
 - Pallid disc edema, severe vision loss, may affect second eye shortly
 - ROS: headache, jaw claudication, weight loss, fever, cranial neuropathies
 - FA may show patchy choroidal filling; GCA may also present w/CRAO
 - Check CBC, ESR, CRP (not hs), consult rheumatology
 - Admit for IV methylprednisolone, then ~1 year of prednisone or steroid sparing alternatives such as tocilizumab
 - Temporal artery biopsy is gold standard for Dx but may be inconclusive still
 - Rarely other forms of arteritis/vasculitis may present similarly

Anterior Ischemic Optic Neuropathy (AION)

- Non-arteritic (NAION)
 - Disc at risk morphology, PMH of OSA, HTN, HL, DM, hypotension and/or vasodilating medications, anemia, smoking, migraine, coagulopathies, age usually >50yrs
 - May have any VF defect but classically inferior altitudinal corresponding to superior segmental hyperemic optic disc edema
 - 40% of pts have spontaneous improvement
 - 15-20% risk of NAION in the fellow eye in the next 5 years
 - No treatment other than modifying risk factors

Diabetic Papillopathy (formerly papillitis)

- Presents in patients with DM1 or 2, painless unilateral or bilateral disc edema, usually without severe vision loss
- Exact pathophysiology is unclear—some consider it a form of AION, but generally has less severe impact on final vision and different FA features
- No widely accepted treatment but anti-VEGF intravitreal injection has been published in case reports

(Atypical) Optic Neuritis

- In pediatric patients, isolated optic neuritis (ON) tends to present with disc swelling more often than adults who classically have retrobulbar ON (“Typical ON”)
- Adults with ON with disc edema should be worked up for infectious and inflammatory causes, based on PMH and risk factors
 - E.g. Lyme disease, syphilis, TB, Sarcoidosis, MOGAD, etc.



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Neuroretinitis

- Optic disc edema and macular star/hemistar with lipid exudates
- Usually unilateral, most have RAPD and central VF defect
- Infectious:
 - Bartonella henselae, cat scratch fever—can be associated with encephalitis
 - Also syphilis, Lyme, RMSF, toxoplasma, toxocara, etc
- Idiopathic—still often viral prodrome/URI; may be recurrent
 - Final vision better than 20/40 in >90% of patients



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Other assorted causes

- Toxic optic neuropathy (e.g. methanol)
- Optic perineuritis
- Hypertensive emergency
- Meningitis
- Infiltrative lesions (e.g. leukemia, sarcoid)
- CRVO



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CASES



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Summary

- Optic disc edema may result from a wide spectrum of disease, from anatomic anomaly to life-threatening emergency
- Thorough history and exam guide work up and management
- When in doubt please call/refer—if there is concern for raised ICP safest route is ER for emergent imaging



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Thank you!

- Questions?
- Jessica.chang@med.usc.edu
- 323-442-6335 (office)
- My clinic locations:
 - USC HSC Campus east of downtown
 - USC Roski in Arcadia



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Herpetic Eye Disease

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Herpetic Uveitis Management

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


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
No relevant financial disclosures



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Herpes family viruses account for a large proportion of infectious uveitis

Table 4. Causative organisms of infectious ocular inflammation by diagnosis and medication codes.

Specified by ICD-9 Diagnosis Code	Sample size, n
Parasite	
<i>Histoplasma</i>	5976
<i>Toxoplasma</i>	1104
Other	28
Viral	
VZV	1129
HSV	442
Other	577
Unspecified	
infectious endophthalmitis	1610
infectious iridocyclitis	733
Mycobacterial	36
Syphilis	36
Specified by NDC or J Medication Code	
Viral	3854
Bacterial	2943
<i>Toxoplasma</i>	786
Fungal	542
Mycobacterial	49
Parasite (excluding toxoplasma)	33

ICD = International Classification of Diseases; VZV = varicella zoster virus; HSV = herpes simplex virus
NDC = National Drug Code

<https://doi.org/10.1371/journal.pone.0237995.t004>

PLOS ONE




RESEARCH ARTICLE

Incidence, prevalence, and risk factors of infectious uveitis and scleritis in the United States: A claims-based analysis


Youning Zhang¹, Sarina Amin¹, Kristina I. Lung², Seth Seabury², Narsing Rao¹, Brian C. Toy^{1*}

¹ Department of Ophthalmology, Roski Eye Institute, Keck School of Medicine, University of Southern California, Los Angeles, California, United States of America, ² Leonard D. Schweitzer Center for Health Policy & Economics, University of Southern California, Los Angeles, California, United States of America


(Zhang et al, PLOS One 2020)

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


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


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The Wages of SIN! by Keith Brown



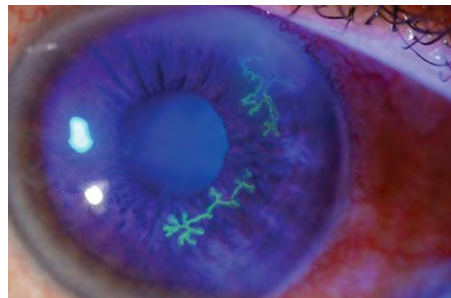
Objectives

- Recognize the clinical presentation of herpetic uveitides
- Review updates in the medical and surgical management of herpetic uveitis

4

Case: 60M painful right facial vesicular rash for 1 week

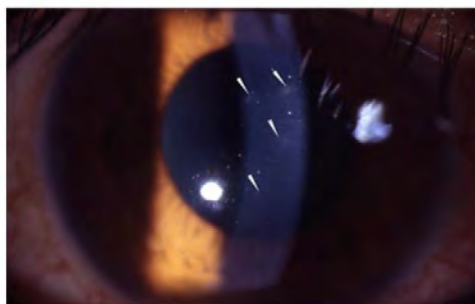


5

Herpetic keratouveitis can have multiple phenotypes

- unilateral, high IOP, iris atrophy
- stellate KP (diffuse, not in Arlt triangle)
- VZV > HSV-1 > HSV-2 >> CMV >> EBV

Other clinical manifestations include epithelial/stromal/endothelial keratitis with sequelae including immune ring (CMV), decreased corneal sensation and scarring (HSV and VZV), and corneal neovascularization



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Herpesvirus serologies have limited utility in uveitis evaluation

- HSV-1 seroprevalence increases from 54% in adolescence to 90% after age 50
- Thus, positive IgG not helpful in confirming diagnosis, and positive IgM does not prove ocular infection
- Negative viral serology may be helpful in excluding viral etiology
- Aqueous PCR demonstrates over 90% sensitivity and specificity

(Pleyer and Chee, Clin Ophthalmol 2015)



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Landmark HEDS results guide clinical management

- Stromal keratitis:
 - Topical steroids speed resolution of stromal keratitis, with slow taper to minimize recurrence. They are contraindicated in the setting of epithelial keratitis. (Wilhemus et al, Ophthalmology 1994)
 - Oral acyclovir did not improve acute outcomes (Barron et al, Ophthalmology 1994) but did reduce rates of recurrence by 50% (Wilhemus et al, NEJM 1998)
- Anterior uveitis
 - Oral acyclovir may improve treatment success in conjunction with topical steroids (HEDS Study Group, Arch Ophthalmol 1996)

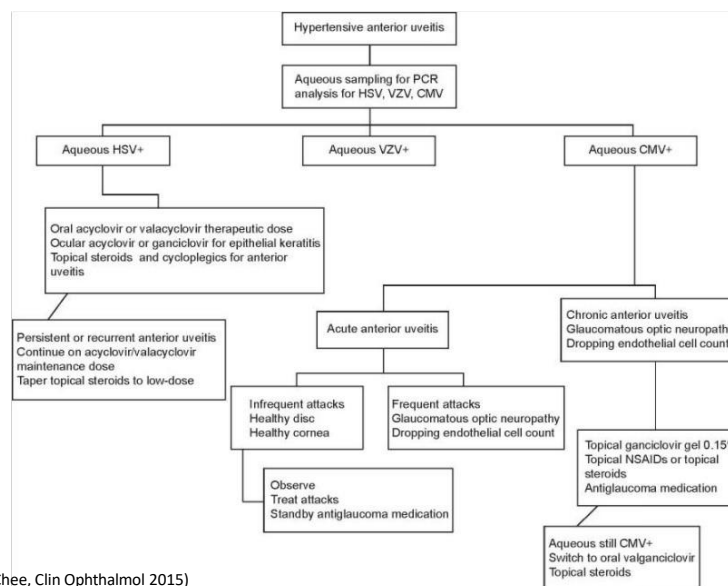


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Modern treatment considerations

- Oral antivirals preferred over topical treatment
- In patients unable to tolerate oral tx (renal dz), topical ganciclovir gel 0.15% less toxic than trifluridine used in HEDS study
- When topical steroids are indicated, consider starting no higher than QID to decrease risk of recurrence with tapering
- In chronic or recurrent inflammation, consider maintenance systemic antivirals + low-dose steroid drops

Herpetic Keratouveitis Management Summary

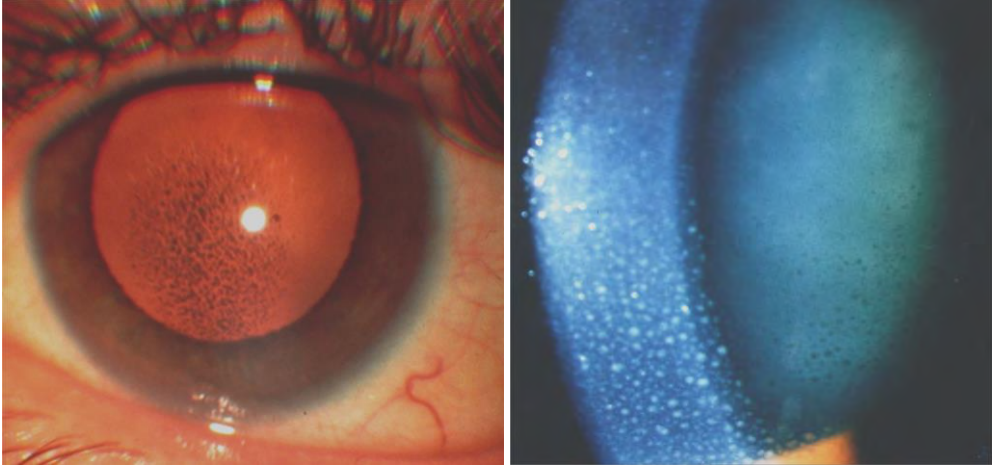


(Pleyer and Chee, Clin Ophthalmol 2015)

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Case: 55M with ocular pain, photophobia and redness and decreased vision OS. VA was 20/20 and 20/50; IOP 10 and 31; Large KPs; 3+cells in AC and **3+ cells in vitreous**



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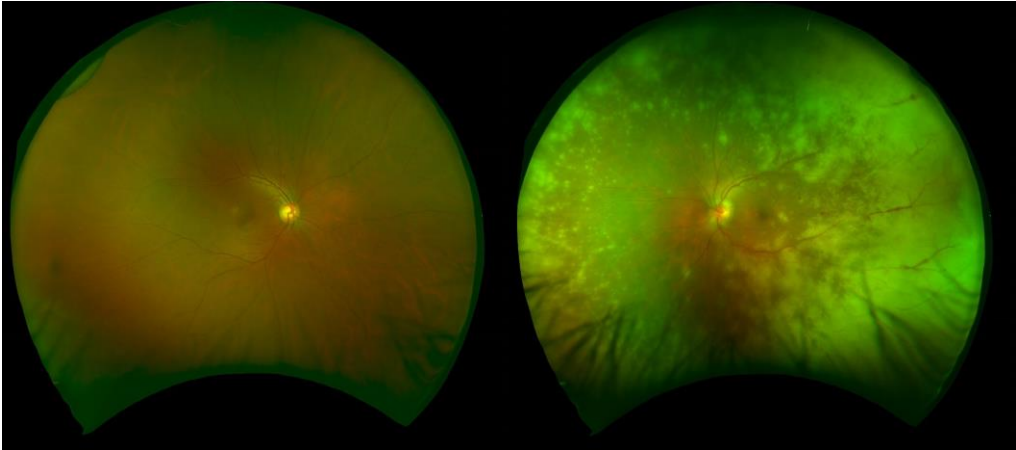
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20/20 20/50

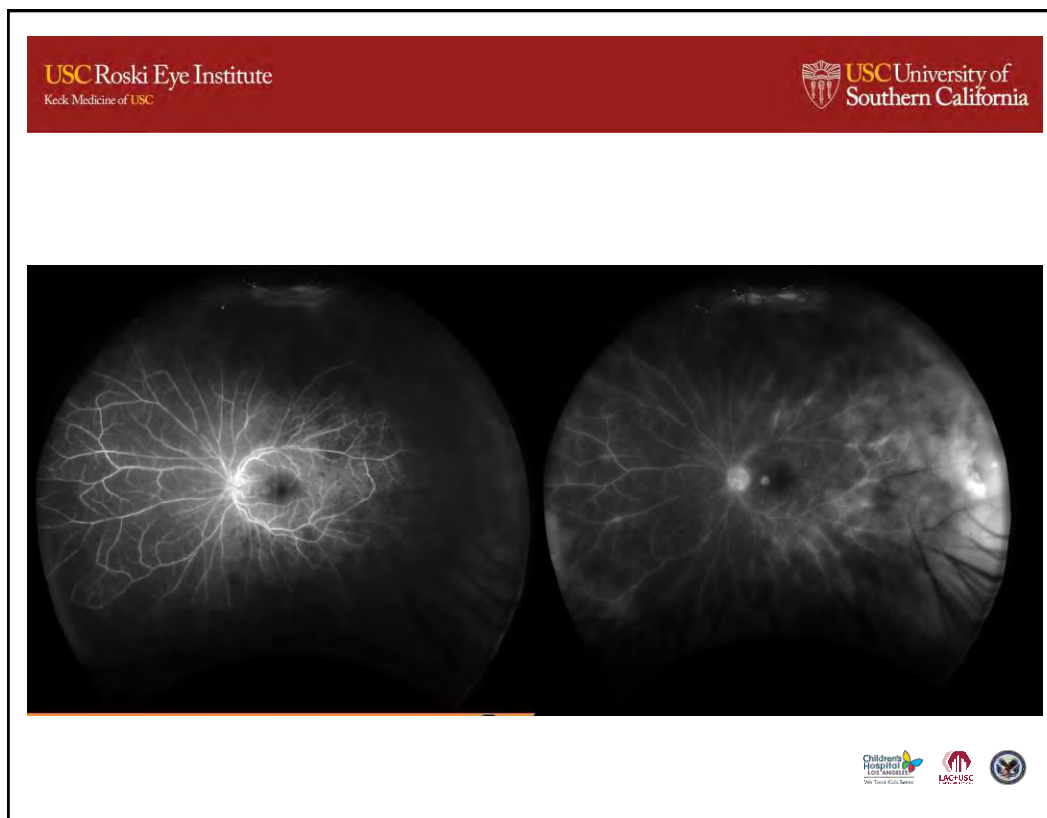


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ARN generally presents as an acute unilateral panuveitis in immunocompetent patients

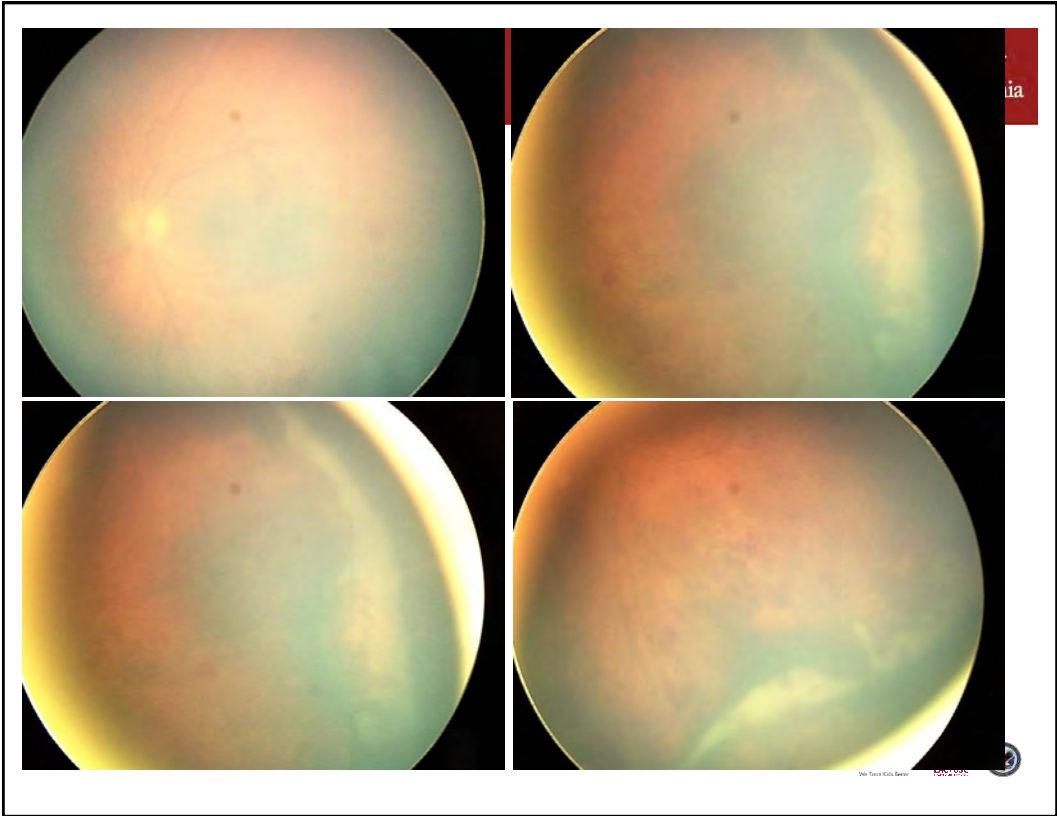
- Sx: acute onset blurred vision, floaters, pain, photophobia
- Initially unilateral, but can become bilateral in 36% of patients within 6 weeks
- VZV > HSV-1 > HSV-2 >> CMV >> EBV
- DDx:
 - CMV retinitis
 - Syphilis
 - Atypical toxoplasmosis
 - Behçet

(Fisher et al, Ophthalmology 1992)

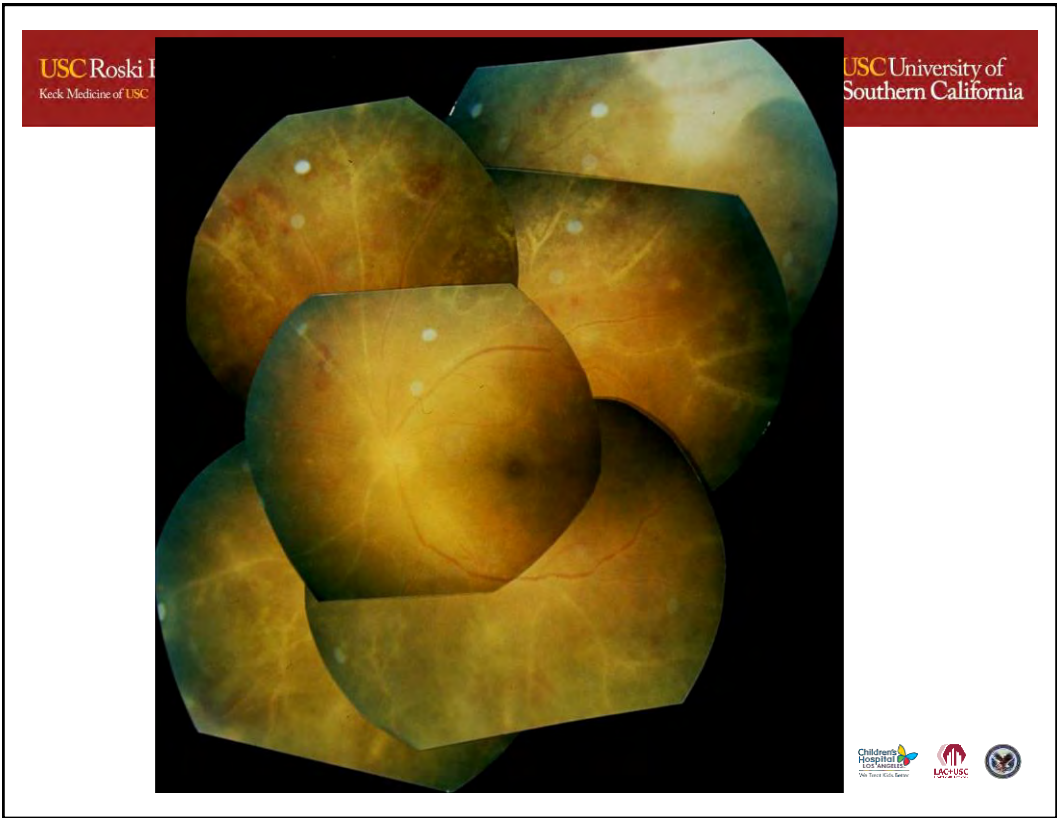
Children's Hospital Los Angeles
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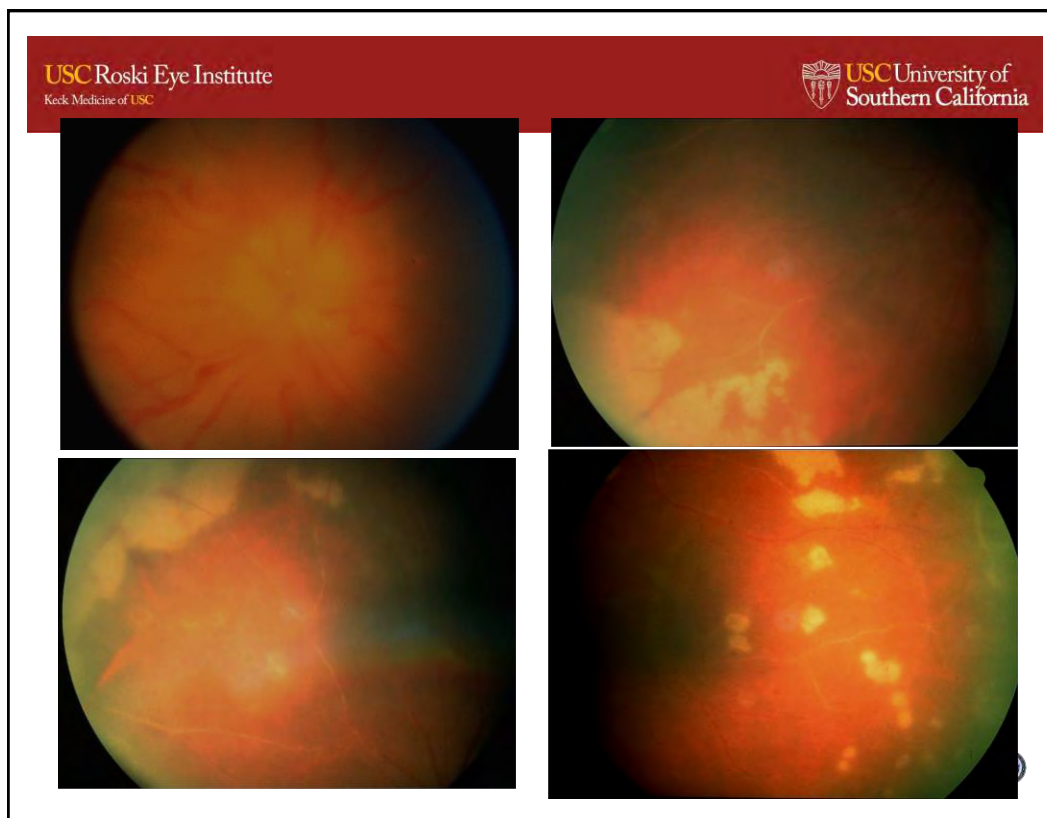
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ARN is primarily a clinical diagnosis

AUS diagnostic criteria

- Peripheral retinal necrosis with circumferential spread
- Rapid progression in absence of antiviral therapy
- Occlusive arteriolitis
- Prominent AC rxn, vitritis
- +/- scleritis, optic neuropathy

(Holland/AUS Executive Committee, AJO 1994)

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AC PCR can be a useful adjunctive diagnostic test

- AC PCR sensitivity and specificity > 95% for HSV/VZV/CMV viral retinitis
- Prognostication value: VZV>HSV ARN demonstrated greater loss of vision, higher risk of final VA worse than 20/200, and 2.5-fold increased risk of secondary RD
- Procedure:
 - Patient supine, aseptic technique
 - Indirect ophthalmoscope for illumination and magnification
 - 0.05-0.1ml collected using a 30g needle/1cc syringe via the limbus, over iris

(Takase et al, Jpn J Ophthalmol 2015;
Blumenkranz et al, Retina 1989;
Wong et al, Ophthalmology 2010)



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Systemic antivirals form the mainstay of ARN treatment

Induction (~10 days)

- Inpatient admission for acyclovir IV 10mg/kg q8h
- ValACV 2g PO TID
- Famciclovir 500mg PO TID

Treatment (~6 weeks)

- ACV 800mg PO 5x daily
- ValACV 1g PO TID
- Famciclovir 500mg PO BID

Maintenance/Secondary Prophylaxis (3-6 months)

- ACV 800mg PO TID
- ValACV 1g PO daily
- Famciclovir 250mg PO BID

Adjunctive treatments:

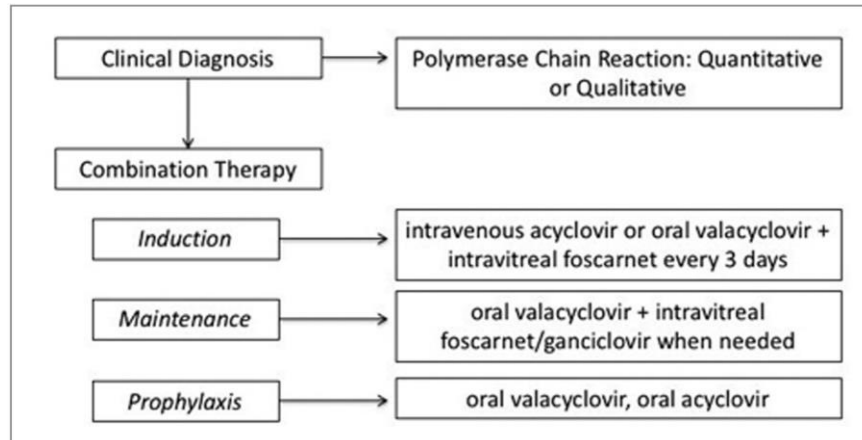
- Intravitreal ganciclovir 4mg or foscarnet 2.4mg
- Oral corticosteroids (prednisone 0.5mg/kg/day) starting 24-48 hours after initiating antivirals
- ASA 325

(Tibbetts et al, Ophthalmology 2010; Schoenberger et al, Ophthalmology 2017)



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ARN Management Summary



(Li et al, OSLI Retina 2019)

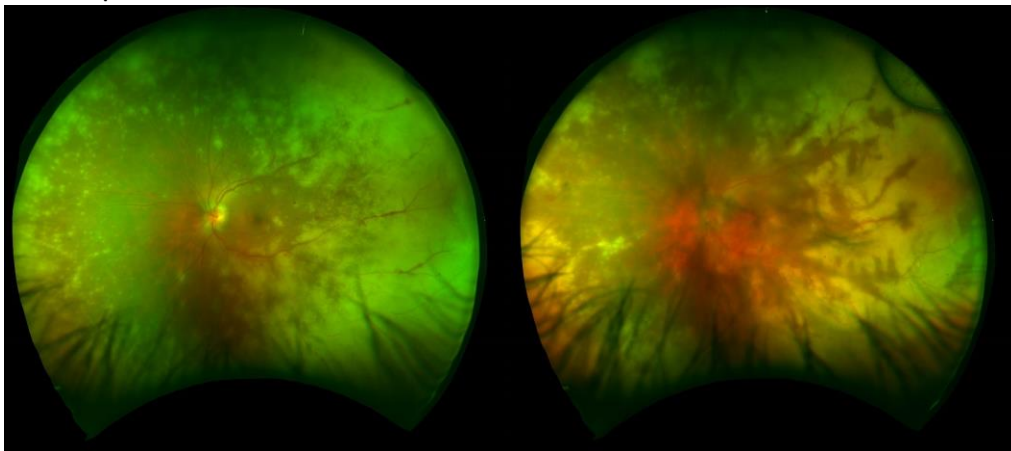


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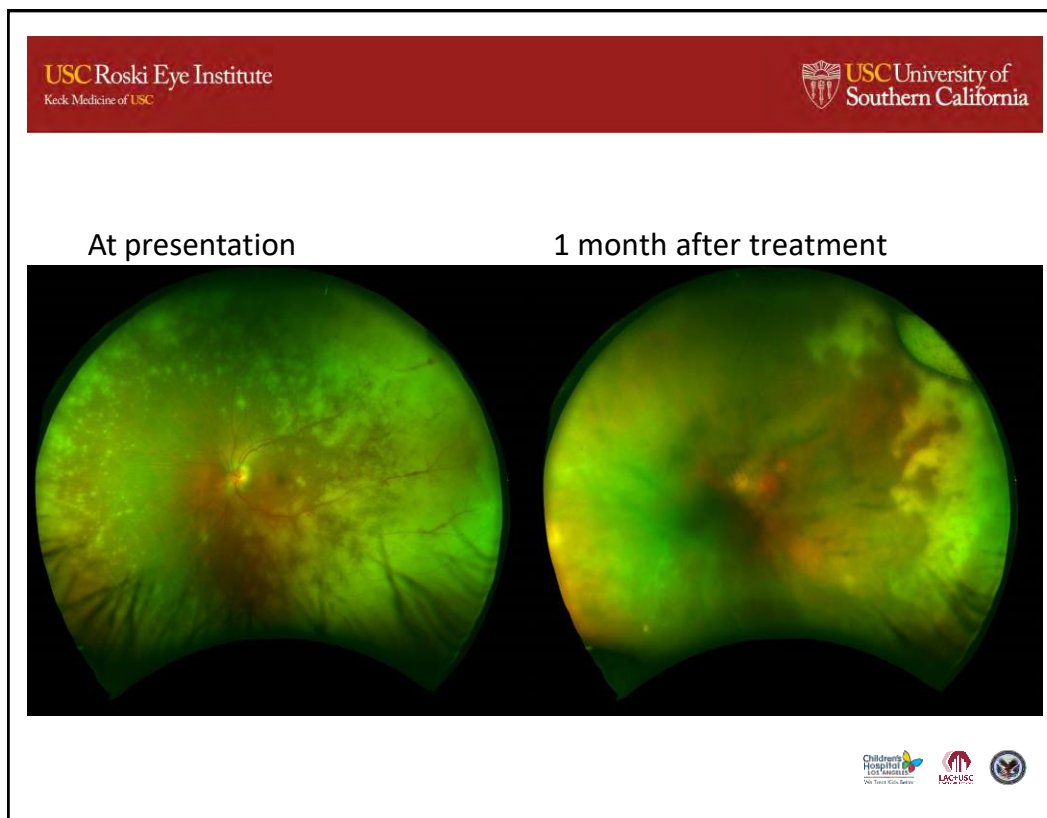
Back to our patient...

At presentation

1 week after treatment



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
Sequelae of ARN cause significant visual morbidity

- RRD (24% in HSV, 62% in VZV), usually within 1-2 months
 - 40% risk reduction with IVI antivirals (VZV)
 - Prophylactic laser barricade or early vitrectomy have not demonstrated efficacy
 - Surgical management usually involves scleral buckling, vitrectomy, and silicone oil tamponade
- Optic atrophy
- Contralateral eye involvement is seen in up to 36% of cases and usually occurs within 6 weeks

(Hillenkamp et al, Ophthalmology 2009;
Lau et al, Ophthalmology 2007)

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



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Acute Retinal Necrosis

	HSV	VZV
Eyes	33	48
Age	34	51
1 year VA better than 20/60	52%	35%
1 year VA worse than 20/200	35%	60%
Retinal Detachment	24%	62%


Intravitreal Foscarnet 40% lower rate of RD in VZV group

(Wong et al, Ophthalmology 2010)

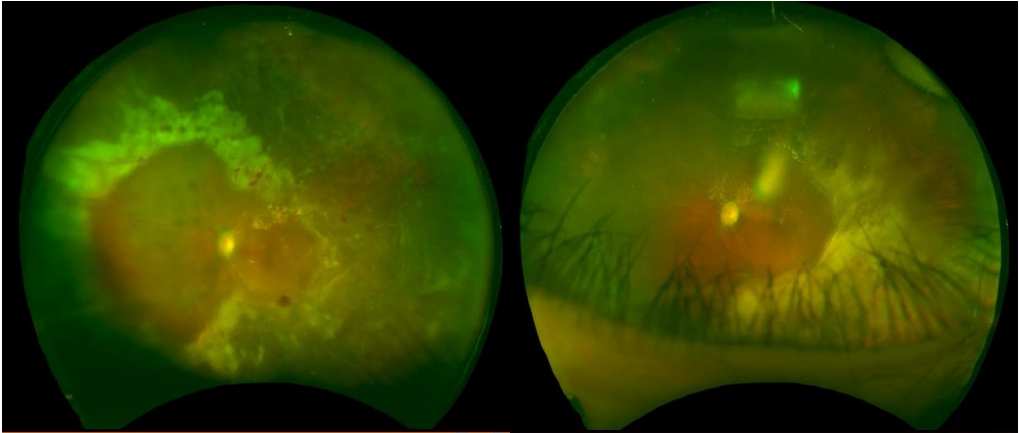
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


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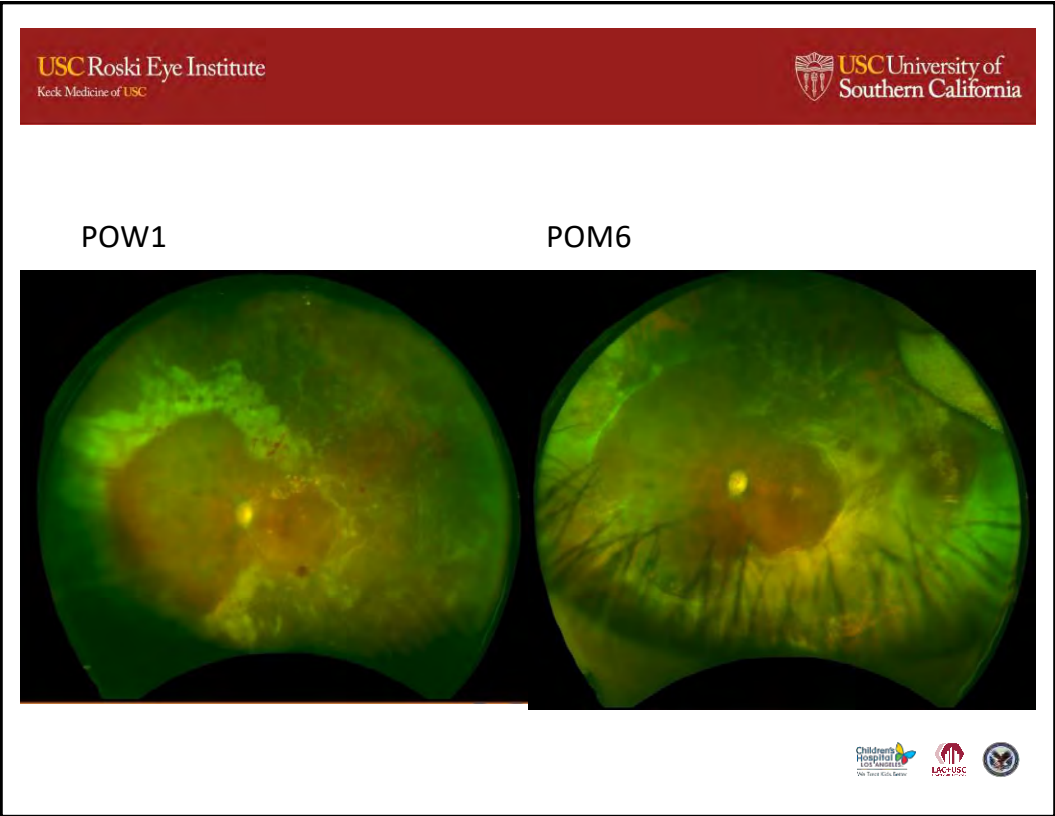
POW1 after SB/PPV/EL/SO

POM3

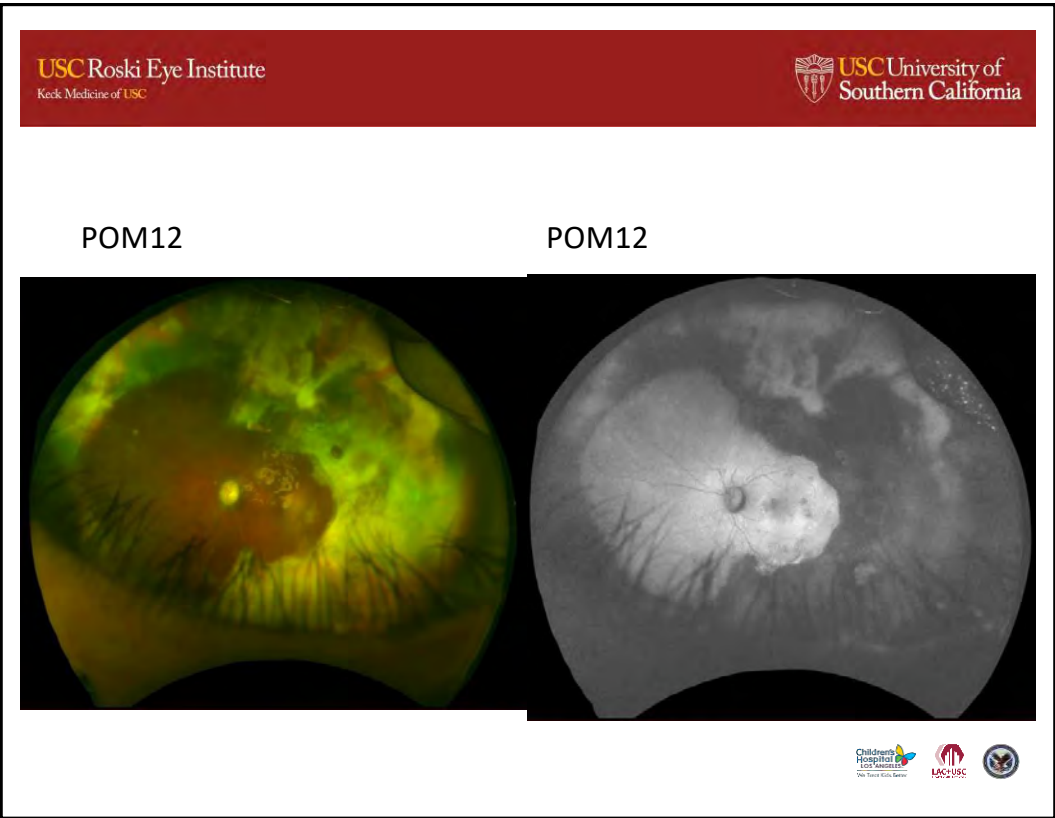


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27



28

ARN Management Summary

- AC PCR has an important diagnostic and prognostic role
- Treatment includes combination systemic antivirals with intravitreal therapy to decrease subsequent risk of RD
- Adjunctive PO steroids and aspirin may be beneficial to minimize sequelae of vitreous condensation and occlusive vasculitis
- Monitor disease activity with exam findings and changes in autofluorescence, secondary RD is common
- Surgical management of ARN-associated RD: SB/PPV/SO

(Li et al, OSLI Retina 2019)

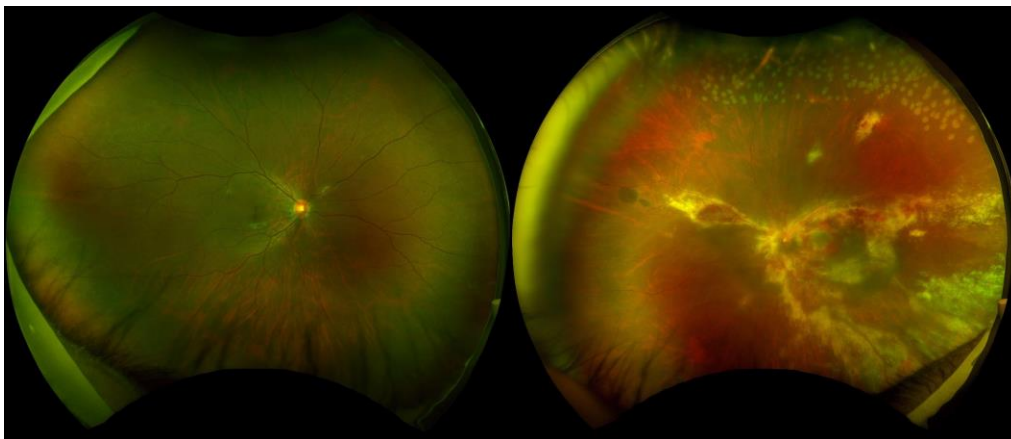


29

Case: 30 year old man newly diagnosed HIV, VL 300k, CD4 36

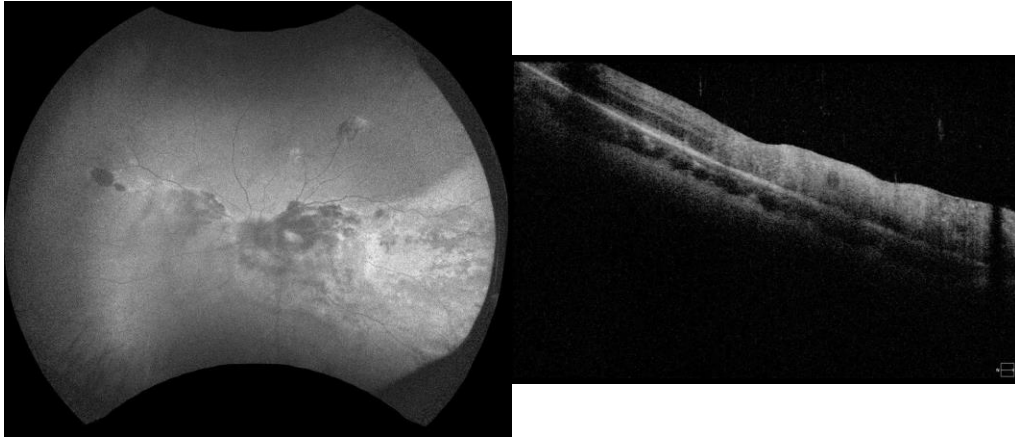
OD 20/20

OS CF



30

CMV Retinitis: OS



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CMV Retinitis

- Immunocompromised host
 - AIDS, CD4<50
 - Immunosuppression post-transplant
- 3 patterns
 - classic / fulminant retinitis
 - posterior pole
 - hemorrhagic
 - granular / indolent retinitis
 - peripheral
 - minimal heme
 - frosted branch / perivascular

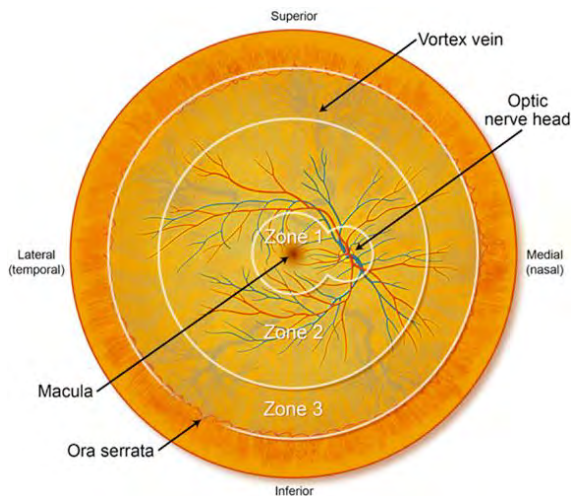


(Images courtesy of Dr. Damien Rodger)

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Staging disease

- **Zone 1:**
1500um (1 disc) diameter surrounding optic nerve and 3000um diameter around the fovea
- **Zone 2:**
Anterior to Zone 1 and posterior to vortex veins
- **Zone 3:**
Anterior to Zone 2 and posterior to ora



(Holland et al, Arch Ophthalmol 1989)



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Systemic antivirals form the mainstay of CMV retinitis treatment

- Diagnosis: clinical, AC PCR
- Treatment: induction and maintenance
 - IV ganciclovir / PO valganciclovir (myelosuppression)
 - IV foscarnet (renal toxicity)
 - IV cidofovir (anterior uveitis, hypotony)
 - PO letermovir (rescue therapy in transplant patients)
 - ~~intravitreal implant ganciclovir (sustained release 6-8 months)~~
 - Serial intravitreal injections ganciclovir or foscarnet
 - semiweekly injections x 3 weeks, then weekly
- ARV in HIV patients
 - Active retinitis recurs in 50% patients within 6-8 weeks of maintenance tx without ARV
 - Can consider d/cing maintenance tx once on ARV, CD4>100 q6months x 2, no active lesions



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Systemic antivirals form the mainstay of CMV retinitis treatment

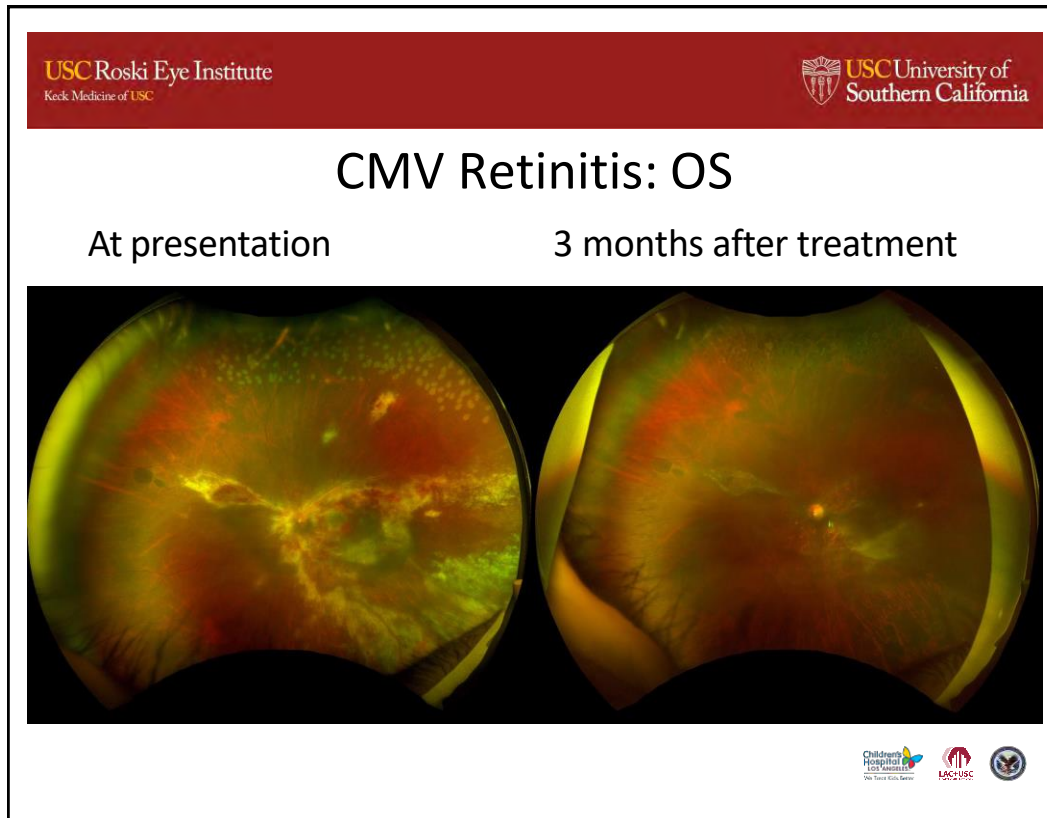
- IV ganciclovir
 - Induction: 5mg/kg BID x 21d
 - Maintenance: 5mg/kg daily
- PO valganciclovir
 - Induction: 900mg BID x 21d
 - Maintenance: 900mg daily
- PO letermovir
 - Maintenance: 480mg PO daily
- IV foscarnet
 - Induction: 60mg/kg TID x 21d
 - Maintenance: 90-120mg/kg daily

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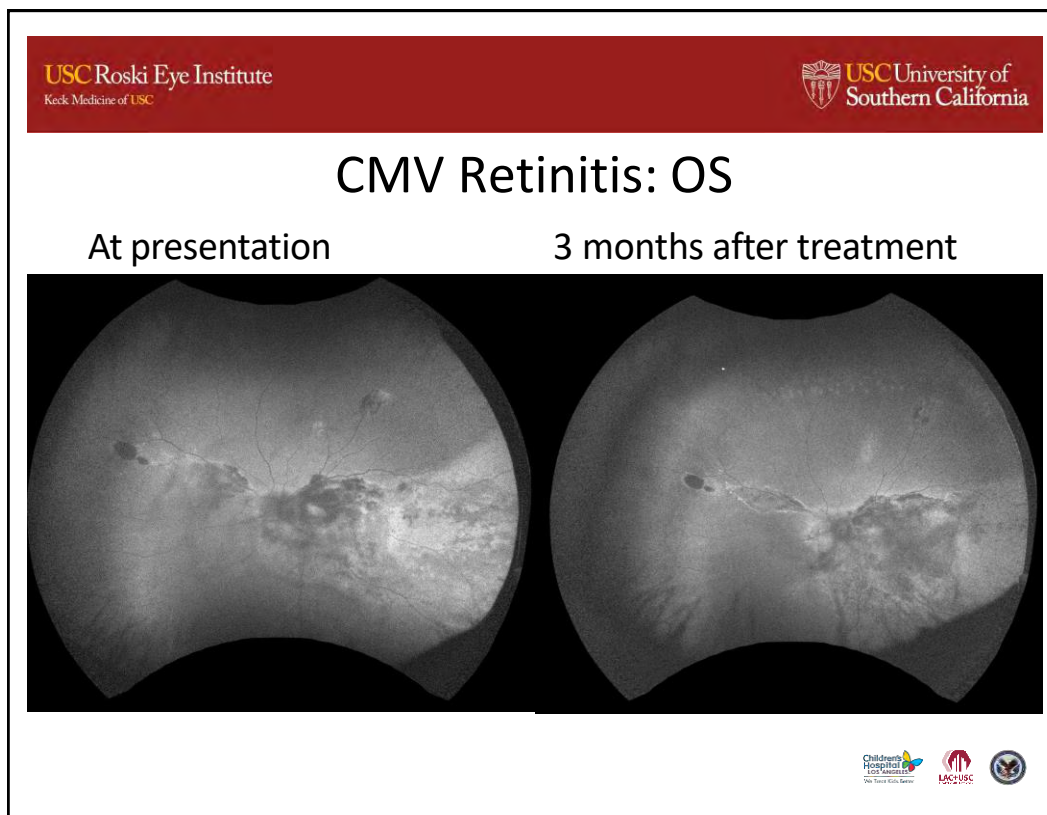
Intravitreal therapy is a useful adjunct for CMV retinitis treatment

- Ganciclovir
 - Nucleoside analogue that inhibits viral polymerases
 - 2-3 mg x 2 injections/week then maintenance of weekly injections
- Foscarnet
 - Pyrophosphate analog which inhibits viral polymerases
 - 1.2 - 2.4 mg x 2 injections/week then maintenance of weekly injections
- No comparison studies between foscarnet and ganciclovir have been reported, however there have been multiple case reports showing the use of foscarnet in ganciclovir resistant CMV cases.
- No differences in visual outcomes and CMV viral load in the aqueous between ganciclovir monotherapy or combined ganciclovir and foscarnet.

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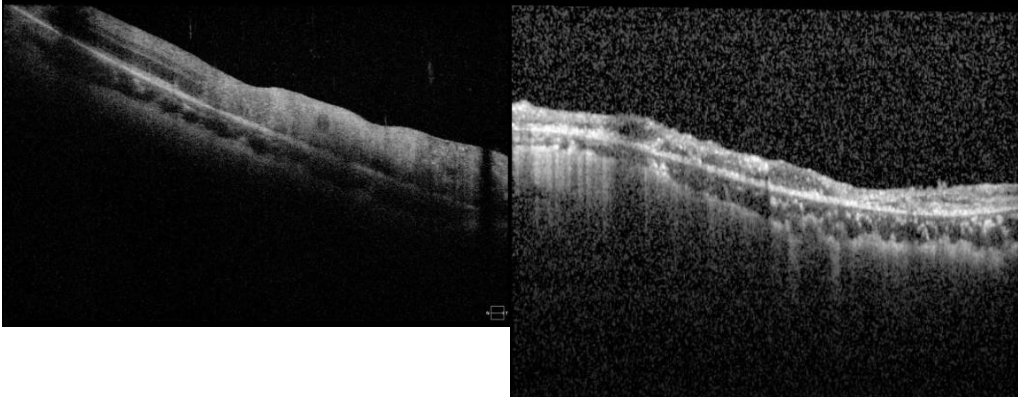
38

USC Roski Eye Institute
Keck Medicine of USC

USC University of Southern California

CMV Retinitis: OS

At presentation 3 months after treatment



Children's Hospital Los Angeles
We Treat Kids Better

LAC+USC

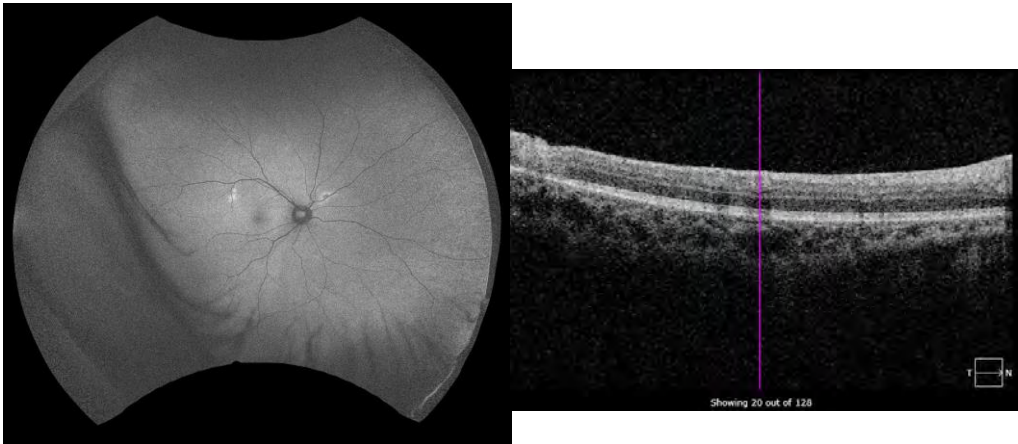
USC

39

USC Roski Eye Institute
Keck Medicine of USC

USC University of Southern California

CMV Retinitis: OD



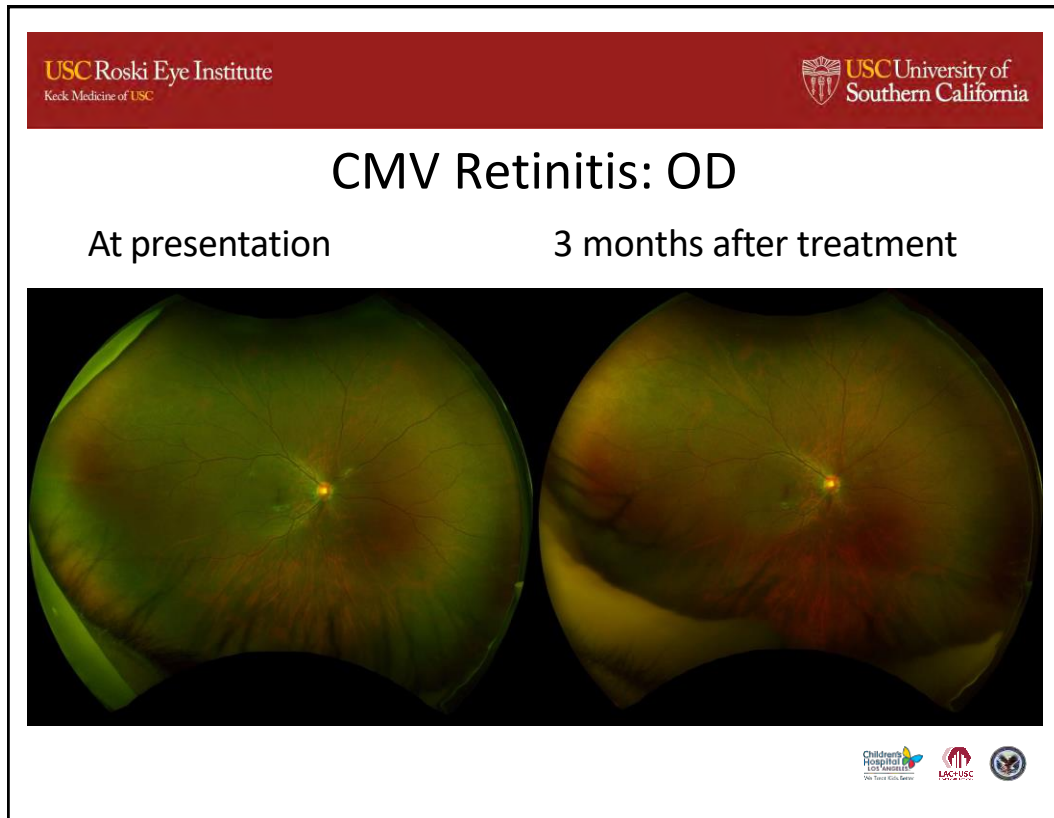
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Children's Hospital Los Angeles
We Treat Kids Better

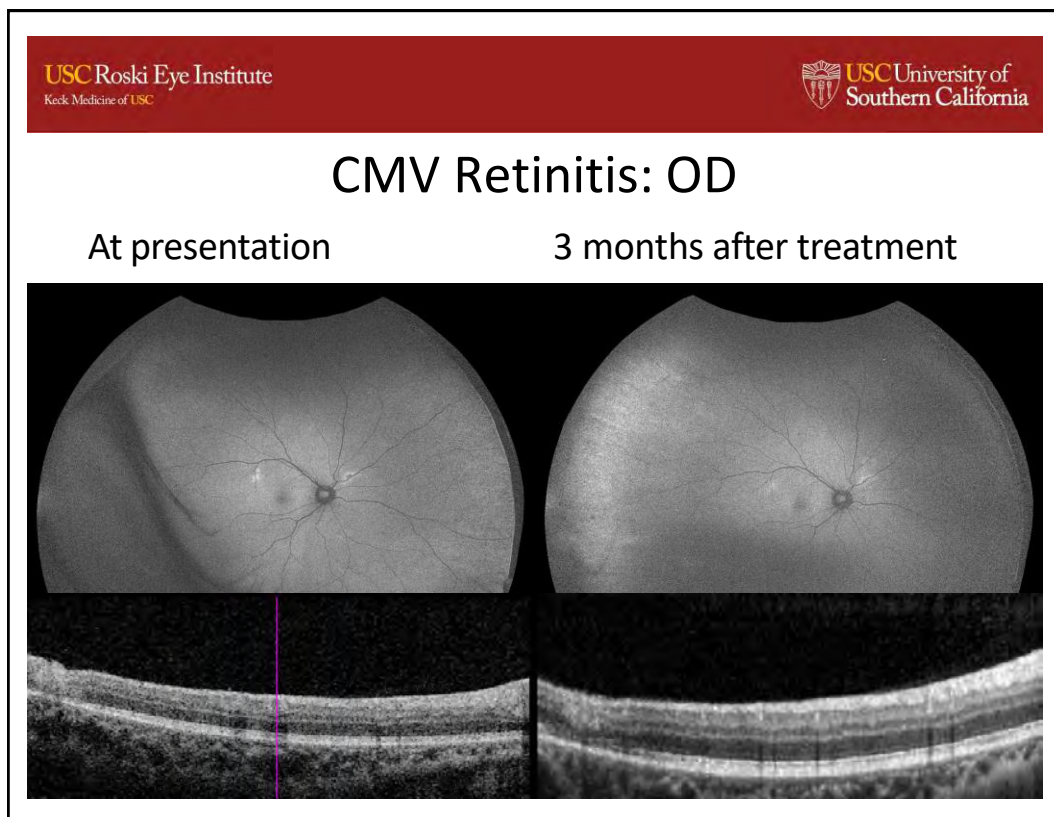
LAC+USC

USC

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Immune recovery uveitis is common among patients starting HIV therapy

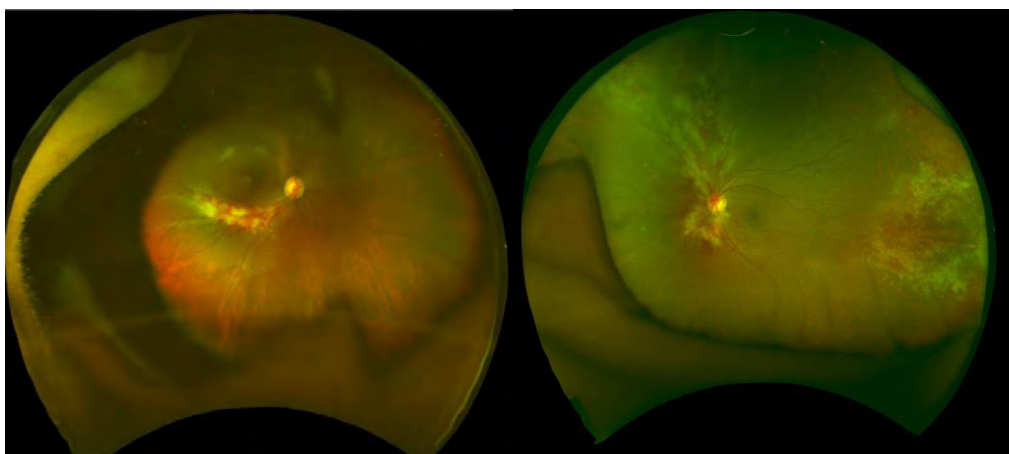
- As the CD4 counts rise with ARV, cell-mediated response to CMV antigens cause anterior or intermediate uveitis
- Risk can be reduced by starting ARV after completion of CMV induction
- Manage IRU with topical steroids

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CMV retinitis: another case

OD

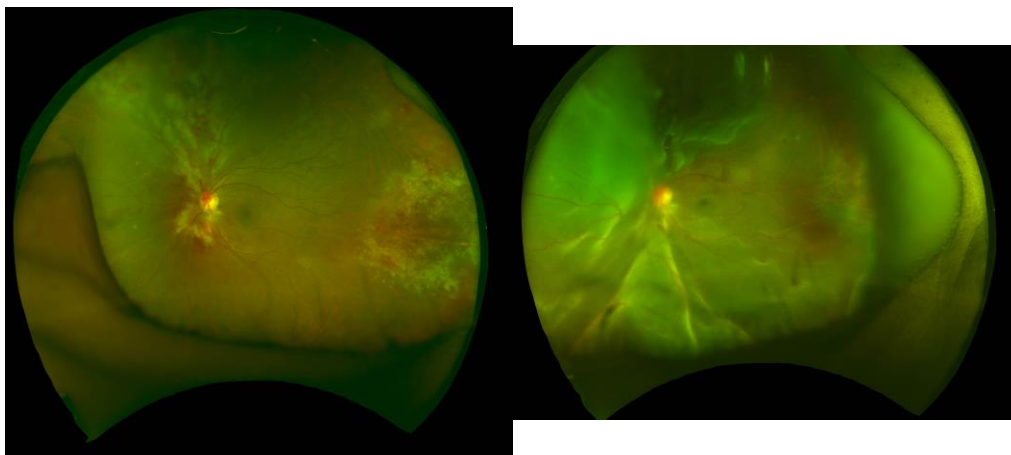
OS



44

CMV retinitis: another case

1 month after initiating treatment



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Retinal detachment can occur in 40% of patients with CMV retinitis

- Breaks occur near thin, necrotic retina
- Risk factors for RRD include
 - Larger lesion size
 - Zone 3 involvement
 - Bilateral disease
- PPV ± SB ± SO / Gas tamponade ± clear lens extraction

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Results of Rhegmatogenous Retinal Detachment Repair in Cytomegalovirus Retinitis with and without Scleral Buckling

Roberto F. García, MD, Marisa Flores-Aguilar, MD, Jose I. Quiceno, MD, Edmund V. Capparelli, PharmD, David Munguia, Baruch D. Kuppermann, MD, PhD, Fernando Arevalo, MD, William R. Freeman, MD

- 22 eyes with PPV/no buckle
 - 86% total reattachment rate
 - 91% macular reattachment rate
- 56 eyes with PPV/buckle
 - 84% total reattachment rate
 - 91% macular reattachment rate
- Unbuckled eyes had 1 D more of hyperopia
- No difference in visual outcome
- *in this study, the SB eyes had a larger percentage of macular involving RDs (59% vs 45%)

(Garcia et al, Ophthalmology 1995)



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Surgical repair of cytomegalovirus-related retinal detachment without silicone oil in patients with AIDS.

Canzano JC¹, Morse LS, Wendel RT.

- 6 eyes in 5 patients with CMV related RDs without SO
- RDs were repaired with SB/PPV/EL/Gas
- 83% of patients had total reattachment; macular reattachment in all eyes.
- Preoperative vision ranged from 20/40 to HM
- Mean postoperative visual acuity was 20/40
- Mean postoperative follow-up was 12 months; one patient detached 7 months after initial repair and was reattached without SO.

(Canzano et al, Retina 1999)

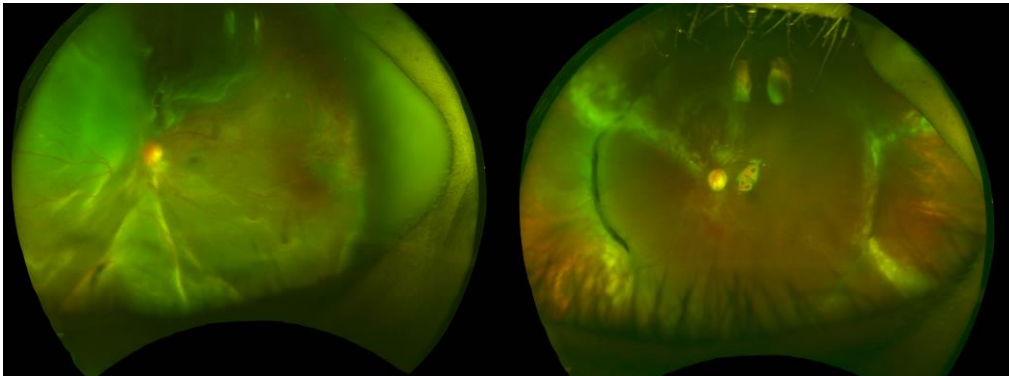


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The patient underwent SB/PPV/EL/SO

Pre-op

POW1



49

POM6

- Doing well, retina attached under oil
- 20/50 | CF ft (3+ PSC OS)
- Combined CEIOL/SOR associated with high risk of redetachment

TABLE 3. Summary of Additional Procedures Performed on Patients With Cytomegalovirus Retinitis Related Retinal Detachments who Underwent Surgical Repair Consisting of Trans Pars Plana Vitrectomy and Oil, and Then Later Underwent Additional Surgery Consisting of Trans Pars Plana Vitrectomy and Oil Removal

	All Eyes (n = 15)	Detached (n = 8)	Attached (n = 7)	P Value*
Additional Procedures at Time of Initial				
RD Repair				
Scleral buckle	4 (27%)	1 (13%)	3 (43%)	.28
Additional Procedures at Time of Oil				
Removal				
Scleral buckle	7 (47%)	5 (63%)	2 (29%)	.31
ERM peel (total # of pts)	3 (20%)	1 (13%)	2 (29%)	.57
Cataract extraction	8 (53%)	7 (88%)	1 (14%)	.01
Pars plana lensectomy	5	5	0	
Phacoemulsification	3	2	1	
Gas tamponade	13 (87%)	7 (88%)	6 (86%)	>.99
C3F8 gas	9	5	4	
SF6 gas	4	2	2	
No gas	2	1	1	

ERM = epiretinal membrane; RD = retinal detachment.
*Fisher's exact test.

(Morrison et al, AJO 2015)

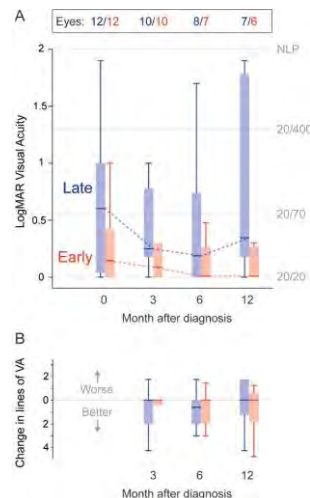


50

Early detection is key to reducing morbidity from CMV retinitis

Active CMV retinitis is often asymptomatic

- Patients with CD4 <50 cells/mm³ should be seen at least every 3 months to screen for CMV retinitis (AAO Preferred Practice Pattern)
- Earlier detection of CMV retinitis within 4 months after CD4<100 resulted in better visual outcomes
- Role for widefield tele-ophthalmology screening



(Ausayakhun et al, BJO 2018)



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CMV Retinitis Management Summary

- Evaluate for etiology of immunosuppression in patients with CMV retinitis
- Screen for other opportunistic infections
- Systemic antivirals form the mainstay of CMV retinitis treatment, with adjunctive intravitreal therapy
- Antivirals are virostatic, so longstanding control of CMV will depend on reconstitution of cellular immunity, which may be accompanied by immune reconstitution uveitis
- Monitor disease activity with exam findings and changes in autofluorescence
- Surgical management of CMV retinitis-associated RRD is patient-dependent and options include:
 - Laser barricade
 - PPV ± SB ± SO / Gas tamponade ± clear lens extraction



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THANKS!

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**Marshall B.
KETCHUM UNIVERSITY**
Southern California College of Optometry

The Calm in the Eye of the Storm: Re-Purposed Medications for COVID-19

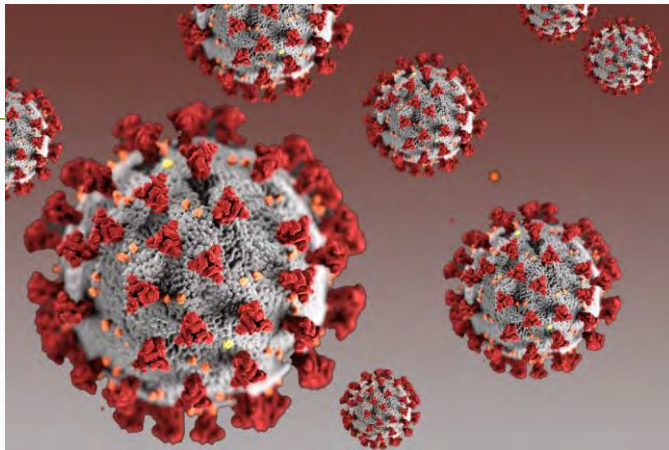
Judy Tong, OD



The Calm in the Eye of the Storm: Re-Purposed Medications for COVID-19

Judy Tong, OD, FAAO
Associate Professor
Assistant Dean of Residencies

1



2

Financial Disclosures

- I have no financial interest in any drug companies, medical or ophthalmic products that are discussed in this presentation.



3

Lecture Objectives

- Timeline
- Epidemiology – Incidence
- COVID-19 Infection – Symptoms
- Eye Involvement
- Cytokine Storm
- Vaccinations
- **Re-Purposed Medications**

4

Timeline

- **January 19, 2020** - First case confirmed in Washington State
- **January 31st** - WHO issues global health emergency
- **February 3rd** - US declares public health emergency
- **March 6th** - 21 passengers test positive on cruise ship off of the SF bay
- **March 11th** - WHO declares COVID-19 pandemic
- **March 13th** – Former president declares COVID-19 a national emergency and travel ban
- **March 19th** – California issues statewide stay at home order

<https://www.ajmc.com/view/a-timeline-of-covid19-developments-in-2020>

5

Epidemiology – Incidence

(worldometer.info)

Last updated: April 22, 2021, 03:52:08ET



United States

Coronavirus Cases:

31,918,591

Deaths:

575,829

Recovered:

24,480,522

6

California Stats

Now Yesterday Search:

#	USA State	Total Cases	New Cases	Total Deaths	New Deaths	Total Recovered	Active Cases	Tot Cases/ 1M pop	Deaths/ 1M pop	Total Tests	Tests/ 1M pop	Population
	USA Total	31,918,591	+47,864	575,829	+276	24,480,522	6,862,240	96,430	1,740	419,387,284	1,267,021	
1	California	3,700,774	+3,512	60,444	+63	1,960,395	1,679,935	93,661	1,530	56,363,675	1,426,487	39,512,223
2	Texas	2,830,578	+1,413	49,437	+26	2,693,355	87,786	97,620	1,705	26,650,395	919,110	28,995,881
3	Florida	2,124,233	+5,520	34,021		1,638,610	451,602	98,904	1,584	26,192,642	1,219,525	21,477,737
4	New York	2,000,173	+6,849	51,391	+75	1,284,382	664,400	102,818	2,642	47,470,275	2,440,184	19,453,561

7

7 Types of Coronavirus

- **SARS-CoV-2**
- **Middle East Respiratory Syndrome (MERS)**
 - 858 deaths. Started in Saudi Arabia in 2014. Seen in Indiana and Florida.
- **Sudden Acute Respiratory Syndrome (SARS)**
 - 774 deaths. Started in Guangdong, China in 2003. No cases reported since 2015.
- **Others – common colds and flus**

8

Symptoms

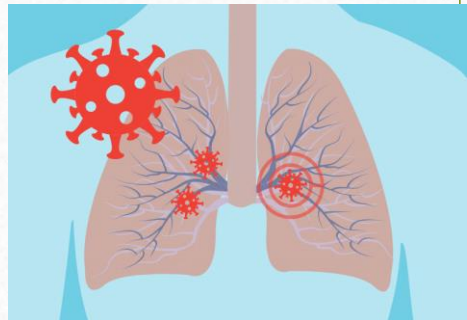
- Fever
- Coughing
- Shortness of Breath (SOB)
- Difficulty Breathing
- Fatigue
- Chills (with or without shaking)
- Body Aches
- Headache
- Sore Throat
- Congestion/Runny Nose
- *Loss of Smell*
- Nausea/Vomiting
- Diarrhea



9

COVID-19 Infection

- Can affect the upper respiratory tract
 - Nose, sinuses, and throat
- Can affect the lower respiratory tract
 - Trachea and lungs

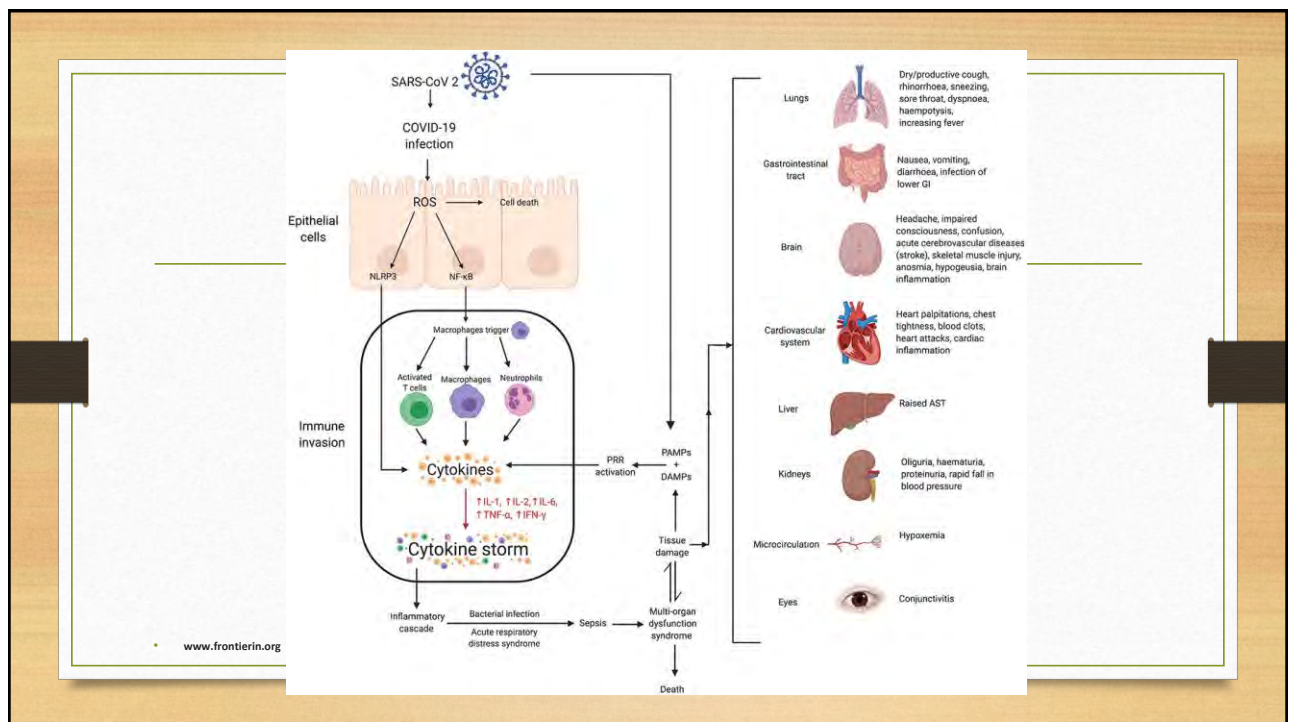


10

Cytokine Storm

- COVID-19 infection triggers innate immune system
- Floods bloodstream with inflammatory proteins called cytokines
- Causes acute respiratory distress syndrome (ARDS).
- Dysregulated hyper-inflammation in response to viral infection (not the virus or the infection) leads to tissue damage and death
- Exaggerated, rapid stimulation of the innate immune response that triggers activation of the Nod-like receptor family, pyrin domain-containing 3 (NLRP3) inflammasome pathway and release of its products including the proinflammatory cytokines IL-6 and IL-1 β .

11



12

COVID-19 Can Be Transmitted Through the Eye

- Ophthalmologist and whistleblower Li Wen Liang MD contracts COVID -19 from his glaucoma patient. Dies in Feb. 2020.
- JAMA Ophthalmology publishes finding that one can catch the COVID – 19 virus through the eye even though prevalence in tears is low.
- JAMA Ophthalmology (Sept. 2020) reports that habitual glasses wearer less likely to be infected by COVID-19. 276 patients hospitalized. 5.8% were myopes while China's population is 80% myopic. Greater than 8 hours wearing time.

13

Conjunctivitis

- Affects anywhere from 1-3% to 10% of Covid – 19 cases
- Accompanies fever, cough, and shortness of breath
- Appears early on in the course of the disease



14

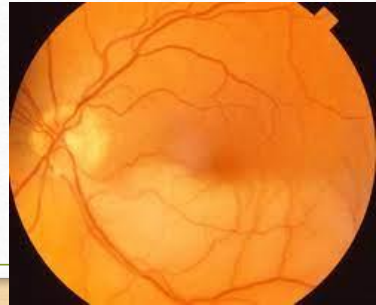
Other Possible Eye Complications

- **Optic Neuritis**

- Khalid Sawalha MD, etal. COVID-19 Induced Acute Bilateral Optic Neuritis. Journal of Investigative Medicine High Impact Case Report. October 31, 2020

- **BRAO**

- Judy Tong OD and Brandon Grove MD.
 - 27 year old Asian Female – 4th year Pharm Student – seen on 6/22/2020



15

Connect Blood Type and COVID-19 Risk

- Blood Advance (March 2021) – SARS-CoV-2 attracted to blood group A antigen on respiratory cells
- No preference of SARS-CoV-2 for blood group A red blood cells, or other blood groups found on respiratory or red cells
- Some reports suggest a 50% greater risk of infection if blood type is A and 50% less risk if blood type is O
- Researchers caution that the results do not point to any blood type being completely protective or vulnerable to the virus

16

Emergency Use Authorization (EUA)

- Emergency mechanism employed to make available the use of “medical countermeasures” in the midst of a public health crisis
- FDA evaluates an EUA request but not through the normal approval channels
- “FDA may allow the use of UNAPPROVED medical products, or unapproved uses of APPROVED medical products in an emergency to diagnose, treat, or prevent serious or life-threatening diseases or conditions when certain statutory criteria have been met, including that there are no adequate, approved, and available alternatives.”

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COVID-19 Vaccines

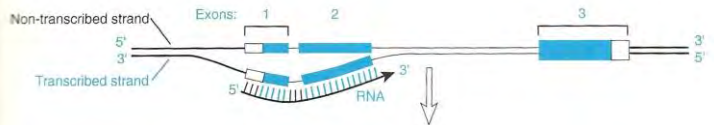
- Moderna = Modified + RNA (Nov. 16th)
 - Reduces risk of COVID-19 infection by 94.5%
 - Works by using mRNA
 - Authorized for people 18 years or older and is given in two (0.5 ml) doses, 28 days apart
- Pfizer-BioNTech = Biopharmaceutical New Technologies (Nov. 18th)
 - 44,000-person trial demonstrate that the COVID-19 vaccine is 95% effective
 - As effective as vaccines for shingles and measles
 - Authorized for people 16 years or older and is given in two (0.3 ml) doses, 21 days apart
- Johnson & Johnson (Feb. 2021)
 - Authorized for people 18 years or older and is given in a single dose

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Steps to Immunity

- For Pfizer and Moderna vaccines

- Is a messenger RNA (mRNA)
- Not a part of the virus
- Will not cause disease
- It is a molecule that contains instructions for making coronavirus's infamous "spike protein."
- Expectation is that the cells in the recipient's body are instructed to make copies of the fake spike protein
- This reaction triggers an immune system response that acts as a "fire drill"
- When subsequent exposure to the real coronavirus in the future, cells in the body are already well-equipped and trained to defend itself



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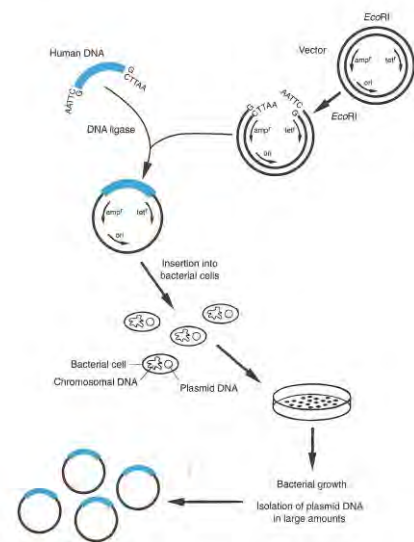
Vaccine Differences

- The Pfizer and Moderna vaccines

- Messenger RNA, or mRNA
- A segment of genetic code to cells to make the surface protein (known as spike) on the SARS-2 virus
- Activation of the immune system to recognize the spike protein as foreign
- Subsequent development of antibodies and other immunity weapons to defend itself

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- **The J&J vaccine**
 - Uses a viral vector – a harmless adenovirus
 - Vector carries genetic code for SARS-2 spike protein
 - Instructs human cells to make the SARS-2 spike protein which then triggers an immune response
 - Same technology used for the development of the Ebola vaccine



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	Johnson & Johnson	Pfizer	Moderna
Type of vaccine	Viral vector	RNA	RNA
How it works	Teaches the immune system to attack the protein the virus uses to infect other cells. The instructions are carried by a non-dangerous virus.	Uses RNA to teach the immune system to target the virus's surface, preventing infection.	Uses RNA to teach the immune system to target the virus's surface, preventing infection.
Effectiveness*	66%	95%	94.5%
Storage conditions	At least three months at refrigerator temperatures	Two weeks at freezer temperatures (-4°F), five days in the refrigerator (36° to 46°F)	One month at refrigerator temperatures
Doses needed per person	One shot	2 shots, three weeks apart	2 shots, four weeks apart
Status of availability	FDA authorized	FDA authorized	FDA authorized

*Note: The Johnson & Johnson vaccine was tested at a time when faster-spreading viral variants were common and in countries where these strains are known to exist.

Sources: Pfizer; Moderna; Johnson & Johnson; U.S. Food and Drug Administration; World Health Organization

U-T

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Signs of Anaphylaxis

Recognizing and Responding to Anaphylaxis

How to recognize anaphylaxis

Healthcare personnel should consider anaphylaxis when patients present with generalized signs or symptoms such as **hives, serious or life-threatening symptoms** (e.g., hypotension, respiratory distress, or significant swelling of the tongue or lips), or **symptoms that involve more than one body system**.



Respiratory:

- sensation of throat closing
- stridor (high-pitched sound while breathing)
- shortness of breath
- wheeze, cough



Gastrointestinal:

- nausea
- vomiting
- diarrhea
- abdominal pain



Cardiovascular:

- dizziness
- fainting
- tachycardia (abnormally fast heart rate)
- hypotension (abnormally low blood pressure)



Skin/mucosal:

- generalized hives
- itching
- swelling of lips, face, or throat



Neurological:

- agitation
- convulsions
- acute change in mental status
- sense of impending doom (a feeling that something bad is about to happen)

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How do you know that you have immunity after an asymptomatic series of COVID vaccination?

- **SARS CoV-2 Spike Protein Antibody Test**
 - Should be (+) **POSITIVE**
 - Neutralizing antibodies that target the spike protein of SARS-CoV-2
 - Blocks the binding of the virus to the ACE2 receptor site
- **IgG Nucleocapsid Protein Antibody Test**
 - Should be (-) **NEGATIVE**
 - Indicates past infection by detecting IgG antibodies

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Re-Purposed Drugs

- **Oleandrin**
 - Plant found in Africa
 - Digoxin like properties
 - Anti-cancer?
 - Toxic



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Chloroquine or Hydroxychloroquine

- **Chloroquine or Hydroxychloroquine with or without Azithromycin**
 - Immunosuppressive agent and anti-parasite
 - Disease modifying antirheumatic drugs (DMARDs)
 - Better known as Plaquenil
 - Prescribed for rheumatoid arthritis and lupus skin problems
 - **NOT RECOMMENDED.** EUA rescinded June 15th due to heart arrhythmias

26

Remdesivir (Veklury)

- The only FDA drug to be APPROVED for treatment of COVID-19
- Broad spectrum anti-viral
- Shorter time to recovery
- IV injection only



• John H. Beigel, M.D., et al. Remdesivir for the Treatment of Covid-19 — Final Report . NEJM. Oct 9, 2020

27

Tocilizumab (Actmera)

- A recombinant humanized anti-interleukin (IL)-6 receptor monoclonal antibody
- FDA approved treatment for
 - moderate to severe rheumatoid arthritis
 - polyarticular juvenile idiopathic arthritis (PJIA) and systemic juvenile idiopathic arthritis (SJIA)
 - cytokine release syndrome
- Controls level of pro-inflammatory (IL)-6 thus reducing effects, duration, and severity of COVID-19 infection.

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Collective Treatment Guidelines for COVID-19 from the REMAP-CAP and RECOVERY Trials

• Trials

- Randomized, Embedded, Multifactorial Adaptive Platform Trial for Community-Acquired Pneumonia (REMAP-CAP)
- Randomized Evaluation of COVID-19 Therapy (RECOVERY)

• Recommendations

- Hospitalized patient with rapid respiratory decompensation (invasive or non-invasive mechanical ventilation or high flow nasal canula) – tocilizumab (single dose) with dexamethasone
- Hospitalized patient with hypoxemia on conventional O2 supplementation – remdesivir, dexamethasone with remdesivir, dexamethasone alone

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Regen-Cov (Casirivimab with imdevimab)

- Regeneron in Phase III drug cocktail
- Monoclonal antibodies
- Subcutaneous Injection
- Reduced risk of symptomatic infection by 81%
- Speed up recovery to 1 week vs 3 weeks

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Other “Re-Purposed” Medications

- Convalescent plasma – plasma containing antibodies to COVID virus from previously infected person
- Lopinavir/Ritonavir and other HIV Protease Inhibitors - HIV anti-viral. NOT RECOMMENDED.
- Ivermectin – Anti-parasite. NO RECOMMENDATION FOR OR AGAINST.

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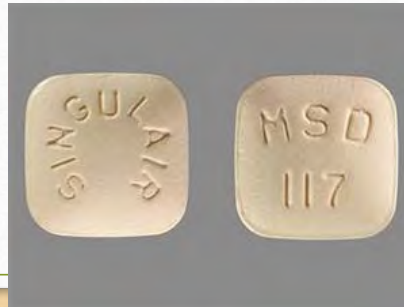
Fluvoxamine (Luvox)

- Selective Serotnine Uptake Inhibitor
- Endoplasmic reticulum chaperone protein involved in the regulation of cytokine production
- Decreases inflammation of the brain
- 2 randomized studies

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Montelukast, Zafirlukast and Pranlukast (Singulair, Accolate, and Onon)

- Cysteinyl leukotriene receptor antagonists
- Likely to inhibit SARS-CoV-2 main protease (Huynh et al. 2020)

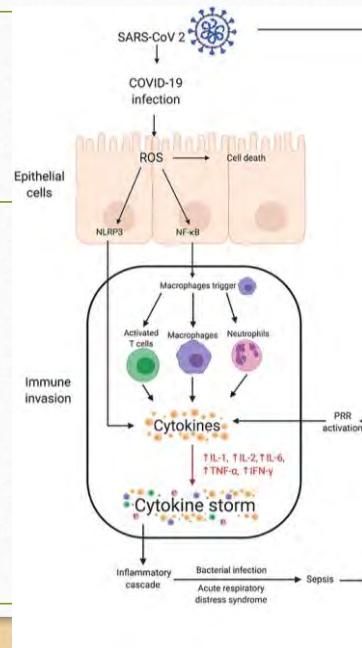


33

Properties of Montelukast (MK)

- Endotheliitis Induced by SARS-CoV-2 Infection
 - Antagonizes the inflammatory cascade induced by angiotensin II in vascular smooth muscle cells decreasing inflammation.
- Neurological Disorders Induced by SARS-CoV-2 Infection
 - Shown to reduce damage induced on the blood-brain barrier, observable anti-convulsant properties, reduced neuro-inflammation
- Atherogenic Vascular Inflammation
 - Exhibit anti-atheromatous properties reducing COVID-19 mortality in atheromatous patients
- Ischemia/Reperfusion
 - May alleviate the ischemia/reperfusion phenomenon reducing risk of arterial and venous thromboembolism
- Asthma, Hyper-Reactivity Bronchitis, and Post-Infectious Cough

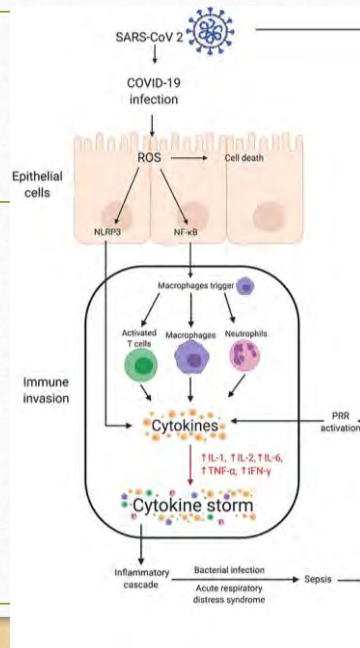
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- **Cytokine Storm**
 - Antagonist action of ZK on CystLT1 receptor protects the endothelium from inflammatory lesions induced by TNF- α (Zhou, etal 2019)
- **Acute Respiratory Distress Syndrome**
 - Decrease in the intensity of the induced cytokine cascade and a lesser activation of neutrophils in the bronchoalveolar fluid
- **Antioxidant Properties**
 - Demonstrable upregulation of mitochondrial genes and genes responding to oxidative stress (Shao, etal. 2006)
- **Anti-Fibrosis Properties**
 - May limit the residual extent of COVID-19 sequelae of pulmonary damage (Peng, etal. 2017)
 - May regulate the extracellular remodeling matrix and inhibits the formation of fibrosis (Debelleix, etal. 2018)

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Vitamin D

- Provides a certain degree of protection against respiratory infections, unsure about COVID-19
- Augments host barrier epithelial integrity by reinforcing intracellular junctions
- Triggers antimicrobial peptide production
- Receptors expressed on surface of immune cells types (B cells, T cells, and antigen presenting cells)
- Influences the expression and secretion of pro-inflammatory chemokines and cytokines

• Lancet Diabetes & Endocrinology

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- Low levels of 1,25-dihydroxyvitamin D3 seen in COVID-19 patients
- Involved in pulmonary angiotensin-converting enzyme 2 expression which in turn reduces lung surface tension in COVID-19
- Exerts inhibitory effects on inflammation namely cytokine IL-6
- African American, Obese, and Seniors are prone to low levels
- Africans American with low levels of Vitamin D more at risk - <40 ng/ml
- Low vitamin D levels with higher COVID-19-related mortality

• Derbyshire EJ, Calder PC. "Respiratory Tract Infections and Antibiotic Resistance: A protective role for Vitamin D". *Frontiers in Nutrition*

39

Trivia but not so Trivial

- Vaccination may provide 6 months or greater protection
- Pfizer vaccination likely will require a 1 year booster
- COVID – 19 infection may confer 3-4 months of protection. Rare to get 2nd time – but there have been reports of re-occurrences
- Is it safe to go back to eat out, church, fly, or stay in a hotel?
- How long will we likely have to wear a mask?
- More symptoms upon second vaccination and more symptoms for those that had COVID.

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IPC: A Case for Collaboration

John Nishimoto, OD and Julie Tyler, OD





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IPC: A Case for Collaboration

John Nishimoto, OD, FAAO

Julie Tyler, OD, FAAO



1

Disclosures

Dr. John Nishimoto

•None

Dr. Julie Tyler

•None

***"Alone we can do so little;
together we can do so
much." – Helen Keller***

2

Learning objectives

1. To emphasize the value of Optometry's role within interprofessional collaborative practice
2. To provide resources for optometrists that will build greater relationships with other health professionals
3. To recognize several common clinical conditions that most benefit from collaboration and management within health professionals

3

Our roles at SCCO/MBKU

- Clinical Education of SCCO students
- Developing Interprofessional Collaborative Practice (ICP) programs
- Collaborating with faculty & students from the School of PA Studies and College of Pharmacy
- Provide students with skills – *including team building*- needed to develop relationships with other health providers

4

Ketchum Health Clinic

To provide co-management services for patients

- *Family Medicine*
 - Includes rotations/consultations with PA and Pharm
- *Pediatrics and Vision Therapy*
- *Low Vision Rehabilitation*
- *Cornea and Contact Lenses*
- *Ocular Disease and Ophthalmology*
- *Telehealth*



5

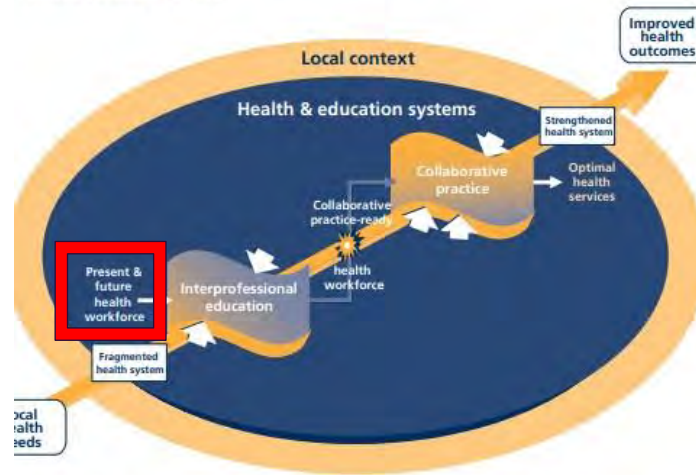
IPC-

World Health Organization (WHO) KEY messages

- Collaborative practice happens when **multiple health workers from different professional backgrounds work together** with patients, families, carers, and communities to deliver the **highest quality of care.**
- The goal is to prepare all health professions students **for deliberately working together with the common goal of building a safer and better patient-centered and community/population-oriented** U.S. health care system.

6

FIGURE 2: Framework for Action on Interprofessional Education & Collaborative Practice



Reprinted with permission from: World Health Organization (WHO). (2010). *Framework for Action on Interprofessional Education & Collaborative Practice*. Geneva: World Health Organization.

7

Who do you collaborate with?

8

Providers that we can collaborate with:

Family Medicine/Practice	Nutritionists
Pediatricians	Audiologists
Emergency Department (ED)	Speech-Language Pathologists
PAs	Occupational Therapists
Nurse Practitioners	Physical Therapists
Pharmacists	Neurologists
Rheumatologists	Psychologists
Endocrinologists	Social Workers
Dermatologists	Schools (Nurses, Teachers, Psychologists)
Allergists	

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How to enhance relationships with providers – *examples from Daniel May, MD*

- **Making introductions & discussing your role**
 - *As an optometrist, we can assist with anything related to the eye and vision, and can refer as a point of contact for any suspected health problem*
 - ❖ *Visual fields*
 - ❖ *Headaches*
 - ❖ *Allergiesand many more*
- **Communication on a relatable level with other professionals**
- **Minimize optometry (or eye specific) jargon**
- **Keep short and concise**

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Give to health care associates – SIMPLE request letter courtesy of Dr. Alissa Nagel-Esposito. Also, SIMPLE consult response

University Eye Center at Ketchum Health
5400 E. La Palma Ave
Anaheim, CA 92807
714-440-7409

March 23, 2021

RE: Patient Name, Diabetic Eye Exam

To Whom It May Concern:

Thank you for referring Patient Name to our office. She is a pleasant 53 year old female that entered our office on March 13, 2021 for a diabetic eye exam. She did not have any vision complaints.

Best Corrected Visual Acuity at 6 Meters (Distance)

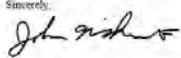
Right Eye: 20/20

Left Eye: 20/20

Our findings on Dilated Eye Examination revealed that she has proliferative diabetic retinopathy, right and left eye. Our recommendation would be to reassess the control of her Type II diabetes. She is also referred to a retinal specialist for consideration of laser treatment.

If you have any questions feel free to contact me. Thank you very much for your time in this matter and allowing us to examine your patient.

Sincerely,



John Nashimoto, O.D.

Diabetic Eye Exam Request

Patient's Name: _____ Date: _____

Patient's Phone: _____

Insurance Company: _____

Type: IDDM or NIDDM Last HbA1c and date: _____

Referring Doctor: _____ MD OD DO

Address: _____

Phone: _____ Fax: _____

Please send results via: ☐ Fax or ☐ E-mail to: _____

8937 W. Sahara Ave. Ste A
Las Vegas, NV 89117
phone: 702-254-3558 fax: 702-254-4012

We will gladly see your patient within 48 hours of receiving your referral.

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CHALLENGES to Interprofessional Collaboration

- INSURANCE
- Availability of services

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Case Example

IPE Clinic at SCCO/Ketchum Health

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"Diabetic Day": March 16, 2021

* 53 YO Hispanic Female - Pt entering with "no ocular or visual complaints" with (+) history of Diabetes Mellitus, Type II

Ocular Hx: Does not wear any glasses (although prescribed)

Med Hx: Type II DM (dx 2018), hyperlipidemia, HTN

Medications: Invokana (DM with kidney issues), Lipitor (Cholesterol), Humalog (Insulin)

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Diabetics

Specific additional history --

Blood Sugar: 112

HgA1c: unknown

OPPORTUNITIES: *Additional Testing In Office*
 Random Blood Sugar (RBS) Glucometry
 Hemoglobin A1c Testing

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Data – Diabetic Eye Exam

Refraction and Best Corrected Distance VAs:

OD: -0.75-0.50x90 (20/20)

OS: -1.00-0.75x92 (20/20)

Preliminary Test: Normal results for pupils, CF, motility

Blood Pressure: 130/80mmHg seated

Anterior Seg & Tonometry:

- **IOP:** 10mmHg/12mmHg OD/OS @16:34
- **Lids/Lashes, Conj, Cornea:** Without pathology OU
- **Iris:** (-) Neovascularization of iris, normal iris appearance
- **Lens:** Trace NS

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Posterior Seg: Optos and Dilated Fundus OD

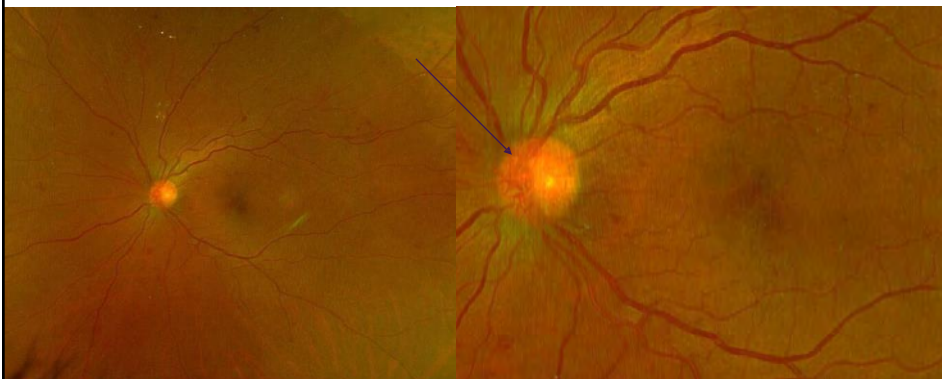


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Posterior Seg: Optos and Dilated Fundus OD



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Management – That Day

PA – Completed **Diabetic foot exam** and counseling of pt regarding shoes and nutrition

Pharmacist – Reviewed medications with patient, proper dosing and timing of medications; discussed considerations of other options for pt based on findings

Team – Consulted with patient; With permission also consulted with family member to educate on visual risks and need for FU

Who can we collaborate with?

- OMD --> Vision
- Primary Care Physician (/whoever managing DM)
- PA
- Nutritionist
- Endocrinologist

Management next steps

Consult PCP- Update status of DM/BS control

- Recommend update of medications to help with protection of kidneys
- If Blood Sugar/HgbA1c not consistent with findings of diabetic ret --> Recommend r/o blood dyscrasias and other causes of neovascularization of retina

Retina Consult – FA & Treatment considerations

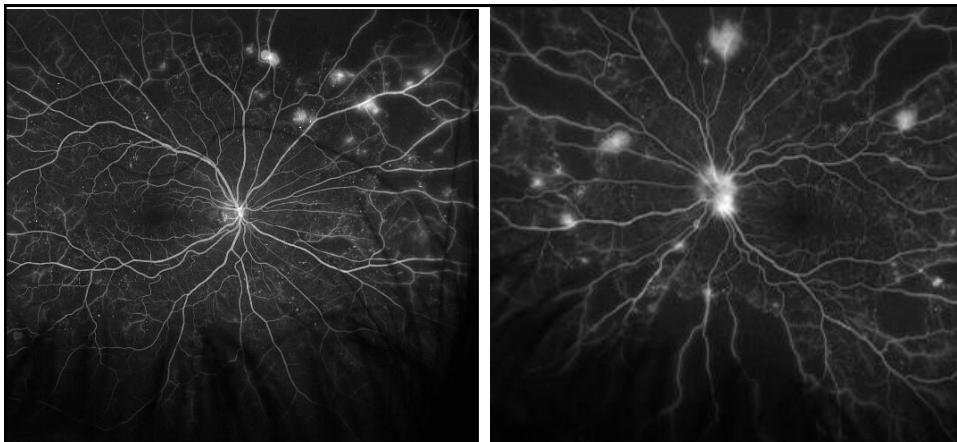
- Anti-VEGF injection vs PRP

Note – PCP follow up visit added Lisinopril (HTN)

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**NVD &
NVE:**
Large
areas
of
drop-
out



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**PRP
OS 7-1-21**

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Diabetes Care - HEDIS Measures – NCQA

Comprehensive Diabetes Care (CDC)

Assesses adults 18–75 years of age with diabetes (type 1 and type 2) who had each of the following:

- Hemoglobin A1c (HbA1c) testing.
- HbA1c poor control ($>9.0\%$).
- HbA1c control ($<8.0\%$).
- HbA1c control ($<7.0\%$) for a selected population. *
- Eye exam (retinal) performed.
- Medical attention for nephropathy.
- BP control ($<140/90$ mm Hg).



Posterior Segment Considerations

Conditions

*Diabetes Mellitus/
Diabetic Retinopathy*

*Hypertension/
Hypertensive
Retinopathy*

*Ocular Ischemic
Syndrome (OIS)*

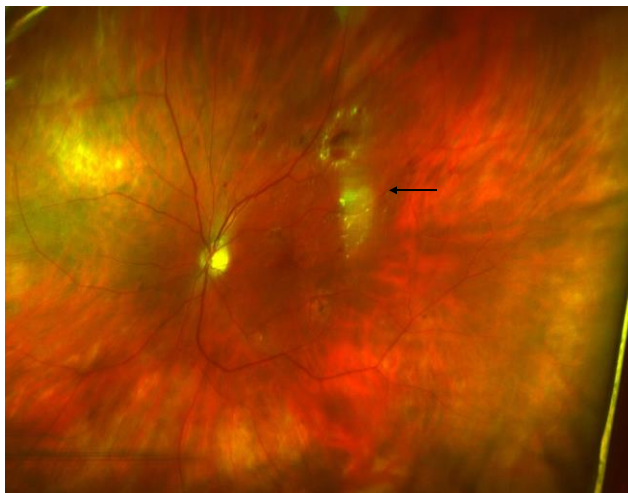
*Central Retinal Artery
Occlusion (CRAO)*

Collaborators

**Who do you
generally think
of "first" in
this area?**

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Hypertensive Retinopathy



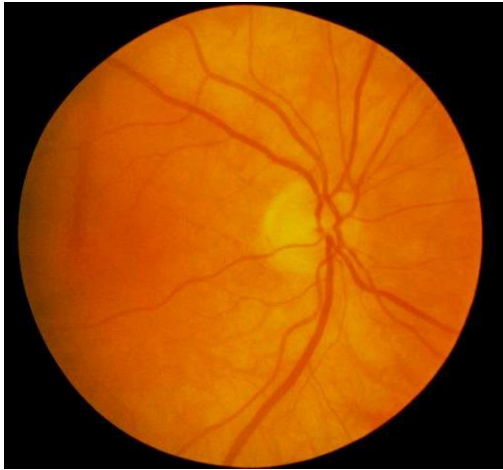
This image shows an artery macroaneurysm with associated exudates in hypertensive retinopathy (arrow)

Communication:
PCP, PA

OMD if needed

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Hypertensive Retinopathy



This image shows changes in the A/V ratio

Communication:
PCP, PA

***Are you
checking BP in
office?***

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Hypertensive Retinopathy – What is significant here?



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BRVO Early Stages

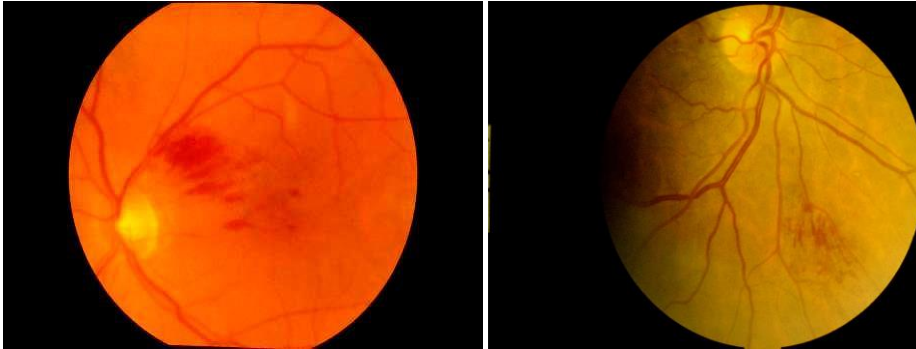


Image examples of small Branch Retinal Vein Occlusions (BRVO)

Testing in office: OCT to assess for mac edema based on area

Communication: PCP, PA, OMD if needed

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BRVO with concurrent AION



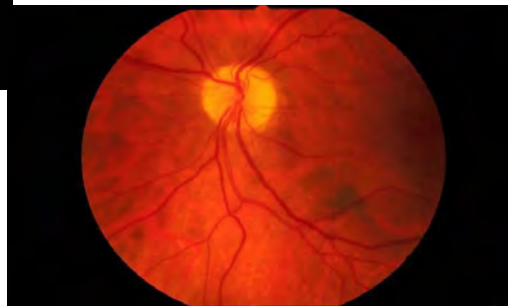
This image shows an OD with inf BRVO (ischemic/CWS) and OS with ON pallor due to past AION

Communication:

PCP, PA

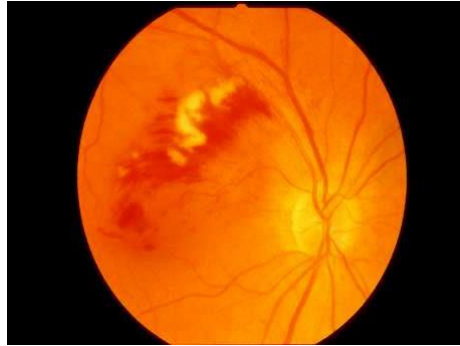
Endocrinologist (DM too!)

OMD if needed



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BRVO "Longstanding" with increased risk for vision



These image shows eyes with increased risk for additional complications --> Ocular and otherwise

**** Neovascular concerns**

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Hollenhorst Plaque



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Who can you collaborate with?

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Carotid Artery Disease (CAD)

Visual concerns with CAD *beyond* Hollenhorst plaques:

- Blackouts to vision (TVO)
- Peripheral signs of Ocular Ischemic Syndrome (OIS), include blot hemes in the "mid-periphery"
- Other non-retinal findings
 - ❖ *Visual Field Defects*
 - ❖ *Decreased persistent VA*
 - ❖ *Corneal folds*

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Carotid Occlusive Disease - Ocular Ischemic Syndrome (OIS)

Major obstruction of Carotid artery --> Look at retina as well as other signs and symptoms

CC/Oc Hx/ROS:

- Pts may complain of decreased and/or "darkened" vision
- **May also have complaints related to:**
 - *Achy, soreness of eye*
 - *Ringling in the ears or heartbeat sounds in the ear*

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Retina in OIS

- Generally, mid-peripheral retina involved
- Dot-blot hemes
- Dilated non-tortuous veins

Outside of the eye:

- Bruits

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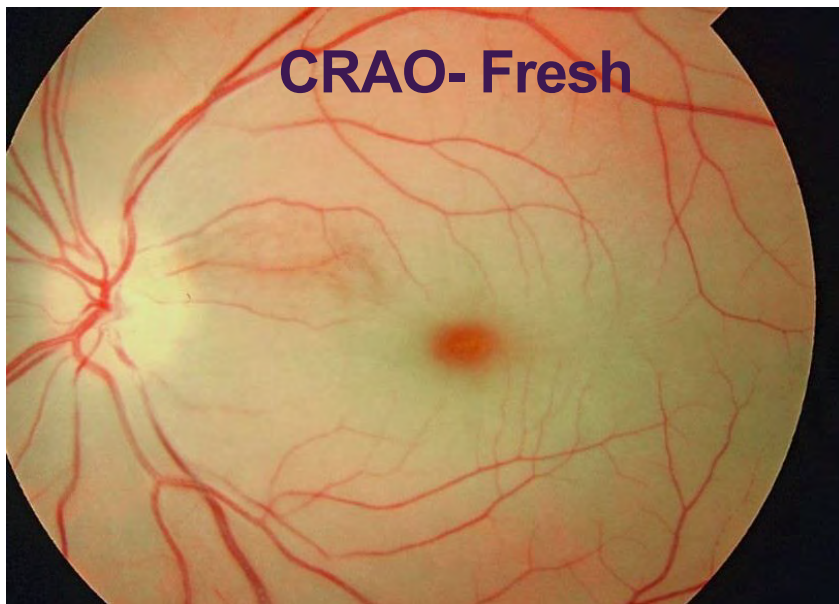
What else to consider with plaques AND?

Differentials and finally, considerations
for collaboration?

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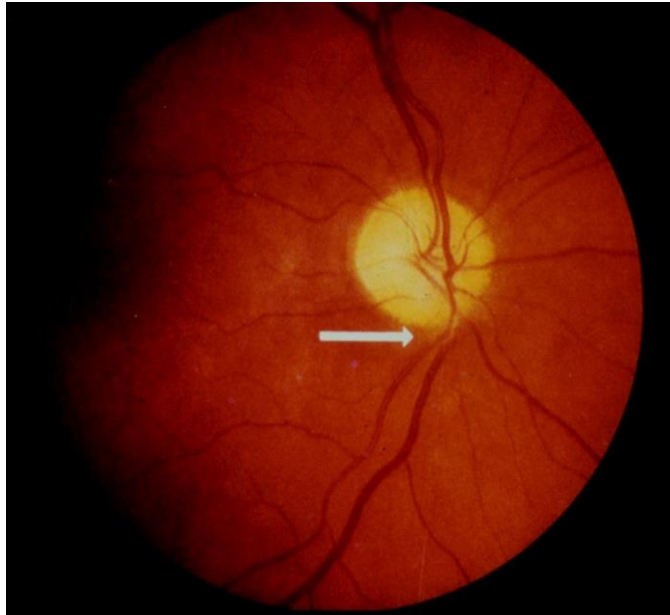
41



By sidthedoc - Own work, CC BY-SA 4.0,
<https://commons.wikimedia.org/w/index.php?curid=84360138d> text

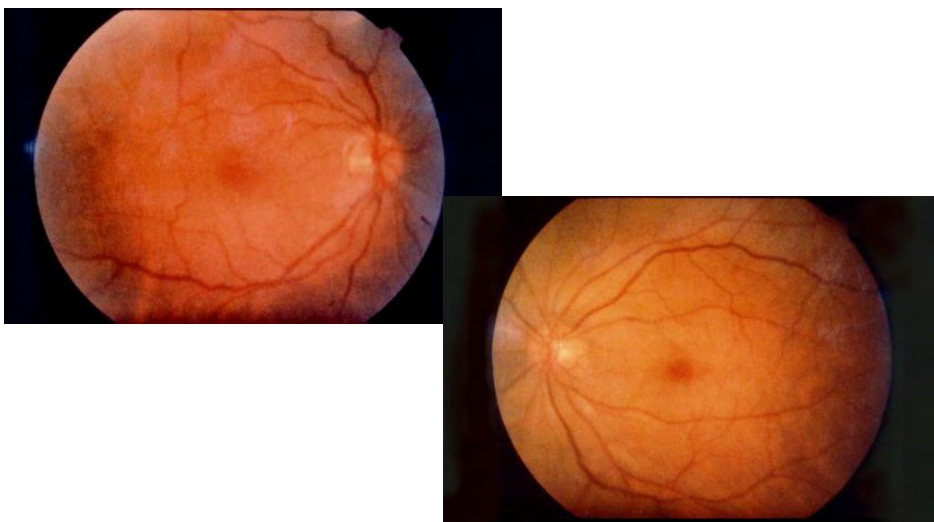
42

CRAO - Old



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CRAO – Incomplete (*compare each picture*)



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CRAO – Current Management Strategies

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CRAO – Collaboration/Considerations

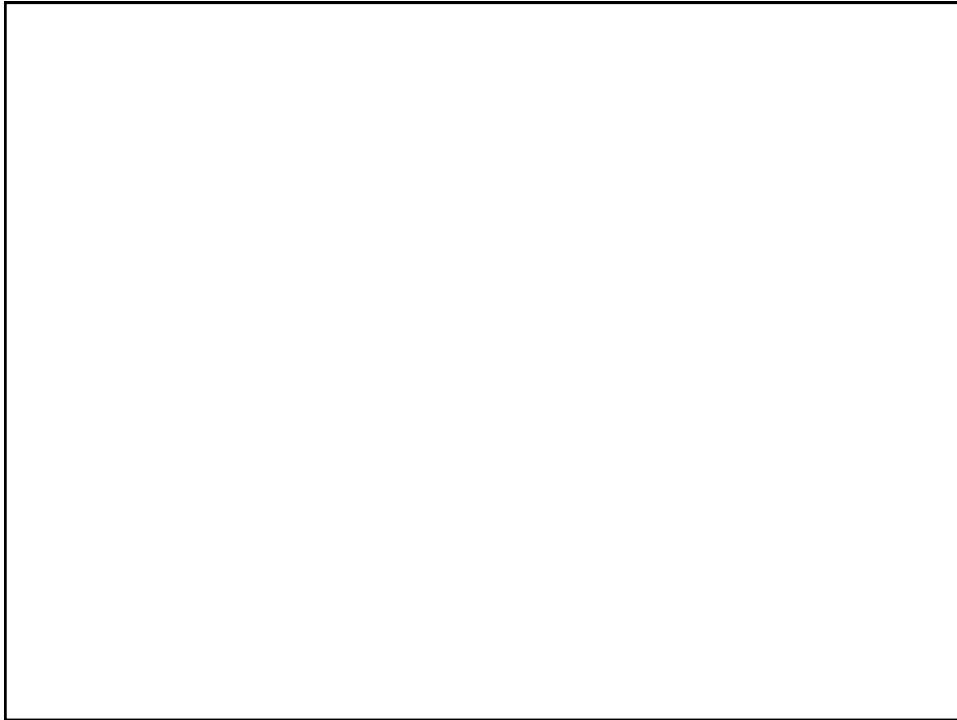
Standards for management in 2021

STROKE centers: Primary vs. Comprehensive

- Primary: Stabilize pts and **are able to provide** victims of ischemic stroke by using a **clot busting drug**
- Secondary: Stabilize pts and **able to perform catheter-based procedures to remove blood clots**, as well as provide neurosurgical services

Availability based on location may be restricted

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Neovascular Glaucoma

Depends on:

- Amount of retinal ischemia and "opportunity" to release VEGF

Vascular endothelial growth factor (VEGF) plays central role in angiogenesis

- Also needs, viable tissue support

Found in:

- Up to 60% of Hemorrhagic CRVO's
- Ocular ischemic syndrome
- Leads to angle closure
- Pupillary margin development

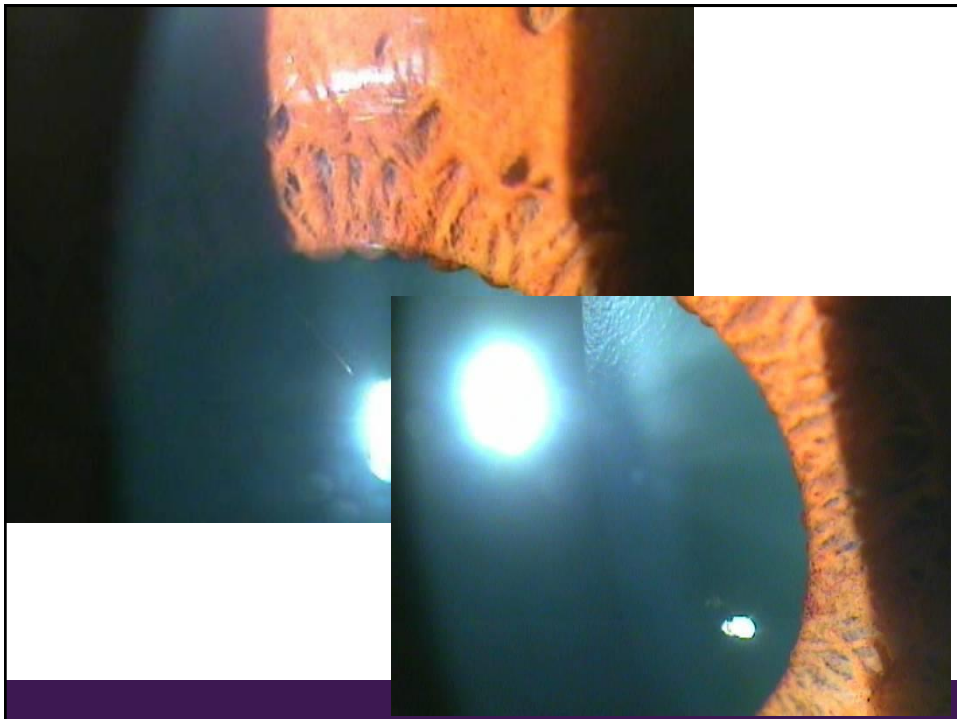
48

Neovascular Glaucoma

Starts as OPEN ANGLE type of glaucoma, but

- Leads to angle closure
- Pupillary margin development often first with most pts who develop, but not all (depends on systemic/ocular etiology)

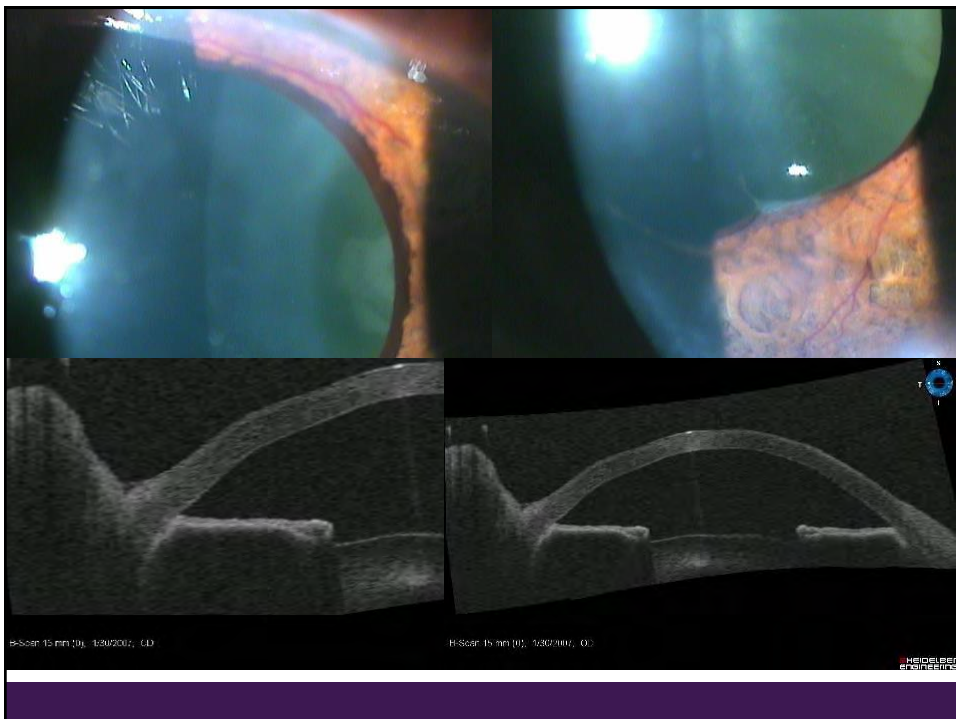
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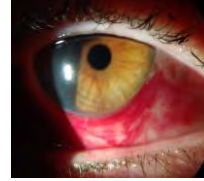
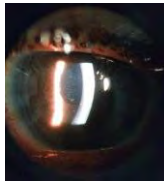
More "typical" anterior segment collaboration opportunities

Conditions

- Uveitis
- Adult inclusion Conjunctivitis (AIC)
- Allergic Conjunctivitis
- Dry Eye
- Thyroid Eye Disease

Collaborators

- Primary care
- Rheumatology
- Infectious disease
- Allergist
- Endocrinology



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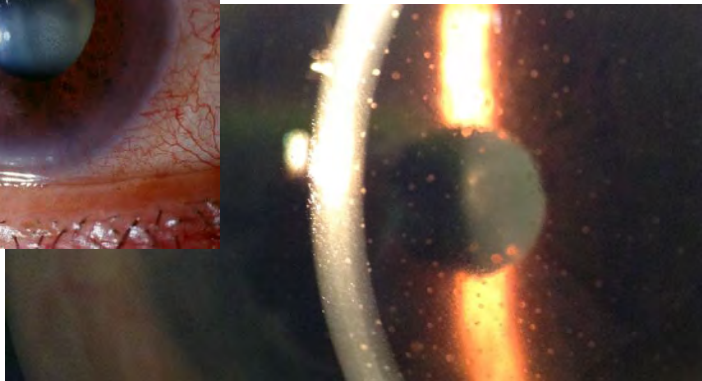
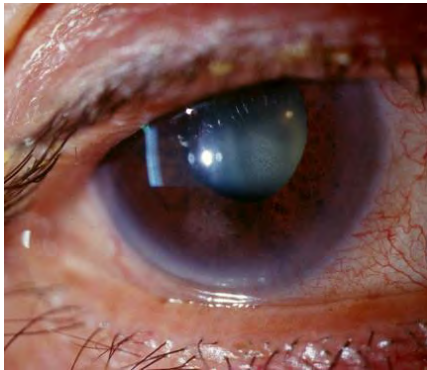
Uveitis -

Communication:

PCP, PA

OMD if needed (*based on surgical hx, seriousness*)

Rheumatologist



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Uveitis: Collaboration opportunities

When do we want to consider a work-up for patients?

- SEVERE presentations without other known etiology
- BILATERAL uveitis → OR evidence/suggestion that condition was previous in other eye
- Granulomatous uveitis
- Significant ocular/med hx or ROS findings/co-morbidity
 - *New rashes, swelling, aches/pains or malaise*; ALSO, (+) *Family hx, contagious disease exposure, risky social habits, travel hx, etc...*

Or if additional risk findings noted during exam with careful health evaluation, tonometry and DFE

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Lab Guide

Referral for Serology and Imaging

From Primary Eye Care _____

Patient's Name: _____ Date: / /

DIAGNOSIS:

- ☐ Anatomic signs
- ☐ Cornea wood spigot)
- ☐ Dry eye disease/Inflammation
- ☐ Keratic cell keratitis
- ☐ Hollenhorst plaque
- ☐ Optic disc edema
- ☐ Optic atrophy/atrophic
- ☐ Optic atrophy/ischemic
- ☐ Optic neuropathy, traumatic
- ☐ Retinal artery occlusion
- ☐ Retinal hemorrhage
- ☐ Retinal pigment epithelial detachment
- ☐ Retinal vasculitis
- ☐ Retinal vein occlusion
- ☐ Uveitis, anterior
- ☐ Uveitis, posterior
- ☐ Vitritis
- ☐ Other _____

DIAGNOSTIC STUDIES REQUESTED:

- ☐ Blood pressure check
- ☐ Central diplop/diplexia, lateral
- ☐ CT brain, with/without contrast
- ☐ CT orbit, with/without contrast
- ☐ Chest X-ray, PA/lateral
- ☐ EEG
- ☐ Cardiac echo
- ☐ Ocular exam
- ☐ Holter monitor
- ☐ MRI brain, with/without contrast
- ☐ MRI orbit, with/without contrast
- ☐ PPD skin test with control
- ☐ Temporal artery biopsy
- ☐ X-ray (other one: _____)

LABORATORY PANELS REQUESTED:

- ☐ Comprehensive Metabolic Panel
- ☐ Lipid Panel
- ☐ Thyroid panel (TSH, T3, free T4)

LABORATORY TESTS REQUESTED:

- ☐ Angiotensin converting enzyme (ACE)
- ☐ Antismoking cytotoxic antibodies (ANCA)
- ☐ Autoantibody antibody (ANA)
- ☐ Auto-oxidative antibody, IgM, IgG
- ☐ Aspartate aminotransferase II
- ☐ Antiplatelet antibodies
- ☐ CBC with differential and platelet count
- ☐ CHV titer, IgM, IgG
- ☐ Cryoglobulin
- ☐ C-reactive protein (quantitative)
- ☐ C-reactive protein (qualitative)
- ☐ D5 DNA
- ☐ Epstein-Barr virus (EBV) immunoglobulin (IgA)
- ☐ Erythrocyte sedimentation rate (ESR)
- ☐ Fasting blood sugar
- ☐ Glycated hemoglobin A1c
- ☐ Hepatitis surface titer, IgM, IgG
- ☐ Hepatitis core titer, IgM, IgG
- ☐ HLA-B27
- ☐ Homocysteine level
- ☐ Hemoglobin electrophoresis
- ☐ Hemogram/tissue
- ☐ HIV test, IgM, IgG
- ☐ Immunofluorescence assay (IFA)
- ☐ Latex coagulopathy
- ☐ Lyme test - ELISA
- ☐ Nuclear acid amplification test (NAAT)
- ☐ Oral glucose 1 hour
- ☐ Protein - C deficiency
- ☐ Protein - S deficiency
- ☐ PT/PTT
- ☐ Quantiferon-TB Gold (QFT)
- ☐ Rheumatoid factor (RF)
- ☐ EPR-ATA-HSA-TP-VDEL
- ☐ Serum bromide
- ☐ Sickie prep
- ☐ Serum protein electrophoresis (SPEP)
- ☐ Transaminase titer, IgM, IgG
- ☐ Typhus multilocus particle agglutination assay

OTHER:

- ☐ Blood cultures:
 - ☐ Aerobic & Anaerobic bacteria
 - ☐ Fungal
 - ☐ Mycobacterium

Dr. _____ License _____

1091 _____

General Lab Order Referral Form
May 2020

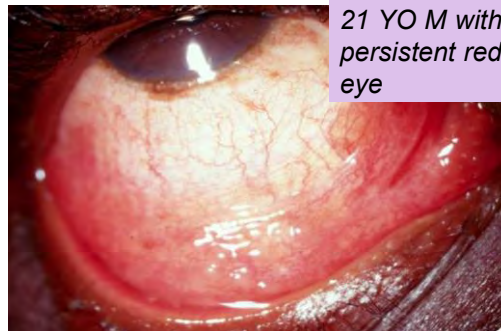
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Adult Inclusion Conjunctivitis

History:

- Recent “new” sex partner, ~4-8 weeks
- Incubation ~5 to 12 dys
- Females often asymptomatic

Often diagnosed because pt Non-responsive, persistent red eye with traditional antibiotic and supportive treatment



21 YO M with
persistent red
eye

Communication:

PCP, PA
Infectious disease/Community
health centers

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Allergic conjunctivitis

24 YO F with
itchy red eye

History:

- Chronic or acute symptoms of redness, itching, irritation

ROS:

- Rashes, chronic rhinitis



Communication:

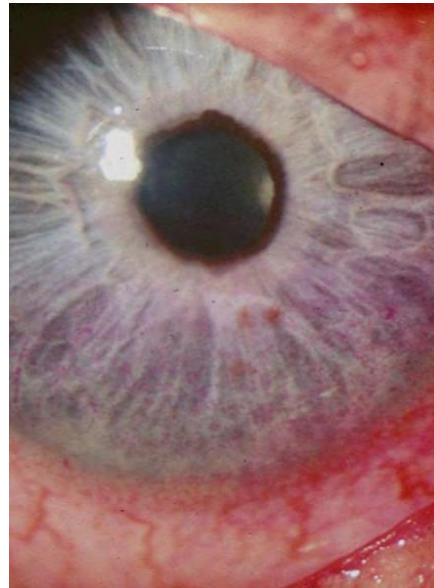
PCP, PA

Allergist (*even if in office testing for therapy*)

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Scleritis

History:

- First time but especially if recurrent

ROS:

- Skin, joints, muscles



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Scleritis

CHECK to see if recent systemic medication changes

Communication:

PCP, PA

Rheumatologist



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Thyroid Eye Disease (TED)/ Thyroid Associated Orbitopathy



75% of pts with thyroid
hyperactivity develop
ocular signs

* Of pts with thyroid
hyperactivity, 15% of
those pts will develop
serious functional
impairment

Communication:

PCP, PA

*** IMAGING!**

Endocrinologist

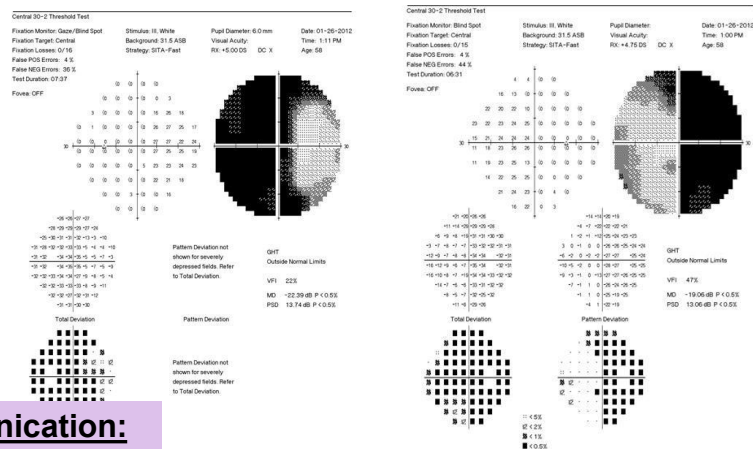
Smoking is the most
important risk factor
for complications
associated with TED

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Thyroid Associated issues may also be associated with



Communication:

PCP, PA

*** IMAGING!**

Endocrinologist

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HFA 1 750-13647-5.1/5.1

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HFA 1 750-13647-5.1/5.1

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Who can you collaborate with?

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More Neurologic Conditions & Collaborations

Conditions

- New onset double vision/ motility issues
- Migraines
- Stroke
- Multiple sclerosis (MS)/ Parkinsons disease
- Progressive supranuclear palsy

Collaborators

- Neuro-ophthalmologist/ Neurologist/Neuro-surgeon
- Therapists – VT, PT, OT
- Psychologist

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Who can you collaborate with?

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THANK YOU!!

None of us is as smart as ALL of us!

- Ken Blanchard

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