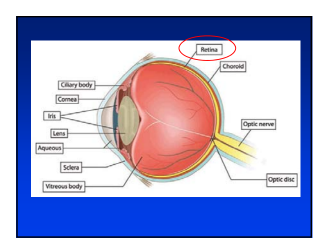
# **Ophthalmologic Issues in VHL** Mark W. Johnson, MD University of Michigan Kellogg Eye Center

#### A Brief History of VHL Disease

- von Hippel (1904)
  - retinal capillary hemangioblastomas
  - several generations of family members
  - several pedigrees
- Lindau (1926)

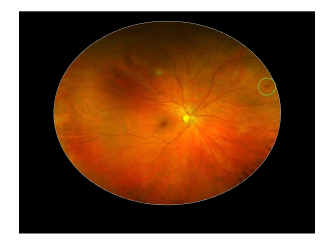
  - described familial syndromehemangioblastomas (retina and cerebellum)
    - cysts (kidney, pancreas, epididymis)
- Melmon and Rosen (1965)
  - criteria for clinical diagnosis



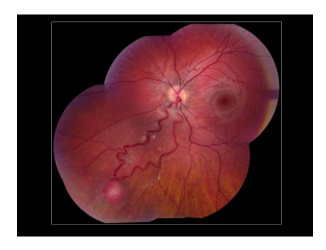
#### **Ocular Manifestations** Retinal Capillary Hemangioblastoma

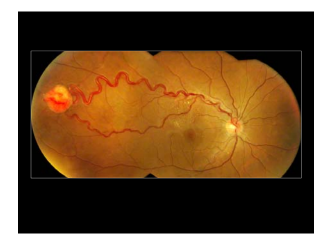
- May be the first manifestation of VHL disease
- Range from tiny lesions to large tumors with major visual impairment
- Located predominantly in retinal periphery (85%)
- Initial appearance
- subtle red or gray dot With growth, appears as distinct nodule
  - dilated feeding and draining vessels

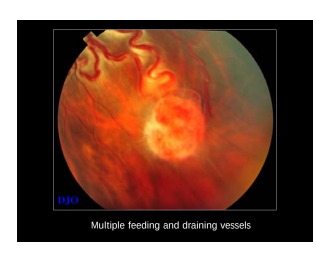


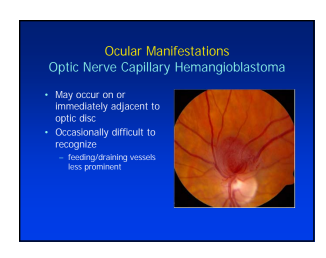


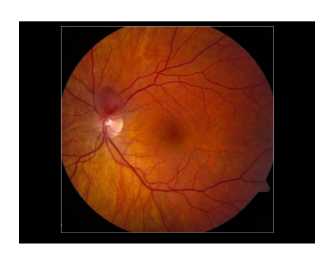


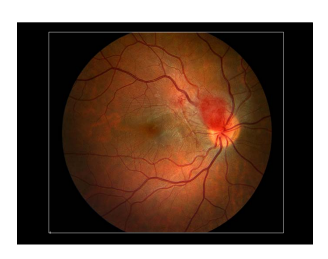














# Retinal & Optic Nerve Hemangioblastoma Natural History

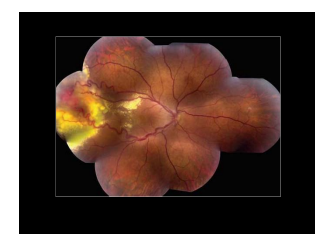
- Can appear at any age
   patients typically have no symptoms initially
  - often discovered on routine or screening exam
- Without treatment

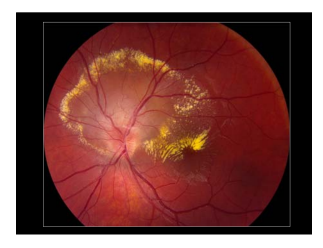
  - rarely regress spontaneously
    usually grow slowly and progressively
    often begin leaking as they enlarge
    eventually displace normal structures
    may completely fill the eye

# Retinal & Optic Nerve Hemangioblastoma Natural History/Secondary Complications

- Leakage
  - retinal edema (swelling)
  - lipid (yellow) exudates
- Fibrosis (scar tissue)
- · Retinal detachment

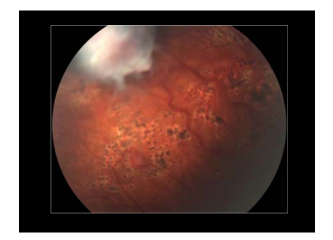
  - exudative (from leakage) tractional (from fibrosis and vitreous traction)
- Bleeding
- Neovascular glaucoma

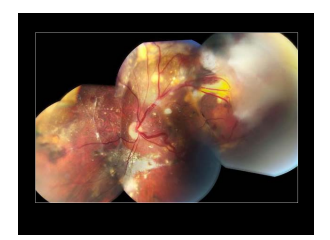








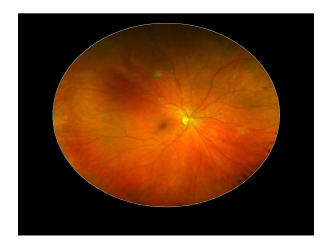


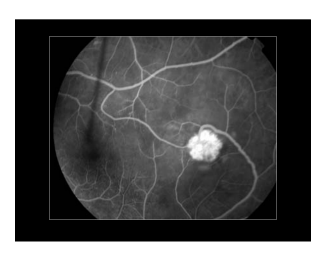


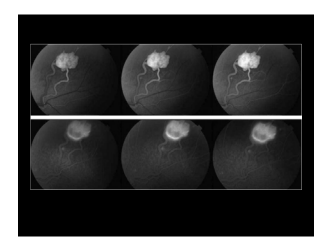
# **Diagnosis**Ocular Hemangioblastoma

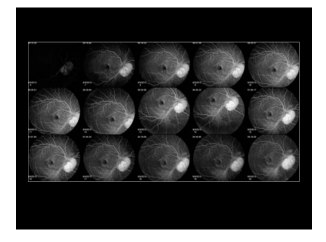
- Diagnosis typically based on clinical appearance
- No definitive diagnostic tool
- Confirmatory/useful studies

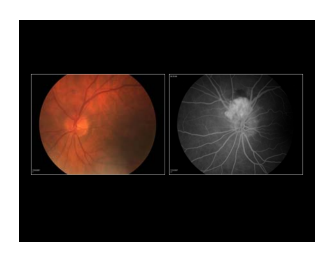
  - wide-angle fundus photography
     fluorescein angiography
     ultrasonography
     optical coherence tomography
     detection of associated macular edema

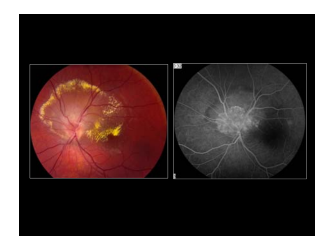


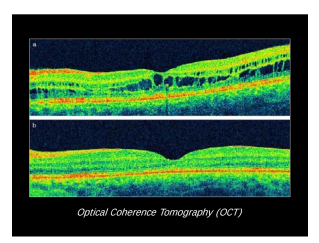












# Diagnosis VHL Disease CLINICAL CRITERIA Family History + CNS\* hemangiobastoma, Pheochromocytoma, or Clear cell renal carcinoma Family History - 2 or more CNS hemangioblastomas or CNS hemangioblastoma + visceral tumor • Up to 20% of cases arise de novo (first affected member of family)--genetic testing extremely helpful in such patients • Family members with mutations should have regular clinical screening studies - ophthalmoscopy yearly starting in infancy

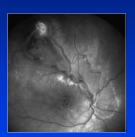
## **Epidemiology** Ocular Manifestations

- Large NEI study (Wong WT, et al, 2008)
  - 38% of patients had ocular involvement
    - mean age 36 years (range, 7 to 84)
    - 47% male
    - 95% white

  - laterality42% unilateral58% bilateral
  - location
    - 85% peripheral
    - 15% optic nerve

#### Vision Loss in VHL Prevalence

- NEI study
  - 77% had 20/20 vision
  - 5.7% legally blind
  - 20% had visual impairment in one eye



#### Vision Loss in VHL Causes

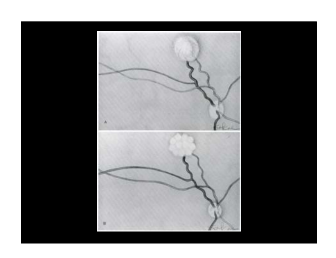
- Tumor exudation (leakage)
  - macular edema
  - exudative retinal detachment
- Glial proliferation (scar tissue) retinal distortion
  - traction retinal detachment
- Neovascularization
  - vitreous hemorrhage or retinal traction
- Neurological lesions
  - increased intracranial pressure leading to optic atrophy
  - hemangioblastomas affecting RB optic nerve or optic tract

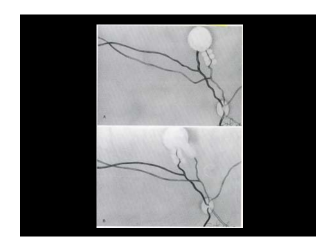


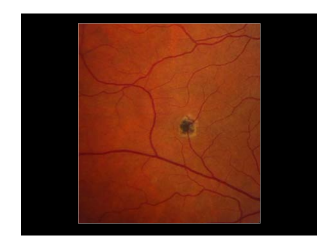
#### **Ablative Treatment** Retinal Hemangioblastomas

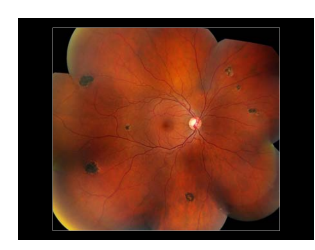
Lesion size/location	Treatment modality
Very small (1-2 mm)	Laser (direct)
Small (3-5 mm)	Laser (feeder vessel + direct)
Small, very peripheral	Cryotherapy
Moderate to large (> 5 mm)	Cryotherapy (consider adjunctive steroid or anti-VEGF)
Complicated (traction, retinal detachment, vitreous hemorrhage)	Vitrectomy and/or scleral buckling surgery (with laser, diathermy and/or cryotherapy)

The smaller the lesion, the easier and safer it is to treat

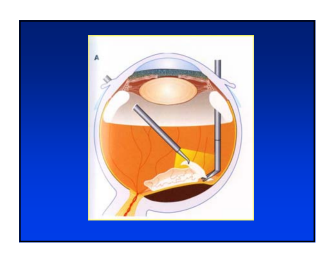












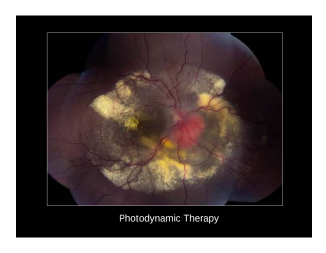
### **Ablative Treatment**

Optic Nerve Hemangioblastomas

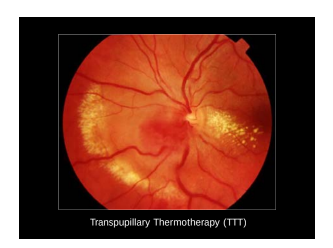
- Treatment difficult—no consensus
  - Laser treatment
    - risk of visual acuity and/or visual field loss
       serial, low-intensity treatments promising
  - Photodynamic therapy
    - mixed results

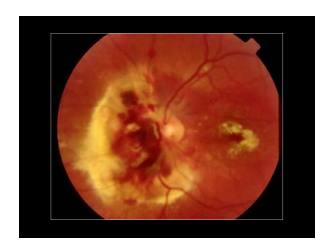
  - risk of optic nerve injury

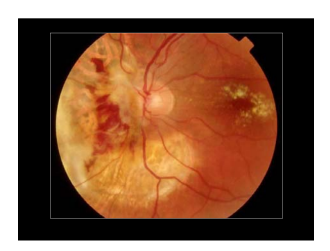
     Transpupillary thermotherapy
     risk of significant nerve injury (little data)
  - Radiation
    - should be avoided (increases VEGF production)

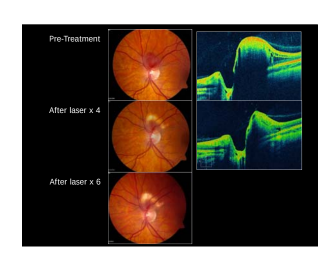


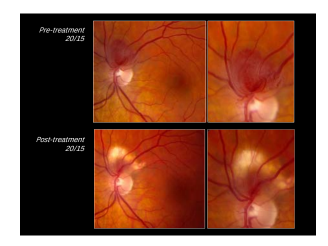












# Pharmacologic Treatment Anti-Angiogenic Agents

- VHL involves high levels of vascular endothelial growth factor (VEGF)
  - drives tumor growth and vessel leakage
- Anti-VEGF treatment is rational approach
- Studies to date
  - decreased leakage

  - no change in tumor size
     anti-VEGF treatment alone appear inadequate
- Successful pharmacologic approaches may need to target multiple proteins upregulated in VHL

