# REFERRAL GUIDELINES for the PRIMARY CARE PHYSICIAN:

Visual symptoms 1,2

Fadi El Baba, MD<sup>3</sup> and Patrick Sibony, MD<sup>4</sup>

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<u>Preferred Practice Patterns</u> of the American Academy of Ophthalmology (AAOO);
Trobe JD <u>The Physician's Guide to Eye Care</u> 1993 AAOO;
Berson FG <u>Basic Ophthalmology</u> 1993 AAOO;
Collins JF, Donnenfeld ED, Perry HD, Wittpenn JR, (ed) <u>Ophthalmic Desk Reference</u>, , Raven Press 1991.

<sup>&</sup>lt;sup>1</sup>Note: These guidelines are intended to help the primary care physician decide if and when a patient needs to be referred for a variety of visual complaints. Hopefully this might reduce the need for specialty care. Needless to say it is impossible to anticipate every possible clinical circumstance and distill the problem into a one page summary per symptom that applies in all instances. There will be exceptions to every recommendation in this handout. Ultimately the decision must be based on clinical judgement and experience in dealing with eye problems. In some instances you may want to call and discuss the case by phone for advice. If there still remains some doubt about how to proceed then we suggest that you refer the patient.

<sup>&</sup>lt;sup>2</sup> Sources:

<sup>3,4</sup> Department of Ophthalmology State University of New York at Stony Brook School of Medicine and Ophthalmology Section, Surgical Service, Northport Veterans Administration Hospital

## **ASYMPTOMATIC PATIENT**

#### A. LOW RISK ADULT

AGE 20-40 Every 3 years

Check visual acuity. Refer if abnormal or if the patient has visual symptoms.

AGE > 40 Every 2 years

Complete examination every 2 years. Every 2-4 years thereafter for presbyopic corrections and check for glaucoma.

#### **B. HIGH RISK ADULT**

- H/O RETINAL DETACHMENT, OCULAR TRAUMA, VISION LOSS
- HYPERTENSION, SICKLE CELL DISEASE
- FH GLAUCOMA OR OTHER HERITABLE DISEASE
- BLACK PATIENTS (RISK OF GLAUCOMA IS MUCH HIGHER)
- > 65
- DIABETES (SEE BELOW)

Refer non urgently if risk factors present Exam every 1-2 years thereafter, unless otherwise indicated

#### C. DIABETICS

| Risk for       | $\Rightarrow$ | background           | proliferative        |
|----------------|---------------|----------------------|----------------------|
|                |               | diabetic retinopathy | diabetic retinopathy |
| diabetes 3 - 4 | years         | 18%                  | 0 %                  |
| diabetes >15   | years         | 80%                  | 25%                  |

#### I. DIABETES ONSET ages 0 - 30

Recommendation: Examination 5 years after onset, yearly thereafter.

#### II. DIABETES ONSET age > 30

Recommendation: Examination at the time of diagnosis, yearly thereafter

#### **III. DIABETES PRIOR TO PREGNANCY**

Recommendation: prior to or early in the first trimester; every 3 m thereafter

# **CHRONIC or PROGRESSIVE VISION LOSS**

#### **DIFFERENTIAL DIAGNOSIS**

|   | •   |     |             |            |            |    |    |
|---|-----|-----|-------------|------------|------------|----|----|
| _ | rat | ra  | <b>\ti\</b> | $\sim$     | $\alpha$ r | ra | ro |
| • | ref | ıaı | JUV         | <b>'</b> C | CI         | ı  | 5  |

- cataracts
- · diabetic retinopathy
- age related macular degeneration (ARMD)
- glaucoma

- optic neuropathies
- maculopathies
- corneal diseases
- psychogenic

#### **HISTORY**

| One eye or both.                                | Refractive problems usually bilateral and symmetrical |
|---|---|
| Blur at near or distance.                       | Refractive usually affects one or other               |
| Selective visual field loss.                    | Optic neuropathies, keratopathies                     |
| Blur improves by squinting or pinhole.          | Refractive  |
| Loss of color vision, color desaturation        | Optic neuropathy, maculopathy                         |
| Flare or halos with headlights or street lights | Posterior subcapsular cataracts, keratopathy          |
| Metamorphosia(wavy distortion of straight line) | Maculopathy   |

#### **EXAMINATION:**

| Visual acuity improves with pinhole or glasses           | Refractive                           |
|--|--------------------------------------|
| Corneal or lens opacification                            | Corneal scar                         |
| Afferent pupillary defect (swinging flashlight sign)     | Retinal or optic nerve dysfunction   |
| No red reflex or difficulty viewing posterior pole       | Cataract                             |
| Optic disc edema or pallor                               | Optic neuropathy                     |
| Pale nerve with cupping                                  | Glaucoma                             |
| Drusen of the retina (soft yellow exudate-like deposits) | Armd                                 |
| Retinal hemorrhages, exudates                            | Diabetes                             |
| Monocular field cuts                                     | Optic neuropathies, maculopathies    |
| Bitemporal hemianopsias                                  | Chiasmal syndrome, pituitary adenoma |
| Homonymous hemianopsia                                   | Hemispheric stroke or tumor          |

#### **REFER NON URGENTLY**

• All patients with unexplained or undiagnosed chronic progressive visual loss 5

<sup>&</sup>lt;sup>5</sup> slow, progressive decline in vision not otherwise explained by refractive errors, glaucoma or other funduscopically visible process (e.g. diabetes, ARMD, maculopathy) is tumor (due to compressive optic neuropathy) until proven otherwise. All patients with unexplained vision loss must be carefully evaluated.

# **SUDDEN MONOCULAR BLINDNESS**

#### **DIFFERENTIAL DIAGNOSIS:**

| • | Retinal detachment (RD)                 | • | Anterior ischemic optic neuropathy (AION)      |
|---|---|---|--|
| • | Vitreous hemorrhage                     | • | Optic neuritis                                 |
| • | Arterial occlusions (CRAO)              | • | Choroidal neovascular membranes                |
| • | Vein occlusions                         | • | Psychogenic                                    |
| • | Age related macular degeneration (ARMD) | • | Sudden appreciation of long-standing blindness |

#### **HISTORY:**

| Floaters and photopsia                 | Retinal detachment, vitreous hemorrhage,                 |  |  |
|--|--|--|--|
| Chromatopsia                           | Retinal artery occlusion (green or blue), vit heme (red) |  |  |
| Headaches, jaw pain, polymyalgia (GCA) | Retinal artery occlusion, AION                           |  |  |
| Painful eye movements                  | Optic neuritis   |  |  |
| Hypertension                           | Retinal artery occlusion, vein occlusion, AION           |  |  |
| Diabetes                               | Vitreous hemorrhages                                     |  |  |
| FH of retinal detachment               | Retinal detachment                                       |  |  |
| Prior H/O neurological symptoms        | Optic neuritis/MS; TIA/stroke (CRAO, AION)               |  |  |

#### **EXAMINATION:**

| Afferent pupil defect                      | CRAO, AION, retinal detachment, optic neuritis |
|--|--|
| Retinal edema, cherry red spot             | CRAO   |
| Macular hemorrhage                         | ARMD, Choroidal neovascular membrane           |
| Drusen (soft yellow exudate like deposits) | ARMD   |
| Numerous, scattered hemorrhages throughout | Vein occlusions                                |
| Optic disc edema                           | Optic neuritis (papillitis), Vein occlusions   |
| Normal posterior pole                      | optic neuritis, psychogenic, peripheral RD     |
| No red reflex, no view of fundus           | vitreous hemorrhage, small pupil               |
| Embolus                                    | CRAO, Branch retinal artery occlusion          |

| REFER IMMEDIATELY:                                    |   |
|---|---|
|   |   |
| Central retinal artery occlusion:                     | <ul> <li>painless, retinal edema, cherry red spot, afferent pupilary defect; consider<br/>carotid disease, cardiogenic emboli and giant cell arteritis</li> </ul> |
| <ul> <li>Branch retinal artery occlusion :</li> </ul> | <ul> <li>same as CRAO but confined to one quadrant <u>+</u> embolus</li> </ul>  |
| <ul> <li>Ischemic optic neuropathy:</li> </ul>        | <ul> <li>painless, pale optic disc edema, APD,</li> </ul>   |
| (i.) <u>Non-arteric</u>                               | <ul> <li>normal ESR, H/O atherosclerosis, hypertension or diabetes</li> </ul>   |
| (ii.) <u>Arteritic</u> :                              | <ul> <li>question carefully for symptoms of GCA, obtain stat ESR, any suspicion of</li> </ul>   |
|   | GCA start steroids, schedule temporal artery biopsy.  |
| Retinal detachment:                                   | <ul> <li>elevated retina, H/O photopsia and floaters</li> </ul>   |
| Vitreous hemorrhage:                                  | <ul> <li>without diabetes may be due to retinal tear or detachment</li> </ul>   |

| RE | FER URGENTLY (within 48 hour | rs)   |  |  |
|----|------------------------------|---|--|--|
| •  | Optic neuritis:              | <ul> <li>young patient, painful eye movements, normal or swollen optic disc,<br/>apd, symptoms of MS</li> </ul>   |  |  |
| •  | Retinal vein occlusion:      | <ul> <li>numerous retinal hemorrhages confined to one quadrant (branch vein<br/>occlusion) or the entire posterior pole (central vein occlusion), optic disc</li> </ul> |  |  |
|    | edema                        |   |  |  |
| •  | ARMD Vitreous hemorrhage:    | <ul> <li>localized hemorrhage confined to macular region, elderly</li> <li>w/ diabetes indicative of proliferative retinopathy.</li> </ul>                              |  |  |

# TRANSIENT VISION LOSS (TVL)

#### TRANSIENT BINOCULAR VISION LOSS (TBVL)

- Optic disc edema (Transient visual obscurations) [def : TVOs are momentary blackouts lasting seconds]
- Vertebrobasilar TIA (1-10 min)
- Migraine (15-45 min)

# B.TRANSIENT MONOCULAR BLINDNESS (TMB) THROMBOTIC/EMBOLIC

- Carotid (1 10 min) TIA
- · Cardiogenic: valvular, dysrhythmia
- Vasculitis: Temporal arteritis, Lupus, etc.
- Hyperviscosity: P Vera, Essential thrombocythemia
- Hypercoagulability: Estrogens, Antiphospholipid Antibody syndromes, Protein C or S deficiency

#### NON THROMBOTIC

- Optic disc edema (TVOs)
- Retinal migraine
- Angle closure, epithelial keratopathies
- Optic disc anomaly (optic disc drusen)
- · Benign, idiopathic of the young
- Demyelinating (Uhthoffs)
- Compressive

#### HISTORY:

| Associated cerebral ischemic symptoms                |  |
|--|--|
| diplopia, dysarthria, vertigo, ataxia                | Vertebrobasilar TIA (cardiac, Atheroemboli)                          |
| ipsilateral hemispheric symptoms                     | Carotid, cardiogenic   |
| Atherosclerotic risk factors                         | Carotid TMB, Posterior TIA   |
| Rheumatic, prosthetic valves, atrial fib, sick sinus | Cardiogenic emboli   |
| Constitutional symptoms                              | Vasculitis, hyperviscosity   |
| Birth control pill, pregnancy, post partum           | Migraine, hypercoagulability   |
| Head or neck trauma                                  | Carotid or vertebrobasilar dissection                                |
| Postural induced                                     | TVOs, high grade carotid stenosis, orthostatic                       |
| Altitudinal pattern of vision loss (like a curtain)  | Embolic mechanism: carotid or cardiogenic                            |
| Precipated by hot shower or exertion?                | Uhthoff's, (old optic neuritis)                                      |
| Palpitations, chest pain ?                           | Cardiogenic emboli   |
| Headache   | Migraine, giant cell arteritis                                       |
| Syncope, lightheadedness                             | Orthostatic hypotension, valvular                                    |
| Gaze induced TMB                                     | Compressive, hematoma or tumor of the orbit                          |
| Light induced TMB                                    | Carotid stenosis   |
| Scintillations                                       | Migraine, Vasculitis, AVM, Focal occipital seizures, occipital tumor |
|  | (see page 7)   |

#### **EXAMINATION:**

Needless to say a the patient needs complete physical examination specifically looking for a murmer, carotid, ocular or cranial bruits, diminished pulses, tenderness over the temporal arteries, hypertension, postural hypotension, focal neurological signs etc. The eye examination is oftentimes normal, however, there are some helpful findings which when present may support a specific diagnosis. The eye exam might be notable for an afferent pupillary defect (optic neuritis, Uhthoffs), retinal emboli (carotid, cardiogenic), retinal vasculitis, optic disc edema (transient visual obscurations), narrow angles, ocular hypertension (angle closure glaucoma).

#### REFER URGENTLY<sup>6</sup> (within 24 hours)

- Amaurosis fugax with elevated ESR or symptoms of GCA, start prednisone then refer
- Frequent episodes of TVL in rapid succession,
- TVL followed by persistent visual field loss (see sudden monocular blindness p 4)
- Transient visual obscurations with optic disc edema

#### **REFER NON URGENTLY**

• Rule out thrombotic-embolic causes , then refer if the etiology remains uncertain.

## **RED EYE**

#### **DIFFERENTIAL**

<sup>6</sup>Note: Transient vision loss is a complaint that does not lend itself to simple universal recommendations. So much depends on the clinical setting. In many instances the patient requires a medical or neurological workup rather than an eye exam. Ultimately it is a judgement call. In general <u>patients can be referred of an eye exam non urgently (within 1-3 weeks</u>). While TVL can be the harbinger of sudden and permanent blindness or stroke, this outcome is fortunately rare.

| • | Conjunctivitis | • | Angle closure glaucoma             | • | Orbital pseudotumor      |
|---|----------------|---|------------------------------------|---|--------------------------|
| • | Blepharitis    | • | Uveitis                            | • | Thyroid orbitopathy      |
| • | Stye           | • | Keratitis (herpes, corneal ulcers) | • | Orbital cellulitis       |
| • | Subconj heme   | • | Neovascular glaucoma               | • | Scleritis , episcleritis |

#### **HISTORY**

| Visual acuity  |                          | Vision normal in conjunctivitis                               |
|----------------|--------------------------|---|
| Pain           |                          | Angle closure, keratitis, scleritis, episcleritis are painful |
| Photophobia    |                          | keratitis, uveitis  |
| Halos          |                          | Sign of corneal edema in angle closure                        |
| Itchy          |                          | Allergic conjunctivitis                                       |
| Discharge ?    | Purulent                 | Bacterial conjunctivities                                     |
|                | Serous                   | Viral conjunctivitis  |
| Eyelids matted | and stick together in AM | Bacterial conjunctivitis                                      |
| Floaters       |                          | Uveitis   |

#### EXAM:

| Vision abnormal in angle closure, uveitis, keratitis,  |
|--|
| Fixed/mid dilated (angle closure), small/fixed or irregular (uveitis)  |
| Elevated in angle closure, may be low in uveitis   |
| Keratitis  |
| Thyroid, orbitopathy, orbital pseudotumor, scleritis   |
| Thyroid, orbitopathy, orbital pseudotumor, scleritis   |
| Episcleritis, scleritis  |
| Thyroid, orbitopathy, orbital pseudotumor, scleritis allergic conjunctivitis   |
| Marginal erythema (blepharitis), upper lid retraction (thyroid), ptosis and swelling(pseudotumor, scleritis, orbital cellulitis) |
| Angle closure, neovascular glaucoma, keratitis, (uveitis)  |
| Bacterial corneal ulcer  |
|  |

| RE | FER IMMEDIATELY:        |  |
|----|-------------------------|--|
| •  | Angle Closure Glaucoma: | painful red eye, hazy cornea, mid dilated fixed pupil, elevated pressure |
| •  | Corneal Ulcer:          | opacified, white corneal infiltrate, red eye, purulent discharge         |

#### **REFER URGENTLY (within 24 - 48 hours)**

| K | EFER URGENILI (WILIIII) 24 - 40 NOI | urs) |                   |   |                         |
|---|-------------------------------------|------|-------------------|---|-------------------------|
| • | Pain                                | •    | Photophobia       | • | Blurred vision          |
| • | Proptosis                           | •    | Ophthalmoplegia   | • | Ciliary flush           |
| • | Irregular corneal refex             | •    | Epithelial defect | • | Pupil fixed or sluggish |
| • | Worsenig after 3 d treatment        | •    | Compromised host  |   |                         |

#### TREAT:

**Blephartis:** gritty, burning, matting, scaling or flaking of lid, mild conjunctival injection. Apply Bacitracin ophthalmic to eyelid HS, Commercial lid hygiene solution (e.g. Eye-scrub qAM) Refer non urgently if symptoms persist.

#### Conjunctivitis:

**Bacterial:** topical antimicrobial medications (e.g. Polytrim QID), refer if redness fails to resolve after 3 days **Viral:** frequent handwashing, non communal activity, no antibiotics needed. Refer urgently if vision blurs, photobic or other signs of keratitis develop.

**Stye:** warm compresses, antibiotic eyedrops, Bacitracin ophthalmic ointment at bedtime. Refer non urgently if it fails to resolve after 1 week. for incision and drainage

Allergic conjunctivitis: topical decongestants (e.g. Naphcon A QID) for symptomatic relief of itch.

Subconjunctival hemorrhage: spontaneous, benign, no treatment required.

# FLASHES, PHOTOPSIA AND SCINTILLATIONS

#### **DIFFERENTIAL**

| RETINAL PHOTOPSIA                                       | CORTICAL SCINTILLATIONS                            |
|---|--|
| momentary bright flashes of light                       | scintillating zig zag lines or colored lights      |
| lasting seconds at most                                 | lasting 2-45 minutes +/- scotomas                  |
| Retinal traction  | Migraine (15-45 min)                               |
| Retinal tear  | <ul> <li>Vertebrobasilar TIA (2-10 min)</li> </ul> |
| <ul> <li>Posterior vitreous detachment (PVD)</li> </ul> | Seizure  |
| Retinal detachment                                      | Arteriovenous malformation                         |

#### **HISTORY and EXAM**

| Duration is single most helpful clue                            | Seconds: retinal                                    |
|---|---|
|   | 2-10 min: TIA                                       |
|   | 15-45 min: migraine                                 |
| Scintillations march across the visual field ("spectral march") | Migraine ( seizures are stereotyped and stationary) |
| Induced by eye or head movement                                 | Retinal photopsia                                   |
| Floaters  | Retinal hole, retinal detachment, PVD               |
| Headache (typically throbbing, unilateral etc)                  | Migraine  |
| Vertigo, diplopia, ataxia, speech etc                           | TIĀ   |
| H/O myopia, FH retinal detachment or trauma                     | Retinal tear, retinal detachment                    |
| Audible cranial bruits, h/o seizures                            | AVM   |
| Associated homonymous hemianopsia                               | Migraine, TIA, AVM                                  |

#### REFER EMERGENTLY

- Observed retinal detachment, absent red reflex or vitreous hemorrhage,
- Photopsia associated with decreased vision, visual field cut or floaters.
- Cortical scintillations with persistent neurological deficits: hemianopsias, hemiparesis (obtain MRI); refer to neurology.

#### **REFER URGENTLY (within 48 hours)**

New onset photopsia or marked worsening of pre-existant chronic photopsia

#### **REFER NON URGENTLY**

- Chronic or recurrent flashes
- Vertebrobasilar TIA: start antiplatelets, neurovascular workup, R/O cardiogenic or vasculitis

#### **TREAT**

Migraine

# **FLOATERS**

Grey spots, cobwebs, black spots that appear to drift or lag with eye movement

#### **DIFFERENTIAL**

| • | Physiologic entopic phenomena       | • | Retinal detachment              |
|---|-------------------------------------|---|---------------------------------|
| • | Posterior vitreous detachment (PVD) | • | Vitreous hemorrhage             |
| • | Retinal tear, hole                  | • | Vitreous inflammation (uveitis) |

#### **HISTORY**

| Sudden onset in an elderly or a high myope                           | PVD, vitreous degeneration       |
|--|----------------------------------|
| Showers of floaters, associated with flashes and/or decreased vision | Retinal tear, retinal detachment |
| New onset floaters in a diabetic                                     | Vitreous hemorrhage              |
| Red eye, pain, photophobia, blurred vision                           | Vitreous inflammation            |

## REFER URGENTLY

- New onset floaters associated with vision loss (see SUDDEN MONOCULAR BLINDNESS)
- New onset floaters in diabetics, vitreous hemorrhage
- Red eye and floaters

## **REFER NON URGENTLY**

Chronic floaters

# **TEARING (EPIPHORA)**

#### **DIFFERENTIAL**

| OVERPRODUCTION   | POOR DRAINAGE  | REFLEX TEARING   |
|--|--|--|
| <ul> <li>Blepharitis</li> <li>Conjunctivits</li> <li>Keratitis</li> <li>Uveitis</li> <li>Orbital inflammatory disease</li> <li>Thyroid orbitopathy</li> <li>Orbital cellulitis etc.</li> </ul> | Eyelid deformity (poor apposition of the lower eyelid)      cicatricial lid retraction     facial nerve palsy     ectropion     others | Dry eyes     - idiopathic     -Keratitis Sicca     -Corneal foreign body     -Trichiasis (eyelash) |
| See red eye p. 6   | Nasolacrimal outflow obstruction:     -congenital     -dacryocystitis     -trauma     -nasolacrimal tumor     -sinus tumor             |  |

#### **HISTORY and EXAM**

| Red eye, pain, photophobia                                      | Inflammatory (see RED EYE)   |
|---|------------------------------|
| Tenderness , swelling, erythema over lacrimal sac               | Dacryocystitis               |
| Purulent reflux from canaliculus induced by pressure on the sac |                              |
| History of Bell's palsy, facial burn, trauma                    | Appositional lid deformity   |
| Unilateral, since birth   | Congenital nasolacrimal duct |
|   | obstruction                  |
| Dry mouth, rheumatic disease                                    | Keratitis sicca              |

#### **REFER URGENTLY**

- See RED EYE if this appears to be inflammatory in origin.
- Dacryocystitis
- Embedded foreign bodies not removable with cotton swab

#### **REFER NON URGENTLY**

- · Refer newly acquired cases, if due to eyelid deformity
- Dry eyes that fail to respond to topical lubricants
- Progressive or intolerable epiphora

#### TREAT:

- Foreign body, if easily removed
- Symptomatic dry eye with topical lubricants
- See guidelines for RED EYE

## **DIPLOPIA**

#### **DIFFERENTIAL**

#### MONOCULAR DIPLOPIA: **BINOCULAR DIPLOPIA** persistent diplopia with monocular occlusion, diplopia with both eyes viewing, resolves with monocular occlusion of either localizes to one eye due to an optical aberration eye; due to an ocular motor misalignment Cataracts Ocular myopathy: thyroid, myasthenia Refractive error Orbital tumor or fracture Vitreous opacity Cranial neuropathy: iii, iv, vi Corneal scar Central: nuclear, internuclear or supranuclear e.g. Internuclear ophthalmoplegia, skew deviation due to Retinal elevation (rare) midbrain, pontine, cerebellar or medullary dysfunction. Cerebral polyopia (rare) Vergence disorders: e.g. convergence insufficiency Psychogenic Decompensated strabismus Convergence spasms (psychogenic)

#### **HISTORY:**

| Monocular "ghost" image              | Refractive or cataract   |
|--------------------------------------|--|
| Vertical or horizontal separation    | Distinguishes between horizontal vs vertical recti                           |
| Worsens at distance or near          | Abduction weakness worse at distance, adduction weakness worse at            |
|                                      | near. Convergence insufficiency symptomatic when reading.                    |
| Worsens with left or right gaze      | Strabismus constant in all directions of gaze, ophthalmoplegias worsen       |
|                                      | when looking towards the field of action of a paretic muscle.                |
| Worsens with head tilt left or right | Superior oblique palsies typically worsen on ipsilateral head tilt.          |
| Ptosis                               | III rd nerve palsies, myasthenia, orbital tumors                             |
| Headache                             | Ischemic cranial neuropathies, aneurysmal iii n palsies, orbital             |
|                                      | pseudotumor, concurrent trigeminal neuropathy (cavernous sinus               |
|                                      | syndrome).   |
| Red eye or proptosis                 | Orbital pseudotumor, thyroid orbitopathy, carotid cavernous fistula, orbital |
|                                      | tumors   |
| Blown pupil                          | Pupil involving iii n palsies often due to aneurysms but less commonly can   |
|                                      | also be ischemic   |
| H/O amblyopia, eye muscle surgery    | Strabismus   |
| History of trauma                    | Cranial neuropathy, orbital fractures, convergence insufficiency             |
| Other neurological complaints        | Cranial neuropathy, central  |
| Diurnal variation: worse in AM       | thyroid orbitopathy  |
| worse in PM                          | ocular myasthenia, decompensated strabismus                                  |

#### **Examination:**

In addition to a careful evaluation of eye movements in all the cardinal positions of gaze, the patient must be careful examined for signs of ptosis, anisocoria, pupil reactivity, lid swelling, proptosis, redness, corneal sensation, facial sensation and bruits.

#### REFER URGENTLY 7

- Acquired and persistent binocular diplopia
- Acquired, painful, pupil involving III n palsy (without a history of diabetes) is aneurysmal or neoplastic
  until proven otherwise. Obtain MRI/MRA urgently.

| REFER NON URGENTLY                 |   |
|------------------------------------|---|
| Monocular diplopia,                | <ul> <li>transient diplopia,</li> </ul> |
| intermittent diplopia when reading | chronic binocular diplopia.             |

<sup>&</sup>lt;sup>7</sup>Note: Imaging studies in recently acquired cases of diplopia are not always necessary e.g. IV n palsies, thyroid orbitopathy, many disorders of vergence, decompensated phoria, ocular myasthenia, pupil sparing diabetic III nerve palsies.

## **ANISOCORIA**

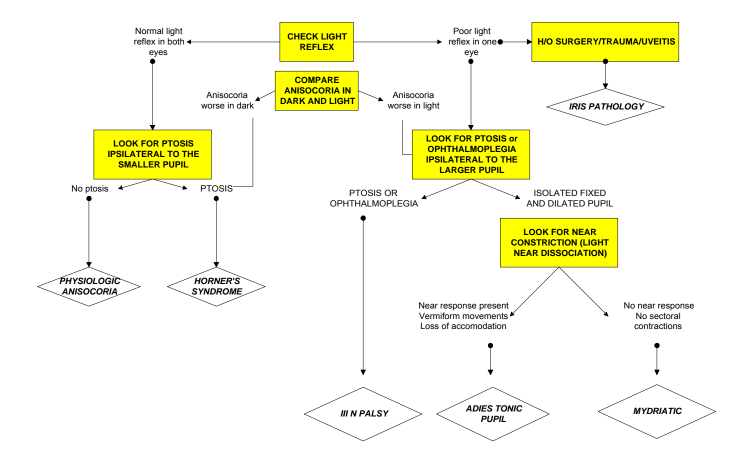
#### **DIFFERENTIAL**

#### SMALL PUPIL

- Horner's syndrome
- Iris synechia: old uveitis, previous surgery
- Chronic Adies tonic pupil
- Physiologic anisocoria

#### **DILATED**, FIXED PUPIL

- Iris pathology: sphincter tear, iris atrophy
- Mydriatics: atropine, scopalamine, mydriacil, cyclogyl
- Adies tonic pupil
- III rd nerve palsy
- Physiologic anisocoria



#### **REFER URGENTLY**

Anisocoria with ptosis or ophthalmoplegia

#### **REFER NON URGENTLY**

Isolated anisocoria

## **OCULAR TRAUMA**

#### TREAT ON SITE AND REFER IMMEDIATELY

Acid or alkalai burn

#### REFER IMMEDIATELY

- severe pain
- · deformed globe
- · eyelid lacerations which
- -involve the lid margin
- -canaliculus
- -deep, prolapsed fat

- new onset subnormal acuity
- · corneal or scleral laceration
- hyphema
- ? intraocular foreign body
- loss of red reflex

- irregular pupil
- · corneal clouding
- · severe lid swelling
- severe conjuctival chemosis
- proptosis

#### **REFER URGENTLY (within 48 hours)**

- Pain
- foreign body sensation
- suspected orbital wall fracture
- photophobia
- large corneal abrasion
- moderate eyelid swelling or chemosis with normal vision
- diplopia
- suspected contusion of globe

#### **TREAT**

- minor corneal abrasions
- removable foreign bodies (note if there is a history of risk of high velocity foreign body patient needs dilated exam to check for occult penetration of the eye)
- superficial brow and lid lacerations that do not involve the lid margin or canaliculus
- periorbital soft tissue injury without change in vision or evidence of ocular contusion

# **SYSTEMIC DRUGS: OCULAR TOXICITY**

(RECOMMENDATIONS FOR MONITORING)

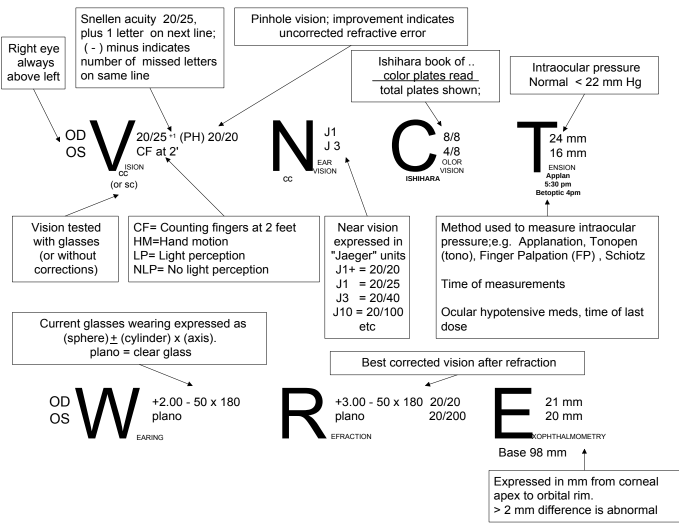
| DRUG            | Complications  | Recommendations   |  |  |
|-----------------|--|---|--|--|
| AMIODARONE      | All corneal deposits ( "whorls")     Reversible when stopped     Symptoms of halos, blur are unusual     Optic neuropathy (rare)   | Refer patients with subnormal vision or symptoms.     Discontinue if symptomatic.     The mere presence of deposits is not in and of itself a reason to discontinue |  |  |
| ANTICHOLINERGIC | Loss of accomodation     Angle closure glaucoma  | Refer for refraction if symptomatic     Refer if angle is narrow or for painful red eye     Open angle glaucoma is not a contraindication                           |  |  |
| CHLOROQUINES    | <ul> <li>&gt;300 g total cumulative dose (3 yrs)</li> <li>"bulls eye" maculopathy</li> <li>Corneal deposits</li> </ul>   | Baseline exam     Follow up q 6 months  |  |  |
| CORTICOSTEROIDS | Cataracts,     Glaucoma     Pseudotumor cerebri  | Refer for slow, decline in vision or transient visual obscurations.     Eye exam q6 months  |  |  |
| DIGITALIS       | Xanthopsia (yellow vision)     Flickering or snowy distortion     Rarely optic neuropathy  | Check blood level and adjust accordingly.     Refer if blood level is normal with symptoms or subnormal vision.   |  |  |
| DILANTIN        | Vestibulocerebellar signs and symptoms     Diplopia, oscillopsia, blurring     Gaze evoked nystagmus   | Check dilantin level and adjust accordingly if in the toxic range.  |  |  |
| ETHAMBUTOL      | <ul> <li>Dose related optic neuropathy as early as 1 m after starting the drug. Reversible early on.</li> <li>At 15 mg/kg incidence &lt; 1%</li> <li>At 20 mg/kg incidence 5%</li> </ul> | <ul> <li>Refer for baseline exam</li> <li>Follow-up every 6 months.</li> <li>Refer urgently for any visual decline.</li> </ul>                                      |  |  |
| THIORIDAZINE    | Pigmentary retinopathy at doses of >1000mg /d  | Maximum dose recommendation 800mg/d     Refer for symptoms  |  |  |

# **OPHTHALMIC MEDICATIONS**

## SYSTEMIC AND OCULAR SIDE EFFECTS

| CLASS            | DRUG   | OCULAR  | SYSTEMIC   |  |  |
|------------------|--|---|--|--|--|
| ANESTHETICS      | Proparicaine     Tetracaine  | Epithelial keratopathy     should be restricted for exam only, never to be used as an analgesic                                   | • none   |  |  |
| ANTIMICROBIALS   | Neomycin ( many brands)     Gentamicin ( many brands)     Tobramycin (Tobrex)                                  | Eyelid or facial dermatitis     Keratitis with long term use  | none   |  |  |
|                  | Erythromicin (Ilotycin)     Ciprofloxicin (Ciloxan)  | none     corneal deposits   | • none • none  |  |  |
|                  | Norfloxacin (Chibroxin)  | ·   |  |  |  |
|                  | Polymixin     Trimethoprim-polymixin (Poly trim)   | <ul><li>none</li><li>none</li></ul>   | none     none  |  |  |
|                  | Sulfacetamide  | eyelid dermatitis   | Stevens Johnson  |  |  |
| ANTIVIRALS       | Trifluridine (Viroptic) Vidarabine (Vira A) Idoxiuridine (Herplex, Stoxil, Dendrid) Acyclovir (Zovirax)        | epithelial keratopathy     conjunctivitis     lacrimal punctal stenosis   | • none   |  |  |
| ARTIFICIAL TEARS | many brands  | • none  | • none   |  |  |
| GLAUCOMA         | Epinephrine (Epifren, Glaucon)     Dipivefrin (Propine)  | conjunctival hyperemia     black conjunctival deposits  | <ul> <li>tachycardia</li> <li>PVCs</li> <li>hypertension</li> <li>tremor</li> <li>anxiety</li> <li>Bradycardia</li> <li>Bronchospasm</li> <li>hypotension, syncope</li> <li>reduced libido</li> <li>lethargy and depression</li> </ul> |  |  |
|                  | Timilol (timoptic) Betaxalol (betoptic) Levobunolol (Betagan) Carteolol (Ocupress) Metipranolol (Optipranolol) | no significant complications  |  |  |  |
|                  | Acetozolamide (Diamox)   | induced myopia  | <ul> <li>Stevens Johnson</li> <li>Renal stones</li> <li>Paresthesias</li> <li>Nausea</li> <li>Dysgeusia</li> <li>Anorexia</li> <li>lassitude</li> <li>Loss of libido , Impotence</li> <li>Acidosis</li> <li>Aplastic anemia</li> </ul> |  |  |
| CHOLINERGICS     | Pilocarpine  | <ul><li>constriction</li><li>conjunctival injection</li><li>induced myopia</li></ul>  | <ul> <li>Headache or brow ache</li> <li>cramping, vomiting</li> <li>diarrhea</li> <li>diaphoresis</li> <li>bronchospasm</li> <li>unstable BP</li> </ul>  |  |  |
| STEROIDS         | Prednisilone (many brands)     Dexamethasone (many brands)     Medrysone (HMS)     Fluoromethalone (FML)       | ocular perforations in patients with necrotizing inflammation     glaucoma     cataract     exacerbate viral and fungal keratitis | • none   |  |  |

# HIEROGLYPHICS OF THE EYE EXAM



LIDS: LF = lid fissure

PUPILS: APD = afferent pupillary defect

SLE: = (SLIT LAMP EXAMINATION)

**CONJ:** (= CONJUNCTIVA)

CORNEA: (= K)

A/C: (= ANTERIOR CHAMBER)

IRIS: PI = peripheral iridectomy

LENS: PSC=posterior subcapsular cataract, NS=nuclear sclerotic cataract

GRADING CATARACT DENSITY: 1+ (mild) to 4+(severe) PCIOL = POSTERIOR CHAMBER INTRAOCULAR LENS,

ACIOL = ANTERIOR CHAMBER IOL

#### **MOTILITY:**

**OCULAR MISALIGNMENT EXPRESSED IN PRISM DIOPTERS (PD)** 

1 PD = light displaced by 1cm at 1 m

- PHORIA is a latent misalignment
- TROPIA is a manifest misalignment.
- NOTATION USED TO QUANTITATE MISALIGNMENT:
  - 1. ORTHO = both eyes aligned

EX = 0

#### 2. AT DISTANCE -

a. ESODEVIATIONS (eyes crossed)

E = esophoria

ET = esotropia

**b. EXODEVIATIONS** 

X = exophoria

XT= exotropia

c. HYPERDEVIATONS (one eye higher relative to the other; by convention lateralize to the upper eye even if the lower eye is abnormal)

RH = right hyperphoria

RHT = right hypertropia

LH = left hyperphoria

LHT = left hypertropia

#### 3. AT NEAR

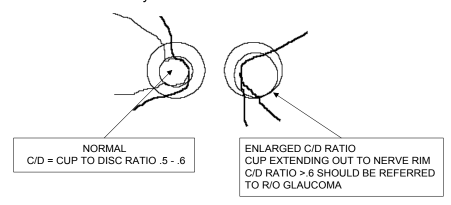
same as above with PRIME e.g. ET', X', LHT'

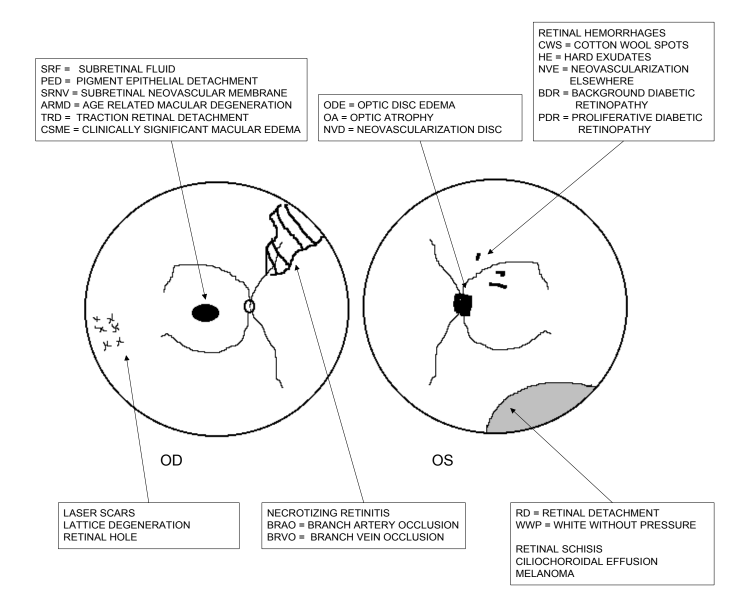
4. Example: Grid shows misalginement in patient's cardinal positions of gaze i.e. 12 prism diopters of left hypertropia in right gaze, 2 prism diopters of left hyperphoria in left gaze, etc. This particular example demonstrates an incomitant misalignment that worsens when looking down and to the right which is typical of a IV nerve palsy. This grid can also be used to document the direction of the fast phase of nystagmus in various positions of gaze by using arrows of varying size to also document its amplitude or intensity.

| RIGHT LEFT |               |      | _              |      |
|------------|---------------|------|----------------|------|
|            | 2 LHT<br>2 XT |      |                | UP   |
| 12 LHT     | 4 LHT         | 2 LH | 1 <sup>0</sup> |      |
| 16 LHT     | 5 LHT<br>4 ET |      |                | DOWN |

## FUNDUS EXAMINATION: (dilated; undilated)

Diagrams are often used to document fundus findings. Examples of common abbreviations and notations used to document a variety of abnormalities are shown below.





# **COMMON ABBREVIATIONS:**

AION Anterior ischemic optic neuropathy

**ALT** Argon laser trabeculoplasty

AMD or Age related macular degeneration

ARMD

APD Afferent pupillary defect

BDR Background diabetic retinopathy
BRAO Branch retinal artery occlusion
BRVO Branch retinal vein occlusion
CRAO Central retinal artery occlusion
CRVO Central retinal vein occlusion

**CSME** Clinically significant macular edema

CWS Cotton wool spot

FRP Focal retinal photocoagulation

HE Hard exudate

LTG Low tension glaucoma
 NVD Neovascularization at disc
 NVE Neovascularization elsewhere
 PACG Primary angle closure glaucoma
 PDR Proliferative diabetic retinopathy
 POAG Primary open angle glaucoma
 PPDR Preproliferative diabetic retinopathy

PRH Preretinal hemorrhage

PRP Panretinal photocoagulation
PVD Posterior vitreous detachment

RD Retinal detachment

RPE Retinal pigment epithelium

SRF Subretinal fluid

SRNV Subretinal neovascularization TRD Traction retinal detachment

VH Vitreous hemorrhage