Lumps and Bumps and Dots – Oh My!

A Posterior Segment Review
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Patient here for “routine exam.” We see… this!

And then this…
So, let’s start with: “Oh my...”

Challenges
- Chronic (Ex. Glaucoma)
  - You have time.
  - You have time to second guess yourself.
  - You have time to second guess yourself and come up with a worse decision.
  - Also action taken will naturally depend on the doctor's comfort level and the case at hand.
- Acute (Ex. Sudden onset anything!)
  - Often, little time to ponder possibilities, but...
  - Many cases will require secondary/tertiary care
  - Second guessing occurs later.
  - Patient education will take on immediacy, just like all the other aspects of care.

And On to Dots...

APMPPE
Acute Posterior Multifocal Placoid Pigment Epitheliopathy
**HISTORY**

- 18 year-old woman
- "Black spot blocking my vision RE and several in LE."
- Neighbor’s house has many bird nests; rashes on legs after an "outing."
- Recent bronchitis and UTI
- PMHx: Asthma
- Meds: Prednisone for asthma from Health Center (25 mg, tapering dose)

**OCULAR EXAMINATION**

- VA: OD CF of 5’ OS 20/40
- Pupils: PERRLA; no APD
- EOM: Full
- VF defect: OD central defect OS superior defect
- IOP: OD 18 OS 18
- SLEx: OD NL state OS NL state

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18 year old woman; OD

OD

OD: 17 sec.

OD: 4:22 min.
Subjectively, "no significant change."

VA: OD 20/20, OS 20/20
Pupils: PERRLA
IOP: OD 14, OS 14
SLEs: OD NL, OS NL

F/U 2 weeks; OD
FOUR WEEK FOLLOW-UP

- New visual loss OS “for 5 days.”
- “Left eye is now worse.”
- Note: OD improved in the third week; statement reflects any change after the initial improvement
- Meds: Prednisone [increased initially to 60 mg, tapering dose]

Here we go again

- VA: OD 20/25 OS 20/100
- Pupils: PERRL
- IOP: OD 15 OS 14
- SLEx: NL State OU

OS

Finally... seven weeks from initial presentation

VA: OD 20/15 OS 20/50
All externals NL state OU
VF 30-2
Nothing wrong
Patient’s imagination

39-Year-Old Man

- “I’ve lost vision in my left eye. Comes and goes.”
- Long-standing Hx of ophthalmic migraine; has never had the headache until recently
- Borderline high BP
- No medications

Physical Exam

- VA: OD 20/15 OS 20/50
- All externals NL state OU
- VF 30-2
- Nothing wrong
- Patient’s imagination
Internal
- Case profile atypical.
- Moderate vitritis noted OS
- OD P slightly elevated OS
- OD NL state
- OS nerve fiber layer swelling
- Patient referred to retinologist for evaluation and co-management.

One week later
- "Vision much worse."
- VA: OD 20/15  OS 20/100
- Pupils: Subtle A.FD OS
- Fundus evaluation:
Remember the headaches?
- Referral for neurological consult and imaging
- Rectangular mass 17X6 mm at the level of the Pons/CN V, left side
- Treated as a "Schwannoma" or a "meningioma."
- No impingement on optic nerve/chiasm
- Is there an association with retinal findings?
  - Seemingly unrelated
  - However, literature tends to support a possible association (not necessarily causative)

Cerebellopontine Angle

Facts About CPA Tumors
- 6-10% of intracranial tumors
- Vestibular schwannomas and meningiomas two most common types
- Acoustic neuroma: a misnomer
- Arise from Schwann cells
- Connection between vestibular schwannoma and neurofibromatosis type 2 (central)
- Genetic
  - Connection with development of retinal lesions
Our patient

- Final examination 11 months after the initial presentation and after removal of the tumor
- VA: OD 20/15  OS 20/400
- Screening Visual Field: OS nasal VF loss with partial crossing of midline
- Patient never returned for threshold testing
- Patient moved out of area and was lost to follow-up

CNS and Retinitis

- Althaus et al. case report suggests a full neurological evaluation may be indicated with APMPPE cases due to possibility of cerebral vasculitis, which may be manifested later (up to years)
- University of Kentucky Study (O’Halloran et al.)
  - 21 males, 10 females ranging in age 8-54, median of 27
  - 38% of patients had antecedent viral illness
  - VA variable from 20/20 to CF
  - CNV findings ranged from HA to sagittal sinus thrombosis
  - Recommendations: LT/CT imaging along with Oo of APMPPE
  - To immunosuppressive agents along with steroids if case is complicated by CNS arteritis

Segway into...

Inflammatory
- Iatrogenic-post-surgical
- Sarcoid
- Ocular sarcoidosis
- Tuberculosis
- Histoplasmosis
- Toxoplasmosis
- Syphilis

Tumors
- Choroidal
- melanoma
- metastatic/melanotic
- Metastatic
- Scleral hemangioma
- Osteoma
- Retinal hemangioma

How to tell the difference?

- Case history
- Physical exam
- Other accompanying characteristic ocular manifestations
- Lab testing/biopsy/imaging

Irvine-Gass

- 72 Year Old male
  - In Jan of 2014 patient reported for annual exam (established)
  - In 2015 had complained of glare, but tolerable
    - General health good
    - O/D: “I have problems with glare, especially at night, which has become more noticeable”
      - In addition: “I have been having dizzy spells; they keep changing my lip medication dose, but no answers yet.”
    - Noted 2+ NS OU same as previous visit, but patient more symptomatic
  - BAT 2015: 20/30 OD, OS; 2016: 20/100 OD, OS
  - Refer to OMD for cataract evaluation

Post-Op 1 week (Feb 2016)

- Patient feels OD vision is worse now than it was before surgery
  - Love hearing that, don’t you?
  - VA: OD 20/15  OS 20/20
  - No problem; must be the healing process
  - Moderate A/C reaction/Od
  - Internal health appears normal
  - Continue with steroid as 6/cld 4/cld three weeks or one month post-op or as needed
Post-Op 1 month (March, 2016)

- Patient unhappy with right eye’s vision
- VA 20/40 and 20/25
- A/C reaction still present (fewer cells, but as much or perhaps an increase of flare)
- Macular reflex affected

Macular Edema

- Exact cause unknown; most likely inflammatory
- Can be associated with:
  - Retinal vein occlusion
  - Uveitis
  - Diabetes
  - Most commonly seen post cataract surgery
  - 1/2 of cases
  - 50% chance of second eye involvement
  - Can involve any area (of literature)
  - My experience has been same

- Treatment varies from topical to injection of steroid or anti-VEGF

Continued

- Increased dose of Durezol
- Included NSAD
- RTC 2 weeks

Two weeks later

- “I have double vision and my right eye is no better.”
- VA 20/50, 20/30
- A/C continues to be mildly to moderately inflamed
- Refer back to surgery center to consider injection given symptoms and OCTs
Two-week follow-up

OD

OS

May 2016

- Dr. Kohne’s healing touch:
  - Due to improvement, no injection was given, but the treatment was continued using only topical means.
  - Patient notices improvement of vision and all the other symptoms.
  - VA 20/40, 20/25.
May 2016

- Patient missed a two-week follow-up
- We called the patient, who stated eye is better
- Rescheduled visit for June

June 2016

- "My vision is bad again, like it was right after the surgery."
- Compliant with medication, steroid now tapered to only once a day
- VA: 20/40 20/25
- Could this be the endpoint of the incident? Could this be as good as we can make it (considering the decrease in CME in May)?
- We are not done

Continued

- Called the surgery center immediately and spoke to one of the doctors (the surgeon was not available).
- Requested another evaluation and re-emphasized consideration of injection.
- As of writing this, patient is at the surgery center being prepared for the first injection.

Sarcoidosis

- African American b/w 20-50 y/o in US
- Mostly females
- Multifaceted disease
- 27-50% have ocular involvement
- Systemic: cutaneous, parotid involvement, facial palsy, arthritis, liver, spleen, etc.
Sarcoid

- Ocular manifestations
  - Granulomatous anterior uveitis
  - Conjunctival granuloma
  - Band keratopathy
  - "Candlewax" drippings on veins
  - Chorioid granuloma
  - Peripapillitis
- Laboratory testing
  - ACE, serum lysozyme, chest X-ray revealing hilar adenopathy, gallium scan, increased calcium

Parotid Gland Enlargement R > L

Ocular Toxocariasis

- Parasite – infects dogs and cats
- Occurs between 2-40 y/o, but predominantly between 7-8
- Usually from eating dirt or under-cooked meat
- Found in eye secondary to decreased VA, leukocoria or strabismus
- Mild-severe iritis/vitritis
- Diagnosed by ELISA for Toxocara
Tuberculosis

- Chronic infection caused by *Mycobacterium tuberculosis*
- Rare, but can cause uveitis
- Ocular manifestations
  - Yellow-white, multifocal choroidal lesion
  - Granulomatous/non-granulomatous anterior uveitis
- Laboratory testing
  - PPD test, chest X-ray
Histoplasmosis
- Fungus that presents in 20-50 y/o whites living near or visited the Ohio-Mississippi River Valley (and several other river valleys throughout the world)
- 60% of cases bilateral
- **lacks any uveitic (anterior and posterior) involvement**
- Affected patients over 30 have higher risk for macular involvement
- Predominantly a clinical diagnosis

Ocular manifestations
- Classic triad
  - Peripheral punched-out lesions
  - Peri- or circum-papillary atrophy
  - Neovascularization of the macular (sub-retinal neovascularization)
- **lacks any uveitic manifestations**
- Testing
  - FA if net is present or suspected
  - OCT
  - HLA-B27
Toxoplasmosis

- Obligate intracellular protozoan
- Most common cause of posterior uveitis
- Usually passed from mother to child
- The mother usually gets the parasite by exposure to feces
- Can be caused by under-cooked meat, breathing in spores or drinking contaminated water
- Ocular manifestations
  - Unilateral, multifocal chorioretinal lesions
  - Severe posterior uveitis response
  - Classic “headlights in the fog”
- Laboratory testing
  - Complement fixation test (positive active disease), Sabin-Feldman methylene blue test, ELISA, HIV test

Syphilis

- Ocular manifestations
  - Depends on stage
    - Stage 1: chancre
    - Stage 2: chorioretinitis, uveitis, optic neuritis, conjunctivitis, episcleritis, scleritis
    - Stage 3: optic atrophy, A. Robertson pupil
- Lab testing
  - RPR, VDRL (current infection), FTA-ABS, MHA-TP (past infection)

Retinal and Choroidal Tumors

- Usually distinguished by their characteristic appearance
- Ocular testing done such as FA, ICG, OCT, and B-scan, as well as laboratory and systemic testing to rule out markers

Choroidal Melanoma

- Most common primary intraocular tumor in adults
- Accounts for 85% of all uveal malignant melanomas – mainly due to vascular supply
- Women slightly more at risk
- More often unilateral and diagnosed in people over 50 years of age
- More common in whites
Continued

- Typically asymptomatic initially. However, patient may present with one or more of the following:
  - Blurred vision, VF loss, and flashes/floaters
- Two varieties:
  - Circumscribed
  - Diffuse
- Radiation is the most common treatment modality
- Enucleation is an option for high risk melanomas
- Concern about seeding must be weighed

Amelanotic Choroidal Melanoma

- Appearance
  - Indistinct borders
  - Yellowish-white
  - Fairly large diameter and thickness
  - Can produce overlying drusen
  - Brown mottling surrounding the lesion
- Testing
  - Ultrasound, FA, ICG
  - Systemic testing should be done to completely rule out metastasis
  - Colonoscopy, Gastrointestinal exam, mammogram, test of abdomen and pelvis
47-Year-Old Woman

- Long-term patient at the clinic
- Long-standing melanotic nevus OS
  - 2 x 1.5 DD, approximately 2 DD superior temporal to macula
  - Stable since first exam in early 2000s.
- Generally healthy, non-smoker, mother of three, no Ptsx.
- April 2016 exam
  - No change in health status
  - No new medications
  - But no one said a thing about OD, including your truly.

Referral

- The lone retinal specialist in town did not accept the patient's insurance
- Referred to OMD, general practitioner
  - Recommended more &-scan
  - Recommended blood work
  - Awaiting results and communication from OMD
- As of July 20, 2016
  - OMD recommends observation only.

Choroidal Metastasis

- More common than primary malignancies
- Choroid most common site for metastasis (80%), due to high metabolic rate and perfusion, as stated before
- Common primary sites
  - Females: breast
  - Males: lung
Continued

- Ocular manifestations
  - Creamy yellow in appearance
  - Can cause RD
  - Vitrreal hemorrhage
  - Can cause proptosis in extreme cases
  - Treatment will depend on size, location, patient age and health
- Tests:
  - Ultrasound, FA (hyperfluorescence in late arterial and early venous)
  - Systemic testing to find primary site if not already known

Solitary Choroidal Hemangioma

- Ocular manifestations
  - Reddish-orange in color with oval or circular shape (classic “tomato cayup” retina)
  - Can create subretinal fluid in macula
  - Can cause CME, exudative RD and secondary glaucoma
  - Usually no treatment unless subretinal fluid threatens the macular region
- Testing:
  - B-scan (solid dome tumor, high reflectivity, acoustic solidity)
  - FA (early filling, late leakage)

Choroidal Osteoma

- Slow growing intraocular bone-like tumor
- Usually unilateral (~75%)
- Found in healthy young females
- Typically near the optic nerve or in the posterior pole
- Symptoms depend on location
Continued

- Ocular manifestations
  - Yellow-white to red-orange in color
  - Edges are scalloped
  - Can cause macular detachment or neovascularization
  - Prognosis depends on location
  - Any growth suggests malignancy until ruled out

- Tests:
  - B-scan (high reflectivity)
  - CT, X-ray, and FA
    - Mottled early hyperfluorescence that intensifies throughout the angiogram cycle

Retinoblastoma

- Most common intraocular malignancy in infants and children
- 2nd most common intraocular tumor in the general population
- Approximately 30% bilateral
- Average age of diagnosis is 18 months
- Can be inherited (only 7%) or sporadic
- Two types:
  - Endophytic (from retina into vitreous)
  - Exophytic (in subretinal space)

- Leukocoria – most common presenting sign
- Strabismus
- Dome-shaped lesion
- Whitish-pink in color
- Can cause non-rhegmatogenous RD
- Early detection is crucial due to high risk of metastasis
- Enucleation is considered as a last resort

- Tests:
  - B-scan (high reflectivity)
  - CT, MRI (to rule out metastasis to brain)
46-Year-Old Man

- Another long-standing patient, who presents for annual exam and to have his previously diagnosed retinal hole evaluated (date of initial Dx 2001 by yours truly).
- Today’s exam unusual except on casual observation followed by a question or two, patient informs us that he has developed eczema “for the past 6-8 months.”
- All else proceeds normally. We decide to bring the patient back after looking at the nerves and IOPs for a preglaucoma workup simply because it had not been done before.
- Then, we evaluate fundus with 30 diopters for the retinal hole and we see this:

Continued

- I diagnose three pars plana exudative cysts.
- Associated with multiple myelomas in older African-Americans (our patient is Caucasian and only 46). He has symptoms of weight loss (patient has gained weight), bone and joint pain (patient complains of no aches or pains), and some anecdotal evidence of association with eczema.
- Refer to retinal specialist with recommendation for bloodwork to check markers (increased immunoglobulins).

Multiple Myeloma

- Early symptoms
  - Fatigue and weakness
  - Weight loss
  - Bone pain
  - Renal problems (build-up of toxins)
  - Anemia
  - Kidney problems (build-up of toxins)
  - High protein (immunoglobulins)
  - Anemia
  - Weakened bones (lytic lesions)
  - CT scans, MRI, PET

Treatment

- Most effective is Stem Cell Transplantation (SCT)
  - Recipient cells removed, frozen, and stored
  - Suitable donor cells are a possibility also
  - Existing bone marrow destruction with chemotherapy
  - Replacement with SCT through venous injection
  - Not a cure, but prolongs life expectancy
Conclusion

- There is a lot to know. It would be difficult to give the patient a clear-cut answer from one visit, but knowing what the possibilities are makes it easier to discuss the findings. It also makes it easier to discuss the case with the OMD involved.

Whatever happened to our other “routine” exams?
What is SIC Anyway?

- Solitary Idiopathic Choroiditis
- Yellowish-white lesion of unknown etiology
- Occurs at any age (usually 20-50)
- No predilection for gender or age
- VA usually good unless in juxtapapillary or foveal area
- IOP and AC normal
- Patient usually lacks any contributory medical, systemic or ocular history, which makes this a diagnosis of exclusion
- It contains both active and inactive phases

And this one?

- 13-Year-Old African-American Female
- Previous patient of our pediatric clinic between the ages of 5 and 8
- Last exam, however, 5 years ago
- No other exams, no other location according to mother
- No indication of the "lesion" anywhere in previous encounters
References

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