

Systemic Diseases

Presented by: Dr. Ahmed Abu El-asrar

Objectives:

- To know the spectrum of ocular involvement secondary to common systemic diseases such as diabetes mellitus.
- Understand the pathogenesis and importance of screening for diabetic retinopathy.
- To know and differentiate between different causes of uveitis from systemic causes.
- To know the indications for ophthalmic referral in patients with systemic diseases, For the following: Endocrine disorders (Diabetic Retinopathy), Cardiovascular disorders (Hypertensive Retinopathy), Infectious disorders (Tuberculosis), Hereditary and Hematopoietic disorders.

Prof. Abu El-asrar emphasized on the importance of (Kanski's Clinical Ophthalmology) book "just search for a disease mentioned in the lecture & look at its pictures" Don't worry we read it for you and added the important notes (Link)

Color index:

Overview:

- By far the most common disease that can affect the eyes (particularly the retina) and even can cause blindness is diabetes if not diagnosed and treated at the time.
- Now, it is considered to be an irreversible cause of blindness.
- Diabetic retinopathy is the commonest cause of legal blindness in individuals between the ages of 20 and 65 years.
 - These are relatively young people which means that blindness due to DR has a major impact on the country because the cost of taking care of blind people is very huge. So, it has a major socioeconomic problem.
- The risk of blindness is about 25 times greater in diabetics than in non-diabetics.
- The incidence of diabetic retinopathy is related more to the **duration** of diabetes than to any other factor. (The longer the duration the higher the risk of developing diabetic retinopathy)
- Diabetic retinopathy is the most common microvascular complication of diabetes mellitus, it is also a neurodegenerative disease > studies showed that diabetic retinopathy in the very early stages of the disease showed neurodegeneration > there is loss and apoptosis of the retinal neurons very early in the course of disease even before the onset of vascular disease.
- So keep in mind diabetic retinopathy has two pathways:
 - 1) neurodegenerative.
 - 2) microvascular.

Risk Factors:

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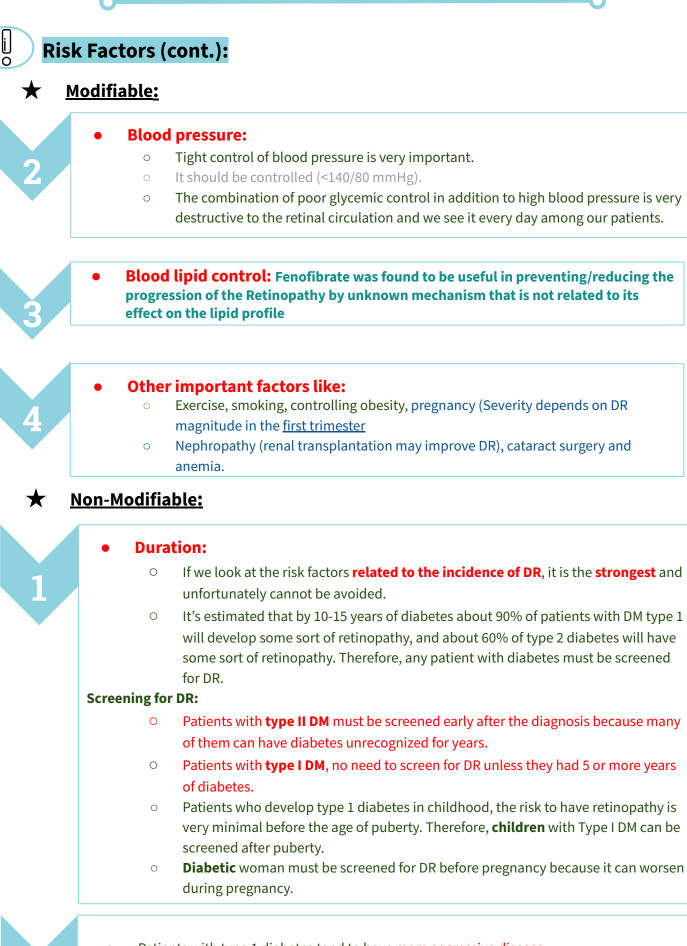
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<u>Modifiable</u>:

Glycemic control:

- **Tight control** of blood sugar especially if started early in the course of diabetes is very beneficial to prevent and stop progression of diabetic retinopathy, not only diabetic retinopathy but also other microvascular diseases such as: nephropathy and neuropathy.
- This evidence came from many studies, one of the oldest is diabetes control and complications study that was multi-centered study, patients were followed for 6 and half years and was mainly focusing on type 1 diabetes. Young diabetics were randomized to receive conventional insulin treatment (1 or 2 injections per day) vs tight control of blood sugar in the form of insulin pump or several insulin injections per day. The group who had tight control they **had mean HA1C 7.2%**. Then at the end of follow up, it was clear and obvious that tight control of blood sugar protected against development and progression of diabetic retinopathy.

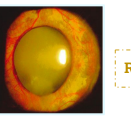
Another big observation that after termination of study, all the patients resumed the previous medication, so those patients who were tightly controlled are no longer tightly controlled. Then it was found that even the 2 groups have equal blood sugar levels, those who had a **tight control early in the course of diabetes were still protected**. So early tight control after the onset of diabetes is very important. This phenomenon is known as "metabolic memory".



- Patients with type 1 diabetes tend to have more aggressive disease
- more aggressive fibrovascular proliferation compared to type 2 → they tend to have aggressive disease and need particular attention to prevent blindness.

Diabetes Mellitus





Rubeosis Iridis

Iris: Rubeosis Iridis. due to the ischemia the retina will secrete many angiogenic factors, most importantly VEGF, which will lead to the formation of new vessels in the iris. Also the new vessels can obstruct the angle of anterior chamber and this will lead to neovascular glaucoma (very aggressive leads to blindness).

Treated by Pan-Retinal Photocoagulation,

Advanced proliferative diabetic retinopathy is the cause of Rubeosis Iridis (SAQ)

Lens: cataract; Diabetics are more prone to have it (glucose affects osmolarity \rightarrow the lens gets opacified).

Anterior Uveitis: inflammation of the iris and/or the ciliary body. Also called Iridocyclitis (inflamed iris + ciliary body) or Iritis (inflamed iris). Iridocyclitis and Iritis are used interchangeably because they are undistinguished clinically

Retinopathy: the most common and major problem is retinopathy.

Optic neuropathy and **3rd**, **4th & 6th nerve palsies**: in poorly controlled diabetes diplopia might happen because 3,4,6 supply the extraocular muscles.

A SPACE FOR YOUR NOTES

Classification:

- Background Diabetic retinopathy (BDR): Characterised by Microaneurysms, dot and blot hemorrhage, and exudates. They constitute the earliest signs of DR. (Especially <u>Microaneurysms</u>, they are the <u>first ever sign</u>)
- **2. Diabetic Maculopathy:** Strictly refers to any retinopathy at the macula
- 3. **PreProliferative Diabetic Retinopathy (PPDR):** Manifests as cotton-wool spots, venous changes, intraretinal microvascular anomalies (IRMA). These signs indicate severe ischemia, and that retinal neovascularization is coming very soon
- **4. PDR:** Characterised by neovascularization (Whether NVD or NVE)

5. Advanced diabetic eye disease:

Characterised by tractional retinal detachment. Vitreous hemorrhage, and neovascular glaucoma (Rubeosis Iridis)

Pathogenesis

- The story behind Diabetic Retinopathy has 2 components:
 - Neuropathy: Very early in the course of diabetes, the retinal neurons are suffering even without vascular retinopathy. So, there is a very early phenomenon of neuropathy that retinal neurons are suffering and many of them die early because of apoptosis as result of hyperglycemic exudative stress "centers of apoptosis are expressed by retinal neurons even in subjects without DR".
 - **Microvascular disease** (what we see clinically) which has 2 major changes:

Progressive vasculopathy



Progressive vasculopathy:

Characterized by **leakage** of blood retinal barrier due to the damage of endothelial cells lining the retinal capillaries by increased adhesion of leukocytes, (vessels of retina are lined by endothelial cells with tight junctions lying on basement membrane and surrounded by pericytes. The tight junctions of endothelial cells are responsible for integrity of blood retinal barrier). Very early in the course of diabetes, you have disruption of tight junction proteins such as occludin and cadherin, and thickness of BM and loss of supporting pericytes, this means that blood vessels become **leaky**. So, they leak fluid and lipoprotein and this will cause edema, and **edema** of macula is an important cause of moderate visual loss in diabetic retinopathy. Exudates seen in ophthalmoscope are basically a chronic oedema

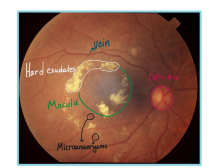
- This will lead to retinal hypoxia and ischemia. Then retinal ischemia will activate transcriptional factors such as HIF-1-alpha "hypoxia- inducible- factor 1 alpha".
- This HIF-1-alpha will upregulate many angiogenic factors to be secreted by the retina and the most famous is vascular endothelial growth factor "VEGF".

Progressive microvascular occlusion:

- VEGF is known to be hyperpermeability factor. It will cause breakdown of blood retinal barrier. In addition, it's an angiogenic factor (it induces proliferation, migration and tube formation of endothelial cells) which ends with formation of new vessels (proliferative disease).
- These new vessels are not healthy and always accompanied by fibrous tissue. At the end, this fibrous tissue will cause traction of retina → tractional retinal detachment and the new vessels can bleed → the patient comes to ER with dramatic visual loss "suddenly he/she woke up and he/she cannot see", the first change that cause dramatic visual loss is vitreous hemorrhage.

Non Proliferative Diabetic Retinopathy (NPDR)

- Retinopathy is a disease of small blood vessels in the retina.
- This is the right eye of a patient with diabetes.
- If you look at the retinal vessels, the veins are always darker than arteries and broader
- You can see in the **macula**, the collection of **hard exudates** (<u>composed</u> of lipoprotein and lipid laden macrophages).
- These hard exudates are <u>the result of leakage</u>, it's a sign of macular edema (commonly causes deterioration of vision in diabetic patients [moderate visual loss] because it affects the macula).
- You can see small red dots, theses dots are **microaneurysms** and these are the **source of leakage**
- <u>Management</u> here is to occlude the aneurysms with <u>focal</u> laser photocoagulation
 - **what are the targets of focal laser?** red dots. Focal laser has been shown to be associated with better outcomes compared to no treatment.
- Nowadays we have anti-VEGF agents that can be injected to the eyes, so if the edema is involving the center of macula like in this patient, we can **enhance the effect of laser by giving injection of anti-VEGF agents.**
 - bevacizumab (Avastin), aflibercept and ranibizumab.
- In summary Breakdown in capillaries → leak → edema → chronic accumulation of fluid and lipoprotein → hard exudate!



Diabetic Retinopathy DR

Non Proliferative Diabetic Retinopathy (NPDR)

- This is another example of macular edema, you can see the **hard exudates** (hard exudates are the result of leakage "breakdown of blood retinal barrier"). and you can see the red spots that we need to treat with focal laser photocoagulation.
- To control the edema we need to close the aneurysm.

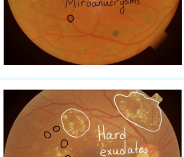
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- Another example of more extensive **hard exudates**, and you can see the red dots "we should close by focal laser photocoagulation". and we can also combine it with anti-VEGF
 - bevacizumab (Avastin), aflibercept and ranibizumab. 0
- This cartoon shows you how we do focal photocoagulation:
 - Hard exudates are the rings. 0
 - Microaneurysms are the small dots inside the rings. 0
- This is a patient who had an enlarged ring of hard exudates and in the center are laser scars, usually it takes up to 6 months for hard exudates to be absorbed.
- Then after 6 months, as you can see in the second picture, there is a **complete resolution** of hard exudates
 - (we occlude the aneurysms not ablate them!). 0
- There are signs in the retina that can tell me that the retina is very ischemic before the development of new vessels
- These signs are classified as severe non-proliferative retinopathy, which means that within 1 year, the patient will develop proliferative diabetic retinopathy and he has a high risk of blindness
 - Treat with pan retinal photocoagulation.
 - One of these signs is venous looping.

(more details about this in the next page)

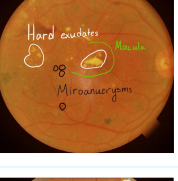




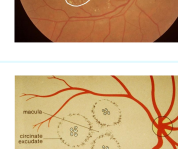




Hard exudate







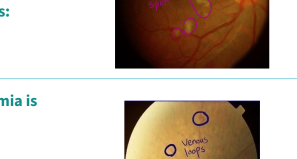
Microanuery

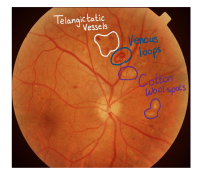
Non Proliferative Diabetic Retinopathy (NPDR)

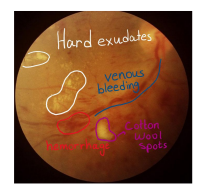
- Most of the previous pictures showed the leakage component of retinopathy.
- The other component as we said is progressive occlusion of retinal arterioles and this will cause ischemia.
- One of the <u>early</u> signs of infarction of retina [retinal ischemia] is: cotton wool spots (due to occlusion of retinal arterioles), which are different from hard exudates (sign of leakage)!
- The most important and dangerous sign of <u>severe</u> retinal ischemia is venous changes
- What are these changes? venous loops and Beading .
- **Venous Beading:** look at the course of this vein, there are dilated areas and constricted areas
- So **venous looping** and **beading** are the most reliable and important signs of **severe retinal ischemia.**
- Venous loops. They call it sometimes omega sign.
- You can see the **cotton wool spots(infarction(** and **intra-retinal hemorrhage.**
- We have <u>another sign of retinal ischemia</u> called **Intra-retinal microvascular abnormalities (or IrMAs);**

these are dilated **telangiectatic vessels** within the retina, the origin is not well known it can be collaterals, it can be new vessels still within retina. You can see here the dilated telangiectatic vessels.

- We tend to treat with **pan retinal photocoagulation in this stage.**
- You can see **venous beading*** (dilated and constricted parts), **cotton wool spots, hard exudates** (lipoprotein)
- The patient has signs of ischemia and leakage and lots of intraretinal hemorrhages
- Presence of intraretinal hemorrhages in 4 quadrants is a sign of severe non-proliferative retinopathy.
- **★** TO SUMMARIZE: the signs of <u>severe</u> non- proliferative retinopathy are:
- 1-Retinal infarction (cotton wool spots)
- 2-Venous looping and beading (most reliable sign)
- 3-Intra-retinal microvascular abnormalities (telangiectatic vessels)
- 4- Intra-retinal hemorrhage in 4 quadrants







Pathology	Hard Exudates	Cotton wool
Cause	Oedema and Leakage	Neuronal debris (due to ischemia)
Character	Waxy yellow, distinct margins, and <u>might</u> <u>involve the fovea</u>	histologically seen as cystoid bodies. Fluffy whitish, <u>seen only post equatorial retina,</u> obscure the underlying blood vessels

Diabetic Retinopathy DR

If the retina is ischemic (severe non-proliferative) and the patient was

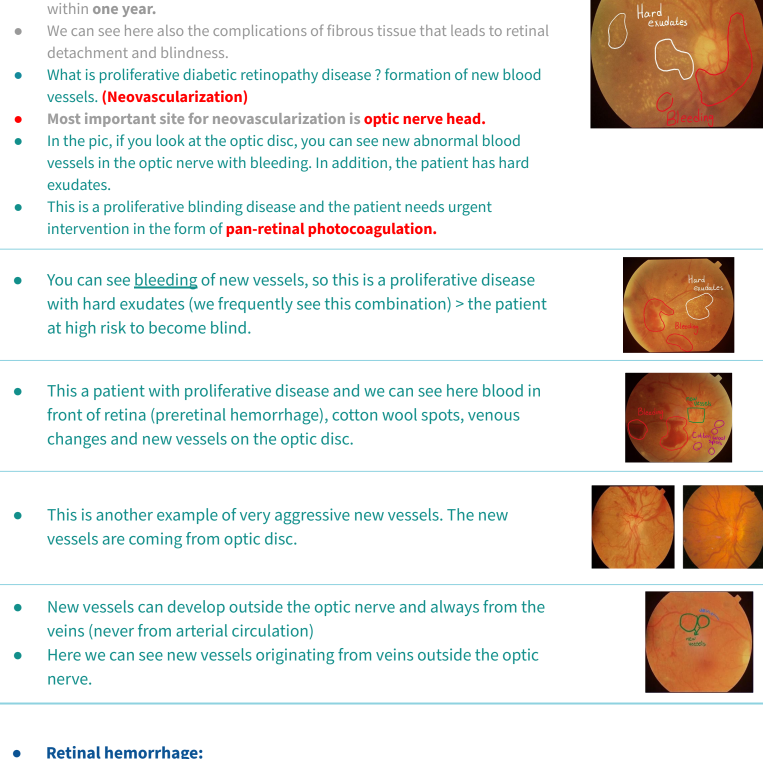
ischemia \rightarrow VEGF release \rightarrow **neovascularization** \rightarrow bleeding.

This is called **PROLIFERATIVE diabetic retinopathy**

capillaries are occluded):

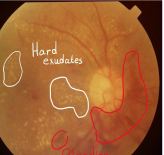
neglected this leads to \rightarrow Progressive microvascular occlusive disease (the

45% of patients with severe non-PDR will progress into proliferative disease



Retinal nerve fiber layer hemorrhage arises from superficial precapillary arterioles, wherase Intraretinal hemorrhage arises from the venous system

Proliferative Diabetic Retinopathy (PDR)

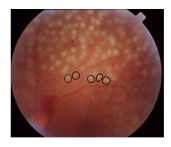


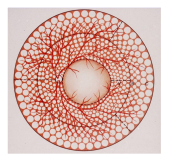


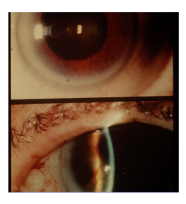
Proliferative Diabetic Retinopathy (PDR)

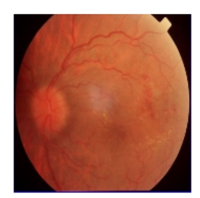
- Proliferative disease.
- Patient now has bleeding 'subhyaloid hemorrhage'.
- You can see the neovascularization, hard exudates and cotton wool spots.
- **Picture above:** now the treatment of proliferative DR or severe non-PDR is by laser (pan-retinal photocoagulation).
- Why is it call like that? because you apply scattered laser burns throughout the retina sparing the optic nerve and macula and this automatically will be followed by regression of new vessels. Mechanism: destroys the ischemic retina that releases the angiogenic material.
- Another modality for treatment is to inject antibodies into the eye to block vascular endothelial growth factor (VEGF) → helps control edema.
- **Picture below:** example of pan-retinal photocoagulation. These are laser burns.
- If there is extensive exudates and hemorrhages, PRP "panretinal photocoagulation" is done (the whole periphery gets cauterized except the posterior pole). results in loss of rods → loss of vision at night.
- Another big complication of retinal ischemia is formation of new vessels on the iris and the angle of anterior chamber, this is called rubeosis iridis (neovascular glaucoma).
- As a result of ischemia, the new vessels will not only develop on the retina, they also develop on the iris and it involves the angle "will close the angle by fibrous tissue", this will cause very aggressive type of glaucoma called neovascular glaucoma and it's a very serious complication and blinding disease.
- The angiogenic factors like VEGF will move into the anterior chamber and cause neovascularization of iris and the angle. This happens with any retinal ischemia like central retinal vein occlusion, but mainly with DR.
- Patients with DR can have vision threatening diseases such as macular edema and proliferative diseases but they remain asymptomatic as long as there's no hemorrhage. and once they develop hemorrhage it's too late! so how can we prevent blindness?
- Regular screening of a patient with diabetes to examine the retina is very important before it becomes symptomatic and if we find it we treat it with laser coagulation, so the ONLY way to prevent blindness is to have national screening programs for diabetic retinopathy











• Most common cause of both bilateral and unilateral proptosis in an adult.





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Pathogenesis:

- An autoimmune disease characterized by serum IgG antibodies bind to TSH receptors in the thyroid and causes overstimulation and high thyroid hormone production.
- The autoimmune antibodies infiltrate the eye causing inflammation of extraocular muscles and associated with increased secretion of glycosaminoglycans and osmotic imbibition of water.

Risk Factors:



• Smoking (most important)

• Family history

• Pretibial myxedema.

Systemic manifestation:

- Heat intolerance.
- Weight loss.



- Thyroid function test: high T3, T4 and low TSH.
- Visual evoked potential: to exclude optic neuropathy.
- CT scan if suspecting thyroid eye disease



Treatment:

- Anti-thyroid medications or thyroid ablation with radioactive iodine (for disease itself).
- Steroid, lubricants and eye protection before sleep (for eye symptoms).
- If there is restrictive myopathy surgical intervention is required.

Ocular Manifestations

- Eyelid retraction and Lid lag
- Infiltrative ophthalmopathy
- Dysthyroid optic neuropathy
- Proptosis and exophthalmos
- Chemosis, exposure keratopathy, ophthalmoplegia
- Restrictive thyroid myopathy
 - Most commonly involve the inferior rectus, <u>causing elevation defect</u> (due to fibrotic contractures of the inferior rectus).
 - Second most common is <u>abduction defec</u>t due to the involvement of medial rectus.
 - Third is superior rectus (depression defect)
 - Fourth is lateral rectus (adduction defect)
 - The muscle movement is not much affected. However, the opposite muscle movement is affected due to the fibrotic contractures of the antagonist muscle
 - Thyroidectomy doesn't help. Steroids must be administered



Hertel ophthalmometer To measure the degree of proptosis or exophthalmos.



Common here. It is due to extrapulmonary TB. If diagnosed and treated early, you will save the patient

- TB is a chronic granulomatous infection usually caused in humans by mycobacterium tuberculosis.
- TB is primarily a pulmonary disease but may spread by the bloodstream to other sites; ocular involvement (TB can involve any part of the eye) commonly occurs without clinically overt systemic disease.
- Extrapulmonary TB when you have an eye infection without pulmonary infection in 60% of cases.
- Tubercles uveitis is an important cause of blindness.
- TB may be indolent and the first manifestation in the eye.
- It can be: 1. direct infection 2. immune response to tubercular protein.
- TB is the second most common cause of uveitis in KSA, after Vogt- koyanagi-Harada disease, and the third cause is Behçet disease.
- Granulomatous inflammation that is the disposition of mutton-fat keratic precipitate, iris nodules, infiltration of the choroids, and retinal vasculitis; These are the most important manifestation of TB in the eye.
- Mutton-fat keratic precipitation: collection of inflammatory cells on the corneal endothelium that appear large with yellowish color (can be seen as white dot inferiorly, mostly due to staph but could be caused by TB).



How to diagnose ocular tuberculosis?

- Clinical findings consistent with tuberculosis.
- Rule out other specific uveitic entities (e.g. Behcet disease, sarcoidosis, etc).
- Investigations (tuberculin test).
- If the patient is treated early for at least 9 months with anti-TB drugs blindness can be prevented.

Investigations:

- First you should take a good history (family history or history of exposure will increase the chance that the eye inflammation is caused by TB).
- CXR to roll out that the patient has previous infection in the chest.
- We rely more on tuberculin skin test, if it was strongly positive, 15 mm or more induration, this will support the diagnosis.
- PCR and the interferon-gamma release assay (IGRA).
- Aqueous or vitreous sampling rarely yields demonstrable (smear – acid-fast bacilli in Ziehl–Neelsen staining – or culture – Lowenstein–Jensen medium).



Treatment:

- Prolonged Anti-TB therapy (multi-drug therapy):
 - 4 drugs in 2 months, then continuefor 6 months with 2 drugs.
 - Isoniazid with Vitamin B6 (pyridoxine) to prevent the development of peripheral neuropathy, rifampicin, pyrazinamide and ethambutol.
 - Ethambutol can cause optic neuropathy.
- Topical and systemic steroids may be used concomitantly to reduce inflammation-induced damage.

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💽 Ocular manifestations:

Uveitis

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Phlyctenular keratoconjunctivitis

A hypersensitivity reaction of the cornea and conjunctiva to bacterial antigens, is characterized by discrete nodular areas of corneal or conjunctival inflammation.

it is divided into granulomatous & non-

are big "mutton-fat" it indicates a

like iris \rightarrow called anterior uveitis.

If it's involving the posterior part like

choroid \rightarrow called posterior uveitis. If it's involving the whole uvea \rightarrow

granulomatous uveitis.

pan-uveitis.

granulomatous, if the keratic precipitates

If uveitis is only involving the anterior part

TB is an important cause of uveitis, uveitis

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Interstitial keratitis and Vitritis



Retinal vasculitis 'Eales disease'

If a patient is presented with intraocular inflammation and signs suggestive of tuberculosis you must treat the patient because it's a blinding disease.

-Tb Induced vasculitis is preferentially Venous



Granuloma of the choroid (Tubercle)

If large enough→ tuberculoma

in TB can mimic anything.



- This is a 16-years-old patient. Notice the pinkish nodules in the angle, she had granulomas 'Phlyctenular keratoconjunctivitis'.
- When you see such granulomas, you have to think about 2 conditions: either TB or sarcoidosis.
- This patient had many members of her family with TB and she was managed with systemic anti-tuberculous treatment with improvement.
- The patient came to the ER. You can see white veins (due to . inflammatory exudate around the blood vessels) + hemorrhages "tuberculous retinal vasculitis".
- It responds to anti-TB treatment, if you don't give anti-tuberculous treatment it will end by losing the eye.



Granuloma of the anterior chamber





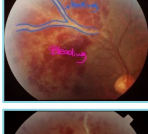
Ocular manifestations (cont.):

- Another common manifestation of ocular TB is retinal vasculitis (inflammation of blood vessels of the retina).
- If you look here to the retinal vessels, this is a vein, but you can see it turned white with a lot of intra-retinal hemorrhages meaning that this vein is inflamed and leaking blood. This white appearing-vein is called *cheesing* and it's a sign of retinal vasculitis(it's more severe than retinal vasculitis found in patients with sarcoidosis), it's due to peri-venous accumulation of **inflammatory exudates** and cells.
- TB is a common cause of occlusive retinal vasculitis. We have the advantage in the retina that we don't need biopsy to diagnose vasculitis, we see it clinically.

Sarcoidosis

- Sarcoidosis is an important cause of uveitis. It's not common here but in a country like Japan, sarcoidosis is the most common cause of uveitis.
- Sarcoidosis causes non-caseating granulomas when compared to TB that causes caseating granulomas.
- Retinal vasculitis can also be seen in sarcoidosis.
- When we suspect sarcoidosis as a cause of uveitis, we always ask for CT of the chest.
- What do you expect to see in CT? Hilar lymphadenopathy and also granulomatous infiltration of the lungs.

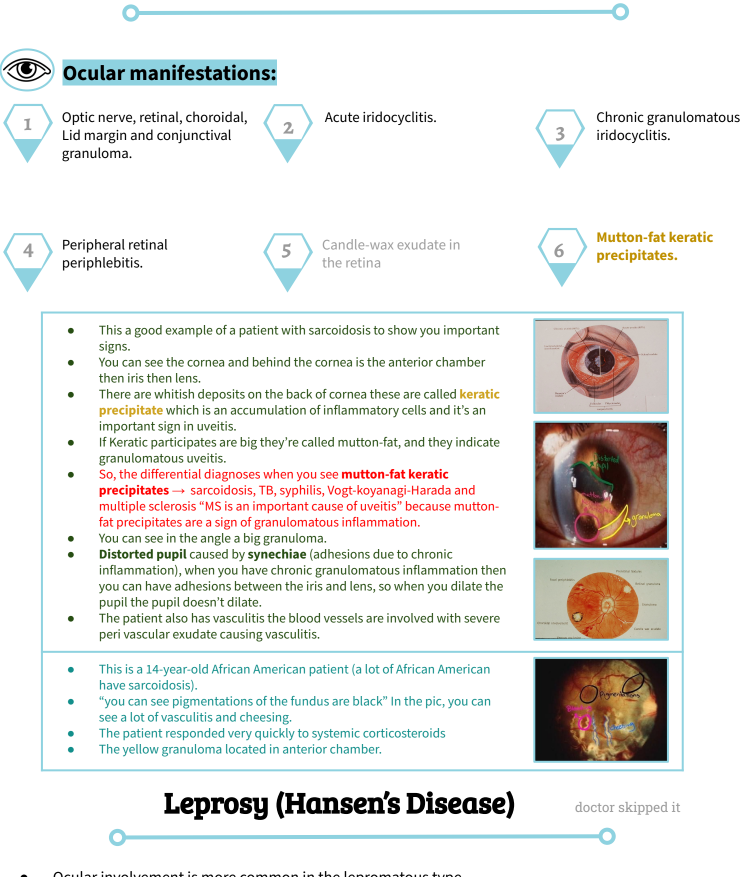
Systemic manifestations Treatment Tuberculin skin test: negative in The triad: Immunosuppressive with -erythema nodosum sarcoidosis. systemic steroids and **NSAIDs. Chest X-ray:** showing bilateral hilar -bilateral hilar lymphadenopathy (BHL) lymphadenopathy (DIAGNOSTIC). -polyarthralgia. **Patients respond to Elevated serum ACE levels and/or** steroids + anti elevated serum lysozyme. Could be the same as TB modulatory agents such manifestation. as mycophenolate Abnormal liver enzyme tests. mofetil or CellCept which is an antimetabolite **Biopsy** should be taken to confirm drug. the diagnosis, if we were in doubt. (Transbronchial biopsy)



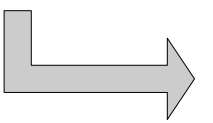


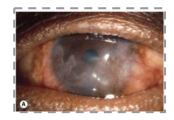
Inflamed vessels (retinal vasculitis). due to peri-vascular inflammatory infiltrates

Sarcoidosis



- Ocular involvement is more common in the lepromatous type.
- Signs: Facial nerve affection, Loss of the lateral portions of the eyebrows and eyelashes (madarosis), Interstitial keratitis with beading of nerves, iritis.





Syphilis

- When you see a patient with uveitis you always have to rule out syphilis
- It develops at the secondary stage of syphilis (this is international recommendation) by doing serological testing of syphilis "VDRL, fluorescent treponemal antibody absorption (FTA-ABS)".
- Despite this, we diagnose syphilis very rarely here (now is getting common), but when we look to western countries like UK there are a lot of cases of syphilis "syphilitic uveitis" which means that until now we are protected against this bad disease.
- Nowadays, we see syphilitic uveitis more with HIV+ patients
- London is a city full of syphilis, the patient comes with syphilitic uveitis → receives treatment → cured then he will come again with another attack of syphilitic uveitis due to another exposure.

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• Signs: Uveitis with/out granuloma, retina with **ground glass appearance**

Congenital

- Interstitial keratitis.
- Chorioretinitis.
- Bilateral
- Transplacental infection.

- Acquired
- Ocular chancre.
- Iridocyclitis.
- Interstitial keratitis.
- Chorioretinitis.
- Neuro-ophthalmic.
- Unilateral

Rubella

- Cataract.
- Microphthalmos (small eyes).
- Retinopathy (pigmentary retinopathy: salt and pepper).
- Glaucoma.
- Anterior uveitis: unresponsive to steroids.
- They use VERY big glasses & hearing aid also.
- If the mother is infected with rubella virus, the baby can be born with congenital rubella syndrome.
- Systemically, they have congenital heart disease and deafness.

Wilson's Disease

(Hepatolenticular degeneration)

• Systemic manifestations: liver disease, basal ganglia dysfunction, psychiatric disturbances.

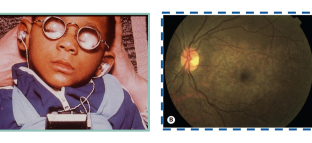
Ocular manifestations:

- **Kayser-Fleischer ring** (Copper deposit in peripheral part of the cornea) consists of a brownish-yellow zone of fine copper dusting in peripheral descemet membrane detected with gonioscopy (Important sign).
- Green sunflower cataract (copper deposited in the lens).
- There is excessive copper deposition in the tissues due to deficiency of the carrier protein which is called alpha 2 globulin "ceruloplasmin".
- So, in the eye, the copper can be deposited at the peripheral part of Descemet's membrane and this will cause Kayser-Fleischer ring (the presence of this ring is diagnostic for Wilson's disease).
- Treatment: Penicillamine.

by can be born with congenital rubella syndrome.







Marfan's Syndrome

- It is an autosomal dominant disease.
- Systemic manifestation: arachnodactyly
- (Long fingers), heart diseases, bone deformities.

Ocular manifestations:

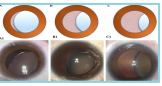
- Lens subluxation bilaterally, which occurs due to weak zonules. It is the most common and severe ocular feature in Marfan's syndrome.
- Subluxation is usually superotemporal.
- Angle anomaly.
- Glaucoma.
- Hypoplasia of the dilator M.
- Axial myopia.
- Retinal detachment.

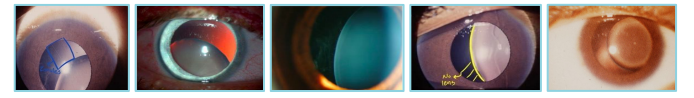




This is the systemic manifestation with arachnodactyly.

e systemic ation with odactyly.





What do you see behind the pupil? subluxated lens, you can see the equator of the lens visible through pupil.

Systemic Lupus Erythematosus

- If you see a patient with these facial features, what's your diagnosis? SLE.
- SLE is an autoimmune disease associated with multiple autoantibodies, **What are they?** Antinuclear Antibodies, and Anti-ds DNA Antibodies. The patient has high ESR, low C3 and C4.
- It can affect the eye, particularly retina, but only If the disease is active
- It causes multiple retinal infarcts due to occlusion of retinal arterioles. The retinal effects would **increase** if the patient was also positive for antiphospholipid antibodies, **What are they?** Lupus Anticoagulants, Anticardiolipin antibodies, and Anti-B2-glycoprotein Antibodies
- What would you see in the retina? the disease mainly manifests as retinopathy (cotton wool spots) meaning it causes occlusion of retinal arterioles.
- So, the main pathology is micro-thrombosis of retinal arterioles causing multiple retinal infarctions (cotton wool spots), with or without hemorrhage.
- Scenario: young lady with bilateral multiple cotton wool spots (SLE).







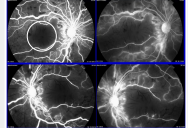
Good example of Cotton Wool spots. This is a mild disease Yellow: Cotton Wool Spots

More advance disease Blue: Occlusion of this arteriole causing infarction Yellow: Cotton Wool Spots Green: Hemorrhage

Systemic Lupus Erythematosus

- This is a typical example of a patient who unfortunately lost her vision (20/200). SLE is blinding if not treated properly.
- **The white areas are multiple areas of infarction.** The treating physician thought they were retinitis.
- The moment we saw the patient in the ER, we made a diagnosis of possible SLE. The patient was admitted, and positive for Antinuclear Antibodies, and Anti ds-DNA antibodies.
- She was referred to Rheumatology for treatment.
- Same patients under Fluorescein Angiography (imaging of the retina with white dye).
- Black lines (circled in yellow) are occluded retinal vessels with extensive ischemia and infarction. The macula is severely ischemic.
- Active SLE uncontrolled medically can cause this presentation.
- Patient's vision will remain poor after treatment (irreversible damage) because all the macula's circulation is occluded. Some drugs, however, can damage the retinal pigment epithelium, e.g. Chloroquine, and therefore, any patient who is taking such medications should be examined regularly.

(Infarctions all over the retina involving the macula; which leads to central vision loss)



(Extensive occlusion of retinal arterioles involving the macula)

Rheumatoid Arthritis

- A seropositive disease.
- Which factor do you need in order to diagnose RA? Rheumatoid factor.



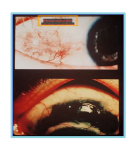
Ocular manifestations:

- K.C.S (Keratoconjunctivitis Sicca; dryness of the eye):
 - Autoimmune disorder attacking the lacrimal gland.
 - Positive Rose Bengal staining \rightarrow K.C.S.
- Scleritis: (patient complains of severe eye dryness)
 - An important cause of sclerites and melting of the sclera if not controlled (Scleromalacia Perforans; severe form of Scleritis and the Blinding complication of this disease).
- Keratitis.



RA deformity. Radial deviation, swollen fingers and elbow nodules.





Melting of the sclera with Scleromalacia Perforans exposing the underlying different tissues. **Ankylosing Spondylitis**

- Seronegative.
- X-rays of sacroiliac joints shows juxta-articular osteoporosis in the early stages.
- Acute recurrent non-granulomatous iridocyclitis.
- This is an interesting disease for us. Whenever we see a young male patient in the ER having a unilateral acute non-granulomatous anterior uveitis, ankylosing spondylitis is the most important differential diagnosis, and we must rule it out.
- What kind of test should we do? HLA-B27 typing is the most specific test.
- This type of uveitis (recurrent, acute, non-granulomatous) is the type that happens in patients who are positive for HLA-B27.
- It can be positive in systemic diseases like:
 - Ο **Ankylosing Spondylitis**
 - 0 **Psoriasis**
 - 0 **Reiter's disease**
 - Ο IBD
- This type of uveitis can also occur without systemic disease.
- What do you mean non-granulomatous? You don't see mutton-fat precipitates (unlike T.B and Sarcoidosis) which is sign of granuloma.
- Whenever we see a young male patient with acute recurrent non-granulomatous iridocyclitis we have to rule out Ankylosing by HLA-B27 typing.
- Refer the patient to Rheumatology because at this stage, if the patient has Ankylosing Spondylitis, you have to start systemic treatment early to prevent spinal deformity. Frequently, eye manifestations as uveitis and acute recurrent non granulomatous uveitis are the first manifestations of Ankylosing Spondylitis.
- Systemic manifestation: pain and stiffness in the lower back with limitation of movement, calcification of spinal ligaments gives rise to a 'bamboo spine'.
- Ocular manifestation: acute recurrent non-granulomatous anterior uveitis.
- Complications: synechiae.

Sjogren's Syndrome

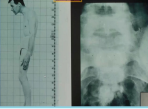
doctor skipped it

- Autoimmune disease.
- Involvement of salivary glands (leading to dry mouth), bronchial epithelium & vagina.
- Ocular features: K.C.S. keratoconjunctivitis sicca "dryness of the eye".
- **Systemic manifestations:** dryness of skin and mouth, arthralgia and polyneuropathy.
- **Investigations:**
 - Schirmer tear test.
 - Positive Rose Bengal staining (for keratoconjunctivitis sicca).
 - ANA, RF positive. > Associated with HLA-B8/DR3.

Reiter's Syndrome (Reactive Arthritis)

doctor skipped it

- A triad of: urethritis, conjunctivitis, and seronegative arthritis.
- Ocular features: conjunctivitis, keratitis, and iridocyclitis





Juvenile Chronic (idiopathic) Rheumatoid Arthritis

🚽 🛨 The patterns of disease (important):

• **Systemic onset "Still's disease":** uveitis is extremely rare.

The child would present with fever, maculopapular rash, lymphadenopathy, hepatosplenomegaly and pericarditis. With this presentation, uveitis is very rare.

- **Polyarticular onset:** uveitis is fairly rare (5 or more joint are affected).
- Pauciarticular onset: 20% develop uveitis at onset (common), and ≤4 joints are affected.
 In western countries, this is the most common cause of uveitis in children.
 This is a disease of children. The eye can be affected by blinding inflammation.
- The problem here is that children cannot complain, and the eye looks quiet, but the chronic inflammation can destroy the eye.
- Such condition can lead to blindness.
- That's why we have to know **The risk factors for a child with juvenile chronic arthritis to develop uveitis (refer to ophthalmologist):**
 - Female.
 - Pauciarticular onset.
 - Arthritis developed <4 years old.
 - Positive Antinuclear Antibodies.
- Complications are common, mainly glaucoma and cataract.
- **Ocular manifestations:** chronic non-granulomatous uveitis, band keratopathy, and posterior synechiae.
- Investigations:
 - Anti-ANA antibodies will be +ve in majority of pauciarticular type.
 - Rheumatoid Factor is +ve in some polyarticular type.
 - HLA-B27 will be +ve in some patients.
- **Treatment:** usually they need biologic therapy like infliximab. Topical and systemic steroid with a Mydriatic Agent to prevent posterior synechiae.



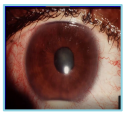
Child with multiple joints affected; keep this case in mind.

- Blinding disease due to recurrent uveitis affecting the retina.
- Very common disease. 3rd most common causes of uveitis in our country.
 The country having the highest incidence of the disease is in Turkey. The disease is highly prevalent along the Silk Road (East Asia and Southeast Asia with South Asia, Persia, the Arabian Peninsula, East Africa and Southern Europe). Very common around Mediterranean Basin, China, Japan, Korea, and Turkey. You don't see it in Caucasians.
- It is a disease of a multisystem vasculitis.
- The major cause of visual loss in patients with Behçet's disease is recurrent episodes of vaso-occlusive retinal vasculitis.
- Patients with Behçet's disease have a very important involvement of polymorphonuclear leukocytes in the pathogenesis of the disease.
- We see many patients who present with ulcers but after having recurrent episodes of DVT.
- There is no specific lab study to diagnose Behçet's disease, the diagnosis is a clinical one.
- The criteria required:
 - **Recurrent painful oral ulcer** (mouth ulcers should be in all patients because if you look at epidemiological studies, mouth ulcer was the most common manifestation of the disease in about 97% of the patients).
 - In addition to the mouth ulcer, you need 2 of the followings:
 - Skin lesions.
 - Recurrent genital ulcers.
 - **Eye manifestation (**Anterior nongranulomatous **uveitis).**





- 1st line is Biological therapy; anti TNFs. (This disease do not respond to steroids)
- Most effective drug is **Infliximab** (and Adalimumab) biological agents targeting TNF-Alpha.



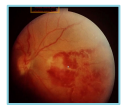
Hypopyon is a whitish material (puss) involving inflammatory cells in the anterior chamber of the eye. Accumulation of Neutrophils.



Aphthous ulcer



It causes retinitis and it's a blinding explosive disease. In the past, if you look into the literature, they were telling whatever you do, the patient becomes blind.



Occlusive retinal vasculitis + hemorrhage

- Toxoplasmosis cause frequent retinitis and can lead to blindness if the central of macula involved.
- Caused by Toxoplasma gondii after eating raw meat, are obligatory intracellular protozoan parasite, can be:
 - **Congenital** (if the mother is infected for the first time during pregnancy):
 - Convulsions
 - Chorioretinitis
 Chorioretinitis
 Chorioretinitis
 Chorioretinitis
 Chorioretinitis
 Chorioretinitis

vision-threatening complications.

- Intracranial calcification.
- Acquired:
 - Reactivation of old lesion, manifest manly as retinitis.
- Pharmacological therapy (equally effective) to treat toxo-retinitis if it needs to be treated are:
 - Clindamycin
 - Sulphonamides
 - Pyrimethamine (daraprim)
 - Steroids
 - Sulphadiazine, Cotrimoxazole, Erythromycin, Minocycline, Azithromycin.
- The **most common** cause of <u>focal retinitis</u> worldwide
- The fourth most common cause of uveitis in the country. It is an infectious cause of uveitis.
- The severity of infection of a baby depends on the timing of infection by mother.
 - So, if it happens in the **first trimester** what will happen to the baby? **Abortion.**
 - If it happens in the **third trimester**, the baby will end up with **congenital toxoplasmosis**.
- If the mother is infected for the first time in her life, the baby will become infected (no antibodies to protect the baby), but if she is infected as a recurrent infection, the baby is protected.
 - That's why at the start of pregnancy they always order antibodies screening for toxoplasma.
 - If the mother has IgG positive antibodies meaning that she was exposed before, so there is no fear.
 - But if the mother was seronegative at the beginning of pregnancy and then during pregnancy became positive then the risk is very high.
- The **acquired toxoplasmosis** affects retina causing toxo-retinitis (focal necrotizing retinitis which is usually located to an adjacent scar, and this is the classical appearance of the disease).



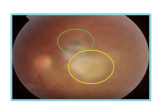
Cerebral calcification with a child with toxoplasmosis



classic example Green: Old scar. Yellow:Active retinitis. an old scar meaning that the patient had previous infection of the retina. Nearby the old scar is an active retinitis (this is a recurrent disease).



Aggressive retinitis



Green: Old scar. Yellow: Active retinitis. Loss of vision is because the center of the macula in involved and losing its function

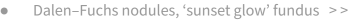
Vogt-Koyanagi-Harada Syndrome

- VKH is an idiopathic multisystem T cells mediated autoimmune disease featuring inflammation of **melanocyte-containing tissues** such as the uvea, ear and meninges.
- Clinical Manifestations :
 - Cutaneous signs (alopecia, vitiligo, poliosis)
 - Poliosis: absent or decreased melanin in head hair, eyebrows or eyelashes.
 - Neurological signs (vertigo , tinnitus, deafness, headache)
 - Anterior uveitis, Posterior uveitis can happen (granuloma).
- You should know about this disease because the disease is **very common** and it is a **multisystem disease**.
- This is the **most common cause** of autoimmune uveitis in **Saudi Arabia**. More common in **young women.**
- It tends to affect pigmented individuals (you will not see it in Caucasians).
- The disease is blinding, but if we **treat the patient early** in the course of the disease by a large dose of systemic corticosteroid combined with immunomodulatory agent such as mycophenolate mofetil (an anti-metabolite like azathioprine and methotrexate but much safer) then we can prevent all of these complications.
- A big problem in the country that not many ophthalmologists know how to diagnose it early so when the patient comes with headache and inflammation of the optic nerve (optic nerve disc swelling), they make wrong diagnosis of pseudotumor cerebri and they refer patient to neurologist (a lot of investigations done to the patient: MRI, lumbar puncture) then the patient will become blind. So we have to have high index of suspicion for the diagnosis of Vogt-Koyanagi-Harada disease.
- VKH is sometimes subdivided into
 - A. Vogt–Koyanagi disease, characterized mainly by skin changes and anterior uveitis
 - B. Harada disease, in which neurological features and exudative retinal detachments predominate.
- Phases:
 - \circ Acute uveitic stage: posterior uveitis with multiple serous retinal detachments \rightarrow panuveitis
 - Convalescent stage: sunset-glow fundus (orange-red discoloration of the choroid due to inflammatory depigmentation, poliosis and vitiligo
 - Recurrent stage: panuveitis with acute exacerbations of anterior uveitis and iris nodules



Ocular manifestations:

- Bilateral granulomatous anterior uveitis.
- Bilateral multifocal posterior uveitis.



Mutton-fat keratic precipitates, chronic manifestation, Acute phase manifested as inflammation of the choroid with exudative retinal detachment (accumulation of fluids under the retina).

Investigations:

- Associated with HLA-DR1 and HLA-DR4
- Lumbar puncture if diagnosis uncertain; CSF shows a transient lymphocytic pleocytosis, and melanin-containing macrophages.



- High-dose steroid &
 immunomodulatory (mycophenolate mofetil)
- infliximab In case of steroid resistance.
 Always high dose.

Vogt-Koyanagi-Harada Syndrome



Complications if untreated:

- Poliosis.
- Vitiligo.
- Posterior synechiae.
- Choroidal neovascularization.
- Glaucoma, Cataract.
- Subretinal fibrosis, Retinal atrophy.
- Sunset glow fundus, or orange fundus.







(Adhesion between posterior iris and the Anterior lens surface

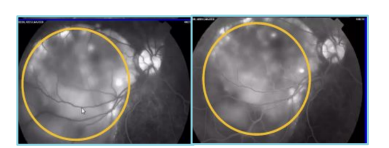
or iris face



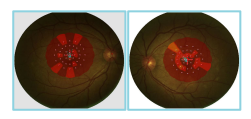
Sunset glow fundus Vitiligo

In Acute uveitic phase:

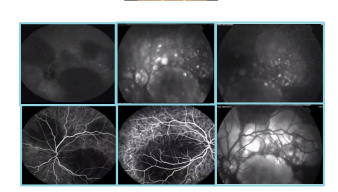
- Uveitis typically will cause granulomatous inflammation of choroids and this will cause secondary exudative retinal detachment.
- Notice here there is a lot of **fluid under the retina (Yellow).** The fluid does not contribute to the color (dark or light), rather, fluid causes change of position, i.e. elevation of the retina.
- Treat early aggressively to prevent blindness
- **Green:** Normal retina.



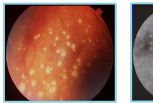
- Retinal angiography.
- Yellow: the dye accumulated "pooling" under the detached retina, exactly at the area filled with fluid (choroid).

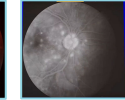


This is a test we use to assess retinal sensitivity called microperimetry.



- Also here, multiple areas of exudative retinal detachment.
- Typical for Vogt-Koyanagi-harada syndrome.
- This patient needs urgent treatment, in the form of large dose systemic corticosteroid, and other immunomodulatory agents like mycophenolate.



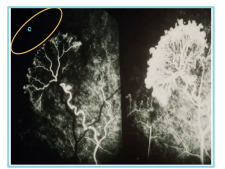




- Conjunctival, comma-shaped capillaries.
- Retinal changes & Major complications :
 - \circ Arterial occlusions \rightarrow ischemia of peripheral retina \rightarrow neovascularization
 - Peripheral retinal ischemia.
 - Neovascular patterns.
 - Capillary closure.
 - Vitreous hemorrhage.
 - Traction retinal detachment.
 - Salmon patches + black sunbursts
- What you should know about SCD here is: As result of sickle cell there is occlusion in the periphery of the retina (causing peripheral retinal ischemia), and can be complicated by neovascularization of the retina then lead to bleeding in the eye that looks like "sea fans".
- How to prevent bleeding? By applying laser to ischemic retina. (Same as DR management)
- In fluorescein angiography there is no circulation at all because it is completely occluded.
- SCD retinopathy is differentiated from diabetic retinopathy by the location of the new vessels, DR will be around the center, while SCD retinopathy in the periphery.

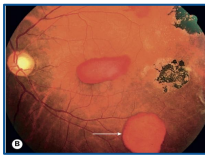


Yellow: New vessels with bleeding. Green: Retinal ischemia. Occlusion of peripheral retinal circulation causing ischemia. You can see new vessels with bleeding. To prevent bleeding, you have to apply laser (scattered laser to the area of retinal ischemia).





Fluorescein angiography shows retina is not vascularized. The picture on the left is early fluorescein angiogram showing massive ischemia, and on the right is delayed fluorescein angiogram which shows a big patch of complete filling of the **new vessels** with leakage around it (new vascular tufted filled with fluorescein).





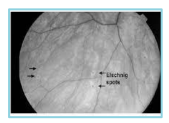


Keith Wagener grouping:

- Stage I & II: Arteriolar attenuation (silver and copper wire in the artery), increased light reflex. Because of the high pressure
- State III: Cotton wool spots (infarctions), hard exudates, hemorrhages, macular star, retinal edema.
- **Stage IV:** All of the above + edema of the optic disc.

Ischemic choroidal infarcts (Elschnig's Spots):

- Retinal arterial macroaneurysm.
- Ischemic optic neuropathy.
- As a compensatory phenomenon for high blood pressure, the first thing to happen is that the retinal arterioles become smaller (they attenuate) then the walls of arterioles become thicker, so it will reflect more light. This will create what we call "copper wire and silver wire arteries". Then we see occlusion of retinal arterioles which appears as cotton wool spots and exudates then we can see hemorrhages as a result of severe hypertension and **macular edema** then the last stage we expect to see **edema** of the optic nerve head.





- Yellow: Hard exudates (radial distribution)
- Green: Nicking "nipping" Feature of hypertensive Retinopathy Retinal vein is deviated and narrowed by overlying retinal artery.
- what is special about these hard exudates compared to diabetes? Here it is arranged radially & this is pathognomonic for hypertensive retinopathy (radial distribution, macular star, fan shaped) very IMP.
- You don't see this in other conditions; In diabetic retinopathy, the hard exudate tend to form rings.
- You can see the artery with area of silver wiring, and this is where the artery is crossing over the vein.
- There is a vein under the artery that becomes attenuated and this is called nicking "nipping" (narrowing of the lumen of the vein under the artery) because the artery is becoming so thick due to the arteriosclerosis, and the vein will become constricted because the artery is pressing on it.
- Sometimes it can cause changes in the course of the vein causing deflection. Here you can see the silver wire . arteries, copper wires and radially arranged hard exudates.
- Notice the vessel it is called silver wire and the end of the vessels copper wire due to thickening of the walls of the arteries.

CLASSIC CASE



- Yellow: Optic Disk Swelling.
- Green: Silver Wire.
- Blue: Copper Wire.
- You can see optic nerve head swelling (edema), look at the arteriole the color there is white (this is what they call silver wire) & the rest is reddish (this is what they call copper wire).



- Yellow: Hard exudates.
- Green: Cotton Wool spots.
- This is another young patient with pheochromocytoma. There are many cotton wool spots, and you can see the radial distribution of hard exudates.

• Large & medium sized vessels are affected (e.g. temporal artery).

- Over 60 years old.
- Sudden visual loss due to anterior ischemic optic neuropathy profound unilateral visual loss.
 - Why is it important? Because Patients can present with sudden loss of vision.
- Amaurosis fugax which means recurrent attacks of loss of vision before complete visual loss.
- Anterior ischemic optic neuropathy "main reason for blindness" occlusion of small blood vessels supplying the optic nerve (there is another type of ischemic optic neuropathy not related to giant cell arteritis called NAION, we usually see it in diabetic hypertensive patients).

Giant Cell Arteritis

- **Non-arteritic anterior ischaemic optic neuropathy (NAION):** more common, caused by occlusion of the short posterior ciliary arteries resulting in partial or total infarction of the optic nerve head. Patient complains of sudden painless monocular visual loss; this is frequently discovered on awakening, suggesting a causative role for nocturnal hypotension.
- Arteritic anterior ischaemic optic neuropathy (AAION): caused by giant cell arteritis (GCA). About 50% of patients with GCA have polymyalgia rheumatica (PMR)" pain and stiffness in proximal muscle groups, typically the shoulders and biceps, that is worse on waking, scalp tenderness and jaw claudication".



- Central retinal artery occlusion.
- Cotton wool spots.
- Anterior segment necrosis.
- Oculomotor palsies and Cortical blindness.
- Scalp tenderness and jaw claudication



- Over 60 years, females.
- Smoking, low body mass index.
- early menopause

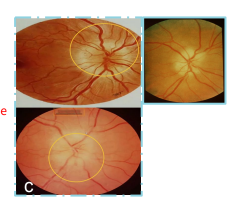
- Gangrene of scalp because of temporal arteritis.
- This is what happens: patient can present to the ER with blindness in one eye (no light perception). When we look to the optic nerve we see white optic nerve and the margins are ill-defined "means it's swollen" and we call this pale disc swelling 'chalky white' edematous disc <u>"characteristic feature"</u> and this is a sign of ischemic optic neuropathy because the disease will cause occlusion of the small blood vessels that supply the optic nerve. Admission is required in such patients, because it's a life-threatening disease.
- Confirmed by temporal artery biopsy, ESR and CRP at the ER
- Why treated by systemic corticosteroids therapy? To protect the normal eye, while the affected eye will not resolved anymore because of the occlusion of arteries that supply optic nerve.



- Yellow: Pale optic disk swelling with ill-defined margin (typical ischemic neuropathy).
- Pic C: Patient presented to the ER and you can see white disc. In this situation we always admit the patient. We need to confirm the diagnosis, so we do ESR (high ESR) and we do temporal artery biopsy (confirmatory) then if the diagnosis is confirmed you have to give the patient a large dose of systemic corticosteroids. We do an urgent temporal artery biopsy but if the patient has high ESR and C-

reactive protein we can start systemic corticosteroids immediately. **Why?** To protect the other eye because in the affected eye you cannot reverse





Summary

Diabetes Mellitus

• Risk factors:

- <u>Non-Modifiable:</u>
 - Duration (strongest):
 - Patients with type II DM must be screened early after the diagnosis
 - Patients with type I DM, no need to screen for DR unless they had 5 or more years
 - Type: Patients with type 1 diabetes tend to have more aggressive disease
- <u>Modifiable:</u>
 - Glycemic control
 - Blood pressure
 - Blood lipid control
 - Others: smoking, obesity, pregnancy
- Ocular manifestations:
 - Iris: rubeosis Iridis (neovascular glaucoma, Treated by Pan-Retinal Photocoagulation
 - Lens: cataract
 - o Iridocyclitis: inflammation of the iris and of the ciliary body
 - Retinopathy: the most common and major problem
 - Optic neuropathy and 3rd, 4th & 6th nerve palsies

Non Proliferative Diabetic Retinopathy (NPDR)

- Retinal vessel microangiopathy → blood leaks → retinal hemorrhages → retinal infiltration with lipids and fluid → macular edema
- Ocular manifestations:
 - Signs of leakage:
 - Microaneurysms (red dots): source of leakage
 - Hard exudates: result of leakage
 - Macular edema: may cause moderate visual loss
 - Signs of occlusion (ischemia) severe non- proliferative retinopathy (progress to PDR within 1 year):
 - **Cotton-wool spots:** early sign of infarction of the nerve fiber layer
 - Venous looping and beading: most reliable and important signs of retinal ischemia
 - Intra-retinal microvascular abnormalities (IrMAs): dilated telangiectatic vessels
 - intraretinal hemorrhages in 4 quadrants

Proliferative Diabetic Retinopathy (PDR)

- Retinal vessel microangiopathy → chronic retinal hypoxia → abnormal proliferation of blood vessels → traction on retina → retinal detachment
- Findings of nonproliferative retinopathy are usually present
- Neovascularization: the most important site is optic nerve head

Management

- Laser photocoagulation: in nonproliferative retinopathy to occlude the aneurysms (red dots)
- Anti-VEGF agents: to prevent further neovascularization in patients with proliferative retinopathy and macular edema and enhance the effect of laser
- Laser panretinal photocoagulation: in severe nonproliferative retinopathy or proliferative retinopathy

Summary

Graves' disease

• Ocular manifestations:

- Most common cause of both bilateral and unilateral proptosis in an adult.
- Eyelid retraction and Lid lag
- Infiltrative ophthalmopathy
- Dysthyroid optic neuropathy
- Proptosis and exophthalmos
- Chemosis, exposure keratopathy, ophthalmoplegia
- Restrictive thyroid myopathy
- Diagnosis:
 - Thyroid function test: high T3, T4 and low TSH.
 - \circ $\;$ Visual evoked potential: to exclude optic neuropathy
 - $\circ~$ CT scan if suspecting thyroid eye disease
 - \circ $\;$ Hertel ophthalmometer To measure the degree of proptosis or exophthalmos.

Tuberculosis

• Ocular manifestations:

- Phlyctenular keratoconjunctivitis
- Interstitial keratitis and Vitritis
- Uveitis: granulomatous & non- granulomatous, if the keratic precipitates are big "mutton-fat" it indicates a granulomatous uveitis (mutton-fat keratic precipitate)
- Retinal vasculitis 'Eales disease'
- $\circ~$ Granuloma of the choroid (Tubercle), If large enough \rightarrow tuberculoma
- Diagnosis:
 - Clinical findings consistent with tuberculosis.
 - Rule out other specific uveitic entities (e.g. Behcet disease, sarcoidosis, etc).
 - Investigations (tuberculin test).
- Management: Prolonged Anti-TB therapy

Sarcoidosis

• Ocular manifestations:

- Optic nerve, retinal, choroidal, Lid margin and conjunctival granuloma (non-caseating granulomas)
- Mutton-fat keratic precipitates
- Acute iridocyclitis and Chronic granulomatous iridocyclitis
- Peripheral retinal periphlebitis

• Diagnosis:

- Tuberculin skin test: negative in sarcoidosis.
- Chest X-ray\CT: showing bilateral hilar lymphadenopathy (BHL)
- Management: Immunosuppressive with systemic steroids

Leprosy (Hansen's Disease)

- Ocular involvement is more common in the lepromatous type.
- Signs: Facial nerve affection, Loss of the lateral portions of the eyebrows and eyelashes (madarosis), Interstitial keratitis with beading of nerves, iritis.



Summary

Vogt-Koyanagi-Harada Syndrome

idiopathic multisystem T cells mediated autoimmune disease featuring inflammation of

melanocyte-containing tissues

- Ocular manifestations:
 - Mutton-fat keratic precipitates
 - most common cause of autoimmune uveitis in the country
 - \circ $\,$ Poliosis: absent or decreased melanin in head hair, eyebrows or eyelashes $\,$

• Systemic manifestations:

- Vitiligo
- \circ deafness
- neurological signs

• Management:

- High-dose steroid
- o immunomodulatory agent such as mycophenolate mofetil

• Complications if untreated:

- Poliosis
- Vitiligo
- Posterior synechiae
- Glaucoma, Cataract

Q1- What is the effect of vascular leakage in diabetic retinopathy?

- A. Cotton wool spots
- B. Neovascularization
- C. Hard exudate
- D. Venous beading

Q2- Young male presented to the ER with a unilateral acute non-granulomatous anterior uveitis patient complains of back pain that is worse in the morning, what is the appropriate test to order for this patient?

A. Temporal artery biopsyB. HLA-B27C. Antiphospholipids antibodiesD. HLA-DR4

Q3- A 27 year old male presented to ophthalmology clinic found to have cataract, long fingers, bone deformities and cardiac abnormalities which was diagnosed with Marfan syndrome, which eye manifestation could found in this picture :

- A. Glaucoma
- B. Granuloma formation
- C. Posterior synechiae
- D. Lens subluxation



Q4- A 4 year old child presented to ophthalmology clinic, on examination patient found to have cataract, pigmented retina and microphthalmos, his mother mentioned that he have deafness and heart anomaly since birth, what is the most likely diagnosis?

- A. Toxoplasmosis
- B. Sarcoidosis
- C. Rubella
- D. Marfan syndrome



Q5- A 76 year old hypertensive male presented to the ER with sudden visual loss and jaw claudication on examination patient has scalp tenderness what is the most likely diagnosis?

A. Giant cell arteritis B. Hypertensive retinopathy C.Diabetic retinopathy D. Graves' disease

Short Answer Questions





A: what is your finding?

B: if the patient diagnosed with Vogt-Koyanagi-Harada Syndrome what other complications you might find?

Case 2



A: Yellow circle indicates?

B: Green circle indicates?

C: What is the difference between hard exudate in hypertensive patients compared to diabetics?

Answers:

Case 1 A: Posterior synechiae B: Poliosis,Vitiligo,Mutton-fat keratic precipitates in the eye

Case 2 A: Hard exudates (radial distribution) B: Nicking "nipping" (Retinal vein is deviated and narrowed by overlying retinal artery) C: It is arranged radially in hypertensive patients & this is pathognomonic for hypertensive retinopathy (radial distribution , macular star, fan shaped). This work was originally done by **438 and 439 Ophthalmology Team**

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