



Pharmacology of eye

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Sympathetic Innervation:

- 1- to dilator pupillae muscle radial muscles of the iris, leading to pupil dilation (mydriasis)
- 2- to blood vessels within the eye, influencing ocular blood flow and intraocular pressure
- 3- to Müller's muscle leading to eyelid retraction
- 4- Beta-2 adrenoceptors in the ciliary body increase the secretion of aqueous humor but Alpha-2 adrenoceptors in the ciliary body suppress it.

Parasympathetic innervation:

- 1-To constrictor pupillae muscle, narrowing the pupil in response to bright light (light reflex).
- 2-To the ciliary muscle, causing it to contract, leading to lens accommodation.

Drainage of aqueous humor:

Aqueous humor flows from the posterior chamber → anterior chamber → exits via two routes:

1- Conventional Pathway (90% of outflow):

Fluid traverses the trabecular meshwork → Schlemm's canal → episcleral veins.

2-Unconventional Pathway (10% of outflow):

Fluid drains through the ciliary muscle, suprachoroidal space, and sclera (uveoscleral route).

Drugs

1- Drugs affecting pupil size

2-Treatment of Glaucoma

3-Drugs that ↑↑ IOP

A- Drugs affecting pupil size

1- Miotics drugs

Drug Class	Examples	Effect on Pupil	Mechanism
Opioids (systemic)	Morphine, Heroin, Fentanyl	Miosis	Activates μ -opioid receptors, inhibiting sympathetic tone.
Cholinergic Agonists (local)	Pilocarpine, Carbachol	Miosis	Stimulates parasympathetic system (muscarinic receptors).
Acetylcholinesterase Inhibitors (local)	Physostigmine, Neostigmine,	Miosis	Increases acetylcholine levels, activating muscarinic receptors
Guanthiden		Miosis	Reduces Release of NE in the eye:
α 1-Adrenergic Blockers	Prazosin, Tamsulosin	Miosis	Blocks sympathetic stimulation of the dilator muscle
Sedatives / Barbiturates	Benzodiazepines (high doses)	Miosis	CNS depression reduces sympathetic tone.

Locally acting miotics

(parasympathomimetics): stimulate M3 receptors in

1- CPM → miosis + wide angle of filtration & space of Fontana.

2- Ciliary muscle → accommodation to near vision + open canal of Schlemm.

3- Some stimulate Nm receptors in upper eye lid → upper eyelid twitches.

1- Direct parasympathomimetics:

-Choline esters: bethanichol(M only) &carbachol (M+N).

- Alkaloid:pilocarpine(M only).

2- Indirect parasympathomimetics:

-Reversible:physostigmine(eserine) &demecarium.

-Irreversible :organophosphorus→ecothiophate&isofluorophate: Long-lasting strong effect with extreme miosis, but produce irritation &cataract

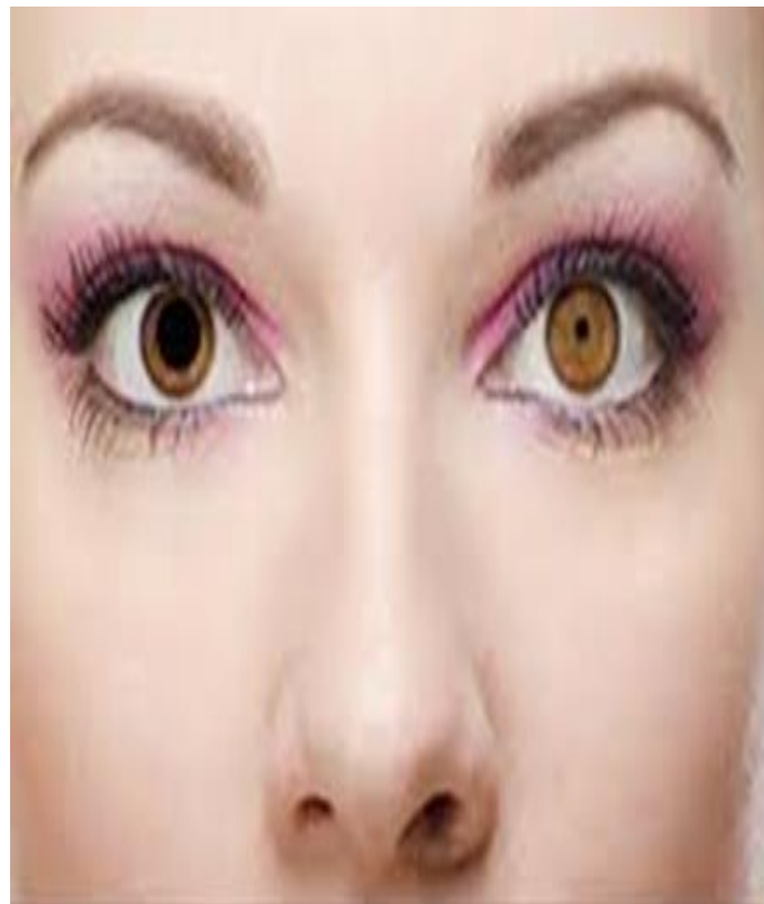
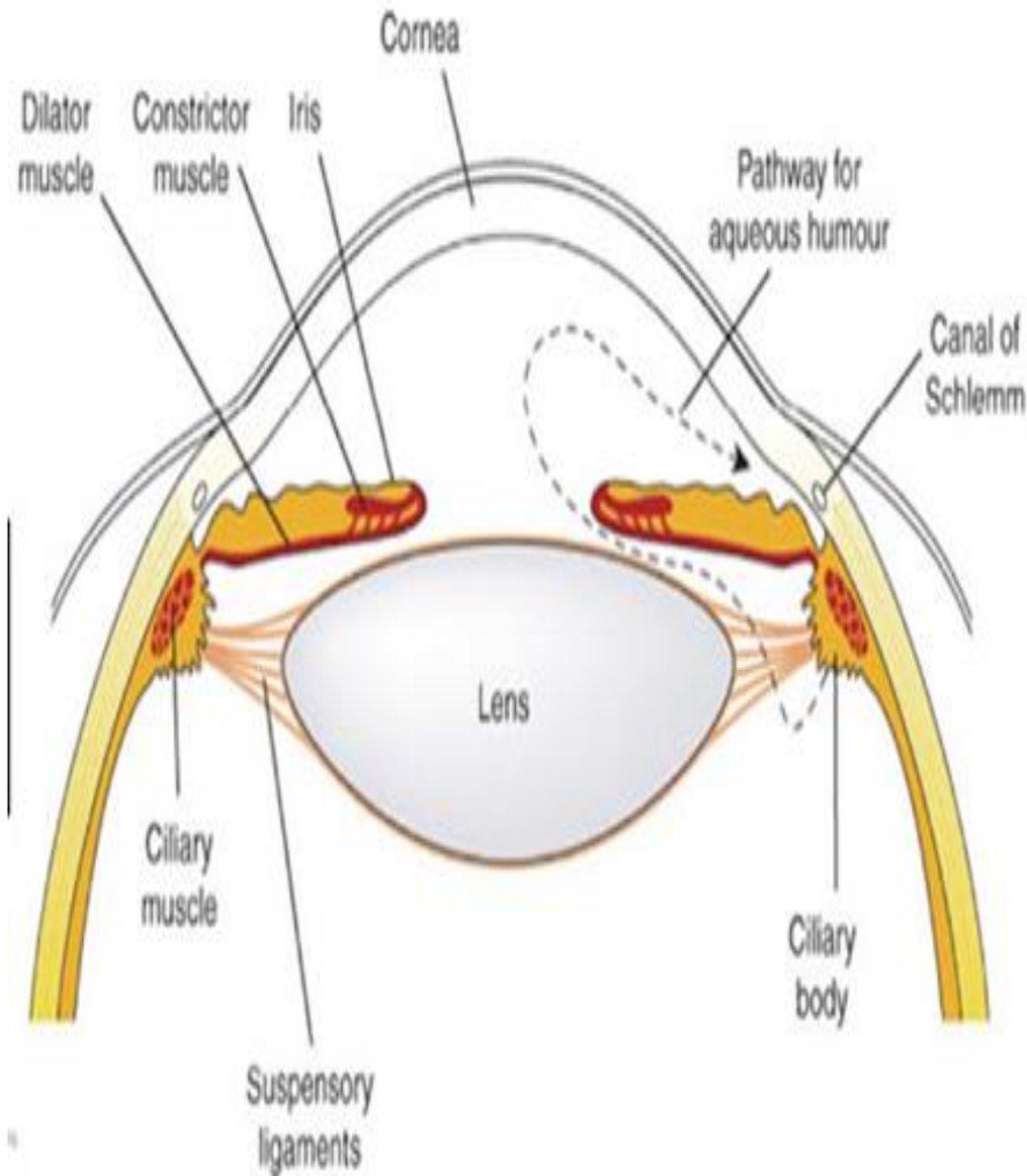
Therapeutic uses:

1-Glaucoma.

2-Counteract mydriatics after fundus examination.

3-Alternatively with mydriatics to cut adhesion between iris & lens.





Opiate use or overdose is one of the most common causes of pinpoint pupils.

Guanthidine:

Paralysis of Dilator Pupillae Muscle → miosis + ↓↓ IOP

Relaxation of levator palpebrae superioris → ↓↓ exophthalmos of hyperthyroidism.

Morphine stimulates **opiate receptor** in **3rd cranial nerve nucleus** → stimulates oculomotor nerve → **ciliary ganglia (Nn)** → eye → ACh → stimulates **M3** receptors of **CPM** → marked **miosis** (pin point pupil).

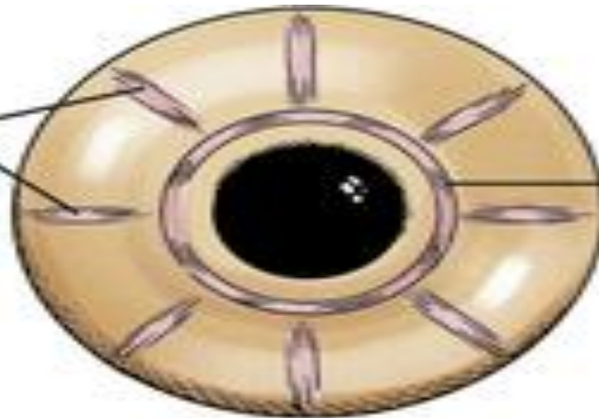
Pin-point pupil of morphine can be antagonized by:

- 1) Systemic naloxone** → block opiate μ receptors in CNS.
- 2) Systemic ganglion blocker** → block Nn of ciliary ganglia.
- 3) Topical or systemic atropine** → block M3 receptors on CPM

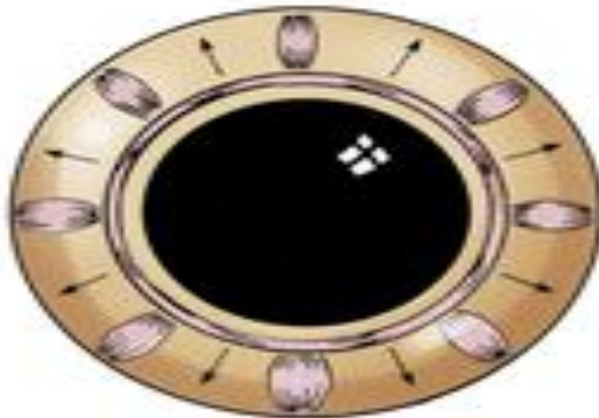
2- Mydriatics drugs

Drug Class	Examples	Effect on Pupil	Mechanism
Sympathomimetics (indirect).	Epinephrine, Cocaine, Amphetamines	Mydriasis	Stimulates adrenergic receptors, enhancing sympathetic activity
Anticholinergics	Atropine, Tropicamide, Scopolamine	Passive Mydriasis	Blocks parasympathetic innervation to the constrictor pupille muscle.
α 1-Adrenergic Agonist	Phenylephrine	Mydriasis	Stimulates dilator muscle via α 1 receptors.
SSRIs & SNRIs	Fluoxetine, Venlafaxine	Mydriasis	Increased serotonin activity affects autonomic control
Tricyclic Antidepressants	Amitriptyline, Imipramine	Mydriasis	Strong anticholinergic effects block pupil constriction.
Hallucinogens	LSD, MDMA	Mydriasis	Serotonin and dopamine effects increase sympathetic tone
Dopaminergic Drugs	Levodopa, Bromocriptine	Mydriasis	Enhances dopamine signaling, indirectly increasing sympathetic effects

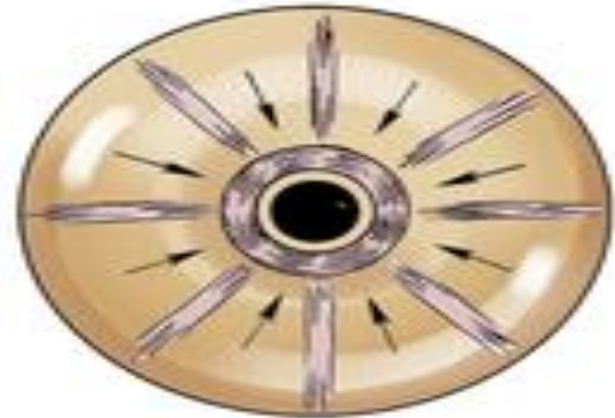
Dilator pupillae muscle (sympathetic)



Constrictor pupillae muscle (parasympathetic)



DILATION (mydriasis)
Dilators contract
Constrictors relax



CONSTRICTION (miosis)
Constrictors contract
Dilators relax

A- Sympathomemetic:

mechanism: Stimulate **$\alpha 1$ receptors** leading to:

Contraction of **DPM** \rightarrow **Active mydriasis** (intact light reflex) & no cycloplegia ,
BV \rightarrow VC \rightarrow decongestion & \downarrow IOP.

○ **Examples:** Direct: phenylephrine., Indirect: amphetamine, Mixed: ephedrine.

○ **Therapeutic uses:** fundus examination especially in elderly patients liable for glaucoma.

B- Cocaine:

□ **Surface anesthesia** \rightarrow loss of sensory reflex (corneal & conjunctival reflex)

□ **Indirect sympathomimetic:** \downarrow neuronal uptake (1) $+$ MAO inhibitor \rightarrow \uparrow endogenous NA \rightarrow stimulates **$\alpha 1$ receptors** \rightarrow **active mydriasis & decongestion.**

C)Parasympatholytics:

Mechanism: Block M3 receptors in:

- 1) **CPM** → passive mydriasis → lost light reflex & narrow angle of filtration.
- 2) **Ciliary muscle** → cycloplegia (loss of accommodation) + closing canal of Schlemm.
 - Result is **lost light reflex + cycloplegia + ↑↑ IOP. Examples:**
 - 1) **Natural belladonna alkaloids: atropine & hyoscine.**
 - 2) **Synthetic:** homatropine, cyclopentolate , tropicamide & eucatropine.
 - **Therapeutic uses:**
 - 1) **Atropine** is used in iritis and corneal ulcer (to prevent adhesions), and measurement of refraction in children.
 - 2) **Synthetic substitutes:** in fundus examination.

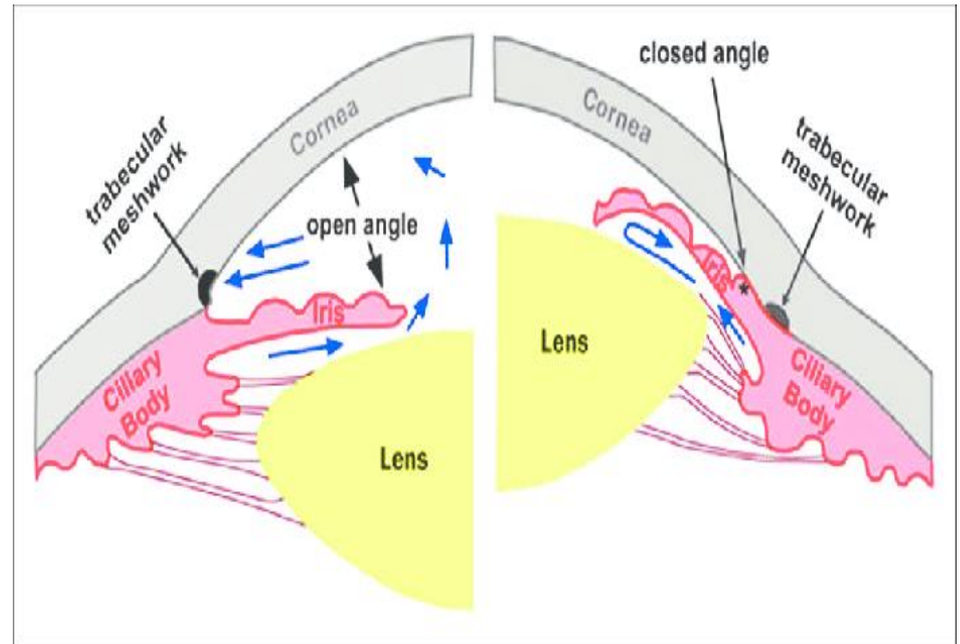
All are contraindicated in glaucoma

Treatment of Glaucoma

Normal Intra-Ocular Pressure (IOP) = 15-25 mmHg.

Glaucoma may be :

- 1) **Closed angle** (narrow-angle) glaucoma
- 2) **Open-angle glaucoma**
(Chronic glaucoma)



I- Closed angle (narrow-angle) glaucoma:

- ❑ Needs **surgical intervention** (iridectomy).
- ❑ Due to **occlusion of angle of filtration** by iris root coming in contact with periphery of the cornea (Acute congestive glaucoma).

Drugs used to decrease I.O.P before surgery are:

- 1) **Miotic eye drops:** a) **Pilocarpine** (of choice) with low concentration.
b) **Physostigmine** (not preferred due to congestion & extreme miosis).
- 2) **Carbonic anhydrase inhibitors:** acetazolamide ($\downarrow\downarrow$ aqueous secretion)
- 3) **Osmotic agents (dehydrating agent):** mannitol (20%) IV, MgSO₄ rectally & Glycerine (50%) orally: they produce rapid reduction of IOP.
- 4) **Brimonidine & apraclonidine** (α_2 agonists).
- 5) **Recently β -Blockers can be used with pilocarpine**

II- Open-angle glaucoma (Chronic glaucoma):

• Drugs used are:

1) Miotic eye drops

(Pilocarpine & Physostigmine).

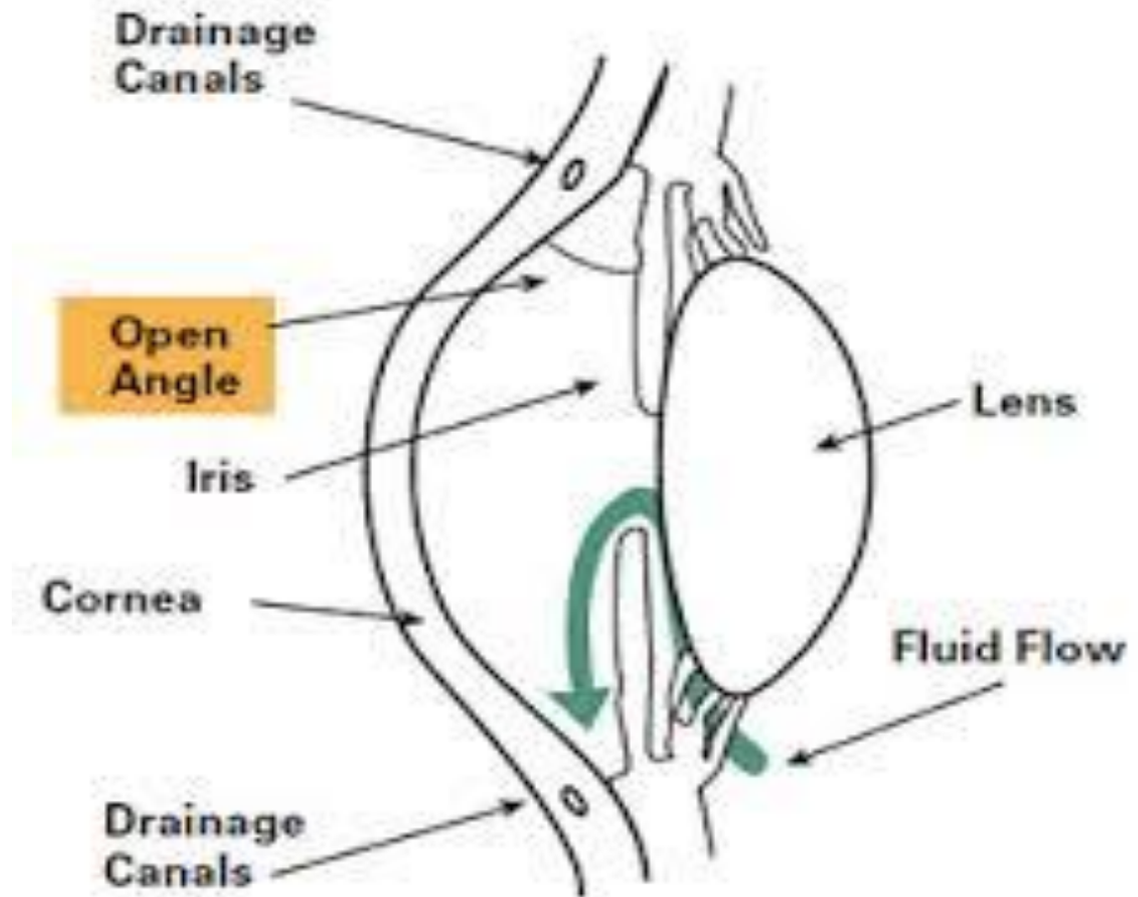
1) Carbonic anhydrase inhibitors:

(inhibit aqueous formation):

a- Acetazolamide orally

b- Dorzolamide & Brinzolamide

(locally)



3- Sympathomimetic eye drops: (Adrenaline & Dipivefrin) → VC → decrease synthesis of aqueous humor.

4- B-blockers: decrease cAMP → decrease aqueous humor e.g. timolol & betaxolol. Side effects: tolerance, systemic absorption.

5) α_2 agonists: a- Apraclonidine: used only for short time due to tachyphylaxis. b- Brimonidine: decrease aqueous secretion & ↑ uveoscleral outflow.

Side effects: allergic conjunctivitis, dry mouth & fatigue.

6) PGF 2α analogues e.g. Latanoprost, travoprost & bimatoprost:

They decrease IOP by ↑↑ uveoscleral outflow.

The most potent ocular hypotensives.

Side effects: conjunctival hyperpigmentation & hyperemia, and headache.

7) Guanethidine

Drugs that ↑↑ IOP:

1. **Parasympatholytics** (atropine).

2. **Drugs with atropine-like effect:**

a) Some H1 blockers (Diphenhydramine).
(Disopyramide).

b) Some antiarrhythmics

1. **Ganglion blockers.**

2. **Corticosteroids.**

3. **Nitrates.**

Thank you