



☒ Sheet

☐ Slides

number : 6

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This sheet is dependant mainly on slides – with some notes – rather than depending on the record

Cardiovascular System

Atropine causes tachycardia by vagal block.

Lower doses often result in initial **bradycardia** before the effects of peripheral vagal block is seen.

This slowing may be due to block of **M1 autoreceptors on vagal postganglionic fibers**.

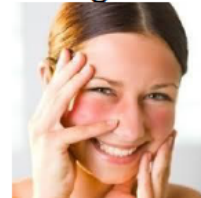
The ventricles are less affected

In toxic concentrations, it can cause intraventricular conduction block due to a local anesthetic action.

All blood vessels contain endothelial muscarinic receptors that mediate vasodilation.

These receptors are blocked by antimuscarinic drugs.

At toxic doses, antimuscarinic agents cause **cutaneous vasodilation**, especially in the the blush area. The mechanism is unknown.



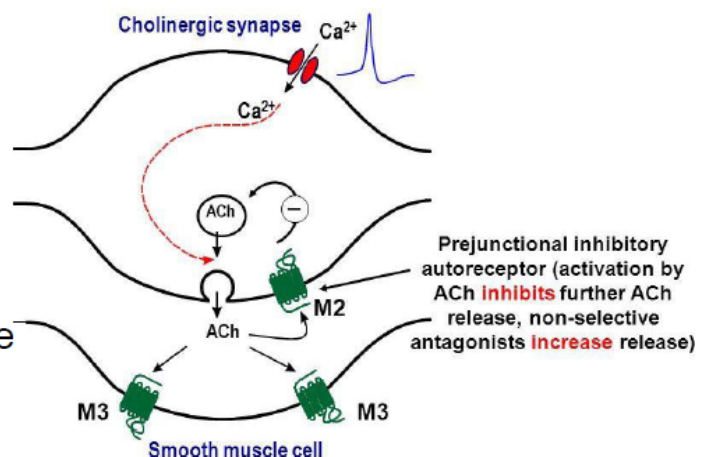
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-(the blush area) is "the face and the neck"

-The mechanism of this vasodilation is unknown but maybe it is due to sweat inhibition which leads to increase the body temperature so we will have vasodilation in that area in order to decrease the temperature.

Respiratory System

Atropine causes some **bronchodilation** & **reduce secretion**.



The effectiveness of nonselective antimuscarinic drugs in treating **bronchial asthma** is limited because block of **autoinhibitory M2** oppose the bronchodilation caused by block of **M3** receptors on airway.

Antimuscarinic drugs are frequently used before the administration of **inhalant anesthetics** to reduce the accumulation of secretions in the trachea.

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Gastrointestinal Tract

Complete muscarinic block cannot totally abolish activity of GIT, since local hormones in the enteric nervous system also modulate GI functions.

Antimuscarinic drugs have marked effects on salivary secretion causing **dry mouth**

Gastric secretion is blocked less effectively: the volume and amount of acid, pepsin, and mucin are all reduced, but large doses of atropine may be required.

Basal secretion is blocked more effectively than that stimulated by food, nicotine, or alcohol.

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Gastrointestinal drugs : Pirenzepine, telenzepine are mentioned below

Gastrointestinal Tract cont..

Pirenzepine and telenzepine

M1 blockers. Reduce gastric acid secretion with fewer adverse effects than atropine

GI smooth muscle **motility** is affected from the stomach to the colon and both tone and propulsive movements are diminished.

Gastric emptying time is prolonged, and **intestinal transit time is lengthened**.

Diarrhea due to overdosage with muscarinic agents is readily stopped.

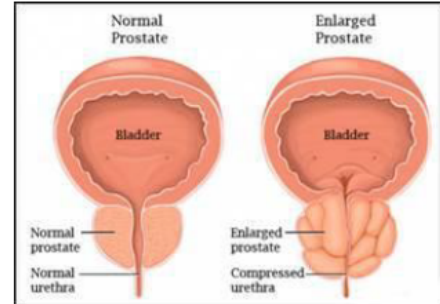
Diarrhea caused by nonautonomic agents can usually be temporarily controlled.

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Genitourinary Tract

Relaxes smooth muscle of the ureters and bladder wall and slows voiding.

Useful in the **treatment of spasm** induced by mild inflammation, surgery, and certain neurologic conditions, but it can precipitate urinary retention in men who have **prostatic hyperplasia**



Sweat Glands

Atropine suppresses **sweating**.

In adults, body temperature is elevated only with large doses, but in infants and children even ordinary doses may cause "**atropine fever**."

Therapeutic Applications of the anti-muscarinic agents:

1. Central Nervous System Disorders, "Parkinson's Disease"

Usually the treatment of Parkinson disease is by giving dopaminergic drugs like L-dopa that is converted in the brain into dopamine. But sometimes they add an **adjunctive** therapy in some patients but with all of the adverse effects.

2. Motion Sickness "nausea and vomiting caused by over stimulation of the Body Balance Center"

☐ **Scopolamine** is one of the oldest remedies & is as effective as any more recently introduced agent.

☐ Given by injection or by mouth or as a transdermal patch.

☐ The **patch formulation** produces significant blood levels over 48–72 hours.

☐ Useful doses by any route usually cause significant sedation and dry mouth.

Antimuscarinic Drugs Used in Ophthalmology.

Drug	Duration (days)	Usual Concentration(%)
Atropine	7–10	0.5–1
Scopolamine	3–7	0.25
Homatropine	1–3	2–5
Cyclopentolate	1	0.5–2
Tropicamide	0.25	0.5–1

Ophthalmologic Disorders

Antimuscarinic agents, as eye drops or ointment, produce **mydriasis and cycloplegia** are very helpful in doing a complete examination.

The shorter-acting drugs are preferred

Should never be used for mydriasis unless cycloplegia or prolonged action is required.

Alpha- adrenoceptor stimulant drugs, **phenylephrine**, produce a short mydriasis sufficient for funduscopy examination.

Antimuscarinics also used to prevent **synechia**.



A **synechia** is an eye condition where the iris adheres to either the cornea or lens. The longer-lasting preparations, especially homatropine, are preferred.

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Respiratory Disorders

Atropine was routinely used as a **preoperative** medication when anesthetics such as ether were **used to decrease airway secretions and to prevent laryngospasm**. Newer inhalational anesthetics are far less irritating to the airways.

Scopolamine also produces significant **amnesia** for the events associated with surgery and **obstetric delivery**.

Urinary retention and intestinal hypomotility following surgery are exacerbated by antimuscarinic drugs.

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Ipratropium

a synthetic analog of atropine, is used as an inhalational drug in **asthma** with reduced systemic effects.

Ipratropium is also useful in chronic obstructive pulmonary disease (COPD) a condition that occurs more frequently in older patients, particularly chronic smokers.

Tiotropium

has a longer bronchodilator action and can be given once daily.

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Ipratropium and Tiotropium are Drugs work on the lungs

Cardiovascular Disorders

Marked **reflex vagal discharge** sometimes accompanies the pain of **myocardial infarction** (e.g., vasovagal attack) and may depress sinoatrial or atrioventricular node function sufficiently to impair cardiac output.

Atropine is used in this situation.

Rare individuals have **hyperactive carotid sinus reflexes** and may experience faintness or even syncope as a result of vagal discharge in response to pressure on the neck, e.g., from a tight collar.

Such individuals may benefit from the use of **atropine** or a related antimuscarinic agent.

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Gastrointestinal Disorders

Antimuscarinic agents can provide some relief in the treatment of common **traveler's diarrhea** and other mild hypermotility.

They are often combined with an **opioid antidiarrheal drug**.

Atropine with **diphenoxylate**, (**Lomotil**) is available in both tablet and liquid form.

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Urinary Disorders

Provide symptomatic relief in the treatment of **urinary urgency** caused by minor inflammatory bladder disorders.

Oxybutynin

more selective for M3 receptors, is used to relieve bladder spasm after urologic surgery.

It reduce involuntary voiding in patients with neurologic disease.

Darifenacin

has greater selectivity for M3 receptors & long half-life used in adults with urinary incontinence.

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Oxybutynin and Darifenacin are Urogenital medications

An alternative treatment for urinary incontinence refractory to antimuscarinic drugs is intrabladder injection of **botulinum toxin A**.

By interfering with the release of neuronal acetylcholine, **botulinum toxin** is reported to reduce urinary incontinence for several months after a single treatment.

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Cholinergic Poisoning

Caused by cholinesterase inhibitor & wild mushrooms

Atropine is used to reverse the muscarinic effects, to treat the CNS effects as well as the peripheral effects of the organophosphate inhibitors.

Large doses of atropine may be needed to oppose the muscarinic effects of extremely potent agents like **parathion** and chemical warfare nerve gases.

1–2 mg of atropine sulfate may be given **IV** every 5–15 minutes until signs of effect (**dry mouth, reversal of miosis**) appear.

The drug is **repeated many times**, since the acute effects of the anticholinesterases may last 24–48 h.

1 g of atropine per day may be required for **one month** for full control of muscarinic excess.

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Adverse Effects

Treatment with atropine or its congeners induces undesirable effects.

At higher concentrations, atropine causes block of all parasympathetic functions.

Poisoned individuals manifest:

dry mouth, mydriasis, tachycardia, hot and flushed skin, agitation, and delirium for as long as 1 week.

Children, especially infants, are **very sensitive** to the hyperthermic effects of atropine.

Deaths have followed doses as small as 2 mg.

Overdoses of atropine are treated **symptomatically**

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Adverse Effects cont..

When **physostigmine** is used, **small** doses are given *slowly* intravenously.

Symptomatic treatment may require temperature control with cooling blankets and **seizure** control with **diazepam**.

Poisoning by high doses of **quaternary antimuscarinic** drugs is associated with all of the peripheral signs but few or none of the CNS effects of atropine.

They may cause **ganglionic blockade** with marked orthostatic hypotension

Treatment of the antimuscarinic effects can be carried out with a quaternary cholinesterase inhibitor such as **neostigmine**.

Control of hypotension may require the administration of a sympathomimetic drug such as **phenylephrine**.

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Contraindications

Glaucoma

Even systemic use of moderate doses may precipitate angle closure (and acute glaucoma) in patients with shallow anterior chambers.

Prostatic hyperplasia

In **elderly men**, antimuscarinic drugs should always be used with caution and should be avoided in those with a history of prostatic hyperplasia.

Nonselective antimuscarinic agents should never be used to treat acid-peptic disease.

Because the antimuscarinic drugs slow gastric emptying, they may *increase* symptoms in patients with gastric ulcer.