## Subject Name: Pharmacology

## WINTER – 2023 EXAMINATION

Model Answer – Only for the Use of RAC Assessors

# Subject Name: Pharmacology

## **Important Instructions to examiners:**

- 1) The answers should be examined by key words and not as word-to-word as given in the model answer scheme.
- 2) The model answer and the answer written by candidate may vary but the examiner may try to assess the understanding level of the candidate.
- 3) The language errors such as grammatical, spelling errors should not be given more Importance (Not applicable for subject English and Communication Skills.
- 4) While assessing figures, examiner may give credit for principal components indicated in the figure. The figures drawn by candidate and model answer may vary. The examiner may give credit for any equivalent figure drawn.
- 5) Credits may be given step wise for numerical problems. In some cases, the assumed constant values may vary and there may be some difference in the candidate's answers and model answer.
- 6) In case of some questions credit may be given by judgement on part of examiner of relevant answer based on candidate's understanding.
- 7) For programming language papers, credit may be given to any other program based on equivalent concept.
- 8) As per the policy decision of Maharashtra State Government, teaching in English/Marathi and Bilingual (English + Marathi) medium is introduced at first year of AICTE diploma Programme from academic year 2021-2022. Hence if the students write answers in Marathi or bilingual language (English +Marathi), the Examiner shall consider the same and assess the answer based on matching of concepts with model answer.

Q. No.	Sub Q. N.	Answer	Marking Scheme
1		Answer any <u>SIX</u> of the following:	30 M
1.	a	Classify antibiotics. Write note on anthelmintics.	3M Classify 2M
		Classification:	Note
		1) Effective against gram-positive bacteria	
		a) Used for systemic infections eg. Penicillin, Erythromycin	
		b) Used topically e.g. Bacitracin.	
		2) Effective against Gram Negative bacteria	



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a) Used for systemic infections eg Streptomycin, Kanamycin		
b) Used locally in intestinal infections eg. Paromomycin		
3) Effective against both gram positive and Gram Negative bacteria		
a) Used for systemic infections eg. Ampicillin, Amoxicillin, Carbenicillin		
b) Used topically eg. Neomycin, Framycetin		
4) Effective against gram-positive and gram-negative bacteria ,rickettsiae		
and Chlamydia eg Tetracycline, Chloramphenicol		
5) Effective against acid fast bacilli eg. Rifampicin, Streptomycin		
1. Inhibitors of cell wall synthesis eg Penicillins		
2. Inhibitors of cell membrane function eg. Polymixin		
3. Inhibitors of protein synthesis eg. Tetracyclins		
4. Inhibitors of nucleic acid synthesis/ function; eg. Rifampicin		
5. Inhibitors of metabolism eg. Sulpha drugs.		
Or		
Effective against gram +ve bacteria: Penicillin etc		
Effective against gram -ve bacteria: Streptomycin etc		
Effective against both gram +ve & gram -ve bacteria:		
Tetracycline, Chloramphenicol.etc		
Effective topically :Framycetin ,Polymyxin B,neomycin etc		



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		Any other correct classification can be considered.	
		Note on Anthelmintics:	
		Anthelmintics are the pharmacological agents which kill or expel the worms and are used in treatment of helminthiasis.	
		Classification:	
		<ol> <li>For roundworm, hookworm and pinworm: Mebendazole, albendazole, piperazine, levamisole, pyrantel pamoate</li> <li>For whipworm, Trichinella spiralis: Mebendazole, albendazole.</li> </ol>	
		3. For tapeworms: Praziquantel, niclosamide, albendazole.	
		4. For Hydatid disease: Mebendazole.	
		5. For threadworm: Ivermectin, albendazole	
		6. For filariasis: Diethylcarbamazine, ivermectin, albendazole.	
		OR any other correct classification	
1.	b	What are different routes of drug administration? Write the advantages and	3M ROA
		disadvantages of IV route.	1M Adv. 1M Disadv.
		Enteral	
		– Parenteral	
		– Local applications	
		Enteral -	
		drug placed directly in the GI tract	
		Sublingual - placed under the tongue	
		i oral - swallowing	
		rectum - Absorption through the rectum (enema)	
		Parenteral:	
		Inhalation	
		Injections: Intravenous, Intramuscular ,Intradermal, Subcutaneous ,Intrathecal ,	
		Intraperitoneal, Intramedullary, Intraarticular etc	
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Local A	Application					
Or tal	bular format	Enteral	Pare	enteral	Local applicati	ions
Oral	Sublingual	Enema	Injections	Inhalations		
		Retention Evacuant	Intravenous Intraarterial Intramuscular			
			Subcutaneous Intraperitoneal Intrathecal			
			Intramedullary Intraarticular			

## Advantages :

- 1. Has rapid onset of action
- 2. 100% bioavailability
- 3. It is useful in medical emergencies and so is a life saving route
- 4. It can be employed in unconscious, uncooperative patients.
- 5. Drugs which irritate the stomach can be given by this route.
- 6. It avoids drug degradation by digestive juices in GIT ,or by



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		mist pass metabolism in the liver.		
		7. Accuracy of dose		
		8. It is useful in case of vomiting and diarrhoea		
		Disadvantages		
		1. Once administered, difficult to withdraw, hence less safe		
		2. More expensive		
		3. Self-medication difficult		
		4. Aseptic technique and skill required, Proper care should be taken to avoid infection		
		5. only aqueous solution can be given.		
1.	c	Describe oral hypoglycemic agents with reference to it's classification and examples.	Descript	ion
		Why insulin is not used by oral route.	3M 2M Fo	r
		Oral hypoglycemics are pharmacological agents when administered orally decrease blood	Justifica	tion
		glucose level.		
		Classification:-		
		1) Sulfonylureas		
		a) First generation:- Ex. Tolbutamide, Chlorpropamide.		
		b) Second generation:-Ex. Glibenclamide, Glipizide, Gliclazide.		
		2) Biguanides: Metformin, Phenformin.		
		3) Thiazolidinediones: Pioglitazone.		
		4) Meglitinides: Repaglinides.		
		5) Alpha Glucosidase inhibitors: Acarbose.		
		6) Newer agents: Sitagliptin, Extenaide, Canagliflozin etc.		
		1. Sulfonylureas		
		i) First generation:- Ex. Tolbutamide		
		ii) Second generation:-Ex. Glibenclamide, glipizide, gliclazide.		
		2. Meglitinides		
		Ex. Repaglinide, Nateglinide.		
		3. Glucagon like peptide-1 receptor agonists		



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	Ex. Exenatide, Liraglutide.	
	4. Dipeptidyl peptidase-4 inhibtors	
	Ex. Sitagliptin, Vildagliptin, Saxagliptin.	
	B. Overcome insulin resistance	
	I) Biguanide: Ex. Metformin.	
	II) Thaizolidinediones: Ex. Pioglitazone.	
	C. Miscellaneous antidiabetic drugs	
	a) alpha glucosidase inhibitors: Ex. Acarbose, Miglitol.	
	b) Sodium glucose cotransport-2:- Dapagliflozin.	
	Why insulin is not used by oral route?	
	I. Insulin is a polypeptide hormone secreted by beta cells of islets Langerhans	
	of pancreas.	
	II. Commercially it is extracted from pancreas of cattle or pigs or made by biotechnology	
	methods	
	III. When given orally proteolytic enzymes and gastric juice, hydrochloric acid in	
	stomach causes its degradation, polypeptide is fragmented and therapeutic effect is lost.	
	Hence Insulin is not given by oral route.	
u	Discuss different drugs used in the treatment of angina pectoris. Write mechanism	3M
	of action of any one drug.	Classification
	Angina pectoris: - It is a symptom resulting due to cardiac ischemia. It is described as a	2M MOA
	condition in which there is a compressing type of pain in the chest.	
	1. Nitrates:-	
	1. Muates	
	a) Short acting:- eg. Glyceryl trinitrate, Nitroglycerine.	
	b) Long acting:- eg. Isosorbide dinitrate, Isosorbide mononitrate, Pentaerythritol	
	tetranitrate.	
	2. Beta blockers:- eg. Propranolol, Metoprolol, Atenolol	
	3. Calcium channel blockers:-eg.Verapamil, Diltiazem ,Nifedipine, felodipine, amlodipine	
	4. Potassium channel blockers:-eg. Nicorandil	



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		5. Miscellaneous-eg. Ranolazine	
		Mechanism of action of nitrates:	
		Organic nitrates are rapidly denitrated enzymatically in the smooth muscle cells to release	
		the reactive free radical nitric oxide which causes a direct relaxant effect on vascular	
		smooth muscles, and the dilation of coronary vessels improves oxygen supply to the	
		myocardium.	
		Note: Any other drug mechanism can be considered.	
1.	e	Define General anaesthetics. What is Schizophrenia? Mention drugs used in	1M def.
		treatment of Schizophrenia.	1M Schi. 3M Treat.
		Definition:	
		General anaesthetics are the pharmacological agents which produce reversible loss of	
		consciousness by depressing CNS	
		Ex. Thiopentone sodium, Nitrous oxide, Propofol, Diethyl ether	
		Schizophrenia is a major psychotic disorder of split personality in which person is	
		detached from reality. Schizophrenia may result in some combination of hallucinations,	
		delusions, emotional outbursts, lack of interest in surrounding and overall disturbed	
		thinking. Excess of dopamine in brain is associated with Schizophrenia.	
		1) Phenothiazines	
		a) aliphatic side chain: Chlorpromazine, Triflupromazine	
		b) piperidine side chain: Thioridazine	
		c) piperazine side chain: Trifluoperazine, Fluphenazine	
		2) Butyrophenones: Haloperidol, Trifluperidol, Penfluridol	
		3) Thioxanthenes : Flupenthixol	
		4) Other heterocyclics: Pimozide, Loxapine.	
		5) Atypical antipsychotics: Clozapine, Olanzapine etc.	
1.	f	<ul> <li>i) Give the pharmacological action of Adrenaline.</li> <li>Heart: - adrenaline with its action on B-receptors of heart increases heart rate, force</li> </ul>	2.5M Each
		of contraction and cardiac activity.	
		• Blood vessels and blood pressure:- the blood vessels of skin and mucous	



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memorane are constructed. Adrenatine unates blood vesser	is of skeletal muscles	Uy	
acting on B-receptors. The net result is thus decrease in	peripheral resistance.	It	
shows biphasic response on blood pressure in moderate do	se		
Smooth muscles: -			
a) Bronchial smooth muscles: - adrenaline is a powerful br	onchodilator		
■ b) Smooth muscles of GIT: - The muscles of GIT are relax	ed and peristaltic		
movement becomes sluggish.			
🖆 c) Central Nervous system: - Therapeutic doses of adrenali	ne may give rise to		
tremors, restlessness, palpitation and apprehension			
Metabolism: - it produces hyperglycaemia by accelerating	glycogenolysis in the		
liver.			
Antiallergic action: - adrenaline is a physiological antag	gonist of histamine a	nd	
counters the bronchoconstriction and hypotension of	anaphylactic shock.	If	
combined with local anaesthetic, prolongs its action.			
🖆 Action on eye: Adrenaline causes mydriasis.			
ii) Classify Parasympathomimetics with examples.			
A)Ester of Choline-Methacholine, Carbachol, Acetylcholine.			
B) Cholinomimetic alkoloids- Piolcarpine, Muscarine.			
C) Cholinesterase inhibitor-			
a) Reversible :-Neostigmine, physostigmine, pyridostigmin	e.		
b) Irreversible:- Organophosphorus compounds, (DFP, mal	athion, parathion)		



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write treatment of peptic ulcer by classifying	the drugs used and mention	3111	
mechanism of action of each category. Classif	ication:	Treatm 2M For Me	
A. Drugs that neutralize acid :			
Antacids- sodium bicarbonate ,Aluminiur	n hydroxide.		
<b>B. Drugs that reduce acid secretion:</b>			
H2 receptor antagonist - Eg. Cimetidine, R	anitidine.		
Proton pump Inhibitor : Eg. Omeprazole,pa	antoprazole.		
Anticholinergics: E.g. Pirenzepine.			
Prostaglandins: Eg. Misoprostol.			
C. Drugs acting on ulcer: Sucralfate, Colloid	al bismuth.		
<b>D. Antibacterials</b> :. Eg. Amoxicillin, Clarithron	mycin, Metronidazole.		
OR			
I.Reduction of gastric acid secretion.			
• H2-antihistamines: Cimetidine, R	anitidine, and Famotidine.		
<ul> <li>Proton pump inhibitors: Omepraz</li> </ul>	cole, Lansoprazole, and Pantoprazole.		
• Anticholinergics: Pirenzepine, Ox	kyphenonium.		
<ul> <li>Prostaglandin analogues: Misopro</li> </ul>	ostol.		
II. Neutralization of gastric acid (Antacids).			
• Systemic: NaHCO3 and Sodium	citrate.		
• Nonsystemic: Mg(OH)2, CaCO3,	, Aluminium hydroxide gel and		
Magnesium trisilicate.			
III. Ulcer protectives: Sucralfate, Colloida	l Bismuth Subcitrate (CBS).		
IV. Anti-Helicobacter pylori drugs: Amoxicill	in, Clarithromycin, Metronidazole,		
Tinidazole, and Tetracycline.			
H2 antihistamines-These drugs block hi	stamine- induced gastric acid secretions.		
Inhibit acid production by reversible com	peting with histamine for binding with		



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	H2 receptor on the basolateral membrane of parietal cells.			
	Proton pump inhibitors (PPIs) block the gastric H,K-ATF acid secretion.	Pase, inhibiting gast	ric	
5	Antacids are basic substances which neutralise gastric acid gastric contents. Peptic acidity is reduced if pH rises above	1		
	<ul> <li>Ulcer protectives like sucralfate does not inhibit gastric ac with existing stomach acid to form a thick coating that cover ulcer, protecting the open area from further damage.</li> <li>Anti H pylori drugs-drugs like Amoxicillin, an antibiotic a bacterial cell wall synthesis, leading to cell death.</li> </ul>	ers the surface of an	ı	

Q. No	Su b Q. N.	Answer	Marking Scheme
2		Answer any <u>TEN</u> of the following:	30M
2	a	Describe the factors influencing absorption of drugs. Explain any one factor.	Enlist 1.5M
		Note : Question might be Enlist and explain anyone. Consider it as Enlist and explain anyone.	Explanat1.5 M
		I) Biological factors/ physiological factors / Intrinsic factors:	
		a) pH of drug and pH of GIT:pH of GIT and Blood may interfere with the absorption of	
		the drug. E.g. Salicylate and barbiturates (acidic drugs) remain in unionized form in the	
		stomach, and in the acidic pH of the stomach, they are rapidly absorbed.Basic drugs like	
		pethidine and ephedrine are only absorbed in the small intestine, as these drugs exist in un-	
		ionised form in an alkaline environment. Unionized drugs are lipid soluble while ionized	
		drugs are water-soluble agents. Hence unionized drugs are better absorbed than ionized	
		drugs.	



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b) Presence of food and other agents. Most drugs are better absorbed in an empty stomach. The presence of food in the stomach dilutes the drug and retards absorption of drugs. E. g. Ampicillin, Aspirin, tetracycline etc. The presence of other drugs in the gut may increase or decrease the absorption of drug by drug-drug interaction e.g. Presence of vitamin C increases the absorption of iron salts. from the gut. presence of calcium, magnesium decreases the absorption of tetracycline by forming a poorly absorbed chelate complex. Liquid paraffin reduces the absorption of fat-soluble vitamins like A, D, E and K.

c) Surface area :The larger the surface area the better the absorption. Drugs are better absorbed in the intestine than in the stomach because of the large surface area of the intestine. Gastrointestinal surgery reduces the absorption of drugs because of decrease in surface area.

**d)Functional integrity of gastrointestinal tract**/ **Gastrointestinal transit time:** The motility of the stomach is important to the rate at which orally administered drug is passed on to the intestine. Delayed gastric emptying reduces the absorption of orally administered aspirin. Food also affects gastric emptying time. Absorption of amoxicillin, ampicillin and cephalexin is reduced in the presence of food. This is due to enhanced gastric emptying. An increase in peristalsis reduces the residence time of the drug in GIT, So reduces absorption.

**e)Blood flow in GIT:**The increase of blood flow in the gut due to vasodilation increases the absorption of drugs. Decrease of blood flow in the gut due to vasoconstriction as in haemorrhagic shock decreases absorption of drugs.

## II) Pharmaceutical factors/ Extrinsic factors:

**a) Physical state of the drug**: Liquids are better absorbed than solid medications. Aqueous solutions are more quickly absorbed than oily solutions. Soluble drugs are more easily absorbed than insoluble ones.

**b) Particle size:** Smaller the particle size, the better the absorption since it provides a greater surface area for absorption. Small particle size is useful in the absorption of corticosteroids and antibiotics like Chloramphenicol, Griseofulvin and oral anticoagulants.



•		$\mathbf{W} \mathbf{H} \mathbf{H} \mathbf{E} \mathbf{K} = \mathbf{Z} \mathbf{U} \mathbf{Z} \mathbf{U} \mathbf{Z} \mathbf{U} \mathbf{U} \mathbf{H} \mathbf{U} \mathbf{H} \mathbf{U} \mathbf{H} \mathbf{U} \mathbf{U} \mathbf{U}$		
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		c) Formulation: The method of formulation influences absorption. Substances like lactose, sucrose, starch, calcium phosphate, and calcium lactate are used as inert diluents in		
		formulating tablets and powders. These are the agents that may interfere with the active		
		drug and affect its absorption.		
		Eg. calcium and magnesium ions reduce the absorption of tetracycline.		
		d) Disintegration time of drug: It is the time taken for the solid dosage form of a drug to		
		disintegrate into fine particles in the gut completely. It depends on the type of drug and		
		excipients used in it. If the disintegration time is longer, the absorption of the drug is delayed.		
		e) Dissolution time of drug : It is the time taken for a solid dosage form of a drug to go		
		into the solution in the gut after it has been disintegrated. The solution is absorbed faster		
		than the solid dosage form. If dissolution time is longer, the absorption of the drug is		
		delayed.		
2	b	Classify NSAIDS with examples.	3 M	
		A) Non selective COX inhibitors (traditional NSAIDS)		
		1.Salicylates and analogs		
		eg. Aspirin		
		2. Propionic acid derivatives:		
		eg. Ibuprofen, Naproxen, Ketoprofen, Flurbiprofen.		
		3. Anthranilic acid derivatives:		
		eg. Mefenamic acid.		
		4. Aryl-acetic acid derivatives:		
		eg. Diclofenac, Aceclofenac.		
		5. Oxicam derivatives:		
		eg. Piroxicam, Tenoxicam.		
		6. Pyrrolo-pyrrole derivative:		
	1			



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	eg. Retorolac.			
	7. Indole derivative:			
	eg. Indomethacin.			
	8. Pyrazolone derivatives:			
	eg. Phenylbutazone, oxyphenbutazone.			
	B) Preferential COX-2 inhibitors			
	Eg. Nimesulide, Meloxicam, Nabumetone			
	C) Selective COX-2 inhibitors			
	Eg. Celecoxib, Etoricoxib, Parecoxib			
	D) Analgesic-antipyretics with poor anti-inflammatory action			
	1. Para aminophenol			
	derivative:-Eg. Paracetamol			
	2. Pyrazolone derivatives:-			
	Eg. Propyphenazone.			
	3. Benzoxazocine			
	derivative:-Eg. Nefopam.			
	<u>OR</u>			
	1) Salicylates			
	Eg. Aspirin, Sodium salicylate			
	2) Para aminophenol derivatives			
	eg. Paracetamol, Phenacetin			
	3) Indole acetic acid derivatives			



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eg. Indomethacin			
4) Anthranilic acid derivatives			
eg. Mefenamic acid			
5) Propionic acid derivatives			
eg Ibuprofen, Naproxen			
6) Oxicam derivatives			
eg Piroxicam			
7) Pyrazolone derivatives			
eg Phenylbutazone, Oxyphenbutazone			
8) Phenyl acetic acid derivatives			
eg Diclofenac			
9) COX 2 inhibitors			
eg. Rofecoxib			
10) Miscellaneous			
eg. Nimesulide, Metamizol etc.			
c Classify antifungal drugs with examples. 1. Antibiotics:		JIVI LAC	1
Ex. Amphotericin B, Nystatin, Hamycin, Griseofulvin.			
2. Antimetabolite: Ex. Flucytosine (5-FC).			
3. Azoles:			
(a) Imidazoles:			
Topical: Ex.Luliconazole, Clotrimazole, Miconazole, Oxico	onazole.		
Systemic: Ex. Ketoconazole.			
(b) Triazoles (systemic): Ex. Fluconazole, Itraconazole, Vorico	onazole, Posaconazole.		
4. Allylamine: Terbinafine.			
5. Other topical agents: Tolnaftate, Undecylenic acid, Benzoid	e acid.		



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2 u	Define and classify antihypertensive drugs with examples.	1WI Del. 2M Class	si.
	Antihypertensive drugs are the agents used in treatment of hypertension or abnormal elevation in blood pressure.		
	Classification (According to site of action):		
	1. Centrally acting Drugs: Clonidine, Methyldopa		
	2. Drugs acting on autonomic ganglia: Hexamethonium		
	3. Drugs acting on post ganglionic sympathetic nerve		
	endings a) Adrenergic neuron blockers; Guanethidine		
	b) Catecholamine depletors: Reserpine		
	4. Drugs acting on adrenergic receptors:		
	a) Alpha adrenergic blockers: Phentolamine		
	b) Beta adrenergic blockers: Propranolol		
	5. Vasodilators: Hydralazine		
	6. Drugs acting reflexly by stimulating baroreceptors: Veratrum		
	7. Oral Diuretics: Thiazides, Frusemide, spironolactone, amiloride etc		
	8. Calcium Channel Blockers: Nifedipine, Amlodipine, Felodipine		
	9. Drugs acting on renin angiotensin system:		
	a) ACE inhibitors: Enalapril, Ramipril		
	b) Angiotensin Receptor Blockers: Losartan, Telmisartan		
	10.Miscellaneous: MAO inhibitors (Pargyline).		



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2	Subje e f	<ul> <li>What is haematinics? Give uses of anticoagulant drugs.</li> <li>Haematinics: Are the drugs which when administered favour erythropoiesis i.e. synthesis of red blood cells and increase the oxygen carrying capacity of the blood.</li> <li>Eg: cyanocobalamin, folic acid, iron etc.</li> <li>Uses of anticoagulant: <ul> <li>A. In treatment and secondary prophylaxis of:</li> <li>(a) Deep vein thrombosis (DVT).</li> <li>(b) Pulmonary embolism (PE).</li> <li>B. Prophylaxis of thromboembolism (e.g., stroke) in patients with the following:</li> <li>(a) Valvular atrial fibrillation and non-valvular atrial fibrillation.</li> <li>(b) Heart valve replacement.</li> <li>(c) Heart failure.</li> <li>(d) Myocardial ischemia.</li> </ul> </li> <li>Explain mechanism of action and side effects of sulphonamides.</li> <li>Mechanism - Many microorganisms require Para amino benzoic acid (PABA) for the synthesis of folic acid. PABA &amp; sulphonamides are similar in chemical structure such that bacteria are not able to differentiate them. There is also competition between these two substances for the same receptor site. Bacteria take up sulphonamide instead of PABA &amp; inhibit formation of folic acid which is required for the bacterial growth and has bacteriostatic action.</li> </ul>	20221 1M defin with example Uses 2M MOA 1M Side effects	2 M
		<b>Side effects</b> - crystalluria, haematuria, agranulocytosis, renal impairment, allergic reactions		
2	g	<ul> <li>What are diuretics? Explain thiazides as diuretics.</li> <li>These are the pharmacological agents which when administered, increase rate of formation of urine as well as excretion of urine.</li> <li>Examples: Mannitol, Theophylline, Acetazolamide, Furosemide, Spironolactone, Chlorothiazide etc.</li> <li>Explanation:</li> </ul>	Intro. 1 Explain	



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		and chloride ions with water in the distal tubule. This causes excr chloride ion, water and little potassium ions also and produces diuretion	etion of sodium			
2	h	<ul> <li>Write a note on Antithyroid drugs. Antithyroid drugs</li> <li>These are the pharmacological agents which are used in the treatment</li> <li>Classification <ol> <li>Thiourea derivative</li> <li>Propylthiouracil, Carbimazole</li> </ol> </li> <li>Popultion inhibitors</li> <li>Eg. Potassium thiocyanate, Potassium perchlorate</li> <li>Jodides</li> <li>Potassium iodide, Sodium iodide</li> <li>Radioactive Iodine</li> <li>eg.1311 (iodine isotopes).</li> </ul>	of hyperthyroidi:	sm.	3M	
2	i	<ul> <li>Explain triple response of Histamine.</li> <li>TRIPLE RESPONSE: When histamine is applied locally or injected skin histamine produces a typical response known as "triple resp characterized by three distinguish sign:</li> <li>i. Flush- it is redness at the site of application because of hyperaemia</li> <li>ii. Flare- Patch formation in the vicinity of 1.5 cm of flush occurs due this is called flare.</li> <li>iii. Wheal- around 1.5cm of flare permeation of fluid occurs, raising th called as wheal (swelling formation).</li> </ul>	onse" which is to vasodilation of	ı &	M	
2	j	<b>Define Hypnotics. Classify with suitable examples.</b> <b>Hypnotics</b> are the pharmacological agents which act on CNS, resembling natural sleep Eg. Phenobarbitone, Amylobarbitone etc.	and produce s	C	Def. 1M, Classi. 21	М



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	Classification-			
	I) Barbiturates			
	a) Long acting barbiturates e.g. Phenobarbitone			
	b) Intermediate acting barbiturates e.g. Cyclobarbitone, Amylobarbito	one		
	c) Short acting barbiturates e. g. Hexobarbitone, Secobarbitone			
	d) Ultra short acting barbiturates e. g. Thiopentone, Methohexitone			
	II) Non barbiturates			
	a) Benzodiazepine e.g. Diazepam, Chlordiazepoxide			
	b) Alcohols e.g. Chloral hydrate, Ethyl alcohol			
	c) Aldehydes e. g. Paraldehyde			
	d) Miscellaneous e.g. Hyoscine ,Meprobamate			
	OR			
	I) Derivatives of urea:			
	i) Barbiturates			
	a) Long acting barbiturates (duration 8hr or more)			
	e.g. Phenobarbitone.			
	b)Intermediate acting barbiturates ( duration 4hrs)			
	e.g. Amylobarbitone.			
	c) Short acting barbiturates (duration less than			
	4hrs) e. g. Hexobarbitone, Secobarbitone.			
	d) Ultra short acting barbiturates( These agents give instanta duration of action is less than 1hr.)	neous action and the	ne	



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		e. g. Thiopentone, Methohexitone.		
		ii) Related diureides: eg. Glutethimide, Methyprylon.		
		II) Benzodiazepines		
		e.g. Diazepam,Chlordiazepoxide.		
		III) Alcohols		
		e.g. Chloral hydrate, Ethyl alcohol.		
		IV) Aldehydes		
		e. g. Paraldehyde.		
		V) Acetylated carbinols		
		e.g. Ethomidate, Ethchlorvynol.		
		VI) Miscellaneous		
		e.g. Hyoscine, Meprobamate.		
		VII) Inorganic ions e.g. Bromide		
2	k	Write use and one example of: i) Expectorant-	1.5M e	_
		Use: These are the pharmacological agents which are used to remove mucus from the	one exai	nple)
		respiratory tract and are used in productive cough associated with bronchitis, Asthma,		
		Emphysema		
		Eg. Guaifenesin, Potassium iodide, Vasaka, Ammonium chloride.		
		ii) Bronchodilator-		
		Use: It is used in the treatment of bronchial asthma.		
		Used in bronchospasm		
		Used in COPD (Chronic pulmonary obstructive disease)		
		Eg. Salbutamol, Terbutaline, Adrenaline, Ephedrine, Isoprenaline, Theophylline,		
		Aminophylline etc.		
3		Attempt ALL of the following	1M Each	1



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3	a	Drugs administered through the following route are most likely to be subjected to first pass metabolism.	1M	
		i) Oral ii) Sublingual iii) Subcutaneous iv) Rectal Ans:- Oral		
3	b	Give two examples of neuromuscular blocking agents.	1M	
		<ul> <li>Ans: D.tubocurarine,Pancuronium,Gallamine,Alcuronium,Succinylcholine, Vecuronium,</li> <li>Mivacurium, Atracurium etc.</li> <li>(Any other correct examples can be considered).</li> </ul>		
3	c	The B1 receptor are located in	1M	
		i) Heart		
		ii) Lungs		
		iii) Kidney		
		iv) Adrenal gland		
		Ans: i) Heart , iii) Kidney (Any one of these two answers can be considered).		
3	d	Local anaesthetics produce :	1M	
		i) Analgesia, amnesia, loss of consciousness		
		ii) Blocking pain sensation without loss of consciousness		
		iii) Alleviation of anxiety and pain with an altered level of consciousness		
		iv) A stupor or somnolent state.		
3	e	Ans: ii) Blocking pain sensation without loss of consciousness. Give two examples of mydriatics.	1M	
		Ans: Phenylephrine, Adrenaline, Atropine, Homatropine, Cyclopentolate, Tropicamide.		
		(Any other correct examples can be considered).		
3	f	If the pressure inside eye is higher than normal person,	1M	
		i) May Have cataract.		
		ii) May be at risk of glaucoma.		



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		III) Neeu eye glasses	
		iv) Have infection	
		Ans: May be at risk of glaucoma.	
3	g	Give two examples of drugs used as nootropic agents.	1M
		Ans: Piracetam, Pyritinol ,Modafinil, Caffeine, L-Theanine, Panax Ginseng,	
		Gingko biloba, Brahmi.	
		(Any other correct examples can be considered).	
	h	Phenytoin is used in the treatment of	1M
3		Ans: Epilepsy, Cardiac arrhythmia	
		(anyone correct Ans, can be considered)	
3	i	Statins are used in atherosclerosis. state true or false.	1M
		Ans: True	
3	i		1M
U	J	Give an example of one drug used in the treatment of manic depressive illness.	
		Ans: Lithium, Valproic acid, Divalproex sodium, Carbamazepine, Lamotrigine	
		(Any other correct example can be considered)	
3	k	Give an example of one thrombolytic agent.	1M
		Ans:Streptokinase,Alteplase,Reteplase.Tenecteplase.Urokinase.Prourokinase.	
		Anistreplase.	
		(Any other correct example can be considered)	
	1	Define laxative	1M
3		Ans: These are the agents which when administered relieve constipation without gripping	
		pain and loss of water. OR	
		Laxatives are substances that loosen stools and increase bowel movements.	
3	m	Antidiuretics are the drugs which have following action	1M
		i) Increase urine output	
		ii) Decrease urine output	
		iii) Stop urine formation	
		iv) Cause drowsiness	
		Ans: ii) Decrease urine output	
3	n	is caused by deficiency of Vitamin D.	1M



	Subj	ect Name: Pharmacology Subject Code: 2	0221
		Alls: Alexels of Osteolilaiacia (ally one call be considered)	
3	0	The main hormones secreted by the thyroid gland is	1M
3	p	<ul><li>Ans: Thyroxine or Tetraiodothyronine (T4) and Triiodothyronine (T3).</li><li>Give any two examples of anti-histamines.</li></ul>	1M
-	ſ	Ans:Diphenhydramine,Cetirizine,Chlorpheniramine,Cyclizine,Dimenhydrinate	
		Doxylamine, Hydroxyzine, Meclizine.	
		(Any other correct examples can be considered) Ranitidine, famotidine etc.	
3	q	Mention two drugs used in the treatment of cancer.	1M
		Ans: Nitrogen Mustards, Cyclophosphamide, Methotrexate, Vinblastine, Vincristine,	
		Uracil Mustards, Fluorouracil	
		(Any other correct examples can be considered)	
3	r	Name one anti TB drugs used in resistant TB.	1M
		Ans: Bedaquiline, Pretomanid, Moxifloxacin ,PAS, Clycloserine, Ethionamide	
3	s	(Any other correct example can be considered)Give an example of any one biological drug.	1M
5	3	Ans:	1 IVI
		Asparginase, Abatacept, Belatacept, Cetuximab, Daclizumab, Elotuzumab, Golimumab, Pegasp	
		argase.	
		(Any other correct example can be considered)	
3	t	Define biologicals.           Biologicals are those classes of medicines which are grown and then purified from large-	1M
		scale cell cultures of bacteria or yeast, or plant or animal cells.	
		OR	
		Biologicals are a diverse group of medicines which includes vaccines, growth factors,	
		immune modulators, monoclonal antibodies, as well as products derived from human blood	
		and plasma.	