



Code No.: H-8150/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy (PCI) VI-Semester (Backlog) Examination, February / March 2026**

**Subject: Medicinal Chemistry - III**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. Write the structure and uses of Amoxicillin
2. Write the structure and uses of Chloramphenicol.
3. Differentiate Penam & Cephem with structures.
4. Write the mechanism of action of Macrolide antibiotics.
5. Write the structure of Chloroquine & Ethambutol?
6. Define and classify Prodrugs and give two examples.
7. Write the mechanism of action of folate reductase inhibitors.
8. Define and Write the applications of Combinatorial chemistry
9. Write the mechanism of action of Aminoglycosides?
10. Give the synthesis and uses of Dapsone.

**PART B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. a) Write the structures and uses of three drugs each from Cephalosporins and Pencillins  
b) Discuss in detail about SAR, Mechanism of action and side effects of Tetracyclins
12. What are antifungal agents, give the classification and Discuss in detail about synthetic anti fungal agents.
13. a) Classify Sulphonamides with structural examples and write the SAR  
b) Write the synthesis and mechanism of action of Sulfacetamide and Sulfamethoxazole

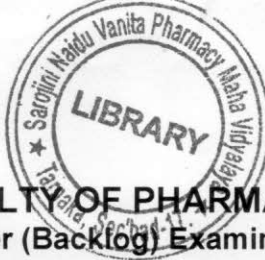
**PART C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Write in detail about beta lactamase inhibitors.
15. Write in detail about quinolones.
16. What is docking and write in detail about various docking techniques.
17. Give the structure Mechanism of Action and uses of a) Itraconazole b) Praziquantal
18. Write the synthesis and uses of Nitrofurantoin and Ciprofloxacin.
19. Explain various physicochemical parameters used in QSAR studies.
20. Define combinatorial chemistry and write in detail about solid phase and solution phase synthesis.
21. Write the structures of any five synthetic anti tubercular agents and write their mechanism of action.
22. What are anthelmintics? Write the synthesis, Mechanism of action and uses of Mebendazole.

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Code No.: H-8151/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy (PCI) VI-Semester (Backlog) Examination, February/March 2026**

**Subject: Pharmacology-III**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. What are nasal decongestants? Give examples.
2. Classify antiemetics with examples.
3. What are superinfections and give any two examples?
4. What are the therapeutic uses of sulfonamides?
5. Write the MOA and therapeutic uses of metronidazole.
6. Write the MOA and therapeutic uses of dapsone.
7. Classify immunosuppressants.
8. Mention any two organisms causing UTIs and STDs.
9. Define circadian rhythm and write any two applications of chronotherapy.
10. What are the clinical features of organophosphate poisoning.

**PART B**

**Note: Answer any two questions.**

**(2 x 10= 20 Marks)**

11. Classify anti-asthmatic drugs. Discuss about bronchodilators.
12. Classify anticancer drugs and describe the general toxicity of cytotoxic drugs.
13. Discuss about general principles of management of poisoning.

**PART C**

**Note: Answer any seven questions.**

**(7 x 5= 35 Marks)**

14. Write notes on treatment of constipation.
15. Write the MOA, adverse drug reactions and therapeutic uses of aminoglycosides.
16. Discuss the pharmacology of cotrimoxazole.
17. Discuss about antimicrobial resistance.
18. Classify antifungal drugs. Write the pharmacology of ketoconazole.
19. Classify anti-tubercular drugs. Write the pharmacology of isoniazid.
20. Classify anti-retroviral drugs. Discuss about protease inhibitors.
21. Write notes on protein drugs.
22. Write notes on acute and chronic toxicity studies.

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Code No: H-8152/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy (PCI) VI - Semester (Backlog) Examination, February/March 2026**

**Subject: Herbal Drug Technology**

**Time: 3 Hours**

**Max. Marks: 75**

**PART - A**

**Note: Answer All the Questions.**

**(10 x 2 = 20 Marks)**

1. Define "Herbal medicine" and "herbal drug preparation" as per WHO.
2. Explain Bio-prospecting emphasizing on its merits and demerits.
3. Fenugreek is a health food. Justify.
4. What are the principles of Homeopathic system of medicine.
5. Write about any two natural sweeteners.
6. Write applications of Bacillus thuringensis (Bt).
7. What are the objectives of schedule Z.
8. What are the different types of pheromones.
9. Write about guar gum and saffron.
10. Write about CIMAP and CSIR.

**PART - B**

**Note: Answer any Two Questions.**

**(2 x 10 = 20 Marks)**

11. Explain ICH guidelines for quality assessment of herbal drugs.
12. What is TKDL. Explain its role in prevention of wrongful patenting of natural products.
13. Discuss about processing techniques of herbal raw materials.

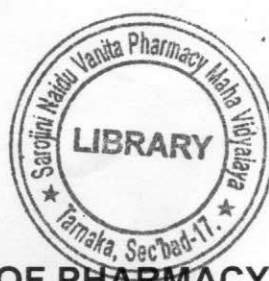
**PART - C**

**Note: Answer any Seven questions**

**(7 X 5 = 35 Marks)**

14. Write a note on organic farming.
15. Write a note on nutraceuticals in prevention and management CVS disorders.
16. Explain herb-drug interactions of Hypercium and Garlic.
17. Elaborate on the patenting aspects of curcuma.
18. Write a note on antioxidants in herbal cosmetics.
19. Explain the methods of preparation and standardization of Bhasmas.
20. Give the composition and functions of ASU DTAB.
21. Write about 'Farmers right' and Breeders' right.
22. Write a note on regulations of manufacture of ASU drugs in India.

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Code No: H-8154/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy (PCI) VI - Semester (Backlog) Examination, Feb/March 2026**

**Subject: Pharmaceutical Biotechnology**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer All the Questions.**

**(10 x 2 = 20 Marks)**

1. Define Biosensor and Immobilization.
2. Write the role of DNA ligase and polymerase in DNA replication.
3. List the Linkers used in gene manipulation.
4. What are the ideal properties of Vector?
5. Write the mechanism of conversion of toxin to toxoids.
6. Write the uses of human fibrinogen.
7. Define Mutation. Explain about point mutations.
8. Write about spargers used in a fermentor.
9. What are restriction endonucleases.
10. What are plasmids? Explain how plasmids offer extra characters to the host.

**PART – B**

**Note: Answer any Two Questions.**

**(2 x 10 = 20 Marks)**

11. Explain the methods of horizontal gene transfer from one bacterium to another bacterium.
12. Explain the general methods of vaccine preparation.
13. What are plasma substitutes and ideal properties of plasma substitutes? Explain the production of plasma substitutes.

**PART – C**

**Note: Answer any Seven questions**

**(7 X 5 = 35 Marks)**

14. Give an account of collection, processing & storage of whole human blood.
15. Write about Structure and function of MHC class I and II.
16. Explain the genetic organization of Eukaryotes.
17. Explain about western blotting technique.
18. Explain the production of Insulin by fermentation technology.
19. Write about polymerase chain reaction and its applications.
20. Write about the production enzyme catalase.
21. Explain type I and type II hypersensitivity reactions.
22. What are Immunoglobulins? Name the Immunoglobulin that can cross the placental barrier. Write about IgG and IgE antibodies.

**\*\*\*\*\*End\*\*\*\*\***



Code No. H-8153/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy (PCI) VI - Semester (Backlog) Examination, Feb/March 2026**  
**Subject: Biopharmaceutics and Pharmacokinetics**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. Define Noyes and Whitney equation and its application.
2. Define ADME.
3. Differentiate Phase I and Phase II metabolic pathways.
4. Define Absolute bioavailability and Relative bioavailability
5. Explain Apparent volume of distribution.
6. What is Flip-Flop phenomenon and how it is useful in method of residual.
7. What is protein binding. How it effects kinetics.
8. Write a note on excretion of drugs through skin.
9. Write a note on enterohepatic circulation.
10. Define Bioequivalence.

**PART – B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. Discuss about factors influencing drug absorption through GIT.
12. Derive mathematical equations to calculate pharmacokinetic parameters for a drug administered by IV infusion, given blood data, assuming drug follows one compartment open model.
13. Derive Michaelis-Menten equation and how do you estimate Km and Vmax.

**PART – C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Discuss about in-vitro-in-vivo (IVIVC) correlations.
15. Write a note on Carrier mediated transport.
16. Explain factors affecting renal excretion of drugs.
17. Explain methods of adjustment of dose and dosage regimen in patients with renal and hepatic failure systems.
18. Write in detail about pH partition hypothesis and its limitation.
19. Describe about physiological barriers to the distribution of drugs.
20. Write methods to enhance the bioavailability of poorly soluble drugs?
21. Describe factors causing non-linearity.
22. Explain the various methods for assessment of bioavailability.

**\*\*\*\*\*End\*\*\*\*\***



Code No. H-8153/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy (PCI) VI - Semester (Backlog) Examination, Feb/March 2026**  
**Subject: Biopharmaceutics and Pharmacokinetics**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. Define Noyes and Whitney equation and its application.
2. Define ADME.
3. Differentiate Phase I and Phase II metabolic pathways.
4. Define Absolute bioavailability and Relative bioavailability
5. Explain Apparent volume of distribution.
6. What is Flip-Flop phenomenon and how it is useful in method of residual.
7. What is protein binding. How it effects kinetics.
8. Write a note on excretion of drugs through skin.
9. Write a note on enterohepatic circulation.
10. Define Bioequivalence.

**PART – B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. Discuss about factors influencing drug absorption through GIT.
12. Derive mathematical equations to calculate pharmacokinetic parameters for a drug administered by IV infusion, given blood data, assuming drug follows one compartment open model.
13. Derive Michaelis-Menten equation and how do you estimate Km and Vmax.

**PART – C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Discuss about in-vitro-in-vivo (IVIVC) correlations.
15. Write a note on Carrier mediated transport.
16. Explain factors affecting renal excretion of drugs.
17. Explain methods of adjustment of dose and dosage regimen in patients with renal and hepatic failure systems.
18. Write in detail about pH partition hypothesis and its limitation.
19. Describe about physiological barriers to the distribution of drugs.
20. Write methods to enhance the bioavailability of poorly soluble drugs?
21. Describe factors causing non-linearity.
22. Explain the various methods for assessment of bioavailability.

\*\*\*\*\*End\*\*\*\*\*



Code No: H-8155/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy (PCI) VI - Semester (Backlog) Examination, Feb/March 2026**

**Subject: Quality Assurance**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer All the Questions.**

**(10 x 2 = 20 Marks)**

1. Define the terms QA & GMP
2. Mention different ICH guidelines.
3. Why hygiene maintenance is needed in Pharma Industry?
4. Name different personnel records in a pharmaceutical industry.
5. Name few equipment used in pharmaceutical industry.
6. What are the reports issued by QC department?
7. What are to be mentioned in a complaint to a pharma industry?
8. What is waste disposal?
9. Write the significance of validation.
10. Mention different distribution records.

**PART – B**

**Note: Answer any Two Questions.**

**(2 x 10 = 20 Marks)**

11. Discuss about different elements of TQM.
12. Explain about the premises of a Pharmaceutical Industry.
13. What is Calibration? Write its significance and explain calibration of pH meter.

**PART – C**

**Note: Answer any Seven questions**

**(7 X 5 = 35 Marks)**

14. Write informative notes on QbD.
15. Give the procedure for NABL accreditation.
16. Write notes on maintenance of stores for raw materials.
17. Write quality control tests for rubber closures.
18. Explain the contents of a SOP.
19. Discuss on recalling of products.
20. What is validation. Write about types of validation.
21. Discuss on qualification of UV-Visible spectrophotometer.
22. Give informative notes on good warehousing practices.

**\*\*\*\*\*End\*\*\*\*\***

**FACULTY OF PHARMACY**  
**B. Pharmacy (PCI) VI - Semester (Main & Backlog) Examination, September 2025**  
**Subject: Medical Chemistry - III**

Time: 3 Hours

Max. Marks: 75

**PART - A**

Note: Answer all the questions.

(10 x 2 = 20 Marks)

1. Define and classify  $\beta$ -Lactam antibiotics?
2. Write the mechanism of action of Aminoglycosides?
3. What are  $\beta$ -Lactamase inhibitors, give examples?
4. Define Prodrugs?
5. Write the structure and uses of Chloroquine?
6. Write the mechanism of action of monobactams?
7. What are folate reductase inhibitors, give few examples?
8. Give the structure and uses of Para amino salicylic acid?
9. Write the applications of combinatorial chemistry?
10. Define QSAR?

**PART - B**

Note: Answer any two questions

(2 x 10 = 20 Marks)

11. (a) Define antibiotics? Write the classification and SAR of Penicillins?  
(b) Write the synthesis and uses of Chloramphenicol?
12. (a) Give the classification of anti-tubercular agents with examples?  
(b) Write the SAR of quinolones?
13. (a) Write the classification of Anti-fungal agents?  
(b) Give the synthesis and mode of action of Sulfacetamide?

**PART - C**

Note: Answer any seven questions.

(7 x 5 = 35 Marks)

14. Give a note on epimerization of Tetracyclines?
15. Write a note on chemical degradation of cephalosporins?
16. Write the synthesis, mode of action and uses of Metronidazole?
17. Give the classification of Antimalarial agents with examples (write any one structure for each class)?
18. Write the synthesis MOA and uses of Isoniazid?
19. What are anthelmintics? Write the synthesis of Mebendazole?
20. Give a note on combinatorial chemistry?
21. Write the SAR and general synthesis of Sulphonamides?
22. Write the structure, synthesis and uses of Dapsone?

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**FACULTY OF PHARMACY**  
**B. Pharm (PCI) VI - Semester (Main & Backlog) Examination, September 2025**  
**Subject: Pharmacology-III**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. Define Asthma and COPD.
2. What are Biosimilars? Give examples.
3. Write the mechanism of action of Co-trimoxazole.
4. Write a note on lead poisoning.
5. What are common adverse drug effects of chemotherapeutic agents?
6. Differentiate between tolerance and Resistance.
7. Write the mechanism of action of Chloramphenicol.
8. Classify antidiarrhoeals with examples.
9. What is MRSA?
10. What is circadian rhythm? Mention its importance in pharmacotherapy.

**PART – B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. Classify Cephalosporins with examples. Write the mechanism of action, pharmacokinetics, adverse effects and uses of cephalosporins.
12. Write in detail about the pharmacological therapy of COPD.
13. Classify broad spectrum antibiotics with examples. Write in detail about mechanism, adverse drug reactions and uses of penicillins.

**PART – C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Classify anti-ulcer drugs. Write a note on proton pump inhibitors.
15. Write the mechanism of action and uses of sulphonamides.
16. Write the prophylaxis of malaria.
17. Classify anti tubercular drugs with examples and explain the pharmacology of the drugs.
18. Brief up the mechanisms of antifungal drugs.
19. Write the toxicity of quinolones.
20. What are immunostimulants? Write a note on immunoglobulins and their uses.
21. What are cell cycle inhibitors? Explain with examples.
22. Write the mechanism of action and uses of digestants and carminatives.

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**FACULTY OF PHARMACY**

**B. Pharmacy VI - Semester (PCI) (Main & Backlog) Examination, September 2025**

**Subject: Herbal Drug Technology**

**Time: 3 Hours**

**Max. Marks: 75**

**PART - A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. Define Herbal medicinal products.
2. What are bhasma and lehya?
3. Write the side effects and interaction of pepper as health food.
4. Define breeder's right.
5. Name any two natural sweeteners with biological source.
6. Mention the evaluation of herbal syrups.
7. What are the source, active constituents and uses of alfalfa?
8. What are ideal requirements of bhasma?
9. Significance of preparation of herbarium.
10. Mention health benefits of spirulina.

**PART - B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. Explain the good agriculture practices in the cultivation of medicinal plants.
12. Explain in detail about the scope and types of Nutraceutical products available in the market.
13. Explain in detail the concept of Ayurvedic system of medicine.

**PART - C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. What are Nutraceuticals? Discuss on the present market scenario and scope of Nutraceuticals.
15. Explain the method of preparation of Aristas.
16. Discuss the objective and components of GMP.
17. Describe the role of honey as health food.
18. Give the side effects and interactions of Ginkgo Biloba.
19. Define herb. Explain the method of processing of Herbal raw materials.
20. Write a short note on binders and diluents used as herbal excipients.
21. Describe the role of herbs in dental care.
22. Discuss in brief on "Sodhana" process.

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## FACULTY OF PHARMACY

B. Pharmacy (PCI) VI - Semester (Main & Backlog) Examination, September 2025  
Subject: Biopharmaceutics and Pharmacokinetics

Time: 3 Hours

Max. Marks: 75

## PART – A

Note: Answer all the questions.

(10 x 2 = 20 Marks)

1. Write Henderson Hasselbach equation and explain the terms.
2. Differentiate tissue binding and protein binding of drugs.
3. List markers used in renal clearance.
4. List the official dissolution apparatus according to USP.
5. Explain Apparent volume of distribution.
6. If equation of the curve is  $C=10.e^{-0.69t}$  for a drug administered by IV route and following one compartment open model, then calculate its biological half-life.
7. What is Flip-Flop phenomenon and how it is useful in method of residual.
8. Write formulas for calculating loading dose and maintenance dose.
9. Define non-linear pharmacokinetics.
10. How do you calculate Creatinine Clearance.

## PART – B

Note: Answer any two questions.

(2 x 10 = 20 Marks)

11. Derive mathematical equations to calculate pharmacokinetic parameters for a drug administered by IV infusion, given blood data, assuming drug follows one compartment open model.
12. Discuss about factors affecting protein-drug binding and clinical significance of protein binding of drugs.
13. Derive Michaelis-Menten equation and how do you estimate  $K_m$  and  $V_{max}$ .

## PART – C

Note: Answer any seven questions.

(7 x 5 = 35 Marks)

14. Discuss about Passive absorption and Active absorption of drugs.
15. Describe tissue permeability of drugs.
16. Write a note on kinetics of protein-drug binding.
17. Describe non-renal excretion of drugs.
18. Discuss about *in-vitro-in-vivo* (IVIVC) correlations.
19. Explain methods of adjustment of dose and dosage regimen in patients with hepatic failure.
20. A 650mg I.V. dose of a drug is administered to a 65kg subject, the plasma drug concentration was found to decline biexponentially. The equation that best describes the drug kinetics  $C=67.e^{-14t}+ 33.e^{-3t}$ ; C is in mg/lit Calculate the different volumes of distribution  $V_c, V_p, V_d\beta, V_{darea}, V_{dss}$ .
21. How do you calculate absorption rate constant,  $K_a$  by using Wagner Nelson method.
22. Describe factors causing non-linearity.

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**FACULTY OF PHARMACY**

**B. Pharmacy VI - Semester (PCI) (Main & Backlog) Examination, September 2025**

**Subject: Pharmaceutical Biotechnology**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. Define Immobilization. What are the types of immobilization?
2. What is protein engineering?
3. Write a note on DNA ligases.
4. What are nucleases? Explain the types nucleases.
5. What are vaccines? Enlist types of vaccines.
6. Write the preparation and uses of human Thrombin.
7. What are mutants? Types of mutants.
8. Write about foam control equipment.
9. Write a note on transposons.
10. Write the organisms responsible for the production of Amylases and Lipases.

**PART – B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. Explain the production of insulin by rDNA technology.
12. Explain the production of penicillin by fermentation technology.
13. What is hybridoma technology? Explain the production of monoclonal antibodies.

**PART – C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Explain pBR322 and pUC vectors.
15. Write the applications of genetic Engineering in medicine.
16. Explain the stability of official vaccines.
17. Explain Enzyme linked immunosorbent Assay.
18. What is recombination? Explain general mechanism of recombination.
19. Explain the collection, processing and storage of whole human blood.
20. Explain the preparation of dried human plasma.
21. Explain type II and type III hypersensitivity reactions.
22. Explain the manufacture of dextran. Add a note on methods of size reduction of dextran.

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**FACULTY OF PHARMACY**

**B. Pharmacy VI - Semester (PCI) (Main & Backlog) Examination, September 2025**  
**Subject: Quality Assurance**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. What is the objective of NABL accreditation?
2. What is the purpose of ICH guidelines?
3. Differentiate between primary and secondary packaging materials giving suitable examples.
4. How will you calibrate a pH meter?
5. Explain the concept of QSEM.
6. What are the objectives of sanitation in a drug manufacturing area?
7. Define analytical method validation and list the parameters.
8. Explain the elements of QBD.
9. What are the advantages GMP?
10. What is Quality audit. Write different types of audits.

**PART – B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. Describe the concept of TQM?
12. Define validation and explain the importance of validation.  
What are the different types of validation? Write a note on validation master plan.
13. Explain good warehousing practices (GWP).

**PART – C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Classify the Complaints and write about the evaluation of complaints.
15. Explain the maintenance of sterile areas in pharma industry.
16. Explain the GLP protocol for the conduct of a nonclinical laboratory study.
17. Define and explain the contents of batch formula record.
18. Write about the personnel responsibilities in QA department.
19. Explain the QC tests for glass as a packaging material.
20. What are the benefits of ISO 9000? Add a note on ISO 14000.
21. What are the sources of contamination and mix up in pharmaceutical manufacturing?  
How one can control this type of problems?
22. Explain the location, construction and sanitation of plant.

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Code No: G-13105/PCI

**FACULTY OF PHARMACY**  
**B. Pharmacy (PCI) VI - Semester (Backlog) Examination, March 2025**  
**Subject: Quality Assurance**

**Time: 3 Hours**

**Max. Marks: 75**

**PART - A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. What is TQM?
2. Mention elements of QbD.
3. List out the benefits of ISO accreditation.
4. Name different personnel records in a pharmaceutical industry.
5. Name few equipment used in pharmaceutical industry.
6. Why GLP is necessary?
7. What are to be mentioned in a complaint to a pharma industry?
8. What is quality audit? Write different types of audits?
9. Write the significance of validation.
10. Mention different distribution records?

**PART - B**

**Note: Answer any two questions**

**(2 x 10 = 20 Marks)**

11. Discuss about ICH guidelines.
12. Discuss about different components of master formula.
13. What is Calibration? Write its significance and explain calibration of pH meter.

**PART - C**

**Note: Answer any seven questions**

**(7 x 5 = 35 Marks)**

14. Write the procedure for NABL accreditation.
15. Explain the location, construction and sanitation of plant.
16. Write notes on maintenance of stores for raw materials.
17. Write quality control tests for glass containers.
18. Write notes on general provisions required to maintain GLP.
19. Discuss on recalling and waste disposal in pharma industry.
20. Explain about validation master plan.
21. Discuss on qualification of UV-Visible spectrophotometer.
22. Give informative notes on good warehousing practices.

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Code No: G-13102/PCI

**FACULTY OF PHARMACY**  
**B. Pharmacy (PCI) VI - Semester (Backlog) Examination, March 2025**  
**Subject: Herbal Drug Technology**

**Time: 3 Hours**

**Max. Marks: 75**

**PART - A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. What are the objectives of IPR?
2. Define the term Nutraceuticals.
3. What are antioxidants and give their importance.
4. Write the significance of natural excipient.
5. Define Aristas and Asawas.
6. List the plant based the research institutes in India.
7. What are the advantages of Farmers rights?
8. Give the source and health benefits of Amla
9. Give the source and interactions of Pepper.
10. Write a note on Authentication of plants.

**PART - B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. Explain the Good Agricultural practices in cultivation of Medicinal plants.
12. Elaborate the health benefits and role of Nutraceuticals in management of Diabetes.
13. List the skin care products. Explain the raw materials of herbal origin used in skin care products.

**PART - C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Define the term patent. Give its objectives and criteria for patent award.
15. What are the objectives of Schedule T. Write a note on Infrastructural requirements?
16. Write a note on patenting aspects of Traditional knowledge.
17. Classify the Excipients. Write the advantages and disadvantages of herbal Excipients.
18. Explain the Curcumin case study.
19. Give an informative note on scope and future prospects of Herbal Industry.
20. Write the health benefits of Spirulina and Honey.
21. Describe the role of colorants. Elaborate different colorants of natural origin.
22. Give the sources and side effects and interactions of Hypericum.

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Code No. G-13100/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy VI - Semester (PCI) (Backlog) Examination, March 2025**

**Subject: Medicinal Chemistry - III**

**Time: 3 Hours**

**Max. Marks: 75**

**PART - A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. Write the mechanism of action of Penicillins?
2. Write the structure and uses of Chlorotetracycline?
3. What are Macrolides, give few examples?
4. Give the applications of Prodrugs?
5. Write the structure and uses of Ciprofloxacin?
6. Write the mechanism of action of Acyclovir?
7. What are folate reductase inhibitors, give few examples?
8. Give the structure and uses of metronidazole?
9. Write the applications of combinatorial chemistry?
10. Define Partition coefficient, Hansch analysis?

**PART - B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

1. (a) Define antibiotics? Write the classification and SAR of Cephalosporins?  
(b) Write the synthesis and uses of Chloramphenicol?
2. (a) Give the classification of antiviral agents with examples?  
(b) Write the synthesis and uses of Nitrofurantion?
13. (a) Write the classification and SAR of Sulphonamides?  
(b) Give the synthesis and mode of action of Diethylcarbamazine citrate?

**PART - C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Give a note on epimerization of Tetracyclines?
15. Write a note on  $\beta$ -Lactamase inhibitors?
16. Write the synthesis, mode of action and uses of Chloramphenicol?
17. Give the classification of Antimalarial agents with examples (write any one structure for each class)?
18. Write the synthesis, MOA and uses of Isoniazid?
19. What are antifungal agents? Write the synthesis of Miconazole?
20. Give a note on combinatorial chemistry?
21. Write the classification of Anti-protozoal agents? Write the structure and uses of Tinidazole and Ornidazole?
22. Write the structure, synthesis and uses of Dapsone?

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Code No. G-13104/PCI

**FACULTY OF PHARMACY**

**B. Pharmacy (PCI) VI - Semester (Backlog) Examination, March 2025**  
**Subject: Pharmaceutical Biotechnology**

**Time: 3 Hours**

**Max. Marks: 75**

**PART – A**

**Note: Answer all the questions.**

**(10 x 2 = 20 Marks)**

1. Define biosensors. Write the main components of biosensors.
2. Write a brief note on penicillinase.
3. Write about types of aerators in Fermenter.
4. What is protein engineering?
5. Differentiate exonucleases and endonucleases.
6. Describe the importance linkers and adapters.
7. Differentiate between exotoxins and endotoxins.
8. Define the following:  
a. Cosmid b. Toxoid.
9. Write a note on DNA ligase.
10. What are monoclonal antibodies? Mention its uses.

**PART – B**

**Note: Answer any two questions.**

**(2 x 10 = 20 Marks)**

11. Write the significance of microbial biotransformation, Explain various methods of biotransformation.
12. Discuss the production of Penicillin by fermentation process.
13. Discuss the preparation & purification of Dextran, Plasma substitute.

**PART – C**

**Note: Answer any seven questions.**

**(7 x 5 = 35 Marks)**

14. Discuss about PCR.
15. Describe in brief about cloning vectors.
16. Discuss type II Hypersensitivity and type III Hypersensitivity reactions.
17. Explain basic principles of genetic engineering.
18. What are mutations? Explain the types of mutations.
19. Write about IgG and IgE antibodies.
20. Describe the process of conjugation.
21. Explain the preparation of dried human plasma.
22. Explain the methods for immobilization of enzymes.

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