B. Pharmacy VII - Semester (PCI) (Main & Backlog) Examination, March 2025 Subject: Instrumental Methods of Analysis

Time: 3 Hours Max. Marks: 75

#### PART - A

## Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. State and explain Beer-Lambert equation.
- 2. What are the different types of fundamental modes of vibration in molecules after absorption of IR radiations?
- 3. Define fluorescence and Phosphorescence phenomena.
- 4. Write the principles of Flame photometry technique.
- Define the term Retention time and Resolution in HPLC?
- 6. Write the principles of partition and adsorption chromatography.
- 7. Write the applications of gel permeation chromatography.
- 8. What are the different types of Ion exchange resins used in Ion-exchange chromatography?
- 9. Write the principles of separation in Electrophoresis.
- 10. Write about the different types of columns used in GC.

#### PART - B

#### Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Describe different components of UV spectrophotometer with a neat labelled diagram.
- 12. Explain the principles and experimental details of paper chromatography for Quantitative analysis.
- 13.a) Describe the different sampling preparation techniques in IR spectroscopy.
  - b) Describe different types of detectors used in HPLC instruments.

#### PART - C

#### Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Discuss the different factors influencing intensity of fluorescence of molecules.
- 15. Explain the theoretical principles and applications of affinity chromatography.
- 16. Explain in brief about Paper electrophoresis technique.
- 17. Describe different methods for quantitative analysis of single component samples by UV spectrophotometry.
- 18. Explain the principle and measurement of Interferences in Atomic Absorption spectroscopy.
- 19. Explain the principles, advantages and disadvantages, and applications of thin layer chromatography.
- 20. Write about the Spectrophotometric titrations with examples?
- 21. Explain the different derivatization techniques used in Gas Chromatography?
- 22. Explain the instrumentation of Nephelotubiodmetry.

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## B. Pharmacy (PCI) VII - Semester (Main & Backlog) Examination, March 2025 Subject: Novel Drug Delivery Systems

Time: 3 Hours Max. Marks: 75

PART - A

Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Define the following terms?
  - a) Controlled drug delivery system
  - b) Sustained drug delivery system
- 2. Distinguish between matrix and reservoir systems?
- 3. List out the methods used for liposomes?
- 4. Define the following
  - a) Osmotic drug delivery system
- b) Transdermal drug delivery system
- 5. Classify gastro retentive drug delivery systems?
- 6. Define the following?
  - a) Implants
- b) Niosomes
- 7. Differentiate between Zero Order and First Order release kinetics?
- 8. List out the different types of nanoparticles?
- 9. Applications of monoclonal antibodies?
- 10. Discuss the advantages of Ocusert?

PART - B

Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Discuss the formulation and evaluation of floating drug delivery systems?
- 12. Write in detail about the coacervation phase separation technique?
- 13. Write in detail about the following?
  - a) Explain about the push pull systems?
  - b) Mucoadhesive drug delivery system?

PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Discuss about the factors influencing formulation of sustained release system?
- 15. Write the polymerization techniques?
- 16. Explain the Wuster process for microencapsulation with an example?
- 17. Explain the different theories of mucoadhesion?
- 18. Describe the formulation of Buccal drug delivery systems?
- 19. Discuss about the metered dose inhalers?
- 20. Write about ocular controlled drug delivery systems? Describe the methods to overcome the ocular barriers?
- 21. Write about the applications Intrauterine devices?
- 22. Write about the elementary osmotic pump?

Code No: G-13108/PCI

## **FACULTY OF PHARMACY**

## B. Pharmacy VII Semester (PCI) (Main & Backlog) Examination, March 2025 Subject: Pharmacy Practice

Time: 3 Hours Max. Marks: 75

#### PART - A

## Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Classify Hospitals based on the system of medicine and speciality.
- 2. What is Idiosyncrasy? Give examples.
- Define rational use of medicines.
- 4. Enlist the types of drug distribution systems.
- 5. Mention the different sources of drug information.
- 6. What do you mean by automatic stop orders?
- 7. Define Clinical Pharmacy. Mention its objectives.
- 8. Explain the significance of OTC drugs.
- 9. Define inventory. Mention the objectives of inventory control.
- 10. Define and classify ADR.

#### PART - B

## Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Define Medication Adherence. Mention the methods to measure it. What is the role of a Pharmacist in promoting medication adherence in patients.
- 12.a) Explain in detail the objectives of Pharmacy and Therapeutic Committee (PTC).
  - b) Discuss the role of PTC in adverse drug monitoring.
- 13. Define Therapeutic Drug Monitoring (TDM). Mention its objectives and explain the process involved in TDM.

#### PART - C

## Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Define hospital and explain its organization.
- 15. Describe the various systems involved in the dispensing of drugs to inpatients.
- 16. Define hospital formulary and explain its need.
- 17. Explain the role and reponsibilities of community pharmacist.
- 18. Explain why communication skill is important for a pharmacist.
- 19. Discuss the role of Pharmacist in the education and training program in the hospital.
- 20. Discuss the role of Pharmacist in the interdepartmental communication and community health education.
- 21. Explain hospital budget preparation and implementation.
- 22. Mention the various laboratory blood tests. Explain their significance.

**Code No: G-13107/PCI** 

## **FACULTY OF PHARMACY**

B. Pharmacy (PCI) VII - Semester (Main & Backlog) Examination, March 2025 Subject: Industrial Pharmacy - II

Time: 3 Hours Max.Marks:75

#### PART - A

#### Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. What is scale up?
- 2. Write a note on documentation in pilot plant.
- 3. What is technology transfer?
- 4. Write a note on legal issues in technology transfer.
- 5. What is qualification and validation?
- 6. Write a note on Investigator's Brochure (IB).
- 7. What is quality assurance?
- 8. Why informed consent procedure is important in clinical trials?
- 9. Write the role of ISO in quality management,
- 10. Write a note on state licensing authority responsibilities.

#### PART - B

## Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. What is pilot plant? Write the general considerations for pilot plant and scale up for Tablets and Liquid dosage forms.
- 12. Write a note on the (i) IND and NDA application (ii) Clinical research protocol.
- 13. (a) Write a note on Indian drug regulatory. Write CDSCO functions.
  - (b) Explain about Central Drugs Laboratory and its function.

#### PART - C

## Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Write the SUPAC guidelines for solid and liquid dosage forms.
- 15. Write a note on documentation in pilot plant and scaleup.
- 16. Write general principles of technology transfer.
- 17. Write the role and responsibility of regulatory affairs professionals.
- 18. Write a note on APCTD, NRDC, TIFAC technology transfer agencies in India.
- 19. Write the Principles and applications of QBD.
- 20. Write a note on TQM.
- 21. Write a note on NABL and GLP.
- 22. Write a note on regulatory requirements and approval procedures for new drugs.

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Code No: F-7191/PCI

## **FACULTY OF PHARMACY**

B. Pharmacy VII - Semester (PCI) (Main & Backlog) Examination, March 2024 Subject: Instrumental Methods of Analysis

Time: 3 Hours Max. Marks: 75

PART - A

Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Explain bathochromic shift and Hypsochromic shift with examples.
- 2. What are chromophores and auxochromes. Give examples.
- 3. Write the principles of absorption in IR spectroscopy.
- 4. Write the principles of partition and adsorption chromatography.
- 5. Write the different fuel gases and oxidants used in the flame photometry technique.
- 6. Write the different types of stationary phases used in gel permeation chromatography separations.
- 7. Write the ion exchange mechanism in ion exchange chromatography.
- 8. Define the Capacity factor.
- 9. Write the effect of solvent on the absorption maximum of compounds.
- 10. Write the applications of affinity chromatography.

PART - B

Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Describe different components of IR spectrophotometer with a neat labelled diagram.
- 12. Explain the principles and experimental details of Paper chromatography.
- 13. Explain the principles and instrumentation of the HPLC technique.

PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Describe the Jablonski diagram and explain different internal and external processes in fluorescence emission.
- 15. Explain the factors affecting Ion exchange Chromatography and applications of the technique.
- Explain different sample handling techniques used in IR spectroscopy.
- 17. Write the Instrumentation and applications of the flame photometry technique.
- 18. Write short notes on nepheloturbidometry.
- 19. Describe the different types of detectors used in UV spectrophotometers.
- 20. Explain the different development techniques used in paper chromatography.
- 21. Write the principles and applications of Atomic absorption spectroscopy.
- 22. Discuss the theory and principles of separation in capillary electrophoresis.

Code No: F-7194/PCI

## **FACULTY OF PHARMACY**

B. Pharmacy VII - Semester (PCI) (Main & Backlog) Examination, March 2024 Subject: Novel drug delivery systems

Time: 3 Hours Max. Marks: 75

#### PART - A

## Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Differentiate between matrix and reservoir system.
- 2. Define polymers. Classify them with examples.
- 3. Define microencapsulation. Write its applications.
- 4. Write the advantages and disadvantages of mucoadhesive drug delivery system.
- 5. Define microspheres and microcapsules.
- 6. Write note on permeation enhancers used in Transdermal drug delivery system with examples.
- 7. What is floating time and floating lag time.
- 8. Write the applications of targeted drug delivery system.
- 9. Explain the basic structural components of liposomes.
- 10. Explain about intra ocular barriers.

### PART - B

## Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Explain the approaches used in development of gastro retentive drug delivery systems.
- 12. Explain in detail any two methods of microencapsulation.
- 13. Explain the basic components and formulation approaches used in transdermal drug delivery system.

#### PART - C

## Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Discuss the physicochemical factors affecting controlled drug delivery system.
- 15. Explain the principles of mucoadhesion.
- 16. Write a note on nebulizers.
- 17. Discuss about intra uterine devices.
- 18. Explain about preparation methods of liposomes.
- 19. Write about production of monoclonal antibodies.
- 20. Explain about ocular inserts.
- 21. Explain about osmotic pump.
- 22. Explain about the inflatable and gastroadhesive systems.

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Code No: F-7193/PCI

## **FACULTY OF PHARMACY**

## B. Pharmacy VII - Semester (PCI) (Main & Backlog) Examination, March 2024 Subject: Pharmacy Practice

Time: 3 Hours Max. Marks: 75

#### PART - A

## Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Define Hospital.
- 2. Define Hospital Pharmacy.
- 3. Define ADR.
- 4. Define controlled drugs.
- 5. Define Hospital formulary.
- 6. Mention few drugs which require TDM.
- 7. What do you mean by automatic stop order?
- 8. Define OTC drugs.
- 9. What do you mean by investigational new drug?
- 10. What is the significance of ESR?

#### PART - B

## Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Define clinical pharmacy, explain the functions and responsibilities of clinical pharmacist.
- 12. Define inventory control? Explain in detail any one method of inventory control technique used in the procurement of drugs
- 13. Explain in detail therapeutic drug monitoring.

## PART - C

## Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Explain functions of hospital pharmacy.
- 15. Describe different types of adverse drug reactions.
- 16. What are the legal requirement for establishing a community pharmacy?
- 17. Explain in detail procedure for dispensing of controlled drugs.
- 18. What do you mean by rational use of drugs? How the concept of Rational use can be implemented for OTC drugs
- 19. Explain salient features of hospital budget preparation.
- 20. Explain in detail various drug purchasing procedure in a hospital pharmacy
- 21. Explain in detail any four blood tests and their significance.
- 22. What do you mean by adherence? What are the methods to improve the patient adherence towards chronic therapy.

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Code No: F-7192/PCI

## **FACULTY OF PHARMACY**

## B. Pharmacy VII - Semester (PCI) (Main & Backlog) Examination, March 2024 Subject: Industrial Pharmacy

Time: 3 Hours Max. Marks: 75

#### PART - A

## Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Write a note on SUPAC guidelines.
- 2. What is pilot plant and scale-up?
- 3. Explain the importance of validation.
- 4. What is Technology transfer?
- 5. Write the role of regulatory affairs.
- 6. Mention five important data documents for ANDA.
- 7. Write a note on different stages of clinical trials.
- 8. What is informed consent?
- 9. Write a note on ISO 9000.
- 10. Write the role of CDL.

PART - B

Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Write about pilot plant and scale up requirements for Tablets and Capsules.
- 12. (a) What is technology transfer? Write general principles of Technology Transfer.
  - (b) Write the role and responsibility of regulatory affairs professionals.
- 13. (a) Explain the principles of QBD and applications of QbD.
  - (b) Write a note on NABL and GLP.

PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Write a note on pilot plant scale-up for liquid dosage forms.
- 15. Write a note on Technology Transfer procedure from R&D to production (Process, packaging and cleaning).
- 16. Write briefly on Investigational New Drug (IND) Application.
- 17. Write the role of biostatistics in pharmaceutical product development
- 18. What is QRM? Describe the principle and process of QRM.
- 19. Write a note on six sigma concept.
- 20. Write briefly on TQM.
- 21. Write a note on Indian Regulatory. Write CDSCO functions.
- 22. Write a note on COPP.

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Code No: E-12328/PCI

## **FACULTY OF PHARMACY**

## B. Pharmacy VII Semester (PCI) (Backlog) Examination, July-2023

Subject: Novel drug delivery systems

Time: 3 Hours Max. Marks: 75

PART - A

Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Define terms sustained, controlled and targeted release dosage forms.
- List out pharmacokinetic parameters suitable for selection of drug for controlled drug delivery system.
- 3. Explain about inflatable systems.
- 4. Explain the Nasal and Pulmonary routes of drug delivery.
- 5. Write the advantages and disadvantages of gastroretentive drug delivery system.
- 6. Describe various coating materials used in microencapsulation.
- 7. Write note on transmucosal permeability.
- 8. Explain advantages and development of intrauterine devices.
- 9. Write the applications of monoclonal antibodies.
- 10. Differentiate between liposomes and niosomes

PART - B

Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Explain in detail physicochemical and biological factors affecting controlled release formulations.
- 12. Explain in detail coacervation phase separation method with suitable examples.
- 13. Discuss about in detail a) Alzet osmotic pump b) Dry powder inhaler

PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Discuss the classification and applications of polymers used in controlled drug delivery system.
- 15. Explain the theories of mucoadhesion.
- 16. Explain about factors affecting permeation in transdermal drug delivery system..
- 17. Discuss the approaches used in development of gastroretentive drug delivery system.
- 18. Explain about nasal sprays and nebulizers.
- 19. Write a note on niosomes.
- 20. Discuss the ocusert with neat sketch.
- 21. Explain the preparation methods of nanoparticles.
- 22. Explain metered dose inhalers.

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Code No: E-12327/PCI

## **FACULTY OF PHARMACY**

## B. Pharmacy VII Semester (PCI) (Backlog) Examination, July 2023 Subject: Pharmacy Practice

Time: 3 Hours Max. Marks: 75

PART - A

Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Classify Hospitals based on the system of medicine and speciality.
- 2. Define community Pharmacy
- 3. Mention any two pharmacokinetic drug interactions
- 4. Define rational use of medicines
- 5. Mention few principles for the inclusion of drugs in hospital formulary
- 6. Define TDM.
- 7. Mention any two functions of DTC
- 8. Define DIC.
- 9. Mention two tertiary references used for drug information center?
- 10. What is the significance of C-reactive protein?

PART - B

Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Define clinical pharmacy, What are the roles and responsibilities of a clinical pharmacy department.
- 12. Define adherence. What are the factors affecting patient adherence. How adherence can be improved?
- 13. Explain in detail organisation and functions of DTC

PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Explain schedule N of drugs and cosmetics act 1940.
- 15. Describe different types of drug interactions with examples
- 16. What are the legal requirement for establishing a community pharmacy
- 17. Explain economic order quantity.
- 18. What are OTC drugs? How OTC drugs to be counselled?
- 19. Explain salient features of hospital budget preparation.
- 20. Explain the importance of communication skill for the pharmacist
- 21. Explain various urine test and its significance
- 22. What is patient counselling? What are the barriers of patient counselling?

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Code No: E-12325/PCI

## **FACULTY OF PHARMACY**

B. Pharmacy VII Semester (PCI) (Backlog) Examination, July / August 2023 Subject: Instrumental Methods of Analysis

Time: 3 Hours Max. Marks: 75

PART - A

Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Explain different types of electronic transitions.
- 2. What is fluorescence quenching? Give examples.
- 3. Write the applications of Nephelometry and turbidometry techniques.
- 4. What are the different types of molecular vibrations in IR spectroscopy?
- 5. Write the principles of separation in Electrophoresis.
- 6. What is chromophore and auxochrome. Give examples.
- 7. What are the different types of Ion exchange resins used in Ion-exchange chromatography?
- 8. Define theoretical plate and give the formula for calculating theoretical plates.
- 9. Write the principle involved in affinity chromatography.
- 10. State and explain Beer-Lamberts law.

PART - B

Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Describe different components of a UV-Visible spectrophotometer.
- 12. Explain the principles and experimental details of thin layer chromatography.
- 13. Explain the principles and instrumentation of Gas chromatography technique.

PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Explain in brief the gel electrophoresis technique.
- 15. Describe different methods for quantitative analysis of single component samples by UV spectrophotometry.
- 16. Explain the principles, advantages and disadvantages, and applications of paper chromatography.
- 17. Describe different types of detectors used in HPLC.
- 18. Write the principles and applications of atomic absorption spectroscopy.
- 19. Explain different sample handling techniques used in IR spectroscopy.
- 20. Explain the principles of fluorescence and Phosphorescence with the help of the Jablonski diagram.
- 21. Explain the principles and applications of partition and adsorption chromatography.
- 22. Write the different factors affecting electrophoresis separartions G.Pulla Reddy College of Pharmacy Hyderabad

Code No: E-12326/PCI

## **FACULTY OF PHARMACY**

B. Pharmacy VII Semester (PCI) (Backlog) Examination, July 2023 Subject: Industrial Pharmacy

Time: 3 Hours Max. Marks: 75

PART - A

Note: Answer all the questions.  $(10 \times 2 = 20 \text{ Marks})$ 

- 1. What is technology transfer?
- 2. Write a note on raw materials importance in pilot plant.
- 3. Write a note on analytical method transfer procedure in technology transfer.
- 4. Write a note on legal issues in technology transfer.
- 5. What is Investigator's Brochure (IB)?
- 6. What is quality assurance?
- 7. Write a note on GLP.
- 8. Write a note on role of ISO in quality management.
- 9. Write a note on COPP.
- 10. Write a note on state licensing authority responsibilities.

PART - B

Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. What is pilot plant? Write the general considerations for pilot plant and scale up for Tablets and Liquid dosage forms.
- 12. Write a note on the i) Principles of QBD ii) Six sigma concept.
- 13.(a) Write a note on Indian Regulatory. Write CDSCO functions.
  - (b) Explain about Central Drugs Laboratory and its function.

PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Write the significance of personnel requirements and space requirements in pilot plant and scale up.
- 15. Write a note on documentation in pilot plant and scaleup.
- 16. Write general principles of technology transfer.
- 17. Write a note on technology transfer agencies in India.
- 18. Write the role and responsibility of regulatory affairs professionals.
- 19. Write about IND and NDA application.
- 20. Write the role of biostatistics in pharmaceutical product development.
- 21. Write the applications of QbD in formulation development.
- 22. Write the SUPAC guidelines for various dosageforms.

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Code No: E-12264/PCI

## FACULTY OF PHARMACY

B.Pharmacy VII-Semester (PCI) (Main & Backlog) Examination, March 2023.

Subject: Instrumental Method of Analysis

Time: 3 Hours Max.Marks:75

#### PART - A

## Note: Answer all the questions.

(10 x 2 = 20 Marks)

- Explain the principle involved in Pyroelectric detector in IR spectroscopy?
- 2. What are Chromophores, mention some example?
- Define the term Quanching and their types?
- 4 Explain the principle involved in Bolometer Detector?
- Define the term Electrophoratic mobility?
- Write adsorbents used in column chromatography?
- Mention different types of detectors used in Gas chromatography?
- B. Write about the types of pumps used in HPLC?
- Explain Spectrophotometric Titrations?
- 10. Derive the Beer Lambert's law?

#### PART - B

#### Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Compare the methodology of Thin Layer and Paper Chromatography?
- 12. Explain in detail about the working principle and technique of Capillary Electrophoresis?
- Explain the principle and applications of Nepheloturbidometry?

#### PART - C

## Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Explain the theory and procedure involved in Affinity Chromatography?
- 15 Write the Applications of Flame Photometry with examples?
- 16. Write in detail about the factors affecting Electrophoretic Mobility?
- Describe the methodology of PartitionColumn Chromatography?
- 18. Write about the Interferences and their types in Atomic Absorption Spectroscopy?
- 19. Write a note on different sources of IR spectroscopy?
- 20 Describe the Factors affecting Ion Exchange Process in Ion Exchange Chromatography?
- 21 Write about the applications of UV with respect to single and multi-component analysis?
- 22. What about the stationary Phases used in Gel Chromatography?

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Code No: E-12265/PCI

## FACULTY OF PHARMACY

## B. Pharmacy VII Semester (PCI) (Main & Backlog) Examination, March-2023

Subject: Industrial Pharmacy

Time: 3 Hours Max. Marks: 75

#### PART-A

#### Note: Answer all the questions.

(10x2=20marks)

- What is pilot plant and scale-up?
- Write a note on SUPAC.
- Dofino validation.
- Name few approved regulatory bodies.
- 5. Write the role of regulatory affairs
- Wirte the importance of ANDA
- Write a note on informed consent.
- 8. Write a note on GLP.
- 9 Write a note on COPP.
- 10. What is the role of Drug control laboratory?

#### PART-B

## Note: Answer any two questions

(2x10=20Marks)

- 11. Write about pilot plant and scale up requirements for solid dosage forms.
- 12. a) What is technology transfer? Write general principles of Technology Transfer.
  - b) Write a note on Technology Transfer agencies in India.
- 13. a) Explain the principles of QBD
  - b) Write a note on six sigma concept.

#### PART-C

#### Note: Answer any seven questions

(7x5=35Marks)

- 14. Write a note on pilot plant scale up for semi solid dosage forms.
- 15. Write a note on Technology Transfer related documentation.
- 16. Write the role of regulatory affairs department in drug approval.
- Write briefly on Investigational New Drug (IND) Application.
- 18. What is ORM? Describe the principle and process of QRM.
- 19 Write a note on ISO 14000.
- Write briefly on TQM.
- Write a note on Indian Regulatory, Write COSCO functions.
- Explain about Central Drugs Laboratory and its function.







Code No: E-12266/PC)

## FACULTY OF PHARMACY

B. Pharmacy VII-Semester (PCI) (Main & Backlog) Examination, March-2023 Subject: Pharmacy Practice

TIMe: 3 hours

Max.Marks: 75

#### PART-A

Note: Answer all the questions.

(10x2=20marks)

- Define and classify hospitals.
- 2. Define hospital pharmacy
- Define ADR?
- 4. Mention the different types of community pharmacy outlet
- 5. Define hospital formulary.
- Define TDM
- 7. Define medication adherence.
- 8. What do you mean by automatic stop orders?
- Define drug information center.
- 10 Give a general patient counselling information for antibiotics

#### PART-B

## Note: Answer any two questions

(2x10=20Marks)

- 11 Describe organisational structure of hospital and its functions. Add a note on responsibilities of a hospital pharmacist.
- 12. Describe legal requirements to establish a community pharmacy outlet. What are the records to be maintained in community pharmacy outlet?
- 13 Explain the causes of medication adherence? What is the role of a pharmacist in promoting medication adherence in patients with chronic diseases?

#### PART-C

## Note: Answer any seven questions

(7x5=35Marks)

- 14. What is the need for TDM. What are factors to be considered during TOM?
- Explain the importance of ipatient history and patient medication history interview.
- 16. Explain the organisation and functions of pharmacy and therapeutic committee.
- Mention the sources of drug information. Explain the advantages and disadvantages of the same.
- Explain the role of a pharmacist in interdepartmental communication and health education in community
- 19 Explain why communication skill is important for a pharmacist.
- Explain drug related problems with examples.
- Describe economic order quantity.
- Explain various urine tests and their significance

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Code No: E-12267 / PCI

## FACULTY OF PHARMACY

## B. Pharmacy VII-Semester (PCI) (Main & Backlog) Examination, April-2023

Subject: Novel Drug Delivery Systems

Time: 3 Hours

PART - A

Max. Marks: 75

Note: Answer all questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- Define the following dosage forms:
  - (a) Controlled drug delivery systems
  - (b) Sustained release drug delivery systems
- Differentiate between matrix and reservoirs systems.
- 3 List out the methods used for microencapsulation technique
- 4 Write the advantages of buccal drug delivery systems.
- Types of permeation enhancers used in TDDS with examples.
- 6 Define the following
  - (a) Lipsomes (ii) Nipsomes
- 7 Differentiate between Zero Order and Frist Order release kinetics
- 8 List out the different types of nanoparticles.
- 9 Enumerate the applications of monoclonal antibodies.
- 10 Write the advantages of Ocuserts

PART - B

Note: Answer any two questions.

 $\{2 \times 10 = 20 \text{ Marks}\}$ 

- 11 Discuss the formulation of Transdermal drug delivery systems with the components and examples? Add a note on factors affecting permeation
- 12 Describe in detail any two methods for preparation of microencapsulation.
- 13 Write in detail about the following:
  - (a) Explain about the Alzet osmotic pump.
  - (b) Mucoadhesive drug delivery system.

PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14 Classify the polymers with examples.
- 15 Discuss about the factors influencing of controlled drug delivery system.
- 16 Explain the different theones of mucoadhesion
- 17 Write about the elementary osmotic pump.
- 18 Describe the formulation of floating drug delivery systems.
- 19 Discuss about the dry powder inhalers
- 20 Write a note on intraocular barriers. Describe the methods to overcome the problem.
- 21 Intrauterine devices and their applications.
- 22 Explain the Wuster process for microencapsulation with an example

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## B. Pharmacy VII-Semester (PCI) (Backlog) Examination, August 2022 Subject: Industrial Pharmacy

Time: 3 Hours Max. Marks: 75

Note: Answer all questions PART – A, any two questions from PART – B and any seven question from PART – C.

## $PART - A (10 \times 2 = 20 Marks)$

- 1 Write a note on SUPAC.
- 2 What is validation?
- 3 Write a note on DQ, IQ, OQ and PQ.
- 4 What is QRM?
- 5 Define API and excipient.
- 6 What are various phases of clinical trials?
- 7 What is the aim of NDA?
- 8 Define Bioavailability and Bioequivalence.
- 9 Write a note on CDSCO.
- 10 What is RDTL and its functions?

## $PART - B (2 \times 10 = 20 Marks)$

- 11 (a) Write the General considerations for pilot plant and scale up.
  - (b) Write a note on platform technology.
- 12 (a) Write a note on six sigma concept.
  - (b) Write a note on ISO 14000.
- 13 (a) Discuss Regulatory requirements and approval procedures for New Drugs.
  - (b) Write the responsibilities of State Licensing authorities.

## PART - C (7 x 5 = 35 Marks)

- 14 Explain the procedure for pilot plat scale-up for semisolid dosage forms.
- 15 What is technology transfer? Write general principles of Technology Transfer.
- 16 Write the role and responsibility of regulatory affairs professionals.
- 17 Write a note on technology transfer agencies in India.
- 18 Write briefly on Investigational New Drug (IND) Application.
- 19 Write about QbD and its applications.
- 20 Write about the Certificate of Pharmaceutical Product (COPP).
- 21 Write a note on the principle and process of QRM.
- 22 Write NDA Review process.

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## B. Pharmacy VII-Semester (PCI) (Backlog) Examination, August 2022 Subject: Instrumental Methods of Analysis

Time: 3 Hours Max. Marks: 75

Note: Answer all questions PART – A, any two questions from PART – B and any seven question from PART – C.

## $PART - A (10 \times 2 = 20 Marks)$

- 1. Define auxochrome and chromophore with example.
- 2. What is Quenching and types of quenching?
- 3. Write the interferences in Flame photometry and types of interference.
- 4. Name the Infra-Red radiation source.
- 5. Define the term chromatography and the general principle involved in it.
- 6. Mention the factors affecting Electrophoretic Mobility.
- 7. Write about the temperature program in Gas chromatography.
- 8. Explain different types of pumps used in HPLC and their brief working principle.
- 9. Explain the principle involved in Ion Exchange Chromatography.
- 10. Write the theory involved in Gel Chromatography.

## $PART - B (2 \times 10 = 20 Marks)$

- 11.(a) Explain in detail about the construction and working principle of detectors used in UV-Vis spectroscopy.
  - (b) Write about the Methodology involved in Paper Chromatography.
- 12.(a) Describe the sources and sampling techniques in IR spectroscopy.
  - (b) Explain the factors affecting in exchange methodology in ion exchange chromatography.
- 13.(a) Explain the applications of HPLC with examples.
  - (b) Write about the Instrumentation of Affinity chromatography.

## PART – C $(7 \times 5 = 35 \text{ Marks})$

- 14. Explain the technique of Capillary Electrophoresis.
- 15. Write about electronic transitions and solvent effect on absorption spectra.
- 16. Describe the theory involved in fluorimetric technique.
- 17. Explain the instrumentation of Nephelotubiodmetry.
- 18. Write the factors affecting vibration in IR spectroscopy.
- 19. Differentiate between single and multi-component analysis in UV-Vis spectroscopy with examples.
- 20. Explain the principle and Interference in Atomic Absorption spectroscopy.
- 21.(a) Write the principle involved in column chromatography.
  - (b) Explain the working principle of Thermocouple Detector.
- 22. Write about the Detectors used in HPLC.

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B. Pharmacy VII - Semester (PCI) (Backlog) Examination, September 2022 Subject: Novel Drug Delivery Systems

Time: 3 Hours Max. Marks: 75

#### PART - A

Note: Answer all questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Define terms sustained, controlled and targeted release dosage forms.
- 2. Write ideal characters suitable for selection of drug for controlled drug delivery system.
- 3. Explain about inflatable systems.
- 4. Explain the Nasal and Pulmonary routes of drug delivery.
- 5. Write the advantages and disadvantages of gastroretentive drug delivery system.
- 6. Explain various coating materials used in microencapsulation.
- 7. Write a note on transmucosal permeability.
- 8. What is floating time and floating lag time.
- 9. Write the applications of monoclonal antibodies.
- 10. Compare and contrast liposomes and niosomes.

### PART - B

Note: Answer any two questions.

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Explain in detail physiochemical and biological factors affecting controlled release formulations.
- 12. Explain in detail coacervation phase separation method with suitable examples.
- 13. Discuss about advantages and disadvantages and development of intra uterine devices and applications.

#### PART - C

Note: Answer any seven questions.

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Discuss the classification and applications of polymers used in controlled drug delivery system.
- 15. Explain the theories of mucoadhesion.
- 16. Explain about factors affecting permeation in transdermal drug delivery system.
- 17. Discuss the approaches used in development of gastroretentive drug delivery system.
- 18. Explain about nasal sprays and nebulizers.
- 19. Explain about osmotic pump.
- 20. Discuss the ocusert with neat sketch.
- 21. Explain the preparation methods of nanoparticles.
- 22. Explain dry powder and metered dose inhalers.

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## B. Pharmacy VII-Semester (PCI) (Backlog) Examination, September 2022 Subject: Pharmacy Practice

Time: 3 hours Max Marks: 75

PART - A

Note: Answer all the questions.

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Define Primary, Secondary and Tertiary hospital.
- 2. Mention the functions of hospital pharmacy
- 3. Mention the classification of ADR
- 4. Define idiosyncrasy.
- 5. Mention few examples of pharmacokinetic drug interactions
- 6. Mention few drugs which require TDM
- 7. Define patient counselling.
- 8. Define lead time.
- 9. Define investigational drug.
- 10. Give a general patient counselling information for NSAIDs

PART - B

Note: Answer any two questions

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Describe different types of drug interactions. Add a note on reporting and management of ADR
- 12. Describe organisation, structure, type and design of wholesale and community pharmacy outlet
- 13. Explain different types of drug distribution system in a hospital. What do you mean by satellite pharmacy?

PART - C

Note: Answer any seven questions

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Define hospital formulary. What are the contents of hospital formulary? What is the difference between hospital formulary and essential drugs list?
- 15. Explain the role of pharmacist in improving medication adherence and highlight few counselling barriers.
- 16. Describe schedule N of drugs and cosmetics act rules 1945.
- 17. Describe the policies of pharmacy and therapeutic committee.
- 18. Explain the systematic approach of handling a drug information query.
- 19. Explain the role of a pharmacist in training and education.
- 20. Explain hospital budget preparation and implementation.
- 21. Define OTC drugs. What is the role of pharmacist in implementing rational use OTC drugs.
- 22. Classify investigational drugs. Explain haematological tests and its significance.

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## B. Pharmacy VII - Semester (PCI) (Main & Backlog) Examination, February / March 2022

**Subject: Instrumental Methods of Analysis** 

Time: 3 Hours Max. Marks: 75

#### PART - A

Note: Answer all questions:

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. State and explain Beer-Lambert equation.
- 2. What is fluorescence quenching? Give examples.
- 3. Write the principles of Flame photometry technique.
- 4. Write the applications of Nephelometry and turbidometry techniques.
- 5. Write different types of stationary phase column packing materials used in HPLC.
- 6. Write Van Deempter equation.
- 7. What are the different types of Ion exchange resins used in Ion-exchange chromatography?
- 8. Define theoretical plate and give formula for calculating theoretical plates.
- 9. What is an electronic transition and types?
- 10. Write the principle involved in affinity chromatography.

## PART - B

Note: Answer any two questions:

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Describe different components of IR spectrophotometer.
- 12. Explain the principles and experimental details of paper chromatography for Quantitative analysis.
- 13. Explain the principles and instrumentation of Gas chromatography technique.

#### PART - C

## Note: Answer any seven questions:

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Explain in brief about Paper electrophoresis technique.
- 15. Describe different methods for quantitative analysis of single component samples by UV spectrophotometry.
- 16. Explain the principles, advantages and disadvantages and applications of thin layer chromatography.
- 17. Write about Gel Permeation chromatography.
- 18. Write the principles and applications of Atomic absorption spectroscopy.
- 19. Explain different sample handling techniques used in IR spectroscopy.
- 20. Explain the principles of fluorescence and Phosphorescence with help of Joblonski diagram.
- 21. Explain the principles and applications of partition and adsorption chromatography.
- 22. Write the different factors affecting electrophoretic mobility of ions in electrophoresis separations.

## B. Pharmacy VII-Semester (PCI) (Main & Backlog) Examination, February / March 2022

**Subject: Novel Drug Delivery Systems** 

Time: 3 Hours Max. Marks: 75

PART - A

**Note: Answer all questions:** 

 $(10 \times 2 = 20 \text{ Marks})$ 

- 1. Define terms sustained, controlled and targeted release dosage forms.
- 2. Enlist ideal characters suitable for selection of drug for controlled drug delivery system.
- 3. Define microencapsulation, write its applications.
- 4. What are implantable drug delivery system with examples?
- 5. Write the advantages and disadvantages of mucoadhesive drug delivery system.
- 6. Explain various coating materials used in microencapsulation.
- 7. Write a note on permeation enhancers with examples.
- 8. What is floating time and floating lag time?
- 9. Write the applications of monoclonal antibodies.
- 10. Write the methods of evaluation of liposomes.

PART - B

**Note: Answer any two questions:** 

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Explain in detail physicochemical and biological factors affecting controlled release formulations.
- 12. Explain the methods of microencapsulation.
- 13. Discuss the basic components, formulation approaches for development of transdermal drug delivery system.

PART - C

Note: Answer any seven questions:

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Discuss the classification and applications of polymers used in controlled drug delivery system.
- 15. Explain the theories of mucoadhesion.
- 16. Write a note on osmotic pump.
- 17. Discuss the approaches used in development of gastroretentive drug delivery system.
- 18. Explain about nasal sprays and nebulizers.
- 19. Write a note on niosomers.
- 20. Discuss the ocuserts with neat sketch.
- 21. Explain the applications of intrauterine devices.
- 22. Explain the formulation considerations of buccal drug delivery system.

## B. Pharmacy VII - Semester (PCI) (Main & Backlog) Examination, February / March 2022

**Subject: Pharmacy Practice** 

Time: 3 Hours Max. Marks: 75

#### PART - A

Note: Answer all questions:  $(10 \times 2 = 20 \text{ Marks})$ 

- 1. What are the roles of clinical pharmacist in ward rounds?
- 2. Write the classification of drug related problems.
- 3. Mention the requisites & objectives for management of materials in hospital pharmacy.
- 4. Describe the significance of Drug Information Center.
- 5. Explain the important considerations for Therapeutic Drug Monitoring.
- 6. Give a brief note on the factors affecting drug variability.
- 7. Write a short note on material requirement for community pharmacy.
- 8. Give definition of drug integrations and classify them accordingly.
- 9. Enumerate the types of drug ADRs with examples.
- 10. Write a note on rational use of drugs.

#### PART - B

## Note: Answer any two questions:

 $(2 \times 10 = 20 \text{ Marks})$ 

- 11. Define P & T Committee and write its objectives, organization and various functions.
- 12. Define Hospital and enumerate the organization and functions of hospital.
- 13. What is meant by clinical pharmacy? Explain functions and responsibility of clinical pharmacy.

#### PART - C

## **Note: Answer any seven questions:**

 $(7 \times 5 = 35 \text{ Marks})$ 

- 14. Give a comprehensive note on factors affecting Therapeutic Drug Monitoring.
- 15. Explain the roles and responsibility of hospital pharmacist.
- 16. Describe the procurement or purchasing procedure for pharmaceuticals in detail.
- 17. Explain various hematologic tests and their significance.
- 18. Explain the steps involved in the preparation of hospital formulary.
- 19. Elaborate the requirements for establishment of Drug Information Center.
- 20. Provide the detailed role of pharmacist in medication adherence.
- 21. Write all the inclusive steps involved in patient counseling.
- 22. Define Inventory Control. Specify the methods of Inventory Control.

## B. Pharmacy VII - Semester (PCI) (Main & Backlog) Examination, February / March 2022

Subject: Industrial Pharmacy - II

Time: 3 Hours Max. Marks: 75

PART - A

Note: Answer all questions:  $(10 \times 2 = 20 \text{ Marks})$ 

- 1 What is the need of pilot plant studies in pharmaceutical industries?
- 2 Write the level of changes expected under SUPAC.
- 3 Explain the quality risk management to technology transfer.
- 4 Describe the role of project team in the technology transfer.
- 5 Enlist at least four names of regulatory authorities functioning all around the world.
- 6 Enumerate the categories and type of INDs.
- 7 What are the benefits of NABL accreditation?
- 8 Mention the difference between corrective actions and preventive actions in quality system.
- 9 Write the functions of state regulatory authority.
- 10 What are the regulatory requirements for new drug approval?

PART - B

Note: Answer any two questions:  $(2 \times 10 = 20 \text{ Marks})$ 

- 11 Explain the steps involved in scale-up technology.
- 12 Define TQM and explain its key elements.
- 13 Discuss IND approval process in detail with help of flow diagram.

PART - C

Note: Answer any seven questions:  $(7 \times 5 = 35 \text{ Marks})$ 

- 14 Discuss the scale-up considerations for liquid oral pharmaceuticals.
- 15 Define the following: (a) Quality (b) QC (c) QA (d) Technology transfer (e) QbD
- 16 Discuss business process benchmarking as a tool of quality management.
- 17 What are the roles of regulatory affairs personnel in pharmaceutical industry?
- 18 Describe different models for the statistical design of clinical trials.
- 19 Discuss transfer of technology between R & D and manufacturing unit.
- 20 Differentiate between GMP and GLP.
- 21 Discuss importance of non-clinical drug development.
- 22 Describe the terms "QTPP" and "CQA" concerning QbD. Library,

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## **FACULTY OF PHARMACY**

# B. Pharmacy VII-Semester (PCI) (Backlog) Examination, September 2021

**Subject: Instrumental Method of Analysis** 

Time: 2 Hours Max. Marks: 75

## PART - A

Note: Answer any seven questions. (7 X 3 = 21 Marks)

- Explain the principle involved in Silicon photodiode detector in UV-Vis spectroscopy?
- 2. What are Singlet, Doublet and Triplet electronic states in Fluorimetry?
- 3. Define the term Retention time and Resolution in HPLC?
- 4. Explain the principle involved in Bolometer Detector?
- 5. Define the term Rf value.
- 6. Write the principles involved in Gel electrophoresis?
- 7. Mention different types of columns used in Gas chromatography?
- 8. Write different detectors compatible to HPLC?
- 9. Classify the Ion exchange chromatography?
- 10. Write about the deviations of Beer-Lamberts Law?

## PART - B

Note: Answer any one questions.

(1 X 14 = 14 Marks)

- 11. (a) Explain Theory and Instrumentation of Affinity Chromatography?
  - (b) Derive Beer-Lamberts Law?
- 12. Explain in detail the Instrumentation and Derivatization technique in Gas Chromatography?
- 13. (a) Write about the Spectrophotometric titrations with examples?
  - (b) Explain the Internal and External conversions in fluorimetry?

## PART - C

Note: Answer any five questions.

(5 X 8 = 40 Marks)

- 14. Write about the fundamental modes of Vibrations in polyatomic molecules?
- 15. Explain the Applications of Atomic Absorption spectroscopy with example?
- 16. Write in detail about the factors affecting Electrophoretic Mobility?
- 17. Describe the methodology of Adsorption Column Chromatography?
- 18. Write about the Interferences and their types in Flame Photometry?
- 19. Write a note on Wavelength selectors and sources of IR spectroscopy?
- 20. Describe the Principle involved in different sources of radiation of UV-Vis spectroscopy?
- 21. Write the methodology and application of Thin Layer Chromatography?
- 22. What is Quenching and explain the types of Quenching with examples?

# B. Pharmacy VII-Semester (PCI) (Backlog) Examination, September 2021

Subject: Industrial Pharmacy - II

Time: 2 Hours Max. Marks: 75

PART - A

Note: Answer any seven questions.  $(7 \times 3 = 21 \text{ Marks})$ 

- 1. What is Pilot Plant?
- 2. Write a note on SUPAC.
- 3. What is Technology Transfer?
- 4. Name few approved regulatory bodies and Technology Transfer agencies in India.
- 5. What is the role of regulatory affairs?
- 6. What are various phases of clinical trials?
- 7. What is Quality Assurance?
- 8. Write a note on GLP.
- 9. Write a note on Indian regulatory.
- 10. What is the role of Drug control laboratory?

#### PART - B

## Note: Answer any one questions.

(1 X 14 = 14 Marks)

- 11. What is Pilot plant and scale-up? Explain in detail about the scale up techniques for Solid dosage forms (Tablets/Capsules).
- 12. (a) Write a note on Indian Regulatory. Write C D S C O functions.
  - (b) Write short note on State Licensing authorities.
- 13. (a) Write the principles of TQM.
  - (b) Explain the principles of QBD.

PART - C

Note: Answer any five questions.  $(5 \times 8 = 40 \text{ Marks})$ 

- 14. Explain the procedure for pilot plant scale-up for liquid dosage form.
- 15. What is technology transfer? Write general principles of Technology Transfer.
- 16. Write the role of regulatory affairs department in drug approval.
- 17. What is QRM? Describe the principle and process of QRM.
- 18. Write briefly on Master Formula Record and its importance.
- 19. Write a note on ICH guidelines.
- 20. Explain about Central Drugs Laboratory and its function.
- 21. Write brief note on (i) IND (ii) NDA.
- 22. Write protocol for technology transfer.

## B. Pharmacy VII-Semester (PCI) (Backlog) Examination, September 2021

**Subject: Pharmacy Practice** 

Time: 2 Hours Max. Marks: 75

PART – A

Note: Answer any seven questions. (7 X 3 = 21 Marks)

- 1. Define Hospital. Classify it based on clinical ground.
- 2. What is Idiosyncrasy? Give examples.
- 3. Differentiate hospital formulary and drug list.
- 4. Enlist the types of drug distribution systems.
- 5. Mention the specific objectives of health education.
- 6. Discuss the interpretation of the prescription.
- 7. Define Budget. Mention the approaches involved in the budget preparation.
- 8. Explain the significance of OTC drugs.
- 9. Classify drug store based on design.
- 10. Mention the role of hospital pharmacist in the investigational use of drugs.

#### PART - B

Note: Answer any one questions.

(1 X 14 = 14 Marks)

- 11. Define Therapeutic Drug Monitoring (TDM). Mention its objectives and explain the process involved in TDM.
- 12. (a) Explain in detail the objectives of Pharmacy and Therapeutic Committee (PTC).
  - (b) Discuss the role of PTC in adverse drug monitoring.
- 13. Define Clinical Pharmacy. Explain in detail the functions and responsibilities of clinical pharmacist.

## PART - C

Note: Answer any five questions.

(5 X 8 = 40 Marks)

- 14. Discuss in detail the functions of hospital pharmacy.
- 15. Explain the role and responsibilities of community pharmacist.
- 16. Mention the role of Pharmacist in the medication adherence.
- 17. Describe the various systems involved in the dispensing of drugs to inpatients.
- 18. Illustrate the criteria for addition or deletion of drugs from hospital formulary.
- 19. Define patient counseling. Enlist the steps involved in patient counseling.
- 20. Discuss the role of Pharmacist in the interdepartmental communication and community health education.
- 21. Describe in brief the rational use of common over the counter medications.
- 22. Mention the various laboratory blood tests. Explain their significance.

# B. Pharmacy VII-Semester (PCI) (Backlog) Examination, September 2021

**Subject: Novel Drug Delivery Systems** 

Time: 2 Hours Max. Marks: 75

PART – A

Note: Answer any seven questions. (7 X 3 = 21 Marks)

- 1. Define the following dosage forms?
  - (a) Controlled drug delivery systems (b) Targeted drug delivery system.
- 2. Differentiate between matrix and reservoir systems?
- 3. List out the methods used for microencapsulation?
- 4. Define the following: (a) Implants (b) Transdermal drug delivery system.
- 5. Types of permeation enhancers used in TDDS with examples?
- 6. Define the following: (a) Liposomes (b) Niosomes
- 7. Differentiate between Zero Order and First Order release kinetic?
- 8. List out the different types of nanoparticles?
- 9. Enumerate the applications of monoclonal antibodies?
- 10. Write the advantages of Ocuserts?

PART - B

Note: Answer any one questions.

(1 X 14 = 14 Marks)

- 11. Discuss the formulation of Transdermal drug delivery systems with the components and examples? Add a note on factors affecting permeation?
- 12. Write in detail about the coacervation phase separation technique with examples?
- 13. Write in detail about the following:
  - (a) Explain about the Alzet osmotic pump?
  - (b) Mucoadhesive drug delivery system?

PART - C

Note: Answer any five questions. (5 X

(5 X 8 = 40 Marks)

- 14. Discuss about the factors influencing formulation of controlled drug delivery system?
- 15. Write the polymerization techniques?
- 16. Explain the Wuster process for microencapsulation with an example?
- 17. Explain the different theories of mucoadhesion?
- 18. Describe the formulation of floating drug delivery systems?
- 19. Discuss about the metered dose inhalers?
- 20. Write a note on intraocular barriers? Describe the methods to overcome the problem?
- 21. Write about the different types and applications of Intra-uterine devices?
- 22. Write about the elementary osmotic pump?

## B. Pharmacy VII-Semester (PCI) (Main) Examination, March 2021

## **Subject: Instrumental Methods of Analysis**

Time: 2 Hours Max. Marks: 75

#### PART - A

## Note: Answer any seven questions.

(7 X 3 = 21 Marks)

- 1. Define chromophore and Auxochrome and give examples.
- 2. Explain the phenomenon of Fluorescence and Phosphorescence.
- 3. What are the different types of fundamental modes of vibration in molecules after absorption of IR radiations?
- 4. Write the principles of partition and adsorption chromatography.
- 5. Write the different fuel gases and oxidants used in flame photometry technique.
- 6. Write the applications of gel permeation chromatography.
- 7. Write the ion exchange mechanism of ion exchange chromatogramphy.
- 8. Define retardation factor.
- 9. What is Bathochromic and Hypsochromic shift?
- 10. Write the principle involved in affinity chromatography.

#### PART - B

## Note: Answer any one questions.

(1 X 14 = 14 Marks)

- 11. Describe different components of UV spectrophotometer with a labeled diagram.
- 12. Explain the principles and experimental detail of thin layer chromatography for Quantitative analysis.
- 13. Explain the principles and instrumentation of HPLC technique.

## PART - C

Note: Answer any five questions.

(5 X 8 = 40 Marks)

- 14. Discuss the factors influencing intensity of fluorescence and applications of Fluorimetry technique.
- 15. Explain about gel electrophoresis.
- 16. Explain different sample handling techniques used in IR spectroscopy.
- 17. Write the theory and principle involved in flame photometry technique.
- 18. Write short notes on nepheloturbidometry.
- 19. Describe the different types of detectors used in Gas Chromatography.
- 20. Explain the different techniques used in paper chromatography.
- 21. Write the principles and applications of Atomic absorption spectroscopy.
- 22. Write the different factors affecting electrophoretic mobility of ions in electrophoresis separations.

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## B. Pharmacy VII-Semester (PCI) (Main) Examination, March 2021

**Subject: Novel Drug Delivery Systems** 

Time: 2 Hours Max. Marks: 75

## PART - A

#### Note: Answer any seven questions.

(7 X 3 = 21 Marks)

- 1. Write the advantages and disadvantages of controlled release dosage forms.
- 2. Explain various pharmacokinetic properties for selection of drug for controlled drug delivery system.
- 3. What are niosomes, write its structural components.
- 4. What are transdermal drug delivery system. Write its applications.
- 5. Write the advantages and disadvantages of mucoadhesive drug delivery system.
- 6. Define microspheres and microcapsules.
- 7. Write note on permeation enhancers with examples.
- 8. What is floating time and floating lag time.
- 9. Write the applications of targeted drug delivery system.
- 10. Write about classification of liposomes.

#### PART - B

#### Note: Answer any one question.

(1 X 14 = 14 Marks)

- 11. Explain the approaches used in development of gastro retentive drug delivery systems.
- 12. Explain in detail coacervation phase separation with suitable examples.
- 13. Discuss classification, properties and applications of polymers used in controlled drug delivery system.

## PART - C

## Note: Answer any five questions.

(5 X 8 = 40 Marks)

- 14. Discuss the physicochemical factors affecting controlled drug delivery system.
- 15. Explain the principles of mucoadhesion.
- 16. Write a note on metered dose inhaler.
- 17. Discuss the basis used in development of transdermal drug delivery system.
- 18. Explain about intra-uterine devices.
- 19. Write about production of monoclonal antibodies.
- 20. Discuss the ocular barriers, methods to overcome barriers.
- 21. Explain the approaches used in development of controlled drug delivery systems.
- 22. Explain the formulation considerations of buccal drug delivery system.

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## B. Pharmacy VII-Semester (PCI) (Main) Examination, March 2021

**Subject: Pharmacy Practice** 

Time: 2 Hours Max. Marks: 75

PART - A

Note: Answer any seven questions.

(7 X 3 = 21 Marks)

- 1. Describe the role of clinical pharmacist in health care setting?
- 2. Enumerate the types of drug related problems.
- 3. Mention the requisite Objectives for management of materials in hospital pharmacy.
- 4. Indicate the advantages and disadvantages of Unit Dose Distribution System.
- 5. Provide four examples of TDM drugs with their therapeutic range.
- 6. Give a brief note on Factors which influence drug variability?
- 7. Write a short note on the Material requirement for community pharmacy.
- 8. Define ADR and classify.
- 9. Explain types of drug interactions with example.
- 10. Write a note on rational use of drugs.

PART - B

Note: Answer any one question.

(1 X 14 = 14 Marks)

- 11. Define hospital formulary and elaborate the stepwise procedure involved in the preparation of hospital formulary.
- 12. What is clinical pharmacy? Elucidate functions and responsibility of clinical pharmacy.
- 13. Give a detailed account on the factors affecting Therapeutic Drug Monitoring.

PART - C

Note: Answer any five questions.

(5 X 8 = 40 Marks)

- 14. Explain the roles and responsibility of hospital pharmacist.
- 15. Write down the legal requirements for establishment and maintenance of drug store.
- 16. Enumerate the organization and functions of hospital.
- 17. Explain in detail about the role of pharmacist in medication adherence.
- 18. Define Pharmacy and Therapeutic Committee & explain the objectives, organization and functions.
- 19. Give comprehensive note on the steps involved in patient counseling.
- 20. Define Inventory Control. Specify the methods involved in Inventory Control.
- 21. Describe the procurement or purchasing procedure for pharmacists in detail.
- 22. Explain the various hematologic tests and their significance.

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## B. Pharmacy VII-Semester (PCI) (Main) Examination, March 2021

Subject: Industrial Pharmacy - II

Time: 2 Hours Max. Marks: 75

#### PART - A

Note: Answer any seven questions.

(7 X 3 = 21 Marks)

- 1. What is platform technology?
- 2. Define: (a) Pilot Plant (b) Scale-up.
- 3. 'Technology transfer means physical transfer of goods'. True or false, explain.
- 4. Write the roles of regulatory affairs department.
- 5. Explain the term "Technology transfer".
- 6. Differentiate between IND and NDA.
- 7. Write the applications of Quality by Design.
- 8. What is OOS? How does OOS apply only to finished products?
- 9. Enlist functions of regulatory authorities.
- 10. Write the vision and mission of CDSCO.

#### PART – B

Note: Answer any one question.

(1 X 14 = 14 Marks)

- 11. Explain the process of Change control with the help of flow-chart.
- 12. Discuss the NDA approval process in detail, illustrate with the help of flow diagram.
- Explain the features of finished product technology transfer as per WHO quidelines.

#### PART - C

Note: Answer any five questions.

(5 X 8 = 40 Marks)

- 14. Discuss the stages of pharmaceutical product life-cycle.
- 15. Explain the principles of Good Laboratory Practice (GLP).
- 16. Describe in detail the barriers to technology transfer.
- 17. What is Investigator's Brochure (IB)? Comment on the content of IB.
- 18. Discuss the objectives of pilot plant.
- 19. Explain SUPAC guidelines.
- 20. Write about ISO 9000 series.
- 21. Describe the phases of clinical trials.
- 22. Enlist the key elements of TQM and explain any one of them.

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