

FACULTY OF PHARMACY

**M.PHARMACY (PCI) (Pharma Analysis) II SEMESTER (Backlog)
Examination, June-2026**

SUBJECT: Modern Bio-Analytical Techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions.

(5 x 15 = 75 Marks)

1. Explain the principles and procedures involved in bioanalytical sample preparation methods such as protein precipitation, liquid–liquid extraction, solid phase extraction, and other novel sample preparation approaches
2. Discuss the basic considerations of drug interactions, including pharmacokinetic (PK) and pharmacodynamic (PD) interactions, with special emphasis on Cytochrome P450–based drug interactions.
3. Demonstrate the role of LC–MS in bioactivity screening and proteomics with suitable examples.
4. Explain the principles and applications of in vitro dissolution and drug release testing, along with the experimental methods used for solubility determination
5. Illustrate the biopharmaceutical factors affecting drug bioavailability and explain their impact on drug absorption and therapeutic response
6. Discuss the instrumentation of flow cytometry and explain the various types of cell culture, general procedures for cell culture, cell isolation, subculture, and cryopreservation with suitable diagrams.
7. Elaborate on the design, evaluation, and clinical significance of bioequivalence studies.
8. Explain the design and evaluation of bioequivalence studies with special emphasis on crossover study designs and generic biologics

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FACULTY OF PHARMACY
M. Pharmacy (PCI) II Semester (Pharmaceutical Analysis) (Backlog)
Examination, June 2026

Subject: Advanced Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions.

(5 x 15 = 75)

1. Explain about method development and trouble shooting process in HPLC.
2. a) Discuss about Ion-exchange chromatography?
b) Explain about head space sampling and columns used in Gas chromatography
3. a) Write the principle and applications of Super critical fluid chromatography
b) Explain about characteristics and methods of capillary electrophoresis?
4. Explain about fragmentation modes in mass spectrometry?
5. a) Write about a) spin-spin coupling and b) Relaxation process in NMR?
b) Write in detail about COSY?
6. a) Write about Nano Liquid Chromatography?
b) Discuss in detail about detectors used in Gas chromatography?
7. a) Explain about various parameters used in HPLC.
b) Discuss about 2D NMR.
8. a) Explain about Quadrupole and Time of flight in MS analysis?
b) Write about ¹³C-NMR?

FACULTY OF PHARMACY
M. Pharmacy (PCI) (Pharmaceutical Analysis) II-Semester (Backlog)
Examination, June 2026

Subject: Herbal & Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions.

(5x15=75Marks)

1. (a) Describe WHO guidelines for standardization of herbal drugs.
(b) Discuss the efficacy of herbal medicinal products.
2. (a) Explain the determination of heavy metals, pesticide residues and microbial contamination in herbal formulations?
(b) Define adulteration and explain various types of adulteration of herbal drugs?
3. (a) Explain the effect of herbal medicine on clinical laboratory testing?
(b) Explain with an example the Ayurvedic Pharmacopoeia of India?
4. (a) Explain AYUSH guidelines for safety monitoring of natural medicine.
(b) Write notes on bio food-drug interactions with suitable examples.
5. (a) Explain the procedure involved in determination of saponification value and Iodine value of cosmetic products.
(b) Write a note on analysis of skin care products as per BIS.
6. Discuss the sampling and testing of baby care products and lipsticks as per BIS.
7. Write about Indian patent law applicable for herbal drugs and natural products.
8. Write a note on
 - (a) Global marketing management.
 - (b) Regulation and dispensing of Herbal drugs.

FACULTY OF PHARMACY
M. Pharmacy (PCI) (Pharmaceutical Analysis) II-Semester (Backlog)
Examination, June 2026

Subject: Quality Control and Quality Assurance

Time: 3 Hours

Max Marks: 75

Note: Answer any five Questions.

(5 x 15= 75Marks)

1. Discuss Good Laboratory Practices (GLP) in detail, including scope, principles, and Quality Assurance Unit.
2. Explain cGMP guidelines as per Schedule M with respect to organization, personnel, and facilities.
3. Explain quality control tests for parenterals and ophthalmic products as per pharmacopoeias.
4. Discuss preparation and importance of SOPs, Master Formula Record, and Batch Formula Record.
5. Explain manufacturing operations and controls with emphasis on sanitation and contamination control.
6. Write short notes on:
 - a) Cross-contamination and mix-ups.
 - b) Quality audit plans and reports
7.
 - a) Discuss CPCSEA guidelines.
 - b) Write short notes on purchase specifications and store maintenance.
8.
 - a) Explain report preparation and documentation in GLP.
 - b) Write short notes on training and hygiene of personnel.

FACULTY OF PHARMACY
M. Pharmacy (Phar. Analysis) II - Semester (Main & Backlog)
Examination, December 2025

Subject: Modern Bio analytical techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five following questions.

(5 x 15 = 75 Marks)

1. a) Summarize the general principle and procedure involved in protein precipitation sample treatment methods.
b) Explain the Bioanalytical method validation as per USFDA guidelines.
2. a) Describe the compendial methods of dissolution testing.
b) Explain different experimental methods for solubility determination.
3. a) Discuss about Cytochrome P450 based drug interactions.
b) Explain about different Pharmacokinetic and Pharmacodynamic drug interactions with examples.
4. a) Write about different cell culture media.
b) Discuss different techniques for the characterization of cells along with their applications.
5. a) Write about the clinical significance of Bioequivalence studies.
b) Explain different methods for assessment of the bioavailability of new drug products.
6. a) Write about basic equipment used in cell culture lab.
b) Discuss about Biopharmaceutical factors affecting drug bioavailability.
7. a) Explain about in-vivo and in- vitro methods for checking cellular permeability of new drug products.
b) Explain about principles, instrumentation and applications of flow cytometry.
8. a) Write about Rat liver microsomes and Human Liver microsomes.
b) Discuss about different approaches for quantification of metabolites.

FACULTY OF PHARMACY

**M. Pharm II-Semester (Pharma. Analysis) (PCI) (Main & Backlog)
Examination, December 2025**

Subject: Advanced Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any Five following questions.

(5X15=75 Marks)

1. Write the principle and Instrumentation of HPLC and explain the following parameters (a) Peak shape (b) Capacity factor (c) Resolution.
b) Write about Chiral analysis and explain about enantiomeric separation?
2. a) Discuss about Size-Exclusion Chromatography and Affinity chromatography?
b) Explain about head space sampling and derivitization in Gas chromatography?
3. a) Write the instrumentation and applications of Super critical Fluid Chromatography?
b) Explain about Crown ethers and buffer additives in capillary electrophoresis?
4. Explain the following Ionization techniques in mass spectrometry a) Electron Impact b) Chemical Ionization C) Field Ionization
5. a) What is chemical shift? Explain the various factors influencing it?
b) Write about 2DNMR?
6. a) Write about Preparative chromatography?
b) Discuss the detectors used in Gas chromatography?
7. a) Explain about columns and column problems in HPLC?
b) Discuss the instrumentation of HPLC.
8. a) Explain about LC-MS analysis?
b) Write about a) coupling constant b) Shielding and de shielding in NMR spectroscopy?

FACULTY OF PHARMACY

M. Pharmacy (PCI) (Pharma Analysis) II - Semester (Main & Backlog) Examination, Dec 2025

Subject: Herbal and Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any Five questions.

(5 x 15=75 Marks)

1. (a) Discuss the standardization of herbal drugs according to AYUSH guidelines. [8]
(b) Explain validation of herbal therapies in detail. [7]
2. Discuss the protocol of Indian patent law as applicable to herbal drugs and natural products. [15]
3. (a) Write a note on Stability testing of natural products. [10]
(b) Explain the effect of herbal medicine on clinical laboratory testing. [5]
4. Explain the spontaneous reporting schemes for bio drug adverse reactions and bio drug-food interactions with suitable examples. [15]
5. (a) Explain the procedure involved in determination of acid value and Iodine value of cosmetic products. [8]
(b) Discuss types of adulteration of herbal drugs? [7]
6. Discuss the sampling and testing of baby care products and lipsticks as per BIS. [15]
7. (a) Explain WHO guidelines for safety monitoring of natural medicine. [10]
(b) Explain the challenges in monitoring the safety of herbal medicine. [5]
8. Compare the herbal drug monographs of IP and Ayurvedic parhacopoeias? [15]

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FACULTY OF PHARMACY
M. Pharmacy (Pharma. Analysis) II-Semester (PCI) (Main & Backlog)
Examination, December 2025

Subject: Quality control and Quality assurance

Time: 3 Hours

Max Marks: 75

Note: Answer any Five Questions.

(5X15=75 Marks)

1. Write a short note on the following
 - (a) Quality control. (5)
 - (b) Quality assurance. (5)
 - (c) Non clinical testing. (5)
2. Explain the various CPSCEA guidelines for laboratory animal facility. (15)
3. Define IPQC. Explain in detail about various IPQC tests for
 - (a) Capsules. (8)
 - (b) Parenterals. (7)
4. Give a brief note on
 - (a) Quality audit plan. (5)
 - (b) Protocols and reports. (5)
 - (c) Distribution records. (5)
5. Discuss the Good laboratory practices for a quality control laboratory in detail. (15)
6. (a) Explain the various documents to be maintained by the quality control department. (7)
(b) Explain Master formula and Batch formula records. (8)
7. Explain various cGMP guidelines according to schedule M. (15)
8. Write a note on
 - (a) Sanitation of manufacturing premises (5)
 - (b) Drug product inspection. (5)
 - (c) Production record review. (5)

FACULTY OF PHARMACY

M. Pharmacy (PCI) II - Semester (Pharm. Analysis) (Backlog) Examination, June 2025

Subject: Modern Bioanalytical techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks. (5 x 15 = 75 Marks)

1. (a) Explain the general principle and procedures involved in extraction of drugs from biological matrices by Solid phase extraction method. (10)
(b) Write about Liquid-Liquid extraction as sample preparation technique. (5)
2. (a) Explain the Bioanalytical method validation as per USFDA guidelines. (10)
(b) Explain the advantages and disadvantages of membrane filtration. (5)
3. (a) What is Bioavailability? Discuss the Biopharmaceutical Factors affecting drug Bioavailability. (10)
(b) Write about different cell culture media. (5)
4. (a) Mention the different alternative methods of dissolution testing. (10)
(b) Define solubility & permeability based on biopharmaceutics classification system. (5)
5. (a) Explain different methods for assessment of bioavailability of new drug product. (10)
(b) Write the clinical significance of bioequivalence studies. (5)
6. Write notes on the following
(a) Cytochrome P450 drug interactions. (8)
(b) Cryopreservation techniques. (7)
7. Discuss about the design and evaluation of bioequivalence studies. (15)
8. Write brief notes on
(a) Cell viability assays (8)
(b) LC-MS in bioactivity screening and proteomics (7)

FACULTY OF PHARMACY

M. Pharmacy (PCI) II - Semester (Pharm. Analysis) (Backlog) Examination, June 2025

Subject: Advanced Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

(5 x 15 = 75 Marks)

1. (a) Explain about various parameters in HPLC.
 - (i) Peak shape
 - (ii) Capacity factor
 - (iii) Plate number and plate height
 - (iv) Resolution.(b) Write about Chiral analysis?
2. (a) Discuss about Size-Exclusion Chromatography and Affinity chromatography?
(b) Explain about head space sampling in Gas chromatography.
3. (a) Write the instrumentation and applications of SFC.
(b) Explain about Crown ethers and buffer additives in capillary electrophoresis.
4. Explain about Electron impact, CI, FAB, ESI Ionization techniques in mass spectrometry.
5. (a) What is chemical shift? Explain the various factors influencing it?
(b) Write about 2DNMR.
6. (a) Write about Preparative chromatography.
(b) Discuss the derivatization methods of Gas chromatography.
7. (a) Explain about columns and column problems in HPLC.
(b) Discuss about NOESY.
8. (a) Explain about LC-MS analysis?
(b) Write about
 - (i) Coupling constant
 - (ii) Shielding and deshielding in NMR spectroscopy

FACULTY OF PHARMACY

M. Pharmacy (Pharma Analysis) II - Semester (PCI) (Backlog) Examination, June 2025
Subject: Herbal and Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. (a) Discuss the standardization of herbal drugs according to AYUSH guidelines. [10]
(b) Explain the pharmacokinetic issues of herbal drugs. [5]
2. Discuss the protocol of Indian patent law as applicable to herbal drugs and natural products. [15]
3. (a) Write a note on Stability testing of natural products. [10]
(b) Explain the effect of herbal medicine on clinical laboratory testing. [5]
4. (a) Describe the spontaneous reporting schemes for bio drug adverse reactions? [8]
(b) Write a note on bio drug-food interactions with suitable examples? [7]
5. (a) Explain the procedure involved in determination of acid value and Iodine value of cosmetic products. [8]
(b) Write the analysis of Skin creams preparations as per BIS. [7]
6. Discuss the sampling and testing of baby care products and lipsticks as per BIS. [15]
7. (a) Explain AYUSH guidelines for safety monitoring of natural medicine. [10]
(b) Explain the challenges in safety monitoring of herbal drugs. [5]
8. Explain Siddha and Unani pharmacopoeias. [15]

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FACULTY OF PHARMACY

M. Pharmacy (PCI) II - Semester (Pharm. Analysis) (Backlog) Examination, June 2025

Subject: Quality control and Quality assurance

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks. (5 x 15 = 75 Marks)

1. (a) Differentiate Quality control and Quality assurance (3)
(b) Discuss Good laboratory practices for quality control laboratory in detail. (12)
2. (a) Write a short note on Pharmaceutical inspection convention (5)
(b) Explain the various CPCSEA guidelines for laboratory animal facility (10)
3. (a) Define IPQC. (2)
(b) Explain in detail IPQC tests for Ophthalmic products. (13)
4. (a) Write a short note on SOP (5)
(b) Explain Master formula and Batch formula records (10)
5. (a) How do you calculate Expiry date (5)
(b) Write about Packaging operation (5)
(c) Add a brief note on Production record review (5)
6. (a) Add a brief note on QSEM (5)
(b) Write in detail about ICH Q series guidelines (10)
7. (a) Define and classify Packaging. (5)
(b) Explain the Quality control tests for Glass as packaging material. (10)
8. (a) Write about Organization and personal responsibilities. (5)
(b) Explain Quality audit plan (5)
(c) Add a note on Electronic data (5)

FACULTY OF PHARMACY

**M. Pharmacy (PCI) II - Semester (Pharm. Analysis) (Main & Backlog) Examination,
December 2024**

Subject: Advanced Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks. (5 x 15 = 75 Marks)

1. Explain about method development and trouble shooting process in HPLC.
2. (a) Discuss about Ion-exchange chromatography?
(b) Explain about head space sampling and columns used in Gas chromatography.
3. (a) Write the principle and applications of Super critical fluid chromatography.
(b) Explain about characteristics and methods of capillary electrophoresis.
4. Explain about fragmentation modes in mass spectrometry?
5. (a) Write about
 - (i) spin-spin coupling and
 - (ii) Relaxation process in NMR
(b) Write in detail about COSY.
6. (a) Write about Nano Liquid Chromatography.
(b) Discuss in detail about detectors used in Gas chromatography.
7. (a) Explain about various parameters used in HPLC.
(b) Discuss about 2D NMR.
8. (a) Explain about Quadrupole and Time of flight in MS analysis?
(b) Write about ^{13}C -NMR?

FACULTY OF PHARMACY
M. Pharmacy (PCI) II - Semester (Pharm. Analysis) (Main & Backlog) Examination,
December 2024
Subject: Modern Bioanalytical techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks. (5 x 15 = 75 Marks)

1. (a) Explain the general principle and procedures involved in extraction of drugs from biological matrices by liquid-liquid extraction method. (10)
(b) Write a note on protein precipitation method. (5)
2. (a) Explain different validation parameters for bioanalytical methods according to USFDA guidelines. (10)
(b) Write a note on SPE sorbents. (5)
3. (a) Discuss Biopharmaceutical factors affecting drug bioavailability. (10)
(b) Write the Biopharmaceutics classification system defined by FDA. (5)
4. (a) Explain about different Pharmacokinetic and Pharmacodynamic drug interactions with examples. (10)
(b) Write the importance and applications of Toxicokinetic studies. (5)
5. (a) Write about principles, instrumentation, and applications of flow cytometry. (9)
(b) Write about cryopreservation and storage of cells. (6)
6. (a) Write about in-vivo and in- vitro methods for checking the cellular permeability of new drug products. (8)
(b) Discuss about Cytochrome P450 based drug interactions. (7)
7. (a) Explain different study designs in bioequivalence studies. (10)
(b) Differentiate absolute and relative bioavailability with illustrative examples and equations. (5)
8. (a) Write about Rat liver microsomes and Human Liver microsomes. (5)
(b) Discuss about different approaches for identification of metabolites. (10)

FACULTY OF PHARMACY

**M. Pharmacy II - Semester (PCI) (Pharma Analysis) (Main & Backlog) Examination,
December 2024**

Subject: Herbal and Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. (a) Discuss the pharmacokinetic and pharmacodynamics issues related to herbal drugs. [10]
(b) Differentiate between herbal drugs and conventional drugs. [5]
2. (a) Explain the determination of pesticide residues and microbial contamination in herbal formulations? [8]
(b) Define adulteration and explain various types of adulteration of herbal drugs? [7]
3. (a) Explain DNA Finger printing techniques in identification of drugs of natural origin? [7]
(b) Explain with an example the Ayurvedic Pharmacopoeia of India? [8]
4. (a) Explain WHO guidelines for safety monitoring of natural medicine. [10]
(b) Write notes on bio drug-drug interactions with suitable examples. [5]
5. (a) Explain the Indian standard specification laid down for sampling and testing of dental products. [8]
(b) Write a note on analysis of Lipsticks as per BIS. [7]
6. Write notes on
(a) Explain the Comparative study of IP and USP with an example? [10]
(b) Determination of Saponification value of cosmetic products. [5]
7. Write about International patent law applicable for herbal drugs and natural products. [15]
8. Discuss the quality of raw materials and general methods of analysis of raw materials used in cosmetic manufacture as per BIS? [15]

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FACULTY OF PHARMACY

**M. Pharmacy (PCI) II - Semester (Pharm. Analysis) (Main & Backlog) Examination,
December 2024**

Subject: Quality control and Quality assurance

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks. (5 x 15 = 75 Marks)

1. (a) Write about Total quality management. (5)
(b) Describe concept and components of Quality control and Quality assurance. (10)
2. (a) Write about CDER and CBER. (5)
(b) Write a detailed note on requirements and guidelines of GMP (schedule M) in Pharma industries. (10)
3. (a) Define IPQC. (2)
(b) Explain in detail IPQC tests for Tablets. (13)
4. (a) Write a short note on good documentation practice guidelines. (5)
(b) What are the different types of audits? Explain in detail about audit methods and techniques involved in it. (10)
5. (a) Write about mix-up and cross contamination. (5)
(b) Add a note on Processing of intermediates and bulk products. (5)
(c) Explain Aseptic process control. (5)
6. Write about the following
(a) Protocol for conduct of non-clinical testing (5)
(b) Quality control of creams (5)
(c) Calculation of yields (5)
7. (a) Explain Master formula and Batch formula records. (10)
(b) Write a short note on SOP. (5)
8. (a) Discuss Good laboratory practices for quality control laboratory in detail. (12)
(b) Add a note on Electronic data. (3)

FACULTY OF PHARMACY

**M. Pharmacy (Pharma. Analysis) II - Semester (PCI) (Backlog) Examination,
June 2024**

Subject: Quality control and Quality assurance

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. Write a detailed note on requirements and guidelines of GMP (schedule M) in Pharma industries? [15]
2. Write a short note on the following
 - (a) Quality control. [5]
 - (b) Quality assurance. [5]
 - (c) Non clinical testing. [5]
3. Define IPQC. Explain in detail about various IPQC tests for
 - (a) Tablets [8]
 - (b) Ophthalmics [7]
4. Explain
 - (a) Batch formula Record [8]
 - (b) Master formula Record [7]
5. Write a short note on the following
 - (a) Expiry date calculation [5]
 - (b) Limitations of production [5]
 - (c) Calculation of yields [5]
6. Explain the various CPCSEA (CCSEA – New non enclosure) guidelines for laboratory animal facility. [15]
7. Describe the quality control test for containers, closures and secondary packing materials? [15]
8. Write a note on
 - (a) Sanitation of manufacturing premises [5]
 - (b) Drug product inspection. [5]
 - (c) Production record review. [5]

FACULTY OF PHARMACY

**M. Pharmacy (Pharm. Analysis) II-Semester (PCI) (Backlog) Examination,
June 2024**

Subject: Modern Bio Analytical Techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks. (5 x 15 = 75 Marks)

1. (a) Write about the following sample preparation techniques. [6]
(i) Solid phase extraction
(ii) Liquid liquid extraction
(b) Explain the Bioanalytical method validation as per USFDA guidelines. [9]
2. (a) Discuss Biopharmaceutical factors affecting drug bioavailability. [10]
(b) Write the Biopharmaceutics classification system defined by FDA. [5]
3. (a) What is enzyme inhibition? Discuss drug interactions due to enzyme inhibition with examples. [7]
(b) Discuss drug-protein binding interaction with examples. [8]
4. (a) Write about principles, instruments, and applications of flow cytometry. [9]
(b) Write about cryopreservation and storage of cells. [6]
5. (a) Explain different study designs in bioequivalence studies. [10]
(b) Differentiate absolute and relative bioavailability with illustrative examples and equations. [5]
6. (a) Discuss the importance and applications of Toxicokinetic studies. [8]
(b) Write about the basic equipment used in the cell culture lab. [7]
7. (a) Discuss different approaches for the identification of metabolites. [10]
(b) Write a short note on the clinical significance of bioequivalence studies. [5]
8. (a) Describe the compendial methods of dissolution testing. [7]
(b) Write about in-vivo and in-vitro methods for checking the cellular permeability of new drug products. [8]

FACULTY OF PHARMACY
M. Pharmacy (Pharma. Analysis) II-Semester (PCI) (Backlog) Examination,
June 2024

Subject: Advanced Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. (a) Explain about method development and trouble shooting in HPLC.
(b) Write about Chiral analysis of Pharmaceuticals using HPLC
2. (a) Discuss about Ion-exchange chromatography?
(b) Explain about head space sampling and columns used in Gas chromatography
3. (a) Write the principle and applications of Super critical fluid chromatography
(b) Explain about characteristics and methods of capillary electrophoresis?
4. Explain about fragmentation modes in mass spectrometry?
5. (a) Write about a) spin-spin coupling and b) relaxation process in NMR?
(b) Write in detail about COSY?
6. (a) Write about Nano Liquid Chromatography?
(b) Discuss in detail about detectors used in Gas chromatography?
7. (a) Explain about various parameters used in HPLC.
(b) Discuss about 2D NMR.
8. (a) Explain about Quadrupole and Time of flight in MS analysis.
(b) Write about ^{13}C -NMR?

FACULTY OF PHARMACY

M. Pharmacy (Pharm Analysis) II - Semester (PCI) (Backlog) Examination, June 2024

Subject: Herbal and Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions.

(75 Marks)

1. (a) How Herbal medicines are differentiated from Conventional Drugs?
(b) Discuss about Standardization of Herbal drugs as per WHO guidelines.
2. (a) What is Adulteration? Write about different types of Adulteration with suitable examples.
(b) Explain the procedure involved in determination of foreign matter pesticide residue in Herbal drugs.
3. (a) Discuss on Adulterant screening using advanced Analytical Techniques.
(b) Give the protocol for Stability Testing of natural products.
4. (a) Explain bio-drug drug interactions with suitable examples.
(b) Write notes on challenges in monitoring the safety of Herbal Medicines.
5. Write the procedure involved in determination of
(a) Acid value
(b) Moisture Content
6. Write short notes on
(a) Validation of Herbal Therapies.
(b) Global Marketing Management of Herbal Drugs
7. (a) Compare the monographs of Herbal Dugs mentioned in different Pharmacopoeia.
(b) Explain the determination of Saponification Value.
8. (a) Explain the general methods of analysis of raw materials used in cosmetics manufacturing as per BIS.
(b) Brief out the testing of baby care products.

FACULTY OF PHARMACY

**M. Pharmacy (Pharm. Analysis) II Semester (PCI) (Main & Backlog) Examination,
October 2023**

Subject: Modern Bio Analytical Techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.(5 x 15 = 75 Marks)

1. (a) Explain different sample preparation approaches involved in bioanalytical methods. [10]
(b) Explain the following validation parameters in bioanalytical method validation as per USFDA guidelines. [5]
(i) Linearity
(ii) Recovery studies
2. (a) Describe the compendial methods of dissolution testing. [8]
(b) Write about different experimental methods for solubility determination. [7]
3. (a) Discuss drug-protein binding interaction with examples. [8]
(b) What is enzyme induction? Discuss drug interaction due to enzyme induction. [7]
4. (a) Write about the basic equipment used in the cell culture lab. [7]
(b) Describe different techniques for the characterization of cells along with their applications. [8]
5. (a) Write about the clinical significance of Bioequivalence studies. [5]
(b) Explain different methods for assessment of the bioavailability of new drug products. [10]
6. (a) Discuss Biopharmaceutical factors affecting drug bioavailability. [10]
(b) Write about cryopreservation and storage of cells. [5]
7. (a) Discuss different approaches for the quantification of metabolites. [9]
(b) Write about different cell culture media. [6]
8. (a) Write about in-vivo and in-vitro methods for checking the cellular permeability of new drug products. [9]
(b) Write in brief about drug interactions linked to transporters. [6]

FACULTY OF PHARMACY

**M. Pharmacy (Pharma. Analysis) II-Semester (PCI) (Main & Backlog) Examination,
November 2023**

Subject: Advanced Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. (a) Explain about various parameters in HPLC.
(i) Peak shape (ii) Capacity factor
(iii) Plate number and plate height (iv) Resolution.

(b) Write about Preparative HPLC.
2. (a) Discuss about Size-Exclusion Chromatography and Affinity chromatography?
(b) Explain about head space sampling in Gas chromatography
3. (a) Write the instrumentation and applications of SFC.
(b) Explain about Crown ethers and buffer additives in capillary electrophoresis?
4. Explain about Electron impact, CI, FAB, ESI Ionization techniques in mass spectrometry?
5. (a) What do you mean by chemical shift? Explain the various factors influencing it?
(b) Write about 2DNMR?
6. (a) Write about Chiral Chromatography?
(b) Discuss the derivatization methods of Gas chromatography?
7. (a) Explain about columns and column problems in HPLC?
(b) Discuss about NOESY.
8. (a) Explain about LC-MS analysis?
(b) Write about (i) coupling constant (ii) LC-NMR?

FACULTY OF PHARMACY

**M. Pharmacy II Semester (Ph. Analysis) (PCI) (Main & Backlog) Examination,
November 2023**

Subject: Herbal & Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. (a) Discuss the standardization of herbal drugs according to WHO guidelines. [10]
(b) Differentiate between herbal drugs and conventional drugs. [5]
2. (a) Explain the determination of pesticide residues and microbial contamination in herbal formulations? [8]
(b) Write a note on Global marketing management? [7]
3. (a) Discuss adulterant screening of herbal drugs using HPLC? [7]
(b) Explain with an example the Ayurvedic Pharmacopoeia of India? [8]
4. (a) Explain WHO guidelines for safety monitoring of natural medicine. [10]
(b) Write notes on bio drug-food interactions with suitable examples. [5]
5. (a) Explain the Indian standard specification laid down for sampling and testing of dental products. [8]
(b) Write a note on analysis of skin creams as per BIS. [7]
6. Write notes on
(a) Global marketing management. [6]
(b) Determination of Acid value of cosmetic products. [4]
(c) Analysis of dental preparations. [5]
7. Write about Indian patent law applicable for herbal drugs and natural products. [15]
8. Discuss the quality of raw materials and general methods of analysis of raw materials used in cosmetic manufacture as per BIS? [15]

FACULTY OF PHARMACY

**M. Pharmacy (Pharma. Analysis) II Semester (PCI) (Main & Backlog) Examination,
November 2023**

Subject: Quality Control and Quality Assurance

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. (a) Write in detail about ICH Q series guidelines. [8]
(b) Explain about Quality control and Quality assurance. [7]
2. Write about the following
(a) Organization and personnel responsibilities. [5]
(b) Maintenance of sterile areas. [5]
(c) Personal records and environmental control. [5]
3. Define IPQC. Write in detail about different IPQC tests for tablets and parenterals. [15]
4. (a) What is SOP? Write about different techniques to write SOP. [8]
(b) Write a note on Quality audit plan. [7]
5. (a) Write about mix-up and cross contamination. [8]
(b) Explain about Expiry date calculation and calculation of yields. [7]
6. Explain various quality control tests for Glass as a packaging material. [15]
7. (a) Write a note on Production record review. [7]
(b) Aseptic process control. [8]
8. Discuss Good laboratory practices for quality control laboratory in detail. [15]

FACULTY OF PHARMACY

M. Pharmacy (Pharma Analysis) II-Semester (PCI) (Backlog) Examination, April / May 2023

SUBJECT: Modern Bio Analytical Techniques

Time: 3 Hours

Max Marks: 75

Note: Answer Any Five Questions. ALL Questions carry Equal Marks.

1. a) Write about solid phase extraction technique. [5]
b) Explain the different validation parameters in bio-analytical method validation as per USFDA guidelines. [10]
2. a) What is Bioavailability? Give the Biopharmaceutical Factors affecting drug Bioavailability. [10]
b) Write the Biopharmaceutics classification system defined by FDA. [5]
3. a) Explain about different Pharmacokinetic and Pharmacodynamic drug interactions with examples. [10]
b) Write the importance and applications of Toxicokinetic studies. [5]
4. a) Write about principles, instrumentation and applications of flow cytometry. [9]
b) Write about basic equipment used in cell culture lab. [6]
5. a) Explain different study designs in bioequivalence studies. [10]
b) Write the clinical significance of Bioequivalence studies. [5]
6. a) Describe the principles and applications of Cell viability assays. [8]
b) Write about Rat liver microsomes and Human Liver microsomes. [7]
7. a) Discuss about different approaches for identification of metabolites. [10]
b) Differentiate absolute and relative bioavailability with illustrative examples and equations. [5]
8. a) Describe the compendial methods of dissolution testing. [7]
b) Write about *in-vivo* and *in-vitro* methods for checking cellular permeability of new drug products. [8]

Code No: E-12251PCI

FACULTY OF PHARMACY

**M. Pharmacy (Pharma Analysis) II-Semester (PCI) (Backlog) Examination,
May 2023**

SUBJECT: Herbal and Cosmetic Analysis

Time: 3 Hours

Max Marks: 75

Note: Answer any five questions. All questions carry equal Marks

1. a) Explain validation of herbal therapies in detail. [10]
b) Compare herbal drugs with conventional drugs. [5]
2. a) Explain HPTLC as a DNA finger printing technique for the identification of Herbal drugs. [10]
b) Write notes on pesticide residue determination in herbal drugs. [5]
3. a) Write informative notes on stability testing of natural products. [8]
b) Compare the herbal drug monographs of IP and USP. [7]
4. a) Write the spontaneous reporting schemes for bio drug adverse reactions and bio drug-food interactions. [10]
b) Explain the challenges in monitoring the safety of herbal medicines. [5]
5. a) Explain the procedure involved in determination of acid value and iodine value of cosmetic products. [5]
b) Briefly write about Indian standard specification laid down for sampling of herbal drugs. [5]
6. Discuss the testing of skin care products in detail. [15]
7. Write notes on [3x5=15]
 - a) Efficacy of herbal medicine products
 - b) Causes of adulteration
 - c) General methods of analysis of raw materials used in cosmetics preparation.
8. Discuss on Indian patent law applicable for herbal drugs and natural products. [15]

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Code No: E-12250/PCI

FACULTY OF PHARMACY

**M. Pharmacy (Pharma Analysis) II Semester (PCI) (Backlog) Examination,
April / May 2023**

Subject: Quality Control and Quality Assurance

Time: 3 Hours

Max Marks: 75

Note: Answer Any Five Questions. ALL Questions carry Equal Marks.

1. Write a short note on the following
 - a) Quality control [5]
 - b) Quality assurance. [5]
 - c) Non clinical testing. [5]
2. Explain the various CPCSEA guidelines for laboratory animal facility. [15]
3. Define IPQC. Explain in detail about various IPQC tests for
 - a) Capsules. [8]
 - b) Parenterals. [7]
4. Give a brief note on
 - a) Quality audit plan [5]
 - b) Protocols and reports. [5]
 - c) Distribution records. [5]
5. Discuss the Good laboratory practices for a quality control laboratory in detail. [15]
6.
 - a) Explain the various documents to be maintained by the quality control department. [7]
 - b) Explain Master formula and Batch formula records. [8]
7. Explain various cGMP guidelines according to schedule M.
8. Write a note on
 - a) Sanitation of manufacturing premises. [5]
 - b) Drug product inspection. [5]
 - c) Production record review. [5]

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Code No: E-12248/PCI

FACULTY OF PHARMACY

**M. Pharmacy (Pharma Analysis) II-Semester (PCI) (Backlog) Examination,
May 2023**

SUBJECT: Advanced Instrumental Analysis

Time: 3 Hours

Max Marks: 75

Note: Answer any five questions. All questions carry equal marks.

(5 x 15 = 75 Marks)

1. a) Explain about various parameters in HPLC
(i) Selectivity (ii) Capacity factor (iii) Plate number plate height (iv) Resolution
b) Write about UPLC
2. a) Discuss about Size-Exclusion Chromatography and Affinity chromatography?
b) Explain about head space sampling in Gas chromatography
3. a) Write the instrumentation and applications of SFC
b) Explain about Crown ethers and buffer additives in capillary electrophoresis?
4. Explain about Electron impact, CI, FAB, ESI Ionization techniques in mass spectrometry?
5. a) What do you mean by chemical shift? Explain the various factors influencing it?
b) Write about 2DNMR?
6. a) Write about Chiral Chromatography?
b) Discuss the derivatization methods of Gas chromatography?
7. a) Explain about columns and column problems in HPLC?
b) Discuss about NOESY
8. a) Explain about LC-MS analysis?
b) Write about a) coupling constant b) LC-NMR?

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FACULTY OF PHARMACY

M. Pharmacy (Pharma Analysis) II Semester (PCI) (Main & Backlog)

Examination, December 2022

SUBJECT: Modern Bio Analytical Techniques

Time: 3 Hours

Max Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. a) Write the general principle and procedure involved in protein precipitation methods. [5]
b) Explain the Bioanalytical method validation as per USFDA guidelines. [10]
2. a) Describe the compendial methods of dissolution testing. [8]
b) Write about different experimental methods for solubility determination. [7]
3. a) Discuss about Cytochrome P450 based drug interactions. [7]
b) What is enzyme inhibition? Discuss about drug interactions due to enzyme inhibition with examples. [8]
4. a) Write about cryopreservation and storage of cells. [6]
b) Describe different techniques for characterization of cells along with their applications. [9]
5. a) Discuss in detail about Bioequivalence protocol. [10]
b) Write about clinical significance of Bioequivalence. [5]
6. a) Write about basic equipments used in cell culture lab. [6]
b) Discuss about Biopharmaceutical factors affecting drug bioavailability. [9]
7. a) Discuss about different approaches for quantification of metabolites. [9]
b) Write about different cell culture media. [6]
8. a) Write about *in-vivo* and *in-vitro* methods for checking cellular permeability of new drug products. [9]
b) Write in brief about drug interactions linked to transporters. [6]

FACULTY OF PHARMACY

M. Pharmacy (Pharma Analysis) II - Semester (PCI) (Main & Backlog)

Examination, December 2022

Subject: Herbal and Cosmetic Analysis

Time: 3 Hours

Max Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. a) Discuss Pharmacodynamic & Pharmacokinetic issues of Herbal drugs. [10]
b) How can we differentiate herbal drugs from conventional drugs? [5]
2. a) Explain the determination of heavy metals in herbal drugs. [8]
b) Write notes on global marketing management trends of Herbal drugs. [7]
3. a) Discuss HPLC as modern technique of adulterant screening of Herbal drugs. [8]
b) Write notes on different herbal pharmacopeia. [7]
4. a) Explain AYUSH guidelines for safety monitoring of natural medicine. [7]
b) Write notes on bio drug-food interactions with suitable examples. [8]
5. a) Explain the Indian standard specification laid down for sampling and Testing of baby care products. [10]
b) Write the tests for lip sticks. [5]
6. Write notes on [3 x 5 = 15]
 - a) Efficacy of Herbal medicine products.
 - b) Determination of foreign matter in herbal drugs
 - c) Challenges in safety monitoring of herbal drugs.
7. Discuss the determination of peroxide value and moisture content in herbal drugs. [7.5 + 7.5 = 15]
8. Explain the testing procedures for hair products and skin creams [7.5 + 7.5 = 15]

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FACULTY OF PHARMACY

M. Pharmacy (Pharma Analysis) II - Semester (PCI) (Main & Backlog)

Examination, December 2022

SUBJECT: Advanced Instrumental Analysis

Time: 3 Hours

Max Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. a) Explain about trouble shooting in HPLC
b) Write about preparative HPLC?
2. a) Discuss about Ion-exchange chromatography?
b) Explain about head space sampling and columns used in Gas chromatography
3. a) Write the principle and applications of Super critical fluid chromatography
b) Explain about characteristics and methods of capillary electrophoresis?
4. Explain about fragmentation modes in mass spectrometry?
5. a) Write about a) Spin-spin coupling and b) Relaxation process in NMR?
b) Write in detail about COSY?
6. a) Write about Nano Liquid Chromatography?
b) Discuss in detail about detectors used in Gas chromatography?
7. a) Explain about various parameters in HPLC?
b) Discuss about meta stable ions in Mass spectrometry.
8. a) Explain about Quadrupole and Time of flight in MS analysis?
b) Write about ^{13}C -NMR?

FACULTY OF PHARMACY

M. Pharmacy (Pharma. Analysis) II Semester (PCI) (Main & Backlog)

Examination, December 2022

Subject: Quality Control and Quality Assurance

Time: 3 Hours

Max Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1) a) Explain about Quality control and Quality assurance. [8]
b) Write in detail about Total Quality management. [7]
- 2) a) Explain the controls on environmental pollution. [8]
b) Explain the maintenance of sterile areas [7]
- 3) Write in detail about inprocess quality control (IPQC) testing of Tablets and Parenterals. [15]
- 4) a) Explain the various documents to be maintained by the quality control department. [7]
b) Explain Master formula and Batch formula records. [8]
- 5) Discuss about
a) Mix-up's and cross contamination. [8]
b) Aseptic process control. [7]
- 6) Discuss the Good laboratory practices for a quality control laboratory in detail. [15]
- 7) Explain the following
a) Non clinical testing. [5]
b) Controls on animal house. [5]
c) Report Preparation. [5]
- 8) Explain various quality control tests for Glass as a packaging material. [15]

FACULTY OF PHARMACY
M. Pharmacy (Pharmaceutical Analysis) II Semester (PCI) (Supply)
Examination, May 2022

Subject: Herbal and Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five of the following questions.

- 1 (a) Write differences between herbal and conventional drugs.
(b) Discuss the standardization of herbal drugs according to AYUSH guidelines.
- 2 (a) What is adulteration? Explain types with suitable examples.
(b) Write notes on DNA finger printing technique used for identification of drugs on natural origin.
- 3 (a) Describe adulterant screening using modern analytical techniques.
(b) Write a note on effect of herbal medicine on clinical laboratory testing.
- 4 (a) Write the spontaneous reporting schemes for bio drug adverse reactions and bio drug-food interactions.
(b) Explain the challenges in monitoring the safety of herbal medicines.
- 5 (a) Explain the procedure involved in determination of acid value of cosmetic products.
(b) Discuss the sampling and testing of baby care products as per BIS.
- 6 Write the analysis of personal hygiene preparations as per BIS.
- 7 Write notes on:
(a) Causes of adulteration
(b) Monographs of herbal drugs
(c) Determination of saponification value of cosmetic products.
- 8 Write about Indian patent law applicable for herbal drugs and natural products.

FACULTY OF PHARMACY
M. Pharmacy (Pharmaceutical Analysis) II Semester (PCI) (Supply)
Examination, May 2022

Subject: Advanced Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 (a) Explain about various parameters in HPLC.
(a) Peak shape (b) Capacity factor
(c) Plate number and plate height (d) Resolution.
(b) Write about Preparative HPLC.
- 2 (a) Discuss about Size-Exclusion Chromatography and Affinity chromatography.
(b) Explain about derivatization in Gas chromatography.
- 3 (a) Write the instrumentation of SFC.
(b) Explain about Crown ethers and buffer additives in capillary electrophoresis.
- 4 Explain about different types of Ionization techniques and analyzers in mass spectrometry.
- 5 (a) What do you mean by chemical shift? Explain the various factors influencing it.
(b) Write about NOESY.
- 6 (a) Write about Ultra Liquid Chromatography.
(b) Discuss the principle and Instrumentation of Gas chromatography.
- 7 (a) Explain about columns and column problems in HPLC.
(b) Discuss about C13 NMR.
- 8 (a) Explain about DART-MS analysis.
(b) Write about (a) coupling constant (b) Nuclear magnetic double resonance.

FACULTY OF PHARMACY
M. Pharmacy (Pharmaceutical Analysis) II Semester (PCI) (Supply)
Examination, May 2022

Subject: Modern Bio Analytical Techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 (a) Write the general principle and procedure involved in protein precipitation method.
(b) Explain the Bioanalytical method validation as per USFDA guidelines.
- 2 (a) Describe the compendial methods of dissolution testing.
(b) Write about different experimental methods for solubility determination.
- 3 (a) Discuss about Cytochrome P450 based drug interactions.
(b) Write about clinical significance of Bioequivalence studies.
- 4 (a) Write about cryopreservation and storage of cells.
(b) Describe different techniques for characterization of cells along with their applications.
- 5 (a) Discuss in detail about Bioequivalence protocol.
(b) Write about clinical significance of Bioequivalence studies.
- 6 (a) Write about equipment used in cell culture lab.
(b) Discuss about Biopharmaceutical factors affecting drug bioavailability.
- 7 (a) Discuss about different approaches for quantification of metabolites.
(b) Write about different cell culture media.
- 8 (a) Write about in-vivo and in-vitro methods for checking cellular permeability of new drug products.
(b) Write in brief about drug interactions linked to transporters.

FACULTY OF PHARMACY
M. Pharmacy (Pharmaceutical Analysis) II Semester (PCI) (Supply)
Examination, May 2022

Subject: Quality Control and Quality Assurance

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 Write a short note on the following:
 - (a) Quality control
 - (b) Quality Assurance
 - (c) Non clinical testing.
- 2 Explain the various CPSCEA guidelines for laboratory animal facility.
- 3 Define IPQC. Explain in detail about various IPQC tests for
 - (a) Capsules
 - (b) Parenterals.
- 4 Give a brief note on:
 - (a) Quality audit plan
 - (b) Protocols and reports
 - (c) Distribution records.
- 5 Discuss the Good laboratory practices for a quality control laboratory in detail.
- 6
 - (a) Explain the various documents to be maintained by the quality control department.
 - (b) Explain Master formula and Batch formula records.
- 7 Explain various CGMP guidelines according to schedule M.
- 8 Write a note on:
 - (a) Sanitation of manufacturing premises.
 - (b) Drug product inspection.
 - (c) Production record review.

FACULTY OF PHARMACY

**M. Pharmacy (Pharmaceutical Analysis) II Semester (PCI) (Main & Backlog)
Examination, December 2021**

Subject: Quality Control and Quality Assurance

Time: 2 Hours

Max. Marks: 75

Note: Answer any three of the following questions. (3 x 25 = 75 Marks)

- 1 Describe the concept, components of Quality control and Quality assurance.
- 2 Explain the various CPSCEA guidelines for laboratory animal facility.
- 3 Define IPQC explain in detail various IPQC tests for
 - (a) Tablets.
 - (b) Ointments.
- 4 Write a brief note on:
 - (a) Quality audit plan.
 - (b) Batch formula record.
- 5 Write a note on:
 - (a) Sanitation of manufacturing premises.
 - (b) Drug product inspection.
 - (c) Production record review.
- 6 Describe sources of contamination and methods of contamination control.
- 7 Write in detail about
 - (a) SOP
 - (b) Protocols and reports.
- 8 Discuss the Good laboratory practices for a quality control laboratory in detail.

FACULTY OF PHARMACY
M. Pharmacy (Pharmaceutical Analysis) II Semester (PCI) (Main & Backlog)
Examination, November 2021

Subject: Modern Bio Analytical Techniques

Time: 2 Hours

Max. Marks: 75

Note: Answer any three of the following questions. (3 x 25 = 75 Marks)

- 1 (a) Explain about different sample preparation approaches involved in bio-analytical methods.
(b) Explain the following validation parameters in bio-analytical method validation as per USFDA guidelines.
Linearity
Specificity
- 2 (a) What is Bioavailability? Give the Biopharmaceutical Factors affecting drug Bioavailability.
(b) Write the Biopharmaceutics classification system defined by FDA.
- 3 (a) What is enzyme inhibition? Discuss about drug interactions due to enzyme inhibition with examples.
(b) Discuss about drug-protein binding interactions with examples.
- 4 (a) Write about principles, instrumentation and applications of flow cytometry.
(b) Write about basic equipments used in cell culture lab.
- 5 (a) Explain different study designs in bioequivalence studies.
(b) Differentiate absolute and relative bioavailability with illustrative examples and equations.
- 6 (a) Write about cryopreservation and storage of cells.
(b) Discuss the importance and applications of Toxicokinetic studies.
- 7 (a) Discuss about different approaches for identification of metabolites,
(b) Write short note on clinical significance of bioequivalence studies.
- 8 (a) Describe the compendial methods of dissolution testing.
(b) Write about *in-vivo* and *in-vitro* methods for checking cellular permeability of new drug products.

FACULTY OF PHARMACY
M. Pharmacy (Pharmaceutical Analysis) II Semester (PCI) (Main & Backlog)
Examination, December 2021

Subject: Advanced Instrumental Analysis

Time: 2 Hours

Max. Marks: 75

Note: Answer any three of the following questions. (3 x 25 = 75 Marks)

- 1 (a) Explain about method development and trouble shooting in HPLC.
(b) Write about Chiral analysis of Pharmaceuticals using HPLC.
- 2 (a) Discuss about Ion-Pair chromatography.
(b) Explain about head space sampling and columns used in Gas chromatography.
- 3 (a) Write the principle and applications of Super critical fluid chromatography.
(b) Explain about principles and methods of capillary electrophoresis.
- 4 Explain about the following ionization techniques in mass spectrometry.
(a) FAB (b) Electron impact (c) MALD (d) ESI.
- 5 (a) Write about spin-spin coupling and coupling constant.
(b) Write in detail about COSY.
- 6 (a) Write about Nano Liquid Chromatography.
(b) Discuss the principle and detectors use din Gas chromatography.
- 7 (a) Explain about various parameters used in HPLC.
(b) Discuss about 2D NMR.
- 8 (a) Explain about Quadrpole and Time of flight in MS analysis.
(b) Write about LC-NMR.

FACULTY OF PHARMACY
M. Pharmacy (Pharmaceutical Analysis) II Semester (PCI) (Main & Backlog)
Examination, December 2021

Subject: Herbal and Cosmetic Analysis

Time: 2 Hours

Max. Marks: 75

Note: Answer any three of the following questions. (3 x 25 = 75 Marks)

- 1 (a) Write a notes on efficacy of herbal medicines products.
(b) Discuss the validation of herbal therapies.
- 2 (a) How can we determine microbial contamination in herbal formulations?
(b) How foreign matter is determined in herbal drugs?
- 3 (a) Explain the adulterant screening of herbal drugs and their products using modern analytical techniques.
(b) Write notes on WHO guidelines on quality assessment of herbal drugs.
- 4 (a) Explain WHO guidelines for safety monitoring of natural medicine.
(b) Write notes on bio drug-food interactions with suitable examples.
- 5 (a) Explain the Indian standard specification laid down for sampling and testing of dental products.
(b) Write a note on analysis of skin creams as per BIS.
- 6 Write notes on
(a) Global marketing management.
(b) Determination of ester value of cosmetic products.
(c) Analysis of personal hygiene preparations.
- 7 Write about Indian patent law applicable for herbal drugs and natural products.
- 8 (a) Write notes on pharmacokinetic issues related to herbal remedies.
(b) Discuss on an herbal monograph.

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutical. Analysis) II-Semester (PCI) (Suppl.)

Examination, August 2021

Subject: Advanced Instrumental Analysis

Time: 2 Hours

Max. Marks: 75

Note: Answer any Three Questions.

(3 x 25 = 75 Marks)

1. a) Explain the following.
i) Capacity factor ii) Plate height iii) Resolution
b) Explain briefly about
i) UPLC ii) Chiral analysis in HPLC
2. a) Explain the following
i) Ion pair chromatography ii) Affinity chromatography
b) Explain principle and derivitization techniques involved in gas chromatography?
3. a) Explain the principle and instrumentation of super critical fluid chromatography
b) Explain characteristics, Principles, methods and modes of capillary electrophoresis
4. a) Explain the instrumentation and fragmentation rules of mass spectrometry
b) Explain the following ionization techniques
i) Electron impact ii) Field ionization
5. Explain the following
i) Chemical shift ii) Spin – spin coupling iii) Double resonance
6. Explain instrumentation, Solvents and various trouble shooting methods in HPLC
7. Explain about isotopic peaks, metastable ions and various mass analysers used in mass spectrometry
8. Explain the following techniques?
i) FT-NMR ii) ¹³CNMR iii) Cosy

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutical. Analysis) II-Sem. (PCI) (Suppl.)

Examination, July 2021

Subject: Herbal & Cosmetic Analysis

Time: 2 Hours

Max. Marks: 75

Note: Answer any Three Questions.

(3 x 25 = 75 Marks)

1. a) How can we differentiate herbal drugs from conventional drugs?
b) Explain the validation protocol for herbal therapies.
2. a) What is adulteration and deterioration? Write the causes and measures of it.
b) Explain the DNA finger printing technique in identification of drugs of natural origin.
3. a) Give brief explanation on adulterant screening using modern analytical instruments.
b) Write the protocol for stability testing of herbal drugs.
4. a) Explain the bio-drug and bio-food interactions with suitable examples.
b) Write a note on challenges in monitoring the safety of herbal medicines.
5. Explain the general methods of analysis of raw materials used in cosmetic manufacture as per BIS.
6. Write the analysis of baby care products and dental products as per BIS.15
7. Write notes on:
 - a) Efficacy of herbal medicine products
 - b) Global marketing management of herbal drugs
 - c) Determination of acid value of cosmetic products.
8. Compare the monographs of herbal drugs of different pharmacopoeias.

FACULTY OF PHARMACY

M. Pharmacy (Pharma. Analysis) II-Semester (PCI) (Suppl.)

Examination, July 2021

Subject: Quality control and Quality assurance

Time: 2 Hours

Max. Marks: 75

Note: Answer any Three Questions.

(3 x 25 = 75 Marks)

- 1) Write a detailed note on requirements and guidelines of GMP (schedule M) in Pharma industries?
- 2) Write a short note on the following
 - a) Quality control.
 - b) Quality assurance.
 - c) Non clinical testing.
- 3) Define IPQC. Explain in detail about various IPQC tests for
 - a) Tablets
 - b) Ophthalmics
- 4) Explain
 - a) Batch formula Record
 - b) Master formula Record
- 5) Write the detail notes on the following
 - a) Expiry date calculation.
 - b) Limitations of production.
 - c) Calculation of yields.
- 6) Explain the various CPSCEA guidelines for laboratory animal facility.
- 7) Describe the quality control test for containers, closures and secondary packing materials?
- 8) Write a note on
 - a) Sanitation of manufacturing premises.
 - b) Drug product inspection.
 - c) Production record review.

FACULTY OF PHARMACY

**M. Pharmacy (Pharmaceutical. Analysis) II – Semester. (PCI) (Suppl.) Examination,
July 2021**

Subject: Modern Bio analytical techniques

Time: 2 Hours

Max. Marks: 75

Note: Answer any Three Questions.

(3 x 25 = 75 Marks)

1. a. Write about Liquid-Liquid extraction as sample preparation technique.
b. Explain the Bioanalytical method validation as per USFDA guidelines.
2. a. Describe the compendial methods of dissolution testing.
b. Write about different experimental methods for solubility determination.
3. a. Explain about different pharmacokinetic drug interactions.
b. Write the importance and applications of Toxicokinetic studies.
4. a. Write about cryopreservation and storage of cells.
b. Describe different techniques for characterization of cells along with their applications.
5. a. Explain different study designs in bioequivalence studies.
b. Differentiate absolute and relative bioavailability with illustrative examples and equations.
6. a. Write about basic equipments used in cell culture lab.
b. Write about principles, instrumentation and applications of flow cytometry.
7. a. Discuss about different approaches for identification of metabolites.
b. Write short note on clinical significance of bioequivalence studies.
8. a. Describe the principles and applications of Cell viability assays.
b. Write about Rat liver microsomes and Human Liver microsomes.

Examination, October 2020

Time: 2 Hours**Max. Marks: 75**

(3 x 25=75 Marks)

1. a) Explain about different sample preparation approaches in bioanalytical methods.
b) Explain the following validation parameters in bioanalytical method validation as per USFDA guidelines.
 - i) Linearity
 - ii) Precision
2. a) Discuss about Biopharmaceutical factors affecting drug bioavailability.
b) Write the Biopharmaceutics classification system defined by FDA.
3. a) Explain different types of PK-PD drug interactions with suitable example.
b) Discuss the role of LC-MS in bioactivity screening and proteomics.
4. a) Write about basic equipments used in cell culture lab.
b) Write about principles, instrumentation and applications of flow cytometry.
5. a) Explain different methods for assessment of bioavailability of new drug product.
b) Write the clinical significance of bioequivalence studies.
6. a) Discuss the importance and applications of Toxicokinetic studie.
b) Write about different cell culture media.
7. a) Write about *in-vivo* and *in- vitro* methods for checking cellular permeability of new drug products.
b) Write in brief about drug interactions linked to transporters.
8. a) Describe the principles and applications of Cell viability assays.
b) Write about Rat liver microsomes and Human Liver microsomes.

FACULTY OF PHARMACY

M. Pharmacy (Pharm. Analysis) II-Sem. (PCI) (Main & Backlog)

Examination, October 2020

Subject: Herbal & Cosmetic Analysis

Time: 2 Hours

Max. Marks: 75

Note : Answer any Three questions

(3 x 25=75 Marks)

1. (a) Write the WHO guidelines for herbal drug standardization.
(b) Compare the herbal drugs with conventional drugs.
2. (a) Explain the different types adulteration of herbal drugs with suitable examples
(b) How foreign matter is determined in herbal drugs?
3. Explain the adulterant screening of herbal drugs and their products using modern analytical techniques.
4. (a) Write the WHO guidelines for safety monitoring of natural medicine.
(b) Explain bio-drug interactions with suitable examples.
5. Write notes on determination of
(a) Saponification value
(b) Moisture content.
(c) Heavy metals
6. Write notes on
(a) DNA finger printing technique.
(b) Effect of herbal medicine on clinical laboratory testing
(c) Analysis of personal hygiene preparations.
7. Write about Indian patent law applicable for herbal drugs and natural products.
8. (a) Write the spontaneous reporting schemes for bio-adverse reactions.
(b) Write the general methods of analysis of raw materials used in cosmetic manufacture as per BIS.

FACULTY OF PHARMACY

M. Pharmacy (Pharma. Analysis) II-Semester (PCI) (Main & Backlog)

Examination, October 2020

Subject: Quality Control and Quality Assurance

Time: 2 Hours

Max. Marks: 75

Note : Answer any Three questions

(3 x 25=75 Marks)

- 1) Write a detailed note on requirements and guidelines of GMP (schedule M) in Pharma industries?
- 2) Write a short note on the following
 - a) Quality control.
 - b) Quality assurance.
 - c) Non clinical testing.
- 3) Define IPQC. Explain in detail about various IPQC tests for
 - a) Tablets
 - b) Ophthalmics
- 4) Explain
 - a) Batch formula Record
 - b) Master formula Record
- 5) Write the detail notes on the following
 - a) Expiry date calculation.
 - b) Limitations of production.
 - c) Calculation of yields.
- 6) Explain the various CPSCEA guidelines for laboratory animal facility.
- 7) Describe the quality control test for containers, closures and secondary packing materials?
- 8) Write a note on
 - a) Sanitation of manufacturing premises.
 - b) Drug product inspection.
 - c) Production record review.

FACULTY OF PHARMACY**M. Pharmacy (Pharma Analysis) II-Semester (PCI) (Suppl.) Examination, January 2020****Subject: Quality Control and Quality Assurance****Time: 3 Hours****Max Marks: 75****Note: Answer Any Five Questions. ALL Questions carry Equal Marks.**

- | | | |
|---|--|----|
| 1 | a) Explain about Quality Control and Quality Assurance. | 8 |
| | b) Write in detail about Total Quality Management. | 7 |
| 2 | a) Explain the control on environmental pollution. | 8 |
| | b) Explain the maintenance of sterile areas. | 7 |
| 3 | Write in detail about inprocess Quality Control (IPQC) testing of Tablets and parenterals. | 15 |
| 4 | a) Explain the various documents to be maintained by the quality control department. | 7 |
| | b) Explain Master formula and Batch formula records. | 8 |
| 5 | Discuss about | |
| | a) Mix-up's and cross contamination. | 8 |
| | b) Aseptic process control | 7 |
| 6 | Discuss the Good laboratory practices for a quality control laboratory in detail. | 15 |
| 7 | Explain the following | |
| | a) Non-clinical testing. | 5 |
| | b) Controls on animal house | 5 |
| | c) Report Preparation. | 5 |
| 8 | Explain various quality control tests for Glass as a packaging material. | 15 |

FACULTY OF PHARMACY**M. Pharmacy (Pharma. Analysis) II-Semester (PCI) (Suppl.) Examination,****January 2020****Subject : Advance Instrumental Analysis****Time: 3 Hours****Max. Marks: 75****Note:** Answer Any Five Questions. All Questions Carry Equal Marks.

1. a) Explain about various parameters like peak shape. Capacity factor, plate number plate height and resolutions to be considered in HPLC chromatogram 10
b) Write about HPLC importance in chiral analysis of pharmaceuticals? 5
2. a) Discuss about ion pair chromatography 5
b) Explain the instrumentation and pharmaceutical applications of HPTLC 10
3. a) Write the principle and instrumentation of SFC? 7
b) Explain about CE-MS Hyphenation? 8
4. a) Elaborate with neat sketch diagram different types of ionization techniques and analyzers in mass spectrometry? 15
5. a) What do you mean by chemical shift? Explain the various factors influencing it? 10
b) Write about correlative spectroscopy? (COSY) 5
6. a) Write about various columns used in GLC? 8
b) Discuss the principle and applications of size exclusion chromatography? 7
7. a) Explain about HILIC approach in HPLC? 7
b) Discuss about C¹³ NMR 8
8. a) Explain about Q-TOF hyphenation (MS.MS) 7
b) Write the principle and stationary phases used in affinity chromatography? 8

FACULTY OF PHARMACY

**M. Pharmacy (Pharma. Analysis) II-Semester (PCI) (Suppl.) Examination,
January 2020**

Subject: Herbal & Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 Explain the following: (15)
 - (a) Iodine value
 - (b) Peroxide value
 - (c) Ester value
- 2 Explain the following in the evaluation of cosmetic products. (15)
 - (a) Moisture content
 - (b) Viscosity
 - (c) Heavy metals
- 3 What are the different sampling and testing procedures of the following cosmetics products. (15)
 - (a) Baby care products
 - (b) Dental products
 - (c) Skin care products
- 4 Explain briefly the DNA finger printing techniques in identification of drugs. (15)
- 5 Briefly explain the WHO and AYUSH guidelines for safety monitoring of natural products. (15)
- 6 (a) Explain briefly the adulteration screening using modern analytical instruments. (8)
(b) Briefly explain the protocols for stability testing of natural products. (7)
- 7 (a) Describe different measures used in monitoring the safety of herbal products. (7)
(b) Explain with suitable examples about: (8)
 - (i) bio drug –drug interactions
 - (ii) bio drug-food interactions
- 8 Explain the protocols of Indian and International patent laws applicable in herbal drugs and natural products. (15)

FACULTY OF PHARMACY

M. Pharmacy (Pharma Analysis) II- Semester (PCI) (Suppl.) Examination,

January 2020

Subject: Modern Bio Analytical Techniques

Time: 3 Hours

Max Marks: 75

Note: Answer Any Five Questions. ALL Questions carry Equal Marks.

1. a) What is the importance of extraction of drugs and metabolites from biological matrices? 5
b) Describe the bioanalytical method procedure for liquid and solid phase extraction? 10
2. a) Mention the different alternative methods of dissolution testing. 11
b) Define solubility & permeability based on biopharmaceutics classification system. 4
3. Describe various drug (pk-pd) interactions)? 15
4. Discuss the principles and applications of flow cytometry. 15
5. Write the different methods for the assessment of bioavailability and bioequivalence? 15
6. a) Explain the drug permeability by in-vivo method? 8
b) Write notes on cross over design. 7
7. Write notes on the following
a) Drug interaction linked to transporters. 8
b) Cryopreservation techniques. 7
8. Discuss about the design and evaluation of bioequivalence studies. 15

FACULTY OF PHARMACY

**M. Pharmacy (Pharm. Analysis) II-Semester (PCI) (Main) Examination,
August 2019**

Subject : Advanced Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any Five Questions. All Questions Carry Equal Marks.

1. (a) Explain the following chromatographic parameter (i) Capacity factor (ii) Selectivity (iii) Resolution 9
(b) Explain the principle involved in UPLC and compare it with HPLC in terms of different parameters? 7
2. (a) Explain the Principle involved in size exclusion chromatography and write about commercially available columns and their properties. 7
(b) Explain in detail about derivatisation in Gas chromatography 8
3. (a) Explain the principle and applications of super critical fluid chromatography? 7
(b) What is capillary electrophoresis? Explain its principle, methods and modes of CE? 8
4. (a) What is the theory involved in mass spectrometry and explain the following ionization techniques (i) Electron impact (ii) field ionization (iii) MALDI ionization 10
(b) Explain Mc. Lafferty arrangement with example. 5
5. (a) Define chemical shift? Explain the factors influencing chemical shift. 7
(b) Draw a schematic NMR spectra and explain the interpretation for the following compounds (i) Diethylether (ii) Ethoxyacetic acid (iii) n- propyl formate
6. (a) Explain the following techniques 8
1. NOESY 2. COSY
(b) Explain the following mass analyzers in detail
1. Quadrupole 2. Time of flight
7. (a) What is enantiomeric separations? Explain role of HPLC in chiral analysis? 7
(b) Write the principle, head space sampling and columns used in gas chromatography
8. (a) Explain the principle involved in the following hyphenated techniques 7
(i) LC-MS (ii) LC-NMR (iii) CE-MS
(b) Write the applications of
(i) LC-MS (ii) LC-NMR (iii) CE-MS 8

FACULTY OF PHARMACY

**M. Pharmacy (Pharma. Analysis) II-Semester (PCI) (Main) Examination,
August 2019**

Subject: Herbal & Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 (a) Write a note on efficacy of herbal medicine products. (5)
(b) Explain the pharmacodynamic and pharmacokinetic issues of herbal medicines. (10)
- 2 (a) Write about sampling procedures of drugs of natural origin. (7)
(b) How foreign matter is determined in herbal drugs? (8)
- 3 (a) Explain the adulterant screening of herbal drugs and their products using modern analytical techniques. (10)
(b) Write a note on effect of herbal medicine on clinical laboratory testing. (5)
- 4 (a) Write the spontaneous reporting schemes for bio drug adverse reactions and bio drug –drug interactions. (10)
(b) Give the challenges in monitoring the safety of herbal medicine. (5)
- 5 (a) Explain the Indian standard specification laid down for sampling and testing of baby care products. (10)
(b) Write a note on analysis of skin creams as per BIS. (5)
- 6 Write notes on : (3x5)
(a) Global marketing management
(b) Determination of ash value of cosmetic products
(c) Analysis of personal hygiene preparations
- 7 Write about Indian patent law applicable for herbal drugs and natural products. (15)
- 8 (a) Write about DNA finger printing techniques in identification of natural drugs. (7)
(b) Discuss the stability testing of natural products. (8)

FACULTY OF PHARMACY

M. Pharmacy (Pharma Analysis) II- Semester (PCI) (Main) Examination, Aug. 2019

Subject: Quality Control and Quality Assurance

Time: 3 Hours

Max Marks: 75

Note: Answer Any Five Questions. ALL Questions carry Equal Marks.

1. Write a detailed note on requirements and guidelines of GMP(schedule M) in Pharma industries? 15
2. Write brief notes on
 - a) Good warehousing practice 7
 - b) Pharmaceutical inspection convention 8
3. Describe the quality control test for containers, closures and secondary packing materials? 15
4. a) Write a short note on good documentation practice guidelines. 6
b) What are the different types of audits? Explain in detail audit methods and techniques involved in it. 9
5. Describe the guidelines of CPCSEA 15
6. a) Explain the quality control test for ointments according to IP 8
b) Release of finished product. 7
7. Write brief notes on following
 - a) Change control 7
 - b) SOP 8
8. Describe sources of contamination and methods of contamination control? 15

FACULTY OF PHARMACY

M. Pharmacy (Pharma Analysis) II- Semester (PCI) (Main) Examination, Aug. 2019

Subject: Modern Bio Analytical Techniques

Time: 3 Hours

Max Marks: 75

Note: Answer Any Five Questions. ALL Questions carry Equal Marks.

1. a) What is the importance of extraction of drugs and metabolites from biological matrices?
b) Describe the bioanalytical method procedure for liquid and solid phase extraction? 15
2. a) Mention the different alternative methods of dissolution testing transport models 11
b) Define solubility & permeability based on biopharmaceutics classification system. 4
3. Describe various drug interaction (pk-pd) interactions)? 15
4. Discuss the principles and applications of flow cytometry. 15
5. Write the different methods for the assessment of bioavailability and bioequivalence? 15
6. a) Explain the drug permeability by in-vivo method? 8
b) Write notes on cross over design. 7
7. Write notes on the following
a) Drug interaction linked to transporters. 8
b) Cryopreservation techniques. 7
8. Discuss about the design and evaluation of bioequivalence studies. 15

FACULTY OF PHARMACY

**M. Pharmacy (Pharm. Analysis) II-Semester (PCI) (Suppl.) Examination,
February 2019**

Subject: Herbal & Cosmetic Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

1. Write a short note on the following: (15)
 - a) Herbal and Conventional drugs
 - b) Adulteration and Deterioration
 - c) Types of adulteration
2. Write a short note on the following: (15)
 - a) WHO guidelines
 - b) AYUSH guidelines
3. Explain briefly about: (15)
 - a) acid value
 - b) saponification value
 - c) rancidity
4. Explain briefly the evaluation of the following cosmetic products according to Bureau of Indian Standards. (15)
 - a) Hair products
 - b) Skin creams
 - c) Lip sticks
5. Write a note on effect of herbal medicine on clinical lab testing? (15)
6. Explain briefly the stability testing of natural products? (15)
7. Explain briefly about bio drug adverse reactions, bio drug-drug and bio drug-food interactions with suitable examples? (15)
8. Explain briefly the WHO guidelines in quality assessment of herbal drugs? (15)

FACULTY OF PHARMACY

**M. Pharmacy (Pharmaceutical Analysis) II-Semester (PCI) (Suppl.) Examination,
February 2019**

Subject: Quality Controls and Quality Assurance

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 Write a short note on the following
 - a) Quality control. (5)
 - b) Quality assurance. (5)
 - c) Non clinical testing. (5)
- 2 Explain the various CPSCEA guidelines for laboratory animal facility. (15)
- 3 Define IPQC. Explain in detail about various IPQC tests for
 - a) Capsules. (8)
 - b) Parenterals. (7)
- 4 Give a brief note on
 - a) Quality audit plan. (5)
 - b) Protocols and reports. (5)
 - c) Distribution records. (5)
- 5 Discuss the Good laboratory practices for a quality control laboratory in detail. (15)
- 6
 - a) Explain the various documents to be maintained by the quality control department. (7)
 - b) Explain Master formula and Batch formula records. (8)
- 7 Explain various cGMP guidelines according to schedule M.
- 8 Write a note on
 - a) Sanitation of manufacturing premises (5)
 - b) Drug product inspection. (5)
 - c) Production record review. (5)

FACULTY OF PHARMACY

**M. Pharmacy (Pharmaceutical Analysis) II-Semester (PCI) (Suppl.) Examination,
February 2019**

Subject: Advance Instrumental Analysis

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 Write the principle involved in HPLC and explain the following.
(a) Peak shapes (b) Plate number (c) Plate height (10)
(d) Explain various pumps used in HPLC. (5)
- 2 Explain the principle and stationary phases of the following: (2x7½)
(a) Ion Exchange chromatography (b) Affinity chromatography
- 3 Write in detail about Instrumentation, columns and detectors used in Gas chromatography. (15)
- 4 (a) Explain the instrumentation and applications of super critical fluid chromatography. (7)
(b) Explain characteristics and pharmaceutical analysis of capillary electrophoresis. (8)
- 5 (a) Explain the following ionization techniques
(a) chemical ionization (b) FAB (c) ESI (9)
(b) Explain fragmentation pattern of
(a) Alcohols (b) Aldehydes (c) aliphatic acids (6)
- 6 Explain the following: (3x5)
(a) Spin-spin coupling
(b) Coupling constant
(c) Nuclear magnetic double resonance
- 7 Write about the principles instrumentation and applications of : (2x7½)
(a) TLC
(b) Size exclusion chromatography
- 8 (a) Explain in detail about chiral stationary phases (CSP's). (6)
(b) Explain principle and applications of HPTLC. (9)

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FACULTY OF PHARMACY

**M. Pharmacy (Pharmaceutical Analysis) II-Semester (PCI) (Suppl.) Examination,
February 2019**

Subject: Modern Bio Analytical Techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 Write about the following sample preparation techniques. 6
 - (a) Solid phase extraction
 - (b) Liquid Liquid extraction
 - (c) Explain the Bioanalytical method validation as per USFDA guidelines. 9
- 2 (a) Discuss about Biopharmaceutical factors affecting drug bioavailability. 10
 - (b) Write the Biopharmaceutics classification system defined by FDA. 5
- 3 (a) What is enzyme inhibition? Discuss about drug interactions due to enzyme inhibition with examples. 7
 - (b) Discuss about drug-protein binding interactions with examples. 8
- 4 (a) Write about principles, instrumentation and applications of flow cytometry. 9
 - (b) Write about cryopreservation and storage of cells. 6
- 5 (a) Explain different study designs in bioequivalence studies. 10
 - (b) Differentiate absolute and relative bioavailability with illustrative examples and equations. 5
- 6 (a) Discuss the importance and applications of Toxicokinetic studies. 8
 - (b) Write about basic equipments used in cell culture lab. 7
- 7 (a) Discuss about different approaches for identification of metabolites. 10
 - (b) Write short note on clinical significance of bioequivalence studies. 5
- 8 (a) Describe the compendia methods of dissolution testing. 7
 - (b) Write about *in-vivo* and *in- vitro* methods for checking cellular permeability of new drug products. 8

FACULTY OF PHARMACY**M. Pharmacy (Pharm. Analysis) II-Semester (PCI) (Main) Examination,****August 2018****Subject: Advance Instrumental Analysis****Time: 3 Hours****Max. Marks: 75****Note: Answer any five questions. All questions carry equal marks.**

- 1 a. Explain about various types of columns and column problems in HPLC. (9)
b. Write the principle and advantages of Ultra and Nano liquid chromatography? (6)
- 2 a. Discuss about ion exchange chromatography and write in detail about its applications? (7)
b. Explain the various components of HPTLC and write its advantages over column chromatography? (8)
- 3 a. Write about various detectors used in GLC? (10)
b. Explain the principle and basic configuration of capillary electrophoresis? (5)
- 4 Elaborate with neat sketch, the instrumentation of mass spectrometry? (15)
- 5 a. What do you mean by chemical shift? Explain the various factors influencing it? (10)
b. Explain about nuclear double resonance and its applications? (5)
- 6 a. Mention various tandem MS/MS systems and explain any one briefly with neat sketch? (9)
b. Discuss the principle and applications of size exclusion chromatography? (6)
- 7 a. Explain about preparative HPLC? (7)
b. Discuss about FT NMR with reference to C¹³ NMR (8)
- 8 a. Explain about LC-NMR hyphenation. (9)
b. Write about fragmentation ruleS in MS? (6)

FACULTY OF PHARMACY

**M. Pharmacy (Pharm. Analysis) II-Semester (PCI) (Main) Examination,
August 2018**

Subject: Modern Bio Analytical Techniques

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 Write notes on bio analytical method validation as per FDA Guidelines? (15)
- 2 Explain the factors effecting for enhancement of bioavailability of drugs? (15)
- 3 Describe the Cytochrome P450-based drug interactions ? (15)
- 4 Write brief notes on
 - a) Various types of cell culture (8)
 - b) LC-MS in bioactivity screening and proteomics (7)
- 5 Describe the principles and applications of cell viability assays of MTT assays?(15)
- 6 Write the alternate methods for dissolution testing? (15)
- 7
 - a) Define and explain bioavailability, bioequivalence and biosimilar. (6)
 - b) Write about various design to conduct bioavailability studies. (9)
- 8
 - a) Discuss about the bioanalytical methods such as protein precipitation. (7)
 - b) Describe the various solubility techniques. (8)

FACULTY OF PHARMACY**M. Pharmacy (Pharm.Analysis) II-Semester (PCI) (Main) Examination, August 2018****Subject: Herbal & Cosmetic Analysis****Time: 3 Hours****Max. Marks: 75****Note: Answer any five questions. All questions carry equal marks.**

- 1 (a) Write note Herbal medicines Vs Conventional drugs. 5
(b) Explain the standardization of herbal drugs according to WHO guidelines. 10
- 2 What is adulteration and deterioration? Explain types, causes and measure of adulteration. 15
- 3 (a) Describe the stability testing of natural products with suitable examples. 8
(b) Write a note on effect of herbal medicine on clinical laboratory testing. 7
- 4 (a) Write the spontaneous reporting schemes for bio drug adverse reactions and bio drug-food interactions. 10
(b) Write about AYUSH guideline on safety monitoring of natural medicine. 5
- 5 Explain the general methods of analysis of raw materials used in cosmetic manufacture as per BIS. 15
- 6 Write the analysis of lipsticks and hair products as per BIS. 15
- 7 Write notes on 3x5=15
(a) Determination of pesticide residues in herbal formulations.
(b) Challenges in monitoring the safety of herbal medicines.
(c) Determination of iodine value of cosmetic products.
- 8 Write about Indian patent law applicable for herbal drugs and natural products. 15

FACULTY OF PHARMACY

M. Pharmacy (Pharm.Analysis) II-Semester (PCI) (Main) Examination, August 2018

Subject: Quality Controls and Quality Assurance

Time: 3 Hours

Max. Marks: 75

Note: Answer any five questions. All questions carry equal marks.

- 1 Describe concept, components of Quality Assurance and Quality control. (15)
- 2 What are the requirements of an organization and personnel as per USFDA? (15)
- 3 Describe the in process quality control and finished products quality control of tablet according to Indian pharmacopeia. (15)
- 4 Write a brief notes on a) Quality audit plan (8)
b) Batch formula record (7)
- 5 Write the detail notes on the following (5)
(a) Expiry date calculation (5)
(b) Limitations of production (5)
(c) Calculation of yields (5)
- 6 a) Describe the overview of ICH Guidelines with Q series (8)
b) Write notes on SOP. (7)
- 7 a) Write note on the aseptic process control. (8)
b) Write about the organization and personnel responsibilities as per WHO. (7)
8. a) Describe the onsite sanitation of manufacturing premises (8)
b) Write note on finished product (7)

FACULTY OF PHARMACY**M. Pharmacy (Pharm. Analysis) II-Semester (PCI) (Main) Examination,****August 2018****Subject: Advance Instrumental Analysis****Time: 3 Hours****Max. Marks: 75****Note: Answer any five questions. All questions carry equal marks.**

- 1 a. Explain about various types of columns and column problems in HPLC. (9)
b. Write the principle and advantages of Ultra and Nano liquid chromatography? (6)
- 2 a. Discuss about ion exchange chromatography and write in detail about its applications? (7)
b. Explain the various components of HPTLC and write its advantages over column chromatography? (8)
- 3 a. Write about various detectors used in GLC? (10)
b. Explain the principle and basic configuration of capillary electrophoresis? (5)
- 4 Elaborate with neat sketch, the instrumentation of mass spectrometry? (15)
- 5 a. What do you mean by chemical shift? Explain the various factors influencing it? (10)
b. Explain about nuclear double resonance and its applications? (5)
- 6 a. Mention various tandem MS/MS systems and explain any one briefly with neat sketch? (9)
b. Discuss the principle and applications of size exclusion chromatography? (6)
- 7 a. Explain about preparative HPLC? (7)
b. Discuss about FT NMR with reference to C¹³ NMR (8)
- 8 a. Explain about LC-NMR hyphenation. (9)
b. Write about fragmentation ruleS in MS? (6)
