M. Pharmacy I - Semester (Common to All) (PCI) (Main & Backlog) Examination, May 2022

Subject: Modern Pharmaceutical Analytical Techniques

Time: 3 Hours Max. Marks: 75

Note: Answer any five questions.

 $(5 \times 15 = 75 \text{ Marks})$

- 1 (a) State and explain Beer-Lambert's Law. Add a note on the deviations from Beer's law.
 - (b) Explain the concept of chromophore, auxochrome and bathochromic shift with suitable examples.
- 2 (a) Explain the instrumentation of FTIR with a neat labelled diagram. Add a note on the advantages of FTIR.
 - (b) Explain the mplecular vibrations in IR.
- 3 (a) What is the principle AAS? Explain the instrumentation.
 - (b) List the differences between AAS and flame photometry.
- 4 What is the significance of chemical shift? What are the factors affecting chemical shift? Name the internal standard and justify its selection as internal standard in NMR spectroscopy.
- 5 What is the principle of Mass Spectrometry? With a neat labelled diagram briefly explain the components of MS instrumentation.
- 6 (a) Classify the ionization techniques in MS. Explain any three methods in detail.
 - (b) Define Base peak, molecular ion peak and metastable ion.
- 7 (a) Explain the principle of X-ray diffraction.
 - (b) Explain HPLC instrumentation with a labelled diagram.
- 8 (a) Explain the experimental set up required for gel electrophoresis.
 - (b) Describe the principle and applications of RIA.

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Main & Backlog) Examination, May 2022

Subject: Regulatory Affairs

Time: 3 Hours Max. Marks: 75

Note: Answer any five questions.

 $(5 \times 15 = 75 \text{ Marks})$

- 1 (a) Write a note on documentation in pharmaceutical industry.
 - (b) Write a note on Hatch Waxman Act amendments.
- 2 Explain SUPAC for Immediate release and Modified-release dosage forms.
- 3 Describe the regulatory approval process for ANDA and NDA.
- 4 (a) Explain the regulations for medical devices.
 - (b) Describe Q8, Q9 and Q10 ICH Quality guidelines.
- 5 (a) Write a note on CTD and eCTD.
 - (b) Explain the ways and means of US registration for foreign drugs.
- 6 Discuss about
 - (a) Investigation of Medicinal Products Dossier (IMPD)
 - (b) Investigation brochure.
- 7 Write a note on
 - (a) Clinical trial protocol
 - (b) Regulatory requirements of MHRA
- 8 Write a note on
 - (a) Pharmacovigilance safety monitoring
 - (b) Institutional review board.

M. Pharmacy (Pharmaceutics) I - Semester (PCI) (Main & Backlog) Examination, May 2022

Subject: Modern Pharmaceutics

Time: 3 Hours Max. Marks: 75

Note: Answer any five questions.

 $(5 \times 15 = 75 \text{ Marks})$

- 1 (a) Write the determination of shelf-life of formulations?
 - (b) Discuss the evaluation of parenteral dosage forms?
- 2 (a) Discuss the ICH guidelines for calibration and validation of equipment?
 - (b) Explain the terms DQ, IQ, OQ and PQ?
- 3 (a) Write a note on inventory management and production control management?
 - (b) Write a note on sale forecasting and budget planning in industries?
- 4 (a) Explain the phases of compaction profile?
 - (b) Discuss the solubility enhancement techniques of drugs?
- 5 (a) Write a note on ANOVA test?
 - (b) Write about the pharmacokinetic parameters?
- 6 (a) Write a note on response surface method for optimization of formulation?
 - (b) Describe f1 and f2 factors and their calculations?
- 7 (a) Discuss the evaluation of dispersion systems?
 - (b) Describe the preparation and evaluation of SMEDDS?
- 8 (a) Write note on Total quality management?
 - (b) Discuss students t-test and it's applications?

M. Pharmacy (Pharmaceutics) I - Semester (PCI) (Main & Backlog) Examination, May 2022 Subject: Drug Delivery System

Time: 3 Hours Max. Marks: 75

Note: Answer any five questions.

 $(5 \times 15 = 75 \text{ Marks})$

- 1 (a) What are the differences between sustained release and controlled release formulations?
 - (b) Explain in detail various physiochemical and biological approaches for controlled release formulations.
- 2 (a) Classify polymers? What are biodegradable and non biodegradable polymers?
 - (b) Explain in detail applications of polymers in controlled release formulations.
- 3 (a) Define pharmacogenetics. What are personalized medicines?
 - (b) Write in detail about bioelectronics medicines and telepharmacy.
- 4 (a) What are rate controlled drug delivery system?
 - (b) Explain in detail osmotic drug delivery system.
- 5 Classify and explain in gastro-retentive drug delivery systems.
- 6 Explain about mucosal stransdermal delivery of vaccines.
- 7 (a) What are permeation enchancers, explain the mechanism of permeation enhancers with examples.
 - (b) Explain the permeation barriers of ocular drug delivery system.
- 8 (a) What are the barriers of protein drug delivery system.
 - (b) Explain stability of proteins.

M. Pharmacy I Semester (PCI) (Suppl) Examination, December 2021 (COMMON TO ALL)

Subject: Modern Pharmaceutical Analytical Techniques

Time: 2 Hours Max. Marks: 75

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

 $(3 \times 25 = 75 \text{ Marks})$

- 1 (a) State and explain Beer-Lambert's law. Add a note on the deviations from Beer's law.
 - (b) Explain the electronic transitions in UV spectroscopy.
- 2 (a) Explain the principle and instrumentation of FTIR with a neat labelled diagram.
 - (b) Explain the named advantages of FTIR.
 - (c) What are the major differences between Dispersive instruments and FTIR?
- 3 (a) What is the principle of Fluorescence? Explain the radiative and non radiative pathways of relaxation.
 - (b) Add a note on the factors affecting fluorescence.
- 4 (a) Explain NMR instrumentation with a diagram.
 - (b) Briefly explain shielding and deshielding with suitable example.
- 5 (a) What is the principle of MS? With a neat labelled diagram briefly explain the components of MS instrumentation.
- 6 (a) Classify the ionization techniques in MS. Explain any three methods in detail.
 - (b) Define Base Peak, molecular ion peak and metastable ion.
- 7 (a) Explain GC instrumentation with a labelled diagram.
 - (b) What are the applications of HPLC?
- 8 (a) Explain the experimental set up required for capillary electrophoresis.
 - (b) Describe the principle and application of ELISA.

M.Pharmacy (Pharmaceutics) I Semester (PCI) (Suppl) Examination, December 2021

Subject: Drug Delivery System

Time: 2 Hours Max. Marks: 75

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

 $(3 \times 25 = 75 \text{ Marks})$

- 1 (a) Define personalized medicine. Explain telepharmacy and 3D printing of pharmaceuticals.
 - (b) What are non-biodegradable polymers? Explain the mechanism of polymer degradation?
- 2 (a) Mention different types of rate controlled drug delivery system. Explain in detail rate preprogrammed drug delivery system.
 - (b) Describe in detail osmotic drug delivery system.
- 3 (a) Explain various theories of mucoadhesion.
 - (b) Explain in detail types of gastro retentive drug delivery system.
- 4 (a) What are the barriers of ocular drug delivery system?
 - (b) Describe in detail evaluation of ocular DDS.
- 5 Define vaccine. Write a note on uptake of antigens and permeation of vaccines through mucosal and transdermal route.
- 6 (a) Classify transdermal drug delivery system. What are the ideal properties of a drug to be formulated as TDDS?
 - (b) Describe in detail Iontophoretic DDS.
- 7 (a) What are stability issues of proteins and peptides?
 - (b) Explain various non-invasive routes of administration of proteins and peptides.
- 8 (a) Mention different types of transdermal permeations enhancers and write about its mechanism of improving permeation.
 - (b) Write a note on Bioelectronic medicines.

M.Pharmacy (Pharmaceutics) I Semester (PCI) (Suppl) Examination, December 2021

Subject: Modern Pharmaceutics

Time: 2 Hours Max. Marks: 75

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

 $(3 \times 25 = 75 \text{ Marks})$

- 1 (a) Discuss the preparation and evaluation of SMEDDS.
 - (b) List out the methods used for determination of drug-excipient interactions and discuss any three methods in detail with examples.
- 2 (a) Discuss the ICH guideline for calibration and validation of equipment with an example.
 - (b) Write a note on sale forecasting and budget planning in industries.
- 3 (a) Describe the layout of buildings, services in industries according to GMP.
 - (b) Write a note on inventory management and production control management.
- 4 (a) Write the Heckel equation and draw the Heckel plots for determination of Porosity of tablet during compression process.
 - (b) Discuss the methods for enhancement of aqueous solubility of drugs.
- 5 (a) Discuss about the pharmacokinetic parameters required for determination of bioavailability.
 - (b) Describe the comparison of dissolution profiles of dosage forms using similarity and difference factors.
- 6 (a) Write a note on response surface method for optimization of formulation.
 - (b) Explain the terms DQ, IQ, OQ and PQ.
- 7 (a) What is compaction profile? Explain the phases of compaction profile with a suitable examples.
 - (b) Write a note on Total quality management.
- 8 (a) Write in brief about the determination of shelf-life of formulations.
 - (b) Write about the variance and standard deviation and its application in pharmaceutical formulations.

M. Pharmacy (Pharmaceutics) I Semester (PCI) (Suppl) Examination, December 2021

Subject: Regulatory Affairs

Time: 2 Hours Max. Marks: 75

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

 $(3 \times 25 = 75 \text{ Marks})$

- 1 (a) Write a note on Hatch Waxmann Act.
 - (b) Write a note on SUPAC guidelines.
- 2 Explain ANDA regulatory approval process.
- 3 What are the regulatory requirements for approval of API?
- 4 (a) Explain the objectives of CMC considerations during drug development.
 - (b) Enlist ICH quality guidelines.
- 5 Explain the regulatory requirements of Europe Union (EU).
- 6 Discuss about
 - (a) Investigation of medicinal products dossier (IMPD).
 - (b) Bio-equivalence studies for generic drugs assessment.
- 7 Write a note on
 - (a) HIPAA.
 - (b) Pharmacovigilance safety monitoring.
- 8 Write a note on
 - (a) Informed consent process and procedures.
 - (b) Institutional review board.

FACULTY OF PHARMACY M.Pharmacy I Semester (PCI) (Main & Backlog) Examination, July 2021 (COMMON TO ALL)

Subject: Modern Pharmaceutical Analytical Techniques

Time: 2 Hours Max. Marks: 75

Note: Answer any three from the following questions. $(3 \times 25 = 75 \text{ Marks})$

- 1 (a) With a neat labelled diagram explain UV/Visible instrumentation.
 - (b) Briefly explain the electronic transitions with examples.
- 2 (a) Explain the molecular vibrations in IR.
 - (b) Write the sampling methods in IR spectroscopy.
- 3 (a) Explain the principle of flame photometry.
 - (b) With a diagram explain the instrumentation for flame photometry.
 - (c) List some metals that can be analysed by flame photometry.
- 4 (a) Explain the principle of proton NMR spectroscopy.
 - (b) What is the significance of chemical shift? What are the factors affecting chemical shift?
 - (c) What is the internal standard used in NMR spectroscopy? Why it is selected as internal standard?
- 5 (a) List and explain the steps in MS.
 - (b) What are the mass analysers used in MS? Explain any two in detail.
- 6 (a) Explain HPLC instrumentation with a labelled diagram.
 - (b) List and explain any 2 GC detectors.
- 7 (a) Explain Bragg's equation and derive the equation.
 - (b) Explain the principle and the materials required for Paper electrophoresis.
- 8 (a) Explain the principle and types of RIA?
 - (b) Briefly explain Zone electrophoresis and Moving boundary electrophoresis.

FACULTY OF PHARMACY M.Pharmacy (Pharmaceutics) I Semester (PCI) (Main & Backlog) Examination, July 2021

Subject: Modern Pharmaceutics

Time: 2 Hours Max. Marks: 75

Note: Answer any three from the following questions. $(3 \times 25 = 75 \text{ Marks})$

- 1 (a) Discuss about the preparation and evaluation of SMEDDS.
 - (b) Discuss the evaluation tests for parenteral dosage forms
- 2 (a) Discuss the ICH guidelines for calibration and validation of equipment with an example.
 - (b) Explain the terms DQ, IQ, OQ and PQ.
- 3 (a) Write a note on inventory management and production control management.
 - (b) Write note on Total quality management.
- 4 (a) What is compaction profile? Explain the phases of compaction profile with a suitable examples.
 - (b) Discuss the methods for enhancement of aqueous solubility of drugs.
- 5 (a) Write a note on ANOVA test.
 - (b) Write about the variance and standard deviation and its application in pharmaceutical formulations.
- 6 (a) Write a note on response surface method for optimization of formulation.
 - (b) Describe the comparison of dissolution profiles of dosage forms using similarity and difference factors.
- 7 (a) Write in brief about the determination of shelf-life of formulations.
 - (b) Discuss the evaluation tests for both dispersion systems.
- 8 (a) Write a note on sale forecasting and budget planning in industries.
 - (b) Discuss students t-test and its applications.

FACULTY OF PHARMACY M.Pharmacy (Pharmaceutics) I Semester (PCI) (Main & Backlog) Examination, July 2021

Subject: Drug Delivery System

Time: 2 Hours Max. Marks: 75

Note: Answer any three from the following questions. $(3 \times 25 = 75 \text{ Marks})$

- (a) Define sustained release, delayed release and controlled release drug delivery system with examples.
 - (b) Define pharmacogenetics. Explain Bioelectronic medicines.
- 2 (a) Classify activation modulated system. Explain osmotic drug delivery system.
 - (b) What are rate controlled drug delivery system? Explain feedback regulated systems.
- 3 (a) Mention the various barriers of buccal mucoadhesive drug delivery system.

 Mention two exmaples of marketed mucoadhesive drug delivery system.
 - (b) What are the drug properties to formulate as a FDDS? Explain in detail floating drug delivery system.
- 4 (a) Describe formulation and evaluation of ocular drug delivery system.
 - (b) Write a note on ocular insitu gels.
- 5 (a) Explain structure of skin. What are barriers of transdermal permeation? What are the ideal properties of a drug candidate to permeate through the skin?
 - (b) Classify transdermal permeation enhancers. Explain the mechanism of permeation enhancers.
- 6 (a) Describe in detail different strategies to formulate a stable protein and peptide drug formulations.
 - (b) Describe physical and chemical stability of proteins and peptides.
- 7 (a) Explain in detail mucosal and active transdermal methods of delivering vaccines through mucosa and skin.
 - (b) Explain single shot vaccines.
- 8 (a) Explain various physicochemical and biological factors that influences the formulation of SR and CR formulations.
 - (b) Classify polymers. Describe properties of any two synthetic non-biodegradable polymers.

FACULTY OF PHARMACY M.Pharmacy (Pharmaceutics) I Semester (PCI) (Main & Backlog) Examination, August 2021

Subject: Regulatory Affairs

Time: 2 Hours Max. Marks: 75

Note: Answer any three of the following questions. $(3 \times 25 = 75 \text{ Marks})$

- 1 (a) Explain CFR with respect to Pharmaceutical product development.
 - (b) Write a note on generic drugs product development.
- 2 (a) Explain the importance of documentation and documents to be maintained in Pharmaceutical industry.
 - (b) What is the impact of outsourcing Bioavailability and Bioequivalence studies to Contact Research Organisations (CRO).
- 3 (a) Explain SUPAC guidelines for Immediate release dosage form.
 - (b) Explain evaluation of drug product performance by *invitro* studies.
- 4 Write a note on
 - (a) CTD and eCTD.
 - (b) Regulations for combination products.
- 5 Explain the regulatory requirements of TGA.
- 6 Discuss about
 - (a) Health Insurance Portability and Accountability Act.
 - (b) Institutional review board/ independent ethics committee.
- 7 Write a note on investigation of medicinal products dossier (IMPD) and Investigator brochure.
- 8 Write a note on
 - (a) Pharmacovigilance and safety monitoring in clinical trials.
 - (b) Enlist ICH Efficacy guidelines.

Code No: 6330/PCI

FACULTY OF PHARMACY

M. Pharmacy (pharmaceutics) I-semester (PCI) (Suppl.) Examination, October 2020

Subject: Drug Delivery System

Time: 2 hrs Max Marks: 75

Note: Answer any three questions.

(3x25=75 Marks)

- 1. Explain different mechanisms of drug delivery from sustained or controlled release formulations? Add a note on application of polymers in sustained release dosage forms?
- 2. What do you mean by personalized medicines? Describe in detail 3D printing of pharmaceuticals and telepharmacy.
- 3. Explain the principles of rate-controlled drug delivery systems? Write a note on feedback regulated drug delivery systems?
- 4. a) Mention different types of gastro-retentive drug delivery system?
 - b) Describe in detail floating drug delivery system and its evaluation?
- 5. a) Describe in detail formulation and evaluation of buccal drug delivery system?
 - b) Explain the different factors affecting mucosal drug permeation?
- 6. a) Describe barriers of ocular drug delivery system and what are the methods to overcome the same?
 - b) Describe ideal properties of a drug to formulate as a transdermal drug delivery system?
- 7. a) Describe different routes of administration of protein drug delivery and its barriers of permeation?
 - b) Explain stability of protein pharmaceuticals?
- 8. What do you mean by single shot vaccine delivery systems? Explain mucosal delivery of vaccines?

Code No: 6332/PCI

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Suppl.) Examination, November 2020

Subject : Regulatory Affairs

Time: 2 Hours Max. Marks: 75

Note: Answer any Three questions.

(3 x25=75 Marks)

- 1. a) Explain evaluation of drug product performance by *invitro* studies
 - b) Write a note on generic drugs product development
- 2. Explain Hatch-Waxmann Act and its amendments.
- 3. Explain SUPAC guidelines for immediate release dosage form
- 4. Write a note on
 - a) CTD and eCTD
 - b) Regulations for combination products.
- 5. Explain the regulatory requirements of TGA
- 6. Discuss about
 - a) ICH guidelines for quality & safety
 - b) Institutional review board / independent ethics committee
- 7. Write a note on investigation of medicinal products dossier (IMPD) and Investigator brochure.
- 8 Write a note on
 - a) Pharmacovigilance and safety monitoring in clinical trials.
 - b) Enlist ICH Efficacy guidelines.

Code No: 6331/PCI

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (PCI)

(Suppl.) Examination, November 2020

Subject: Modern Pharmaceutics

Time: 2 Hours Max. Marks: 75
Note: Answer any Three questions. (3 x25=75 Marks)

- 1. Explain the about drug Excipient interactions and methods of determination
- 2. Describe accelerated stability testing of solution and solid dosage forms.
- 3. What is validation. Discuss the validation & calibration of any two equipment
- 4. Discuss WHO good manufacturing practices in a pharmaceutical Industy.
- 5. Discuss about IQ, OQ, PQ & DQ by taking an example.
- 6. a) Explain the types of compaction profiles.
 - b) Write the advantages and disadvantages of strain gauges.
- 7. a) Describe Heckle plots and its significance with necessary equations and graphs.
 - b) Write Biopharmaceutics Classification System (BCS) of drugs with examples.
- 8. a) Explain the reasons for conducting the stability studies of drugs.
 - b) Explain formulation and dosage form related factors influencing the dissolution of tablets.

CODE NO: 6102/PCI

Max. Marks: 75

FACULTY OF PHARMACY

M. Pharmacy I – Semester (Main & Backlog) Examination, January 2020 (Common Paper for all Except Pharmacy Practice)

Subject : Modern Pharmaceutical Analytical Techniques

Time: 3 Hours

	Note: Answer any Five Questions. All Questions Carry Equal Marks.	
1.	(a) State and explain Beer- Lambert's law. Add a note on the deviations from Beelaw.(b) Explain solvents and the selection criteria for UV/Visible spectroscopy.(c) What is solvent shift?	er's 8 4 3
2.	(a) Explain the principle and instrumentation of FTIR with a neat labelled diagram.(b) Explain about the sampling techniques and applications of FR spectroscopics.	8 py 7
3.	(a) What is the principle of Fluorescence? Explain the radiative and non radial pathways of relaxation.(b) Add a note on the factors affecting fluorescence and quenchers in fluorescence.(c) What are the criteria for a molecule to exhibit the phenomena of fluorescence	tive 7 6
4.	(a) Explain the principle of proton NMR spectroscopy.(b) What is the significance of chemical shift. What are the factors affecting chemical shift?(c) Explain about spin-spin crippling and it's importance in NMR	5 6 4
5.	(a) Classify the ionization techniques in MS. Explain any three methods in detail.(b) Differentiate between Base peak and molecular ion peak.	12 3
6.	(a) Explain HPLC instrumentation.(b) What are the applications of HPLC?	10 5
7.	(a) Explain Braggs equation and derive the equation.(b) What is the principle involved in rotating crystal technique?	8 7
8.	Explain the principle, working and applications of (a) Capillary electrophoresis (b) Gel electrophoresis	71/2 71/2

Code No: 6103/PCI

FACULTY OF PHARMACY

M.Pharmacy (pharmaceutics) I-Semester (PCI) (Main & Backlog) Examination, February 2020

Subject: Drug Delivery System

Time: 3hrs Max Marks: 75 Note: Answer any five questions, all questions carry equal marks. 1 Explain physiochemical and biological approaches for designing SR formulations? (15) 2 What do you mean by personalized medicines? Describe in detail bioelectronic medicines and telepharmacy. (15)3 Explain the principles of rate controlled drug delivery systems? Write a note on osmotically controlled drug delivery systems? (15)4 a) Mention different types of gastroretentive drug delivery system? Mention its advantages and disadvantages? (5) b) Describe in detail buccal drug delivery system? (10)5 Explain the principles and theories of mucoadhesion? Add a note on mechanisms of mucosal drug permeation? (15)6 a) Describe barriers of ocular drug delivery system and what are methods to overcome the same? (8) b) Write a note on transdermal permeation enhancers? (7) 7 What are the barriers of protein drug delivery? Explain various modified formulations of proteins and peptides to overcome the absorption barriers? (15)8 What are the advancements in vaccine delivery systems? Explain transdermal delivery of vaccines? (15)

Code No: 6105/PCI

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Main & Backlog) Examination, January 2020

Subject: Regulatory Affairs

Ti	Time: 3 Hours Max. N	
	Note: Answer Any Five Questions. All Questions Carry Equal Marks	
1	(a) Write a note on CFR (code of federal regulation).(b) Write a note on distribution records and master formula record.	(8) (7)
2	Explain NDA regulatory approval process.	(15)
3	What are the regulatory requirements for approval of API?	(15)
4	(a) Explain the objectives of CMC considerations during drug development.(b) Enlist ICH Quality guidelines.	(9) (6)
5	Explain the regulatory requirements of TGA	(15)
6	Discuss about : a) global submission of ANDA b) Bio-equivalence studies for generic drugs assessment.	(9) (6)
7	Write a note on: (a) HIPAA (b) Pharmacovigilance safety monitoring	(6) (9)
8	Write a note on: (a) Investigator brochure (b) Clinical trial protocol	(8) (7)

Code No: 6104/PCI

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Main & Backlog) Examination, January 2020

Subject: Modern Pharmaceutics

Time: 3 Hours Max. Marks: 75

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

a) Explain about distribution of forces during tablet compression with diagrams and equations.	10M
b) Write the applications of force displacement curves of tablet compression.	5M
2. Explain about Heckell and peppas plots	15M
a) Write various drug excipients interactions with necessary examples.b) Describe the salient features of accelerated stability testing of solution dosage	10M
forms.	5M
4. Describe calibration and validation of equipment as per ICH and WHO guidelines.	15M
5. Explain the WHO good manufacturing practices.	15
6. Write the formulation considerations and evaluation of parentral dosage forms.	15M
7. Explain about Material management inventory control in pharmaceutical industry.	15M
8. a) Define factorial designs and write its applications in formulations.	5M
 b) Describe the methods of comparision of dissolution of two products. 	10M

M. Pharmacy (Common paper for all Specialization) I-Semester (PCI) (Suppl.) Examination, August 2019

Subject: Modern Pharmaceutical Analytical Techniques

Time: 3 Hours Max. Marks: 75 Note: Answer any five questions. All questions carry equal marks. 5 1 a) Write Beer-Lambert's law and derive the expression b) Mention the different methods of quantitative analysis by uv-visible spectroscopy. Explain any one method in detail. 10 2. a) Explain the interpretation procedure of IR spectra of different organic compounds in detail. With examples of schematic IR spectra. b) What is fluorescence? Write the factors affecting fluorescence. 5 3. a) What is chemical shift? Write the factors influencing chemical shift? 8 b) Write a note on FT-NMR 7 4. a) Explain the instrumentations and working of mass spectrometer with schematic diagram. 8 b) Write the fragmentation patterns of different organic compounds observed in mass spectroscopy. With the help of schematic mass spectra of a few compounds 7 5. Describe the components and working procedure of HPLC with a neat labeled block diagram. 15 6. a) Write the principle, instrumentation and working of zone electrophoresis. 8 7 b) Write the principle and theory of X-ray diffraction study using Brag's law 7. a) Write the principle and instrumentation of flame photometry 7 8 b) Write notes on any two GC detectors 8. Explain the principle, equipment, procedure, advantages and applications of IR Spectrophotometer 15

Code. No: 13306/PCI

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (Suppl.) Examination, August 2019

Subject : Regulatory Affairs

111	me: 3 Hours	viax. Marks: 75
No	ote: Answer any Five Questions. All Questions Carry Equal Marks.	
1.	(a) Write a note on importance and types of Drug Master file(b) Explain the contents of Hatch-Waxman Act(c) Write about the importance of Post marking surveillance	7 5 3
2.	Explain the regulatory requirements for ANDA approval process in US	15
3.	(a) Describe the importance, preparation and organization of CTD(b) Describe the objectives and structure of Harmonization guidelines (ICH)	10 5
4.	Explain in brief a. The regulations for medical devices b. Regulatory requirements of MHRA	8 7
5.	Give a brief note on each part of the contents of Investigational New Drug Ap (IND)	oplication 15
6.	(a) What is Investigational Medicinal Product dossier (IMPD)? Explain the requirement contents of IMPD.(b) Write a note on Scale up process	ents and 7 8
7.	Explain briefly various phases of clinical trials and design of clinical trials for submission of data to FDA for getting NDA approval.	the 15
8.	Give a brief note on the following: a. Institutional Review Board (IRB) b. Informed Consent	8

Code. No: 13304/PCI

7

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (Suppl.) Examination, August 2019

Subject : Drug Delivery System

Time: 3 Hours Max.	Marks: 75
Note: Answer any Five Questions. All Questions Carry Equal Marks.	
1. (a) Define personalized medicine? Explain bioelectronic medicines and 3D printing	g of 10
pharmaceuticals? (b) What are biodegradable polymers? Explain the mechanism of polymer degrad	_
(a) Classify rate controlled drug delivery system? Explain in detail rate preprogram drug delivery system?(b) Explain osmotic drug delivery system?	nmed 8 7
3. (a) Describe in detail theories of mucoadhesion?(b) Explain in detail evaluation of gastro retentive drug delivery system?	10 5
4. (a) What are the barriers of ocular drug delivery system?(b) Describe in detail formulation of ocular DDS?	5 10
5. Define vaccine? Write a note on uptake of antigens and permeation of vaccines the mucosal & transdermal route?	hrough 15
6. Classify transdermal drug delivery system? Describe in detail active and passive methods of transdermal DDS?	15
7. (a) What are the stability issues of proteins and peptides?(b) Explain non-invasive routes of administration of protein and peptides?	7 8
8. (a) Explain in trandermal permeations enhancers and write about its mechanism improving permeation?	of 8

(b) Write a note on telepharmacy?

Code. No: 13305/PCI

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Suppl.) Examination, August 2019

Subject : Modern Pharmaceutics

Tir	Time: 3 Hours Max. Mark	
No	ote: Answer any Five Questions. All Questions Carry Equal Marks.	
1.	(a) Explain the procedure of drug excipient compatibility studies(b) Write a note on response surface methodology	8 7
2.	(a) Write the approaches of process validation and mention their significance(b) Describe different steps involved in equipment qualification	8 7
3.	Explain different elements of Total Quality Management	15
4.	(a) Explain the distribution of forces and consolidation during compaction(b) Write a note on compaction profiles	10 5
5.	(a) Explain about various diffusion parameters	7 1/2
6.	(b) Write about the Higuchi & poppas plots(a) Discuss in detail about ICH guidelines for validation and calibration of equipment(b) Discuss management of materials and inventory control in industries	7 ^{1/2} 8 7
7.	(a) Explain about various optimization techniques in formulation processing with suitab examples(b) Explain physiological and formulation considerations for parenterals	le 8 7
8.	Write in brief about (a) Heckel plots (b) Solubility (c) Good manufacturing practices	

CODE NO: 13147/PCI

Max. Marks: 75

FACULTY OF PHARMACY

M. Pharmacy (Common Paper for all Specialization) I – Semester (Main & Backlog) Examination, January 2019 Subject : Modern Pharmaceutical Analytical Techniques

Time: 3 Hours

No	ote: Answer any Five Questions. All Questions Carry Equal Marks.	
1)	a) With a neat labeled diagram explain UV/Visible instrumentation.	8
2)	b) Briefly explain the electronic transitions with examples	8 8
۷)	a) Explain the factors affecting vibrational frequencies in IR.	o 7
	(b) Write the sampling methods in IR spectroscopy.	,
3	(a) Briefly explain the source of AAS.	8 Pharmacy
	(b) List and explain the interferences.	5
	(c) List some metals that can be analysed by AAS.	2
4	(a) Explain NMR instrumentation.(b) Briefly explain spin-spin coupling with a suitable example.	8 7
Pulla	components of MS instrumentation. (b)Explain Quadrupole and time of flight analysers in detail.	8 7
.6.	(a) What are [−] the column efficiency parameters?	7
	(b) List and explain any 2 GC detectors.	8
7.	Explain the principle and application of capillary electophoresis. Give a labelled diagram	ı
	to indicate the components of the instrument.	
8	(a) Discuss the principle, instrumentation working and application	
	of a. Paper electrophoresis	
	b. Gel electrophoresis	7+8

Code. No: 13150/PCI

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (Main & Backlog) Examination, February 2019

Subject : Regulatory Affairs

Tir	me: 3 Hours	Max. Marks: 75
No	ote: Answer any Five Questions. All Questions Carry Equal Marks.	
1.	(a) Describe various parts of master formula record and write its importance.(b) Explain salient features of Hatch Waxman Act and its amendments.	. 7 8
2.	Enlist different sections of NDA and Write a note on NDA approval process.	15
3.	(a) Explain regulatory requirements of US registration for foreign durgs.(b) Explain SUPAC guidelines specific to manufacturing changes	8 7
4.	(a) Describe the objectives of harmonization guidelines. Enlist ICH quality gu (b) Explain the objectives of CMC considerations during drug development.	uidelines. 10 5
5.	(a) Explain the regulatory requirement for biologics product approval.	8
	(b) What is the purpose of Investigator's Brochure? Give a brief note on the Information to be filled in each part of the IB.	7
6.	(a) Write a note on eCTD.	7
7	(b) Write different designs of BE studiesfor Generic drugs assessment.	8
1.	(a) Give an outline of factors that must be addressed in the clinical trial proto per USFDA check list.	8 8
	(b) Give a Reddy brief note on Pharmacovigilance and safety monitoring in trials.	clinical 7
8.	Write brief notes on:	
	a. Regulatory requirements 1701 of EU	7
	b. Health Insurance- Portability and Accountability Act.	8

Code. No: 13149/PCI

7

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Main & Backlog) Examination, February 2019

Subject: Modern Pharmaceutics

Time: 3 Hours	Max. Marks: 75
 Note: Answer any Five Questions. All Questions Carry Equal Marks. (b) Explain the terms DQ, IQ, OQ, & PQ 1701 1. (a) Write about stability testing of pharmaceuticals (b) What is optimization? Explain various optimization techniques used for pharmaceutical formulations 	7 8 7
(a) Define validation. What is the importance of validation? Write a note on types of validation ———————————————————————————————————	different 8
3. (a) Discuss about Higuchi & poppas plots(b) Explain in brief about Industrial and Personnel Relationship.(b) Write a note on heckle plots	10 5 5
4. (a) Describe in detail about physics of tablet compression	10
5. (a) Explain about various parameters influencing dissolution (b) Write in brief about students T test 1701Hyderabad (b) What areReddySMEDDS? Give note on importance and formulation of SMEDDS	10 5 7
6. (a) Explain about the production planning and control with suitable examples	8
	_
(b) Give an account on various approaches for inventory management and	7 control 8
8. Write a note on (a) Similarity, dissimilarity factors (b) Total quality management (c) Factorial design	

. (a) Expla

Code. No: 13148/PCI

7

FACULTY OF PHARMACY

M. Pharmacy (Pharmaceutics) I-Semester (PCI)(Main & Backlog) Examination, January 2019

Subject : Drug Delivery System

	me: 3 Hours ote: Answer any Five Questions. All Questions Carry Equal Marks.	Max. Marks: 75
Note. Answer any Five Questions. All Questions Carry Equal Marks.		
1.	(a) Compare sustained release, delayed release and controlled drug delivery with examples?(b) Define pharmacogenetics? Explain telepharmacy?	system 6 9
2.	(a) What do you mean by activation modulated system? Explain osmotic drug system?(b) Classify rate controlled drug delivery system? Explain feedback regulated	8
3.	(a) What are the barriers of buccal mucoadhesive drug delivery system? Menexamples of marketed mucoadhesive drug delivery system?(b) Explain in detail floating drug delivery systems?	tion two 7
4.	(a) Describe formulation and evaluation of ocular drug delivery system?	10
5.	(a) Explain structure of skin? What are barriers of transdermalPharmacypermeation? What are (b) What are ocular insitu gels? the ideal properties of a drugLibrarycandidate to permeate through the skin? (b) Classify transdermal permeation enhancers? Explain the mechanism of permeation of permeation in the mechanism of permeating in the mec	5 10 ermeation
	enhancers?	5
6.	Reddy (a) Explain in detail various strategies to formulate stable protein and p formulations?	eptide drug 8
7.	(a) Explain in detail mucosal and active transdermal methods of delivering vactoring mucosa-and skin?(b) Write a note on single shot vaccines?	ccines 8 7
8.	(a) Explain various physiochemical and biological factors that influences the S formulations?	SR and CR 10
	(b) Classifypolymers? Explain properties of any two synthetic biodegradable p	oolymers? 5

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M. Pharmacy (Common to All) I-Semester (PCI) (Suppl.) Examination, August 2018

Subject: Modern Pharmaceutical Analytical Techniques Time: 3 Hours Ma	ax. Marks: 75
 Note: Answer any five questions. All questions carry equal mand 1 (a) Discuss the instrumentation of double beam UV visible spectropho with a neat labeled diagram. (b) What is Isobestic point? Explain with a labeled UV spectrum giving examples. 	tometer (10)
 (a) Compare the instrumentation and working a dispersive and foruie IR spectrometers. Write the advantages and disadvantages of the techniques. (b) Draw a schematic IR spectrum for any one compound and indicat absorption wave number regions for any four functional groups in compound. 	e two (10) e the
 (a) Explain (ii) Spin-spin coupling and coupling constant. (b) Draw a schematic HNMR spectrum for any onecompound and explain coupling constant. (i) Chemical shift values (ii) Nature of protons (iii) Number of 	-
 4 (a) Discuss the theory and principle of mass spectroscopy and explain instrumentation and working of mass spectrometer with a neat lab diagram. (b) What is fragmentation? Explain the following by taking a simple explain. 	peled (10)
 (i) Fragmentation peaks (ii) Molecular ion peak (iii) Base peal 5 (a) Discuss the theory of HPLC. Describe the instrumentation and wor of HPLC with a neat labelled diagram. (b) Draw a schematic HPLC chromatomgram and explain (i) Retention time (ii) Resolution (iii) Peak Asymmetry 	` ,
6 (a) Discuss the theory and principle of electrophoresis. Explain the me capillary electrophoresis and its applications with examples.(b) What is isoelectric focusing?	ethod of (12) (3)
 7 (a) Discuss the theory and principle of Gas chromatography. Explain the instrumentation and working of Gas chromatography and explain var stationary and mobile phases used in GC. (b) How non voralile compounds can be analysed by GC. Explain the with few examples? 	ious (11)
8 Write a note on : (a) Flame emission spectroscopy (b) Instrumentation and application of Florescence spectroscopy	(6) (9)

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Suppl.) Examination, August 2018

Subject: Drug Delivery Systems

Time: 3 Hours Max. Marks: 75 Note: Answer any five questions. All questions carry equal marks. 1 Discuss in detail physicochemical and biological factors influencing design of SRDDS. (15)2 (a) Classify polymers and write the applications of polymers in controlled drug delivery systems. (8)(b) Write a note on 3D printing of pharmaceuticals. (7) 3 Write a short note on a) Osmotic activated drug delivery systems. (8)b) Mechanically activated drug delivery system (7) 5 structure of oral mucosa and buccal absorption. (15)(10)b) Write about mucosal and transdermal delivery of vaccines 6 a) Explain various essential components of transdermal drug delivery system. (5) b) Explain in detail various evaluation methods for TDDS. (10)7 Define proteinand peptide delivery systems. Add note on barriers for protein delivery. (15)8 a) What are methods to enhance drug permeation through transdermal route? (7) b) Describe in brief-preparation and evaluation of gastro retentive floating tablets. (8)

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Supple.) Examination, August 2018

Subject: Regulatory Affairs

Time: 3 Hours Max. Marks: 75

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

1	(a) Explain the importance of documentation in pharmaceutical industry and add a not on master formula record, distribution records.(b) Write a note on Code Of Federal Regulation.	ote (10) (5)
2	Explain ANDA regulatory approval process.	(15)
3	What are the regulatory requirements for approval of an API?	(15)
4	Write a note on (a) CTD and eCTD (b) ICH Quality guidelines	(9) (6)
5	Explain the regulatory requirements of EU	(15)
6	Discuss about regulations for Combination products andofMedical devices.	(15)
7	Write a note on (a) informed consent processand procedures (b) Pharmacovigilance safety monitoring	(6) (9)
8	Write note on (a) Investigator broothure	(7)
	(b) investigation of medicinal products dossier	(8)

M. Pharmacy (Pharmaceutics) I-Semester (PCI) (Suppl.) Examination, August 2018

Subject: Modern Pharmaceutics

Ti	me: 3 Hours Max. Marks: 7	75
	Note: Answer any five questions. All questions carry equal marks.	
1	(a) Explain the methods for testing of drug excipient compatability studies.(b) Describe the preparation of SMEDDS and their evaluation tests.(c) Define factorial design. Describe the applications and limitations.	6 6 3
2	(a) Discuss the ICH guidelines for calibration and validation of equipment with an example.(b) Describe the validation of tableting process.	8 7
3	(a) Discuss the sales forecasting and budget planning in pharmaceutical industries.(b) Describe the layout of buildings, services, equipment and their maintenance in industries.	. 8 7
4	 (a) What is compaction profile? Explain the phases of compaction profile with a suitable examples. (b) Define the term intrinsic solubility. How to determine? it's significance? (c) Explain the properties of granules effecting the compression behavior of tablets. 	5 5 .5
5		5
	 (b)Describe the comparison of dissolution Uprofiles of dosage forms using similarity and difference factors. (c) Explain the variance and standardCollegedeviation with it's significance. 	5 5
6	(a) Describe the methods of evaluation of physical stability of emulsions.	5
	(b) Describe the sterilization procedures for evaluation of parenterals.	5 ₅
7	(a) Write the methods for determination of the order of a reaction.(b) Explain the photo degradation and it's testing procedure.	8 7
8	(a) Discuss the methods for improvement of aqueous solubilty of drugs.(b) Write Heckle equation and draw Heckle plots for porosity and explain them.	8 7