

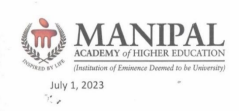
## Academic Program Regulations – 2017

Based on PCI Notification in the Gazette of India, No 362, dated 11 December, 2014

## **<u>Program Title</u>: MPharm (Master of Pharmacy) CBCS (Choice Based Credit System)**

**Specialization:** Pharmaceutical Analysis

Manipal College of Pharmaceutical Sciences Manipal Academy of Higher Education Manipal-576 104, Karnataka, India



#### Academic Program Regulations – 2017 : MPharm, CBCS – Approval

The Master of Pharmacy (MPharm) program, CBCS of Manipal Academy of Higher Education being offered at Manipal College of Pharmaceutical Sciences under the title "Academic Program Regulations – 2017: MPharm, CBCS" has been duly approved by the Academic Council of Manipal Academy of Higher Education.



REGISTRAR



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The Master of Pharmacy (M.Pharm) Course Regulations, 2014

No. 14-136/ 2014-PCL-In exercise of the powers conferred by Sections 10 and 18 of the Pharmacy Act,

1948 (8 of 1948), the Pharmacy Council of India, with the approval of the Central Government hereby makes the following regulations; namely-

#### **CHAPTER I: REGULATIONS**

#### 1. Short title and commencement

These regulations shall be called as "Master of Pharmacy (MPharm) Degree Program-Regulations - Choice Based Credit System (CBCS). The program regulations are based on the PCI notification in the Gazette of India, No. 362, dated 11 December 2014.

They shall come into effect from the academic year 2017-18. The regulations framed are subject to modifications from time to time by the authorities of the Manipal Academy of Higher Education.

#### 2. Minimum qualification for admission

A pass in the following examinations

- a) BPharm degree examination of an Indian university established by law in India from an institution approved by Pharmacy Council of India and has scored not less than 55% of the maximum marks (aggregate of 4 years of BPharm).
- b) Every student, selected for admission to postgraduate pharmacy program in any PCI approved institution should have obtained registration with the State Pharmacy Council or should obtain the same within one month from the date of his/her admission, failing which the admission of the candidate shall be cancelled.
- c) Any foreign pharmacy degree approved by the Pharmacy Council of India.

Note: It is mandatory to submit a migration certificate obtained from the respective university where the candidate had passed his/her qualifying degree (BPharm)

#### **3. Duration of the program**

The program of study for MPharm shall extend over a period of four semesters (two academic years). The curricula and syllabi for the program shall be prescribed from time to time by Manipal Academy of Higher Education based on the inputs from Pharmacy Council of India, New Delhi.

#### 4. Medium of instruction and examination

Medium of instruction and examination shall be in English.

#### 5. Working days in each semester

Each semester shall consist of not less than 100 working days. The odd semesters shall be conducted from the month of July/August to November/December and the even semesters shall be conducted for the month of December/January to May/June in an academic year.

#### 6. Attendance and progress

A candidate is required to put in at least 80% attendance in individual courses considering the theory and practical separately. The candidate shall complete the prescribed course satisfactorily to be eligible to appear for the respective examinations.

#### 7. Program/Course credit structure

As per the philosophy of Credit Based Semester System, certain quantum of academic work viz. theory classes, practical classes, seminars etc., are measured in terms of credits. On satisfactory completion of the courses, a candidate earns credits. The amount of credit associated with a course is dependent upon the number of hours of instruction per week in that course. Similarly, the credit associated with any of the other academic, co-curricular activities is dependent upon the quantum of work expected to be put in for each of these activities per week/per activity.

#### 7.1. Credit assignment

#### 7.1.1. Theory and laboratory courses

Courses are broadly classified as Theory and Practical. Theory courses consist of Lecture (L) and tutorial (T). Practical (P) courses consist of hours spent in the laboratory. Credits (C) for a course are dependent on the number of hours of instruction per week in that course, and is obtained by using a multiplier of one (1) for lecture/tutorial and a multiplier of half ( $\frac{1}{2}$ ) for practical (laboratory) hours. Thus, for example, a theory course, having four lectures per week throughout the semester carries a credit of 4. Similarly, a practical having four laboratory hours per week throughout the semester carries a credit of 2. The contact hours of seminars, and research work and journal club shall be treated as that of practical courses for the purpose of calculating credits, i.e., the contact hours shall be multiplied by  $\frac{1}{2}$ .

#### 7.2. Minimum credit requirements

The minimum credit points required for the award of MPharm degree is 95. However, based on the credit points earned by the students under the heads of co-curricular activities and choice based inter/multidisciplinary courses, a student shall earn a maximum of 100 credit points. These credits are divided into theory courses, practical, seminars, research work, journal club and co-curricular activities over the duration of four semesters. The credits are distributed semester-wise as shown in Table 14. Courses generally progress in sequence, building competencies and their positioning indicate certain academic maturity on the part of the learners. Learners are expected to follow the semester-wise schedule of courses given in the syllabus.

#### 8. Academic work

A regular record of attendance both in theory, practical, seminar, assignment, journal club, discussion with the supervisor, research work presentation and dissertation shall be maintained by the department/teaching staff of the respective courses.

#### 9. Course work of study

The specializations in MPharm program are given in Table 1.

Table 1.	Table 1. List of MPharm specializations and their codes						
S. No.	Specialization	Code					
1	Pharmaceutics	MPH					
2	Industrial Pharmacy	MIP					
3	Pharmaceutical Chemistry	MPC					
4	Pharmaceutical Analysis	MPA					
5	Pharmaceutical Quality Assurance	MQA					
6	Pharmaceutical Regulatory Affairs	MRA					
7	Pharmaceutical Biotechnology	MPB					
8	Pharmacy Practice	MPP					
9	Pharmacology	MPL					
10	Pharmacognosy	MPG					

The course work of MPharm specializations shall include semester wise theory and practical as given in Table 2 to 13. The number of hours to be devoted to each theory lectures (L), tutorials (T) and practical (P) course in any semester shall not be less than that shown in Table 2 to 13.

Table 2.	Course work of MPharm – P	harmace	eutics (MF	PH) special	ization	
Course	Course Title	Cre	dit hours	Credit	Marks	
Code		Lecture	Tutorial	Practical	Points	
		(L)	<b>(T)</b>	<b>(P</b> )		
Semester I						
PQA-MPH101T	Modern Pharmaceutical	4			4	100
	Analytical Techniques					
PCE-MPH102T	Drug Delivery Systems	4	1		5	100
PCE-MPH103T	Modern Pharmaceutics	4	1		5	100
PRM-MPH104T	Regulatory Affairs	4	1		5	100
PCE-MPH105P	Pharmaceutics Practical I			12	6	150
PCE-MPH106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II						
PCE-MPH201T	Molecular Pharmaceutics	4	1		5	100
	(Nano Tech and Targeted					
	DDS)					
PCE-MPH202T	Advanced Biopharmaceutics	4	1		5	100
	and Pharmacokinetics					
PCE-MPH203T	Computer Aided Drug	4	1		5	100
	Delivery Systems					
PCE-MPH204T	Cosmetic and	4	1		5	100
	Cosmeceuticals					
PCE-MPH205P	Pharmaceutics Practical II			12	6	150
PCE-MPH206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous i	node				

Table 3. Course work of MPharm –Industrial Pharmacy (MIP) specialization						
Course	Course Title	Cre	dit hours	/week	Credit	Marks
Code		Lecture	Tutorial	Practical	Points	
		(L)	<b>(T)</b>	<b>(P)</b>		
Semester I						
PQA-MIP101T	Modern Pharmaceutical	4			4	100
	Analytical Techniques					
PCE-MIP102T	Pharmaceutical Formulation	4	1		5	100
FCE-IVIIF 1021	Development					
PCE-MIP103T	Novel Drug Delivery	4	1		5	100
FCE-IVIIF 1051	Systems					
PRM-MIP104T	Intellectual Property Rights	4	1		5	100
PCE-MIP105P	Industrial Pharmacy			12	6	150
PCE-WIP103P	Practical I					
PCE-MIP106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II						
PCE-MIP201T	Advanced Biopharmaceutics	4	1		5	100
FCE-IVIIF2011	and Pharmacokinetics					
PCE-MIP202T	Scale-up and Technology	4	1		5	100
PCE-IVIIP2021	Transfer					
PCE-MIP203T	Pharmaceutical Production	4	1		5	100
PCE-IVIIP2051	Technology					
PRM-MIP204T	Entrepreneurship	4	1		5	100
PRM-MIP2041	Management					
PCE-MIP205P	Industrial Pharmacy			12	6	150
PCE-WIP203P	Practical II					
PCE-MIP206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	mode.				

Table 4. Course work of MPharm – Pharmaceutical Chemistry (MPC) specialization						
Course	Course Title	Cre	dit hours	/week	Credit	Marks
Code		Lecture	Tutorial	Practical	Points	
		(L)	<b>(T)</b>	<b>(P</b> )		
Semester I						
PQA-MPC101T	Modern Pharmaceutical	4			4	100
	Analytical Techniques					
PCH-MPC102T	Advanced Organic	4	1		5	100
FCH-MFC1021	Chemistry I					
PCH-MPC103T	Advanced Medicinal	4	1		5	100
FCH-MFC1051	Chemistry					
PCH-MPC104T	Chemistry of Natural	4	1		5	100
FCH-MFC1041	Products					
PCH-MPC105P	Pharmaceutical Chemistry			12	6	150
FCH-MFC103F	Practical I					
PCH-MPC106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II						
PCH-MPC201T	Advanced Spectral Analysis	4	1		5	100
PCH-MPC202T	Advanced Organic	4	1		5	100
FCH-MFC2021	Chemistry II					
PCH-MPC203T	Computer Aided Drug	4	1		5	100
1 CH-IMI C2031	Design					
PCH-MPC204T	Pharmaceutical Process	4	1		5	100
r C11-Ivir C2041	Chemistry					
PCH-MPC205P	Pharmaceutical Chemistry			12	6	150
1 C11-WIF C203F	Practical II					
PCH-MPC206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	node.				

Table 5. Cou	rse work of MPharm – Pharm	naceutica	l Analysis	s (MPA) sp	ecializat	ion
Course	Course Title	Cre	dit hours	/week	Credit	Marks
Code		Lecture	Tutorial	Practical	Points	
		(L)	<b>(T</b> )	<b>(P</b> )		
Semester I						
PQA-MPA101T	Modern Pharmaceutical Analytical Techniques	4			4	100
PCH-MPA102T	Advanced Pharmaceutical Analysis	4	1		5	100
PCH-MPA103T	Pharmaceutical Validation	4	1		5	100
PCH-MPA104T	Food Analysis	4	1		5	100
PCH-MPA105P	Pharmaceutical Analysis Practical I			12	6	150
PCH-MPA106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II					•	•
PCH-MPA201T	Advanced Instrumental Analysis	4	1		5	100
PCH-MPA202T	Modern Bioanalytical Techniques	4	1		5	100
PCH-MPA203T	Quality Control and Quality Assurance	4	1		5	100
PCH-MPA204T	Herbal and Cosmetic Analysis	4	1		5	100
PCH-MPA205P	Pharmaceutical Analysis Practical II			12	6	150
PCH-MPA206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	mode.				

Table 6.	ourse work of MPharm – Pha specializ		cal Qualit	y Assuran	ce (MQA	.)
Course	Course Title	Credit hours/week			Credit	Marks
Code		Lecture (L)	Tutorial (T)	Practical (P)	Points	
Semester I						
PQA-MQA101T	Modern Pharmaceutical Analytical Techniques	4			4	100
PQA-MQA102T	Quality Management Systems	4	1		5	100
PQA-MQA103T	Quality Control and Quality Assurance	4	1		5	100
PQA-MQA104T	Product Development and Technology Transfer	4	1		5	100
PQA-MQA105P	Pharmaceutical Quality Assurance Practical I			12	6	150
PQA-MQA106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II						
PQA-MQA201T	Hazards and Safety Management	4	1		5	100
PQA-MQA202T	Pharmaceutical Validation	4	1		5	100
PQA-MQA203T	Audits and Regulatory Compliance	4	1		5	100
PQA-MQA204T	Pharmaceutical Manufacturing Technology	4	1		5	100
PQA-MQA205P	Pharmaceutical Quality Assurance Practical II			12	6	150
PQA-MQA206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	mode.				

Table 7. C	ourse work of MPharm – Pha specializ		cal Regula	atory Affai	irs (MRA	r)
Course	Course Title	Cre	edit hours	/week	Credit	Marks
Code		Lecture (L)	Tutorial (T)	Practical (P)	Points	
Semester I						
PRM-MRA101T	Good Regulatory Practices	4			4	100
PRM-MRA102T	Documentation and Regulatory Writing	4	1		5	100
PRM-MRA103T	Clinical Research Regulations	4	1		5	100
PRM-MRA104T	Regulations and Legislation for Drugs & Cosmetics, Medical Devices, Biologicals & Herbals and Food & Nutraceuticals in India and Intellectual Property Rights	4	1		5	100
PRM-MRA105P	Regulatory Affairs Practical I			12	6	150
PRM-MRA106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II			•			
PRM-MRA201T	Regulatory Aspects of Drugs and Cosmetics	4	1		5	100
PRM-MRA202T	Regulatory Aspects of Herbal and Biologicals	4	1		5	100
PRM-MRA203T	Regulatory Aspects of Medical Devices	4	1		5	100
PRM-MRA204T	Regulatory Aspects of Food and Nutraceuticals	4	1		5	100
PRM-MRA205P	Regulatory Affairs Practical II			12	6	150
PRM-MRA206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	mode.				

Table 8. Course	e work of MPharm – Pharmac	eutical B	iotechnol	ogy (MPB)	specializ	zation
Course	Course Title	Cre	dit hours	Credit	Marks	
Code		Lecture	Tutorial	Practical	Points	
		(L)	<b>(T)</b>	<b>(P</b> )		
Semester I						
PQA-MPB101T	Modern Pharmaceutical	4			4	100
	Analytical Techniques					
PBT-MPB102T	Microbial and Cellular	4	1		5	100
FD1-WIFD1021	Biology					
PBT-MPB103T	Bioprocess Engineering and	4	1		5	100
FD1-MFD1031	Technology					
PBT-MPB104T	Advanced Pharmaceutical	4	1		5	100
r D I -Ivir D I 04 I	Biotechnology					
PBT-MPB105P	Pharmaceutical			12	6	150
	Biotechnology Practical I					
PBT-MPB106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II						
PBT-MPB201T	Proteins and Protein	4	1		5	100
	Formulations					
PBT-MPB202T	Immunotechnology	4	1		5	100
	Bioinformatics and	4	1		5	100
PBT-MPB203T	Computational					
	Biotechnology					
PBT-MPB204T	Biological Evaluation of	4	1		5	100
	Drug Therapy					
PBT-MPB205P	Pharmaceutical			12	6	150
	Biotechnology Practical II					
PBT-MPB206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	node.				

Table 9. (	Course work of MPharm – Ph	armacy F	Practice (N	(IPP) speci	alization	
Course	Course Title	Cre	dit hours	/week	Credit	Marks
Code		Lecture	Tutorial	Practical	Points	
		(L)	<b>(T)</b>	<b>(P</b> )		
Semester I						
PPR-MPP101T	Clinical Pharmacy Practice	4			4	100
PPR-MPP102T	Pharmacotherapeutics I	4	1		5	100
PPR-MPP103T	Hospital and Community	4	1		5	100
11 K-WI11 1031	Pharmacy					
PPR-MPP104T	Clinical Research	4	1		5	100
PPR-MPP105P	Pharmacy Practice Practical I			12	6	150
PPR-MPP106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II						
PPR-MPP201T	Principles of Quality Use of	4	1		5	100
11 K-IVII I 2011	Medicines					
PPR-MPP202T	Pharmacotherapeutics II	4	1		5	100
	Clinical Pharmacokinetics	4	1		5	100
PPR-MPP203T	and Therapeutic Drug					
	Monitoring					
PPR-MPP204T	Pharmacoepidemiology and	4	1		5	100
-	Pharmacoeconomics					
PPR-MPP205P	Pharmacy Practice Practical II			12	6	150
PPR-MPP206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	node.				

Table 10	. Course work of MPharm –	Pharmac	ology (M	PL) special	ization	
Course	Course Title	Credit hours/week		/week	Credit	Marks
Code		Lecture	Tutorial	Practical	Points	
		(L)	<b>(T</b> )	<b>(P</b> )		
Semester I						
PQA-MPL101T	Modern Pharmaceutical	4			4	100
	Analytical Techniques					
PHA-MPL102T	Advanced Pharmacology I	4	1		5	100
	Pharmacological and	4	1		5	100
PHA-MPL103T	Toxicological Screening					
	Methods I					
PHA-MPL104T	Cellular and Molecular	4	1		5	100
rna-mrl1041	Pharmacology					
PHA-MPL105P	Pharmacology Practical I			12	6	150
PHA-MPL106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II						
PHA-MPL201T	Advanced Pharmacology II	4	1		5	100
	Pharmacological and	4	1		5	100
PHA-MPL202T	Toxicological Screening					
	Methods II					
PHA-MPL203T	Principles of Drug Discovery	4	1		5	100
PHA-MPL204T	Clinical Research and	4	1		5	100
PHA-MPL2041	Pharmacovigilance					
PHA-MPL205P	Pharmacology Practical II			12	6	150
PHA-MPL206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous a	node.				

Table 11.	Course work of MPharm – P	harmaco	gnosy (M	PG) specia	lization	
Course	Course Title	Cre	dit hours	Credit	Marks	
Code		Lecture	Tutorial	Practical	Points	
		(L)	<b>(T)</b>	<b>(P</b> )		
Semester I						
PQA-MPG101T	Modern Pharmaceutical Analytical Techniques	4			4	100
PCO-MPG102T	Advanced Pharmacognosy I	4	1		5	100
PCO-MPG1021 PCO-MPG103T	Phytochemistry	4	1		5	100
	Industrial Pharmacognostical	4	1		5	100
PCO-MPG104T	Technology	4	1		5	100
PCO-MPG105P	Pharmacognosy Practical I			12	6	150
PCO-MPG106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II						
PCO-MPG201T	Medicinal Plant Biotechnology	4	1		5	100
PCO-MPG202T	Advanced Pharmacognosy II	4	1		5	100
PCO-MPG203T	Indian Systems of Medicine	4	1		5	100
PCO-MPG204T	Herbal Cosmetics	4	1		5	100
PCO-MPG205P	Pharmacognosy Practical II			12	6	150
PCO-MPG206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	mode.				

Table 13. Course work for MPharm III and IV semesters(Common for all specializations)						
Course	Course Title	Credit hours/week Credit Ma				
Code		Lecture (L)	Tutorial (T)	Practical (P)	Points	
PHA-MRM301T	Research Methodology and Biostatistics*	4			4	100
MJC302P	Journal Club*			2	1	100
MRW401P	Research Work			70	35	600
Total 4 72 40 800						
* No end-semester examination. Only continuous mode						

Table 14. Semester wise course work credits distribution				
Semester	Credit Points			
Ι	26			
П	27			
III and IV	40			
Total course work credits	93			
Co-curricular activities (Attending conference, scientific presentations, other scholarly activities and choice based inter/multidisciplinary courses)	Minimum=02* Maximum=07*			
Total credit points	Minimum=95 Maximum=100			

\*Credit points for co-curricular activities (Table 15A) and choice based inter/multidisciplinary courses (Table 15B).

Table 15A. Guidelines for awarding credit points for co-curricular activities			
Name of the Activity	Maximum Credit Points Eligible/ Activity		
Participation in National level seminar/Conference/Workshop/Symposium/Training programs (related to the specialization of the student)	01		
Participation in International level seminar/Conference/Workshop/Symposium/Training programs (related to the specialization of the student)	02		
Academic award/ Research award from State level/National agencies	01		
Academic award/Research award from International agencies	02		
Research/ Review publication in National journals (Indexed in Scopus/Web of Science)	01		
Research/ Review publication in International journals (Indexed in Scopus/Web of Science)	02		

Note: International conference: Held outside India

International journal: The editorial board outside India

\*The credit points assigned for extracurricular and or co-curricular activities shall be given by the principal of the college and the same shall be submitted to the university. The criteria to acquire this credit point shall be defined by the college from time to time.

	Table 15B. List of choice based inter/multidisciplinary courses						
Course Code	Course Title	Credits	Department/Institution offering the Course				
Interdisciplina	ary courses						
<b>PCE-001E</b>	Generic Drug Development	1	Pharmaceutics, MCOPS				
PCE-002E	Pharmaceutical Dissolution Technology	1	Pharmaceutics, MCOPS				
PCE-003E	Particulate Drug Delivery Systems	1	Pharmaceutics, MCOPS				
PCE-004E	3D Printing in Pharmaceutical Manufacturing	1	Pharmaceutics, MCOPS				
PCH-001E	Preparative Separation Techniques	1	Pharmaceutical Chemistry, MCOPS				
PCH-002E	Molecular Modeling and Drug Design	1	Pharmaceutical Chemistry, MCOPS				
PCH-003E	Hyphenated Techniques	1	Pharmaceutical Chemistry, MCOPS				
<b>PCH-004E</b>	Chemicals - Environment, Health and Safety	1	Pharmaceutical Chemistry, MCOPS				
PQA-001E	Theory and Practice of Analytical and Bioanalytical Method Development and Validation	1	Pharmaceutical Quality Assurance, MCOPS				
PQA-002E	Good Documentation Practices and e-Documentation Practices in Pharmaceutical Industry	1	Pharmaceutical Quality Assurance, MCOPS				
PQA-003E	Trouble Shooting in High Performance Liquid Chromatography	1	Pharmaceutical Quality Assurance, MCOPS				

PQA-004E	Professional Development	1	Pharmaceutical Quality Assurance, MCOPS	
PQA-005E	Stability Testing of Drugs and Biologicals	1	Pharmaceutical Quality Assurance, MCOPS	
PQA-006E	USFDA Drug Regulatory Affairs	1	Pharmaceutical Quality Assurance, MCOPS	
PQA-007E	Rest of the World Drug Regulations	1	Pharmaceutical Quality Assurance, MCOPS	
PQA-008E	Evaluation of Medical Devices	1	Pharmaceutical Quality Assurance, MCOPS	
<b>PBT-001E</b>	Clean Room Concepts	1	Pharmaceutical Biotechnology, MCOPS	
<b>PBT-002E</b>	Biosimilars	1	Pharmaceutical Biotechnology, MCOPS	
<b>PBT-003E</b>	Principles of Gene Cloning	1	Pharmaceutical Biotechnology, MCOPS	
<b>PBT-004E</b>	Tissue Engineering	1	Pharmaceutical Biotechnology, MCOPS	
PPR-001E	Retail Pharmacy Practice	1	Pharmacy Practice, MCOPS	
<b>PPR-002E</b>	Fundamentals of Medical Writing	1	Pharmacy Practice, MCOPS	
PPR-003E	Systematic Review and Meta- Analysis	1	Pharmacy Practice, MCOPS	
PPR-004E	Pharmacokinetics Data Analysis (Employing WinNonlin)	1	Pharmacy Practice, MCOPS	
PHA-001E	Cancer Biology	1	Pharmacology, MCOPS	
РНА-002Е	Screening Methods for Drug Development	1	Pharmacology, MCOPS	
РНА-003Е	Free Radical Biology and Medicine	1	Pharmacology, MCOPS	
РНА-004Е	Regulatory Toxicology in Drug Discovery and Development	1	Pharmacology, MCOPS	
PCO-001E	Nutraceuticals	1	Pharmacognosy, MCOPS	
PCO-002E	Extraction, Separation and Purification of Phytoconstituents	1	Pharmacognosy, MCOPS	
PCO-003E	Nanophytopharmaceuticals	1	Pharmacognosy, MCOPS	
PCO-004E	Herbal Monographs	1	Pharmacognosy, MCOPS	
PRM-001E	Retail Business Management	1	Pharmaceutical Regulatory Affairs & Management, MCOPS	
PRM-002E	Intellectual Property Management	1	Pharmaceutical Regulatory Affairs & Management, MCOPS	
PRM -003E	General Management Principles	1	Pharmaceutical Regulatory Affairs & Management, MCOPS	
PRM -004E	Entrepreneurship Development	1	Pharmaceutical Regulatory Affairs & Management, MCOPS	
Multidisciplin	ary courses			
MU-001E	Certificate Course in Bioinformatics	3	School of Life Sciences, MU	
MU-002E	Project Management	4	Department of Humanities and Social Science, MIT	
MU-003E	Certificate Course in Bioethics	2/4	Centre for Bioethics, MU	
MU-004E	Academic Research and Writing	3	Manipal Centre for Philosophy and Humanities, MU	
MU-005E	Certificate Course in Biosecurity	5	Dept. of Public Health, MU	
CR-001E	Any one of the Online courses	1 and	Coursera	

#### **10. Program committee**

1. The MPharm program shall have a program committee constituted by the Head of the Institution in consultation with all the Heads of the Departments.

2. The composition of the program committee shall be as follows: A teacher at the cadre of professor shall be the Chairperson; One teacher from each MPharm specialization and four student representatives (two from each academic year), nominated by the Head of the Institution.

- 3. Duties of the program committee:
  - i. Periodically reviewing the progress of the classes.
  - ii. Discussing the problems concerning curriculum, syllabus and the conduct of classes.
  - iii. Discussing with the course teachers on the nature and scope of assessment for the course and the same shall be announced to the students at the beginning of respective semesters.
  - iv. Communicating its recommendation to the Head of the Institution on academic matters.
  - v. The program committee shall meet at least twice in a semester, preferably at the end of each sessional exam and before the end semester exam.

#### 11. Examinations/Assessments

The schemes for internal assessment and end-semester examinations are given in Table 16.

Table 16. Schemes for internal assessments and end semester examinations							
	Internal Assessment			End-Semester Exams			
Course	Contin	Session	al Exams		Marks	Duration	Total
Course	uous Mode	Marks	Duration	Total			Marks
			Semester I a	and II			
Theory	10	15	1 hr each	25	75	3 hrs	100
Practical	20	30	6 hrs	50	100	6 hrs	150
Seminar				100			100
		S	emester III	and IV			
PHA-MRM301T Research Methodology and Biostatistics*	20	40+40	2 hrs each	100			100
MJC302P Journal Club*				100			100
MRW401P Research Work		100+100	1 hr each	200	400		600
* No end-semester ex	* No end-semester examination. Only continuous mode						

#### 11.1. Internal assessment: Continuous mode

The marks allocated for continuous mode of internal assessment of theory courses shall be awarded based on the students' performance in the assignments/surprise tests, etc., while in the lab course it is based on the practical record, regular viva-voce etc.

#### 11.1.1. Sessional exams

Two sessional exams for each theory course and one sessional examination for a practical course shall be conducted as per the schedule fixed by the college. The schemes of question papers for theory and practical sessional examinations are given below. The average marks of two sessional examinations of theory courses shall be computed for internal assessment of 15 marks as given in Table 16.

Question paper pattern – MPharm Theory sessional examinations						
Manipal College of Pharmaceutical Sciences						
Manipal Academy of Higher Education, Manipal						
MPharm Theo	MPharm Theory Sessional Examinations, Month and Year					
	Course Code. Course Title					
Date: dd-mm-yyyy	Duration: 2 hrs	Max. Marks: 45				
Ins	tructions: Answer ALL questions					
Long Essays (2x 10 marks) = 20	) marks					
1. Question						
2. Question						
Short Essays $(4 \times 5 \text{ marks}) = 20$	marks					
3. Question						
4. Question						
5. Question						
6. Question						
7. Short answers (1 mark $\times$ 5 =	5 marks)					
7A.						
7B.						
7C.						
7D.						
7E.						

Question paper pattern – MPharm practical sessional examinations					
Manipal College of Pharmaceutical Sciences					
Manipal Academy of Higher Education, Manipal					
MPharm Practical Sessional Examinations, Month and Year					
<u>Course Code. Course Title</u>					
Date: dd-mm-yyyy	Duration: 6 hrs	Max. Marks: 60			
Instruc	tions: Answer ALL questions.				
1. Synopsis (10 marks)					
2. Major Experiment (25 marks)					
3. Minor Experiment (15 marks)					

4. Viva-Voce (10 marks)

	MPharm seminar evaluation scheme						
	PRE	SENTATION (	Marks awarded for each criteria				
		Criteria			Те	acher 1	Teacher 2
1	Preparedness	(10 marks)					
2	Response to q	uestions (10 mar	ks)				
3	Audio-visual	aids (10 marks)					
4	Clarity of pres	sentation (10 mai	·ks)				
5	Breadth and d	epth of material	presented (10 m	arks)			
			Marks a	awarded			
		Average mark	s awarded for p	resentatio	n out o	f 50 (A) =	
WR	RITE UP (50 Ma	arks)					
Ma	rks awarded for	each criterion					
	Content	Recent	Organization	Diagr	am,	Originality	Marks
(0	optimum and	information	(sequent and	illustra	tions	(10 marks)	awarded for
rele	evant to topic)	opic) or out of date methodical) & references			write up out of 50 (B)		
	(10 marks)	(10 marks)	(10 marks)	(10 marks) (10 marks)			01 50 (B)
Rer	Remarks if any:						
	Seminar marks awarded out of 100 = (A+B) =						

#### 11.2 End-semester examination

End-semester examinations are conducted for eligible students twice at the end of the semester, namely, the main examination for regular students and the make-up/supplementary examinations for the failed students only before the commencement of the next semester (Table 17). In case a failed student could not clear the course in the make-up/supplementary examination, the student would have his next examination along with the regular students only in the main examination.

Table 17. Tentative schedule of end-semester examinations					
Semester Main Examination Make-up/Supplementary Exams					
I and III	November/December	December/January			
II and IV	May/June	July/August			

Question paper pattern – MPharm theory end-semester examinations						
Manip	Manipal Academy of Higher Education, Manipal					
<u>MPharm The</u>	MPharm Theory End-Semester Examinations, Month and Year					
	Course Code. Course Title					
Date: dd-mm-yyyy	Duration: 3 hrs	Max. Marks: 75				
]	Instructions: Answer ALL questions.					
Answer the following (5 mark	$ks \times 10 = 50$ marks)					
1. Question						
2. Question						
3. Question						
4. Question						
5. Question						
Answer the following with sp	Answer the following with specific answers (5 marks $\times$ 5 = 25 marks)					
6A.						
6B.						
6C.						
6D.						
6E.						

Question paper pattern – MPharm practical end-semester examinations						
<u>MPharm Practi</u>	MPharm Practical End-Semester Examinations, Month and Year					
Manipal Academy of Higher Education, Manipal						
Course Code. Course Title						
Date: dd-mm-yyyy	Duration: 6 hrs	Max. Marks: 100				
In	structions: Answer ALL questions.					
1. Synopsis (15 marks)						
2. Major Experiment (45 marks)						
3. Minor Experiment (25 marks)						
4. Viva-Voce (15 marks)						

#### 12. Pass and award of performance grades

#### 12.1: Minimum for a pass in a course

A student should obtain a minimum of 35% marks in the end-semester exam of each course. A student shall be declared PASS if, the candidate secures E-grade separately in each course, in a 10-Point-Relative-Letter Grading-Scheme. Accordingly, no candidate shall be declared to have passed in any course unless he/she obtains a grade not less than E-grade.

#### 12.2 Award of performance grades

The marks obtained in the end semester and internal assessments in a course are added together and a 10-Point-Relative-Letter Grading Scheme is used to allot an appropriate letter grade to the student's performance in that course.

#### 12.3 The 10-Point-Relative-Letter-Grading- Scheme

The letter grades and grade points that are used to assess the students' performance in a course are given in the Table 18

Table 18. 10-Point-Relative-Letter Grading-Scheme			
Letter Grade	Grade Point	Performance	
A+	10	Outstanding	
А	9	Excellent	
В	8	Good	
С	7	Fair	
D	6	Average	
E	5	Pass	
F/I/DT/ab	0	Fail	

F: Fails, I: Incomplete, DT: Detained, ab: Absent

#### Note the following:

- 1. Internal assessment marks and end-semester examination marks put together are taken into account for the 10-Point-Relative-Letter Grading-Scheme in each course separately. However, the scheme is applied to a student who scores minimum 35% marks in the end-semester examination of each course.
- 2. Appropriate letter grades, from E to A+, are awarded, in each theory and lab courses, to only such candidate who has passed the course in first attempt. However, grades E to C are only awarded to a student who makes multiple attempts to pass a course; except in case of I-grade.
- 3. A candidate who is eligible and registers for the end-semester examination but fails to appear in the end-semester examination or fails in the course gets a grade 'ab', indicating failure.
- 4. A student who is eligible and registers for the end-semester examination but fails to appear in the end-semester examination due to valid reasons will get a grade 'I', indicating incomplete. However, it needs prior approval of the HOI and the Registrar Evaluation, Manipal Academy of Higher Education.
- 5. The grade DT is given to a candidate who fails to put in the minimum required attendance for appearing the end-semester examination for a course.
- 6. A student who earns a grade 'E' or above in a course is declared to have successfully completed the course and earns the credits assigned to that particular course. A course successfully completed cannot be repeated for the purpose of improving the grade.
- 7. Final evaluation of the performance of a student in each course (theory and lab separately) will be carried out on a 10-Point-Letter Grading-Scheme corresponding to the marks obtained in that course. Eventually each letter grade of the course is converted into a specific grade point associated with the letter grade as given in the table above.

# <u>Note</u>: For the calculation of SGPA and CGPA, the credits assigned for course work are only taken for account.

The performance of a student in a semester is indicated by a number called 'Semester Grade Point Average' (SGPA). The SGPA is the weighted average of the grade points obtained in all the courses by the student during the semester. For example, if a student takes four courses (Theory/Practical) in a semester with credits C1, C2, C3 and C4 and the student's grade points in these courses are G1, G2, G3 and G4, respectively, and then students' SGPA is equal to:

$$SGPA = \frac{C1G1 + C2G2 + C3G3 + C4G4}{C1 + C2 + C3 + C4}$$

The SGPA is calculated to two decimal points. It should be noted that, the SGPA for any semester shall take into consideration the F and ab-grade awarded in that semester. For example if a learner has F or ab-grade in course 4, the SGPA shall then be computed as:

$$SGPA = \frac{C1G1 + C2G2 + C3G3 + C4 * ZERO}{C1 + C2 + C3 + C4}$$

#### 12.5. Cumulative Grade Point Average (CGPA)

The CGPA is calculated with the SGPA of all the IV semesters to two decimal points and is indicated in the final grade report card/final transcript showing the grades of all IV semesters and their courses. The CGPA shall reflect the failed status in case of F grade(s), till the course(s) is/are passed. When the course(s) is/are passed by obtaining a pass grade on subsequent examination(s) the CGPA shall only reflect the new grade and not the fail grades earned earlier. The CGPA is calculated as:

$$CGPA = \frac{C1S1 + C2S2 + C3S3 + C4S4}{C1 + C2 + C3 + C4}$$

where C1, C2, C3,.... is the total number of credits for semester I,II,III,.... and S1,S2, S3,.... is the SGPA of semester I,II,III,.....

#### 12.6. Conversion of GPA/CGPA into a percentage

The performance of students who are pursuing pharmacy programs in the Manipal College of Pharmaceutical Sciences, Manipal Academy of Higher Education, Manipal is awarded on a 10-Point-Relative-Letter Grading-Scheme.

In this system the top band (top 10%) of students who score highest marks are placed at A+ which is equivalent to 10 grade point (maximum). Thus, 10 GPA or CGPA is equivalent to 100%. Accordingly, 1 GPA or CGPA is equivalent to 10%.

Based on this, the following formula is applied to convert GPA or CGPA to an appropriate percentage:

Percentage secured by the candidate = GPA or CGPA  $\times$  10

#### 13. Make-up/Supplementary examination

In case, a student fails to secure an E-grade in any theory or practical courses, he/she shall reappear for the end-semester examination of that course. However, his/her marks of the internal assessment shall be carried over, and he/she is entitled for a maximum grade of 'C' only, irrespective of the students' top order performances.

However, the candidates with DT-grade will also be allowed to take this examination provided they meet the eligibility criteria laid for the candidates to appear in the end-semester examinations of the courses of the programs.

**Important to Note:** A student who once failed (F-grade) or DT (Detained) grade in any course, a maximum of C-grade will only be awarded in subsequent end-semester examinations, irrespective of the student's high performances in that particular course. However, those who miss regular examinations due to valid reasons (I-grade) will be allowed to retain whatever the grades they secure in the make-up/supplementary examinations. In case a candidate with DT-grade reregisters for the course to repeat the entire education process in the subsequent academic year, after paying the required course fee, the DT status in that particular course is removed and he/she is allowed to retain the grades that he/she secures in the end-semester examination.

After the results are declared, grade cards will be issued to each student, which will contain the list of courses for that semester and the grades obtained by the student.

#### 14. Improvement of internal assessment

A student shall have the opportunity to improve his/her performance only once in the sessional exam component of the internal assessment. The re-conduct of the sessional exam shall be completed before the commencement of next end-semester theory examinations.

#### 15. Promotion to the next higher class

A student shall be eligible to carry forward all the courses of I and II semesters till the III/IV semester examinations. However, the results of IV semester (Research project) of the student shall be declared only when the student passes in all the courses of I and II semesters.

#### **16. Declaration of class**

The class shall be awarded on the basis of CGPA as follows:

First Class with Distinction	= CGPA of 7.50 and above
First Class	= CGPA of 6.00 to 7.49
Second Class	= CGPA of 5.00 to 5.99

#### 17. Research project work

#### **17.1 Dissertation submission**

All the students shall undertake a research project under the supervision of a teacher in semester III and IV and submit a dissertation at the end of the IV semester. Four copies of the dissertation shall be submitted.

**Note:** If any candidate fails to submit the dissertation on or before the date prescribed, the end-semester examination of such candidate shall be at a later date, which shall not be earlier than 6 months from the date fixed in the first instance.

#### **17.2 Dissertation evaluation**

The performance of student in the dissertation work is assessed as per the scheme given in the Table 19.

Table 19. MPharm dissertation evaluation scheme								
Internal Assessment		University Examination						
Presentation 1	<b>Presentation 2</b>	Total	Dissertation		Viva Voce		Total	Grand
(III semester)	(IV semester)		Evaluation (300)		Joint			Total
			by Examiners		Evaluati	ion by		
			Internal and					
					External			
					Examiners			
					(100)			
			Internal	External	Presenta	Viva-		
					tion	voce		
i	ii	i+ii=A	i	ii	iii	iv	i+ii+i	A+B
							ii+iv	
							=B	
100	100	200	150	150	50	50	400	600

The internal and external examiners appointed by the university shall evaluate the dissertation of the research project separately as per the following evaluation criteria.

Evaluation of Dissertation: For 150 marks each separately by Internal and External Examiners		
	Marks	
Objective(s) of the study	25	
Literature search	25	
Methodology adopted	30	
Results and discussions	30	
Conclusions and outcomes	20	
Bibliography	20	
_		
Total	150	

#### **Evaluation of Presentation and Viva-voce: For 100 marks** jointly by Internal and External Examiners

		Marks
Presentation of work		30
Communication skills		20
	Total	50
Viva-voce		50

#### 18. Award of degree

Candidates who fulfill the requirements mentioned above shall be eligible for award of degree.

#### **19.** Duration for completion of the program

The duration for the completion of the program shall be four academic years (double the duration of the prescribed duration). In case, a student is not able to complete the program within this period, the candidate has to reregister for the program.

#### **20. Revaluation of answer papers**

There is a provision for revaluation of the answer papers of the end-semester examination as per Manipal Academy of Higher Education policy, however, the candidates have to apply separately by paying the prescribed fee.

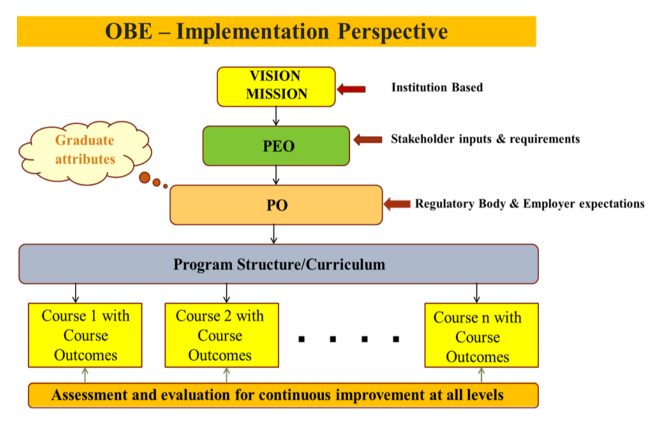
#### 21. Re-admission after break of study

The candidate who seeks re-admission to the program after taking a break of the study, has to get the approval from the university.

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# OUTCOME BASED EDUCATION (OBE) FRAMEWORK

#### **CHAPTER II**



## **MCOPS** Vision Mission

Vision: "Excellence in Pharmaceutical Education and Research"

**Mission:** "Marching with the Times"

## **Quality Policy**

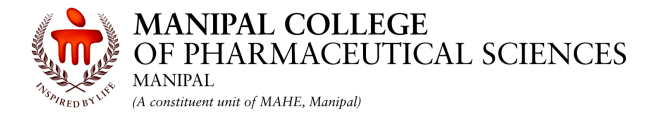
- MCOPS is committed towards providing value-based pharmaceutical education to meet industry, hospital and community needs through continuous improvement of infrastructure and facility for learning, practice and research.
- MCOPS shall provide an environment conducive to the development of staff and students.



### MPharm Pharmaceutical Analysis Program Educational Objectives

The Department of **Pharmaceutical Chemistry**, Manipal College of Pharmaceutical Sciences Manipal, strives to nurture an attitude conducive to self-learning and lifelong learning that would:

PEO	Education Objective
No	
PEO 1	Build an education leading to a Masters' degree in
	Pharmaceutical Analysis with integrated professional
	knowledge and skills with research competencies in the
	analysis and quality control of food, drugs, cosmetics and
	herbal products.
PEO 2	Provide the Masters' students with comprehensive knowledge
	and skills to deliver professional services in the field of
	Pharmaceutical Analysis and enable them to adapt according
	to evolving paradigms in Academia, Pharmaceutical industry
	and Research.
PEO 3	Cultivate an inclination for higher learning and
	entrepreneurship
PEO 4	Foster the best in-class experimental hands-on training in the
	analysis and quality control of drugs, food products, herbals
	and cosmetics by using sophisticated analytical tools and
	techniques.
DEO 5	
PEO 5	Empower and sensitize the pharmaceutical analyst to serve the
	Academia, Pharmaceutical Industry and society with honesty
	and integrity.



# MPharm Pharmaceutical Analysis Program Outcomes (POs)

# After successful completion of M Pharm Pharmaceutical Analysis program, students will be able to:

PO No	Attribute	Competency								
PO 1	Domain knowledge	Provide comprehensive knowledge in the field of pharmaceutical analysis and to integrate professional and research skills in the analysis and quality control of drugs, food, herbal products and cosmetics.								
PO 2	Problem analysis	Recognise and analyze the problems related to analytical /bioanalytical method development, validation, other routine quality control aspects and data analysis leading to meaningful conclusions.								
PO 3	Design/develop solutions	Develop solutions for problems related to analytical/bioanalytical method development, characterization of novel drug molecules and quality control of drugs, food, herbal products and cosmetics through strategies in Pharmaceutical Analysis and interdisciplinary approaches.								
PO 4	Conduct investigations of complex problems	ConductConceptualize and investigate the complex problems related toinvestigations of complexbioanalytical method development, bioavailability and bioequivalence studies quality of drugs, food, herbal products								
PO 5	Modern tool usage	Develop, validate and apply appropriate analytical and bioanalytical methods in the field of drug development and quality control using modern analytical instruments.								
PO 6	Business and society	Develop and facilitate rapid, cost effective and validated analytical/bioanalytical methods, quality control test methods for the benefit of the business and society.								
PO 7	Environment and sustainability	Understand and provide solutions to reduce the hazards from Pharmaceutical Industry through usage of environmental friendly approaches and demonstrate its sustainability.								
PO 8	Ethics	Inculcate and apply ethical principles while discharging professional responsibilities								

PO 9	Individual / Teamwork	Function effectively as an individual and as a member, demonstrate leadership qualities as a leader in diverse teams, and in multidisciplinary settings for team building capacities.
PO No	Attribute	Competency
PO 10	Communication	Possess soft skills and communicate effectively ideas, present the scientific reports in a comprehensive and focused manner to the scientific community, regulatory agencies and society at large.
PO 11	Project management and finance	Demonstrate the knowledge of financial management to evaluate existing and new projects related to method development and quality control for effective decision making.
PO 12	Life-long learning	Comprehend the need to engage oneself as a life-long learner.

# CHAPTER – III

- Course Work
- > COs POs Mapping
- Course Outcomes
- Course Content and Assessment Plan
- > Syllabus in detail

Course wo	rk of MPharm – Pharmace	utical Aı	nalysis (N	(IPA) spec	ializatio	n
Course	Course Title	Cr	edit hours/	week	Credit	Marks
Code		Lecture (L)	Tutorial (T)	Practical (P)	Points	
Semester I						
PQA-MPA101T	Modern Pharmaceutical Analytical Techniques	4			4	100
PCH-MPA102T	Advanced Pharmaceutical Analysis	4	1		5	100
PCH-MPA103T	Pharmaceutical Validation	4	1		5	100
PCH-MPA104T	Food Analysis	4	1		5	100
PCH-MPA105P	Pharmaceutical Analysis Practical I			12	6	150
PCH-MPA106S	Seminar*			2	1	100
	Total	16	3	14	26	650
Semester II					•	
PCH-MPA201T	Advanced Instrumental Analysis	4	1		5	100
PCH-MPA202T	Modern Bioanalytical Techniques	4	1		5	100
PCH-MPA203T	Quality Control and Quality Assurance	4	1		5	100
PCH-MPA204T	Herbal and Cosmetic Analysis	4	1		5	100
PCH-MPA205P	Pharmaceutical Analysis Practical II			12	6	150
PCH-MPA206S	Seminar*			2	1	100
	Total	16	4	14	27	650
* No end-semester	examination. Only continuous	mode.				

	Course work for MPharm III and IV semesters (Common for all specializations)											
Course	Course         Course Title         Credit hours/week         Credit         Marks											
Code		Lecture	Tutorial	Practical	Points							
		(L)	<b>(T)</b>	<b>(P</b> )								
PHA-MRM301T	Research Methodology and	4			4	100						
	Biostatistics*											
MJC302P	Journal Club*			2	1	100						
MRW401P	Research Work			70	35	600						
	Total	4		72	40	800						
* No end-semester	examination.											

S No	Course Code	Course Name	Credits	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11	PO12
1	PQA-MPA101T	Modern Pharmaceutical Analytical Techniques	4	CO1 CO2 CO3	CO1 CO2 CO3	CO1 CO2 CO3	CO1	CO1 CO2 CO3	CO1	CO1 CO2 CO3	CO1 CO2 CO3	CO3	CO1 CO3	CO1 CO2 CO3	
2	PCH-MPA102T	Advanced Pharmaceutical Analysis	5	CO1 CO2 CO3	CO1 CO2 CO3	CO1 CO2 CO3	CO1 CO2 CO3								
3	PCH-MPA103T	Pharmaceutical Validation	5		CO1 CO2 CO3 CO4			CO1 CO2 CO3 CO4	CO1 CO2 CO3 CO4	CO2 CO4					
4	PCH-MPA104T	Food Analysis	5	CO1 CO2 CO3 CO4 CO5	CO5	CO1 CO2 CO3 CO4 CO5	CO1 CO2 CO3 CO4	CO1 CO2 CO3 CO4	CO1 CO2 CO3 CO4	CO5					
5	PCH-MPA105P	Pharmaceutical Analysis Practical I	6	CO1 CO3 CO4	CO1 CO2 CO3 CO4	CO2 CO3 CO4	CO2	CO2 CO3 CO4	CO2			CO3 CO4			
6	PCH-MPA106S	Seminar*	1	CO1	CO1 CO2		CO2 CO5					CO3	CO3 CO4 CO5		CO6
7	PCH-MPA201T	Advanced Instrumental Analysis	5	CO1 CO2 CO3	CO1 CO2	CO1 CO2	CO1 CO2	CO1 CO2			CO1	CO1 CO3			
8	PCH-MPA202T	Modern Bioanalytical Techniques	5		CO1 CO2 CO3	CO1 CO2 CO3	CO1 CO2 CO3	CO1 CO2 CO3	CO1 CO2 CO3				CO3		
9	PCH-MPA203T	Quality Control and Quality Assurance	5	CO1 CO2 CO3 CO4	CO1	CO1 CO3	CO4				CO1				

# PROGRAM OUTCOMES (POs) AND COURSE OUTCOMES (COs) MAPPING

S No	Course Code	Course Name	Credits	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO1O	PO11	PO12
10	PCH-MPA204T	Herbal and Cosmetic Analysis	5	CO1 CO2 CO3 CO4	CO1 CO2 CO3 CO4	CO1 CO3	CO1 CO3	CO1 CO4	CO1 CO2	CO1	CO4	CO1 CO2 CO4		CO1 CO2	
11	PCH-MPA205P	Pharmaceutical Analysis Practical II	6	CO1 CO3 CO4	CO1 CO2 CO3 CO4	CO2 CO3 CO4	CO2	CO2 CO3 CO4	CO2			CO3 CO4			
12	PCH-MPA206S	Seminar*	1	CO1	CO1 CO2		CO2 CO5					CO3	CO3 CO4 CO5		CO6
13	PHA-MRM301T	Research Methodology and Biostatistics*	4	CO1		CO1	CO2	CO2						CO1	
14	MJC302P	Journal Club*	1	CO1	CO1		CO1					CO2 CO3	CO3		CO4
15	MRW401P	Research Work	35	CO1	CO1	CO4	CO5	CO5	CO6	CO3	CO3		CO5 CO6	CO2	

# **CHAPTER III: SYLLABUS**

# MPHARM – PHARMACEUTICAL ANALYSIS (MPA)

# SEMESTER I

# PQA-MPA101T: MODERN PHARMACEUTICAL ANALYTICAL TECHNIQUES

MODERN PHARMACEUTICAL ANALYTICAL TECHNIQUES (Theory)           SCOPE/ SUMMARY         OBJECTIVES/COURSE OUTCOMES           SCOPE/ SUMMARY         OBJECTIVES/COURSE OUTCOMES           This course deals with various advanced analytical instrumentatil techniques for identification, characterization and quantification of drugs. Instruments dealt are NMR, Mass spectrometer, IR, HPLC, GC etc.         After completion of the course, a student will be able to understand:           NMR, Mass spectrometer, IR, HPLC, GC etc.         The theory, instrumentation & applications of Mass spectroscopy.         The theory, instrumentation & applications of of chromatographic technique.         Marks of assessment Plan           Si No.         Distribution of marks of assessment Plan           Si Si Si No.         Course Content         Syllabus of various spectroscopy.         Distribution of marks of assessment/ Si Si S2           1         Will know about theory, instrumentation and application of various spectroscopy.         Unit I (15 hrs)         30         10         20           2         Will know about the theory, instrumentation and applications of various spectroscopy.         Unit II (6 hrs)         13         3         10           3         instrumentation and applications of various spectroscopy.         Unit IV (8 hrs)         19         4         15	COU	RSE CODE	PQA-MPA10	1T								
This course deals with various advanced analytical instrumental techniques for identification, characterization and quantification of drugs. Instruments dealt are NMR, Mass spectrometer, IR, HPLC, GC etc.       After completion of the course, a student will be able to understand:         1       In theory, instrumentation & applications of of chromatographic technique.       The theory, instrumentation & applications of of chromatographic technique.         5       Si No.       Course Content       Syllabus (Chapters) assessment plan         1       Will know about the theory, instrumentation and applications of various spectroscopy.       Unit I (15 hrs)       30       10       20         2       instrumentation and applications of NMR spectroscopy.       Unit II (8 hrs)       11       13       3       10         3       Will know about the theory, instrumentation and applications of Mass spectroscopy.       Unit II (8 hrs)       13       10       20         3       Will know about the theory, instrumentation and applications of Mass spectroscopy.       11       13       3       10         4       Will know about the theory, instrumentation and applications of Mass spectroscopy.       28       28       20         5       Will know about the theory, instrumentation and applications of Mass spectroscopy.       28       28       20         4       Will know about the theory, instrumentation and applications of Mass spectroscopy.	COU	RSE TITLE		IARMAC	EUTICAL	ANALYT	ICAL TI	ECHNIQ	QUES			
advanced analytical instrumental techniques for identification, characterization and quantification of drugs. Instruments dealt are NMR, Mass spectrometer, IR, HPLC, GC etc. SI No. SI SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI No. SI SI SI SI SI SI SI SI SI SI		SCOPE/ S	SUMMARY		OBJE	CTIVES/COURSE OUTCOMES						
SI No.Course ContentSyllabus (Chapters or Units with hours)Marks of assessmentDistribution of marks of assessmentEnd Sem exam (30% of marks) marks of assessment)1Will know about theory, instrumentation and application of various spectroscopic techniques.Unit I (15 hrs)3010202Will know about the theory, instrumentation and applications of NMR spectroscopy.Unit II (8 hrs)3010203Will know about the theory, instrumentation and applications of Mass spectroscopy.Unit III (6 hrs)133104Will know about the theory, instrumentation and applications of various chromatographic techniques.Unit IV (8 hrs)194155Will know about the theory, instrumentation and applications of various chromatographic techniques.Unit IV (8 hrs)194155Will know about the theory, applications and applications of electrophoresis, X-ray crystallography, Potentiometry, thermal techniques and Immuno assays.28820	advand technic charac of drug NMR,	ced analytical in ques for identif terization and c gs. Instruments Mass spectrom	nstrumental ication, quantification dealt are neter, IR,	<ol> <li>understar</li> <li>The trivisible</li> <li>The trivisible</li></ol>	derstand: The theory, instrumentation & applications of U visible spectroscopy, IR, Fluorimetry & AES. The theory, instrumentation & applications of NM spectroscopy. The theory, instrumentation & applications of Ma spectrometry. The theory, instrumentation & applications of chromatographic technique. The theory, instrumentation & applications electrophoresis, XRD, polarimetry, thermal							
SI No.Course ContentSylabus (Chapters or Units with hours)Marks of assessment $\frac{30\% of marks}{of assessment}$ $End Semexam(70\% of marks)marks ofassessment)1Will know about theory,instrumentation and application ofvarious spectroscopic techniques.Unit I(15 hrs)3010202Will know about the theory,instrumentation and applications ofNMR spectroscopy.Unit II(8 hrs)315103Will know about the theory,instrumentation and applications ofNMR spectroscopy.Unit III(6 hrs)133104Will know about the theory,instrumentation and applications ofvarious chromatographic techniques.Unit IV(8 hrs)194155Will know about the theory,instrumentation and applications ofvarious chromatographic techniques.Unit IV(8 hrs)194155Will know about the theory,instrumentation and applications ofvarious chromatographic techniques.Unit IV(15 hrs)288205Will know about the theory,applications and applications ofelectrophoresis, X-raycrystallography, Potentiometry,thermal techniques and Immunoassays.2128420$			Course		<del>_</del>	-						
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2instrumentation and applications of NMR spectroscopy.Unit II (8 hrs)155103Will know about the theory, instrumentation and applications of Mass spectroscopy.Unit III (6 hrs)133104Will know about the theory, instrumentation and applications of various chromatographic techniques.Unit IV (8 hrs)194155Will know about the theory, instrumentation and applications of various chromatographic techniques.Unit IV (8 hrs)194155Will know about the theory, applications and applications of electrophoresis, X-ray crystallography, Potentiometry, thermal techniques and Immuno assays.Unit V (15 hrs)28820	1	instrumentatio	on and applic	ation of		30			20			
3instrumentation and applications of Mass spectroscopy.Unit III (6 hrs)133104Will know about the theory, instrumentation and applications of various chromatographic techniques.Unit IV (8 hrs)194155Will know about the theory, applications and applications of electrophoresis, thermal techniques and Immuno assays.Unit IV (15 hrs)19415	2	instrumentatio	on and applica			15	5		10			
4instrumentation and applications of various chromatographic techniques.Unit IV (8 hrs)194155Will know about the theory, applications and applications of electrophoresis, X-ray crystallography, Potentiometry, thermal techniques and Immuno assays.19415	3	instrumentatio	on and applica	•		13		3	10			
5applications and applications of electrophoresis, crystallography, thermal techniques and Immuno assays.28285Unit V (15 hrs)286820	4	instrumentatio	on and applica	ations of		19		4	15			
	5	applications electrophoresi crystallograph thermal tech	and applicat s, y, Poten	ions of X-ray tiometry,		28		8	20			
		ž	Total M	Aarks of A	ssessment	105	15	15	75			

# PQA-MPA101T: MODERN PHARMACEUTICAL ANALYTICAL TECHNIQUES THEORY 52 hrs

1.a. UV-Visible Spectroscopy: Introduction, theory, laws, instrumentation associated with UV-Visible spectroscopy, choice of solvents and solvent effect and applications of UV-Visible spectroscopy, Difference/ Derivative spectroscopy.

b. IR Spectroscopy: Theory, modes of molecular vibrations, sample handling, instrumentation of dispersive and Fourier-Transform IR spectrometer, factors affecting vibrational frequencies and applications of IR spectroscopy. 5 hrs

c. Spectroflourimetry: Theory of fluorescence, factors affecting fluorescence, quenchers, instrumentation and applications of fluorescence spectrophotometry.
 2 hrs

d. Flame Emission Spectroscopy and atomic absorption spectroscopy: Principle, instrumentation, interferences and applications.3 hrs

2. NMR Spectroscopy: Quantum numbers and their role in NMR, principle, instrumentation, solvent requirement in NMR, relaxation process, NMR signals in various compounds, chemical shift, factors influencing chemical shift, Spin-Spin coupling, coupling constant, nuclear magnetic double resonance, brief outline of principles of FT-NMR and 13C NMR. Applications of NMR spectroscopy. 8 hrs

3. **Mass Spectroscopy**: Principle, theory, instrumentation of mass spectroscopy, different types of ionization like electron impact, chemical, field, FAB and MALDI, APCI, ESI, APPI analyzers of quadrupole and time of flight, mass fragmentation and its rules, meta stable ions, isotopic peaks and applications of mass spectroscopy. **6 hrs** 

4. **Chromatography**: Principle, apparatus/instrumentation, chromatographic parameters, factors affecting resolution and applications of the following:

a) Planar chromatography-Paper, Thin layer chromatography and High performance thin layer chromatography b) Ion exchange chromatography c) Column chromatography d) Gas chromatography e) High performance liquid chromatography and ultra high performance liquid chromatography f) Affinity chromatography g) Gel chromatography. **8 hrs** 

#### 5. Other Analytical Techniques

a. Electrophoresis: Principle, instrumentation, working conditions, factors affecting separation and applications of the following: a) Paper electrophoresis b) Gel electrophoresis
c) Capillary electrophoresis d) Zone electrophoresis e) Moving boundary electrophoresis f) Iso electric focusing.
3 hrs

b. X-ray Crystallography: Different X-ray diffraction methods, Bragg's law	, Rotating
crystal technique, X-ray powder technique, types of crystals and applications	of X-ray
diffraction.	2 hrs
c. Potentiometry: Principle and application of potentiometry.	2 hrs
d. Thermal Techniques: Principle and application of Differential Scanning Ca	alorimetry,
Differential Thermal Analysis and Thermo Gravimetric Analysis.	5 hrs
e. Immunological Assays: RIA (Radio immuno assay), ELISA.	3 hrs

- Spectrometric Identification of Organic Compounds, Robert M Silverstein, 6<sup>th</sup> edition, John Wiley & Sons, 2004.
- Principles of Instrumental Analysis, Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5<sup>th</sup> edition, Eastern Press, Bangalore, 1998.
- 3. Instrumental Methods of Analysis, Willards, 7<sup>th</sup> edition, CBS Publishers.
- Practical Pharmaceutical Chemistry, Beckett and Stenlake, Vol II, 4<sup>th</sup> edition, CBS Publishers, New Delhi, 1997.
- 5. Organic Spectroscopy, William Kemp, 3<sup>rd</sup> edition, ELBS, 1991.
- Quantitative Analysis of Drugs in Pharmaceutical Formulation, PD Sethi, 3<sup>rd</sup> edition, CBS Publishers, New Delhi, 1997.
- 7. Pharmaceutical Analysis-Modern Methods, Part B, J W Munson, Volume11, Marcel Dekker Series.
- 8. Introduction to Spectroscopy, Donald L Pavia. 8th edition, CENGAGE Learning, USA
- 9. Contemporary Practice of Chromatography, Poole, Colin F and Sheila A Schuette, Elsevier Science Publishers.
- 10. Remington: The Science and Practice of Pharmacy, latest edition, Pharmaceutical Press, UK.
- 11. Quantitative Analysis of Pharmaceutical Formulations by HPTLC, PD Sethi, CBS Publishers, New Delhi.

#### **SEMESTER I**

# PCH-MPA102T: ADVANCED PHARMACEUTICAL ANALYSIS

COL	JRSE CODE	PCH-MPA102T								
COL	J <b>RSE TITLE</b>	ADVANCED PH	IARMACEU	TICAL ANAL	YSIS	(Theory)				
	SCOPE/ SU	MMARY	<b>OBJECTIVES/COURSE OUTCOMES</b>							
aspec drug Elem profi degra phyto proto biolo and t	cts of Impurity, I products, in re- nental impuri ling and cha adants, Stabili opharmaceuticals ocol preparation.	racterization of ty testing of s and their It also covers the various vaccines d procedure.	able to under 1. Appropria analytical 2. Principles group ana research applicatio 3. Analysis of stability stability stability stability pontent and Assess	Upon completion of this course the student shou able to understand: 1. Appropriate analytical skills required to analytical method development. 2. Principles of various reagents used in furgroup analysis that renders necessary supresearch methodology and demonstrate application in the practical related problems. 3. Analysis of impurities in drugs, residual solver stability studies of drugs and biological produ- intent and Assessment Plan Syllabus Distribution of material Syllabus Distribution Syllabus Distributi						
Sl No.	( 'ourse ( 'ontent		(Chapters or Units with hours)	Marks of assessment	exam (30% of marks of assessment) S1 S2		exam (70% of marks of assessment)			
1	evaluation incl and quantitativ	e insights on total uding qualitative ve, of impurities ources as per ICH	Unit I (10 hrs)	20	5		15			
2	sources of electricity quantification instrumentation	*	Unit II (8 hrs)	15	5		10			
3	Learn about Impurity profiling, degradants, characterization Accelerated stability testing & 3 shelf life calculation, Learning WHO and ICH stability testing guidelines and guidelines for biological products		Unit III (10 hrs)	20	5		15			
4		conduct Stability opharmaceuticals	Unit IV (7 hrs)	15		5	10			
5		iological assay some biological	Unit V (8 hrs)	15		5	10			

6	Learn the Immunoassays (IA) with Basic principles, procedures , Production of antibodies, Learn the various immunoassay techniques	Unit VI (9 hrs)	20		5	15
	Total Marks of	Assessment	105	15	15	75

#### PCH-MPA102T: ADVANCED PHARMACEUTICAL ANALYSIS

#### THEORY

1. Impurity and stability studies:

Definition, classification of impurities in drug Substance or Active Pharmaceutical Ingredients and quantification of impurities as per ICH guidelines Impurities in new drug products:

Rationale for the reporting and control of degradation products, reporting degradation products content of batches, listing of degradation products in specifications, qualification of degradation products

Residual solvents: General principles, classification of residual solvents, Analytical procedures, limits of residual solvents, reporting levels of residual solvents

2 Elemental impurities:

Element classification, control of elemental impurities, potential sources of elemental impurities, identification of Potential Elemental Impurities, analytical procedures, instrumentation.

Stability testing protocols:

Selection of batches, container orientation, test parameters, sampling frequency, specification, storage conditions, recording of results, concept of stability, commitment etc. Important mechanistic and stability related information provided by results of study of factors like temperature, pH, buffering species, ionic strength and dielectric constant etc. on the reaction rates with practical considerations.

3 Impurity profiling and degradent characterization: 10 hrs

Method development, Stability studies and concepts of validation. Accelerated stability testing & shelf life calculation, WHO and ICH stability testing guidelines, Stability zones, steps in development, practical considerations. Basics of impurity profiling and degradent

10 hrs

52 hrs

#### 8 hrs

characterization with special emphasis. Photostability testing guidelines, ICH stability guidelines for biological products

4 Stability testing of phytopharmaceuticals: 7 hrs
 Regulatory requirements, protocols, HPTLC/HPLC finger printing, interactions and complexity.

- 5 Biological tests and assays of the following: 8 hrs
  a. Adsorbed Tetanus vaccine b. Adsorbed Diphtheria vaccine c. Human anti haemophilic vaccine d. Rabies vaccine e. Tetanus Anti toxin f. Tetanus Anti serum g. Oxytocin h. Heparin sodium IP i. Antivenom. PCR, PCR studies for gene regulation, instrumentation (Principle and Procedures)
- 6 Immunoassays (IA)
   9 hrs
   Basic principles, Production of antibodies, Separation of bound and unbound drug, Radioimmunoassay, Optical IA, Enzyme IA, Fluoro IA, Luminiscence IA, Quantification and applications of IA.

- Vogel's textbook of quantitative chemical analysis Jeffery J Bassett, J. Mendham, R. C. Denney, 5<sup>th</sup> edition, ELBS, 1991.
- Practical Pharmaceutical Chemistry Beckett and Stenlake, Vol II, 4th Edition, CBS publishers, New Delhi, 1997.
- Textbook of Pharmaceutical Analysis K A Connors, 3rd Edition, John Wiley & Sons, 1982.
- 4. Pharmaceutical Analysis Higuchi, Brochmman and Hassen, 2nd Edition, Wiley Inter science Publication, 1961.
- Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi, 3rd Edition, CBS Publishers New Delhi, 1997.
- Pharmaceutical Analysis- Modern methods J W Munson Part B, Volume 11, Marcel Dekker Series.
- The Quantitative analysis of Drugs D C Carratt, 3rd edition, CBS Publishers, NewDelhi, 1964.
- 8. Indian Pharmacopoeia Vol I, II & III 2007, 2010, 2014.
- 9. Methods of sampling and microbiological examination of water, first revision, BIS

- Practical HPLC method development Snyder, Kirkland, Glajch, 2nd edition, John Wiley & Sons.
- 11. Analytical Profiles of drug substances Klaus Florey, Volume 1 20, Elsevier, 2005
- 12. Analytical Profiles of drug substances and Excipients Harry G Brittan, Volume 21 30, Elsevier, 2005.
- The analysis of drugs in biological fluids Joseph Chamberlain, 2nd edition, CRC press, London.
- 14. ICH Guidelines for impurity profiles and stability studies.

#### SEMESTER I

# PCH-MPA103T: PHARMACEUTICAL VALIDATION

COU	URSE CODE	PCH-MPA10	3T								
COU	<b>IRSE TITLE</b>	PHARMACE	UTICAL VA	LIDATION (T	heory)						
	SCOPE/ SUM	MARY	<b>OBJECTIVES/COURSE OUTCOMES</b>								
valid to ind quali cover about	course is to und ation and how it c dustry and thus to ty of the products rs the complete t the validat odology and appl	can be applied o improve the s. The subject information tion, types, ications	able to: 1. Understa and Valid 2. Perform of 3. Perform of manufact 4. Understa of IPR in	<ol> <li>Understand and explain the concepts of qualification</li> <li>Perform qualification of Instruments</li> </ol>							
SI No.	Course C	ontent	Syllabus (Chapters or Units with hours)	Marks of assessment			sment End Sem exam (70% of marks of assessment)				
1	Able to explain validation and of manufacturin Analytical and equipments	Qualification g equipments,	Unit I (10 hrs)	20	5		15				
2	Learn to Qua Analytical Inst glasswares	lification of ruments and	Unit II (12 hrs)	25		7	18				
3	Learn to vali systems includi validation	idate Utility ing Cleaning	Unit III (10 hrs)	20	2	2	16				
4	Learn to valida methods and systems.	-	Unit IV (10 hrs)	20	8		12				
5	Learn the gene of Intellectual pr		Unit V (10 hrs)	20		6	14				
		Total Marks of	Assessment	105	15	15	75				

#### PCH-MPA103T: PHARMACEUTICAL VALIDATION

#### THEORY

52 hrs

1. Introduction: Definition of Qualification and Validation

Advantage of Validation, Streamlining of Qualification & Validation process and Validation Master Plan.

Qualification: User Requirement Specification, Design Qualification, Factory AcceptanceTest (FAT)/ Site Acceptance Test (SAT), Installation Qualification, OperationalQualification, Performance Qualification, Re- Qualification (Maintaining status- CalibrationPreventive Maintenance, Change management), Qualification of Manufacturing Equipments,Qualification of Analytical Instruments and Laboratory equipments.10 hrs

- 2 Qualification of analytical instruments: Electronic balance, pH meter, UV-Visible spectrophotometer, FTIR, GC, HPLC, HPTLC Qualification of Glassware: Volumetric flask, pipette, Measuring cylinder, beakers and burette.
   12 hrs
- 3 Validation of Utility systems: Pharmaceutical Water System & pure steam, HVAC system, Compressed air and nitrogen.

Cleaning Validation: Cleaning Validation - Cleaning Method development, Validation and validation of analytical method used in cleaning. Cleaning of Equipment, Cleaning of Facilities. Cleaning in place (CIP). 10 hrs

4 Analytical method validation: General principles, Validation of analytical method as per ICH guidelines and USP.

Computerized system validation: Electronic records and digital significance-21 CFR part 11 and GAMP. 10 hrs

5 General Principles of Intellectual Property: Concepts of Intellectual Property (IP), Intellectual Property Protection (IPP), Intellectual Property Rights (IPR); Economic importance, mechanism for protection of Intellectual Property–patents, Copyright, Trademark; Factors affecting choice of IP protection; Penalties for violation; Role of IP in pharmaceutical industry; Global ramification and financial implications. Filing a patent applications; patent application forms and guidelines. Types patent applications-provisional and non-provisional, PCT and convention patent applications; International patenting requirement procedures and costs; Rights and responsibilities of a patentee; Practical aspects regarding maintaining of a Patent file; Patent infringement meaning and scope. Significance of transfer technology (TOT), IP and ethics-positive and negative aspects of IPP; Societal responsibility, avoiding unethical practices. **10 hrs** 

- 1. B. T. Loftus & R. A. Nash, "Pharmaceutical Process Validation", Drugs and Pharm Sci. Series, Vol. 129, 3rd Ed., Marcel Dekker Inc., N.Y.
- 2. The Theory & Practice of Industrial Pharmacy, 3rd edition, Leon Lachman, Herbert A. Lieberman, Joseph. L. Karig, Varghese Publishing House, Bombay.
- 3. Validation Master Plan by Terveeks or Deeks, Davis Harwood International publishing.
- 4. Validation of Aseptic Pharmaceutical Processes, 2nd Edition, by Carleton & Agalloco, (Marcel Dekker).
- 5. Michael Levin, Pharmaceutical Process Scale-Upl, Drugs and Pharm. Sci. Series, Vol. 157,2nd Ed., Marcel Dekker Inc., N.Y.
- 6. Validation Standard Operating Procedures: A Step by Step Guide for Achieving Compliance in the Pharmaceutical, Medical Device, and Biotech Industries, Syed Imtiaz Haider
- 7. Pharmaceutical Equipment Validation: The Ultimate Qualification Handbook, Phillip A. Cloud, Interpharm Press
- 8. Validation of Pharmaceutical Processes: Sterile Products, Frederick J. Carlton (Ed.) and James Agalloco (Ed.), Marcel Dekker, 2nd Ed.
- 9. Analytical Method validation and Instrument Performance Verification by Churg Chan, Heiman Lam, Y.C. Lee, Yue. Zhang, Wiley Inter Science.

#### SEMESTER I

# PCH-MPA104T: FOOD ANALYSIS

COU	RSE CODE	PCH-MPA104T								
COU	IRSE TITLE	FOOD ANALYSIS	(Theory)							
	SCOPE/ SU	J <b>MMARY</b>	<b>OBJECTIVES/COURSE OUTCOMES</b>							
This course is to impart knowledge on analysis of food constituents and finished food products. This course also includes application of instrumental analysis in determination of pesticides in variety of food products Course Cor			<ul> <li>Upon completion of this course the student should be able to understand various analytical techniques in the determination of:</li> <li>1. Food Constituents</li> <li>2. Food Additives</li> <li>3. Finished Food Products</li> <li>4. Pesticides in Food</li> <li>5. Also student shall have the knowledge on food regulations and legislations</li> <li>ntent and Assessment Plan</li> </ul>							
		Course Cor	nont and 7355			bution of	marks of			
SI No.	Cour	Syllabus (Chapters or Units with hours)	Marks of assessment	Sessiona (30% of n assessn S1	narks of nent) S2	ent End Sem exam (70% of marks of assessment)				
1	Able to under techniques in carbohydrates, fiber crude fibe	proteins, dietary	Unit I (10 hrs)	20	5		15			
2		stand the analytical determination of mins	Unit II (10 hrs)	20	5		15			
3		stand the analytical etermination of Food	Unit III (12 hrs)	25	3	6	16			
4		stand the analytical etermination of Milk ituents	Unit IV (10 hrs)	20	2	5	13			
5	techniques in	stand the analytical determination of lso on regulations on	Unit V (10 hrs)	20		4	16			
		Total Marks of	Assessment	105	15	15	75			

#### PCH-MPA104T: FOOD ANALYSIS

#### THEORY

52 hrs

1. Carbohydrates: Classification and properties of food carbohydrates. General methods of analysis of food carbohydrates. Changes in food carbohydrates during processing. Dietary fibre, Crude fibre.

Proteins: Chemistry and classification of amino acids and proteins, Physico-Chemical properties of protein and their structure, general methods of analysis of proteins and amino acids. 10 hrs

2. Lipids: Classification, general methods of analysis, refining of fats and oils; hydrogenation of vegetable oils, Determination of adulteration in fats and oils, Various methods used for measurement of spoilage of fats and fatty foods.

Vitamins: classification of vitamins, methods of analysis of vitamins, Principles of microbial assay of vitamins of B-series. 10 hrs

3. Food additives: Introduction, analysis of Preservatives, antioxidants, artificial sweeteners, flavors, flavor enhancers, stabilizers, thickening and jelling agents.

Pigments and synthetic dyes: Natural pigments, their occurrence and characteristic properties, permitted synthetic dyes, Non-permitted synthetic dyes used by industries, Method of detection of natural, permitted and non-permitted dyes. 12 hrs

4. General Analytical methods for milk, milk constituents and milk products like ice cream, milk powder, butter, margarine, cheese including adulterants and contaminants of milk. Analysis of fermentation products like wine, spirits, beer and vinegar.
10 hrs

5. Pesticide analysis: Effects of pest and insects on various food, use of pesticides in agriculture, pesticide cycle, organophosphorus and organochlorine pesticides analysis, determination of pesticide residues in grain, fruits, vegetables, milk and milk products. Legislation regulations of food products with special emphasis on BIS, Agmark, FDA and US-FDA.
10 hrs

- The chemical analysis of foods David Pearson, Seventh edition, Churchill Livingstone, Edinburgh London, 1976
- Introduction to the Chemical analysis of foods S. Nielsen, Jones & Bartlett publishers, Boston London, 1994.
- 3. Official methods of analysis of AOAC International, sixth edition, Volume I & II, 1997.
- 4. Analysis of Food constituents Multon, Wiley VCH.
- 5. Dr. William Horwitz, Official methods of analysis of AOAC International, 18th edition, 2005.

# SEMESTER I

# PCH-MPA105P: PHARMACEUTICAL ANALYSIS PRACTICAL I

COU	URSE CODE	PCH-MPA 105P						
COU	<b>IRSE TITLE</b>	PHARMACEUTICAI	L ANALYSIS F	PRACTICA	L-I			
This	<b>SCOPE/SU</b> s course is d		OBJECTIVES/COURSE OUTCOMES On completion of this course student shall be able to					
<ul> <li>practical knowledge on analysis of drugs and drug intermediates by titrimetric methods and using sophisticated instrumental methods.</li> <li>1. Perform calibration of volumetric analytical instruments.</li> <li>2. Perform qualitative and quantitati Pharmacopoeial compounds, natu products using instrumental techni</li> </ul>					volumetric app d quantitative a bounds, natural	aratus and malysis of and food		
		Course Co	ntent and Asses	sment Plan				
					Distributi assessment			
Sl No.	Cou	rse Content	Syllabus (Chapters or Units with hours)	Total Marks of assessment	Sessional exam (25 % of total marks of			
					S1	assessment)		
1	Learn to calibrate apparatus and analytical instruments, quantitative analysis of Pharmacopoeial compounds/ formulations.		1, 2 and 13	40	08	32		
2	analysis of quantitative functional grou and proteins, saponification Peroxide value and rancidity, p residue, vitami additives in foo	value, Iodine value, a, acid value, fat content preservatives, pesticide n contents and food d products, & analysis ynthetic colors, density	Experiments 5 to 11 and 20 to 29 (84 hrs)	80	18	62		
3	Learn the analyst HPLC and ( impurity profili	sis of compounds using Gas Chromatography, ng of drugs & Cleaning y one equipment	3,4 12 and	10	04	06		
		Total Marks	of Assessment	130	30	100		

## PCH-MPA105P: PHARMACEUTICAL ANALYSIS PRACTICAL I

1. Analysis of Pharmacopoeial compounds and their formulations by UV Vis spectrophotometer

- 2. Simultaneous estimation of multi component containing formulations by UV
- 3. spectrophotometry
- 4. Experiments based on HPLC
- 5. Experiments based on Gas Chromatography
- 6. Estimation of riboflavin/quinine sulphate by fluorimetry
- 7. Estimation of sodium/potassium by flame photometry
- 8. Assay of official compounds by different titrations
- 9. Assay of official compounds by instrumental techniques.
- 10. Quantitative determination of hydroxyl group.
- 11. Quantitative determination of amino group
- 12. Colorimetric determination of drugs by using different reagents
- 13. Impurity profiling of drugs
- 14. Calibration of glasswares
- 15. Calibration of pH meter
- 16. Calibration of UV-Visible spectrophotometer
- 17. Calibration of FTIR spectrophotometer
- 18. Calibration of GC instrument
- 19. Calibration of HPLC instrument
- 20. Cleaning validation of any one equipment
- 21. Determination of total reducing sugar
- 22. Determination of proteins

23. Determination of saponification value, Iodine value, Peroxide value, Acid value in

food products

- 24. Determination of fat content and rancidity in food products
- 25. Analysis of natural and synthetic colors in food
- 26. Determination of preservatives in food
- 27. Determination of pesticide residue in food products
- 28. Analysis of vitamin content in food products
- 29. Determination of density and specific gravity of foods
- 30. Determination of food additives

#### **SEMESTER I**

# PCH- MPA 106S: SEMINAR IN PHARMACEUTICAL ANALYSIS

CO	COURSE CODE PCH- MPA 106S							
СО	COURSE TITLE SEMINAR IN PHARMACEUTICAL ANALYSIS							
	SCOPE/ SUMMARY	OBJECTIVE	S/COURSE OU	JTCOMES				
The subject is designed to create an environment where teachers provide the students a critical eye and openness to fortify the presentation and academic writing skills of students in the field of Pharmaceutical AnalysisUpon completion of the course t be able to:1. Develop skills to gather, or information, and defend a Pharmaceutical Analysis1. Develop skills to gather, or information, and defend a Pharmaceutical Analysis2. Learn to organize analytical audio-visual aids.3. Acquire communication and presentation skills.4. Effectively uestions raised by peers and scrutiny.5. Develop scientific writing skill 6. Cultivate a sense of upgradatio through self and continuous lear			ganize, deliver given topic in concepts using to the tand scientific					
	Course Conter	nt and Assessment Pl	an					
SI No.	Course Content	Hours	Total Marks of assessment	Marks End Sem exam				
1	The students should be able to develop skills to gather, organize, communicate the scientific information, and defend a given topic in Pharmaceutical Analysis	gather, organize, the scientific d defend a given topic 2 hours/week 100 semester 2 hours/week 100 Only continuous						

# MPHARM –PHARMACEUTICAL ANALYSIS (MPA) SEMESTER II

# PCH-MPA201T: ADVANCED INSTRUMENTAL ANALYSIS

COU	JRSE CODE	PCH-MPA-20	)1T					
COU	J <b>RSE TITLE</b>	ADVANCED	INSTRUM	ENTAL ANA	ALYSIS	S (Theor	y)	
This	SCOPE/ SUM	MARY to impart		<b>OBJECTIVES/COURSE OUTCOMES</b> Upon completion of this course the student should be				
fundamental knowledge on various analytical instrumental techniques for identification , characterization and quantification of drugs using LC-MS, GC-MS, ATR-IR and Hypenated techniques.			able to: 1. Unders skills identifi 2. Unders Chroma 3. Unders the hyp	tand the the of IR, NM cation of vari tand the theo atography tec tand the theo henated techr	eoretica IR, an ous org retical hniques retical niques.	al and d Mass ganic con and prac s.	Interpretation spectra and	
		Cours	se Content a	nd Assessmer		ribution	of marks of	
SI No.	Course (	Content	Syllabus (Chapters or Units with hours)	Marks of assessment	Sess ex (30 man	assess sional cam % of rks of sment) S2		
1	Will learn the p max for few compound Woodward Fie learn to in chemical struct IR. Learn the techniques of I	classes of based on ser rule. And terpret the ture based on Hyphenated	Unit I (10 hrs)	20	5		15	
2	Will unders spectroscopy w 1D and 2D N and COSY, Inadequte tec interpretation compounds	vith respect to MR, NOESY HETCOR, hniques and	Unit II (14 hrs)	30	10		20	
3	Will understan Mass Spectro	nd about the oscopy with o mass	Unit III (12 hrs)	25		5	20	

	16 hrs

# Page | 58

	fragmentation and its rules, fragmentation of important functional groups, meta stable ions, Mc Lafferty reaarangement, ring rule, isotopic peaks, interpretation of organic compounds.					
4	Will understand and learn aboutdifferentChromatographic techniques in terms of method developmentand troubleshooting etc.Will also learn Principle, procedure and applications of various hyphenated chromatographic techniques:	Unit IV (16 hrs)	30		10	20
	Total Marks of A	Assessment	105	15	15	75

## PCH-MPA201T: ADVANCED INSTRUMENTAL ANALYSIS

#### THEORY

## 1. UV and IR spectroscopy

Wood ward – Fieser rule for 1,3- butadienes, cyclic dienes and  $\alpha$ ,  $\beta$ -carbonyl compounds and interpretation compounds of enones. NIR, ATR-IR, IR Interpretation of organic compounds.

## 2. NMR spectroscopy

1-D and 2-D NMR, NOESY and COSY, HECTOR, INADEQUATE techniques, Interpretation of organic compounds.

## 3. Mass Spectroscopy

Mass fragmentation and its rules, Fragmentation of important functional groups like alcohols, amines, carbonyl groups and alkanes, Meta stable ions, Mc Lafferty rearrangement, Ring rule, Isotopic peaks, Interpretation of organic compounds.

## 4. Chromatography

# 10 hrs

# 14 hrs

12 hrs

# 10 111 5

**52 hrs** 

**HPLC:** Columns, column problems, gradient HPLC, HPLC solvents, trouble shooting, sample preparation, method development, New developments in HPLC-role and principles of ultra, nano liquid chromatography in pharmaceutical analysis. Immobilized polysaccharide CSP's: Advancement in enantiomeric separations, reverse phase Chiral method development and HILIC approaches. HPLC in Chiral analysis of pharmaceuticals. GC Derivitisation techniques.

Principle, Instrumentation and Applications of the following:

a) GC-MS b) GC-AAS c) LC-MS d) LC-FTIR e) LC-NMR f) CE-MS g) High
 Performance Thin Layer chromatography h) Super critical fluid chromatography i)
 Ion Chromatography j) Ion-Exclusion Chromatography k) Flash chromatography.

- Spectrometric Identification of Organic compounds Robert M Silverstein, Sixth edition, John Wiley & Sons, 2004.
- Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5<sup>th</sup> edition, Eastern press, Bangalore, 1998.
- 3. Instrumental methods of analysis Willards, 7<sup>th</sup> edition, CBS publishers.
- 4. Organic Spectroscopy William Kemp, 3<sup>rd</sup> edition, ELBS, 1991.
- 5. Quantitative analysis of Pharmaceutical formulations by HPTLC P D Sethi, CBS Publishers, New Delhi.
- Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi, 3<sup>rd</sup> Edition, CBS Publishers, New Delhi, 1997.
- Pharmaceutical Analysis- Modern methods Part B J W Munson, Volume 11, Marcel Dekker Series.
- 8. Organic Spectroscopy by Donald L. Paviya, 5th Edition.

#### **SEMESTER II**

# PCH-MPA202T: MODERN BIOANALYTICAL TECHNIQUES

COU	<b>URSE CODE</b>	PCH-	MPA202T					
COU	<b>IRSE TITLE</b>	MOD	ERN BIOAN	ALYTICAI	L TECHNIQU	UES (T	heory)	
	SCOPE/ SU	MMA	RY	OBJI	ECTIVES/C	OURSI	E OUTC	COMES
provide detailed knowledge about the importance of analysis of drugs in biological matricesknow 1. Extra samp valid2. Biop phare devel3. Cell meta				action of dru les and pa ation harmaceution nacokinetic opment. culture te	es and toxicol echniques, in l its identifi	bolites bioana deration kinetic	from bic alytical s in in drug p assay o	ological method cluding product f drug
			-		Assessment P	lan		
Sl No.	Cours	se Cont	ent	Syllabus (Chapter s or Units with	Marks of assessment	Distribution of marks of assessmentSessional exam (30% of marksEnd Sem exam		nent End Sem exam (70% of marks
1	Able to underst in extraction metabolites matrices		e techniques drugs and biological	hours) Unit I (10 hrs)	20	<u>S1</u> 3	<u>\$2</u> 2	of assessment) 15
2	Able to understand various biopharmaceutical considerations during product development			Unit II (10 hrs)	20	5		15
3	Able to ur pharmacokinetic considerations development		toxicokinetic	Unit III (12 hrs)	25	7		18
4	Able to under culture techniqu		various cell	Unit IV (10 hrs)	20		8	12
5	Able to un techniques invo	ndersta olved i		Unit V (10 hrs)	20		5	15

identification, Drug product performance, in vivo bioavailability and bioequivalence					
Total Marks of Assessment		105	15	15	75

#### PCH-MPA202T: MODERN BIOANALYTICAL TECHNIQUES

#### THEORY

**52 hrs** 

1. Extraction of drugs and metabolites from biological matrices:

General need, principle and procedure involved in the Bioanalytical methods such as Protein precipitation, Liquid - Liquid extraction and Solid phase extraction and other novel sample preparation approach.

Bioanalytical method validation: USFDA and EMEA guidelines. 10 hrs

#### 2. Biopharmaceutical Consideration:

Introduction, Biopharmaceutical factors affecting drug bioavailability. In Vitro Dissolution and Drug Release Testing, Alternative methods of dissolution testing. Transport models, Biopharmaceutics Classification System. Solubility: Experimental methods. Permeability: In-vitro, in-situ and In-vivo methods. **10 hrs** 

#### 3. Pharmacokinetics and Toxicokinetics:

Basic consideration, Drug interaction (PK-PD interactions), The effect of protein-binding interactions. The effect of tissue-binding interactions, Cytochrome P450-based drug interactions, Drug interactions linked to transporters. Microsomal assays Toxicokinetics-Toxicokinetic evaluation in preclinical studies. Importance and applications of toxicokinetic studies. LC-MS in bioactivity screening and proteomics. **12 hrs** 

#### 4. Cell culture techniques

Basic equipments used in cell culture lab. Cell culture media, various types of cell culture, general procedure for cell cultures; isolation of cells, subculture, cryopreservation, characterization of cells and their applications. Principles and applications of cell viability assays (MTT assays), Principles and applications of flow cytometry. **10 hrs** 

5. Metabolite identification:

In-vitro / in-vivo approaches, protocols and sample preparation. Microsomal approaches (Rat liver microsomes (RLM) and Human liver microsomes (HLM) in Met –ID. Regulatory perspectives.

In-vitro assay of drug metabolites & drug metabolizing enzymes.

Drug Product Performance, In Vivo: Bioavailability and Bioequivalence:

Drug Product Performance, Purpose of Bioavailability Studies, Relative and Absolute Availability. Methods for Assessing Bioavailability, Bioequivalence Studies. Design and Evaluation of Bioequivalence Studies. Study Designs, Crossover Study Designs. Generic Biologics (Biosimilar Drug Products). Clinical Significance of Bioequivalence Studies. **10 hrs** 

- Analysis of drugs in Biological fluids Joseph Chamberlain, 2<sup>nd</sup> Edition. CRC Press, New York. 1995.
- Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5<sup>th</sup> edition, Eastern press, Bangalore, 1998.
- Pharmaceutical Analysis Higuchi, Brochmman and Hassen, 2<sup>nd</sup> Edition, Wiley Interscience Publications, 1961.
- Pharmaceutical Analysis- Modern methods Part B J W Munson, Volume 11, Marcel Dekker Series
- Practical HPLC method Development Snyder, Kirkland, Glaich, 2<sup>nd</sup> Edition, John Wiley & Sons, New Jersey. USA.
- Chromatographic Analysis of Pharmaceuticals John A Adamovics, 2<sup>nd</sup> Edition, Marcel Dekker, New York, USA. 1997.
- Chromatographic methods in clinical chemistry & Toxicology Roger L Bertholf, Ruth E Winecker, John Wiley & Sons, New Jersey, USA. 2007.
- Good Laboratory Practice Regulations, 2<sup>nd</sup> Edition, Sandy Weinberg Vol. 69, Marcel Dekker Series, 1995.
- Good laboratory Practice Regulations Allen F. Hirsch, Volume 38, Marcel Dekker Series, 1989.
- 10. ICH, USFDA & CDSCO Guidelines.

#### **SEMESTER II**

# PCH-MPA203T: QUALITY CONTROL AND QUALITY ASSURANCE

CO	URSE CODE	PCH-MPA203T					
СО	URSE TITLE	QUALITY CONT	ROL AND (	QUALITY A	ASSUF	RANCI	E (Theory)
	SCOPE/ SUN	MMARY	OBJEC	TIVES/CO	URSE	OUT	COMES
fund aspe assu indu like qua	s course is design damental knowledg ects of quality courance aspects o ustries. It covers the cGMP, QC tests lity certification ulatory affairs.	<ul> <li>be able to k</li> <li>1. The control industry</li> <li>2. The imp</li> <li>3. The scopto pharm</li> <li>4. To under</li> </ul>	now GMP aspe oortance of o pe of quality naceutical in	ects in docum y certif ndustri	n pha entatio ïcation es	student shall armaceutical n is applicable s of QA and	
		Course Cor	ntent and Ass		an		
					Distri	assess	of marks of sment
SI N o.	Course	Content	Syllabus (Chapters or Units with hours)	Marks of assessment	-	am % of ks of	End Sem exam (70% of marks of assessment)
1	the concept and even control and quality includes good la which includes p	be acquainted with volution of quality y assurance which aboratory practice protocol for non- control on animal	Unit I (10 hrs)	20	5		15
2	the cGMP guidel schedule M and U between GMP and		Unit II (12 hrs)	20	7		18
3	material, finis	he analysis of raw shed products, als and in process	Unit III (10 hrs)	20	3	3	14
4	documentation in industry which industry	vill learn about n pharmaceutical clues quality audit specifications and protocols and	Unit IV (10 hrs)	20		6	14

5	Will understand about the drug product inspection, expiry date calculation, production record review, aseptic process control and packaging	Unit V (10 hrs)	20		6	14
	Total Marks of Assessment			15	15	75

#### PCH-MPA203T: QUALITY CONTROL AND QUALITY ASSURANCE

#### THEORY

#### 52 hrs

1. Concept and Evolution of Quality Control and Quality Assurance

Good Laboratory Practice, GMP, Overview of ICH Guidelines - QSEM, with special emphasis on Q-series guidelines.

Good Laboratory Practices: Scope of GLP, Definitions, Quality assurance unit, protocol for conduct of non clinical testing, control on animal house, report preparation and documentation. 10 hrs

- cGMP guidelines according to schedule M, USFDA (inclusive of CDER and CBER) Pharmaceutical Inspection Convention (PIC), WHO and EMEA covering: Organization and personnel responsibilities, training, hygiene and personal records, drug industry location, design, construction and plant lay out, maintenance, sanitation, environmental control, utilities and maintenance of sterile areas, control of contamination and Good Warehousing Practice. CPCSEA guidelines. 12 hrs
- 3. Analysis of raw materials, finished products, packaging materials, in process quality control (IPQC), Developing specification (ICH Q6 and Q3)

Purchase specifications and maintenance of stores for raw materials. In process quality control and finished products quality control for following formulation in Pharma industry according to Indian, US and British pharmacopoeias: tablets, capsules, ointments, suppositories, creams, parenterals, ophthalmic and surgical products (How to refer pharmacopoeias), Quality control test for containers, closures and secondary packing materials. **10 hrs** 

4. Documentation in pharmaceutical industry: Three tier documentation, Policy, Procedures and Work instructions, and records (Formats), Basic principles- How to maintain, retention and retrieval etc. Standard operating procedures (How to write), Master Formula Record, Batch

Formula Record, Quality audit plan and reports. Specification and test procedures, Protocols and reports. Distribution records. Electronic data. 10 hrs

5. Manufacturing operations and controls: Sanitation of manufacturing premises, mix-ups and cross contamination, processing of intermediates and bulk products, packaging operations, IPQC, release of finished product, process deviations, charge-in of components, time limitations on production, drug product inspection, expiry date calculation, calculation of yields, production record review, change control, sterile products, aseptic process control, packaging.

- Quality Assurance Guide by organization of Pharmaceutical Procedures of India, 3<sup>rd</sup> revised edition, Volume I & II, Mumbai, 1996.
- Good Laboratory Practice Regulations, 2<sup>nd</sup> Edition, Sandy Weinberg Vol. 69, Marcel Dekker Series, 1995.
- Quality Assurance of Pharmaceuticals- A compendium of Guide lines and Related materials Vol I & II, 2<sup>nd</sup> edition, WHO Publications, 1999.
- 4. How to Practice GMP's P P Sharma, Vandana Publications, Agra, 1991.
- The International Pharmacopoeia vol I, II, III, IV & V General Methods of Analysis and Quality specification for Pharmaceutical Substances, Excipients and Dosage forms, 3<sup>rd</sup> edition, WHO, Geneva, 2005.
- Good laboratory Practice Regulations Allen F. Hirsch, Volume 38, Marcel Dekker Series, 1989.
- 7. ICH guidelines
- 8. ISO 9000 and total quality management
- The drugs and cosmetics act 1940 Deshpande, Nilesh Gandhi, 4<sup>th</sup> edition, Susmit Publishers, 2006.
- 10. QA Manual D.H. Shah, 1<sup>st</sup> edition, Business Horizons, 2000.
- Good Manufacturing Practices for Pharmaceuticals a plan for total quality control Sidney H. Willig, Vol. 52, 3<sup>rd</sup> edition, Marcel Dekker Series.
- Steinborn L. GMP/ISO Quality Audit Manual for Healthcare Manufacturers and Their Suppliers, Sixth Edition, (Volume 1 - With Checklists and Software Package). Taylor & Francis; 2003.
- Sarker DK. Quality Systems and Controls for Pharmaceuticals. John Wiley & Sons; 2008.

#### **SEMESTER II**

# PCH-MPA204T: HERBAL AND COSMETIC ANALYSIS

COU	URSE CODE	PCH-MPA204T					
COU	URSE TITLE	HERBAL AND COS	METICS AN	NALYSIS	(Theor	ry)	
		UMMARY		CTIVES/C			
Regu inter evalu for equij	course is converted by the course of a course is converted by the course of the course of the course of the course.	<ul><li>be able to I</li><li>1. Regular</li><li>2. Analys</li><li>3. Herbal</li><li>4. Manufa</li></ul>	tory aspects tory aspects is of natura drug-drug acturing and ts as per reg	s of he l produ interac nd eva gulator	erbal rem ucts and ction aluation	monographs of cosmetic	
							of marks of
Sl No		rse Content	· •	Marks of assessment	ex (30 ma	assess sional xam 0% of rks of ssment) S2	sment End Sem exam (70% of marks of assessment)
1	with respect Regulations Conventional	out Herbal remedies to Toxicity and of Herbals vs drugs, Herbal drug based on WHO and ines	1 (10 hrs)	20	5		15
2	Deterioration various met regulatory requ herbal drug ind		2 (10 hrs)	20	5		15
3	products and laboratory, Ad Regulation and drugs, Stability products, proto of Herbal of Pharmacopoeia	Testing of natural drugs in clinical dulterant Screening, dispensing of herbal y testing of natural ocol and Monographs drugs in different WHO guidelines in tent of herbal drugs.	3 (12 hrs)	25	5	3	17

4	Will learn about Herbal drug-drug interaction with respect to WHO and AYUSH guidelines, Spontaneous reporting schemes for adverse reactions and Challenges in monitoring the safety of herbal medicines.	4 (10 hrs)	20		6	14
5	Will understand Evaluation of cosmetic products with respect to the main ingredients used and general methods of analysis of raw material used in cosmetic manufacture as per BIS and Indian Standard specification laid down for sampling and testing of various cosmetics in finished forms.	5 (10 hrs)	20		6	14
	Total Marks of A	Assessment	105	15	15	75

#### PCH-MPA204T: HERBAL AND COSMETIC ANALYSIS

#### THEORY

#### 52 hrs

Herbal remedies- Toxicity and Regulations: Herbals vs Conventional drugs, Efficacy of herbal medicine products. Validation of Herbal Therapies, Pharmacodynamic and Pharmacokinetic issues. Herbal drug standardization: WHO and AYUSH guidelines. 10 hrs
 Adulteration and Deterioration: Introduction, types of adulteration/substitution of herbal drugs, Causes and Measure of adulteration, Sampling Procedures, Determination of Foreign Matter, DNA Finger printing techniques in identification of drugs of natural origin, heavy metals, pesticide residues, phototoxin and microbial contamination in herbal formulations. Regulatory requirements for setting herbal drug industry: Global marketing management, Indian and international patent law as applicable herbal drugs and natural products and its protocol. 10 hrs

3. Testing of natural products and drugs: Effect of herbal medicine on clinical laboratory testing, Adulterant Screening using modern analytical instruments, Regulation and dispensing of herbal drugs, Stability testing of natural products, protocol.

Monographs of Herbal drugs: Study of monographs of herbal drugs and comparative study in IP, USP, Ayurvedic Pharmacopoeia, American herbal Pharmacopoeia, British herbal Pharmacopoeia, Siddha and Unani Pharmacopoeia, WHO guidelines in quality assessment of herbal drugs. 12 hrs 4. Herbal drug-drug interaction: WHO and AYUSH guidelines for safety monitoring of natural medicine, Spontaneous reporting schemes for bio drug adverse reactions, bio drug-drug and bio drug-food interactions with suitable examples. Challenges in monitoring the safety of herbal medicines. **10 hrs** 

5. Evaluation of cosmetic products: Determination of acid value, ester value, saponification value, iodine value, peroxide value, rancidity, moisture, ash, volatile matter, heavy metals, fineness of powder, density, viscosity of cosmetic raw materials and finished products. Study of quality of raw materials and general methods of analysis of raw material used in cosmetic manufacture as per BIS.

Indian Standard specification laid down for sampling and testing of various cosmetics in finished forms such as baby care products, skin care products, dental products, personal hygiene preparations, lipsticks. Hair products and skin creams by the Bureau Indian Standards. 10 hrs

- 1. Pharmacognosy by Trease and Evans
- 2. Pharmacognosy by Kokate, Purohit and Gokhale
- 3. Quality Control Methods for Medicinal Plant, WHO, Geneva
- 4. Pharmacognosy & Pharmacobiotechnology by Ashutosh Kar
- 5. Essential of Pharmacognosy by Dr.S.H.Ansari
- Cosmetics Formulation, Manufacturing and Quality Control, P.P. Sharma, 4<sup>th</sup> edition, Vandana Publications Pvt. Ltd., Delhi
- 7. Indian Standard specification, for raw materials, BIS, New Delhi.
- 8. Indian Standard specification for 28 finished cosmetics BIS, New Delhi
- 9. Harry's Cosmeticology 8th edition
- 10. Suppliers catalogue on specialized cosmetic excipients
- 11. Wilkinson, Moore, seventh edition, George Godwin. Poucher's Perfumes, Cosmetics and Soaps
- 12. Hilda Butler, 10th Edition, Kluwer Academic Publishers. Handbook of Cosmetic Science and Technology, 3rd Edition.

### MPHARM – PHARMACEUTICAL ANALYSIS (MPA)

#### SEMESTER II

### PCH-MPA205P: PHARMACEUTICAL ANALYSIS PRACTICAL II

COU	RSE CODE PCH-MPA205P					
COU	RSE TITLE PHARMACEUTICAI	ANALYSIS PRACTICAL – I				
	SCOPE/ SUMMARY	OBJEC	TIVES/CC	OURSE OUTC	COMES	
practi interp bioan	course is designed to impart cal knowledge on spectral oretation and analytical and alytical techniques of drugs and naceuticals.	<ul> <li>On completion of this course student shall be able to</li> <li>1. Interpret the spectra</li> <li>2. Separate analyte form biological samples by various techniques.</li> <li>3. Do the Qualitative and quantitative analysis of Pharmacopoeial compounds and their formulations in biological samples</li> <li>4. Do Quality control tests</li> </ul>				
	Course Co	ontent and Ass	essment Pla	an		
SI No.	Course Content	Syllabus (Chapters or Units with hours)	Total Marks of assessment	Distribut assessment Sessional exam (25 % of total marks of assessment) S1		
1	Learn to compare the absorption spectra by UV and Wood ward – Fieser rule, interpret spectra of organic compounds by FT-IR, NMR and MS, determine the purity by DSC in pharmaceuticals, identify organic compounds using FT-IR, NMR, CNMR and Mass spectra and quality control tests for primary and secondary packing materials.	Experiments 01 to 6 and 13 (46 hrs)	40	08	32	
2	Learn the separation of biomolecules utilizing various sample preparation techniques and quantitative analysis of components by gel electrophoresis , quantitative analysis of components by HPLC techniques, isolation of analgesics from biological fluids	7,8,9, 14, 15 and 18 to 23 (84 hrs)	80	18	62	

	(Blood serum and urine),assay of raw materials as per official monographs, testing of related and foreign substances in drugs and raw materials, quantitative analysis of rancidity in lipsticks and hair oil,determination of aryl amine content and developer in hair dye,determination of foam height and SLS content of Shampoo, total fatty matter in creams (Soap, skin and hair creams), acid value and saponification value, and determination of calcium thioglycolate in depilatories				
3	Learn the protocol preparation and performance of analytical/Bioanalytical method validation, protocol preparation for the conduct of BA/BE studies according to guidelines, in-process and finished product quality control tests for tablets, capsules, parenteral and creams and preparation of Master Formula Record and Batch Manufacturing Record.	Experiments 10,11 12,16 and17 (14 hrs)	10	4	06
	Total Marks o	of Assessment	130	30	100

### PCH-MPA205P: PHARMACEUTICAL ANALYSIS PRACTICAL II

- 1. Comparison of absorption spectra by UV and Wood ward Fieser rule
- 2. Interpretation of organic compounds by FT-IR
- 3. Interpretation of organic compounds by NMR
- 4. Interpretation of organic compounds by MS
- 5. Determination of purity by DSC in pharmaceuticals
- 6. Identification of organic compounds using FT-IR, NMR, CNMR and Mass spectra

7. Biomolecules separation utilizing various sample preparation techniques and Quantitative analysis of components by gel electrophoresis.

8. Biomolecules separation utilizing various sample preparation techniques and Quantitative analysis of components by HPLC techniques.

9. Isolation of analgesics from biological fluids (Blood serum and urine).

- 10. Protocol preparation and performance of analytical/Bioanalytical method validation.
- 11. Protocol preparation for the conduct of BA/BE studies according to guidelines.

12. In process and finished product quality control tests for tablets, capsules, parenterals and creams

- 13. Quality control tests for Primary and secondary packing materials
- 14. Assay of raw materials as per official monographs
- 15. Testing of related and foreign substances in drugs and raw materials
- 16. Preparation of Master Formula Record.
- 17. Preparation of Batch Manufacturing Record.
- 18. Quantitative analysis of rancidity in lipsticks and hair oil
- 19. Determination of aryl amine content and Developer in hair dye
- 20. Determination of foam height and SLS content of Shampoo.
- 21. Determination of total fatty matter in creams (Soap, skin and hair creams)
- 22. Determination of acid value and saponification value.
- 23. Determination of calcium thioglycolate in depilatories.

# MPHARM –PHARMACEUTICAL ANALYSIS (MPA) SEMESTER II

### PCH-MPA206S: SEMINAR IN PHARMACEUTICAL ANALYSIS

COUR	SE CODE PCH-MPA206S					
COUR	SE TITLE SEMINAR IN PHARMA	CEUTICAL ANALY	YSIS			
	SCOPE/ SUMMARY	OBJECTIVES	/COURSE OU	JTCOMES		
enviro the stu fortify skills	subject is designed to create an nment where teachers provide dents a critical eye and openness to the presentation and academic writing of students in the field of aceutical Analysis.	<ul> <li>Upon completion of the course the student shall be able to:</li> <li>1. Develop skills to gather, organize, deliver information, and defend a given topic in Pharmaceutical Analysis</li> <li>2. Learn to organize analytical concepts using audio-visual aids.</li> <li>3. Acquire communication and presentation skills.</li> <li>4. Effectively respond to the questions raised by peers and stand scientific scrutiny.</li> <li>5. Develop scientific writing skill.</li> <li>6. Cultivate a sense of upgradation of knowledge through self and continuous</li> </ul>				
	Course Content a	and Assessment Plan				
SI No.	Course Content	Hours	Total Marks of assessment	Marks End Sem exam		
1.	The students should be able to develop skills to gather, organize, communicate the scientific information, and defend a given topic in Pharmaceutical Analysis	2 hours/week	100	No end- semester examination. Only continuous mode.		

### MPHARM – PHARMACEUTICAL ANALYSIS (MPA)

#### SEMESTER III

### PHA-MRM301T: RESEARCH METHODOLOGY AND BIOSTATISTICS

COUR	SE CODE	PHA-MRM301T							
COURSE TITLE RESEARCH METHODOLOGY AND BIOSTATISTICS (Theory)									
		UMMARY		CTIVES / C					
This subject is designed to understand the advanced knowledge for research methodology, ethics in research, medical research, design, conduct and interpretation of results. This subject deals with principles of statistics and their applications in biostatistics involving parametric tests, non-parametric tests, correlation, regression, probability theory and statistical hypotheses.Upon completion of the course, the student shall be able to1. Know the various components of research design and methodology.2. Appreciate advanced statistical techniqu in solving the research problems.									
					Distr		f marks of		
Sl No.	Cou	urse Contents	Syllabus (Chapters or Units with hours)	Total Marks of assessment	(80 %	assessm al exam of total of ent)			
					<b>S1</b>	S2			
1	Understand Research study design	Methodology and	Unit I (10 hrs)	20	20		-		
2	and their biostatistics various	. Besides, learning techniques of to interpret the	Unit II (12 hrs)	20	20		-		
3	records and handling experimenta		Unit III (10 hrs)	10		10	-		
4		l learn the history, and concepts of earch.	Unit IV (10 hrs)	20		20	-		
5	for all me additional	ry, basic principles dical research and principles for esearch combined al care.	Unit V (10 hrs)	10		10	-		
		<b>T</b> 11(1)	Assessment	80	40	40			

#### PHA-MRM301T: RESEARCH METHODOLOGY AND BIOSTATISTICS

#### THEORY

52 hrs

#### UNIT – I

General Research Methodology: Research objectives, requirements, practical difficulties, review of literature, study design, types of studies, strategies to eliminate errors/bias, controls, randomization, crossover design, placebo, blinding techniques.

#### $\mathbf{UNIT} - \mathbf{II}$

Biostatistics: Definition, application, sample size, importance of sample size, factors influencing sample size, dropouts, statistical tests of significance, null hypothesis, P values, degree of freedom, interpretation of P values. Type of significance tests, parametric tests (students "t" test, ANOVA, correlation coefficient, regression), non-parametric tests (Wilcoxon rank tests, analysis of variance, correlation, chi-square test),

#### UNIT – III

CPCSEA guidelines for laboratory animal facility: Goals, veterinary care, quarantine, surveillance, diagnosis, treatment and control of disease, personal hygiene, location of animal facilities and laboratories, anesthesia, euthanasia, physical facilities, environment, animal husbandry, record keeping, SOPs, personnel and training, transport of lab animals.

#### $\mathbf{UNIT} - \mathbf{IV}$

Medical Research: History, values in medical ethics, autonomy, beneficence, nonmaleficence, double effect, conflicts between autonomy and beneficence/non-maleficence, euthanasia, informed consent, confidentiality, criticisms of orthodox medical ethics, importance of communication, control resolution guidelines, ethics committees, cultural concerns, truth telling, online business practices, conflicts of interest, referral, vendor relationships, treatment of family members, sexual relationships fatality.

#### $\mathbf{UNIT} - \mathbf{V}$

Declaration of Helsinki: History, introduction, basic principles for all medical research and additional principles for medical research combined with medical care.

# MPHARM –PHARMACEUTICAL ANALYSIS (MPA) SEMESTER III

#### MJC302P: JOURNAL CLUB IN PHARMACEUTICAL ANALYSIS

CO	URSE CODE MJC302P						
CO	URSE TITLE JOURNAL CLUB I	N PI	HARMACEUTIC	CAL ANALYS	SIS		
	SCOPE/ SUMMARY		OBJECTIVES/COURSE OUTCOMES				
The subject is designed to create an environment where students present a published research paper, and critically analyse it, that would enhance the communication, presentation and analytical skills of the students.			<ul> <li>Upon completion of the course the student shall be able to:</li> <li>1. Learn to organize research concepts using audio-visual aids.</li> <li>2. Acquire communication and presentation skills.</li> <li>3. Effectively respond to the questions raised by peers and stand scientific scrutiny.</li> <li>4. Cultivate a sense of upgradation of knowledge</li> </ul>				
	Course Cor	ntent	t and Assessment Plan				
					Marks		
Sl No.	Course Content		Hours	Total Marks of assessment	End Sem exam		
1.	The students should be able to devel skills to gather, organize, deliv information, and defend a giv research topic in pharmaceuti Analysis.	ver ven	2 hours/week		No end-semester examination. Only continuous mode.		

## MPHARM – CHOICE BASED INTERDISCIPLINARY COURSES PCE-001E: GENERIC DRUG DEVELOPMENT

#### (15 hrs)

Introduction to Generic Drug Product Development, API, Analytical Methods Development and Methods Validation for Solid Oral Dosage Forms, Experimental formulation development, Scale-Up, Process Validation, Technology Transfer, Drug stability, QC, QA. Drug product performance in vitro, ANDA Regulatory approval process, BE and drug product assessment in vivo, SUPAC, Outsourcing BA and BE studies to CROs, Legal and legislative hurdles.

#### REFERENCES

- Handbook of Pharmaceutical Generic Development, (oral dosage form volume 24 of Drug development series), Locum publishing house, USA.
- Generic Drug Product Development-Solid Oral dosage form, Leon Shargel and Isadore Kanfer, Marcel Dekker, USA, ISBN: 0-8247-5460-3.

#### PCE-002E: PHARMACEUTICAL DISSOLUTION TECHNOLOGY

(15 hrs)

2 hrs
2 hrs
2 hrs
2 hrs
1 hr
1 hr
2 hrs
1 hr
2 hrs

#### REFERENCES

- Pharmaceutical Dissolution Testing by Umesh V. Banakar; Singapore: CRC Press Taylor & Francis Group.
- Pharmaceutical Dissolution Testing by Jennifer Dressman and Johannes Krämer; Singapore: Taylor & Francis Group.

#### PCE-003E: PARTICULATE DRUG DELIVERY SYSTEMS

#### (15 hrs)

**Microparticulate drug delivery Systems:** Introduction, advantages, disadvantages, types, methods of preparation & characterization, in vitro & in vivo evaluations and applications.

#### 6 hrs

**Nanoparticulate drug delivery Systems:** Introduction, advantages, disadvantages, types, methods of preparation & characterization, in vitro & in vivo evaluations and applications.

#### 9 hrs

#### REFERENCES

- 1. Encyclopedia of controlled delivery, Editor- Edith Mathiowitz, Published by Wiley Interscience Publication, John Wiley and Sons, Inc, New York. Chichester/Weinheim
- N.K. Jain, Controlled and Novel Drug Delivery, CBS Publishers & Distributors, New Delhi, First edition 1997 (reprint in 2001)
- 3. S.P.Vyas and R.K.Khar, Controlled Drug Delivery concepts and advances, Vallabh Prakashan, New Delhi, First edition 2002
- 4. Nanoteachnology in Drug Delivery, Melgardt M. De Villers, Pornanong Aramwit and Glen S Kwon, Springer.
- 5. Nanoparticulate Drug Delivery Systems, Deepak Thassu, Michel Delees and Yashwant Pathak, Informa Healthcare, NY, USA.
- Nanoparticulates as Drug Carriers, Vladimir P Torchilin, Imperial College Press, London.

### PCE-004E: 3D PRINTING IN PHARMACEUTICAL MANUFACTURING (15 hrs)

Introduction, 3D printing technologies, printing based inkjet systems, Nozzle based deposition systems, Laser based writing systems, 3D printing for customized drug delivery systems, Limitations and challenges.

#### REFERENCES

- 1. 3D printing in pharmaceutics: A new tool for designing customized drug delivery systems. Goole Jonathan and Amighi Karim, Int. J Pharm. 499 (2016), 376-394.
- 3D fabricated polymer-based drug delivery systems, Simon Moulton and Gordon G Wallace, J of Cont. Rel. 193 (2014), 27-34.
- Fabrication of printed drug delivery systems. Natalja Genina, Ruzica Kolakovic, Mirja Palo, Daniela Fors, Helka Juvonen, Petri Ihalainen, Jouko Peltonen, Niklas Sandler, NIP 29 and Digital Fabrication (2013), 236-238.

### PCH-001E: PREPARATIVE SEPARATION TECHNIQUES (15 hrs)

- Column chromatography: Introduction, stationary phase, mobile phase selection, column selection, sample loading techniques, elution technique.
   9 hrs
- Flash chromatography: Principle, advantages, instrumentation, mobile phase selection, column selection, sample loading techniques, elution technique.
   6 hrs

### PCH-002E: MOLECULAR MODELLING AND DRUG DESIGN (15 hrs)

1. Molecular Geometry, Molecular mechanics, Quantum mechanics, Molecular dynamics, Pharmcophore, Molecular docking, Library generation and structure-based drug design.

> 12 hrs 3 hrs

2. Database and Software Resources

#### PCH-003E: HYPHENATED TECHNIQUES

(15 hrs)

Principle and applications of following hyphenated techniques

1. GC-MS	4. EC-MS	7. LC-MS-MS	10. GC-AES
2. LC-MS	5. CE-MS	8. GC-MS-MS	
3. LC-NMR	6. GC-IR	9. GC-NMR	

### PCH-004E: CHEMICALS-ENVIRONMENT, HEALTH AND SAFETY

(15 hrs)

Chemical safety, Chemical	hazards,	handling	of	chemicals/gases,	storage	of	chemicals,
chemical waste disposal.							8 hrs
First aid procedures							1 hr

ory practice	es:					2 hrs
tection						1 hr
materials:	Regulatory	requirements,	hazards,	handling,	storage,	disposal,
rocedures.						2 hrs
						1 hr
	tection materials:	materials: Regulatory	tection materials: Regulatory requirements,	tection materials: Regulatory requirements, hazards,	tection materials: Regulatory requirements, hazards, handling,	tection materials: Regulatory requirements, hazards, handling, storage,

# PQA-001E: THEORY AND PRACTICE OF ANALYTICAL AND BIOANALYTICAL METHOD DEVELOPMENT AND VALIDATION (15 hrs)

1.	Introduction to the concept of validation.	1 hr
2.	Development of analytical method using UV/PDA spectroscopy,	
	Fluorimetry, HPLC, LC-MS/MS.	4 hrs
3.	Validation of the analytical method as per ICH-Q2(R1).	3 hrs
4.	Development of bioanalytical method using HPLC and LC-MS/MS.	2 hrs
5.	Validation of bioanalytical method as per USFDA guidance.	3 hrs
6.	Introduction to Novel upcoming technologies in bioanalysis	
	like dry matrix spot analysis.	1 hr
7.	Introduction to Analysis of therapeutic proteins and peptides.	1 hr

#### **Evaluation**

Formative: Development of validation protocols & problem-based learning	g. (30%)
Summative: Open book periodical tests & end semester exam.	(70%)

# PQA-002E: GOOD DOCUMENTATION PRACTICES AND E-DOCUMENTATION PRACTICES IN PHARMACEUTICAL INDUSTRY

#### (15 hrs)

1.	Introduction to GDP and E – documentation	3 hrs
2.	Basic levels of documentation	6 hrs
	a. Level -1, Level-2, Level-3 and Level-4 documentation	
3.	Case studies in each level	3 hrs
4.	Open lab and e-documentation concept	3 hrs

### **PQA-003E: TROUBLE SHOOTING IN HIGH PERFORMANCE LIQUID**

#### **CHROMATOGRAPHY**

#### (15 hrs)

1.	Introducti	on to	HPLC	mod	ules an	nd so	urce o	of errors/malfunction in HPLC	5 hrs
-	~								

- 2. Startup preliminary checks for trouble shooting6 hrs
- 3. Trouble shooting in HPLC module wise including demonstration 4 hrs

#### PQA-004E: PROFESSIONAL DEVELOPMENT

#### (15 hrs)

- 1. Introduction to Professional Career Development
- 2. Introduction to Career Planning: Self Assessment
- 3. Identifying Your Professional Talents
- 4. Introduction to Career Planning: Career Exploration
- 5. Developing Your Professional Resume
- 6. Enhancing Your Professional Resume
- 7. Preparing Your Career and Internship Cover Letters
- 8. Professional Communications
- 9. Preparing for Your Employment an Internship Interviews
- 10. Conducting Your Employment and Internship Interviews
- 11. Introduction to the Career Fair Search Process
- 12. Exploring Internship Options within Your Profession
- 13. Networking Search Strategies
- 14. Developing Your Professional Career Portfolio
- 15. Influencing Your Networking Partners
- 16. Essential practical skills and problem solving
- 17. communication of scientific information
- 18. IT skills.

#### Assessments:

- assignments
- case studies
- portfolios
- presentations

#### PQA-005E: STABILITY TESTING OF DRUGS AND BIOLOGICALS

#### (15 hrs)

- 1. Introduction to drug stability and its importance.
- 2. Stability testing of drug substances and drug products as ICH Q1 series guidelines 11 hrs
- 3. Stability testing of biotechnological/biological products as per ICH Q5C guidelines. 2 hrs

#### PQA-006E: USFDA DRUG REGULATORY AFFAIRS

#### (15 hrs)

- 1. Process of new drug development.
- 2. Contents of IND, NDA, CTD and eCTD, ANDA
- 3. SMF and DMF
- 4. Post marketing regulatory requirements.

#### PQA-007E: REST OF THE WORLD DRUG REGULATIONS

#### (15 hrs)

Legislation and regulations for import, manufacture, distribution and sale of drugs and pharmaceuticals in:

- 1. Brazil
- 2. ASEAN countries
- 3. CIS countries
- 4. GCC Countries.

#### PQA-008E: EVALUATION OF MEDICAL DEVICES

#### (15 hrs)

A. Biological evaluation of medical devices

Scope, General principles applying to biological evaluation of medical devices, categorization of medical devices, testing, selection of biological evaluation tests, Assurance of test methods

B. Clinical evaluation of Medical devices 5 hrsImportance, scope, clinical evaluation in brief

10 hrs

#### **PBT-001E: CLEAN ROOM CONCEPTS**

#### (15 hrs)

#### Unit 1. Fundamental aspects of microbiology

Morphology, Microscopy, Growth and Controlling Growth of Microorganisms.

#### **Unit 2. Clean Room aspects**

Clean Room concepts, layout, facilities, HEPA and ULPA Filter, biosafety cabinets and various classes, biosafety levels, personnel controls.

Unit 3. Microbial monitoring, detection and enumeration of microorganisms 6 hrs Monitoring techniques, air samples, microbiological assessment of liquids, solids and semisolids, disposal of cultures.

#### REFERENCES

- 1. Cleanroom Microbiology for the Non-microbiology by David M. Carlberg, Second Edition CRC press, 2005.
- 2. Biopharmaceutical, Biochemistry and Biotechnology, Second Edition, Gary Walsh, Wiley Publishers, 2003.

#### PBT-002E: BIOSIMILARS

#### (15 hrs)

#### **Unit -I Biosimilars- Introduction**

Definitions, Generics and Branded drugs, biosimilars, introduction to biologics, differences between biosimilars and generics, manufacturing process and technical challenges associated with production of biosimilar molecules, status of biosimilars. The role of patents in the drug industry and protein-based biopharmaceuticals.

#### Unit -II Guidelines on Similar Biologic: Regulatory Requirements for Registration 8 hrs and Marketing Authorization in India

Principles for Development of Similar Biologics, Competent authorities, Selection of reference biologics, Quality consideration of similar biologics, Quality comparability study, Data requirement for preclinical study, clinical application and market authorisation, Post market data for similar biologics.

#### REFERENCES

- 1. Proposed guidelines for similar biologics in India, CDSCO.
- 2. Biosimilars and Interchangeable Biologics: Strategic Elements By Sarfaraz K. Niazi
- 3. Guidelines from Indian regulatory bodies such as CDSCO, DBT etc. issued time to time.
- 4. Relevant journals and periodicals.

7 hrs

3 hrs

#### **PBT-003E: PRINCIPLES OF GENE CLONING**

#### (15 hrs)

Unit I The aims of Gene Cloning: Techniques of gene manipulation, Outline of gene cloning. Unit II 6 hrs Gene Cloning: Gene cloning procedure, tools required for gene cloning, cloning strategy, techniques for selection of clones, preservation of clones. **Unit III** 6 hrs **Applications of Gene Cloning:** In production of therapeutic proteins, diagnostic proteins, transgenic animals and plants. REFERENCES

- 1. Molecular biotechnology, Bernard R. Glick, Jack J. Pasternak, Cheryl L. Patten, 2010, ASM Press.
- 2. Biotechnology: The Biological Principles, M.D. Trevan, S. Boffey, K.H. Goulding and P. Stanbury. 1998, Tata McGraw Hill Edition.

#### **PBT-004E: TISSUE ENGINEERING**

#### (15 hrs)

# Introduction to Tissue Engineering: Animal cell culture fundamentals, concept and need of tissue engineering, historical prospective, current status, industry challenges

Unit II

Biomaterials for Tissue Engineering: Overview, features of scaffold and biomaterials, types of biomaterials, nanofiberous material as biomaterial.

#### **Unit III**

Applications of Tissue Engineering: in Bone tissue regeneration, vascular tissue engineering, skin regeneration, cardiac tissue engineering and neural tissue engineering.

#### REFERENCES

- 1. Principles of Tissue Engineering, 4th Edition, Robert Vanza, Robert Langer, Joseph Vacanti. 2014. Academic Press.
- 2. Exploring Nanotechnology in Healthcare, Edited by N. Udupa, 2013, Manipal University Press.

# Unit I

#### Page | 83

5 hrs

5 hrs

5 hrs

#### PPR-001E: RETAIL PHARMACY PRACTICE

#### (15 hrs)

- Retail Pharmacy Management: Site selection, acquisition of premises for a retail pharmacy, layout of drug store, Legal aspects of retail pharmacy (includes Schedule N requirement), role of retail pharmacist and Code of ethics for practicing pharmacists.
- Purchase and inventory control, stocking, sale promotion, maintenance of records, economics and management
   5 hrs
- 3. Communication skills2 hrs4. Medication therapy management2 hrs5. Patient counselling2 hrs

#### REFERENCES

I. Introduction

- Philip P. Gerbino. Remington: The Science and Practice of Pharmacy. Philadelphia, PA. Lippincott Williams & Wilkins. (latest edition)
- 2. Revikumar KG,Miglani BD. A Text Book of Pharmacy Practice. Career Publications (latest edition)
- G Parthasarathi, Karin Nyfort Hansen, Milep C Nahata. A Textbook of Clinical Pharmacy Practice Essential Concept and Skill. Oriental Longman Private Limited (latest edition)
- 4. Ashley WE, Justin J. Community and clinical pharmacy services: A step-by-step approach. The McGraw-Hill Companies, Inc, USA

#### **PPR-002E: FUNDAMENTALS OF MEDICAL WRITING**

(15 hrs)

	$\triangleright$	Brief overview of scientific writing	
	$\triangleright$	Scope and importance	
	$\triangleright$	Different types and areas of writing	
	۶	Career and opportunities	
2. Bas	sic N	Need To Be A Good	4 hrs
	~		
		Language and Style in Medical Writing	
		Literature search	
		-Data bases (Medline, PubMed, Cochrane)	

- Searching principles (using MeSH, Pub Med)
- Developing searching strategy by PICO
- Cortical Analysis Scientific Paper
- > Ethics in Publication (Plagiarism, Copy Rights etc)
- ➢ Reference Writing
  - Different bibliographic styles
  - -Citation databases
  - -Software used in reference writing

#### 3. Different Types of Medical Writing

- Structured abstract writing
- Report writing and sub-types
- Medication leaflets/pills
- Clinical research form
- Informed consent
- Protocol writing
- Case record form
- > PSUR
- > News letter

#### 4. MANUSCRIPT WRTING AND PUBLICATION

- ➢ ICMJE guidelines
- ➢ How to prepare structured manuscript (IMRA)
- Presentation of data (tables , figures and algorithms)
- Conflict of interest
- Acknowledgement
- Publication issues

**Assignments**: Preparation of power point, poster, abstract writing, review article with hand on training in publication issues

#### REFERENCES

1.Janice R Mathews, Jobin M Bowen and Robert W Mathews .Successful scientific writing-A step by step guide for the biological and medical sciences; 1996

7 hrs

2. Piyush Gupta, Navjeevan Singh. How to write thesis and thesis protocol-A primer for medical, dental & nursing courses; 2014

3. John Kirkman. Good style – Writing for science & Technology; 1994

4. Jennifer peat, Elizabeth Elliot, Louise Bour. Scientific writing-Easy when you know how; 1994.

#### PPR-003E: SYSTEMATIC REVIEW AND META-ANALYSIS

#### (15 hrs)

1.	Study designs: Introduction to Case-control studies, Cohort studies, Randomized	
	controlled trials	1 hr
2.	Applied statistics: Descriptive statistics, Hypothesis, Null-hypothesis, type-1 & ty	pe-2
	errors, power, p-value, Confidence intervals, Odds ratio, Relative risk (risk ratio),	Fixed
	effects & Random effects, Concept of homogeneity & heterogeneity and tests for	
	heterogeneity, Various types bias and methods to detect bias, Funnel plot, Effect s	ize &
	effect size indices, Forest plot	3 hrs
3.	Evidence based clinical practice: Definition, importance, levels of evidence.	1 hr
4.	Systematic review and meta-analysis: Definition, types, importance, applications,	
	Meta-analysis groups (Campbell Collaboration, Cochrane Collaboration)	1 hr
5.	Steps involved in conducting Systematic review and Meta-analysis:	5 hrs
	a. Framing the question	
	b. Literature search	
	c. Assessing the quality of studies	
	d. Selection of studies	
	e. Data synthesis & Analysis	
	f.Summarizing the evidence	
	g. Interpretation of the findings	
6.	Introduction to softwares used in meta-analysis: MedCalc, Comprehensive Meta-	
	analysis software, RevMan, Open meta-analysis	1 hr
7.	Writing a meta-analysis protocol, Literature search, Data synthesis & analysis	
	(Assignments)	3 hrs

#### **REFERENCES**:

 Higgins JPT, Green S, editors. Cochrane Handbook for Systematic Reviews of Interventions 4.2.6 [updated September 2006]. In: The Cochrane Library, Issue 4, 2006. Chichester, UK: John Wiley & Sons, Ltd.  Borenstein, M, Hedges LV, Higgins JPT, Rothstein HR. Introduction to Meta-Analysis. John Wiley & Sons, Ltd., 2009.

**Pre-requisites**: Knowledge of Biostatistics & Research Methodology, Web-based literature search

Evaluation: Based on Assignments

# **PPR-004E: PHARMACOKINETICS DATA ANALYSIS**

#### (Employing WinNonlin)

#### (15 hrs)

- 1. Introduction to pharmacokinetic parameters: Elimination rate constant (ke), Elimination half-life, Bioavailability & AUC, Volume of distribution, Clearance, Bioequivalence 2 hrs
- 2. Bioavailability studies: In animal & human 2 hrs
- 3. PK parameters for Oral & IV administration: Calculation of PK parameters for oral & IV administration by plotting concentration vs time graph (using semi-log graph paper)

		2 hrs
4.	Introduction Phoenix WinNonlin: Data entry and data tools, graphs	2 hrs
5.	Non-compartmental analysis using (NCA) Phoenix WinNonlin: For Oral, IV, IV	
	infusion, Sparse sampling and urinary excretion data	3 hrs
6.	Pharmacokinetic modeling: Compartment modelling, choosing the right compart	nent
	model, Simulating using PK model	2 hrs
7.	Bioequivalence data analysis: Parallel, Cross-over study data analysis	2 hrs
RI	EFERENCES	
	1. Gibaldi M, Perrier D. Pharmacokinetics. 2 <sup>nd</sup> edition. Informa Healthcare; 200	7.
		.1

- Rowland M, Tozer TN. Clinical Pharmacokinetics: Concepts & Applications. 4<sup>th</sup> edition. Lippincott Williams& Wilkins;2011.
- 3. Phoenix WinNonlin examples guide. Pharsight. A Certara Company; 2012
- 4. Phoenix WinNonlin Users guide. Pharsight. A Certara Company; 2012.
- 5. Guidance for Industry: Bioavailability and Bioequivalence Studies Submitted in NDAs or INDs- General Considerations. US FDA. 2014.

Pre-requisites: Knowledge of basic pharmacokinetics.

Evaluation: Based on Assignments.

#### PHA-001E: CANCER BIOLOGY

#### (15 hrs)

#### **Objectives/Course Outcomes**

This course aims to provide students an extensive understanding about cancer and its effects on the human body. This course will also discuss the historical aspect of cancer research, basic chemotherapeutics and future scope in cancer research. All the topics will be presented in a simplified scientific manner with an emphasis on gaining a broad understanding of the disease.

- Advanced Cell Biology- Structural and functional basis of cell, structure of chromosome and DNA, Cell cycle and its regulation. 3 hrs
- Molecular and genetic basis of cancer- Growth signaling, dysfunction of cell cycle leading to cancer, Oncogene, tumor suppressor gene, Apoptosis
   6 hrs
- Cellular aspect of cancer- History of cancer, six hallmarks of cancer, Types of cancer, Metastasis, Cancer Prevention and Treatment.
   3 hrs
- Current trends in Cancer research- Targets for development of cancer chemotherapeutics, its identification and relevance, Recent advancements in cancer therapy.
   3 hrs

### PHA-002E: SCREENING METHODS FOR DRUG DEVELOPMENT (15 hrs)

The course aims at providing the conceptual learning of a few basic pharmacological approaches to elucidate the pharmacological activity of a new chemical entity. The knowledge of screening methods facilitates the understanding of animal models, which closely mimics the human pathology, to screen the promising molecules for developing a new drug.

- 1. Introduction: General Screening techniques
- 2. Screening methods for local anesthetics
- 3. Screening methods for anti-hypertensive drugs
- 4. Screening methods for anti-arrhythmic drugs
- 5. Screening methods for anti-inflammatory drugs
- 6. Screening methods for analgesic drugs
- 7. Screening methods for antipyretic drugs
- 8. Screening methods for anticancer drugs
- 9. Screening methods for antidiabetic drugs

- 10. Screening methods for anti-dyslipidemia drugs
- 11. Screening methods for antiepileptic drugs
- 12. Screening methods for antidepressant drugs
- 13. Screening methods for antianxiety drugs
- 14. Screening methods for antiparkinsonian drugs
- 15. Screening methods for Alzheimer's disease related dementia

### PHA-003E: FREE RADICAL BIOLOGY AND MEDICINE (15 hrs)

#### Objectives

To provide fundamentals which are essential for researchers who wish to pursue problems of human health that involve free radicals, related oxidants, antioxidants, and antioxidant enzymes. Students will be able to understand, interpret, and critically think on issues associated with major causes of health problems.

- 1. Basics of free radicals in biology and medicine
- 2. Concept of oxidative stress in biology and diseases
- 3. Oxidative stress markers
- 4. Radical scavengers: The concept of antioxidants
- 5. Redox signaling and NRF2-ARE signaling mechanisms
- 6. Free radicals in inflammation and immune disorders
- 7. Free radicals in diabetes, obesity and metabolic disorders
- 8. Free radicals in cancer
- 9. Free radicals in cardiovascular diseases
- 10. Free radicals in neurodegenerative disorders
- 11. Free radicals and ageing
- 12. Free radicals in infectious diseases and antimicrobial-resistance
- 13. Free radicals in hepato-biliary diseases
- 14. Antioxidants screening methods: In vitro and in vivo
- 15. Recent advances in antioxidant discovery and development

Study material: Recent journal articles from reputed and Open Access Journals

### PHA-004E: REGULATORY TOXICOLOGY IN DRUG DISCOVERY AND DEVELOPMENT (15 hrs)

#### Objectives

This subject is to project the importance of safety testing in pre-clinical research and harmonized standards that they have to adopt for safety testing procedures.

Introduction: General Principles of Toxicology, Agencies and their guidelines regardingToxicology studies, Concept of Good Laboratory practices.3hrs

#### Guidelines for safety testing

Pharmacological studies:Safety Pharmacology Studies for Human Pharmaceuticals, QTinterval prolongation study in animals.3 hrs

**Toxicity testing**: Toxicokinetics: The Assessment of Systemic Exposure in Toxicity Studies,Pharmacokinetics: Guidance for Repeated Dose Tissue Distribution Studies, Acute, subacuteand Chronic toxicity in animals**4 hrs** 

Special toxicity studies:Non-clinical Carcinogenicity studies, Genotoxicity studies,Reproductive toxicology, Immunotoxicity, Safety evaluation of Biotechnology- derivedproducts.5 hrs

### PCO-001E: NUTRACEUTICALS

#### (15 hrs)

#### Scope

Nutraceuticals: The food or part of food with additional health benefits are emerging as major health supplements for prevention, management and some-times cure of various diseases. In the current fast lifestyle, nutraceuticals are encroaching pharma markets not only in the developed but also in the developing countries. Therefore, an additional knowledge in nutraceuticals or health supplements add an edge to the student.

#### Objectives

The course proposes to offer a comprehensive knowledge on the nutraceuticals and their importance to prepare the student for a pursuit in the health food sector.

1.	Introduction to nutraceuticals: Overview, classification, be	enefits of 3	hrs
	nutraceuticals, functional foods		

2.	In organic supplements and vitamins	1 hr
3.	Probiotics, prebiotics and digestive enzymes	1 hr
4.	Dietary supplements and fibres	1 hr
5.	Antioxidants and PUFAs	2 hrs

- 6. Herbs as health foods: Poly phenols and flavonoids (*Ginkgo biloba*, 5 hrs Tea, Citrus fruits, Grape, Soy); Carotene and fatty acids: (Curcuma, Tomato, Flax seed, Olive oil)
  7. Current market scenario of nutraceuticals 1 hr
- 6. Regulatory requirements for nutraceuticals 1 hr

#### REFERENCES

- Biren Shah and A.K. Seth. Text Book of Pharmacognosy and Phytochemistry, Ed1, Elsevier Health Sciences, Gurgaon, Haryana, 2010. Pp 471-83
- 2. S.S. Agrawal and M. Paridhavi. Herbal Drug Technology, Ed2, Universities Press, Hyderabad, Telangana, 2012. Pp 710-21
- Alberta N.A. Aryee and Joyce Irene Boye, Current and Emerging Trends in the Formulation and Manufacture of Nutraceuticals and Functional Food products, In, Nutraceutical and Functional Food Processing Technology Edt. Joyce Irene Boye, Ed.1. Wiley Blackwell, West Sussex UK, 2015. Pp 01-52
- 4. Rorimi E. Aluko, Functional Foods and Nutraceuticals Springer, New York, USA, 2012.
- 5. C.K Kokate, A.P. Purohot and S.B. Gokhale, Pharmacognosy, Vol 1 and 2, Ed. 46, Nirali Prakashan, Pune, 2010. Pp 6.01-6.16

# PCO-002E: EXTRACTION, SEPARATION AND PURIFICATION OF <u>PHYTOCONSTITUENTS</u>

#### (15 hrs)

#### Scope

Herbal drugs are emerging as major source of biomedicines worldwide. Secondary metabolites or phytoconstituents are responsible for medicinal attributes of a plant. Extraction, separation, and purification of these secondary metabolite is an area of interest for researchers. Therefore, this subject will provide an insight on the topic so as to enable the students to carry their research on natural products effectively.

#### Objectives

To impart stronger scientific base in the area of extraction, isolation and purification principles and methodologies of crude drugs.

- 1. Introduction to plant metabolites.
- Extraction techniques: Principle, merits & demerits, applications of 5 hrs maceration, decoction, infusion, percolation, soxhlet extraction, supercritical fluid extraction, microwave-assisted extraction, ultrasound extraction (sonication), advanced phytonics method of extraction, expression and enfleurage method.
- 3. Phytochemical screening of natural products
- 4. Separation and purification of phytoconstituents: Fractional distillation, **7 hrs** fractional liberation, sublimation, fractional crystallization, gel filtration, counter current extraction and chromatographic techniques like Paper Chromatography, TLC, HPLC, HPTLC, column chromatography, gasliquid chromatography, droplet counter current chromatographyand electro-chromatography (Electrophoresis).

#### REFERENCES

- Evans W. C., Trease G. E., Trease and Evan's Pharmacognosy. W.B. Saunders, 2002. 16th Ed.
- Jean Bruneton, Caroline K. Hatton, Pharmacognosy, Phytochemistry, Medicinal Plants. Lavoisier, 1999
- 3. Kokate C.K., Gokhale S.B. and Purohit A.P., Textbook of Pharmacognosy, Nirali Prakashan, Pune, 2008
- Mukherjee Pulok K., Quality Control of Herbal Drugs: An Approach to Evaluation of Botanicals. Business Horizons, 2002.

### PCO-003E: NANOPHYTOPHARMACEUTICALS

(15 hrs)

#### Scope

Nanotechnology is a cutting-edge technology that provides optimal effectiveness in low doses. The course addresses and provides an overview of phytopharmaceuticals in nano size for better therapeutic benefits along with toxicity. Students will be able to understand the benefits of nano-phytopharmaceuticals.

#### **Objectives**

To study the effectiveness in low doses and enhancement of bioavailability of phytopharmaceuticals for better therapeutic values

1.	Definition and history of nanotechnology			
2.	Properties – optical, electrical and magnetic properties of	2 hrs		
	nanomaterials			
3.	Preparation techniques – Polymeric nanoparticles, liposomes, micelles	6 hrs		
	and herbal nanoparticles			
4.	Toxicity studies	2 hrs		
5.	Applications of phytopharmaceuticals, nanophytopharmaceuticals in			
	the treatment of certain diseases			

#### REFERENCES

- Nano: The Essentials: Understanding Nanoscience and Nanotecnology, T. Pradeep, Tata McGraw-Hill Publishing Company Limited, New Delhi, 2008.
- Nanocrystals: Synthesis, Properties and Applications, C. N. R. Rao, P. J. Thomas and G. U. Kulkarni, Springer (2007)
- Nanostructures & Nanomaterials: Synthesis, Properties & Applications, Guozhong Cao, Imperial College Press (2004).
- 4. Nanoparticles as Drug carriers, Vladimir P Torchilin, Imperial College Press, USA, 2006
- Multifunctional Pharmaceutical Nanocarriers, Vladimir Torchilin, Springer Publishing, New York, NY, 2008.

### PCO-004E: HERBAL MONOGRAPHS

(15 hrs)

#### Scope

A monographs is written document intended to promote information exchange and international harmonised standards for the quality control and use of herbal medicines. It contains scientific information on selected medicinal plants that serves as a summary for quality assurance, clinical applications and dosage forms. The proposed course is designed to study and understand herbal monographs

#### Objectives

To impart knowledge on systematic study of herbal drugs with reference its identity, quality and purity

- Introduction to monographs, purpose and content of the monographs, 3 hrs use of the monographs
- Systematic study of the following important plants for their 12 hrs monographs;

Leaf: Vasaka (*Adhatoda zeylanica*) Root: Shatavari (*Asparagus racemosus*) Rhizome:Rasna (*Alpinia galanga*) Bark: Cinchona (*Cinchona officinalis*) Fruit: Pepper (*Piper nigrum*) Entire herb: Kalmegh (*Andrographis paniculata*).

#### REFERENCES

- WHO monographs on selected medicinal plants. Geneva: WHO. Vol. 1. 1999, Vol. 2. 2002, – Vol. 3: 2004, – Vol. 4. 2005.
- Quality Standards of Indian Medicinal Plants Indian Council of Medical Research, New Delhi.
- Indian Herbal Pharmacopoeia A Joint Publication of RRL, Jammu and IDMA, Mumbai.

#### PRM-001E: RETAIL BUSINESS MANAGEMENT

#### (15 hrs)

#### Scope

This course is formulated to impart knowledge on retail management. It also provides an overview about the nitty-gritties of managing a pharmacy store and an overview of Online Pharmacies.

1. Introduction to Retail Management	3 hrs
2. Strategies in Retailing	3 hrs
3. Retail Marketing in rural areas	3 hrs
4. Pharmacy Store Management	4 hrs
5. Online Pharmacy Retailing	2 hrs

#### REFERENCES

- 1. Retail Management by Barry Berman. Person Education 11th Edition.
- 2. Retail Management by Chetan Bajaj. Oxford 2<sup>nd</sup> Edition.
- 3. Retail Management: Text and Cases by UC Mathur. IK International Publishing House Pvt. Ltd.

#### PRM-002E: INTELLECTUAL PROPERTY MANAGEMENT

#### (15 hrs)

#### Scope

This course deals with Intellectual Property Rights with special emphasis on Patents.

1. Basic Concepts of Intellectual Property Rights	3 hrs
2. Patent Administration in India and Patent Filing	3 hrs
3. Revocation of Patents and Patent Infringement Cases	3 hrs
4. Data Protection and Exclusivity	3 hrs
5. Patent as a business tool	3 hrs

#### REFERENCES

1. Basic Concepts of Intellectual Property Rights by Manthan D Janodia. Manipal University Press, 2015.

2. Intellectual Property Rights by SRS Rosedar. Lexis Nexis 2016.

3. Intellectual Property Rights in India by VK Ahuja. Lexis Nexis, 2015.

4. Law relating to Intellectual Property by BL Wadehra. Universal Law Publishing, 2016.

### PRM-003E: GENERAL MANAGEMENT PRINCIPLES

#### (15 hrs)

#### Scope

This course is designed to facilitate students to inculcate managerial skills. It includes major concepts of management.

1. Introduction to management concepts	3 hrs
2. Decision Making	3 hrs
3. Leadership and Motivation	4 hrs
4. Conflict Management	3 hrs
5. Ethical Issues related to Management	2 hrs

### REFERENCES

1. Organisational Behaviour by Stephen P. Robbins, Prentice – Hall, India

2. Management, A global Perspective by Heinz, Weihrich and Harold Koontz. Mc Graw Hill publishing company.

3. Management, tasks, responsibilities and practices by Drucker. Peter. F., Alfied Publisher Pvt. Ltd.

4. Principles and Practice of Management by L M Prasad, Sultan Chand & Sons.

#### PRM-004E: ENTREPRENEURSHIP DEVELOPMENT

#### (15 hrs)

#### Scope

This course is designed to impart knowledge and skills on entrepreneurship development.

1. Entrepreneur and Entrepreneurship	3 hrs
2. Entrepreneurial Development	3 hrs
3. Launching and Organizing an enterprise	3 hrs
4. Cost and Pricing	3 hrs
5. Project proposal development for start-up	3 hrs

#### REFERENCES

1. Hisrich, R.D. and Peters, M.P. (1995): Entrepreneurship – Starting, Developing and Managing a New Enterprise, Richard D., Inwin, INC, USA.

2. Meredith, G.G. etal (1982): Practice of Entrepreneurship, ILO, Geneva.

3. Entrepreneurship Management by Vasant Desai. Himalaya Publishing House, 2011.

#### MPHARM – CHOICE BASED MULTIDISCIPLINARY COURSE

- MU-001E: Certificate Course in Bioinformatics
- MU-002E: Project Management
- MU-003E: Certificate Course in Bioethics
- MU-004E: Academic Research and Writing
- MU-005E: Certificate Course in Biosecurity

(As prescribed from time to time)