R20 Curriculum and Syllabus Master in Pharmaceutical Technology (M. Pharm- Pharmaceutics)





R21 Curriculum and Syllabus, M.Pharm.Pharmaceutics

			SEM	ESTER-	#					
Sl. No.	Type	Course No.		Course]	Name		\mathbf{L}	Т	Ρ	Credits
THEOI	RY									
PRACT	FICAL									
EMBEI	DDED($\Gamma HEORY + 1$	PRACTICA	AL)						
BLENI	DED(MO	OOC + INTE	RNAL AS	SESSME	NT)					
SESSIC	NAL(C	NLY INTER	NAL EVA	LUATIO	N)		-			
MAND	ATORY	NON-CGPA	A COURSE							
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1										

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				SEMESTER-1					
Sl. No.	Type	Course No.		Course Name	e	\mathbf{L}	T	Р	Credits
THEOR	Ŷ								
1		MPH101T	Moder Technie	n Pharmaceutical ques - Theory	Analytical	4	0	0	4
2		MPH102T	Drug I	Delivery System - Th	leory	4	0	0	4
3		MPH103T	Moder	n Pharmaceutics - T	heory	4	0	0	4
4		MPH104T	Regula	tory Affair - Theory		4	0	0	4
PRACT	ICAL								
5		MPH105P	Pharm	aceutics I - Practica	1	0	0	12	6
SESSIO	NAL								
6		MPH106S	Semina	r / Assignment		0	7	0	4
MANDA	ATORY	COURSE							
7		MSD1861	Semina	r and Group Discus	sion	0	0	0	1
8		MSD1862	Skill X Course	and Other Activitis)	les (MOOCs	0	0	0	1
TOTAL						16	7	12	26



			SEMESTER-2				
Sl. No.	Type	Course No.	Course Name	\mathbf{L}	\mathbf{T}	Р	Credits
THEOR	Y						
1		MPH201T	Molecular Pharmaceutics (Nano Tech and TDDS) - Theory	4	0	0	4
2		MPH202T	Advanced Biopharmaceutics & Phar- macokinetics - Theory	4	0	0	4
3		MPH203T	Computer Aided Drug Delivery System - Theory	4	0	0	4
4		MPH204T	Cosmetic and cosmeceuticals - Theory	4	0	0	4
PRACT	ICAL						
5		MPH205P	Pharmaceutics II - Practical	0	0	12	6
SESSIO	NAL						
6		MPH206S	Seminar / Assignment	0	0	7	4
MAND	ATORY	CREDIT C	OURSE				
7	MC	MSD2861	Seminar and Group Discussion	0	0	0	1
8	MC	MSD2862	Skill X and Other activities (MOOCs courses)	0	0	0	1
TOTAL				16	7	12	26

			SEMESTER-3				
Sl. No.	Type	Course No.	Course Name	\mathbf{L}	Т	Р	Credits
THEOR	łY	•					
1		MRM301T	Research Methodology and Biostatis- tics – Theory [*]	4	0	0	4
SESSIO	NAL					Y	
2		MRM302S	Journal Club	0	1	0	1
3		MRM303S	Discussion / Presentation (Proposal Presentation)	0	2	0	2
4		MRM304S	Research Work	0	0	28	14
MAND	ATORY	CREDIT C	OURSE				
5	MC	MSD3861	Seminar and Group Discussion	0	0	0	1
6	MC	MSD3862	Skill X and Other activities (MOOCs courses)	0	0	0	1
TOTAL				4	3	28	21



				SEN	ЛEST	ER-	4						
Sl. No.	Type	Course No.			Cou	rse I	Name			\mathbf{L}	\mathbf{T}	Ρ	Credits
SESSIO	NAL												
1		MRM401S	Jou	rnal Cl	ub					0	1	0	1
2		MRM402S	Res	earch V	Nork					0	0	31	16
3		MRM403S			/ Pr	esent	ation	(Final	Pre-	0	3	0	3
MAND	ATORY	CREDIT C	OUR	SE									
4	MC	MSD4861	Sem	inar a	nd Gro	oup I	Discuss	ion		0	0	0	1
5	MC	MSD4862	Skill cour		id Oth	ner a	ctivitie	s (MC	OCs	0	0	0	1
TOTAL										0	4	31	20

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Catagony	Credit Allocation	Credit Allocation
Category	As Per PCI	As per University
Semester I	26	28
Semester II	26	28
Semester III	21	23
Semester IV	20	22
Total	98	106
Credit Distribution Details		
Professional Core Courses	48	48
Journal Club	2	2
Discussion and Presentation	5	5
Research Work, Project work and internship in indus-	30	30
try or elsewhere		50
Mandatory Courses [Seminar, Attending Conference,		
Scientific Presentations and Other Scholarly Activi-	13	16
ties, Assignment and Skill X		
Total	98	101

Credit Distribution Ratio:



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Credit Distribution in details:

A. Pr	ofessional Cor	e Courses (PC)							
Sl. No.	Paper Code	Theory & Practical				t Week	Credit Points		
			L	Т	Р	Total			
1	MPH101T	Modern Pharmaceutical Ana- lytical Techniques	4	0	0	4	4		
2	MPH102T	Drug Delivery System	4	0	0	4	4		
3	MPH103T	Modern Pharmaceutics	4	0	0	4	4		
4	MPH104T	Regulatory Affair	4	0	0	4	4		
5	MPH105P	Pharmaceutics Practical I	0	0	12	6	6		
6	MPH201T	Molecular Pharmaceutics (Nano Tech and TDDS)	4	0	0	4	4		
7	MPH202T	Advanced Biopharmaceutics & Pharmacokinetics	4	0	0	4	4		
8	MPH203T	Computer Aided Drug Deliv- ery System	4	0	0	4	4		
9	MPH204T	Cosmetic and cosmeceuticals	4	0	0	4	4		
10	MPH205P	Pharmaceutics Practical II	0	0	12	6	6		
11	MRM301T	Research Methodology and Biostatistics	4	0	0	4	4		
		Total Credit:					48		

Sl. No.	Paper Code	Theory	Contact Hours/Week			Credit Points		
		V III		L	Т	Р	Total	
1	MRM302S	Journal Club		0	1	0	1	1
2	MRM401S	Journal Club		0	1	0	1	1
		Total Credit:						2

C. Dis Sl. No.	scussion and F Paper Code		1	Con Hou		t Week	Credit Points
			L	Т	Р	Total	
1	MRM303S	Discussion/Presentation (Proposal Presentation)	0	2	0	2	2
2	MRM403S	Discussion/Presentation (Fi- nal Presentation)	0	3	0	3	3
		Total Credit:					5



D. Re	search Work,	Project work and	d inter	nship) in	inc	lust	ry or e	lsewhere (PW)			
Sl. No.	Paper Code	Practical			Contact Hours/Week							Credit Points
					L	Т	Р	Total				
1	MRM304S	Research Work			0	0	28	14	14			
1	MRM402S	Research Work			0	0	31	16	16			
		Total Credit:							30			

	-	rses [Attending Conference			ntif	ic Pres	sentati	ions and
Other Sl.	· Scholarly Act	tivities and SkillX Seminar]	<u>`</u>	lC) Con	tac	+		
No.	Paper Code	Theory				Week	Cred	it Points
			L	Т	Р	Total		
1	MPH106S	Seminar/Assignments	0	7	0	4		4
2	MSD 1861	Seminar and Group Discussion	0	0	0	1		1
3	MSD 1862	Skill X and other activities (like MOOCs Courses)	0	0	0	1		1
4	MPH206S	Seminar/Assignments	0	7	0	4		4
5	MSD 2861	Seminar and Group Discussion	0	0	0	1		1
6	MSD 2862	Skill X and other activities (like MOOCs Courses)	0	0	0	1		1
7	MSD 3861	Seminar and Group Discussion	0	0	0	1		1
8	MSD 3862	Skill X and other activities (like MOOCs Courses)	0	0	0	1		1
9	MSD 4861	Seminar and Group Discussion	0	0	0	1		1
10	MSD 4862	Skill X and other activities (like MOOCs Courses)	0	0	0	1		1
		Total Credit:						16



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				SEMESTER-1				
Sl. No.	Type	Course No.		Course Name	L	Т	Ρ	Credits
THEOF	RY							
1		MPH101T		Modern Pharmaceutical Analytical Techniques - Theory	4	0	0	4
2		MPH102T]	Drug Delivery System - Theory	4	0	0	4
3		MPH103T	1	Modern Pharmaceutics - Theory	4	0	0	4
4		MPH104T]	Regulatory Affair - Theory	4	0	0	4
PRACI	TICAL		·					
5		MPH105P		Pharmaceutics I - Practical	0	0	12	6
SESSIO	NAL							
6		MPH106S		Seminar / Assignment	0	7	0	4
MAND	ATORY	COURSE						
7		MSD1861	,	Seminar and Group Discussion	0	0	0	1
8		MSD1862		Skill X and Other Activities (MOOCs	0	0	0	1
		110101002	(Courses)	Ū		0	
TOTAL					16	7	12	26
				EKSII				



Course Code	MPH101T
Course Title	MODERN PHARMACEUTICAL ANALYTICAL TECHNIQUES
Category	
LTP & Credits	L T P Credits
	4 0 0 4
Total Contact Hours	60
Pre-requisites	None

This subject deals with various advanced analytical instrumental techniques for identification, characterization and quantification of drugs. Instruments dealt are NMR, Mass spectrometer, IR, HPLC, GC etc.

Course Objective:

Upon completion of the course the students shall be able to:

- 1. Analyze Chemicals and Excipients
- 2. Analyze various drugs in single and combination dosage forms
- 3. Execute theoretical and practical skills of the instruments

Course Content:

UNIT I:

[11L]

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

IR spectroscopy: 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

Spectroflourimetry: Theory of Fluorescence, Factors affecting fluorescence, Quenchers, Instrumentation and Applications of fluorescence spectrophotometer.

Flame emission spectroscopy and Atomic absorption spectroscopy: Principle, Instrumentation, Interferences and Applications.

UNIT II:

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

UNIT III:

[11L]

[11L]



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

UNIT IV:

Chromatography: Principle, apparatus, instrumentation, chromatographic parameters, factors affecting resolution and applications of the following: a) Paper chromatography b) Thin Layer chromatography c) Ion exchange chromatography d) Column chromatography e) Gas chromatography f) High Performance Liquid chromatography g) Affinity chromatography

UNIT V:

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

b. X ray Crystallography: Production of X rays, Different X ray diffraction methods, Bragg's law, Rotating crystal technique, X ray powder technique, Types of crystals and applications of X- ray diffraction.

UNIT VI:

RIA (Radio immunoassay), ELISA, Bioluminescence assays.

Recommended Books (Latest Editions):

- 1. Spectrometric Identification of Organic compounds Robert M Silverstein, Sixth edition, John Wiley & Sons, 2004.
- 2. Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5th edition, Eastern press, Bangalore, 1998.
 - 3. Instrumental methods of analysis Willards, 7th edition, CBS publishers.
- 4. Practical Pharmaceutical Chemistry Beckett and Stenlake, Vol II, 4th edition, CBS Publishers, New Delhi, 1997.
- 5. Organic Spectroscopy William Kemp, 3rd edition, ELBS, 1991.
- 6. Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi, 3rd Edition, CBS Publishers, New Delhi, 1997.
- 7. Pharmaceutical Analysis- Modern methods Part B J W Munson, Volume 11, Marcel Dekker Series

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[11L]

[5L]



CO	Progr	Program Outcome												
	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11			
MPH101T.1	3	-	2	3	1	-	-	2	-	-	3			
MPH101T.2	3	2	2	3	2	2	-	1	-	1	2			
MPH101T.3	2	3	2	3	1	2	1	1	1	1	2			
MPH101T.4	3	3	1	2	2	-	-	2	-	-	-			
MPH101T.5	1	3	1	-	2	2	3	3	3	3	1			





Course Code	M	PH1	021	ר				
Course Title	DI	RUC	d DI	ELIVERY SYSTEMS - Theory				
Category								
LTP & Credits	L T P Credits							
	4	0	0	4				
Total Contact Hours	60							
Pre-requisites	No	one						

This course is designed to impart knowledge on the area of advances in novel drug delivery systems.

Course Objective:

Upon completion of the course the students shall be able to:

- 1. The various approaches for development of novel drug delivery systems.
- 2. The criteria for selection of drugs and polymers for the development of delivering system
- **3.** The formulation and evaluation of Novel drug delivery systems

Course Content:

UNIT I:

Sustained Release (SR) and Controlled Release (CR) formulations:

Introduction basic concepts, advantages/ disadvantages, factors influencing, Physicochemical biological approaches for SR/CR formulation, Mechanism of Drug Delivery from SR/CR formulation. Polymers: introduction, definition, classification, properties and application Dosage Forms for Personalized Medicine: Introduction, Definition, Pharmacogenetics, Categories of Patients for Personalized Medicines: Customized drug delivery systems, Bioelectronic Medicines, 3D printing of pharmaceuticals, Telepharmacy.

UNIT II:

Rate Controlled Drug Delivery Systems:

Principles Fundamentals, Types, Activation; Modulated Drug Delivery Systems; Mechanically activated, pH activated, Enzyme activated, and Osmotic activated Drug Delivery Systems Feedback regulated Drug Delivery Systems; Principles Fundamentals.

UNIT III:

Gastro-Retentive Drug Delivery Systems:

Principle, concepts advantages and disadvantages, Modulation of GI transit time approaches to extend GI transit. Buccal Drug Delivery Systems: Principle of muco adhesion, advantages and disadvantages, Mechanism of drug permeation, Methods of formulation and its evaluations.

[10L]

[10L]

[10L]



[6L]

[10L]

[10L]

[6L]

UNIT IV:

Occular Drug Delivery Systems: Barriers of drug permeation, Methods to overcome barriers

UNIT V:

Transdermal Drug Delivery Systems: Structure of skin and barriers, Penetration enhancers, Transdermal Drug Delivery Systems, Formulation and evaluation.

UNIT VI:

Protein and Peptide Delivery: Barriers for protein delivery. Formulation and Evaluation of delivery systems of proteins and other macromolecules

UNIT VII:

Vaccine delivery systems: Vaccines, uptake of antigens, single shot vaccines, mucosal and transdermal delivery of vaccines

Recommended Books (Latest Editions

- 1. Y W. Chien, Novel Drug Delivery Systems, 2nd edition, revised and expanded
- Robinson, J. R., Lee V. H. L, Controlled Drug Delivery Systems, Marcel Dekker, Inc., New York, 1992.
- 3. Encyclopedia of controlled delivery, Editor- Edith Mathiowitz, Published by Wiley Interscience Publication, John Wiley and Sons, Inc, New York! Chichester/Weinheim.
- 4. N.K. Jain, Controlled and Novel Drug Delivery, CBS Publishers & Distributors, New Delhi, First edition 1997 (reprint in 2001).
- 5. S.P.Vyas and R.K.Khar, Controlled Drug Delivery concepts and advances, Vallabh Prakashan, New Delhi, First edition 2002

Journals

1. Indian Journal of Pharmaceutical Sciences (IPA)

2. Indian drugs (IDMA).

- 3. Journal of controlled release (Elsevier Sciences) desirable
- 4. Drug Development and Industrial Pharmacy (Marcel & Decker) desirable

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CO	Progr	Program Outcome												
	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11			
MPH102T.1	3	-	2	3	1	-	-	2	-	-	3			
MPH102T.2	3	2	2	1	1	3	-	1	-	1	2			
MPH102T.3	2	1	2	-	2	-	2	2	-	-	2			
MPH102T.4	3	3	2	2	1	3	-	-	-	1	1			
MPH102T.5	2	3	2	3	1	2	2	2	-	-	2			





Course Code	M	PH1	031						
Course Title	M	ODI	ERN	V PHARMACEUTICS - Theory					
Category									
LTP & Credits	L T P Credits								
	4	0	0	4					
Total Contact Hours	60								
Pre-requisites	No	one							

Course designed to impart advanced knowledge and skills required to learn various aspects and concepts at pharmaceutical industries

Course Objective:

Upon completion of the course the students shall be able to:

- 1. The elements of preformulation studies.
- 2. The Active Pharmaceutical Ingredients and Generic drug Product development
- 3. Industrial Management and GMP Considerations
- 4. Optimization Techniques Pilot Plant Scale Up Techniques
- 5. Stability Testing, sterilization process & packaging of dosage forms

Course Content:

UNIT I:

a. Preformation Concepts

Drug Excipient interactions - different methods, kinetics of stability, Stability testing. Theories of dispersion and pharmaceutical Dispersion (Emulsion and Suspension, SMEDDS) preparation and stability Large and small volume parental – physiological and formulation consideration, Manufacturing and evaluation.

b. Optimization techniques in Pharmaceutical Formulation

Concept and parameters of optimization, Optimization techniques in pharmaceutical formulation and processing. Statistical design, Response surface method, Contour designs, Factorial designs and application in formulation

UNIT II:

Validation: Introduction to Pharmaceutical Validation, Scope merits of Validation, Validation and calibration of Master plan, ICH WHO guidelines for calibration and validation of equipments, Validation of specific dosage form, Types of validation. Government regulation, Manufacturing Process Model, URS, DQ, IQ, OQ P.Q. of facilities

UNIT III:

[10L]

[20L]

[10L]



[12L]

[12L]

cGMP Industrial Management: Objectives and policies of current good manufacturing practices, layout of buildings, services, equipments and their maintenance Production management: Production organization, materials management, handling and transportation, inventory management and control, production and planning control, Sales forecasting, budget and cost control, industrial and personal relationship. Concept of Total Quality Management.

UNIT IV:

Compression and compaction: Physics of tablet compression, compression, consolidation, effect of friction, distribution of forces, compaction profiles. Solubility.

UNIT V:

Study of consolidation parameters: Diffusion parameters, Dissolution parameters and Pharmacokinetic parameters, Heckel plots, Similarity factors – f2 and f1, Higuchi and Peppas plot, Linearity Concept of significance, Standard deviation , Chi square test, students T-test , ANOVA test.

Recommended Books (Latest Editions

- 1. Theory and Practice of Industrial Pharmacy By Lachmann and Libermann
- 2. Pharmaceutical dosage forms: Tablets Vol. 1-3 by Leon Lachmann
- 3. Pharmaceutical Dosage forms: Disperse systems, Vol, 1-2; By Leon Lachmann
- 4. Pharmaceutical Dosage forms: Parenteral medications Vol. 1-2; By Leon Lachmann
- 5. Modern Pharmaceutics; By Gillbert and S. Banker
- 6. Remington's Pharmaceutical Sciences
- 7. Advances in Pharmaceutical Sciences Vol. 1-5; By H.S. Bean & A.H. Beckett
- 8. Physical Pharmacy; By Alfred martin
- 9. Bentley's Textbook of Pharmaceutics by Rawlins
- 10. Good manufacturing practices for Pharmaceuticals: A plan for total quality control, Second edition; By Sidney H. Willig
- 11. Quality Assurance Guide; By Organization of Pharmaceutical producers of India.
- 12. Drug formulation manual; By D.P.S. Kohli and D.H.Shah. Eastern publishers, New Delhi
- 13. How to practice GMPs; By P.P.Sharma. Vandhana Publications, Agra
- 14. Pharmaceutical Process Validation; By Fra. R. Berry and Robert A. Nash
- 15. Pharmaceutical Preformulations; By J.J. Wells
- 16. Applied production and operations management; By Evans, Anderson, Sweeney and Williams
- 17. Encyclopaedia of Pharmaceutical technology, Vol I III.



CO	Program Outcome												
	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11		
MPH103T.1	3	2	3	2	1	2	-	1	-	-	2		
MPH103T.2	1	2	2	-	-	2	3	2	-	-	2		
MPH103T.3	2	3	1	-	3	1	2	2	1	2	2		
MPH103T.4	2	3	2	3	1	2	1	-	-	-	1		
MPH103T.5	2	3	2	-	1	2	2	2	1	2	3		





Course Code	MPH104T								
Course Title	RI	EGU	JLA	TORY AFFAIR - Theory					
Category									
LTP & Credits	L T P Credits								
	4	0	0	4					
Total Contact Hours	60								
Pre-requisites	No	one							

Course designed to impart advanced knowledge and skills required to learn the concept of generic drug and their development, various regulatory filings in different countries, different phases of clinical trials and submitting regulatory documents: filing process of IND, NDA and ANDA.

Course Objective:

Upon completion of the course, it is expected that the students will be able to understand:

- 1. The Concepts of innovator and generic drugs, drug development process
- 2. The Regulatory guidance's and guidelines for filing and approval process
- 3. Preparation of Dossiers and their submission to regulatory agencies in different countries
- 4. Post approval regulatory requirements for actives and drug products
- 5. Submission of global documents in CTD/eCTD formats
- 6. Clinical trials requirements for approvals for conducting clinical trials
- 7. Pharmacovigilence and process of monitoring in clinical trials

Course Content:

UNIT I:

a. Documentation in Pharmaceutical industry:

Master formula record, DMF (Drug Master File), distribution records. Generic drugs product development Introduction, Hatch-Waxman act and amendments, CFR (CODE OF FEDERAL REGULATION) ,drug product performance, in-vitro, ANDA regulatory approval process, NDA approval process, BE and drug product assessment, in –vivo, scale up process approval changes, post marketing surveillance, outsourcing BA and BE to CRO.

b. Regulatory requirement for product approval:

API, biologics, novel, therapies obtaining NDA, ANDA for generic drugs ways and means of US registration for foreign drugs

UNIT II:

[12L]

[24L]



CMC, post approval regulatory affairs. Regulation for combination products and medical devices. CTD and ECTD format, industry and FDA liaison. ICH - Guidelines of ICH-Q, S E, M. Regulatory requirements of EU, MHRA, TGA and ROW countries.

UNIT III:

[12L]

Non clinical drug development: Global submission of IND, NDA, ANDA. Investigation of medicinal products dossier, dossier (IMPD) and investigator brochure (IB).

UNIT IV:

[12L]

Clinical trials:

Developing clinical trial protocols. Institutional review board/ independent ethics committee Formulation and working procedures informed Consent process and procedures. HIPAA- new, requirement to clinical study process, pharmacovigilance safety monitoring in clinical trials.

Recommended Books (Latest Editions):

- 1. "Generic Drug Product Development", Solid Oral Dosage forms, Leon Shargel and Isader Kaufer, Marcel Dekker series, Vol.143
- 2. "The Pharmaceutical Regulatory Process", Second Edition Edited by Ira R. Berry and Robert P.Martin, Drugs and the Pharmaceutical Sciences, Vol.185, Informa Health care Publishers.
- 3. "New Drug Approval Process: Accelerating Global Registrations" By Richard A Guarino, MD, 5th edition, Drugs and the Pharmaceutical Sciences, Vol.190.
- 4. "Guidebook for drug regulatory submissions", Sandy Weinberg. By John Wiley and Sons.Inc.
- 5. "FDA regulatory affairs: a guide for prescription drugs, medical devices, and biologics", edited By Douglas J. Pisano, David Mantus.
- 6. "Clinical Trials and Human Research: A Practical Guide to Regulatory" Compliance By Fay A.Rozovsky and Rodney K. Adams.

RSIT

- 7. www.ich.org/
- 8. www.fda.gov/
- 9. europa.eu/index_en.htm
- 10. https://www.tga.gov.au/tga-basics

CO	Program Outcome (PO)												
	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11		
MPH104T.1	2	3	2	1	2	1	-	1	-	1	1		
MPH104T.2	3	2	1	2	3	-	2	2	-	-	2		
MPH104T.3	1	3	1	-	2	1	3	3	3	3	1		
MPH104T.4	3	2	1	-	2	2	3	2	-	-	2		
MPH104T.5	2	3	2	-	1	2	2	1	2	3	1		



Course Code	M	PH1	05P								
Course Title	Ph	narm	naceu	tics Practical-I (Practical)							
Category											
LTP & Credits	L T P Credits										
	0	0	12	6							
Total Contact Hours	18	0									
Pre-requisites	No	one									

Course Objective:

Upon completion of the course the students shall be able to:

- **1.** Know the estimation of the active pharmaceutical ingredients in formulations by using different modern techniques.
- **2.** Gain the knowledge to apply the concepts of preformulation in the formulation development..
- **3.** Evaluate the effects of different critical parameters on quality of tablets such as disintegration, dissolution, drug release profiles etc.
- **4.** Understand the formulation development and evaluation methods of different novel drug delivery system.

Suggestive List of Experiments:

- 1. Analysis of pharmacopoeial compounds and their formulations by UV Vis spectrophotometer. [2 day(s)]
- 2. Simultaneous estimation of multi component containing formulations by UV spectrophotometry [2 day(s)]

3.	Experiments based on HPLC :	$[2 \mathrm{day}(\mathrm{s})]$
4.	Experiments based on Gas Chromatography :	[1 day(s)]
5.	Estimation of riboflavin/quinine sulphate by fluorimetry :	$[1 \mathrm{day}(\mathrm{s})]$
6.	Estimation of sodium/potassium by flame photometry :	$[1 \mathrm{day}(\mathrm{s})]$
7.	To perform In-vitro dissolution profile of CR/ SR marketed formulation :	$[2 \mathrm{day}(\mathrm{s})]$



8.	Formulation and evaluation of sustained release matrix tablets :	$[2 \mathrm{day}(\mathrm{s})]$
9.	Formulation and evaluation osmotically controlled DDS	$[2 \mathrm{day}(\mathrm{s})]$
10.	Preparation and evaluation of Floating DDS- hydro dynamically balanced $day(s)$]:	DDS [2
11.	Formulation and evaluation of Mucoadhesive tablets.	$[2 \mathrm{day}(\mathrm{s})]$
12.	Formulation and evaluation of transdermal patches.	$[2 \mathrm{day}(\mathrm{s})]$
13.	To carry out preformulation studies of tablets.	$[1 \mathrm{day}(\mathrm{s})]$
14.	To study the effect of compressional force on tablets disintegration time	$[2 \mathrm{day}(\mathrm{s})]$
15.	To study Micromeritic properties of powders and granulation.	$[1 \mathrm{day}(\mathrm{s})]$
16.	To study the effect of particle size on dissolution of a tablet.	$[2 \mathrm{day}(\mathrm{s})]$
17.	To study the effect of binders on dissolution of a tablet.	$[2 \mathrm{day}(\mathrm{s})]$
18	To plot Hockal plot, Higuchi and poppas plot and determine similarity fact	$ars \left[1 day(s) \right]$

18. To plot Heckal plot, Higuchi and peppas plot and determine similarity factors [1 day(s)] .

	Progr	Program Outcome												
CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11			
MPH105P.1	2	2	1	3	1	3	-	-	-	-	2			
MPH105P.2	2	3	1	3	-	2	-	-	-	-	2			
MPH105P.3	3	3	1	3	-	3	-	-	-	-	2			
MPH105P.4	2	3	1	3	2	3	-	-	-	-	2			



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			SEMESTER-2									
Sl. No.	Type	Course No.	Course Name	L	\mathbf{T}	Р	Credits					
THEOF	RY											
1		MPH201T	Molecular Pharmaceutics (Nano Tech and TDDS) - Theory	4	0	0	4					
2		MPH202T	Advanced Biopharmaceutics & Phar- macokinetics - Theory	4	0	0	4					
3		MPH203T	Computer Aided Drug Delivery System - Theory	4	0	0	4					
4		MPH204T	Cosmetic and cosmeceuticals - Theory	4	0	0	4					
PRACT	ICAL											
5		MPH205P	Pharmaceutics II - Practical	0	0	12	6					
SESSIO	NAL											
6		MPH206S	Seminar / Assignment	0	0	7	4					
MAND	ATORY	CREDIT C	DURSE									
7	MC	MSD2861	Seminar and Group Discussion	0	0	0	1					
8	MC	MSD2862	Skill X and Other activities (MOOCs courses)	0	0	0	1					
TOTAL				16	7	12	26					
						Y						



[12L]

[12L]

[12L]

[12L]

Course Code	M	MPH201T											
Course Title	M	olec	ular	Pharmaceutics (Nano Technology and Targeted DDS)-Theory									
Category													
LTP & Credits	L	L T P Credits											
	4	4 0 0 4											
Total Contact Hours	60	60											
Pre-requisites	No	one											

Scope:

This course is designed to impart knowledge on the area of advances in novel drug delivery systems.

Course Objective:

Upon completion of the course the students shall be able to understand:

- 1. The various approaches for development of novel drug delivery systems.
- 2. The criteria for selection of drugs and polymers for the development of NTDS.
- 3. The formulation and evaluation of novel drug delivery systems.

Course Content:

UNIT I:

Targeted Drug Delivery Systems:

Concepts, Events and biological process involved in drug targeting. Tumor targeting and Brain specific delivery.

UNIT II:	[12L]
Targeting Methods: introduction preparation and evaluation.	Nano Particles &
Liposomes: Types, preparation and evaluation	

UNIT III:

Micro Capsules /Micro Spheres:

Types, preparation and evaluation, Monoclonal Antibodies; preparation and application, preparation and application of Niosomes, Aquasomes, Phytosomes, Electrosomes

UNIT IV:

Pulmonary Drug Delivery Systems:

Aerosols, propellents, Containers Types, preparation and evaluation, Intra Nasal Route Delivery systems; Types, preparation and evaluation.

UNIT V:	UNIT	V:
---------	------	----



Nucleic acid based therapeutic delivery system:

Gene therapy, introduction (ex-vivo & in-vivo gene therapy). Potential target diseases for gene therapy (inherited disorder and cancer). Gene expression systems (viral and nonviral gene transfer). Liposomal gene delivery systems. Biodistribution and Pharmacokinetics. knowledge of therapeutic antisense molecules and aptamers as drugs of future.

Recommended Books (Latest Editions):

- 1. Y W. Chien, Novel Drug Delivery Systems, 2nd edition, revised and expanded, Marcel Dekker, Inc., New York, 1992.
- 2. S.P.Vyas and R.K.Khar, Controlled Drug Delivery: concepts and advances, Vallabh Prakashan, New Delhi, First edition 2002.
- 3. N.K. Jain, Controlled and Novel Drug Delivery, CBS Publishers & Distributors, New Delhi, First edition 1997 (reprint in 2001).

CO		Progr	am Ou	itcome								
		PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11
MPH201	T.1	3	3	1	-	-	2	-	1	-	2	3
MPH201	T.2	2	3	2	2	1	3	-	1	-	1	2
MPH201	T.3	3	2	1	2	-	2	-	2	-	-	2
MPH201	T.4	2	3	2	1	1	2	1	1	-	2	2
MPH201	T.5	3	3	2	2	1	3	-	-	/ -	1	1





Course Code	М	PH2	2027	۲
Course Title	A	lvan	lced	Biopharmaceutics and Pharmacokinetics
Category				
LTP & Credits	L	Т	Р	Credits
	4	0	0	4
Total Contact Hours	60			
Pre-requisites	No	one		

This course is designed to impart knowledge andskills necessary for dose calculations, dose adjustments and to apply biopharmaceutics theories in practical problem solving. Basic theoretical discussions of the principles of biopharmaceutics and pharmacokinetics are provided to help the students' to clarify the concepts.

Course Objective:

Upon completion of the course the students shall be able to understand:

- 1. The basic concepts in biopharmaceutics and pharmacokinetics.
- 2. The use raw data and derive the pharmacokinetic models and parameters the best describe the process of drug absorption, distribution, metabolism and elimination.
- **3.** The critical evaluation of biopharmaceutic studies involving drug product equivalency.
- 4. The design and evaluation of dosage regimens of the drugs using pharmacokinetic and biopharmaceutic parameters.
- 5. The potential clinical pharmacokinetic problems and application of basics of pharmacokinetic.

Course Content:

UNIT I:

[12L]

Drug Absorption from the Gastrointestinal Tract:

Gastrointestinal tract, mechanism of drug absorption, Factors affecting drug absorption, pH-partition theory of drug absorption. Formulation and physicochemical factors: Dissolution rate, Dissolution process, Noyes–Whitney equation and drug dissolution, Factors affecting the dissolution rate.

Gastrointestinal absorption: role of the dosage form: Solution (elixir, syrup and solution) as a dosage form, Suspension as a dosage form, Capsule as a dosage form, Tablet as a dosage form, Dissolution methods, Formulation and processing factors, Correlation of in vivo data with in vitro dissolution data. Transport model: Permeability Solubility Charge State and the pH Partition Hypothesis, Properties of the Gastrointestinal Tract (GIT), pH Microclimate Intracellular pH Environment, Tight Junction Complex.

UNIT II:

[12L]

[12L]

[12L]

[12L]

Biopharmaceutic considerations in drug product design and In Vitro Drug Product Performance:

Introduction, biopharmaceutic factors affecting drug bioavailability, rate-limiting steps in drug absorption, physicochemical nature of the drug formulation factors affecting drug product performance, in vitro: dissolution and drug release testing, compendial methods of dissolution, alternative methods of dissolution testing, meeting dissolution requirements, problems of variable control in dissolution testing performance of drug products. In vitro–in vivo correlation, dissolution profile comparisons, drug product stability, considerations inthedesign of a drug product.

UNIT III:

Pharmacokinetics: Basic considerations, pharmacokinetic models, **compartment modeling:** one compartment model- IV bolus, IV infusion, extra-vascular. **Multi compartment model:** two compartment - model in brief, **non-linear pharmacokinetics:** cause of non-linearity, Michaelis –Menten equation, estimation of kmax and vmax. **Drug interactions:** introduction, the effect of protein- binding interactions, the effect of tissue-binding interactions, cytochrome p450-based drug interactions, drug interactions linked to transporters.

UNIT IV:

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, evaluation of the data, bioequivalence example, study submission and drug review process. biopharmaceutics classification system, methods. Permeability: In-vitro, in-situ and In-vivo methods. Generic biologics (biosimilar drug products), clinical significance of bioequivalence studies, special concerns in bioavailability and bioequivalence studies, generic substitution.

UNIT V:

Application of Pharmacokinetics: Modified-Release Drug Products, Targeted Drug Delivery Systems and Biotechnological Products. Introduction to Pharmacokinetics and pharmacodynamic, drug interactions. Pharmacokinetics and pharmacodynamics of biotechnology drugs. Introduction, Proteins and peptides, Monoclonal antibodies, Oligonucleotides, Vaccines (immunotherapy), Genetherapies.

Recommended Books (Latest Editions):

- 1. Biopharmaceutics and Clinical Pharmacokinetics by Milo Gibaldi, 4th edition, Philadelphia, Lea and Febiger, 1991
- 2. Biopharmaceutics and Pharmacokinetics, A. Treatise, D. M. Brahmankar and Sunil B Jaiswal, Vallab Prakashan, Pitampura, Delhi
- 3. Applied Biopharmaceutics and Pharmacokinetics by Shargel. Land YuABC, 2nd edition, Connecticut Appleton Century Crofts, 1985



- 4. Textbook of Biopharmaceutics and Pharmacokinetics, Dr. Shobha Rani R. Hiremath, Prism Book
- 5. Pharmacokinetics by Milo Gibaldi and D. Perrier, 2nd edition, Marcel Dekker Inc., New York, 1982
- 6. Current Concepts in Pharmaceutical Sciences: Biopharmaceutics, Swarbrick. J,Leaand Febiger, Philadelphia, 1970
- 7. Clinical Pharmacokinetics, Concepts and Applications 3rd edition by Malcolm Rowland and Thom N. Tozer, Lea and Febiger, Philadelphia, 1995
- 8. Dissolution, Bioavailability and Bioequivalence, Abdou. H.M, Mack Publishing Company, Pennsylvania 1989
- 9. Biopharmaceutics and Clinical Pharmacokinetics, An Introduction, 4th edition, revised and expanded by Robert. E. Notari, Marcel Dekker Inc, New York and Basel, 1987.
- 10. Biopharmaceutics and Relevant Pharmacokinetics by John. G Wagner and M.Pemarowski, 1st edition, Drug Intelligence Publications, Hamilton, Illinois, 1971.
- 11. Encyclopedia of Pharmaceutical Technology, Vol 13, James Swarbrick, James. G.Boylan, Marcel Dekker Inc, New York, 1996.
- 12. Basic Pharmaceutics, 1st edition, Sunil S Jambhekarand Philip J Breen, Pharmaceutical Press, RPS Publishing, 2009.
- 13. Absorption and Drug Development- Solubility, Permeability, and Charge State, Alex Avdeef, John Wiley & Sons,Inc,2003.

CO	Progr	Program Outcome											
	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11		
MPH202T.1	3	2	2	_	_	_	1	-	-	-	3		
MPH202T.2	3	3	3	3	-	1	-	-	-	-	3		
MPH202T.3	3	2	3	3	-	2	1	-	2	-	3		
MPH202T.4	3	3	3	3	-	2	1	-	2	-	3		
MPH202T.5	3	3	3	3	-	3	1	-	2	-	3		



Course Code	M	MPH203T							
Course Title	Co	omp	uter	Aided Drug Delivery System					
Category									
LTP & Credits	L T P Credits								
	4	0	0	4					
Total Contact Hours	60								
Pre-requisites	No	one							

This course is designed to impart knowledge and skills necessary for computer applications in pharmaceutical research and development who want to understand the application of computers across the entire drug research and development process. Basic theoretical discussions of the principles of more integrated and coherent use of computerized information (informatics) in the drug development process are provided to help the students to clarify the concepts.

Course Objective:

Upon completion of the course, it is expected that the students will be able to understand:

- 1. History of Computers in Pharmaceutical Research and Development
- 2. Computational Modeling of Drug Disposition
- **3.** Computers in Preclinical Development
- 4. Optimization Techniques in Pharmaceutical Formulation
- 5. Computers in Market Analysis
- 6. Computers in Clinical Development
- 7. Artificial Intelligence (AI) and Robotics
- 8. Computational fluid dynamics(CFD)

Course Content:

UNIT I:

[12L]

a. Computers in Pharmaceutical Research and Development: A General Overview: History of Computers in Pharmaceutical Research and Development. Statistical modeling in Pharmaceutical research and development: Descriptive versus Mechanistic Modeling, Statistical Parameters, Estimation, Confidence Regions, Nonlinearity at the Optimum, Sensitivity Analysis, Optimal Design, Population Modeling

b. Quality-by-Design In Pharmaceutical Development: Introduction, ICH Q8 guideline, Regulatory and industry views on QbD, Scientifically based QbD - examples of application.

UNIT II:

[12L]



Computational Modeling Of Drug Disposition:

Introduction, Modeling Techniques: Drug Absorption, Solubility, Intestinal Permeation, Drug Distribution ,Drug Excretion, Active Transport; P-gp, BCRP, Nucleoside Transporters, hPEPT1, ASBT, OCT, OATP, BBB-Choline Transporter.

UNIT III:

Computer-aided formulation development:

Concept of optimization, Optimization parameters, Factorial design, Optimization technology Screening design. Computers in Pharmaceutical Formulation: Development of pharmaceutical emulsions, microemulsion drug carriers Legal Protection of Innovative Uses of Computers in RD, The Ethics of Computing in Pharmaceutical Research, Computers in Market analysis.

UNIT IV:

a. Computer-aided biopharmaceutical characterization: Gastrointestinal absorption simulation. Introduction, Theoretical background, Model construction, Parameter sensitivity analysis, Virtual trial, Fed vs. fasted state, In vitro dissolution and in vitro in vivo correlation, Biowaiver considerations.

b. Computer Simulations in Pharmacokinetics and Pharmacodynamics: Introduction, Computer Simulation: Whole Organism, Isolated Tissues, Organs, Cell, Proteins and Genes.

c. Computers in Clinical Development: Clinical Data Collection and Management, Regulation of Computer Systems.

UNIT V:

Artificial Intelligence (AI), Robotics and Computational fluid dynamics:

General overview, Pharmaceutical Automation, Pharmaceutical applications, Advantages and Disadvantages. Current Challenges and Future Directions.

Recommended Books (Latest Editions):

- 1. "Computer Applications in Pharmaceutical Research and Development", Sean Ekins, 2006, John Wiley & Sons.
- 2. "Computer-Aided Applications in Pharmaceutical Technology", 1st Edition, Jelena Djuris, Woodhead Publishing
- 3. "Encyclopedia of Pharmaceutical Technology", Vol 13, James Swarbrick, James. G. Boylan, Marcel Dekker Inc, New York, 1996.

[12L]

[12L]

[12L]



CO	Progr	Program Outcome (PO)												
	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11			
MPH203T.1	3	-	-	1	-	1	1	2	1	-	3			
MPH203T.2	2	1	3	2	2	1	1	2	-	-	2			
MPH203T.3	2	3	3	2	3	3	2	3	2	-	1			
MPH203T.4	1	2	2	2	3	3	2	3	2	-	1			
MPH203T.5	1	1	2	1	1	2	-	2	1	-	2			
MPH203T.6	1	2	3	3	1	3	2	3	1	-	1			





Course Code	MPT204T							
Course Title	Cosmetics and Cosmeceuticals							
Category								
LTP & Credits	L T P Credits							
	4	0	0	4				
Total Contact Hours	60							
Pre-requisites	No	one						

This course is designed to impart knowledge and skills necessary forthefundamental need for cosmetic and cosmeceutical products.

Course Objective:

Upon completion of the course the students shall be able to:

- 1. Key ingredients used in cosmetics and cosmeceuticals.
- **2.** Key building blocks for various formulations.
- 3. Current technologies in the market
- 4. Various key ingredients and basic science to develop cosmetics and cosmeceuticals
- 5. Scientific knowledge to develop cosmetics and cosmeceuticals with desired Safety, stability, and efficacy

Course Content:

UNIT I:

[12L]

Cosmetics – Regulatory

Definition of cosmetic products as per Indian regulation. Indian regulatory requirements for labeling of cosmetics Regulatory provisions relating to import of cosmetics., Misbranded and spurious cosmetics. Regulatory provisions relating to manufacture of cosmetics – Conditions for obtaining license, prohibition of manufacture and sale of certain cosmetics, loan license, offences and penalties.

UNIT II:

[12L]

Cosmetics - Biological aspects

Structure of skin relating to problems like dry skin, acne, pigmentation, prickly heat, wrinkles and body odor. Structure of hair and hair growth cycle. Common problems associated with oral cavity. Cleansing and care needs for face, eye lids, lips, hands, feet, nail, scalp, neck, body and under-arm.

UNIT III:

[12L]

[12L]

[12L]

Formulation Building blocks

Building blocks for different product formulations of cosmetics/cosmeceuticals. Surfactants – Classification and application. Emollients, rheological additives: classification and application. Antimicrobial used as preservatives, their merits and demerits. Factors affecting microbial preservative efficacy. Building blocks for formulation of a moisturizing cream, vanishing cream, cold cream, shampoo and toothpaste. Soaps and syndetbars. Perfumes; Classification of perfumes. Perfume ingredients listed as allergens in EU regulation. Controversial ingredients: Parabens, formaldehyde liberators, dioxane.

UNIT IV:

Design of cosmeceutical products Sun protection, sunscreens classification and regulatory aspects. Addressing dry skin, acne, sun-protection, pigmentation, prickly heat, wrinkles, body odor., dandruff, dental cavities, bleeding gums, mouth odor and sensitive teeth through cosmeceutical formulations.

UNIT V:

Herbal Cosmetics Herbal ingredients used in Hair care, skin care and oral care. Review of guidelines for herbal cosmetics by private bodies like cosmos with respect to preservatives, emollients, foaming agents, emulsifiers and rheology modifiers. Challenges in formulating herbal cosmetics

Recommended Books (Latest Editions):

- 1. Harry's Cosmeticology. 8th edition"
- 2. Poucher'sperfumecosmeticsandSoaps,10th edition"
- 3. Cosmetics Formulation, Manufacture and quality control, PP.Sharma,4th edition".
- 4. Handbook of cosmetic science and Technology A.O.Barel, M.Paye and H.I. Maibach. 3 rd edition"
- 5. Cosmetic and Toiletries recent suppliers catalogue"
- 6. CTFA directory'

CO	Progr	Program Outcome												
	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11			
MPH204T.1	3	-	3	-	-	-	-	-	-	-	-			
MPH204T.2	3	-	-	-	-	-	-	-	-	-	3			
MPH204T.3	-	-	-	-	-	-	3	-	-	-	3			
MPH204T.4	-	-	-	-	-	-	3	-	-	-	-			
MPH204T.5	3	-	-	-	-	-	-	-	3	-	-			



Course Code	MPH205P									
Course Title	Pharmaceutics Practical-II (Practical)									
Category										
LTP & Credits	L T P Credits									
	0	0	12	6						
Total Contact Hours	180									
Pre-requisites	No	one								

Course Objective:

Upon completion of the course the students shall be able to:

- 1. Understand the formulation and evaluation of novel drug delivery systems
- **2.** Understand preparation of various cosmetics and cosmeceuticals and know about the use of various key ingredients to develop cosmetics and cosmeceuticals.
- **3.** Understand the use computers in Preclinical and Clinical Development.
- 4. Understand various optimization techniques in Pharmaceutical Formulation.
- **3.** Understand basic concepts in biopharmaceutics and pharmacokinetics process.
- 4. Understand the use raw data and derive the pharmacokinetic models and parameters the best describes the process of drug absorption, distribution, metabolism and elimination

Suggestive List of Experiments:

:

 1.
 To study the effect of temperature change , non solvent addition, incompatible polymer addition in microcapsules preparation
 [2 day(s)]

 :
 :
 :

2.	Preparation and evaluation of Alginate beads	$[2 \mathrm{day}(\mathrm{s})]$
3.	: Formulation and evaluation of gelatin /albumin microspheres :	$[2 \mathrm{day}(\mathrm{s})]$
4.	Formulation and evaluation of liposomes/niosomes	$[2 \mathrm{day}(\mathrm{s})]$
	÷	
5.	Formulation and evaluation of spherules	$[2 \mathrm{day}(\mathrm{s})]$
	÷	

- 6. Improvement of dissolution characteristics of slightly soluble drug by Solid dispersion technique. [2 day(s)]
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7.	Comparison of dissolution of two different marketed products /brands :	$[1 \mathrm{day}(\mathrm{s})]$
8.	Protein binding studies of a highly protein bound drug poorly protein bou day(s)] :	und drug [1
9.	Bioavailability studies of Paracetamol in animals.	$[2 \mathrm{day}(\mathrm{s})]$
10.	Pharmacokinetic and IVIVC data analysis by Winnoline R software \vdots	$[1 \mathrm{day}(\mathrm{s})]$
11.	In vitro cell studies for permeability and metabolism	$[2 \mathrm{day}(\mathrm{s})]$
12.	DoE Using Design Expert® Software	$[1 \mathrm{day}(\mathrm{s})]$
13.	Formulation data analysis Using Design $\operatorname{Expert}_{\widehat{\mathbb{R}}}$ Software	$[1 \mathrm{day}(\mathrm{s})]$
14.	Quality-by-Design in Pharmaceutical Development :	$[1 \mathrm{day}(\mathrm{s})]$
15.	Computer Simulations in Pharmacokinetics and Pharmacodynamics	$[1 \mathrm{day}(\mathrm{s})]$
16.	Computational Modeling Of Drug Disposition	$[1 \mathrm{day}(\mathrm{s})]$
17.	To develop Clinical Data Collection manual	$[2 \mathrm{day}(\mathrm{s})]$
18.	To carry out Sensitivity Analysis, and Population Modeling.	[1 day(s)]
19.	Development and evaluation of Creams	$[1 \mathrm{day}(\mathrm{s})]$
20.	Development and evaluation of Shampoo and Toothpaste base :	$[1 \mathrm{day}(\mathrm{s})]$
21.	To incorporate herbal and chemical actives to develop products	$[1 \mathrm{day}(\mathrm{s})]$



	Program Outcome											
CO	PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11	
MPH205P.1	3	2	2	1	2	3	1	2	-	1	2	
MPH205P.2	2	3	3	1	2	3	1	2	1	-	2	
MPH205P.3	1	2	3	2	2	3	1	2	1	-	2	
MPH205P.4	2	3	3	2	2	3	1	2	-	-	2	
MPH205P.5	2	1	1	2	2	-	1	1	-	-	3	
MPH205P.6	2	2	3	2	3	3	1	2	2	-	1	





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				SE	MEST						
Sl. No.	Type	Course No.			Cou	rse Name		\mathbf{L}	Т	Ρ	Credits
THEOF	RY										
1		MRM301T		earch – The		dology and	Biostatis-	4	0	0	4
SESSIO	NAL										
2		MRM302S	Jou	rnal C	lub			0	1	0	1
3		MRM303S		cussion sentati	'	resentation	(Proposal	0	2	0	2
4		MRM304S	Res	earch	Work			0	0	28	14
MAND	ATORY	CREDIT C	OUR	SE				1	1		I
5	MC	MSD3861	Sem	ninar a	and Gro	oup Discussi	on	0	0	0	1
6	MC	MSD3862		Skill X and Other activities (MOOCs courses)						0	1
TOTAL								4	3	28	21
UNIVERSITY											



Course Code	MRM301T										
Course Title	RESEARCH METHODOLOGY AND BIOSTATISTICS										
Category											
LTP & Credits	L T P Credits										
	4	0	0	4							
Total Contact Hours	60										
Pre-requisites	No	one									

The course describes the basic methodology to carry out the dissertation work.

Course Objective:

After completion of course student is able to know:

- 1. Evaluate the various statistical techniques to solve statistical problems
- 2. Evaluate research methodology
- 3. Analyze statistical techniques in solving the problems
- 4. Analyze the operation of M.S. Excel and other Microsoft applications
- 5. Analyze the operation of SPSS and other statistical software

Course Content:

UNIT I:

General Research Methodology:

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

UNIT II:

Biostatistics:

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

UNIT III:

[12L]

[12L]

[12L]

Medical Research:

History, values in medical ethics, autonomy, beneficence, non-maleficence, 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.

UNIT IV:

[10L]

[10L]

CPCSEA guidelines for laboratory animal facility:

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

UNIT V:

Declaration of Helsinki:

History, introduction, basic principles for all medical research, and additional principles for medical research combined with medical care.

CO-PO Mapping:

CO		Progr	am Ou	itcome								
		PO1	PO2	PO3	PO4	PO5	PO6	PO7	PO8	PO9	PO10	PO11
MRM30	1T.1	1	2	3	-	-	2	-	-	1	-	3
MRM30	1T.2	-	3	3	-	-	2	-	-	/-	-	2
MRM30	1T.3	-	2	3	3	-	2	-	-	/ -	-	1
MRM30	1T.4	-	2	3	3	-	2	-	1	2	-	1
MRM30	1T.5	-	2	3	3	-	2		1	2	-	1

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				SF	MEST	FR 4						
Sl. No.	Type	Course No.		5E.		rse Name			L	T	P	Credits
SESSIO												
1		MRM401S	Jou	rnal C	lub				0	1	0	1
2		MRM402S	Res	earch	Work				0	0	31	16
3		MRM403S	sent	ation)		esentation	(Final	Pre-	0	3	0	3
MAND	ATORY	CREDIT C	OUR	SE								
4	MC	MSD4861	Sem	Seminar and Group Discussion						0	0	1
5	MC	MSD4862		Skill X and Other activities (MOOCs ourses)							0	1
TOTAL									0	4	31	20

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