

VIVEKANAND EDUCATION SOCIETY'S COLLEGE OF PHARMACY

Hashu Advani Memorial Complex, Behind Collector Colony, Chembur (E), Mumbai – 400 074

Sindhi Linguistic Minority, Approved by AICTE, DTE, Pharmacy Council of India & Govt. of Maharashtra, Affiliated to University of Mumbai

B. Pharm Programme is accredited by NBA, New Delhi from 2016-17 to 2021-22

2.6.1

Teachers and students are aware of the stated Programme and Course outcomes of the Programme offered by the institution



Vivekanand Education Society's College of Pharmacy Hashu Advani Complex, Near Collector's Colony, Chembur (E) Mumbai 400074

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M PHARM CBCS SYLLABUS

VES COLLEGE OF PHARMACY

Hashu Advani Memorial Complex, Behind Collector Colony, Chembur (E), Mumbai - 400 074

Teachers and students are aware of the stated Programme and course outcomes of the Programmes offered by the institution (15)

Describe Course Outcomes (COs) for all courses and mechanism of communication

M PHARM CBCS SYLLABUS COURSE OUTCOMES

	PSO MAPPING	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3
OUTCOMES	PO MAPPING	1, 2, 3, 7, 8, 11	1, 2, 3, 4, 6, 7, 11	1, 2, 4, 6, 7, 8, 11	1, 2, 3, 4, 5, 10, 11	1, 3, 4, 6, 7	1, 2, 3, 4, 6, 7, 8, 10, 11	1, 2, 3, 4, 6, 7, 8, 11	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11
M PHARM CBCS SYLLABUS COURSE OUTCOMES	STATEMENTS	Recall the concept of protein folding, receptors and their types, SAR, mechanism of action of certain class of drugs, enzyme kinetics and principles of enzyme inhibitors	Explain and illustrate the principles and applications of medicinal chemistry to impart knowledge about recent advances in the field of medicinal chemistry at the molecular level including different techniques for the rational drug design.	Application of the gained knowledge in basic research of rational design of enzyme inhibitors along with their metabolic profile and stereochemistry.	Evaluating and interpreting the role of chirality in selective and specific therapeutic agents to realize that stereo-selectivity is a prerequisite for drug evolution.	Understand the concepts of pre-formulation, micromerities, tablet compression, optimization.	Apply the preformulation and excipient knowledge for proper design of safe, efficacious, stable and quality formulations.	Investigate various aspects of solubility, dissolution and stability	Analyze the formulation parameters, apply optimization techniques and devise suitable formulation composition.
Z	Course Outcome	C01	CO2	CO3	CO4	C01	C02	CO3	CO4
	SUBJECT			Modern Pharmaceutica	l and Medicinal Chemistry (CBCS)			Modern	Pharmaceutics (CBCS, 2016-17)
	SEM								

Modern Pharmacology (CBCS	CO1	Explain the mechanisms of drug transport and concepts of Pharmacokinetics, pharmacodynamics.	1, 3, 6, 8, 9	1,2,3
Revised 2013)	C02	Explain the factors affecting drug responsiveness, mechanisms of drug dependence and microbial resistance and apoptosis.	1,3,4, 6, 7, 8, 9	1,2,3
	CO3	Explain Immunopharmacology and advances in the pharmacotherapy of CNS, CVS and Diabetes Mellitus.	1,3,4, 6, 8, 9	1,2,3
Modern Analytical Techniques	C01	Recall with examples the terminologies associated with spectroscopy, chromatography, X-ray diffraction, electrophoresis, potentiometry and thermal analysis	1, 2, 3, 8, 11	1,2,3
; :	C02	Explain and illustrate the theory, instrumentation and applications of various techniques involved in spectroscopy, chromatography, X-ray diffraction, electrophoresis, potentiometry and thermal analysis	1, 2, 3, 4, 6, 8, 11	1,2,3
Appropriate to the state of the	CO3	Apply the knowledge gained to calculate concentration by UV-visible spectroscopy, predict the IR frequencies, number of signals in NMR and fragmentation pattern in MS for simple organic compounds	2, 3, 4, 11	1,2,3
	CO4	Predict the spectroscopic behavior of molecules	2, 3, 4, 8, 11	1,2,3
Study of Natural Products (CBCS)	C01	Define and summarize phytochemicals and herbal drugs used in drug discovery, nutraceuticals, immunoglobulins and related applications	1, 2, 3, 8, 11	1, 2, 2003
	CO2	Explain the use of herbal drugs as excipients, in utraceuticals, as immunoglobulins and related applications	1, 2, 3, 4, 6, 8, 11	1, 2, 2003
	CO3	Apply the knowledge gained to isolate phytochemicals from herbal drugs and carry out standardization	2, 3, 4, 11	1, 2, 2003
	C04	Summarize the information of various herbs from Herbal Pharmacopoeia	2, 3, 4, 8, 11	1, 2, 2003

Research Research methodologies like objectives study design, review of literature, randomization, types of studies 1,2,6,7,8,9,10,11	CO3 Identify and make use of descriptors of molecules to develop an equation to quantitatively establish the structure activity relationship. Apply the acquired knowledge in design of covalently and non-covalently binding enzyme inhibitors,
- a	CO3

Advanced	100	Learn and apply advanced concepts of Stereochemistry	1,3,4,8,10,11	1, 2, 3
	C02	Understand and explain basic concepts of Catalysis, its types and different reactions of organometallic compounds.	1,3,4,8,10,11	1, 2, 3
	CO3	Understand the restrosynthetic methods and apply the knowledge of reactions covered for predicting retrosynthetic pathways of newer drugs	1,3,4,8,10,11	1, 2, 3
	C04	Apply and integrate acquired concepts of asymmetric synthesis in synthesis of chiral medicinal compounds.	1,3,4,8,10,11	1, 2, 3
	c05	Understand the merits and techniques involved in combinatorial synthesis	1,3,4,8,10,11	1, 2, 3
	900	Understand various greener chemistry approaches and compare them against conventional methods of Synthesis	1,3,4,8,10,11	1, 2, 3
Advanced Pharmaceutics CO1 I (CBCS Revised 2016)	C01	Understand the recent advances in tablet technology. Will acquire insight to oral controlled release drug delivery system and machinery used for the same.	1,3,5,6,9,11	1,3
	CO2	Familiarize with the recent advances in particulate drug delivery systems, provide an insight to formulation and evaluation of small volume and large volume Parenterals and study the recent advances in injectable controlled release and long acting formulations	1,3,5,6,9,11	1,3
	CO3	Will be introduced to specialized pharmaceutical disperse phase systems	1,3,5,6,9,11	1,3
	CO4	Understand the recent advances in gastro retentive oral drug delivery systems, concepts and various types of oral controlled release drug delivery system and evaluation methods for the same	1,3,5,6,9,11	1,3

1,3	1,3	1,3	1,3	1,3	1,3	1,3	1
1,3,5,6,9,11	1,3,5,6,9,11	1,3,5,6,9,11	1,3,5,6,9,11	1,3,5,6,9,11	1,3,5,6,9,11	1,3,5,6,9,11	1,2,3,4,8,
Acquire knowledge on site specific drug delivery systems to increase therapeutic efficacy of drug with minimum side-effects. Understand physiology of eye and develop advancements in ocular controlled drug delivery systems.	Acquire knowledge on site specific drug delivery systems to increase therapeutic efficacy of drug with minimum side-effects. Understand in detail biochemistry and anatomy of skin, recent developments in transdermal drug delivery systems and evaluate TDDS as per regulatory guidelines. Perceive knowledge on Quality by design to obtain safe, effective and reproducible formulations (as per ICH guidelines)	Understand site specific drug delivery systems to increase therapeutic efficacy of drug with minimum side-effects. Will know specialized pharmaceutical dispersed systems and recent advances in particulate drug delivery systems.	Understand anatomy and physiology of nasal mucosa and lungs. Have knowledge on recent developments in nasal and pulmonary drug delivery systems and its applications.	Understand the structural complexity and challenges to peptides and protein delivery of drugs and develop recent developments in peptide based drug delivery systems.	Gain knowledge on recent advances in particulate drug delivery systems.	Acquire knowledge on site specific drug delivery systems to increase therapeutic efficacy of drug with minimum side-effects. Understand physiology of brain, its barriers and develop advancements in brain controlled drug delivery systems.	Understand the concept of validation, qualification and calibration
500	30	100	CO2	CO3	CO4	25	100
		Advanced Pharmaceutics II (CBCS Revised 2016)					Quality Assurance System (CBCS)

Control of the contro	C02	Describe procedure for qualification of instruments and equipment	1.2.3.4	1
	CO3	Summarize the parameters of ICH guidelines for analytical method validation.	1,11	3
	C04	Comprehend the concept of process validation of different dosage forms	1,2,3,4,5,	1
	CO5	Gain knowledge of the process of cleaning validation	1,2,3,4,8,	2
	900	Correlate the knowledge of IPR with respect to pharmaceutical products	1,2,3,4,7, 8,11	33
Pharmaceutica I Quality Management	C01	Understand the concept of quality, strategic quality management and define different terms involved in quality management systems.	1,2,5,6	1,3
	CO2	Understand the concept of statistical process control (SPC) and explain the principles involved in SPC like process capability, control chart analysis and process control.	2,3,4,5,11	1,3
	CO3	Recognize the importance of customer, different concepts required to achieve customer satisfaction and desired quality the development of quality culture and define and comprehend the different terms, types and process involved in benchmarking.	3,6,8,9,10	es.
	C04	Comprehend principles involved in pharmaceutical quality management like six sigma, ISO, WHO-GMP and CFR-21	2,3,4,5,6,7,10	1,2
	500	Apply ICH guidelines for drug stability, risk management and quality by design.	1,2,3,5,8,9	1,3
Drug Metabolism	100	Recall the concept of drug metabolism, types of drug metabolism and in silico drug metabolism prediction	1, 2, 3, 7, 8, 11	1,2,3
	C02	Explain and illustrate the mechanism for metabolism of drugs through primary and secondary pathways	1, 2, 3, 4, 6, 7, 11	1,2,3
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2	Andread of control to the Control of	CO3	Application of the gained knowledge in basic studies on metabolism and metabolic profiling	1, 2, 4, 6, 7, 8, 11	1,2,3
	Experimental Techniques in	C01	Design novel drug delivery systems and evaluate them.	1, 3, 4, 6, 7	1,2,3
	Pharmacoufica	CO2	Apply the preformulation and excipient knowledge for proper design of safe, efficacious, stable and quality formulations.	1, 2, 3, 4, 6, 7, 8, 10, 11	1,2,3
	1	CO3	Investigate various aspects of dissolution and its mathematical treatment.	1, 2, 3, 4, 6, 7, 8, 11	1,2,3
	1	CO4	Analyze the formulation parameters, apply optimization techniques and devise suitable formulation composition.	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11	1,2,3
	Rational Drug	100	Recall and relate the different structures of protein along with the structure activity relationship of existing studied drugs and their interactions with the protein residues.	1, 2, 3, 4, 7, 8, 11	1,2,3
	nesign (co.go)	CO2	Classify and explain the different techniques to calculate the potential and kinetic energies of the system using Quantum and Molecular Mechanics, energy minimization and molecular conformational space search in the binding cavity of protein.	1, 2, 3, 4, 5, 8, 9, 11	1,2,3
		CO3	Make use of the minimal energy conformation of protein and ligand to construct and develop a model based on desired techniques like molecular docking, 3D-QSAR, pharmacophore modelling, homology modelling, molecular dynamics, etc.	1, 2, 4, 5, 6, 8, 9, 11	1,2,3
I PALL		CO4	Analyze the results obtained based on the characteristics of different interactions (docking), equations (QSAR), binding energy (dynamics) and interpret the molecular mechanism of how a drug act in a particular manner to be either inhibiting or stimulating the enzyme/receptor.	1, 2, 3, 4, 5, 7, 8, 9, 11	1, 2, 3
	Advanced Biochemistry (CBCS)	C01	Recall the protein subfamilies along with defining the terminologies like metabolism, nucleic acid, enzymes, cofactors, biomolecules, etc.	1, 2, 3, 4, 7, 8, 11	1, 2, 3

	C02	Classify and nomenclature of lipids, carbohydrates and nucleic acids, purification, characterization and complete of professions	1, 2, 3, 4, 5, 8, 9, 11	1,2,3
N I I I I I I I I I I I I I I I I I I I	CO3	Apply the knowledge gained in understanding the effects of drugs on lipid metabolism, protein function, nucleic acid biosynthesis, carbohydrates linkages to improve the pharmacokinetic properties of certain drugs	1, 2, 3, 4, 5, 8, 9, 11	1, 2, 3
Green	C01	Know the terms involved in green chemistry and know various guidelines of the environmental management system.	1, 2, 3, 4, 8, 9, 10, 11	1, 2, 3
(cecs)	C02	Understand the concept and techniques of waste management and illustrate the twelve principles of green chemistry. Make use of the microwave concept in the synthetic reactions.	1, 2, 3, 4, 5, 6, 8, 9, 10, 11	1, 2, 3
	CO3	Outline type of catalysis and their uses, safe solvents, water as reaction solvent.	1, 2, 3, 4, 5, 7, 8, 10, 11	1,2,3
	CO4	Learn greener process designing and future prospects to be applied in their research areas.	1, 2, 3, 4, 5, 7, 8, 10, 11	1, 2, 3
Drug Regulatory Affairs	COI	Understand the concepts of innovator and generic drugs, drug development process and the Regulatory guidance and guidelines for filing and approval process.	1,2,4,6,7,9,1	1,2
(CBCS)	C02	Develop and submit the dossiers in CTD/ eCTD formats and the post approval regulatory requirements for actives and drug products	1,2,3,4,7,8,10	1,2
	C03	Understand the requirements in the clinical trials settings and pharmacovigilance activities	1,2,3,4,5,6,7,8,9,10,11	1,2,3
	C04	professional and practical need of pharmaceutical industry.	1,2,3,4,5,6,7,8,10,11	3

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1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3	1,2,3
1,2,3,4,6,7,9,10,11	1,2,3,4,6,7,9,10,11	1,2,3,4,5,6,7,8,10,11	1,2,3,4,5,6,7,8,10,11	1,2,3,4,6,7,9,10,11	1,2,3,4,6,7,9,10,11	1,2,3,4,6,7,9,10,11	1,2,3,4,5,6,7,8,10,11	1,2,3,4,6,7,8,9,10,11	1, 2, 3, 8, 11	1, 2, 3, 4, 6, 8, 11
Describe the role and functional performance of cosmetic excipients , therapeutics ingredients and perfumes in the formulation of cosmetics for skin, hair, nails and oral care.	Understand the quality evaluation and regulations for the use of colors in cosmetics	Formulate and evaluate cosmetics for skin care and hair care as well as dental and oral care	Design and evaluate herbal cosmetics for skin care, hair care and oral care	Utilize novel approaches of formulation technologies in delivery of functional ingredients to skin, hair nails and oral cavity.	Study the classification and preparation methods of synthetic polymers	Study the characterization of polymers rheologically and thermally.	Know about biocompatibility of polymers and understanding the properties of biocompatible polymers are.	Explain why polymers are used in drug delivery applications	Recall with examples the terminologics associated with in vitro methods available for targeted drug delivery systems	Explain and illustrate the various evaluation techniques available for targeted drug delivery systems, basic principles of drug discovery and estimation of drug from complex media
C01	CO2	CO3	CO4	CO5	001	CO2	603	CO4	C01	CO2
_	Revised 2016)	NOT THE STATE OF THE STATE OF	· ·	4	Polymers in Pharmacy (CBCS	Revised 2016)			Drug Evaluation Techniques	(cgcs)
				9					SEM II	

1,2,3	
2, 3, 4, 11	
Apply the knowledge gained to perform in vitro assays and screening methods for different drugs and novel drug delivery systems	
CO3	
	SEM II

Dr. (M

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M PHARM R 2019 SYLLABUS

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Teachers and students are aware of the stated Programme and course outcomes of the Programmes offered by the institution (15) Describe Course Outcomes (COs) for all courses and mechanism of communication 2.6.1 QIM

M PHARM R 2019 SYLLABUS COURSE OUTCOMES

VNekanand Elucation Society's Callege of Pharmacy HAMC, Behind Collector Colony, Chembur, Mumbai - 400 074. PSO MAPPING 1, 2, 3 1, 2, 31, 2, 3 1, 2, 3 1, 2, 3 1, 2, 3 1, 2, 3 1, 2,3 1,2,3 1,2,3 PO MAPPING 1,2,3,4,7,8,10 1, 2, 3, 8, 11 1, 2, 3, 4, 6, 8, 11 1,2,4,5,6,7, 2, 3, 4, 8, 11 1,2,3,4,5,6,7, 1, 2, 3, 4, 6, 7, 8, 10, 11 1, 3, 4, 6, 7, 2, 3, 4, 11 1, 2, 3, 4, 6, 8.10.11 Apply the preformulation knowledge for proper selection of formulation spectroscopy, predict the IR frequencies, number of signals in NMR and Understand formulation and evaluation of Novel drug delivery systems... Analyze the formulation parameters, apply optimization techniques and Apply the knowledge gained to calculate concentration by UV-visible Recall with examples the terminologies associated with spectroscopy, chromatography, X-ray diffraction, electrophoresis & immunoassaya various techniques involved in spectroscopy, chromatography, X-ray Explain and illustrate the theory, instrumentation and applications of Understand the concepts and various approaches for development of Understand the concepts of pre-formulation, tablet compression, Investigate various qualification parameters for equipments and Understand criteria for selection of drugs and polymers for the fragmentation pattern in MS for simple organic compounds Predict the spectroscopic behavior of molecules diffraction, electrophoresis and immunoassaya STATEMENTS device suitable formulation composition. validation parameters for dosage forms. optimization, validation and cGMP. development of delivery system. novel drug delivery systems. excipients Course Outco me C01 C02 C04 CO3 C04 co_3 C01 C02 C02 CO3 C01 **Drug Delivery Systems** Modern Pharmaceutics Analytical Techniques **Pharmaceutical** SUBJECT Modern SEM

upriya S. Shidhaye

							3			-						Mr.	Supriya S. Shidhaye	Vivekanand Educotion Society's College of Pharmacy HAMC, Behind Collector Colony, Chembur, Mumbal - 400 074.
1,2	1,2	1,2,3	3	1	3	1,3		123	2616			1,2,3	1, 2, 3	1, 2, 3	1, 2, 3	1	Dr. (Ars.)	Vivekanan Colls HAMC, Es Chembu
1,2,4,6,7,9,11	1,2,3,4,7,8,10	1,2,3,4,5,6,7, 8,9,10,11	1,2,3,4,5,6,7, 8,10,11	1,3,4,8,11	1,2,3,4,8,11	1,2,3,4,8,9,10		1,2,3,4,8,9,10	,11			1, 2, 3, 8, 11	1, 2, 3, 4, 6, 8, 11	2, 3, 4, 11	2, 3, 4, 8, 11	1,3,8,11	0 20 8 8 10,111	Society Changlar Chan
Understand the concepts of innovator and generic drugs, drug development process and the Regulatory guidance and guidelines for filing and approval process.	Develop and submit the dossiers in CTD/ eCTD formats and the post approval regulatory requirements for actives and drug products	Understand the requirements in the clinical trials settings and pharmacovigilance activities	To correlate the theoretical knowledge with professional and practical need of pharmaceutical industry.	Estimate the active pharmaceutical ingredients in formulations by using different modern analytical techniques.	Apply the concepts of pre-formulation in formulation development.	Understand the formulation and evaluation methods of different novel drug delivery system.	Plan, execute the experiment using various methodologies	(defined protocol or qualitative or quantitative techniques)	and summarize the findings in systematic way verbally and	in written communication.	PHARM PHARMACEUTICAL CHEMISTRY	Recall with examples the terminologies associated with spectroscopy, chromatography, X-ray diffraction, electrophoresis, potentiometry and thermal analysis	Explain and illustrate the theory, instrumentation and applications of various techniques involved in spectroscopy, chromatography, X-ray diffraction, electronhoresis, notentiometry and thermal analysis	Apply the knowledge gained to calculate concentration by UV-visible spectroscopy, predict the IR frequencies, number of signals in NMR and fragmentation nattern in MS for simple organic compounds	Predict the spectroscopic behavior of molecules	Predict and explain the reaction products based on reaction intermediates and mechanism involved.	thetic routes available for synthesis of	Middla Assertion
CO1	C02	C03	C04	C01	C02	C03			CO4		Σ	C01	C02	CO3	C04	C01	C02	
			Regulatory Affairs	G.					Pharmaceutics	Practicals - I				Modern	Analytical Techniques			-
										SEMI	SEMI							

1,2	1,2,3	1	1	1,2	1,2,3	1,2	1,2	1,2,3	1,2,3	1-6,8,11	1,2,3,5,6,8,11	1,2,3,5,6,8,11		1, 2, 3	1, 2, 3	of "Mrs.) Supriva S. Shidhaye	Vhr.2h.3nand Education Scolety's College of Phermacy HAMC, Beilind Collector Colony, Chembur, Mumbal - 400 074.
1,3,8,11	1,3,8,11	1,11	1,6,11	1,3,11	1,2,3,11	1,6,9,10,11	1,3,6,7,9,10,1	1,4,6,7,9,10,1 1	1,3,4,7.9,10,1 1		1,2,3,5,6,8,11 1,2,	1,2,3,5,6,8,11 1,2,		1, 2, 3, 8, 11 1,	1, 2, 3, 4, 6, 1, 8, 11		Be Circle
1,3	1,3		1,	1,		1,6,5	1,3,6,	1,4,6,	1,3,4,	1-6,8,11	1,2,3,	1,2,3,		1, 2,	1, 2,	A STATE OF THE PERSON OF THE P	4.5.5.4 2.5.4
Apply Concept of protecting and deprotecting groups in synthetic schemes.	Apply the knowledge of reactions covered in syllabus for predicting retrosynthetic pathways of newer drugs.	Summarize Different stages of drug discovery	Explain Role of medicinal chemistry in drug research	Correlate different techniques for drug discovery and medicinal chemistry	Drive or deduce appropriate enzyme inhibitor or peptidomimetic if given the case	Recognize the different types of natural compounds and their chemistry and medicinal importance	Explain the phytochemical importance of alkaloid, flavonoid, steroids, terpenoids and vitamins in drug discovery	Use rDNA technology in new drug discovery	Justify the structural elucidation of natural compound based on its various spectroscopic parameters	perform quantitative analysis of organic compounds	perform the various reactions of synthetic importance	isolate products and interpret the experimental data	M PHARM QUALITY ASSURANCE	Recall with examples the terminologies associated with spectroscopy, chromatography, X-ray diffraction, electrophoresis, potentiometry and thermal analysis	Explain and illustrate the theory, instrumentation and applications of various techniques involved in spectroscopy, chromatography, X-ray diffraction, electrophoresis, potentiometry and thermal analysis	risible	Predict the spectroscopic behavior of molecules
CO3	C04	C01	C02	CO3	CO4	C01	C02	CO3	CO4	100	C02	CO3		C01	C02	CO3	CO4
	Advanced Organic Chemistry –I			9	Advanced Medicinal Chemistry				Chemistry of Natural Products		PHARMACEUTICAL	PRACTICAL-I				Modern	Analytical Techniques
								-				SEMI					

														Stiphya S. Shidhaye	Vivekanand Petucerfor Society's College of Prumacy HAMC, Behind Collector Colony, Chembur, Mumbal - 400 074.
1,3	1,3	33	1,2	1,3	1	1	1	1	1	1,2	1,2,3	1,2,3	3	Dr. (Mrs.)	Vivelransin Colls HAMC, Be Chembu
1,2,5,6	2,3,4,5,11	3,6,8,9,10	2,3,4,5,6,7,10	1,2,3,5,8,9	1,	1,2	1,	1	1	1,2,4,6,7,9,11	1,2,3,4,7,8,10	1,2,3,4,5,6,7, 8,9,10,11	1,2,3,4,5,6,7,8,10,11	2, 3, 4, 6, 8,	Society Chembur
Understand the concept of quality, strategic quality management and define different terms involved in quality management systems.	Understand the concept of statistical process control (SPC) and explain the principles involved in SPC like process capability, control chart analysis and process control.	Recognize the importance of customer, different concepts required to achieve customer satisfaction and desired quality the development of quality culture and define and comprehend the different terms, types and process involved in benchmarking.	Comprehend principles involved in pharmaceutical quality management like six sigma, ISO, WHO-GMP and CFR-21.	Apply ICH guidelines for drug stability, risk management and quality by design.	Understand the roles and responsibilities of Quality Control and Quality Assurance departments in pharmaceutical industry	Understand the significance of cGMP and ICH Guidelines in pharmaceutical industry	Describe the analysis of raw materials, packaging materials, in process quality control (IPQC) and finished products for different pharmaceutical dosage forms	Apply knowledge of regulatory requirements for preparing, maintaining, retaining and retrieving the data and documents in pharmaceutical industry	Understand the scope and importance of Intellectual Property Rights (IPR) in pharmaceutical industry	Understand the new product development process, pilot plant scale up and packaging requirements	Understand the necessary information to transfer technology from R&D to actual manufacturing by sorting out various information obtained during R&D	Understand the requirements in the manufacturing settings and regulatory activities	Correlate the theoretical knowledge with professional and practical need of pharmaceutical industry.	vis spectroscopy, fluorescence spectroscopy arform, analyze, determine and report the tion/sample solution	14 + 602°
C01	C02	CO3	CO4	CO5	C01	C02	CO3	CO4	c05	C01	C02	CO3	C04	C01	
				Quality Management Systems					Quality Control and Quality Assurance			à	Product development and technology transfer	7.0	

		C02	Relate the concept of in process quality control & stability studies to design and develop the protocol for testing of pharmaceuticals		1, 2, 3
SEMI	Quality Assurance Practical-I	C03		1, 2, 3, 4	1, 2, 3
			M PHARM PHARMACEUTICS SEM II		
		C01	Understand concept of drug targeting, its application, pulmonary drug delivery systems and gene therapy.	1, 4, 6, 9, 11	1, 2, 3
		C02	Apply the knowledge for selection of appropriate Nanotechnology and delivery system for given class of drug and route of administration.	1, 3, 4, 6, 10, 11	1, 2, 3
		CO3	very	1, 3, 4, 6, 11	1, 2, 3
	Nano technology and	CO4	st composition of NDDS, encompassing Micro and Nano drug systems.	1, 2, 3, 4, 5, 6, 8, 9, 10, 11	1, 2, 3
		C01	Understand the basic concepts in biopharmaceutics and pharmacokinetics	1,2,4,6,7,10,1 1	1, 2,3
		C02	Understand how to use raw data and derive the pharmacokinetic models and parameters that best describe the process of drug absorption, distribution metabolism and elimination.	1,2,3,4,7,8,9,	1,2,3
		C03	Understand the critical evaluation of biopharmaceutic studies involving drup product equivalency	1,2,3,4,5,6,7,8,10,11	1,2,3
		C04	Understand the design and evaluation of dosage regimens of the drugs using pharmacokinetic and biopharmaceutic parameters.	1,2,3,4,5,6,7,8,10,11	123
	Biopharmaceutics & Pharmacokinetics	500	Understand the potential clinical pharmacokinetic problems and application of basics of pharmacokinetic	1,2,4,6,7,10,1	123
		C01	Recall & relate 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	1,2,3,4,6,8,11	1,2,3
		C02	Outline the principles of more integrated and coherent use of computerized information (informatics) in the drug development process	1,2,3,4,6,9,11	1,2,3
	,	CO3	Construct the simulated model of drug delivery systems based on ADME parameters, use of statistical techniques, clinical data collection and management	1,2,3,4,5,8,9,	1,2,3
	Computer Aided Drug Delivery Systems	C04	applications of artificial intelligence and robotics in al automation, evaluate the current challenges and prections	dict 123.4,5,8,111	1,27,000

		500	Define cosmetics and understand the regulatory requirements for	2678911	2
	•	100	labeling, import, manufacturing and sale of cosmetic products in India.		
	(2)	C02	Understand the biological concepts related to different problems of the skin, hair, oral cavity.	1,3,9,11	3
		CO3	Study and review COSMOS guidelines for different classes of ingredients.	11,2,9,10,11	2
		C04	Classify the key ingredients, building blocks, their chemical classes and types, the herbal ingredients used in skin care, hair care required for	1,2,3,4,10,11	1,3
			making cosmetics and cosmeceuticals.		
	Cosmetics and	500	Apply the key ingredients for design and formulation of cosmeceutical products like sunscreen, antiageing, anti-acne and formulations for oral	1,2,3,4,5,10	1,3
	Cosmeceuticals		cavity problems.		
		5	Learner will gain knowledge in the area of advances in novel drug	1.2.3.4.7.8.9	1,3
		5	delivery systems.	-1-1-1-1-1-1-	
			Learners will have knowledge of methods used to determine and		
		C02	interpret the bioavailability and bioequivalence parameters along with	1,2,3,4,7,8,9	1,2,3
			statistical aspects of bioequivalence study.		
,			Learner will attain knowledge and skills necessary for computer		
			applications in pharmaceutical research and development, preclinical		
		03	and clinical development using statistical models/ software, including	1,2,3,4,7,8,9	1,2,3
			optimization of formulation and process of manufacturing,		
			computational modeling of drug disposition.		
	Pharmaceutics Practical	500	Learner will gain knowledge and skills necessary for the developing	1,2,3,4,7,8,9,	1.3
SEM II	=	50	synthetic and herbal cosmetic and cosmeceutical products.	10	o/-
	•				
			PHARMACEUTICAL CHEMISTRY SEM II		
		C01	Recall with examples the terminologies of advanced chromatographic & hyphenated techniques, thermal analysis & radio immunoassays	1, 2, 3, 8, 11	1, 2, 3
			Explain and illustrate the theory and applications of 1-D & 2-D NMR,	12346	
		C02	જ	8, 11	1, 2, 3
			Apply the knowledge gained and perform mathematical calculations to		,
		C03	NMR; Amax, and DBE of organic compounds; mass to charge ratio of	2, 3, 4, 0, 11	1, 2, 3
			tragments in MS		
	Advanced Spectral	C04	Interpret the spectral data and predict the structure of organic compounds.	2, 3, 4, 6, 7, 8, 9 Dr	1, 2, 3
_	Auarysis			100	Ta.

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-		C01	Understand various greener chemistry approaches and compare them against conventional methods of syntheis	1,3,4,8,10,11	1,2
		C02	Learn and express advanced techniques of peptide synthesis	1,3,4,8,10,11	1
		CO3	Describe and discuss upon photochemical and pericyclic reactions	1,3,8,10,11	П
		C04	Learn type of catalysis, its basic mechanism, and various catalytic named reactions used in industrial manufacturing set up.	1,3,7,8,10,11	1,2
	Advanced Organic Chemistry -II	CO5	Apply and integrate acquired concepts of asymmetric synthesis in synthesis of chiral medicinal compounds.	1,3,7,8,10,11	1,2,3
		C01	Recall and relate the different structures of protein along with the structure activity relationship of existing studied drugs and their interactions with the protein residues	1,2,3,10,11	1,2,3
		C02	Classify and explain the different techniques to calculate the potential and kinetic energies of the system using Quantum and Molecular Mechanics, energy minimization and molecular conformational space search in the binding cavity of protein	1,2,3,10,11	1,2,3
		CO3	Make use of the minimal energy conformation of protein and ligand to construct and develop a model based on desired techniques like molecular docking, 3D-QSAR, pharmacophore modelling, homology modelling, molecular dynamics, etc.	1,2,3,4,6,8,11	1,2,3
	Computer Aided Drug Design	C04	Analyze the results obtained based on the characteristics of different interactions (docking), equation (QSAR), binding energy (dynamics) and 1,2,3,5,7,10,1 interpret the molecular mechanism of how a drug acts in a particular manner to be either inhibiting or stimulating the enzyme/receptor	1,2,3,5,7,10,1	1,2,3
		C01	Describe the strategies of scale up process of APIs and intermediates	1,2,3,4,6	1,3
	Pharmaceutical Process Chemistry	C02	Elaborate various unit operations and various reactions in process chemistry	12,3,4,6,7,10, 11	1,2,3
		CO3	Describe various principles of Industrial Safety	12,3,5,6,8,10, 11	1,2,3
		C01	perform the various reactions of synthetic importance	1-6,8,10,11	1,2,3
	PHARMACEUTICAL	CO2		1-6,8,11	1,2,3
SEM II	CHEMISTRY PRACTICAL - II	C03	Experiment with computer aided techniques, validate the models and interpret and predict the results	1-6,8,10,11	(12,3 F

		QUALITY ASSURANCE SEM II		
	C01	Recall the environmental problems and develop an attitude of concern for the industry environment	1, 2, 3, 8, 10,	1, 2, 3
	C02	Make use of the knowledge gained to ensure safety standards in pharmaceutical industry	1, 2, 3, 4, 6, 8, 10, 11	1, 2, 3
	CO3	Analyze and simplify the mechanism and management in different kinds of hazard management system	2, 3, 4, 10, 11	1, 2, 3
Hazards and safety Management	C04	Propose the method of Hazard assessment, procedure and methodology for safe industrial atmosphere.	2, 3, 4, 8, 10, 11	1, 2, 3
	C01	Understand the concept of validation, qualification and calibration	1,2,3,4,8,	П
			11	
	C02	Describe procedure for qualification of instruments and equipment	1.2.3.4	1
	CO3	Summarize the parameters of ICH guidelines for analytical method validation.	1, 11	3
	C04	Comprehend the concept of process validation of different dosage forms	1,2,3,4,5,	1
	CO5	Gain knowledge of the process of cleaning validation	1,2,3,4,8,	2
Pharmaceutical Validation	900	Correlate the knowledge of IPR with respect to pharmaceutical products	1,2,3,4,7,	3
	C01	Understand the concept of Quality Management System, Quality audits& its role, importance in pharmaceutical manufacturing environment.	1, 5, 6,	1
	C02	Apply the conceptual knowledge gained to design & conducting Audits of various areas in the pharmaceutical manufacturing, packaging, storage, distribution, Quality control and ancillary areas (utilities) to assess compliance to the applicable Regulatory requirements.	1, 2, 6, 8	8
	CO3	Create Audit check lists to conduct audits in the above specified areas in pharmaceutical industries including vendors and suppliers of API Raw & packaging materials.	1, 2, 6, 7	7

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		Evaluate Audit observations into categories such as		
	700	Critical, Major & Minor based on the severity of non-compliance	1 3 6	-
Audits and Regulatory	5	to cGMP aspects and Regulatory requirements. Suggest CAPAs	1, 7,	•
Compliance		for compliance to CGMP aspects & Regulatory requirements		
		Understanding the legal requirements, licenses, plant layout, production		,
	C01	planning for the pharmaceutical industry, process automation with	1,2,3,4,5	1,2
		respect to different dosage forms.		
		Explain the concept of quality by design (QbD) and process analytical		
	C02	technology (PAT) and understand the different terminologies and aspects	1,2,3,4,5,8,9	1,2
		involved in QbD and PAT.		
		Analyze the aseptic and non-sterile process technology including	12345101	
	CO3	manufacturing requirements, new technologies and equipment required	1,01,0,1,0,1	1,3
		at each stage of manufacturing.	1	
Pharmaceutical		Evaluate packaging technology required for different types of dosage	12345101	
Manufacturing	C04	forms, evaluation of product package compatibility and stability aspects	1,01,0,10,1	_
Technology		of packaging material.	1	
		Understand the significance of control of hazardous substances and	27168	
	C01	perform analysis to determine and report the content of hazardous	10 11	1, 2, 3
		substances in air/environment	10, 11	
	C02	Relate the concept of quality assurance to design and develop the	1, 2, 3, 4, 6,	1, 2, 3
		process the choosings for esting of primings and the choosing of the choosing	1 7 2 4 10	
Quality Assurance	CO3	Plan, execute and conclude the experiment using qualitative or	1, 2, 3, 4, 10,	1, 2, 3
Practical-II	3	quantitative techniques	11	- 3 - 3 -
		M PHARM SEM III AND IV		
		Students will be able to explain basic research methodologies like	17678910	
	C01	objectives study design, review of literature, randomization, types of	11.	
		studies	11,	1,2,3
	500	Students will be able to explain, analyze the data and apply the statistical 1,2,3,6,7,8,9,	1,2,3,6,7,8,9,	
	700	principles in the evaluation of the research data	10,11	1,2,3
		Students will be able to explain the basic concepts of medical research		
		including informed consent, , concepts like autonomy, beneficence and	1236789	
	C03	non-maleficence, as well as about the declaration of Helsinki and other	10.11	
		guidelines like ICH GCP, Nuremberg code which govern ethical conduct of clinical trials		1.2.3
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		C04	Students will be able to explain the basic facilities in animal handling and animal house facilities like transport, storage and care of animals. As 1,2,3,6,7,8,9, well as about the basic procedures to be followed to ensure the efficient 10,11 management of animal house facility at the site.	1,2,3,6,7,8,9,	
	Research Methodology and Biostatistics				1,2,3
		C01	Develop knowledge to advance your career, specialize in a particular area and help take career in a promising new direction via experimental learning	1,11	1,2,3
		C02	Acquire skills related to literature survey, planning of experiments, data collection, data interpretation	1,2,3,11	1,2,3
		CO3	Learn handling of modern instruments, equipments or software required in the chosen area of work	1,2,4	1,2,3
		C04	Progress of critical thinking and analytical skills through hands-on learning	1,3,11	1,2,3
		C05	Develop oral and written scientific communication skills	8,11	1,2,3
SFM	: Project Work	900	Create innovative ideas or project which will help in understanding the specialized area in more depth and society in large	1,9,11	1,2,3



