

## ALARD COLLEGE OF PHARMACY

Sr.No 50, Near Rajiv Gandhi Infotech Park, Marunji, Pune 411057

## COURSE OUTCOMES B.Pharm First Year (I Sem) 2018 pattern BP101T. HUMAN ANATOMY AND PHYSIOLOGY-I (Theory)

Upon the completion of the course student shall be able to

CO1	Describe various parts of human body and their roles.						
CO2	Explain the structure of cells, tissues and organs along with its significance, various parts of CNS and PNS.						
CO3	Explain different bones in the human skeleton system, their location and significance, Endocrine system and its importance with the help of charts and models.						

## **BP107P. HUMAN ANATOMY AND PHYSIOLOGY-I (Practical)**

Upon the completion of the course student shall be able to

CO1	Identify, compare and contrast between the microscopy of epithelial,
	connective, muscular, nervous tissue of human body.
CO2	Explain the significance of bleeding time, clotting time, blood group
CO2	detection, hemoglobin detection and measurement of blood pressure.
CO3	Demonstrate procedure of white blood cell count and red blood cell
CO3	count and red blood cell count of blood sample.

## BP102T. PHARMACEUTICAL ANALYSIS-I (Theory)

CO1	Outline the method of expressing the concentration with preparation							
	and standardization of various molar and normal solutions.							
CO2	Recall the sources, type and method of minimizing the errors.							
CO3	Explain the principle involved in volumetric and electrochemical analysis							
	of inorganic compounds							

## **BP108P. PHARMACEUTICAL ANALYSIS (Practical)**

CO1	Prepare and standardize primary and secondary standard solutions of
	various normality and molarity
CO2	Perform various volumetric and electrochemical titrations

## **BP103T. PHARMACEUTICS- I (Theory)**

CO1	Outline the history of profession of pharmacy.							
CO2	Enumerate the basics of different dosage forms.							
CO3	dentify pharmaceutical incompatibilities in prescription ,their							
	manifestation and suggest solution to correct same							
CO4	Describe the professional way of handling prescription.							
CO5	Perform the pharmaceutical calculations required during formulation of							
	dosage form.							

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## **BP109P. PHARMACEUTICS I (Practical)**

Upon the completion of the course student shall be able to

1	Formulate various conventional dosage forms in professional way.						
2	Emphasize on the concepts of prescription like translation, calculation						
	and suitability						

## **BP104T. PHARMACEUTICAL INORGANIC CHEMISTRY (Theory)**

CO1	Explain the sources of impurities and method to determine the impurities						
	in an inorganic drugs and pharmaceuticals.						
CO2	Describe the importance of radiopharmaceuticals.						
CO3	Explain the method of preparation, assay, storage conditions and uses						
	of Inorganic compounds such as acidifiers, antacids, cathartics,						
	electrolyte replenisher, antimicrobials, dental products, medicinal gases						
	and miscellaneous compounds like expectorant, sedative, antidotes and						
	radiopharmaceuticals.						

## **BP110P. PHARMACEUTICAL INORGANIC CHEMISTRY (Practical)**

CO1	Identify the Inorganic compounds through various chemical tests.						
CO2	Perform the limit test for certain impurities like chloride, sulphate, iron,						
	arsenic, lead and heavy metals as per the Indian Pharmacopoeia						

## **BP105T.COMMUNICATION SKILLS (Theory)**

Upon the completion of the course student shall be able to

	Explain need of communication skills, barriers to communicate
CO1	effectively and perspectives of communication required to function
	effectively in areas of pharmaceutical operation
	Apply various elements, styles of communications, Basic listening skills,
CO2	writing skills to communicate effectively and manage team as team
	player
CO3	Apply Interview skills presentation skills and group discussion for
	development of leadership qualities and essentials

## **BP105T COMMUNICATION SKILLS (Practical)**

001	Demonstrate	and	Apply	basic	communication	skills	and	advance
COI	learning skills	}						

## **BP 106RBT. REMEDIAL BIOLOGY (Theory)**

CO1	Describe the classification and salient features of five kingdoms of life.						
CO2	plain the basic component of anatomy and physiology of plant						
CO3	Discuss the basic components of anatomy and physiology of animal with						
003	special reference to Human						



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CO4	Explain various parts of CNS, PNS and their Role
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## **BP112RBP. REMEDIAL BIOLOGY (Practical)**

Upon completion of the course student shall be able to

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CO1	Demonstration of different bone in human skeleton system, their
COI	location and significance.
CO2	Perform blood group detection, measurement of blood pressure and tidal
CO2	volume.
CO3	Able to identify microscopy of tissues pertinent to stem, root, leaf, seed,
CO3	fruit and flower.

## BP 106 RMT. REMEDIAL MATHEMATICS (Theory)

	Know the theory and their application of Partial fraction, Logarithms,
CO1	Function, in Pharmacy Limits and continuity, Matrices and
	Determinant, Calculus in Pharmacy
	Solve the different types of problems by applying theory of Partial
CO2	fraction, Logarithms, Function, in Pharmacy Limits and continuity,
	Matrices and Determinant, Calculus



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## B.Pharm First Year (II Sem) 2018 pattern BP 201T. HUMAN ANATOMY AND PHYSIOLOGY-II (Theory)

Upon the completion of the course student shall be able to

CO1	Explain the gross morphology, structure and functions of various
CO1	organs of the human body.
CO2	Describe various homeostatic mechanisms and their imbalances.
CO3	Discuss the anatomy of lungs and other parts of respiratory system,
CO3	tidal volume, artificial respiration and resuscitation methods.

## BP 207 P. HUMAN ANATOMY AND PHYSIOLOGY-II (Practical)

Upon the completion of the course student shall be able to

CO1	Identify various tissues and organs of different systems of human body.
CO2	Explain construction and working of spirometer for the measurement of
CO2	lung volume and capacities.

## **BP202T. PHARMACEUTICAL ORGANIC CHEMISTRY - I (Theory)**

Upon completion of the course student shall be able to

CO1	Outline the structure, name and the type of isomerism of the organic
COI	compound.
CO2	Describe the reaction name of the reaction and orientation of reactions
CO3	Explain the mechanism, kinetics and reactivity of the certain reactions

## BP208P. PHARMACEUTICAL ORGANIC CHEMISTRY - I (Practical)

Upon completion of the course student shall be able to

CO1	Perform the systematic qualitative analysis of organic compounds
CO2	Prepare the suitable solid derivatives from organic compounds

## **BP203 T. BIOCHEMISTRY (Theory)**

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CO1	Describe the chemistry, biological importance and metabolism pattern of
CO1	Biomolecules.
CO2	Summaries the concept of biological oxidation emphasizing on ETC and
CO2	oxidative phosphorylation and identifying related inhibitors.
CO3	Explain catalytic role of enzymes, importance of enzyme inhibitors in
CO3	design of new drugs, therapeutic and diagnostic applications of enzyme
	Explain the genetic organization of mammalian genome and functions of
CO4	DNA in synthesis of RNAs and proteins.

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## **BP 209 P. BIOCHEMISTRY (Practical)**

Upon completion of the course student shall be able to

CO1	Identify and characterize carbohydrates, proteins by various qualitative
COI	chemical tests in a given sample.
CO2	Determine blood creatinine, sugar, total cholesterol and action of
CO2	salivary amylase.

## **BP 204T. PATHOPHYSIOLOGY (THEORY)**

Upon the completion of the course student shall be able to

CO1	Explain the etiology and pathogenesis and complications of severe
COI	diseases and disorders.
CO2	Discuss the signs and symptoms of different diseases and their
CO2	diagnostic procedures.
CO3	Differentiate between acute and chronic diseases based on etiology,
CO3	signs and symptoms and complications.

## **BP205 T. COMPUTER APPLICATIONS IN PHARMACY (Theory)**

Upon the completion of the course student shall be able to

CO1	know the various types of application of computers in pharmacy
000	Understand Concept of Information Systems and Software, various types
CO2	of databases like MYSQL, MS ACCESS, Pharmacy Drug database,
	Number systems, Web technologies and Bioinformatics
	Apply computer knowledge for Chromatographic dada analysis(CDS),
CO3	Laboratory Information management System (LIMS) and Text
	Information Management System(TIMS)

## **BP210P. COMPUTER APPLICATIONS IN PHARMACY (Practical)**

Upon the completion of the course student shall be able to

		Use MS Word, MS Access for designing questionnaire, form to record
		patient information, creating patient database, mailing labels, invoice
		table, and generate reports
	CO2	Create HTML web page, Export Tables, Queries, Forms and Reports to
		web pages and XML Pages

## **BP 206 T. ENVIRONMENTAL SCIENCES (Theory)**

	Understand Multidisciplinary nature of environmental studies Natural
CO1	Resources Renewable and non-renewable resources, associated
	problems
CO2	Understand, explain and Draw Structure and function of various
CO2	ecosystem.
CO3	Understand Environmental Pollution and its remedial methods to reduce



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## B.Pharm Second Year (III Sem) 2018 pattern BP 301T. PHARMACEUTICAL ORGANIC CHEMISTRY-II (Theory)

Upon completion of the course student shall be able to

CO1	Describe the reaction and mechanism of Benzene, phenols, aromatic
	amines and polynuclear hydrocarbons.
CO2	Explain the stabilities of cycloalkanes through different theories.
CO3	Summarize the chemistry of fats and oils.

## BP305P. PHARMACEUTICAL ORGANIC CHEMISTRY-II (Practical)

Upon completion of the course student shall be able to

CO1	Determine the physical constants like acid value, saponification value					
COI	and Iodine value of organic compounds.					
	Synthesize certain organic compounds through acetylation,					
CO2	halogentaion, nitration, oxidation, hydrolysis, Perkins and claisen					
	condensation reactions					

## BP302T. PHYSICAL PHARMACEUTICS-I (Theory)

Upon the completion of the course student shall be able to

CO1	Explain	various	physicochemical	properties	of	drug	molecules
	applicabl	le in the d	esigning of dosage	forms.			
CO2	Demonst	rate use	of physicochem:	ical propertie	es in	the	formulation
	developm	nent and e	valuation of dosag	e forms			

## BP306P. PHYSICAL PHARMACEUTICS - I (Practical)

Upon the completion of the course student shall be able to

CO1	Determination of various physicochemical properties of drug molecules				
	applicable in the designing of dosage forms.				
CO2	Analyze and interpret the data generated from the experiments.				
CO3	Compare and contrast between different method used in the				
	determination of the same physicochemical parameters.				

## BP 303 T. PHARMACEUTICAL MICROBIOLOGY (Theory)

CO1	Explain methods of identification, cultivation and preservation of	of
	various microorganisms	
CO2	Summarize the importance and implementation of sterilization in	n
	pharmaceutical processing and industry.	



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CO3	Discuss microbiological standardization of Pharmaceuticals.
CO4	Outline cell culture technology and its applications in pharmaceutical industries.

## **BP 307P.PHARMACEUTICAL MICROBIOLOGY (Practical)**

Upon completion of the subject student shall be able to

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CO1	Select and utilize different equipment and processing in experimental
	microbiology
CO2	Identify and isolate various microorganisms
CO3	Perform sterility testing of pharmaceutical products
CO4	Perform microbiological standardization of Pharmaceuticals.

## **BP 304 T. PHARMACEUTICAL ENGINEERING (Theory)**

Upon the completion of the course student shall be able to

CO1	Explain use of various unit operations used in Pharmaceutical
	industries.
CO2	Describe the material handling techniques.
CO3	Discuss various methods of hazards and safety management used in
	Pharmaceutical industry
CO4	Outline the significance of plant layout design for optimum use of
	resources.
CO5	Enumerate the various preventive methods used for corrosion control in
	Pharmaceutical industry

## **BP308P - PHARMACEUTICAL ENGINEERING (Practical)**

CO1	Perform various unit operation process involved in pharmaceutical
	manufacturing
CO2	Perform numerical involved in calculating process related
	determinants.
CO3	Create graphs and illustrate actions for data representation
CO4	Analyze and interpret the data generated from the experiments performed.



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## B.Pharm Second Year (IV Sem) 2018 pattern BP 401 T PHARMACEUTICAL ORGANIC CHEMISTRY III (Theory)

On completion of course, student should be able to,

CO1	understand the methods of preparation and properties of organic compounds
CO2	explain the stereo chemical aspects of organic compounds and stereo chemical reactions
СОЗ	know the medicinal uses and other applications of organic compounds

## **BP 402T MEDICINAL CHEMISTRY-I (Theory)**

Upon completion of the course student shall be able to

CO1	Correlate the physicochemical properties and metabolism of drugs with
	biological activity.
CO2	Explain the chemistry of drugs acting on nervous system, opioid and
002	non opioid receptor.
CO3	Describe the mechanism of action of certain pharmacodynamics agents.

## **BP406P. MEDICINAL CHEMISTRY-I (Practical)**

Upon completion of the course student shall be able to

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CO1	Synthesize and explain reaction mechanism of medicinally important
	compounds by using conventional methods and purify them by using
	TLC and Column Chromatography.
CO2	Perform quantitative analysis of drugs such as Chlorpromazine,
CO2	Phenobarbitone, Atropine, Ibuprofen, Aspirin, Furosemide.

## **BP 403 T. PHYSICAL PHARMACEUTICS-II (Theory)**

CO1	Compare and contrast between colloidal and coarse dispersion based on					
	their general properties, principles of formulation and evaluation.					
CO2	Explain and comprehend the principles of preformulations like rheology,					
	deformation of solid and micromeretics.					
CO3	Explain use of physicochemical properties in the formulation					
	development and evaluation of dosage forms					
CO4	Explain with illustration the principles of chemical kinetics & to use					
	them for stability testing and determination of expiry date of					
	formulations					



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## BP 407P. PHYSICAL PHARMACEUTICS- II (Practical)

Upon completion of the subject student shall be able to

	<u> </u>
CO1	Determine physicochemical properties in the formulation development
	and evaluation of dosage forms.
CO2	Make use of principles of chemical kinetics & to use them for stability
	testing.
CO3	Compare and contrast between different method used in the
	determination of the same physicochemical parameters.
CO4	Demonstrate and explain the effect of different excipients and their
	differing concentration on physicochemical determinants of dosage
	forms.

## **BP404T. Pharmacology I - Theory**

Upon completion of the course student shall be able to

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CO1	Explain various terminologies used in pharmacology like synergism,					
	agonist, antagonist, side effect, adverse effects etc.					
CO2	Describe the pharmacological actions of different categories of drugs.					
CO3	Discuss the mechanism of drug action at organ system/sub					
003	cellular/ macromolecular levels.					
	Classify various drugs used for the treatment of disorders of nervous					
CO4	system according to their mechanism of action and apply the basic					
	pharmacological knowledge in the prevention and treatment of various					
	diseases.					

## **BP408P. Pharmacology I - Practical**

CO1	Handle the laboratory equipments and apply techniques used in					
	experimental pharmacology.					
CO2	Identify various laboratory animals and describe CPCSEA guidelines for					
	care and handling and care of laboratory animals.					
CO3	Explain common laboratory techniques, like blood withdrawal, serum and plasma separation, anesthetics and euthanasia used for animal studies.					
CO4	Describe the different routes of drug administration in mice/rats.					
CO5	Demonstrate the effect of drugs on animals by simulated experiments.					



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## BP 405 T. PHARMACOGNOSY & PHYTOCHEMISTRY-I (Theory)

Upon completion of the course student shall be able to

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CO1	Explain history, scope, development of pharmacognosy, sources of drugs and differentiate between organized and unorganized drugs					
CO2	Understand and explain classification of drugs and quality control of drugs of natural origin.					
CO3	Comprehend and understand cultivation, collection, processing, storage of drugs of natural origin, conservation of medicinal plants, plant tissue culture including its development, applications.					
CO4	Explain and understand morphology and anatomy of plant parts.  Explain classification, properties, identification of Glycosides, Tannins, volatile oil, Flavanoids and Resins					
CO5	Comprehend the biological source, chemical nature, uses of plant fibers, hallucinogens, Teratogens, Natural allergens					
CO6	Understand and explain pharmacognostic study of carbohydrates, Proteins, enzymes, lipids, marine drugs					

## BP 405 P. PHARMACOGNOSY & PHYTOCHEMISTRY-I (Practical)

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CO1	Perform analysis of crude drugs by chemical tests
CO2	Determine and perform stomatal number, stomatal index, vein islet
	number, vein islet termination and palisade ratio of leaf drug
	Understand and determine size of starch grains, calcium oxalate
CO3	crystals, length and width of fiber by eye piece micrometer and number
	of starch grains by Lycopodium spore method
CO4	Perform Ash value, Extractive values, moisture content, swelling and
	foaming index of crude drug



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## B.Pharm Third Year (V sem) (2015 Pattern)

## 3.5.1 T INDUSTRIAL PHARMACY-I (Theory):-

The students will be able to

CO1	Perceive the knowledge of dosage form design & formulation strategies.					
	Learn tablets as a dosage form, physic-chemical principles guiding tablet					
CO2	formulation, various tablet additives, manufacture & evaluation,					
	equipments, defects in tablets & remedies their off.					
CO3	Interpret the concept, types, pharmacopoeial specifications, techniques					
CO3	& equipments used in tablet coating.					
CO4	Describe the capsules, its types, additives, size selection,					
	manufacturing, evaluation and equipments used & its defects.					

## 3.5.1 P INDUSTRIAL PHARMACY-I (Practical):-

The students will be able to

	<u></u>
CO1	Utter the correct use of various equipments in pharmaceutics laboratory
	relevant to tablets, capsules and tablet coating.
CO2	Inculcate the knowledge of formulation, evaluation and labeling of
	tablets & capsules.
CO3	Use the equipments and apparatus needed for the preparation as per
	SOP.
CO4	Perform pharmaceutical calculations to determine evaluation of
	granules.
CO5	Describe use of ingredients in formulation and category of formulation.
CO6	Select the suitable packaging material for the preparation.

## 3.5.2 T PHARMACEUTICAL ANALYSIS-III (Theory)

Upon completion of the course student shall be able to

CO1	Enumerate the different types of instrumental analytical techniques				
	available for quality control of APIs & formulations.				
CO2	Explain sampling techniques used for analysis of solid, semisolid and				
CO2	liquids dosage forms.				
	Describe principles, instrumentation and applications of UV-VIS,				
CO3	Flourimetry, Flame photometry, phosphorimetry and				
	Nepheloturbidimetry.				

## 3.5.2 P PHARMACEUTICAL ANALYSIS-III (Practical)

( `( )	Operate UV	-VIS	Spectrometer,	Flame	Photometer,	Fluorimetry	and
	Phosphorime	eter.					
CO2	Interpret the data obtained through analytical experiments.						

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## 3.5.3 T MEDICINAL CHEMISTRY-I (Theory)

Upon completion of the course student shall be able to

CO1	Explain physicochemical properties and pharmacokinetics affecting						
	drug action						
CO2	Describe concept of Receptor along with drug-receptor mechanism.						
	Classify certain therapeutic agents and outline the synthetic route for						
CO3	the selective medicinal compounds of sympathetic, parasympathetic and						
	cardiovascular system.						
CO4	Explain the structural activity relationship of certain therapeutic agents						
	with their uses, adverse effects and recent developments.						

## 3.5.3 P MEDICINAL CHEMISTRY-I (Practical)

Upon completion of the course student shall be able to

CO1	Synthesize and explain reaction mechanism of medicinally important
	compounds by using conventional methods and purify them by using
	TLC and column chromatography.
CO2	Evaluate physicochemical properties of synthesized acid/basic salts of
	drugs.

## 3.5.4 T. PHARMACOLOGY II (Theory)

Upon completion of the course student shall be able to

CO1	Discuss ANS with respect to various neurotransmitters and their signal
	transduction mechanisms in the body.
CO2	Explain Cholinergic and Anti-cholinergic drugs, their classification and
CO2	pharmacology.
СОЗ	Explain Adrenergic and Anti-adrenergic drugs, their classification and
	pharmacology.
CO4	Describe Pharmacotherapy of CVS disorders and Respiratory tract
	disorders.

## 3.5.4 P. PHARMACOLOGY II (Practical)

CO1	Handle the laboratory equipments and apply techniques used in
	experimental pharmacology.
CO2	Identify various laboratory animals and describe CPCSEA guidelines for
	care and handling and care of laboratory animals.
CO3	Explain various routes of drug administration, various methods for
CO3	collection of blood, body fluids and urine from experimental animals.
CO4	Knowledge of various types of bioassays along with their principals,
	drug agonism and antagonism.
CO5	Perform recording of CRC/DRC of Acetylcholine on suitable isolated
	tissue preparation.



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## 3.5.5T. ANALYTICAL PHARMACOGNOSY & EXTRACTION TECHNOLOGY (Theory)

Upon the completion of the course student shall be able to

CO1	Comprehend principle of mass transfer process, effect of various factors
	on mass transfer & principle, working, merits, demerits and applications
	of various extraction techniques.
CO2	Explain principle & applications of chromatographic &
	nonchromatographic separation methods.
CO3	Understand and describe applications of various extraction techniques
	of phytochemicals by identifying their source, properties, isolation and
	tests.
CO4	Explain types, social relevance, Sampling techniques and quality control
	parameters of herbal drug analysis, WHO guidelines for quality control
	of herbal drugs & Current approaches in standardization.

## 3.5.5P. ANALYTICAL PHARMACOGNOSY & EXTRACTION TECHNOLOGY (Practical)

Upon the completion of the course student shall be able to

CO1	Understand and analyze micrometric data of herbal crude drug, perform
	solvent extractions from various plant crude drug and chemical analysis
	of plant extract.
CO2	Analyze herbal crude drug by applying various quality control
	parameters and adulterants in crude drugs and explain microwave
	extraction and column chromatography technique.

## 3.5.6 T PHARMACEUTICAL BUSINESS MANAGEMENT (Theory)

CO1	Describe the Pharmaceutical business and management strategy.
CO2	Gain knowledge of marketing research, product management.
СОЗ	Discuss about human resource and development needs.
CO4	Explain about the disaster management and preparedness, mitigation.
CO5	Participate in group discussion elocution/Extempore/Debate
CO6	How to crack job interviews.
CO7	Differentiate Management concepts and Marketing concepts.



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## 3.5.7 T ACTIVE PHARMACEUTICAL INGREDIENTS TECHNOLOGY (Theory)

CO1	Describe API and fine chemical industry.
CO2	Explain certain classes of reaction, chemical process, reaction system,
	equipment used in API.
CO3	Explain quality control aspects, material safety data sheet (MSDS),
CO3	health hazards, green chemistry approaches.
CO4	Summarize industrial manufacturing methods of certain APIs.
CO5	Explain polymorphism and the techniques involved in resolution of
	racemates and asymmetric synthesis.
CO6	Apply GMP guidelines like ICH Q7,Q7A AND Q11 IN API manufacturing.



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## B.Pharm Third Year (VI sem) (2015 Pattern)

## 3.6.1 T INDUSTRIAL PHARMACY-II (Theory):-

Upon completion of the course student shall be able to

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CO1	Understand disperse systems, its classification, theories of disperse
	systems, thermodynamics v/s kinetic stability considerations.
CO2	Rationalize suspensions, types, formulation development,
	manufacturing, excipients used, and evaluation of suspensions.
CO3	Determine emulsions, their physic-chemical properties, theory of
	emulsification, HLB value and phase inversion temperature, Kraft point,
	cloud point, excipients, and evaluation of emulsions, cracking,
	coalescence, stability and stress testing.
CO4	Recognize semi-solids, anatomy and physiology of skin, selection of
	bases, penetration enhancers, formulation development, percutaneous
	absorption, flux measurement and evaluation.
CO5	Summarize layout for manufacturing of suspensions, emulsions &
	semisolids as per schedule M.

## 3.6.1 P INDUSTRIAL PHARMACY-II (Practical):-

Upon completion of the course student shall be able to

CO1	Explain the correct use of various equipments in pharmaceutics
	laboratory relevant to suspensions, emulsions and semi-solids,
CO2	Formulate, prepare and evaluate suspensions.
CO3	Formulate, prepare and evaluate emulsions.
CO4	Formulate, prepare and evaluate semisolids preparations.
CO5	Prepare the labels so as to suit the regulatory requirements.

## 3.6.3 T MEDICINAL CHEMISTRY-II (Theory)

	<u> </u>
CO1	Explain general aspects of drug metabolism and drug design aspects of
	important drugs.
CO2	Classify the medicinal compounds and acquire knowledge about IUPAC
	names along with mechanism of action of them or the class to which
	they belong.
CO3	Apply scientific knowledge about relationship between biological activity
CO3	and structure of various CNS acting drugs and drugs acting on blood.
CO4	Outline the synthetic route for selective medicinal compounds along
	with their uses, adverse effects and recent developments in CNS active
	drugs and drugs acting on blood.

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## 3.6.3 P MEDICINAL CHEMISTRY-II (Practical)

Upon completion of the course student shall be able to

CO1	Synthesize and explain reaction mechanism of medicinally important
	compounds by using conventional as well as microwave assisted
	methods and purify them by using recrystallization techniques.
CO2	Explain the principle and procedure for the synthesis of compounds and
	interpret their characterization data obtained by IR/NMR spectroscopy.

## 3.6.4 T. PHARMACOLOGY III (Theory)

Upon completion of the course student shall be able to

CO1	Classify various drugs depending upon their pharmacological role and
	mechanism of action in any disease.
CO2	Describe pharmacology of various anesthetic agents (General and Local),
	psychotropic agents, various drugs used in treatment of CNS disorders,
	Parkinson's disease and Alzheimer's disease.
CO3	Explain pharmacology of drugs used in the treatment of G.I. tract
	disorders, Rheumatoid arthritis, Osteoarthritis and Gout.

## 3.6.4 P. PHARMACOLOGY III (Practical)

Upon completion of the subject student shall be able to

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CO1	Demonstrate the dissection of G.I. tract of chicken to isolate ileum.
CO2	Explain and perform matching point, bracketing and interpolation
	bioassay to find unknown concentration of Acetylcholine.
CO3	Demonstrate and discuss recording of effects of CNS acting drugs in
	rats/mice using Actophotometer and anti-epileptic activity using
	Convulsiometer with the help of software.
CO4	Demonstrate recording of effects of skeletal muscle relaxant drugs in
	rats/mice using Rota-rod apparatus and Analgesic activity using Eddy's
	Hot Plate with the help of software.

## 3.6.5T. NATURAL PRODUCT CHEMISTRY (Theory)

CO1	Understand and explain about natural product based drug discovery &
	their contribution in modern drug discovery.
	Comprehend tools & techniques used in study of biosynthetic pathways
CO2	in plants. Explain cardiovascular-active & anti-cancer agents from
	marine source.
CO3	Explain Natural products used as Pharmaceutical excipients including
CO3	Natural colors & dye, Natural sweeteners, Natural polymers
	Describe Herbal dietary supplements, Natural pesticides and natural
CO4	products as oral as bioavailability enhancers, skin permeation
	enhancers, Radiation protecting agents, wound healing agents,



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## 3.6.5 P NATURAL PRODUCT CHEMISTRY (Practical)

Upon completion of the course student shall be able to

	Perform	extract	ion,	isola	tion	aı	nd	esti	imation	of	var	ious
CO1	phytocons	stituents	and	analy	ysis	of	isola	ted	phytoco	nstitı	ıents	by
	chemical t	tests, ch	romato	graph	y and	spe	ectral	l me	thods (UV	/ and	/IR).	
CO2	Derive p	hysical	consta	nts c	f pu	re :	natur	ral (	compoun	d an	d isc	late
CO2	phytocons	stituents	by col	umn c	hrom	ato	graph	ıy.				

## 3.6.6 T. BIOORGANIC CHEMISTRY AND DRUG DESIGN (Theory)

Upon completion of the course student shall be able to

	<u> </u>
CO1	Generalize bioorganic Chemistry and its relevance in drug design and
	discovery.
	Compare and Contrast various drug targets and their biochemical
CO2	features, physiological & pathophysiological roles and their significance
	in drug design
CO3	Describe various approaches in rational drug design.
CO4	Explain the concept of pro-drug in drug design.

## 3.6.7 T. PHARMACEUTICAL BIOTECHNOLOGY (Theory)

CO1	Impart the basics of biotechnology techniques and the various systems used and the method of genetic engineering for production of rDNA products including monoclonal antibodies.
CO2	Apply the knowledge about the application of genetic engineering in animals.
CO3	Illustrate use of Fermenter for production of fermentation products and information about their purification by downstream process.



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## B.Pharm final Year (VII sem) (2015 Pattern)

## 4.7.1 T Sterile Products (Theory)

Upon the completion of the course student shall be able to

_	1
CO1	Enumerate the basics of sterile products
CO2	Describe various packaging materials used, types, choice of containers, official quality control tests and methods of evaluation.
СОЗ	Explain GMP, design and layout of Parenteral Production Facility.
CO4	Explain sterile products with respect to formulation, processing, manufacturing, packing and Quality control.

## 4.7.1 P Sterile Products (Practical)

Upon the completion of the course student shall be able to

	<del>-</del>
CO1	Demonstrate the manufacturing procedure for sterile preparation
CO2	Formulate, pack, evaluate,& label sterile products
CO3	Evaluate Packaging Material for sterile products as per official procedures
CO4	Evaluate marketed parenteral products like lyophilized products as reconstitutable solution or suspension for injection, suspension or emulsion

## 4.7.2 T Pharmaceutical Analysis-V (Theory)

Upon completion of the course student shall be able to

CO1	Differentiate different types of instrumental analytical techniques
COI	Differentiate different types of instrumental analytical techniques available for quality control of API & formulation
CO2	Describe principles, instrumentation and application of HPLC, IR, GC,
CO2	Describe principles, instrumentation and application of HPLC, IR, GC, NIR, Raman and advantages of UPLC
CO3	Explain principles, and application of Scanning Electron Microscopy
003	Explain principles, and application of Scanning Electron Microscopy and Transmission Electron Microscopy.

## 4.7.2 P Pharmaceutical Analysis-V (Practical)

CO1	Operate, UV-visible spectrophotometer for the assay of various APIs and
	formulation
CO2	Interpret the data obtained through Infrared spectra and report the result

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## 4.7.3 T Medicinal Chemistry-III (Theory)

Upon completion of the course student shall be able to

-	1
CO1	Classify the medicinal compound and acquire knowledge about IUPAC names along with mechanism of action of them or the class to which they belongs.
CO2	Describe the metabolism, adverse effect, therapeutic activity and recent
	developments of drugs.
	developments of drugs.
~ ~ ~	
CO3	Apply scientific knowledge about relationship between biological activity
	and structure of various chemotherapeutic agents and antibiotics.
CO4	Outline the synthetic route for selective medicinal compounds.
	F
CO5	Describe the chemotherapy for cancer and bacterial diseases and
	different anti-viral agents.

## 4.7.3 P Medicinal Chemistry-III (Practical)

Upon completion of the course student shall be able to

	CO1	Synthesize medicinally important compounds and purify them using column chromatography
Ī	CO2	Characterize the synthesized compounds using IR and NMR spectras.

## 4.7.4 T Pharmacology-IV (Theory)

Upon completion of the course student shall be able to

CO1	Describe about general principles of chemotherapy of infections and
	mechanism of drug resistance.
	Classify and discuss various chemotherapeutic agents depending upon
CO2	mechanism of action, antibacterial spectrum, resistance, therapeutic
	uses, adverse effects and contraindications.
	Discuss functions, receptor and mechanisms of hormone actions of
CO3	endocrine system, thyroid and parathyroid glands, androgens,
	estrogens, progestin and oral contraceptives.
CO4	Explain the hormones responsible for the cause of diabetes and
004	pharmacotherapy of diabetes.

## 4.7.4 P. Pharmacology-IV (Practical)

CO1	Perform and explain three point and four point bioassay to find unknown concentration of Acetylcholine.
CO2	Discuss the fixed dose combination of various drugs based on possible



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	indications, dose, route of drug administration, justification of inclusion of each ingredient, adverse reactions, contraindications, precautions and special instruction to patients.								
СОЗ	Illustrate rational drug therapy for treatment of various diseases.								
CO4	Justify whether rational drug therapy is followed from the prescription by RMP for specific patient.								

## 4.7.5 T Natural Drug Technology (Theory)

Upon the completion of the course student shall be able to

	Community of 0 complete regularity forting officet on level of accordance
CO1	Comprehend & explain various factors effect on level of secondary
	metabolites, method of cultivation, harvesting, storage .Factors affecting
	on deterioration of drug. Explain various guidelines issued by WHO in
	relation with cultivation, collection, storage etc.
CO2	Explain applications of Plant Tissue culture in production of secondary
	plant metabolites and invitro screening methods and its applications for
	natural products.
	Understand & explain basic principles of therapy in Ayurveda, Unani,
CO3	Siddha and Homeopathy and various dosages, its preparation,
C03	evaluation used in Ayurveda. Explain various novel drug delivery
	systems for herbal drugs.
	Comprehend & Explain herbal cosmetics, formulation and evaluation.
CO4	Explain physical, chemical, spectroscopic means and methods used in
	structural elucidation of herbal product

## 4.7.5 P Natural Drug Technology (Practical)

Upon the completion of the course student shall be able to

CO1	Formulate And Evaluate Ayurvedic /Herbal /Cosmetic/Nutraceutical
COI	Formulation
CO2	Understand Rationale And Conduct pre-formulation Parameters, <i>In-vitro</i> Assays For Correlation With <i>In-vitro</i> Assays.
	Tiobay of the Correlation with the burner issays.

## 4.7.6. T Biopharmaceutics and Pharmacokinetics-(Theory)

	Understand concept of biopharmaceutics and its application in					
CO1	formulation and development, studying various concept of ADME and					
	various factors affecting related to them.					
CO2	studying compartment and non-compartment modelling evaluate the					
CO2	quantity/concentration of drug in body at any point of time					
CO3	Understanding the concept and mechanism of dissolution and in-vitro					
	Understanding the concept and mechanism of dissolution and in-vitro and in-vivo correlation and Learning concepts of bioavailability and					



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bioequivalence	

## 4.7.7 T Pharmaceutical Jurisprudence-(Theory)

	To understand the basic principles, purpose & dimensions of the laws,					
CO1	significance & relevance of pharmaceutical laws in India, significance of schedule M & schedule Y, regulatory system for safety & effectiveness of					
	medicine & quality of product.					
CO2	To Discuss the purpose of the board, inspections by the board or its					
002	representative.					
	To learn the various laws governing the manufacturing, sale, research&					
CO3	usage of drugs, knowledge about patents, procedure for patent					
	application & IPR.					



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## 4.8.1 T Advanced Drug Delivery System (Theory)

Upon the completion of the course student shall be able to

-	<u> -</u>						
	Describe the Concept of Modified Drug Release, Novel Drug Delivery						
CO1	Systems, Pre requisites of drug candidates, Polymers along with						
	Classification, application, and evaluation approaches.						
CO2	Explain therapeutic Aerosols along with typical formulations from,						
002	metered dose, intranasal and topical applications.						
CO3	Enumerate the concept of microencapsulation and optimization.						

## 4.8.1 P Advanced Drug Delivery System (Practical)

Upon the completion of the course student shall be able to

	-			
CO1	To conduct polymer characterization by various methods, DSC, FTIR,			
	XRD, Viscosity and Swelling index.			
	Formulation development and evaluation of beads, sustained release,			
CO2	transdermal, gastro-retentive, microencapsulation and liposomal			
	formulations.			
CO3	To Study the concept of cosmeceuticals, history, difference between			
	cosmetics & cosmeceuticals & cosmeceuticals agents.			

## 4.8.2 T Cosmetic Science (Theory)

Upon completion of the course student shall be able to

CO1	Recognize the concepts of cosmetics, anatomy of skin Vs hair, general
	excipients used in cosmetics.
CO2	To perform formulation, manufacturing, equipments & evaluation for
	skins cosmetics, for hairs cosmetics, for eyes cosmetics, baby cosmetics.
СОЗ	To Study the concept of cosmeceuticals, history, difference between
	cosmetics & cosmeceuticals & cosmeceuticals agents.

## 4.8.2 P Cosmetic Science (Practical)

Upon completion of the course student shall be able to

$\perp$ $C(C)$	To describe use of ingredients in formulation, category of formulation &	
	<i>.</i> 01	prepare labels as per regulatory equipments.
C	CO2	To perform formulation, evaluation and labeling of cosmetics.

## 4.8.3 T Pharmaceutical Analysis-VI (Theory)

CO1	Differentiate	different	types	of	instrumental	analytical	techniques
COI	available for quality control of API & formulation.						



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CO2	Describe principles, instrumentation and application of NMR, ESR & Mass spectroscopy, Introduction to Flow injection analysis.
СОЗ	Describe principles, instrumentation, and application of Ion exchange chromatography, Flash chromatography, Super Critical Fluid Chromatography.

## 4.8.3 P Pharmaceutical Analysis-VI (Practical)

Upon completion of the course student shall be able to

CO1	Operate, UV –Visible spectrophotometer for method validation of various
	APIs and formulation.
1 (1)	Interpret the data obtained through UV, IR, NMR, Mass spectra and
	report the result.

## 4.8.4 T Medicinal Chemistry-IV (Theory)

Upon completion of the course student shall be able to

CO1	Classify the medicinal compound and acquire knowledge about IUPAC names along with mechanism of action of them or the class to which they belongs.
CO2	Describe about metabolism, adverse effect, therapeutic activity and recent developments of drugs.
CO3	Apply scientific knowledge about relationship between biological activity and structure of various antihistaminic, NSAIDs and Narcotics.
CO4	Outline the synthetic route for selective medicinal compounds.
CO5	Describe the Steroidal drugs, Hormones, Diagnostic agents and serotonergic agents

## 4.8.4 P Medicinal Chemistry-IV (Practical)

CO1	Synthesize medicinally important compounds and purify them using
	column chromatography.
CO2	Characterize the synthesized compounds using IR and NMR spectra's.
CO3	Demonstrate the various software's for physico-chemical property prediction and understand how current drugs were developed by using pharmacophores modeling and docking technique. (CADD)

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## 4.8.5 T Pharmacology-V (Theory)

Upon completion of the course student shall be able to

CO1	Describe important aspects, classification, mechanism of drug-drug
	interaction and ADRs.
CO2	Explain Basic aspects of drug safety and Pharmacovigilance in relation
	to monitoring and reporting of ADRs, functioning and role of hospital
	pharmacy and patient compliance.
CO3	Discuss of clinical trials, ethics and practice of GCP guidelines;
	Schedule Y, involved in clinical trials, explain the process, working and
	personnel involved in clinical data management.

## 4.8.5 P Pharmacology-V (Practical)

Upon completion of the subject student shall be able to

CO1	Demonstrate use of isolated tissue preparations for antagonistic
	bioassay methods.
CO2	Explain Basic aspects to carry out neurobehavioral characterization for
	determination of CNS activities.
CO3	Understanding various parametric and non-parametric tests used in
	biostatistics.
CO4	Perform statistical calculation of the given data using suitable statistical
	method.
	modiod.

## 4.8.6 T Natural Products: Commerce, Industry & Regulations (Theory)

Upon the completion of the course student shall be able to

CO1	Understand and explain commerce of various natural products which
	includes its global and local market size, demand and supply, import
	and export. Explain various aspects of Herbal drug industry.
CO2	Comprehend regulations and patenting of Herbal drugs in India.
CO3	Explain the toxicities of herbals and herbal-drug & herbal-food
	interaction.
CO4	Explain the pharmacovigilance of herbal medicines and plant allergens
	including classification, applications and method of preparation of
	allergenic extract.

## 4.8.7 T Quality Assurance Technique (Theory)

CO1	Explain significance of quality in pharmaceutical manufacturing and
	role of regulatory, explain the concept of QbD.
CO2	Describe quality standards of different agencies, Significance of
	validation in quality assurance.



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CO3 Apply cGMP, GLP and GDP while working in pharmaceutical industry.